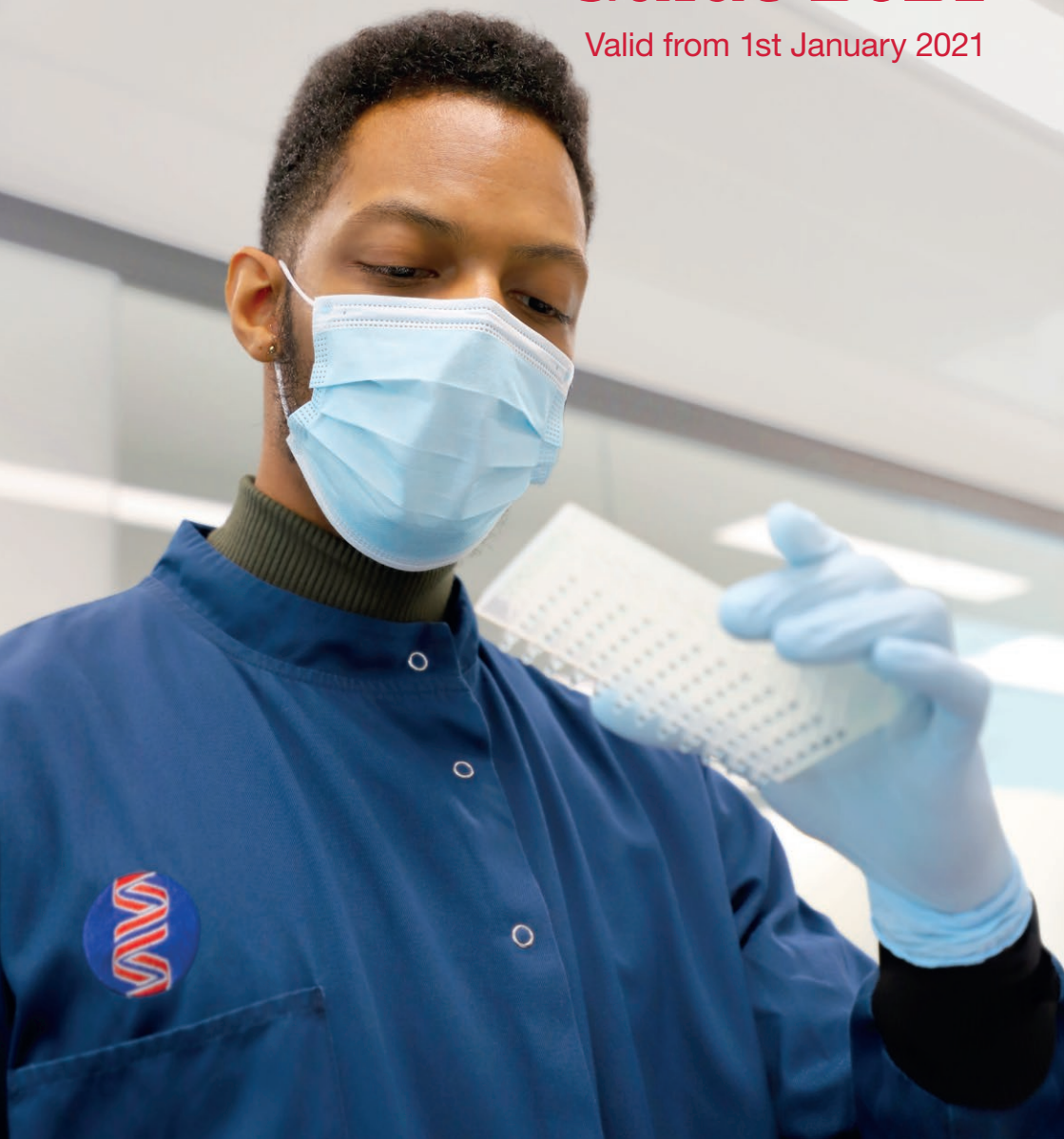




THE DOCTORS
LABORATORY

Laboratory Guide 2021

Valid from 1st January 2021



Cover: Daniel Odukoya, BMS, Laboratory Supervisor within
Molecular Pathology handling PCR samples.

Laboratory Guide 2021

Valid from 1st January 2021

TDL Customer Charter

We are committed to being the most helpful pathology service in the UK. Our goal is always to provide a high level of service to our customers, who request pathology services, for their patients. This is a philosophy shared by all Sonic Healthcare Pathology practices. We are medically led, and patients are our first concern. We always try to look to improve our operational expertise, and we strive to provide professional leadership within our specialities.

We promise to provide easy access to our pathology services

- We will always provide a friendly, helpful service.
- Our automated laboratory departments operate 24 hours a day, 7 days a week, and we aim to achieve, or improve, our published turnaround times.
- Our medical consultants and laboratory teams are available to provide additional clarification, advice or information for tests or results.

We promise to help you

- We invest in technical and operational excellence, with an extensive test repertoire, to ensure access to a leading-edge laboratory service.
- We return results using the reporting method choice, in an as organised and safe way as possible.

We promise to support the communities we work in

- We do our utmost to provide a service, even during extreme external disruptions beyond our control.
- We are committed to our staff's continued professional development.
- We have an organised programme to provide young people with work experience.
- We support our local community.

We promise to listen

- We acknowledge customer issues, and try to resolve them promptly and consistently.
- If our delivery has been adversely affected, we will address and review our procedures so that our service reaches the highest standards.
- We actively ask for feedback so that we can continue to improve our service.

Complaints policy

It is the aim of the company to maintain its core values. Two of these core values are:

- Commit to service excellence.
- Be enthusiastic about continuous improvement.

Where a doctor or patient needs to raise a complaint about service levels they should contact Cyril Taylor, Director of Laboratory Compliance, or Annette Wilkinson, Director of Service at tdlservice@tdlpathology.com giving details of the complaint.

The information forwarded will be treated as confidential and investigated by the above persons. This process will link into Quality Management procedure for incident investigation. Corrective and preventative actions will be introduced where indicated.

Contents

	PAGE
Index of TDL Profiles	2-3
Location maps for TDL London and Manchester	4-5
Helpful information for using The Doctors Laboratory	6-13
Quality assurance	14-20
Special instructions for samples	21
TDL Screening Profiles DL1 – DL12	22-23
Testing for COVID-19 (SARS-CoV-2)	24-25
Biochemistry	27-35
Haematology	36-39
Microbiology	40-47
Endocrinology	49-55
Reproductive health	56-59
TDL Andrology	60-64
Sexual Health: Tests, profiles and detection information	65-76
Immunology: General/Infectious immunology/Serology	77-85
Tropical and travel related immunology	86-88
Virology: Immune status testing	89
Hepatitis testing and hepatitis profiles	90-93
HIV testing	94-95
General	96-98
Tumour markers	99-100
Genetics – Cytogenetics/Molecular genetics	101-128
In-Vivo Tests	129
Antibiotic assays	129
Therapeutic drug assays	130-131
Allergy	133-141
Vitamins, Nutrition and Lifestyle, Omega 3/6	143-145
TDL Tinies™ and Self-collection samples	146-151
Screening for Drugs of Abuse/Alcohol	153-154
Occupational Health	155-156
Cervical Screening	157-165
Histopathology	166-170
Alphabetical test index	172-203
TDL Referral Laboratories	204-206
Terms and conditions of business from 1st Jan 2021	207-213
Forms	215
Downs risk profile (1st & 2nd trimester)	
Leukaemic studies request form (Cytogenetics/Molecular genetics)	
Genetic request form	
Supplies order	
TDL request form	

Index of TDL Profiles

TDL SCREENING PROFILES

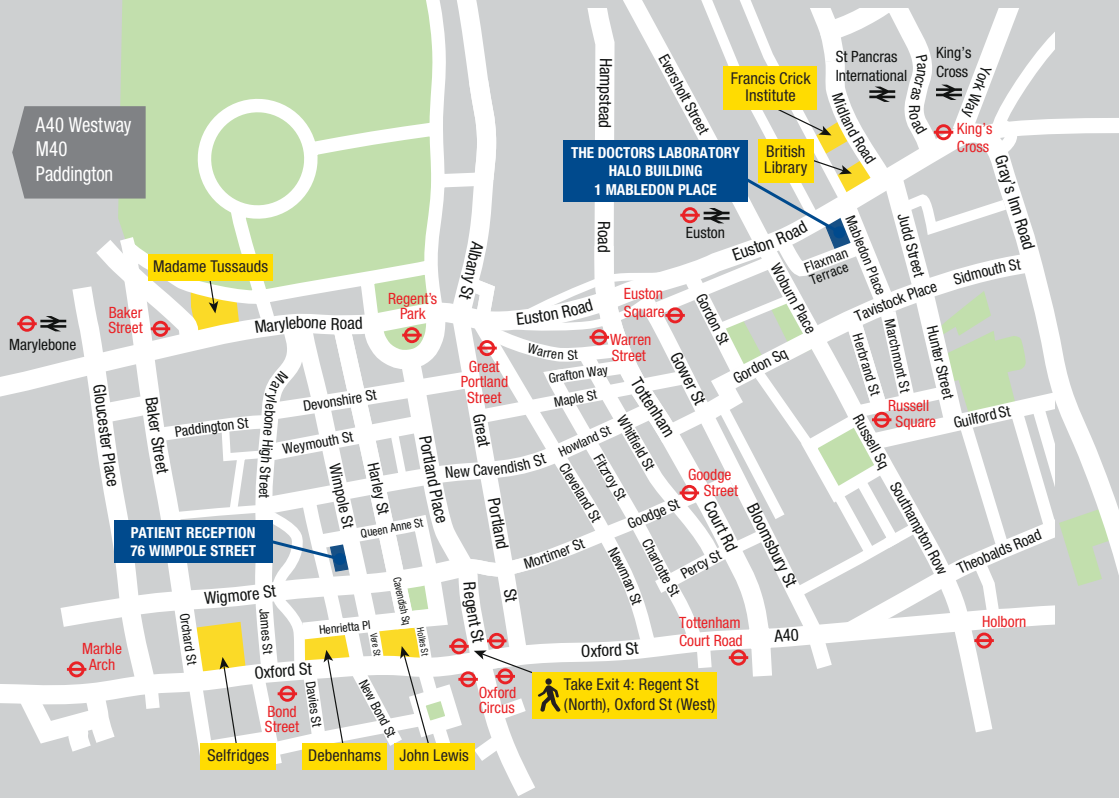
	PAGE
DL1/DL1L Biochemistry Profile	22
DL2/DL2L Haematology and Biochemistry Profile (24 parameters)	22
DL3 Haematology Profile	22
DL4/DL4L Haematology and Biochemistry Profile (16 parameters)	22
DL5/DL5L Postal Haematology and Biochemistry Profile	22
DL6/DL6L General Well Person Profile	22
DL7/DL7L Well Man Profile	23
DL8/DL8L Well Person Profile	23
DL9M Senior Male Profile	23
DL9F Senior Female Profile	23
DL10 Cardiovascular Risk Evaluation Profile	23
DL11 Cardiovascular Risk Plus Profile	23
DL12 Sexual Health Profile 7 STI's by PCR	23

TDL SPECIFIC PROFILES

Alcohol Profiles	153-154
Allergy Screens	133-141
Amenorrhoea Profile	49,55
Anaemia Profile	36,39
Andropause Profile	49,54
Antenatal Profile	36,39
Autoantibody Profiles	77,85
Azoospermia Profile	108, 128
Bone Screens	28,35
Calprotectin/Elastase Profile	77,85
Cardiovascular Risk Profiles	28,35
Chest Pain Profile	28,35
Chlamydia (Species Specific) Antibody Profile	78,85
Chronic Fatigue Syndrome Profile	78,85
Clotting Profiles	36,39
Coeliac Profiles	78,82
NEW COVID-19 (PCR & Antibody testing)	24-25,96-97
Deep Vein Thrombosis (DVT) Profile (Pre-travel screen)	36,39,86-87,111,128
Diabetic Profiles	29,35
Drugs of Abuse/Alcohol Screens	153-154
Enteric Organism Rapid Antigen Detection	86-87
Epstein-Barr Virus Profile	96
Erectile Dysfunction Profile	49,54
Female Hormone Profile	49,54
First Trimester Antenatal Screening Bloods	49,55
Genetic Profiles	128
Haematology Profile	36,39

	PAGE
Hepatitis Profiles	90
Hirsutism Profile	49, 55
HIV Profiles	65, 75-76, 94-95
HRT Profile	49, 55
Impotence Profile	50, 54
Infertility Male Profile	50, 54
Iron Overload Profile	31, 34, 115, 128
Iron Status Profile	31, 34
Jewish Carrier Screen	108, 115, 123, 128
Lipid Profile	31, 34
Liver Function Tests	31, 34
Male Genetic Reproductive Profile	113, 116, 128
Menopause Profile	50, 55
Metabolic Syndrome Profile	50, 55
Mineral Screen	143-144
Myeloma Screen	31, 34
Natural Killer Profile	36, 39
Needle Stick Injury Profile	97
Neurological Viral Screen	97-98
Osteoporosis Screen	32, 35
Pituitary Function Profile	50, 55
Pneumonia (Atypical) Screen	97-98
Polycystic Ovary Syndrome Profile	50, 55
Post-Travel Screens	86-87
Pre-Travel Screen	36, 39, 86-87, 118, 128
Prostate Profile	99
Recurrent Miscarriage Profile	119, 128
NEW Respiratory Viral Screen	25, 97-98
Rheumatology Profiles	80, 84
Rickettsial Species Antibodies	80, 86
NEW SARS-CoV-2 (COVID-19) PCR & Antibody testing	24-25, 96-97
Sports/Performance Profile	143-144
STI/Sexual Health Profiles	65-66, 74-76
Thrombotic Risk/Miscarriage Profile	37, 39, 120, 128
Thyroid Profiles	51, 54
Torch Screen	80, 97-98
Trace Metal Screen	144, 155
Tropical Screen	86-87
Urea and Electrolytes	33-34
Viral Profiles	98
Vitamin Screens	144-145
Von Willebrand Profile	37, 39

Personal Profile (Doctor's own) are available on request.



THE DOCTORS LABORATORY

The Halo Building, 1 Mabledon Place, London WC1H 9AX

Tel: 020 7307 7373

E-mail: tld@tdlpathology.com

Web: www.tdlpathology.com

PATIENT RECEPTION/PHLEBOTOMY SERVICES

76 Wimpole Street, London W1G 9RT

Telephone: 020 7307 7383

Email: patientreception@tdlpathology.com

OPENING TIMES

Monday to Friday 7.00am – 7.00pm

Saturday 7.00am – 1.00pm

Out of hours samples

can be dropped at:

Patient Reception

76 Wimpole Street

London W1G 9RT

Or at any time at

the main laboratory:

The Halo Building

1 Mabledon Place

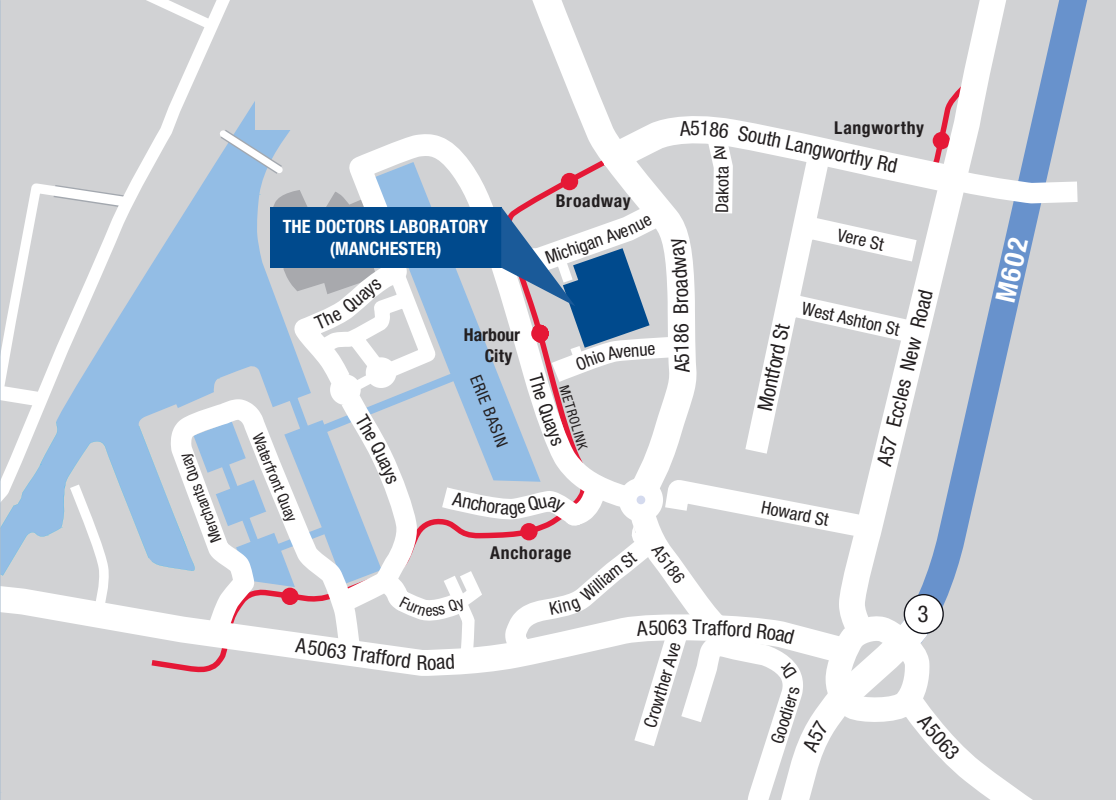
London WC1H 9AX

Samples are taken at

Patient Reception only.

Samples cannot be taken

at The Halo Building.



THE DOCTORS LABORATORY (MANCHESTER)

Michigan House, Michigan Avenue
Salford Quays, Manchester M50 2GY

Tel: 0161 332 7181

Web: www.tdlpathology.com

Samples can be dropped at the
laboratory at any time.

COURIER COLLECTIONS

Tel: 0161 332 7187

Helpful information

The Laboratory Guide is designed to give you an easy-to-use reference for the most regularly requested services, pathology profiles and tests. If you are not able to find details for tests and services, please contact the laboratory on 020 7307 7373. We continue to develop a wide range of test and patient services and our aim is to offer commitment to customer service, strong working relationships and help and support for referring doctors and their practices.

For details about all services, please contact the laboratory on 020 7307 7373, or for more information visit www.tdlpathology.com.

LONDON LABORATORY TIMES: 24 HOURS

A wide range of analytical services are run 24/7 but not all tests or departments operate throughout the night, weekends, or bank holidays.

Outside routine hours the night team provide a wide range of analytical services but not all tests will be run as standard. No surcharges are made unless there are special arrangements for courier collections or investigations requiring additional resources.

Outside Patient Reception hours samples may be dropped off at 76 Wimpole Street, London W1G 9RT or at the laboratory at The Halo Building, 1 Mabledon Place, London WC1H 9AX (see page 4) at any time.

MANCHESTER LABORATORY TIMES: 24 HOURS

Samples may be dropped off at the laboratory at Michigan House, Michigan Avenue, Salford Quays, Manchester M50 2GY (see page 5) at any time.

MANCHESTER TURNAROUND TIMES

Tests not processed at our laboratory in Manchester will be referred to the TDL Main laboratory. If you need information about turnaround times please contact the laboratory.

PATIENT RECEPTION TIMES

Patient Reception is at:

76 Wimpole Street, London W1G 9RT

Monday to Friday 7.00am – 7.00pm, Saturday 7.00am – 1.00pm

Direct line tel: 020 7307 7383

Appointments are only necessary if a patient needs specialised investigations or care. Patients should always bring a request form or referral letter with them. Instructions can be telephoned ahead of the patient's attendance, if this is more convenient.

Patient Reception Direct line tel: 020 7307 7383 Email: patientreception@tdlpathology.com

Sample taking is undertaken by qualified staff for which a standard sample taking fee of £45.00 is charged to patients. A fee of £22.00 is charged to doctors and clinics for each patient.

Sample taking services for extended tests (see page 129) and **Drugs of Abuse with Chain of Custody** are routinely available.

Helpful information

Cervical cytology, HVS and cervical swabs are not taken at 76 Wimpole Street.

Patient Reception sample taking services are not available in Manchester.

SEMEN ANALYSIS

Semen samples need specialist handling within the laboratory. For this reason all requests for Semen Analysis must be made by appointment. Practices or patients can make an online appointment at www.tdlpathology.com/andrologybooking or call **020 7307 7373** to make appointments and confirm instructions for sample collection. There is an attendance fee of £45.00.

- 1 Patients must abstain from ejaculation for at least 2 days but not longer than 5 days before the test.
- 2 Ideally semen samples should be produced at The Doctors Laboratory, 76 Wimpole Street, unless there are exceptional circumstances. In these exceptional circumstances please contact TDL Andrology on 020 7025 7940 for special arrangements and instructions. Refer to Andrology, see page 60.

Semen analysis service is not provided in Manchester.

PATIENT REQUEST FORM

To comply with good clinical practice it is important that there is one request form for each patient's request, and specimens and form are correctly and fully labelled, to include three unique patient identifiers:

- First name, Surname, Date of birth, Hospital/ Clinic number, Medical Record Number (MRN) are examples of patient identifiers
- Time and Date of collection of samples
- Type of sample and Anatomical site, where appropriate (e.g. swabs)
- Relevant clinical information
- Relevant details of medication
- High Risk Samples should be clearly identified on the form and individually packed separately from other samples
- Hazard Group 4 pathogens (such as Ebola or Viral Haemorrhagic Fever) must not be sent to the laboratory – please contact the National Fever Service on 0844 778 8990 for advice before sending samples to the laboratory

If additional tests are required for a sample already received please contact the laboratory on 020 7307 7373 with your request for specific further analysis. Samples are stored within timeframes according to their discipline. Laboratory staff will advise on the ability to undertake further testing from samples already received in the laboratory.

Helpful information

EMAILED REQUESTS FOR ADD ONS

The majority of samples received in the laboratory are kept for one week. If sample type and volume allow, further testing can be requested by telephone (020 7307 7373) or by email to **addons@tdlpathology.com**. Please specify the test details to be added, together with Patient details, and LABORATORY NUMBER need to be given with Emailed requests.

HOME VISITS

This service is available for patients who, for whatever reason, prefer samples to be taken at home or at locations other than a doctor's practice or TDL's Patient Reception at 76 Wimpole Street. This is a service that is used regularly to save time for both doctors and patients and ensures that results can be made available before consultation is undertaken.

There is a visit fee from £120.00 to patients within the M25, from £160.00 for children when two nurses are needed. Home visits outside the M25, for weekends, bank holidays and night fees are by special arrangement. To arrange a Home Visit please telephone Patient Reception on **020 7307 7383** or email homevisits@tdlpathology.com.

TDL COLLECT: SPECIMEN COLLECTION SERVICES BY COURIER

TDL operates a dedicated and extensive specimen collection service. **TDL Collect** provides a 24 hour professional sample collection service on an urgent, regular or random basis. No charge is made for collections from practice within the M25. Sample collection from practices outside the M25 is by arrangement and may incur courier charges.

TDL COLLECT Online Courier Booking is a time saving new service at www.tdlpathology.com/couriers. For your practice's Username and Password please contact Chris Tanalega on 020 7025 7929 or chris.tanalega@tdlpathology.com.

Our couriers are trained to Health and Safety guidelines and maintain our commitment to customer service. For added convenience to doctors and their patients, we also collect samples directly from patients' homes, offices or hotels within the M25.

To arrange courier collection of samples from other areas in the UK please telephone **020 7307 7373**.

High risk samples should be clearly labelled and packed separately from other samples.

TDL Collect cannot transport samples containing Hazard Group 4 pathogens, such as Ebola fever or Viral Haemorrhagic Fever.

TDL COLLECT UK NUMBER: 020 7307 7373

SAMPLE PACKING

Samples need to be transported for subsequent processing and testing. Transport systems will be various and cover both long or short distances.

Samples need to be collected and packed into appropriate sample containers provided by the laboratory in order to maintain integrity of the sample(s). Attention needs to be given to temperature, special transport containers and time limitations.

Helpful information

Clinics, practices and laboratories who are posting or transporting samples by air, sea, rail and road between local, regional and reference laboratories, or between laboratories in other countries, must adhere to a number of regulations. These regulations are designed to deal with transportation accidents and spills, reduce biohazards and keep samples intact for testing.

Regulations are given by several sources including

- National transport regulations
- IATA
- Rail and road traffic agencies
- Postal services

Compliance is mandatory in order to reduce risk to couriers, carrier, laboratory staff and passengers.

Sample transport requirements are based on the category of samples being transported.

Infectious substances are classified as Category A or Category B.

TDL does not arrange for transport of Category A samples (infectious substances capable of causing permanent disability or life threatening or fatal disease to humans or animals).

Instruction and packaging for Category B is provided, covering Biological Substances, UN number UN 3373.

PACKAGING REQUIREMENTS

There are specific packaging instructions and labelling requirements requiring triple packaging.

- 1 Primary leak-proof container – tube or vial containing the sample must be placed inside a ziplock specimen bag with absorbent material
- 2 Secondary watertight container, with absorbent material, intended to protect the primary container
- 3 Outer container protects the secondary container

There are specific packaging instructions for frozen samples requiring shipment using BioFreeze bottles, or Dry Ice.

For information please contact the Referrals Dept (ReferralsOffice@tdlpathology.com)

POSTAL PATHOLOGY

Royal Mail Tracked 24[®]

NEW

Postal pathology services should be considered by all practices in the UK who need a rapid delivery service to the laboratory. Changes with Royal Mail mean that ALL pathology postal packs are now made up with **Tracked 24 returns**. This provides a particularly suitable method of transport for any healthcare organisation. Postal pathology with **Tracked 24 returns** provides:

- Simple and convenient sample handling throughout the UK for most tests.
It is not suitable for microbiology or coagulation samples
- Scope for large and small numbers of samples
- Next morning delivery
- Allows patients and practices to track samples through the Royal Mail system
- Samples can be posted from any Royal Mail post box, including COVID-19 antibodies
- Designated **Priority boxes** for COVID-19 PCR (swab) kits.
- From 1st January there will be a charge of £2.26 for each Royal Mail Tracked 24 pack.
This charge will be itemised in monthly invoices to the practice or patient, as requested.

Helpful information

DX SYSTEM

NEW

DX is a well known next-day courier of Category B specimens – transporting biological samples in compliance with the industry's highest regulations. DX is compliant to IATA regulations, is audited independently by Dangerous Goods Safety Advisors. They work with a combination of large health organisations and smaller, independent laboratories to ensure the safe delivery of specimens every year.

TDL's DX Address is **DX 340201, St Pancras 90 WC**.

PATHOLOGY CONSUMABLES/REQUEST FORMS/POSTAL PACKS

Our Stores Department provides all appropriate sample collection consumables required for sample collection. Orders will be sent same or next day and can be made by telephone (020 7307 7373), e-mail (supplies@tdlpathology.com) or fax 020 7307 7340. There is a Supplies Order Form at the back of this Laboratory Guide.

REQUESTING AND REPORTING OPTIONS

We continually review and update our IT Services for receiving requests and reporting results electronically between practices and the laboratory. A number of innovative report formats are now available.

• Encrypted Email

Results will be sent in encrypted format to any number of predetermined email addresses. Copy reports will be emailed automatically to email addresses on the system.

• Link to Practice Management System

Bidirectional requests and results can be delivered electronically to a number of integrated practice systems. Practice software that accepts data in an HL7 format can be linked to receive results from the laboratory.

All TDL systems are accredited to the latest International Standard for Information Security ISO/IEC 27001:2013.

• TDL e-View

Registered users can view all their results online. This is a secure Login/Password protected look-up system, with a cumulative results reporting function. Results can be accessed any time, from anywhere, through the internet.

• Printed Copy

Results are posted out on the day they are reported.

• TDL Portal

This provides the most accurate option for clinics without a practice management system. For information about this option please contact portal@tdlpathology.com.

Helpful information

EMAILED RESULTS INCORPORATING YOUR LOGO

If your practice or company receives results by email, and would like these personalised with your logo, simply email your company details and logo in GIF format to logo@tdlpathology.com.

TDL WEBSITE: RELAUNCHED FOR 2020/21

The TDL website at www.tdlpathology.com gives updated details of our tests – sample types, turnaround times and special instructions. A new Specialities section provides a new way to find tests you need, and a Services section has additional information for TDL Collect, Postal Pathology and TestGuide app. Reference Ranges are given on the website or can be requested by emailing refranges@tdlpathology.com.

TDL PATHOLOGY HANDBOOK

With more than, 1000 entries and 1100 pages covering pathology tests, methods and disease conditions, the Handbook provides comprehensive detail about the range of tests and services offered by the laboratory. Email handbook@tdlpathology.com for more information.

TDL TESTGUIDE APP

Available for iOS and Android, the TDL TestGuide app covers the full content of the Handbook but is also continually updated with new or revised entries on a regular basis.

If you would like to register for the app, it is straight forward to install this from an appropriate app store for your own device. Email testguide@tdlpathology.com for more details.

User feedback for the TDL website, Pathology Handbook and TestGuide app is always welcome, please send suggestions and comments to tdl@tdlpathology.com.

FEES FOR PATHOLOGY

Fees can be paid directly by patients or by the practice, clinic or requesting organisation. A payment instruction clearly identifying to whom invoices need to be sent must be given with each patient's request.

Patients are normally invoiced within 7 days to the address provided by the patient or practice. Their pathology fees include a standard credit/administration charge.

Receipts for insurance purposes are sent, if requested. Patients visiting Wimpole Street for sample taking have the opportunity to settle their pathology fees at the time of their visit. A credit/administration fee is raised for invoices sent to patients. All normal credit, debit or chargecards are accepted and payment can be made by following the telephone payment instructions given with each invoice.

The Terms and Conditions appearing on pages 207-213 of this Laboratory Guide shall apply to the services we provide to you, unless otherwise agreed.

Helpful information

PROTECTION OF PERSONALLY IDENTIFIABLE INFORMATION

The General Data Protection Regulation (GDPR) came in to force in May 2018 and has had a significant impact upon the way that personal data is managed; placing legal requirements upon data processors and controllers to manage that information securely, maintain records of the processing that is carried out, and report when breaches of the regulation do occur. This has impacted the way many businesses operate, and is not restricted to the healthcare sector.

The GDPR requirements have been implemented within the context of a mature ISO 27001 Information Security Management System – the globally accepted standard by which information is secured. This ensures that senior management have regular visibility of the threats to the confidentiality, availability and integrity of the information that we process, and are able to steer the efforts of their teams to provide an efficient service that places the confidentiality of our customers and their patients at the heart of everything we do.

In order to support our customers compliance with the regulation and as a part of a wider GDPR compliance project TDL has updated its standard terms and conditions to include revised data processing clauses, which are mandatory when providing personal data to another organisation.

Helpful information

WHO TO ASK FOR HELP

24 hour Telephone (Main Switchboard/All Services): 020 7307 7373

Fax: 020 7307 7374

CEO	David Byrne	david.byrne@tdlpathology.com
Group Laboratory Director	Tim Herriman	tim.herriman@tdlpathology.com
Director of Sales/Service	Annette Wilkinson	annette.wilkinson@tdlpathology.com
Director of Genetics & Molecular Pathology	Dr Lisa Levett	lisa.levett@tdlpathology.com
Chief Information Officer (IT)	John Matthews	john.matthews@tdlpathology.com

HEADS OF SUPPORT DEPARTMENTS

Group Laboratory Operations Manager	Lisa Manze	lisa.manze@tdlpathology.com
Director of QMG	Emer Nestor	emer.nestor@tdlpathology.com
Patient/Doctor Invoices	Aneta Kontrova	aneta.kontrova@tdlpathology.com
Logistics/Couriers	Steve Kettle	steve.kettle@tdlpathology.com
Patient Reception/Home Visits	Eileen Flatley	eileen.flatley@tdlpathology.com
Call Centre	Chris Tanalega	chris.tanalega@tdlpathology.com
IT Operations/Customer Service	Rochelle Fakhri	rochelle.fakhri@tdlpathology.com
Sample Reception	Aileen Francis	aileen.francis@tdlpathology.com
Referrals Department	Maulik Trivedi	maulik.trivedi@tdlpathology.com
Human Resources	Matthew Gibbins	matthew.gibbins@tdlpathology.com

HEADS OF LABORATORY DEPARTMENTS (LONDON)

Haematology	Billy Janda	billy.janda@tdlpathology.com
Biochemistry	Dayan Lloyd-Hennie	dayan.lloyd-hennie@tdlpathology.com
Microbiology	Alan Spratt	alan.spratt@tdlpathology.com
Andrology	Andrew Dawkins	andrew.dawkins@tdlpathology.com
Cytology	Margaret Morgan	margaret.morgan@tdlpathology.com
Immunology/Virology	Kushen Ramessur	kushen.ramessur@tdlpathology.com
Cytogenetics	Rebecca Watts	rebecca.watts@hslpathology.com
Molecular Genetics	Dr Stuart Liddle	stuart.liddle@tdlpathology.com
TDL Trials	Abraham Roodt	abraham.roodt@tdlpathology.com
Night Service	Sanjiv Sawock	sanjiv.sawock@tdlpathology.com

TDL MANCHESTER

Operational Site Lead	Diane Benson	diane.benson@tdlpathology.com
Systems Manager	Andy Leeson	andy.leeson@tdlpathology.com
SRA Manager	Georgina Arnold	georgina.arnold@tdlpathology.com
Quality Manager	Eamonn Donnellan	eamonn.donnellan@tdlpathology.com
Courier Control	Marc Rennard	marc.rennard@tdlpathology.com

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Quality assurance



The Doctors Laboratory is committed to providing doctors with pathology of the highest quality. The quality of results is of fundamental importance and the laboratory operates to stringent technical and administrative standards.

Internal quality assurance is achieved by strict adherence to standard operating procedures for all analytical processes. TDL participates in recognised National External Quality Assessment Schemes. These schemes are subscribed to by NHS and private laboratories. Results are subjected to strict internal and external quality control. Details of the laboratories to whom TDL refers specialist testing are available from TDL Referrals. These laboratories are UKAS accredited or of equal accreditation status. Details of the tests that are referred are given on the TDL website. QA is administered by TDL's Quality Management Group (QMG) who also adhere to regulatory and accreditation requirements.

BIOCHEMISTRY: UKNEQAS, WEQAS, RIQAS, BIORAD for

ACE

AFP / CEA & HCG

Antibiotics (Gentamicin, Vancomycin and Amikacin)

Anti-Hbs Detection

Ammonia

Autoimmune (RF and TPO)

B2 Microglobulin

Cardiac Markers

Clinical Chemistry

CMV IgG/IgM

CRP & Ultra-Sensitive CRP

CSF

Cyclosporin and Tacrolimus

DEQAS

Diagnostic Serology Exanthem

Diagnostic Serology Hepatitis

Drugs of Abuse

Ethanol

Faecal Markers for Inflammation (Calprotectin)

Free Beta HCG and PAPP-A

GFR

Glucose / Glucometer

Glycated Haemoglobins

Guildford Peptides

Haematinics

Healthcontrol Therapeutic Drugs Screen (TDM)

Hepatitis A (with B and C)

Hepatitis B Serology

Hepatitis C Serology

HIV Serology

Homocysteine

HTLV

IGF-1

Immunity Screen

Lipase

Lipid Investigations

NT-Pro BNP

Paediatric Bilirubins

Parasitology

Peptide Hormones

PSA, Free PSA

PTH, ACTH and hCT

Rubella IgG Serology

Salicylate and Paracetamol

Specific Proteins

Steroid Hormones

Syphilis Serology

Thyroglobulin Surveys

Thyroid Hormones

Total IgE

Toxoplasma IgG/M Serology

Quality assurance

Tumour Markers
Toxoplasma IgM Serology
Toxoplasma IgG Serology
Trace Elements
Urine Chemistry
Vitamin D (25 OH)

HAEMATOLOGY: UKNEQAS for

Automated Differential Leucocyte Count
Blood Film Morphology
Coagulation (Including PoCT Coagulation)
EBV Mononucleosis
ESR and NRBC (nucleated Rbc)
Flow Cytometry
 Leukaemia immunophenotyping
 Myeloperoxidase
 Iron stain
Full Blood Count
Haematology
Haematology Analysis
Malaria
Parasite Films
Reticulocyte
Sickle Screening
Thrombophilia Screening

Factors assays:

Von Willebrand (vWD) screen
Anti-Xa assays
Plasma viscosities
ADAMTS-13 activity
ADAMTS-13 antibody
Heparin/Platelet Factor 4 Induced Antibodies
Platelet function analysis (RCPA)
Lupus anticoagulant:
 Taipan Venom Time
 DRVWT assay

GENETICS AND MOLECULAR VIROLOGY

MOLECULAR GENETICS

Acquired array (CLL/MDS)
Acute Leukaemia FISH pilot
Acute Lymphoblastic Leukaemia (ALL)
 – G banding and FISH
BoBs Rapid Aneuploidy detection
Chlamydia & Gonorrhoea detection by PCR

Constitutional Clinical Cytogenetics
 (Rounds for Amniocentesis, CVS,
 Solid Tissue, Blood, Array CGH)
Cystic Fibrosis
Duchenne/Becker Muscular Dystrophy
Hereditary Haemochromatosis (C282Y+H63D)
 genotyping + reporting
HLA Class I (HLA-A, HLA-B, HLA-C)
 Tissue Typing (low resolution)
HLA Class II (HLA-DRB1, HLA-DQB1)
 Tissue Typing (low resolution)
HLA-B27 Genotyping
HLA-B57*01 Genotyping
HLA+ Disease Typing
 Cytochrome P450 2D6/2C19 genotyping
Human Papillomavirus DNA
Mature B & T cell Neoplasms –
 FISH for CLL and Lymphoma
Mature B & T cell Lymphoma – G-banding
Myeloid (AML/MDS/CML) – G-banding and FISH
Myeloma – sample FISH set up
 and analysis plus online
NGS AML gene panel
NIPT for aneuploidies
NIPT for sexing
Paternity Testing
Prader-Willi and Angelman Syndromes
QF-PCR Aneuploidy Detection
Sexually Transmitted Diseases (CT/NG/MGEN/TV)
Spinal Muscular Atrophy
Thrombophilia (Factor II, V, MTHFR)
Y Microdeletion PCR Assay

MOLECULAR VIROLOGY

Atypical Mycobacterium
Adenovirus DNA Viral load
Bacterial 16S
B19 virus DNA Viral load
BK virus DNA Viral load
CMV DBS (dried blood spots)
CMV DNA Plasma Viral load
CMV DNA Whole Blood Viral load
CMV Resistance
EBV DNA Plasma Viral load
EBV DNA Whole Blood Viral load
Enterovirus RNA

Quality assurance

Gastroenteritis Virus Panel
Hepatitis B Genotyping
Hepatitis B Drug Resistance Typing
Hepatitis B Viral Load
Hepatitis C genotyping
Hepatitis C Resistance genome detection (NS5a & b)
Hepatitis C Resistance Typing (NS3 & NS5a)
Hepatitis C Viral Load
Hepatitis D Virus Viral load and Qualitative PCR
Hepatitis E Virus Viral load and Qualitative PCR
HIV-1 Drug Resistance (Pol)
HIV-1 Drug Resistance (Integrase)
HIV-1 RNA Viral load
HIV-1 RNA Qualitative PCR
HIV-1 Tropism Genome Detection
HIV-2 Viral load and Qualitative PCR
HSV 1&2 DNA
HSV Drug Resistance
Human Herpes virus 6 DNA
Influenza Haemagglutinin typing
JC virus DNA
Measles and Mumps PCR
MERS Coronavirus
Parechovirus RNA
Respiratory panel I
Respiratory panel II
SARS-CoV-2 (COVID-19) PCR/NAAT
SARS-CoV-2 (COVID-19) antibodies
Syphilis PCR
Transplantation Virus Panel
VZV DNA

MICROBIOLOGY

Laboratory Quality Scheme:

Helicobacter pylori antigen from faeces
Polarising crystal microscopy from synovial fluid
Streptococcus pyogenes (Group A) detection
in pharyngeal samples
Surveillance for multi drug resistant bacteria

UKNEQAS:

Clostridium difficile detection and toxin testing
Faecal parasites
General bacteriology
Genital pathogens
MRSA screening
Microbial susceptibilities

Mycobacterial microscopy
Mycobacterial culture and molecular detection
Antifungal assays
Antifungal susceptibilities
Cryptococcal antigen
Fungal culture
Fungal biomarkers
Urinary antigen

WEQAS POCT:

Urinalysis

QCMD:

Dermatophyte PCR
PCP PCR
Atypical pneumoniae PCR

IMMUNOLOGY

UKNEQAS – General Immunology for:

Allergen Component Testing
Autoimmune Serology ANCA/GBM Antibodies
Bullous Dermatositis Antibodies
Allergen Specific IgE Antibodies
Ner General Autoimmune Serology
Anti-Phospholipid Antibodies (ACAB)
Nuclear and Related Antigens
IGRA TBQ
Intrinsic factor
Islet Cell Antibodies (Diabetic Marker)
Myositis Antibodies
Specific Microbial Antibodies
C1 Esterase Inhibitor
Functional and Complement
Syphilis (TPPA and RPR)
Lyme (IgG and IgM)
Hepatitis C
Hepatitis E (IgG and IgM)

EUROQAS:

Liver Blot

UKNEQAS – Infectious Immunology for:

HIV Serology/POCT
Immunity Screen – VZV, Parvo Viruse, EBV
Chlamydia Detect
Varicella Zoster (IgG) Serology
Parasite Serology
Chlamydia & Gonorrhoea (NAAT/PCR)

Quality assurance

RIQAS Scheme:

Procalcitonin

RCPAQAP Scheme:

Brucella Serology

Legionella (IgG) Serology

Scleroderma Antibodies

Striated Muscle Antibodies

INSTAND Scheme:

Adrenal Antibodies

Hepatitis E Serology

RNAP Antibodies

CSCQ Scheme:

Lyme Serology

Laboratory Quality Scheme:

Herpes Simplex Virus

Cytomegalovirus

Antistreptolysin O Titre

Helicobacter Pylori IgG Antibodies

ENDOCRINOLOGY: UKNEQAS for

Steroid Hormones

Peptide Schemes 1 to 4

Thyroid Scheme

Allergens Scheme

SHBG

Prostate Specific Antigen

Tumour Markers

PTH

Specific IgE/Total IgE

AFP/CEA

CYTOLOGY: EQA, TEQA for

NHSCSP (EQA for Gynaecological Cytopathology)

NHSCSP (TEQA for PAP stain)

Hologic Imager stain (TEQA)

NEQAS:

Urine Cytology

ANDROLOGY: UKNEQAS for

Semen Analysis Scheme

Information security:

Accredited by British Standards Institute

ISO/IEC 27001:2013

LINKS TO THE UKAS SCHEDULES OF ACCREDITATION

(Certain UKAS accreditations can be found under Health Services Laboratories (HSL), which is part of the TDL Group of Laboratories.)

HSL Blood Sciences (8169)

https://www.ukas.com/wp-content/uploads/schedule_uploads/00007/8169%20Medical%20Single.pdf

HSL Infection Sciences (8860)

https://www.ukas.com/wp-content/uploads/schedule_uploads/00007/8860%20Medical%20Single.pdf

HSL Molecular Pathology and Genetics (8059)

https://www.ukas.com/wp-content/uploads/schedule_uploads/00007/8059%20Medical%20Single.pdf

TDL Manchester (8812)

https://www.ukas.com/wp-content/uploads/schedule_uploads/00007/8812%20Medical%20Multiple.pdf

TDL Andrology (10199)

https://www.ukas.com/wp-content/uploads/schedule_uploads/00007/10199%20Medical%20Single.pdf

HSL Cytology (8511)

https://www.ukas.com/wp-content/uploads/schedule_uploads/00007/8511%20Medical%20Single.pdf

Quality assurance

MEASUREMENT UNCERTAINTY

Medical laboratories are responsible for ensuring that test results are fit for clinical application by defining analytical performance goals and selecting appropriate measurement procedures. All types of measurement have some inaccuracy due to bias and imprecision; therefore measurement results can only be estimates of the values of the quantities being measured. To properly use such results, medical laboratories and their clinical users need some knowledge of the accuracy of such estimates.

The complete result of a measurement is a value, a unit and an estimate of uncertainty. This estimate of uncertainty is conventionally referred to as Measurement Uncertainty (MU) and incorporates the cumulative range of factors involved in the testing procedure itself in addition to consideration of the inter-individual and intra-individual biological variation which will potentially influence the overall test result. Evaluating measurement uncertainty is an ISO 15189:2012 accreditation requirement.

In terms of MU determined by the TDL/HSL group of laboratories, it should be noted all assays are performed in strict accordance with the manufacturers' instructions. MU, which has been estimated for each assay during the verification procedure, is reviewed at regular intervals to ensure that MU values do not exceed the pre-defined maximum allowable uncertainty for each assay. Overall assay performance is also regularly monitored through internal quality control (IQC) and external quality assessment (EQA) schemes and incorporated in test result interpretation. MU for individual assays is available upon request.

SAMPLE REJECTION CRITERIA

Sometimes tests cannot be performed in the laboratory if samples fall short of the quality, volume or other eligibility criteria. In these cases, the laboratory may need to reject the samples, and not carry out processing. Sometimes the laboratory is able to rectify a situation – and although turnaround times may be affected, it avoids having to arrange for samples to be taken again.

Summary List for Sample Rejection

- Incorrect sample types received:
 - *Basic incorrect blood tube/other sample.*
 - *Samples without the appropriate preservative (e.g. acidified urine samples).*
 - *Samples that are received ambient, when a frozen sample is required.*
 - *Samples that are received unprotected from light, when they are required to be covered at the point of venepuncture.*
- Samples in incorrect containers (e.g. cervical cytology must be a ThinPrep vial; urine cytology must be in a uricite container).
- Insufficient sample received.
- No sample received.
- Labelling or form issues (mislabeled/unlabelled/no forms/no clinical information).
- Clotted/haemolysed/lipaemic/icteric samples.
- Sample is broken or has leaked in transit.
- Stability time has been exceeded. Stability time is test dependant, and also refers to tests that can only be carried out on certain days of the week.

Quality assurance

- Sample contamination (e.g. being in the same bag as a leaking sample).
- Samples are high risk or infectious.
- Samples that are received in expired tubes.

Department Specific

- Sample Reception will not accept samples packaged with needles of any kind.
- Haematology cannot accept frozen whole blood for testing.
- Coagulation cannot accept over or under filled samples for testing.
- Coagulation cannot accept previously frozen samples that have thawed in transit.
- Biochemistry cannot accept previously frozen samples that have thawed in transit.
- Biochemistry cannot accept samples that display antibody interference.
- Biochemistry cannot accept samples that have had separation delays/un-centrifuged samples that have been stored in the fridge.
- Biochemistry cannot accept paraprotein resulting in viscous samples.
- Biochemistry cannot accept CSF protein that is blood stained.
- Immunology cannot accept TBQ kits that:
 - *Do not contain all of the appropriate tubes.*
 - *Are incubated for more than the specified 16 hours.*
 - *Have passed the incubation time period.*
 - *Are over or under filled.*
- Microbiology cannot accept samples in non-sterile containers or in formalin.
- Referrals cannot accept samples without three points of identification for DRP testing.
- Referrals cannot accept samples that are not labelled by hand for blood group testing.
- Molecular Pathology cannot accept samples for Haemophilia testing without informed consent.
- Cervical Cytology cannot accept over or under filled samples for testing.
- Cervical Cytology cannot accept samples received within three months of the previous test in order to allow epithelial cells to regenerate.
- Urine cytology cannot accept delayed samples unless they have been refrigerated.

Samples deemed to be PRECIOUS (e.g. CSF, fluid, tissue, bone marrow and paediatric samples) will not be discarded by the laboratory. Results will include a comment relating to the condition of the sample (e.g. sample unlabelled).

Quality assurance

CONSULTANT ADVICE AND OPINION

Each department in the laboratory is consultant led. For doctors wanting clinical advice or professional support, TDL consultants can be contacted via the laboratory. Contact the consultant Haematologist to make arrangements for venesections for Haemochromatosis and polycythaemia.

TDL MEDICAL CONSULTANTS

GROUP MEDICAL DIRECTOR

Dr Rachael Liebmann
BSc Hons, MB, BCh, BAO, FRCPath
FAcadMed, SFFMLM

HAEMATOLOGY / BLOOD TRANSFUSION

Professor Marie Scully
MRCP, FRCPath

Dr John Paul Westwood
MD FRCPath

Dr Adrian Bloor
FRCP, FRCPath

Dr Clare Barnes
MRCP, FRCPath

Dr Vivienne Andrews
FRCPath

ANDROLOGY

Dr Sheryl Homa
PhD ARCS FIBMS

BIOCHEMISTRY

Dr Paul Holloway
FRCPath

Dr Royce Vincent
FRCPath

Dr Denise Darby
MRCP, FRCPath

Professor Carel le Roux
FRCPath

Dr Gilbert Weiringa
PhD MRCPPath

Dr Frank Geoghan
MRCPPath

MICROBIOLOGY

Dr Robin Smith
FRCPath

Dr Sophie Collier
FRCPath

IMC

Dr Vanya Gant
FRCPath

Dr Michael Kidd
FRCPath

Dr Damien Mack
FRCPath

Dr Indran Balakrishnan
FRCPath

Dr Simon Warren
FRCPath

Dr Stephen Mephram
FRCPath

Dr Johnathan Lambourne
FRCPath

Dr Emanuel Wey
FRCPath

Dr Edward Kaczmariski
FRCPath

FLOW CYTOMETRY

Dr Raj Gupta
MRCP, FRCP, FRCPath

Dr Geraldine Soosay
FRCPath

IMMUNOLOGY

Dr Scott Pereira
FRCPath

Professor Suranjith Seneviratne
DPhil (Oxon) FRCP, FRCPath

VIROLOGY

Dr Mark Atkins
FRCPath

CYTOLOGY

Dr Mary Falzon
MRCS, LRCP, FRCPath

Dr Rachael Liebmann
BSc Hons, MB, BCh, BAO, FRCPath
FAcadMed, SFFMLM

Dr Geraldine Soosay
FRCPath

GENETICS: MOLECULAR/ CYTOGENETICS

Professor Michael Patton
FRCP, FRCPath
Consultant Clinical Geneticist

Special instructions for samples

- 1 Contact the laboratory for special sample tubes/containers/instructions.
- 2 Confirmation of not negative drug screens by LCMS/MS may take up to 5 days.
- 3 Clinical history essential and protect from light.
- 4 Send to the laboratory without delay.
- 5 Do not send sample to the laboratory between Friday noon and Monday morning.
- 6 Contact the Referrals Department before taking and sending sample to the laboratory.
- 7 Sample should be separated and frozen if sending overnight.
- 8 DRP Form required. DRP Form can be found at the back of the guide.
- 9 Clinical history must be provided.
- 10 Contact the laboratory for special stability tubes for lymphocyte subsets – or take an EDTA sample and ensure same day delivery to the laboratory, Monday to Friday noon (do not send sample between Friday noon and Monday morning).
- 11 Patient consent required. Consent Form can be found at the back of this guide.
- 12 Please provide one sample for each person being tested.
- 13 Protect from light.
- 14 Provide details of travel history.
- 15 Ammonia
Sample: EDTA plasma only. Full tubes and tightly stoppered. On ice, centrifuged and analysed 20-30 mins post venepuncture (or plasma can be frozen). If haemolysed gives falsely high results.
Patient: Fasting. Avoid smoking.
- 16 Lactate
Sample: Fluoride oxalate plasma only.
On ice and separate from cells 15-30 mins, analyse promptly. Handle with care as sweat contains large amounts of lactate. No tourniquet.
Patient: Rest 30 mins prior to test.
- 17 Homocysteine
Should be spun and separated with 1 hour of venepuncture.
- 18 Citrate Samples
Samples should be double spun and separated and frozen within 4-8 hours of sample taking, if a delay is expected with transportation to the laboratory, samples must be transported as frozen.
- 19 Must include patient's age, height and weight.
- 20 Sample types: FCRU or PCR swab or TPV or Semen.
- 21 Urine cytology container, ideally first catch, mid-morning specimen.
- 22 Must be fresh.
- 30 Collect sample at end of exposure.
- 33 Sample must be labelled by hand with first name, family name, gender and date of birth detailed on sample and form. Do not use labels other than the tube label.
- 34 Samples must arrive in the laboratory on the same day of sample taking or contact the laboratory.
- 35 Patient should be fasting and resting for 30 mins before sample taking. Samples need handling urgently.
- 36 Renin: Sample collected either upright/active or resting/supine (3 hours lying).
- 37 Provide sample time and date of collection.
- 38 EDTA sample should not be separated: send whole blood.
- 39 Urgent samples have a 3 day TAT if genotype is required for prenatal diagnosis or two weeks TAT if urgent for other factors.
- 40 Informed Consent is required for these tests.
- 41 Recommendation for patient to attend Patient Reception for sample taking.
- 42 LGV can be added to a positive chlamydia sample using the same swab if requested within 4 days of receipt of result.

Example of profile panel information

Profile name	PRE-TRAVEL SCREEN (DVT)	
Profile content	FBC Factor II Prothrombin Gene Factor V Leiden Anticardiolipin Antibodies	
Turnaround time		TAT 5 DAYS
Sample requirements	DVT1 	Code
	Reference to sample taking and special handling instructions (see above)	

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

TDL Screening Profiles DL1–DL12

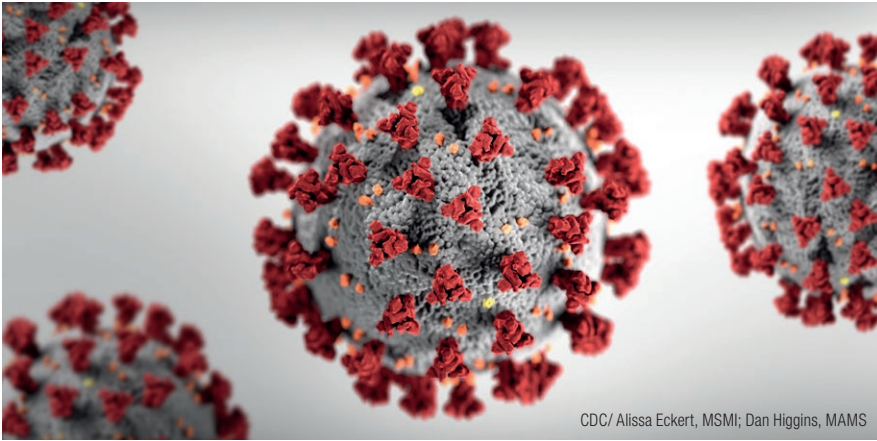
<div>DL1 BIOCHEMISTRY PROFILE</div> <div>Urea and Electrolytes Sodium, Potassium, Chloride, Bicarbonate, Urea, Creatinine, eGFR Liver Function Tests Bilirubin, Alk Phos, AST, ALT, Gamma GT, Total Protein, Albumin, Globulin Cardiac/Muscle Enzymes LDH, CK Bone Markers Calcium, Phosphate, Uric Acid Glucose Triglycerides Cholesterol Iron Total Iron Binding</div> <div><div>TAT 4 HOURS</div><div>DL1</div></div> <div><div><div>plus HDL Cholesterol LDL Cholesterol Non-HDL Cholesterol</div><div>DL1L</div></div><div><div>B</div><div>G</div></div></div>	<div>DL2 BIOCHEMISTRY (24 PARAMETERS) & HAEMATOLOGY PROFILE</div> <div>HAEMATOLOGY FBC with 5-part Diff ESR BIOCHEMISTRY Urea and Electrolytes Sodium, Potassium, Chloride, Bicarbonate, Urea, Creatinine, eGFR Liver Function Tests Bilirubin, Alk Phos, AST, ALT, Gamma GT, Total Protein, Albumin, Globulin Cardiac/Muscle Enzymes LDH, CK Bone Markers Calcium, Phosphate, Uric Acid Glucose Triglycerides Cholesterol Iron/TIBC</div> <div><div>TAT 4 HOURS</div><div>DL2</div></div> <div><div><div>plus HDL Cholesterol LDL Cholesterol Non-HDL Cholesterol</div><div>DL2L</div></div><div><div>A</div><div>B</div><div>G</div></div></div>	<div>DL3 HAEMATOLOGY PROFILE</div> <div>FBC with 5-part Diff ESR</div> <div><div>TAT 4 HOURS</div><div>DL3</div></div> <div><div>A</div></div>
<div>DL5 BIOCHEMISTRY & HAEMATOLOGY POSTAL PROFILE</div> <div>As DL4 DL5/DL5L do not include ESR and Phosphate as these results may be more affected by overnight transit times.</div> <div><div>TAT 4 HOURS</div><div>DL5</div></div> <div><div><div>plus HDL Cholesterol LDL Cholesterol Non-HDL Cholesterol</div><div>DL5L</div></div><div><div>A</div><div>B</div><div>G</div></div></div>	<div>DL6 GENERAL WELL PERSON PROFILE</div> <div>DL2 FT4/TSH Ferritin</div> <div><div>TAT 4 HOURS</div><div>DL6</div></div> <div><div><div>plus HDL Cholesterol LDL Cholesterol Non-HDL Cholesterol</div><div>DL6L</div></div><div><div>A</div><div>B</div><div>G</div></div></div>	<div>BIOCHEMISTRY (16 PARAMETERS) & HAEMATOLOGY PROFILE</div> <div>HAEMATOLOGY FBC with 5-part Diff ESR BIOCHEMISTRY Renal Function Urea, Creatinine, eGFR Liver Function Tests Bilirubin, Alk Phos, AST, ALT, Gamma GT, Total Protein, Albumin, Globulin Bone Markers Calcium, Phosphate, Uric Acid Glucose Triglycerides Cholesterol</div> <div><div>TAT 4 HOURS</div><div>DL4</div></div> <div><div><div>plus HDL Cholesterol LDL Cholesterol Non-HDL Cholesterol</div><div>DL4L</div></div><div><div>A</div><div>B</div><div>G</div></div></div>

TDL Screening Profiles DL1–DL12

DL7 WELL MAN PROFILE DL2 FT4/TSH Ferritin Prostate Profile TAT 4 HOURS DL7 DL7L <i>plus HDL Cholesterol LDL Cholesterol Non-HDL Cholesterol</i> A B G	DL8 WELL PERSON PROFILE DL2 FT4/TSH Ferritin Vitamin D TAT 4 HOURS DL8 DL8L <i>plus HDL Cholesterol LDL Cholesterol Non-HDL Cholesterol</i> A B G	DL9M SENIOR MALE PROFILE 60+ DL2 HDL/LDL Cholesterol HbA1C FT4/TSH Prostate Profile CRP Ferritin QFIT MSU Vitamin D (25 OH) Lp-PLA2 (PLAC) Test TAT 2 DAYS DL9M A B B G RU QFIT ⁴
DL9F SENIOR FEMALE PROFILE 60+ DL2 HDL/LDL Cholesterol HbA1C FT4/TSH CRP Ferritin QFIT MSU Vitamin D (25 OH) HE4 Lp-PLA2 (PLAC) Test TAT 2 DAYS DL9F A B B G RU QFIT ⁴	DL10 CARDIOVASCULAR RISK PROFILE 1 Cholesterol Triglycerides HDL Cholesterol LDL Cholesterol Non-HDL Cholesterol Apolipoprotein A Apolipoprotein B Lipoprotein (a) hsCRP Lp-PLA2 (PLAC) Test TAT 3 DAYS DL10 B B	DL11 CARDIOVASCULAR RISK PROFILE 2 Cholesterol Triglycerides HDL Cholesterol LDL Cholesterol Non-HDL Cholesterol Apolipoprotein A Apolipoprotein B Lipoprotein (a) Fibrinogen hsCRP Lp-PLA2 (PLAC) Test Homocysteine TAT 3 DAYS DL11 B B B C ³⁴
DL12 7 STI PROFILE BY PCR (7 PCR TESTS FROM 1 SAMPLE) Chlamydia trachomatis N. gonorrhoea Mycoplasma genitalium Macrolide Resistance Test (M.gen)* Ureaplasma Trichomonas vaginalis Gardnerella vaginalis Herpes Simplex I/II TAT 2 DAYS DL12 FCRU OR PCR Swab OR TPV OR Semen		

*included if **POSITIVE M.gen** is detected from the same sample

Testing for COVID-19 (SARS-CoV-2)



Illness caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has been termed 'COVID-19', derived from 'coronavirus disease 2019.' The number and range of tests for COVID-19 is developing apace, with emphasis on five key areas:

- Performance of assay: sensitivity, specificity, accuracy
- Speed of processing – how quickly can results be reported
- Sample type and ease of specimen collection
- Choice of tests: which test to use, and when?
- Price: dependant on method and manufacturer: PCR, Antigen and Antibody

TDL will update on testing developments throughout the year – and are currently running:

COVID-19 (SARS-CoV-2) RNA by PCR NEW

Results will be reported as Positive, Not Detected or Invalid if a test has failed. Always provide the patient's address and contact phone number.

Test Code: NCOV

Sample Type	PCR swab (COVID-19 Royal Mail Priority Tracked 24 kits provided)
Performance	Sensitivity 98.0%, Specificity 100%, Accuracy 98.8%
Analysers	Hologic Panther SARS CoV-2 Assay UKAS Accredited Hologic Panther Fusion UKAS Accredited QuantStudio/Reagents Thermofisher UKAS Accredited
Turnaround time	Within 48 hours of receipt of sample

SEE PAGE 96 FOR DETAILS OF COVID TESTS

Testing for COVID-19 (SARS-CoV-2)

COVID-19 (SARS-CoV-2) Abbott IgG Antibody NEW

The Abbott IgG Antibody test is CE marked for venous samples, and one of the PHE selected antibody tests.

Test Code: GCOV

Sample Type	SST/Serum B Venous (>14 days after onset of symptoms)
Performance	Specificity 99.1%, Sensitivity 97.5%
Analysers	Abbott Architect
Turnaround time	24 hours from receipt of sample

COVID-19 (SARS-CoV-2) Roche Elecsys Anti-SARS-CoV-2 Total Antibody NEW

Roche Elecsys Anti-SARS-CoV-2 will additionally detect IgM antibodies, reporting both IgG and IgM as a TOTAL antibody result. The Roche Antibody test is CE marked for both **venous and capillary** samples, and is one of the PHE selected antibody tests.

Test Code: TCOV

Sample Type	SST/Serum B Venous or Capillary (>14 days after onset of symptoms)
Performance	Specificity 100%, Sensitivity 97.4%
Analysers	Roche e801
Turnaround time	24 hours from receipt of sample

Self-collection CE marked IVD for COVID Postal kits NEW

The kits include a Royal Mail Tracked 24 return label.

Respiratory Viral Profile NEW

PCR swabs or Antibody samples can be sent to the laboratory:

- with your practice's courier collection
- delivered to Patient Reception at 76 Wimpole Street W1G 9RT
- delivered to the HALO Laboratory (WC1H 9AX)
- delivered to TDL Manchester (M50 2GY)
- posted to the Halo Laboratory, using next day delivery Royal Mail COVID Priority Tracked 24, provided with the swab pack.

















































RESPIRATORY VIRAL RNA SCREEN BY PCR	
Flu A Flu B Respiratory Syncytial Virus (RSV) COVID-19	NEW 2021 TAT 48 HOURS
FLU4	
PCR nasopharyngeal	

TDL reports all Antibody and PCR activity daily to Public Health England (PHE). It is a statutory requirement that laboratories notify this information and it is therefore essential that the patient's address and postcode are provided so that positive results can be followed by Test and Trace.

Biochemistry

TEST	CODE	SAMPLE REQ	TAT
5 HIAA	RU5H	PU ¹	5 days
5' Nucleotidase	5NT	B	5 days
6-Thioguanine Nucleotides	TGN	A A	2 weeks
21 Hydroxylase Ab's	21HA	B (Frozen)	10 days
Acetylcholine Receptor Autoantibodies	ACRA	B ⁴	5 days
Acetylcholinesterase Isoenzymes	ACEI	AF	7 days
Acid Phosphatase – Total	APT	B	5 days
Adenosine Deaminase	AD	A / B / Fluid	3 weeks
Adiponectin	ADIP	B	2 weeks
Albumin	ALB	B	4 hours
Alcohol (Medical) [Do not use alcohol swab prior to sample taking]	ALCO	G ¹	4 hours
Alcohol (Urine)	UALC	RU	4 hours
Aldolase	ALDO	B	5 days
Alk Phosphatase Isoenzymes	APIE	B	5 days
Alkaline Phosphatase	ALP	B	4 hours
Alpha 1 Antitrypsin (Serum)	A1AT	B	1 day
Alpha 1 Antitrypsin (Stool)	A1AF	RF	10 days
Alpha 1 Antitrypsin Genotype – PI*M, PI*S, PI*Z	GENE	A ⁹	4 weeks
Alpha 1 Glycoprotein	OROS	B (Frozen)	5 days
Alpha 1 Microglobulin	A1MG	RU ^{1,22}	10 days
Alpha 2 Macroglobulins	A2MG	B	5 days
Alpha Feto Protein (Maternal)	AFPM	B	4 hours
ALT (Alanine Aminotransferase) (SGPT)	ALT	B	4 hours
Aluminium (Blood)	ALUM	K	7 days
Amino Acid (Serum/Plasma)	AMIN	B	7 days
Amino Acid Quantitative (Urine)	UAAQ	RU	7 days
Amino-Laevalinic Acid (Urine)	RUAL	100mls PU	5 days
Ammonia	AMMO	A (Frozen) ¹⁵	4 hours
Amylase	AMY	B	4 hours
Amylase (Urine)	UAMY	CU	4 hours
Amylase Isoenzymes	AMYI	B	5 days
Amyloidosis (Amyloid A Protein)	SAA	B	5 days
Androstenediolglucuronide	ANDG	B	3 weeks
Angiotensin II	ANG2	A (Frozen)	2 weeks
Angiotensin Converting Enzyme	ACE	B	4 hours
Angiotensin Converting Enzyme – CSF	ACEF	CSF (Frozen)	2 weeks
Antimony (Urine)	ANTI	RU ³⁰	10 days
Antimullerian Hormone (AMH Plus)	AMH	B	4 hours
AP50 Alternative Hemolytic Complement	AP50	B (Frozen)	2 weeks
Apolipoprotein A1	APOA	B	3 days
Apolipoprotein B	APOB	B	3 days
Apolipoprotein C	APOC	B	3 months
Apolipoprotein E (12 hours fasting)	APOE	B (fasting)	5 days

Biochemistry

TEST	CODE	SAMPLE REQ	TAT
Arsenic (Blood)	ARS	 or 	5 days
Arsenic (Urine)	ARSE	RU ³⁰	5 days
Arylsulphatase A	ARYL	 ^{5,6}	8 weeks
Aspartate Transaminase (AST) (SGOT)	AST		4 hours
Bence-Jones Protein	RBJP	1 x 30mls (RU)	5 days
Beta 2 Microglobulin (Serum)	B2MG		2 days
Beta 2 Microglobulin (Urine)	UB2M	RU	3 days
Beta-Glucuronidase (Sly Disease)	BGLU	  ^{9,4}	8 weeks
Bicarbonate	HCO3		4 hours
Bile Acids – Serum	BILE		4 hours
Bilirubin (Direct/Indirect)	DBIL		4 hours
Bilirubin (Total)	BILI		4 hours
Bilirubin (Urine)	UBIL	RU	1 day
Biotinidase	BIOT	 (Frozen plasma) ⁴	3 weeks
Bismuth	BISM		5 days
BNP (NT-pro BNP)	BNP		4 hours
Bone Alkaline Phosphatase	BALP	 (Frozen)	2 weeks
Bone Screen	BONE	 CU	4 hours
Bone Screen (Bloods only)	BON2		4 hours
BUN (Blood Urea Nitrogen)	BUN		4 hours
C Reactive Protein	CRP		4 hours
C Reactive Protein (High Sensitivity)	HCRP		4 hours
C1 Esterase: Function & Total	FC1E	  (Plasma Frozen) ^{4,18}	10 days
C1q Binding Immune Complex	IMCP		5 days
Cadmium (Blood)	CADM	 or 	5 days
Cadmium (Urine)	URCD	RU ³⁰	5 days
Calcium	CA		4 hours
Calcium (24 hour Urine)	UCA	PU	4 hours
Calcium/Creatinine Ratio	CACR	RU 	4 hours
Carbohydrate Deficient Glycoprotein	CDG		2 weeks
Carbohydrate Deficient Transferrin (CDT)	CDT	 ⁴	3 days
Cardiac Enzymes (not chest pain)	CENZ		4 hours
Cardiovascular Risk Profile 1	PP10	 	3 days
Cardiovascular Risk Profile 2	PP11	     ³⁴	3 days
Carnitine – Free & Total	CARN	  (Frozen Plasma)	10 days
Ceruloplasmin	CERU		1 day
Chest Pain Profile	CPP		STAT
Chloride	CL		4 hours
Cholesterol	CHO		4 hours
Cholesterol (Familial Hypercholesterolaemia)	GENE	  ⁹	4 weeks
Cholinesterase (Blood)	CHRC		5 days
Cholinesterase (Serum/Pseudo)	CHPS		4 hours
Chromium (Blood)	CHRO		5 days

Biochemistry

TEST	CODE	SAMPLE REQS	TAT
Chromium (Urine)	URCR	RU ³⁰	10 days
Chromogranin A	CGA	B	5 days
Chromogranin A & B	MTAB	J ¹	3 weeks
Citrate (Blood)	CITR	B	5 days
Citrate (Urine)	UCIT	CU (Frozen)	5 days
CK (MB Fraction)	CKMB	B	4 hours
CK Isoenzymes	CKIE	B	5 days
Cobalt (Blood)	COB	A	5 days
Cobalt (Serum)	COBB	B	5 days
Cobalt (Urine)	COBA	RU ³⁰	5 days
Coenzyme Q10	CQ10	B	2 weeks
Cold Agglutinin	CAGG	J ¹	5 days
Collagen (Type I, II, IV) Antibodies	COAB	B	10 days
Collagen Type 1 Cross-Linked N-Telopeptide – NTX	NTX	2nd EMU	2 weeks
Complement C1q	C1Q	B	5 days
Complement C2	C2	B	10 days
Complement C5	C5A	B	2 weeks
Complement C6	C6	B (Frozen)*	5 weeks
Complement C7	C7	B (Frozen)*	5 weeks
Complement C8	C8	B (Frozen)*	5 weeks
Complement C9	C9	B (Frozen)*	5 weeks
Complement Factor H	FACH	B	3 weeks
Copper (Serum)	COPP	B	5 days
Copper (Urine)	URCU	CU	5 days
Cortisol Binding Globulin	CBG	B (Frozen)	1 month
Creatine Kinase (CK, CPK)	CKNA	B	4 hours
Creatinine	CREA	B	4 hours
Creatinine (Urine)	UCR	CU	4 hours
Creatinine Clearance	CRCL	B CU	4 hours
Crosslaps (Serum DPD)	SDPD	B (Freeze within 24 hours)	4 days
Cyclic Amp (Urine)	CAMP	CU (Frozen)	5 days
Cyclosporin (Monoclonal)	CYCL	A	1 day
Cystatin C	CYCC	B	5 days
Cystine – Quantitative (Beta-CTX)	QCYS	PU	5 days
Deoxypyridinoline (DPD) – Serum	SDPD	B (Freeze within 24 hours)	4 days
Deoxypyridinoline (DPD) – Urine	DPD	EMU	4 days
Diabetic Profile 1	DIAB	A G	8 hours
Diabetic Profile 2	DIA2	A G RU	2 days
Electrolytes	ELEC	B	4 hours
Electrolytes (Urine)	UELE	CU	4 hours
ELF/Enhanced Liver Fibrosis	ELF	B	5-7 days

* Separate and freeze within 2 hours after collection.









































Key: See page 21 for sample taking and special handling instructions.

Biochemistry

TEST	CODE	SAMPLE REQ	TAT
Eosinophil Cationic Protein	ECP	B	7 days
Faecal Fat (1 Day Collection)	TFFA	LF ⁶	5 days
Faecal Fat (3 day)	FFAT	LF ⁶	5 days
Faecal Lactoferrin	FLAC	RF	5 days
Faecal Sugar Chromatography	FCRO	RF (Frozen)	3 weeks
Faecal Urobilinogen	FURO	RF	5 days
Fat Globules in Faeces	FGLO	RF	1 week
Ferritin	FERR	B	4 hours
Fibrotect (Liver Fibrosis)	FIBT	B	2 weeks
Fluoride (Urine)	UFL	RU	5 days
Folate (Red Cell)	RBCF	A	2 days
Folate (Serum)	FOLA	B	1 day
Free Fatty Acids	FFA	B (Frozen) ¹	10 days
Fructosamine	FRUC	B	1 day
Fructose – Plasma	FRU	G ⁷ (Frozen)	5 days
Galactose-1-Phosphate Uridyltransferase	GAL1	H ^{5,6}	2 weeks
Galactosidase – Alpha*	GALA	J *	6 weeks
Gall Stone Analysis	RSTA	STONE	10 days
Gamma GT	GGT	B	4 hours
Gastrin	GAST	B (Frozen)	5 days
Globulin	GLOB	B	4 hours
Glucagon	GLUG	J ¹	10 days
Glucose	RBG	G	4 hours
Glucose Tolerance Test			See page 129
Haemochromatosis – HFE common mutations C282Y + H63D	HMD	A ⁹	3 days
Haemosiderin (Urine)	HSID	EMU	2 weeks
Haptoglobin	HAPT	B	5 days
HbA1c	GHB	A	6 hours
HDL Cholesterol	HDL	B	4 hours
HDL2 & HDL3 Fractions	HDLF	B	3 weeks
Homocysteine (Quantitative)	HOMO	B ¹⁷	1 day
Homocysteine (Urine)	HCYS	CU	2 weeks
Homovanillic Acid (HVA)	HVA	PU	5 days
Hyaluronic Acid	AHT	B	1 week
Hydroxybutyrate Dehydrogenase	HBD	B (Frozen)	1 week
Hydroxyprolene	UHYD	CU	2 weeks
IgG Subclasses	IGSC	B	4 days
Immunoglobulin A	IGA	B	4 hours
Immunoglobulin D	IGD	B	5 days
Immunoglobulin E – Total	IGE	B	1 day

* Sample must reach TDL Referrals Dept. urgently, to be tested within 24 hours of collection.
Monday–Thursday only. Referrals to send immediately

Biochemistry

TEST	CODE	SAMPLE REQ	TAT
Immunoglobulin G	IGG		4 hours
Immunoglobulin M	IGM		4 hours
Immunoglobulins (IgG, IgM, IgA)	IMM		4 hours
Insulin-Like Growth Factor 2	IGF2	 ⁶	1 month
Iodide – Urine	UIOD	RU	1 week
Iodine – Serum	IODI		1 week
Ionised Calcium	ICPA		5 days
Iron (TIBC included)	FE		4 hours
Iron Overload Profile	IOP	   ⁹	3 days
Iron Status Profile	ISP		4 hours
Lactate (Plasma)	LACT	 ¹⁶	1 day
Lactate Dehydrogenase (LDH)	LDH		4 hours
Lactate Pyruvate Ratio	LPR	J ¹	4-6 weeks
Lactose Tolerance Test			See page 129
LDH Isoenzymes	ISOL		5 days
LDL7 Subfractions	LDL7		10 days
Lead (Blood)	LEAD		5 days
Lead (Urine)	URPB	RU	5 days
Leptin	LEPT	 ¹⁹	5 days
Leucine Amino Peptidase	LAP		5 days
Lipase	LIPA		4 hours
Lipid Profile	LIPP		4 hours
Lipoprotein (a)	LPOA		4 hours
Lipoprotein Electrophoresis	LEL		5 days
Lithium (take 12 hours after dose)	LITH		4 hours
Liver Fibrosis (Enhanced Liver Fibrosis ELF)	ELF		5-7 days
Liver Fibrosis Fibrotest	FIBT		2 weeks
Liver Function Tests	LFT		4 hours
Lp-PLA2 (PLAC) Test	PLA2		2 days
Lysosomal Enzyme Screen	LE	  ⁶	2 months
Lysozyme	LYSO		5 days
Magnesium (Serum)	MG		4 hours
Magnesium (Urine)	URMG	PU	1 day
Manganese (Serum)	MANG		5 days
Mannose Binding Lectin	MBL		3 weeks
Mercury (Blood)	MERC	 or 	5 days
Mercury (Urine)	URHG	RU ¹	5 days
Methaemoglobin	METH		3 days
Methaqualone	METQ	RU	5 days
Methylmalonic Acid – Serum	MMAS		5 days
Methylmalonic Acid – Urine	MMA	CU	2 weeks
Microalbumin (Urine)	UMA	RU	4 hours
Mucopolysaccharides	MPS	RU (Frozen)	3 weeks
Myeloma Screen	MYEL	   RU	5 days

Key: See page 21 for sample taking and special handling instructions.

Biochemistry

TEST	CODE	SAMPLE REQ	TAT
Myoglobin (Serum)	SMYO	B	4 hours
Myoglobin (Urine)	UMYO	RU	5-10 days
Newborn Screening Panel	GUTH	J ¹	2 weeks
Nickel (Serum)	NICK	B	5 days
Nickel (Urine)	NICU	RU	10 days
NMP22 (Bladder tumour)	NMP	J ¹	4 days
Oligosaccharides	UOLI	RU	6 weeks
Orosomucoid (A1AG – Alpha 1 Glycoprotein)	OROS	B (Frozen)	5 days
Osmolality (Serum)	OSMO	B	1 day
Osmolality (Urine)	ROSM	RU	1 day
Osteoporosis Screen	OPS	B B	4 days
Oxalate (Plasma)	POXA	A (Frozen)	7 days
Oxalate (Urine)	UOXA	PU	5 days
Pancreatic Peptide	PP	J	4 weeks
Parathyroid Related Peptide	PTRP	2ml A Plasma frozen (Freeze immediately) ¹	2 weeks
PEth (Phosphatidylethanol)	PETH	A ³⁸	5-7 days
Phencyclidine (PCP)	DUST	RU	5 days
Phosphate	PHOS	B	4 hours
Phosphate (24 hour Urine)	UPH	PU	4 hours
PLAC Test (Lp-PLA2)	PLA2	B	2 days
Plasminogen	PLAS	C (Frozen plasma) ⁴	5 days
Plasminogen Activator Inhibitor – 1	PAI1	C (Frozen plasma)	2 weeks
Porphyria (Blood)	PORP	A ³	15 days
Porphyria (Faeces)	FPOR	RF ³	3 weeks
Porphyria Full Screen (Total: Urine, Stool, Blood)	PORS	A RU , RF ³	3 weeks
Porphyria Screen (Urine)	RPOR	RU ³	3 weeks
Potassium	K	B	4 hours
Pregnancy (Serum) [Quantitative]	QHCG	B	4 hours
Pregnancy Test (Urine)	PREG	RU	4 hours
Procalcitonin	PCAL	B (Frozen) ^{4,7}	1 day
Procollagen 1 Peptide N-Terminal (NTX)	P1NP	B	5 days
Procollagen III Peptide	PRCO	B	5 days
Propoxyphene	DPRO	RU	5 days
Prostatic Acid Phosphatase	PACP	B (Frozen)	3 days
Protein (Urine)	UPRT	CU	4 hours
Protein 14.3.3 (Creutzfeldt–Jakob Disease)	CJD	CSF (Frozen)	5 weeks
Protein Electrophoresis incl. immunoglobulin	PRTE	B	2-4 days
Protein Total (Blood)	PROT	B	4 hours
Protein/Creatinine Ratio (Urine)	UCPR	RU	4 hours
Renal Calculi Screen (Metabolic)	RSPR	J ⁶	5 days
Renal Stone Analysis	RSTA	STONE	10 days
Retinol Binding Protein	RBP	B	3 days
Salicylates	SALI	B	4 hours

Biochemistry

TEST	CODE	SAMPLE REQ	TAT
Selenium (Serum)	SELE	B	4 days
Selenium (Whole Blood)	SELR	A or H	4 days
Serum Free Light Chains	SLC	B	1 week
Silver (Blood)	SILV	B	5 days
Silver (Urine)	USIL	RU	5 days
Sodium	NA	B	4 hours
Superoxide Dismutase Inhibitor	SODI	A / H	5 days
Thiopurine Methyl Transferase	TPMT	A ⁵	5 days
Tissue Polypeptide Antigen	TPA	B	1 week
Total Acid Phosphatase	APT	B	5 days
Total Bile Acid/Bile Salts	BILS	B	1 week
Total IgE	IGE	B	1 day
Transferrin	TRAN	B	1 day
Transferrin Electrophoresis	TREL	B	2 weeks
Triglycerides	TRI	B	4 hours
Trimethylaminuria (Fish Odour Syndrome)	FOS	PU	6 weeks
Troponin T (High sensitive)	TROT	B	4 hours
Tryptase	STRY	B	2 days
Tumour Necrosis Factor – Alpha	TNF	B (Frozen) ⁴	2 weeks
Urate (Uric acid)	UA	B	4 hours
Urea	UREA	B	4 hours
Urea (Urine)	UURE	CU	4 hours
Urea and Electrolytes	U/E	B	4 hours
Urea Electrolytes (Urine)	UELE	CU	4 hours
Uric Acid (Serum)	UA	B	4 hours
Uric Acid (Urine)	UURI	CU	4 hours
Urine Free Light Chains	UFLC	RU	1 week
Urine Organic Acids	UORG	RU (Frozen)	3 weeks
Urine Steroid Screen (Steroid Hormones)	USTE	CU or RU ⁹	2 weeks
Urine Sugar Chromatography	UCRO	RU (Frozen)	3 weeks
Urobilinogen (Urine)	UURO	RU	1 day
Very Long Chain Fatty Acids	VLCF	A or H (Frozen) ⁹	4-6 weeks
Vitamin B12 (Active)	B12	B	1 day
Vitamin B12 (Active)/Red Cell Folate	B12F	A B	2 days
Vitamin B12 (Total)	TB12	B	1 day
Vitamin D (25-OH)	VITD	B	4 hours
VLDL Cholesterol	VLDL	B ¹³	1 week
VMA	UVMA	PU ¹	5 days

LIPID PROFILE

Triglycerides
Cholesterol
HDL Cholesterol
LDL Cholesterol
Non-HDL Cholesterol

TAT
4
HOURS

LIPP

B

UREA AND ELECTROLYTES

Sodium
Potassium
Chloride
Bicarbonate
Urea
Creatinine

TAT
4
HOURS

U/E

B

LIVER FUNCTION TESTS

Bilirubin
ALT
AST
Total Protein
Alkaline Phos
Albumin
Globulin
Gamma-GT

TAT
4
HOURS

LFT

B

IRON STATUS PROFILE

Iron
Total Iron Binding Capacity
Ferritin
Transferrin Saturation

TAT
4
HOURS

ISP

B

IRON OVERLOAD PROFILE

Iron Status Profile
Haemochromatosis
Mutation
H63D/C282Y)

TAT
3
DAYS

IOP

A A B⁹

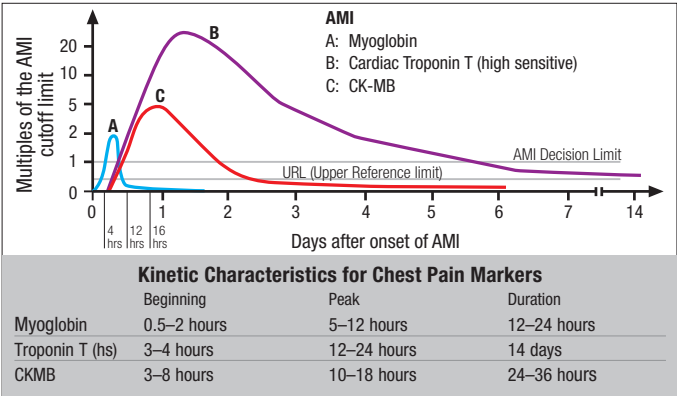
MYELOMA SCREEN

FBC and ESR
Biochemistry Profile
Protein Electrophoresis
Immunoglobulins
(IgA, IgG, IgM)
Bence-Jones Protein

TAT
5
DAYS

MYEL

A B G RU



Troponin T (high sensitive)

This assay can be used to aid in the differential diagnosis of acute coronary syndrome to identify necrosis, e.g. acute myocardial infarction. As a result of its high tissue-specificity, cardiac troponin T is a cardio-specific, highly sensitive marker for myocardial damage. Cardiac Troponin T (hs) increases approximately 3-4 hours after myocardial infarction and may persist for up to 2 weeks.

Biochemistry

BONE SCREEN	
24 hour urinary calcium 24 hour urinary phosphate Urea and Electrolytes Alkaline Phosphatase Total Protein Albumin Globulin Calcium	TAT 4 HOURS
BONE	

B **CU**

BONE SCREEN (BLOODS ONLY)	
Urea and Electrolytes LFT's Calcium Phosphate Vitamin D (25 OH)	TAT 4 HOURS
BON2	

B

OSTEOPOROSIS SCREEN	
Alkaline Phosphatase Calcium Albumin Phosphate Serum Crosslaps (DPD) Vitamin D (25 OH)	TAT 4 DAYS
OPS	

B **B**

CARDIOVASCULAR RISK PROFILE 1	
Cholesterol Triglycerides HDL Cholesterol LDL Cholesterol Non-HDL Cholesterol Apolipoprotein A Apolipoprotein B Lipoprotein (a) hsCRP Lp-PLA2 (PLAC) Test	TAT 3 DAYS
PP10	

B **B**

CARDIOVASCULAR RISK PROFILE 2	
Cholesterol Triglycerides HDL Cholesterol LDL Cholesterol Non-HDL Cholesterol Apolipoprotein A Apolipoprotein B Lipoprotein (a) Fibrinogen hsCRP Lp-PLA2 (PLAC) Test Homocysteine	TAT 3 DAYS
PP11	

B **B** **B** **C** ³⁴

CHEST PAIN PROFILE	
Myoglobin CK MB Fraction Troponin T	STAT
CPP	

B

DIABETIC PROFILE 1	
Glucose HbA1c	TAT 8 HOURS
DIAB	














































A **G**

DIABETIC PROFILE 2	
Glucose HbA1c Microalbumin	TAT 2 DAYS
DIA2	

A **G** **RU**


















































Haematology

All citrate samples  sent by post or with an overnight delay must be double spun and sent frozen.

TEST	CODE	SAMPLE REQS	TAT
Anaemia Profile	ANAE	  	2 days
Antenatal Profile	ANTE	  ³³    	3 days
APTT/KCCT	KCCT	 ¹⁸	4 hours
Atypical Antibody Screen (handwritten tube label)	AASC	 ^{22,33}	2 days
Blood Film Examination	FILM		1 day
Blood Group [†]	ABO	 ^{22,33}	2 days
Carboxyhaemoglobin	CBHB		1 week
Coagulation Profile 1	CLPF	 ¹⁸	4 hours
Coagulation Profile 2	CLOT	  ¹⁸	4 hours
D-Dimers (Fibrinogen Degradation Products)	DDIT	 ⁴	4 hours
DVT/Pre-travel Screen	DVT1	   ⁹	5 days
ESR	ESR		4 hours
Fibrinogen	FIB	 ^{4,18}	4 hours
Full Blood Count	FBC		4 hours
Haematology Profile	PP3		4 hours
Haemoglobin	HB		4 hours
Immune Function Evaluation (Total)	TIE	 +  ^{5,10}	7 days
INR	PTIM	 ¹⁸	4 hours
Lymphocyte Subsets (CD3/CD4/CD8)	LYSS	 ¹⁰	1 day
Malarial Parasites	MALP	 ^{4,9,14}	STAT
Mean Cell Volume (MCV)	MCV		4 hours
Microfilaria Blood Film	MICF		STAT
Natural Killer Profile 2	NKP2		2 days
PAI1 4G/5G Polymorphism	PAIP		10 days
Paul Bunnell (Monospot)	PAUL	 or 	8 hours
Pre-Travel Screen (DVT)	DVT1	   ⁹	5 days
Prothrombin Time	PTIM	 ¹⁸	4 hours
Prothrombin Time + Dose	PT+D	 ¹⁸	4 hours
Reticulocyte Count	RETC		4 hours
Thrombin Time	THRO	 ¹⁸	4 hours
Vitamin K (With PIVKA II)	VITK	 ¹³	10 days

[†] The tube's own label must be completed by hand. This must correspond with same name and date of birth details as given on the request form. Do not affix additional computerised or hand written labels.








SPECIAL HAEMOSTASIS

TEST	CODE	SAMPLE REQ	TAT
Activated Protein C Resistance	APCR	 (Frozen) ^{4,18}	3 days
ADAMTS-13 Activity	CP13	 (Frozen)	3 days
ADAMTS-13 Antibody	A13A	 (Frozen)	1 month
Antithrombin III	A111	 (Frozen) ^{4,9,18}	3 days
Anti-Xa Apixaban monitoring	APIX	 (Frozen)*	3 days
Anti-Xa Fondaparinux monitoring	FOND	 (Frozen)*	3 days
Anti-Xa- LMWH monitoring	LMWX	 (Frozen)*	3 days
Anti-Xa- Rivaroxaban monitoring	RIVA	 (Frozen)*	3 days
Factor II Assay	FAC2	 (Frozen) ^{9,18}	5 days
Factor V Assay	FAC5	 (Frozen) ^{9,18}	5 days
Factor VII Assay	FAC7	 (Frozen) ^{9,18}	5 days
Factor VIII Assay	FAC8	 (Frozen) ^{9,18}	5 days
Factor VIII Inhibiting Antibody	F8IA	  ¹⁸	2 weeks
Factor IX Assay	F1X	 (Frozen) ^{9,18}	5 days
Factor IX Inhibiting Antibody	F9IA	  ¹⁸	2 weeks
Factor X Assay	FX	 (Frozen) ^{9,18}	5 days
Factor XI Assay	FX1	 (Frozen) ^{9,18}	5 days
Factor XII Assay	FX11	 (Frozen) ^{9,18}	5 days
Factor XIII Assay	FA13	 (Frozen) ^{9,18}	5 days
Hughes Syndrome	LUPA	  ^{4,18}	2 days
Lupus Anticoagulant and Anticardiolipin Abs	LUPA	  ^{4,18}	2 days
Lupus Anticoagulant only	LUPC	 ¹⁸	2 days
Miscarriage/Thrombotic Risk Profile	PROP	      ¹⁸	5 days
P2Y12 Receptor Platelet Function Analysis (Clopidogrel Resistance)	P2Y	 (Whole blood)**	1 day
Platelet Aggregation Studies	PLAG	 ^{5,6}	3 days
Protein C	PRC	 (Frozen) ^{4,9,18}	3 days
Protein S Activity	PS1	 (Frozen)	5 days
Protein S Free Ag	FPRS	 (Frozen) ^{4,9,18}	3 days
Taipan Snake Venom Time	TTVT	 ¹⁸	1 week
Thrombotic Risk Profile	PROP	      ¹⁸	5 days
Viscosity (Plasma)	VISC	 ⁴	3 days
Von Willebrand Profile	FVWF	   ^{4,12}	5 days
Von Willebrands Multimers	VWM	   ¹⁸	3 months









* Please state drug and time of dose on request.

** Deliver directly to 60 Whitfield Street, Haemostasis Laboratory

SPECIAL HAEMATOLOGY

TEST	CODE	SAMPLE REQ	TAT
Coombs (Direct Antiglobulin Test)	COOM		2 days
Erythropoietin	ERY		4 days
G6PD	G6PD		3 days
Haemoglobin Electrophoresis	HBEL		4 days
HFE gene (Haemochromatosis) – common mutations C282Y + H63D	HMD	 ⁹	3 days
Sickle Solubility	SSOL		4 days
Thalassaemia Screen	HBEL		4 days

FLOW CYTOMETRY

TEST	CODE	SAMPLE REQ	TAT
Bone Marrow (Aspirate)	BMAS	J ¹	14 days
Bone Marrow (Trephine Biopsy)	BMI	J ¹	3 days
CD3/CD4/CD8	LYSS	 ¹⁰	1 day
CD16	CD16	 ⁴	1 day
CD19 B Cells	CD19	 ⁴	1 day
CD20	CD20	 ¹⁰	2 days
CD25	CD25	 ¹⁰	2 days
CD56	CD56	 ⁴	1 day
CD57	CD57		1 day
Hams Test for PNH (CD59)	HAMS	J ^{34,5}	5 days
Leukaemia Immunophenotyping	LYPT	 ^{4,5}	5 days

Haematology

HAEMATOLOGY PROFILE FBC + 5 part Diff ESR TAT 4 HOURS PP3	COAGULATION PROFILE 1 Prothrombin Time APTT Fibrinogen TAT 4 HOURS CLPF	COAGULATION PROFILE 2 FBC + 5 part Diff Prothrombin Time APTT Fibrinogen TAT 4 HOURS CLOT
A	C ¹⁸	A C ¹⁸
ANAEMIA PROFILE FBC + 5 part Diff ESR Iron, TIBC Ferritin B12 (Active) Folate (RBC) TAT 2 DAYS ANAE	PRE-TRAVEL SCREEN (DVT) FBC Factor II Prothrombin Gene Factor V Leiden Anticardiolipin Antibodies TAT 5 DAYS DVT1	VON WILLEBRAND PROFILE Von Willebrand Factor Von Willebrand Activity (Ristocetin Cofactor) Factor VIII Assay TAT 5 DAYS FVWF
A A B	A A B ⁹	C C C ^{4,12}
THROMBOTIC RISK PROFILE FBC Coagulation Profile Antithrombin III Factor V Leiden gene Factor II Prothrombin gene MTHFR gene Lupus Anticoagulant Protein C Free Protein S Ag Anticardiolipin Abs TAT 5 DAYS PROP	ANTENATAL PROFILE FBC + 5 part Diff Blood Group and Rh Type Atypical Antibody Screen Haemoglobin electrophoresis Syphilis IgG/IgM Glucose FT4/TSH Rubella Antibodies (IgG) Toxoplasma (IgG/IgM) Hepatitis B sAg Hep C Abs Varicella Zoster IgG (Immunity) HIV 1 & 2 Abs TAT 3 DAYS Please ensure the blood group (EDTA) tube label is HANDWRITTEN . Do not affix a secondary label. ANTE	NATURAL KILLER PROFILE 2 CD3 CD4 CD8 CD16/CD56 CD19 TAT 2 DAYS NKP2
A A B C C C ¹⁸	A A ³³ B B B G	A

Microbiology

TEST	CODE	SAMPLE REQ	TAT
16S rRNA Bacterial Gene	16S	J	1 week
18S rRNA Fungal Gene	18S	J	1 week
Aspergillus Precipitins	ASPP	B	5 days
Beta D Glucan	XBDG	B	3 days
Blood Culture [#]	BCUL	2 x BC ⁴	6 days +
Carbapenemase producing organism screen	MDR	STM (rectal)	4-5 days †
Clostridium Difficile Toxin by PCR	CLOS	RF*	2 days
Cryptococcal Antigen	CRYC	Serum or CSF	1 day
Cryptosporidium	CRPO	RF	2 days
CSF for Microscopy and Culture	CSF	CSF	1-3 days
Culture (Any site)	CULT		up to 5 days
Faecal Occult Blood/FOB (immunochemical/FIT)	QFIT	QFIT	1 day
Fluid Culture	FLUD	SC	2-7 days
Fluid for Crystals	FLU2	SC	1 day
Fungal ID + Sens	FUID	Fungal sample/STM	14 days
Galactomanan (Aspergillus Antigen)	SGAL	B	2 weeks
Gonorrhoea (Culture)	GONN	CS ⁺⁺	2-3 days
Group B Strep	GBSX	2 x STM	3-4 days
H. pylori Culture	HPCU	J	3 weeks
HVS	HVS	STM ⁺⁺⁺	2-4 days
IUCD for Culture	IUCD	Send Device	11-12 days
Legionella Urine Antigen	LEGA	RU	1 day
MRSA (Rapid PCR) one swab per site	MRSA	Blue Micro Swab	4 hours
MRSA Culture one swab per site	MRSW	Blue Micro Swab	2 days
Mycology/Skin Scrapings by PCR	DERM	Submit Sample	3-7 days
Nail Clippings	DERM	Nail clippings	3-7 days
Pleural Fluid for Culture	FLUP	SC	7 days
Pneumococcal Antigen	PNAG	RU	1 day
Pneumocystis Jiroveci (PCP) Examination	PCYS	BAL ⁺⁺	2-3 days
Rapid Strep (incl. m/c/s)	RAPS	STM ^{**}	1-3 days ^{**}

[#] Please contact the Phlebotomy at Patient Reception 020 7307 7383 for further details, as needed.

Blood cultures must be taken prior to any other blood samples.

The aerobic bottle must be collected first, followed by the anaerobic bottle.

Each bottle should be filled with 8-10 ml of blood, use the markings on the bottles to achieve this.

- Other bloods can be collected but must be collected **after** the blood cultures.
- Bottles must be labelled with the patient's identification details.
- Bottles and Request Form need to give the **time taken** and the **body site** that the blood was taken from. Ensure that the bottle barcodes are not obscured when adding patient labels.
- Send the blood cultures to the laboratory without delay.

Microbiology

TEST	CODE	SAMPLE REQ	TAT
Schistosoma (Urine)	USCH	Mid-morning terminal urine	8 hours
Sellotape Test	SELL	Send Sample***	1 day
Semen Culture	SPCU	Semen	2-4 days
Skin Scrapings/Mycology by PCR	DERM	Send Sample	3-7 days
Specific Gravity (Urine)	USG	RU	24 hours
Sputum for Routine Culture	SPU1	SC	2-4 days
Sputum for TB Culture (AFB)	SPU2	SC	up to 8 weeks
Stool for OCP and Culture	PENT	RF	2-3 days
Stool for OVA Cysts & Parasites by PCR	OCP	RF	1 day
Stool Reducing Substances	STRS	RF⁷	5 days
Swab (Ear)	EARS	STM	2-4 days (Culture) 8-9 days (Fungal) – same swab
Swab for Culture (Any Site) (see page 44)	SWAB	STM[†]	2-4 days
Synovial Fluid (for microscopy and culture)	FLU2	SC^{†††}	14 days
TB (pleural fluid)	TBCU	SC	up to 8 weeks
TB Culture	SPU2	SC	up to 8 weeks
TB Culture (Urine)	TBUR	3 x EMU	up to 8 weeks
TB Slopes – Confirmation and Sensitivity	TBSL	TB slope (LJ medium-green) ⁶	up to 8 weeks
Tissue for culture	TISS	Tissue sample	up to 14 days
Urine (Microscopy Only)	UMIC	RU	1 day
Urine for Microscopy and Culture	UCEM	MSU^{††††}	1-2 days

* Not performed on formed stool specimens.

** Do not use a black swab for RAPS. Use **Blue** only. Rapid antigen is reported within 4 hours with full culture to follow.

*** Use clear Sellotape only and attach to slide.

**** Culture techniques have been discontinued, please send PCR (see Sexual Health section for full details).

† Presumptive positive isolates will be sent to the PHE reference laboratory for confirmation.

†† BAL: Induced sputum or bronchoalveolar lavage.

††† The optimal sample type from the female genital tract is an endocervical swab. Gonorrhoea does not survive well outside the endocervical epithelium; a negative gonorrhoea culture result from a vaginal swab is not reliable for excluding infection.

†††† Culture for Mycoplasma, Ureaplasma and Trichomonas vaginalis has been discontinued due to the superiority of molecular methods. If investigations for Mycoplasma genitalium, Ureaplasma or Trichomonas vaginalis are required please request PCR testing (see Sexual Health section).

† Please state site of swab collection on **both** request form and swab label.

†† Please provide relevant travel history. If travel history is not provided, stool will be investigated for endemic pathogens only [Campylobacter, Salmonella, Shigella, Shigatoxin-producing E coli (VTEC), Cryptosporidium and Giardia].

††† If prosthetic joint is present please state in clinical details to ensure that enrichment culture is prolonged for 14 days.

†††† Optimal sample type for urine culture is a mid-stream clean catch urine sent in a sterile pot containing boric acid preservative.

Microbiology

URINE CULTURE PROCESSING AND RESULTS

All urine culture testing is performed using manual methods. The culture pathway adheres to national guidance and is a fully UKAS-accredited method.

Manual testing allows a larger amount of urine to be tested than previous automated method, which enables the laboratory to detect lower bacterial counts (as low as 103cfu/mL) and also facilitates the follow up of significant organisms grown from mixed cultures.

If the culture result is indicative of urinary tract infection, antibiotic susceptibilities will be tested from the culture growth and will be available 24 hours after the culture result. 'Direct sensitivities' are no longer performed. Direct susceptibility testing is not inoculum-controlled, produces inaccurate results and is not UKAS-accredited.

Culture results should be interpreted alongside the microscopy WBC count and clinical signs and symptoms. Significant growth on culture in the absence of pyuria may be suggestive of contamination with regional flora rather than true infection. It should be noted, however, that WBC degrade in urine quite rapidly and delays between sample collection and microscopy may lead to falsely low WBC readings which may account for these findings.

What does the result 'No significant growth' mean?

The amount of growth falls below the threshold for urinary tract infection (< 103cfu/mL).

There is no laboratory evidence of urinary tract infection.

Occasionally, this may be seen in very early stages of infection or in a partially treated urinary tract infection. Therefore, please send a repeat specimen if symptoms persist.

What does the result 'mixed growth doubtful significance' mean?

This means that the culture revealed a heavy growth of at least 3 organisms with no predominating organism; this represents contamination of the urine with the patient's flora during collection.

This result does not exclude urinary tract infection but it is not possible to determine the causative organism among the mixture of organisms.

If symptoms persist, please send a repeat urine specimen and ensure that patient understands optimal collection technique.

If you are receiving a lot of 'mixed growth of doubtful significance' results, please consider the following:

- **The instructions that patients are given to collect their urine sample**

Poor collection technique is the most common reason for a heavily mixed growth in a urine sample.

It is almost impossible to collect a urine sample without any contamination from the normal bacterial flora which inhabits the area surrounding the urethral opening, but optimal collection technique will minimise this contamination and allow the true infective cause to stand out and be identified (a patient instruction leaflet is available).

- **Delays between sample collection and laboratory processing**

The time between sample collection and laboratory processing can allow small amounts of contaminating bacterial flora to multiply up to higher amounts prior to laboratory testing, which can result in heavy mixed growth of bacteria on culture. Using a red topped specimen pot containing boric acid preservative will minimise this.

RED TOPPED BORIC ACID CONTAINERS

The preservative reduces the overgrowth of organisms and, to a lesser extent, reduces the degradation of white cells during transit leading to a more accurate laboratory result for both microscopy and culture. UKAS recommends the use of boric acid containers for all urine sample for microscopy and culture (Urine M,C&S) to improve the quality of microbiological results.

Red topped boric acid containers are for requests for urine microscopy and culture (MC&S) ONLY. Boric acid container should NOT be used for:

- Other urine microbiology tests (e.g. investigations for Chlamydia, Mycobacterium, Schistosomiasis, urinary antigen testing)
- Urine samples being analysed by PCR methodology
- Urine samples for non-microbiology tests (e.g. biochemistry, virology, pregnancy testing)
- Very small urine volumes (<20ml) e.g. neonates

Use of urinary dipsticks: boric acid may inhibit leukocyte esterase dipstick readings; dipstick testing performed on a sample in a boric acid container should be interpreted with caution.

If additional tests are required in addition to urine microscopy and culture, **an additional sample in a white-topped universal container should be sent**. In this case, it is advised that the mid-stream clean catch urine is collected in a sterile bowl and then transferred to the necessary specimen containers.

If, despite these measures, a patient has recurrent mixed growth reports from multiple urines, it may suggest that your patient has abnormal urinary tract architecture, immunosuppression or other non-infective cause that requires different laboratory investigations or referral to a specialist. If further information is required, please telephone the laboratory and ask to discuss the case with one of our consultant Microbiologists.

Microbiology

Swabs: Types and Codes

Patient Request Forms AND Swabs should be labelled with the body site from which the sample was taken. **This is important.** The swab site determines the appropriate culture media required to target the most likely pathogens.

SITE	CODE	SAMPLE TYPE
Culture Swabs		
Cervical Swab	CERS	Blue Micro Swab
Eye Swab	EYES	Blue or Orange Micro Swab
Ear Swab	EARS	Blue or Orange Micro Swab
Gonorrhoea	GONN	Black Charcoal Swab
High Vaginal Swab	HVS	Blue Micro Swab
Nasal Swab	NASS	Blue or Orange Micro Swab
Oral Swab	ORSW	Blue Micro Swab
Penile Swab	PENS	Orange Micro Swab
Rectal Swab	RECG	Blue Micro Swab
Skin Swab	SKIS	Blue Micro Swab
Throat Swab	THRS	Blue Micro Swab
Urethral Swab	URES	Orange Micro Swab
Vaginal Swab	VAGS	Blue Micro Swab
Vulval Swab	VULV	Blue Micro Swab
Wound Swab	WOUS	Blue Micro Swab
MRSA by Culture		
	MRSW	Blue Micro Swab x 1 – state site
	MRW2	Blue Micro Swab x 2 – state sites
	MRW3	Blue Micro Swab x 3 – state sites
	MRW4	Blue Micro Swab x 4 – state sites
	MRW5	Blue Micro Swab x 5 – state sites
RAPID MRSA by PCR		
	MRSA	Blue Micro Swab x 1 – state site
Note: This PCR methodology uses Blue Micro Swabs	MRS2	Blue Micro Swab x 2 – state sites
	MRS3	Blue Micro Swab x 3 – state sites
	MRS4	Blue Micro Swab x 4 – state sites
	MRS5	Blue Micro Swab x 5 – state sites

Blue Micro/Transwab are multipurpose, culture swabs in transport medium

Orange Micro/Transwab are small, thin wire culture swabs in transport medium

Black Charcoal Micro/Transwab Wound, skin, urogenital and throat.

Microbiology

PCR METHODS FOR THE DETECTION OF DERMATOPHYTE FUNGAL CULTURES

The detection of Dermatophyte fungal cultures uses High Sensitivity PCR testing. This reduces the overall turnaround time by up to three weeks, and increases the detection of fungal infection compared to combined microscopy and culture. Furthermore the specific targeting pathogens associated with superficial fungal infection is increased which assists in preventing the over reporting of insignificant fungi that are contaminants.

FUNGAL TEST CODES

	Investigation of Superficial Fungal Infection	Investigation of Non-Superficial Fungal Infection
Test Code	DERM*	FUN*
Sample type	Nail, Hair, Skin.	All specimens other than Skin, Hair and Nail.
Turnaround time	72 hours for interim PCR report, and 7 days for final culture (unless the fungal culture needs to be extended for significant growth).	7 days (non-sterile e.g. ear swab) and 3 weeks (sterile i.e. CSF).
Notes	<ul style="list-style-type: none"> • Dermatophyte PCR is replacing microscopy for Nails, Hair and Skin (72 hour TAT). • Non-dermatophyte culture will take 7 days rather than 3 weeks. • Microscopy will be used to confirm significance of rare fungi that may cause infections. • There is no change in the price of this test. 	<ul style="list-style-type: none"> • Non-sterile specimen fungal cultures are performed on Sabouraud's agar plates for 7 days with no microscopy. • Sterile specimen fungal cultures have microscopy (Calcafluor) reported on the day of processing and culture on a Sabouraud's agar slope, incubated for 21 days.

STOOL TEST CODES

Traditional culture methods have been replaced by Real Time PCR for enteric pathogen testing. The benefits are increased sensitivity and a higher detection rate. Once received and processed in the microbiology lab, negative results will be available within 24 hours. Positive results will be followed up with culture and sensitivities for final reporting.

STOOL OCP AND CULTURE		
Sample Type	Please request as PENT	Comments
Stool	Serosep EntericBio PCR Bacteria/Bacterial Toxins • Salmonella • Campylobacter • Shigella • VTEC Parasites • Cryptosporidium • Giardia	All stool samples will be tested for UK Pathogens. Overseas pathogens will only be tested if specifically requested and travel history and clinical details are provided. Samples that are positive for the bacterial pathogens will be cultured to provide sensitivities and, if indicated, for PHE referral. Samples will be kept for 7 days after receipt to allow for additional testing if required.

STOOL FOR OCP		
Sample Type	Please request as OCP	Comments
Stool	Requests for OCP only will include testing for cryptosporidium and giardia by PCR	Overseas pathogens will only be tested if requested and travel history and clinical details are provided.

C. DIFFICILE DETECTION		
Sample Type	Please request as CLOS	Comments
Stool	Serosep Enteric Bio PCR Alere Techlab EIA (Toxin)	Change to PCR and Elisa methods. Two tier PCR & Toxin <i>c. diff</i> screening based on PHE guidance. Improved sensitivity and specificity for both targets tested. Primary <i>c. diff</i> gene screening using Enteric Bio PCR. Secondary sequential testing using Alere EIA to confirm Toxin.

GASTRO VIRUS DETECTION (INCLUDING ROTAVIRUS) SEE VIROLOGY

ENTERIC ORGANISM RAPID DETECTION SEE VIROLOGY

GROUP B STREPTOCOCCUS (GBS)

Group B Streptococcus (GBS or group B Strep) is the most common cause of severe infection in newborn babies, and of meningitis in babies under age 3 months. On average in the UK:

- 2 babies a day develop group B Strep infection
- 1 baby a week dies from group B Strep infection
- 1 baby a week survives group B Strep infection with long term disability

Most GBS infection is of early onset, presenting in babies within the first 6 days of life, and usually within the first 12 hours after birth. Between age 7 days and 3 months, these infections are rare, and in babies over 3 months they are very rare indeed.

Most early-onset GBS infections (in babies aged 0-6 days) can be prevented by giving intravenous antibiotics in labour to women whose babies are at raised risk of developing GBS infection. In the UK, women are offered IV antibiotics in labour based on specific risk factors.

GBS is normal flora of the distal GI tract. Up to 30% of women carry it harmlessly in their vaginal tract. Vaginal carriage at the time of vaginal delivery can result in transmission of GBS to baby. Babies are more vulnerable to infection as their immature immune systems cannot fight off the multiplying bacteria. If untreated, GBS can cause serious infections, such as meningitis and septicaemia, which may lead to stillbirths, and newborn and infant deaths. If they survive, babies can develop permanent problems including hearing or vision loss, or cerebral palsy.

Current GBS prevention focuses on giving intravenous antibiotics to women in labour, aiming to reduce disease in infants at delivery. 2 x **Blue culture swabs** (lower vaginal and lower rectal) should ideally be taken from 35 weeks. Swabs will be placed in enrichment culture in the microbiology laboratory to ensure maximal detection.

Endocrinology

TEST	CODE	SAMPLE REQ	TAT
11 Deoxycorticosterone	DEOX	B	10 days
11 Deoxycortisol	11DC	B (Frozen)	10 days
17 Hydroxyprogesterone	17OH	B	5 days
ACTH (Adreno Corticotrophic Hormone)	ACTH	A (Plasma Frozen) ⁴¹	1 day
Aldosterone	ALDN	B	5 days
Aldosterone (Urine)	UALD	PU	5 days
Alpha Feto Protein	AFP	B	4 hours
Amenorrhoea Profile	AMEN	B	4 hours
Andropause Profile	ANDP	B B	8 hours
Androstenedione	ANDR	B (Frozen)	4 days
Antidiuretic Hormone	ADH	A A (Plasma Frozen) ⁴	10 days
Antimullerian Hormone (AMH Plus)	AMH	B	4 hours
Beta HCG (Quantitative)	QHCG	B	4 hours
BNP (NT-pro BNP)	BNP	B	4 hours
C Peptide	CPEP	B	3 days
Calcitonin	CATO	B (Frozen) ⁴	1 day
Catecholamines (Plasma)	CATE	A A (Plasma Frozen) ⁴	5 days
Catecholamines (Urine)	UCAT	PU ¹	5 days
Cortisol	CORT	B	4 hours
Cortisol (Urine)	UCOR	CU	5 days
DHEA	DHEX	B	7-10 days
DHEA – Urine (Dehydroepiandrosterone)	UDHE	CU	3 weeks
DHEA Sulphate	DHEA	B	4 hours
Dihydrotestosterone	DHT	B B	7 days
Down Syndrome Risk Bloods only (Risk to be calculated by clinician)	HCGF/PAPA	B	4 hours
Down Syndrome Risk Profile (2nd trimester) Quad	DRP	B DRP form ^{7,8}	2 days
Down Syndrome Risk Profile with risk calculation first trimester	DRP	B DRP form + image of scan ^{7,8}	2 days
Erectile Dysfunction Profile	IMPO	A B B G	3 days
Female Hormone Profile	FIP	B	4 hours
First Trimester Antenatal Screen (Risk to be calculated by requesting clinician)	HCGF/PAPA	B	4 hours
Free Cortisol (Urine)	UCOR	CU	5 days
Free T3	FT3	B	4 hours
Free T4	FT4	B	4 hours
FSH	FSH	B	4 hours
Growth Hormone (Fasting)	GH	B ^{7,35}	4 hours
Gut Hormone Profile	GUTP	A A (Frozen within 15 minutes) ⁴¹	3 weeks
Hirsutism Profile	HIRP	B	4 hours
HRT Profile 1	HRT	B	4 hours
HRT Profile 2	HRT2	B G	4 hours
















Endocrinology

TEST	CODE	SAMPLE REQ	TAT
IGF-1 (Somatomedin)	SOMA	B (Frozen) ⁴	1 day
IGF-BP3	IGF3	B (Frozen) ⁴	5 days
Impotence Profile	IMPO	A B B G	3 days
Inhibin A	INIA	B	1 month
Inhibin B	INIB	B (Day 3 of cycle, frozen)	5 days
Insulin	INSU	B	4 hours
Insulin Resistance (Fasting)	FIRI	B G	4 hours
Luteinising Hormone (LH)	LH	B	4 hours
Macroprolactin	PRLD	B	4 days
Male Hormone Profile	MIPR	B	4 hours
Melanin	MELA	RU ¹³	5 days
Melatonin (Serum)	MEL	B (Frozen)	5 days
Melatonin (Urine)	UMEL	CU ¹³	2 weeks
Menopause Profile	MENO	B	4 hours
Metabolic Syndrome Profile	METS	A B B G	9 days
Metanephrines (Plasma)	PMET	A (Frozen plasma)	7 days
Metanephrines (Urine)	UMEX	PU ¹	5 days
Oestradiol (E2)	OEST	B	4 hours
Oestriol (Estriol)	E3	B B	4 days
Oestrone	E1	B B	4 days
Osteocalcin	OST	B (Frozen) ⁴	4 days
Parathyroid Hormone (Whole)	PTHl	B ⁴	1 day
Pituitary Function Profile	PITF	B B	1 day
Polycystic Ovary Syndrome Profile	PCOP	A B B B G G ⁷	5 days
Polycystic Ovary Syndrome SHORT	PCOS	B G	4 hours
Pregnancy (Serum) [Quantitative]	QHCG	B	4 hours
Pregnanetriol (Urine)	UPTR	CU (Frozen)	5 days
Pregnenolone	PREN	B	15 days
Progesterone	PROG	B	4 hours
Proinsulin	PROI	A (Frozen plasma) ⁴	5 days
Prolactin	PROL	B	4 hours
Prolactin (Macro)	PRLD	B	4 days
Renin	RENI	A (Frozen plasma) ³⁶	5 days
Reverse T3	RT3	B ^{7,37}	10 days
Serotonin	SERT	H (Frozen whole blood) ¹	10 days
Serotonin (Urine)	USER	PU 50mls (Frozen) ¹	5 days
Sex Hormone Binding Globulin	SHBG	B	4 hours
Somatomedin (IGF-1)	SOMA	B (Frozen) ⁴	1 day
T3	T3	B	4 hours
T3 (Reverse)	RT3	B ^{7,37}	10 days
Testosterone	TEST	B	4 hours
Testosterone (Bioavailable)	BTES	B	5 days
Testosterone (Free)	FTES	B	3 days

Endocrinology

TEST	CODE	SAMPLE REQ	TAT
Thyroglobulin Abs	TGAB	B	1 day
Thyroglobulin Assay	TGA	B	1 day
Thyroid Abs (incl. Thyroglobulin + Thyroid Peroxidase Abs)	THAB	B	1 day
Thyroid Peroxidase Antibodies/Anti TPO	TPEX	B	1 day
Thyroid Profile 1	TF	B	4 hours
Thyroid Profile 2	TF2	B	2 days
Thyroid Profile 3	TF3	B	4 hours
Thyroxine (T4)	T4	B	4 hours
Thyroxine Binding Globulin	TBG	B (Frozen)	10 days
TSH	TSH	B	4 hours
TSH-Receptor Antibodies	TSI	B	4 days

REPRODUCTIVE IMMUNOLOGY AT
ROSALIND FRANKLIN LABORATORIES, CHICAGO, USA

TEST	CODE	SAMPLE REQS	TAT
Reproductive Immunophenotype Panel	3RF		1 week
NK Assay/Cytotoxicity Panel	4RF		1 week
NK Assay Follow-Up Panel	5RF		1 week
TH1/TH2 Cytokine Ratio	6RF	 ⁵	1 week
Leucocyte Antibody Detection Panel MALE	7RF	 ^{3,4,6}	1 week
Leucocyte Antibody Detection Panel FEMALE	8RF		1 week
HLA DR Antigens	9RF		2 weeks
HLA DQ Alpha Antigens	10RF		2 weeks
HLA DQ Beta Antigens	11RF		2 weeks
NK Assay Panel + Intralipids	16RF		1 week
KIR (Killer-like Immunoglobulin-like Receptors) Genotyping	17RF		2-3 weeks
TH1/TH2 Intracellular Cytokine Ratios with IVIG, Prednisolone	20RF	 ⁵	1 week
TH1/TH2 Intracellular Cytokine Ratios with IVIG	21RF	 ⁵	1 week
TH1/TH2 Intracellular Cytokine Ratios with Prednisolone	22RF	 ⁵	1 week
Endometrial Biopsy Immune Profiling	23RF	J (Contact Referrals)	2 weeks
T Regulatory Cells	25RF		3 days

Patients who have samples taken at TDL’s Patient Reception at 76 Wimpole Street may attend any time during hours of opening on Mondays or Tuesdays, and by **NOON on Wednesdays to allow for same day shipping to Chicago by Fed Ex**. Samples for Rosalind Franklin are not accepted on Thursdays, Fridays or Saturdays. Fed Ex charges are included in these charges.

REPRODUCTIVE IMMUNOLOGY AT ST HELIER, CARSHALTON

TEST	CODE	SAMPLE REQ	TAT
NK (CD69) Cell Assay	CD69	H*	Send Mon-Thurs only
NK Cytotoxicity Assay	HSNK	H H H*	Send Mon-Thurs only
NK (CD69) and NK Cytotoxicity	69C	H H H*	Send Mon-Thurs only
NK Cytotoxicity with suppression, steroid, IVIg & Intralipin	NKCY	H H H*	Send Mon-Thurs only
NK Cytotoxicity with suppression with steroid, IVIg and intralipin, and NK (CD69) cell assay	69CI	H H H*	Send Mon-Thurs only
TH1/TH2 Cytokine Profile	1TH2	H H H*	Send Mon-Thurs only
Suppression with steroid, IVIg and intralipin, NK (CD69) cell assay, TH1/TH2 cytokines	NCIT	H H H*	Send Mon-Thurs only

* Patients need to attend Patient Reception at 76 Wimpole Street by **11.00am latest Mondays – Thursdays**. Samples cannot be accepted on Fridays, Saturdays or Sundays. Allow 2 days for results.

THYROID PROFILE 1

FT4
TSH

TAT
4
HOURS

TF

B

THYROID PROFILE 2

T4 Free T3
TSH Free T4
Thyroglobulin Abs
Thyroid Peroxidase

TAT
2
DAYS

TF2

B

THYROID PROFILE 3

FT3
FT4
TSH

TAT
4
HOURS

TF3

B

FEMALE HORMONE PROFILE

LH
FSH
Prolactin
Oestradiol (17-Beta)

TAT
4
HOURS

FIP

B

MALE HORMONE PROFILE

FSH
LH
Testosterone
Free Androgen Index
Prolactin
SHBG

TAT
4
HOURS

MIPR

B

ANDROPAUSE PROFILE

DHEAs
FSH
Testosterone
Free Androgen Index
LH
SHBG

TAT
8
HOURS

ANDP

B B

ERECTILE DYSFUNCTION/
IMPOTENCE PROFILE

Lipid Profile
Glucose
HbA1C
TSH
Prolactin
Total Testosterone
Free Testosterone
PSA

TAT
3
DAYS

IMPO

A B B G

ANTIMULLERIAN HORMONE/AMH PLUS

Age related reference intervals in women	Age Range	Elecsys AMH (pmol/L)
The reference intervals below are derived from a population of apparently healthy women not taking any contraceptive medication. The reference intervals represent the 10th – 90th percentile values for the women in each age bracket.	20 – 29 years	13.1 – 53.8
	30 – 34 years	6.8 – 47.8
	35 – 39 years	5.5 – 37.4
	40 – 44 years	0.7 – 21.2
	45 – 50 years	0.3 – 14.7

TAT
4
HOURS

AMH

B Samples can be taken, at any time during a patient's monthly cycle. Ambient, unspun sample stability has been validated for up to 5 days. Postal samples are therefore acceptable, and samples can also be collected and posted using TDL TINIES.

More Hormone Profiles
are shown on page 50

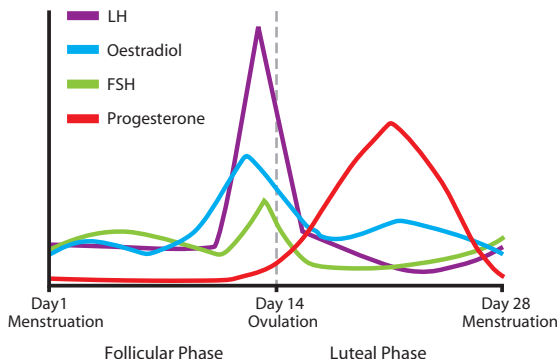
Reproductive health

The tests in this section are drawn from all disciplines of diagnostic pathology and are listed in other appropriate sections in the Laboratory Guide.

PUBERTY

The beginning of the reproductive cycle of life – diagnosis tests may include:

- Oestradiol
- FSH
- LH
- Progesterone
- Androstenedione
- DHEA sulphate
- Testosterone
- SHBG
- Prolactin



THE MENSTRUAL CYCLE/PREGNANCY

This cycle controls female fertility and is influenced by hormone levels which impact bone health and many other aspects of female physiology. Pregnancy lasts 40 weeks and is divided into trimesters.

First Trimester (week 0–13): confirmation of pregnancy and associated tests may include:

- Pregnancy test (urine)
- Quantitated Beta HCG (serum)
- Ectopic Pregnancy assessment (Beta HCG and Progesterone)
- Recurrent Miscarriage Profile
- Antenatal Screen
- Nuchal Scan with Free Beta HCG and PAPP-A or Non-Invasive Prenatal Test (Harmony) for risk assessment of Downs Risk (a DRP request form must be enclosed with samples, see back of guide, and an image of the scan attached to the request form).
Contact TDL Genetics for details of Non-Invasive Prenatal Testing (NIPT)
- Chorionic Villus Sampling (CVS) for chromosomal analysis (PCR for Rapid Trisomy and karyotyping for the rarer abnormalities)
- Toxoplasma/Varicella Zoster/Parvovirus/CMV

Reproductive health

Second Trimester (week 14–26):

testing is primarily directed at evaluating the actual and potential development of the baby and may include:

- Downs Risk Profile (Triple Test +)
- Amniocentesis for chromosomal analysis (AmnioPCR for Rapid Trisomy and karyotyping for the rarer abnormalities)
- Glucose and Protein (urine or serum)

Third Trimester (week 27–40):

testing for foetal wellbeing and the health of the mother may include:

- Glucose and Protein (urine or serum)
- Toxoplasma
- Atypical antibody screening
- Group B Strep (From 35 weeks – rectal and low vaginal swabs)
- Chlamydia

INFERTILITY

Infertility and its management is increasingly implicated in growing numbers of clinical disciplines. More recently, greater emphasis is being given to male infertility. Recent data suggests that approximately 40% of all infertility is ascribed entirely, or in part, to male factors, 40% to female factors with an additional 20% unexplained. Testing at the outset of infertility treatment can reduce some of the emotional and financial costs, as well as allowing couples to pursue other possible options.

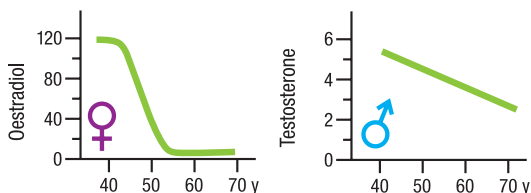
- Hormones
- Lifestyle/Environmental
- Ovarian Reserve
- Unexplained Infertility/Implantation failure
- Male Factors
- Infection
- Chromosomes/Genetics
- Polycystic Ovary Syndrome
- Recurrent/Spontaneous miscarriage

AGEING

Reaching menopause and andropause is a gradual process with modulating hormones as ovarian function declines in women, and the more gradual, less defined and highly variable effect in men. Testing may include:

- Hormones (Menopause/Andropause Profile)
- Testosterone/Free testosterone/Bioavailable Testosterone
- SHBG
- DHEAs
- Thyroid function
- Osteoporosis/Bone Markers

General patterns of age-related decline in estradiol levels in women (left) and total testosterone levels in men (right)



Reproductive health

INFERTILITY

HORMONES	
FEMALE	MALE
FSH – day 2/3	Testosterone/Prolactin/FSH/LH
LH	Sex Hormone Binding Globulin
Oestradiol	Inhibin B (male)
Antimüllerian Hormone (AMH)	Male Hormone Profile
Progesterone – day 21	Andropause Profile
Female Hormone Profile	Insulin Resistance
Prolactin	Erectile Dysfunction
	Impotence Profile

INFECTION	
FEMALE	MALE
High Vaginal swab	Investigations for prostatitis/urethritis
Cervical swab	Mycoplasma Genitalium
Bacterial Vaginosis screen	Ureaplasma
Toxoplasma	Chlamydia/Gonorrhoea
Chlamydia/Gonorrhoea	Chlamydia in Semen
CMV	Hep B sAg/Hep B Core Abs/Hep C/HIV 1&2
Syphilis	Herpes Simplex I/II by PCR
Hep B sAg/Hep B Core Abs/Hep C/HIV 1&2	Semen culture
Herpes Simplex I/II by PCR	Syphilis
STI Profiles	STI Profiles
Infection screening by PCR	Infection screening by PCR

LIFESTYLE/ENVIRONMENT	
FEMALE	MALE
Well Person Profile DL6	Fit for Fertility Male Profile
Zinc, Lead	Well Person Profile DL6
Trace Metal Profile (blood)	Trace Metal Profile (blood)
Antioxidant Activity	Antioxidant Activity
Thyroid Profiles	Thyroid Profiles
Vitamin Profiles	Vitamin Profiles
Vitamin D (25 OH)	Vitamin D (25 OH)
Folate	Folate
Selenium	Selenium
Omega 3/Omega 6	Zinc
	Omega 3/Omega 6
	Oxidative Stress (ROS) in Semen

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Reproductive health

CHROMOSOMES/GENETICS	
FEMALE	MALE
Chromosome/Karyotype (parental) Fragile X (female) Cystic Fibrosis Screen Tay Sachs Jewish Carrier Profile Inherited disorders (specific)	Chromosome/Karyotype (parental) Male Hormone Profile Y-Chromosome microdeletion Fragile X Male Cystic Fibrosis Screen Tay Sachs Jewish Carrier Profile Inherited disorders (specific)
OVARIAN TUMOUR	
FEMALE	
Antimullerian Hormone (AMH)	CA 125/HE4
POLYCYSTIC OVARY SYNDROME	
FEMALE	
Polycystic Ovary Profile	
UNEXPLAINED INFERTILITY/IMPLANTATION FAILURE /RECURRENT MISCARRIAGE	
FEMALE	MALE
Recurrent Miscarriage Profile Reproductive Immunophenotyping (CD 3/4/8, CD 5/19, CD 16/56/69) NK Cell Profile Antiphospholipid Antibodies Lupus anticoagulant and Anticardiolipin Antibodies Thrombotic Profile Antinuclear antibodies Anti-Thyroglobulin Antibodies Chromosome/Karyotype (parental) Infection screening (See Infection)	Chromosome/Karyotype (parental) Y-Chromosome microdeletion Sperm DNA Fragmentation Sperm aneuploidy Infection screening (See Infection) Heavy Metals (Blood) Male Recurrent Miscarriage Profile Oxidative Stress in Semen (Reactive Oxygen Species)
SPERM HEALTH	
MALE	
See TDL Andrology on page 60.	

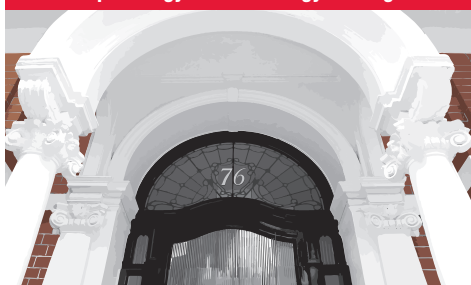
TDL Andrology

The single most important factor determining a man's fertility potential is the production of healthy sperm. A semen analysis has classically been used as the marker of this potential, by providing information about the sperm count, motility and morphology. However, there are other parameters given in a semen analysis that are often neglected or overlooked, which may indicate important pathologies – such as infection, prostatic disease, immunological infertility, retrograde ejaculation, malformation or obstruction of the genital tract, tumour, and congenital or endocrine disorders.

Early diagnosis of the male factor is important in order to detect any underlying pathology, determine the extent of infertility and ensure appropriate treatment. It may also avoid unnecessary investigations for the female partner, particularly if her age is a limiting factor.

For men who have had a vasectomy, clearance should only be given when there is no evidence of presence of sperm in two consecutive semen samples. It is therefore vital to ensure that results are reported according to best practice guidelines. Special clearance may be given at the doctor's discretion when there are persistent non-motile sperm present.

Andrology booking can now be done online at
www.tdlpathology.com/andrologybooking



Guidelines for Producing Samples

Ideally semen samples should be produced on-site at TDL's Patient Reception at 76 Wimpole Street. Ideally patients must abstain from ejaculation for 2-3 days prior to the test, but no less than 2 days and no longer than 5 days before the test. This requirement is important for semen analyses and post vasectomy analyses to ensure reliability of results. It is possible that samples that do not comply with guidelines for abstinence and collection may not be able to be processed. All semen samples must be produced directly into the sterile containers provided by The Doctors Laboratory.

All containers are weighed and batch tested for sperm cytotoxicity. In exceptional circumstances when semen samples are produced off-site, they can only be accepted by the Andrology Department in sample containers provided by TDL.

WHO 2010 guidelines state that two semen analyses should be performed before any diagnosis is confirmed. This may require requests for two (separate) semen analyses.

Appointments

It is important to make an appointment for all semen samples (on or off site) whether for a comprehensive semen analysis or post vasectomy analysis. It may be necessary to give patients who attend without an appointment a specific time to re-attend. The first appointments for post vasectomy samples should usually be 12 weeks and 20 ejaculations after surgery.

Appointments can be made by calling **020 7025 7940**. There is an attendance fee of £45.00 in addition to pathology charges.

Please complete a Pathology Request Form for your patient. If you would like to request other pathology, you can use the same form or complete a second additional form. Results will usually be reported to you within 48 hours.

If you would like to discuss these tests, or any aspect of this service, please contact TDL Andrology on 020 7025 7940 or email andrology@tdlpathology.com for further information.

SEMEN			
TEST	CODE	SAMPLE REQS	TAT
Oxidative Stress in Semen (ROS + MIOXSYS)	SR0S	Semen ¹	1 day
Retrograde Ejaculation	RTRO	Contact Lab	2 days
Semen Analysis, Comprehensive*	SPER	Semen ¹	2 days*
Semen Analysis, Post-Vasectomy**	PVAS	Semen ¹	2 days
Semen Analysis, Vasectomy Reversal*	SPER	Semen ¹	2 days*
Semen Culture	SPCU	Semen	2-4 days
Semen Fructose	SPCF	Semen	2 days
Semen Leucocytes	PMNS	Semen	2 days
Semen Parameters	SPOD	Semen ¹	1 day
Semen Zinc	SPCZ	Semen	up to 10 days
Sperm Aneuploidy	SPPL	Semen ¹	4 weeks
Sperm Antibodies (Serum)	ASAB	B	5 days
Sperm Antibodies/MAR Test (Semen)[†]	ASPA	Semen	1 day
Sperm Comet[®]	CMET	Semen	1-2 weeks
Sperm Count (Post-Vasectomy)	PVAS	Semen ¹	2 days
Sperm DNA Fragmentation (SCSA)	SEXT	Semen ¹	1-2 weeks
Sperm Morphology (Kruger strict criteria)	MRPH	Semen ¹	2 days
Semen parameters may be requested INDIVIDUALLY (eg count only, vitality only, etc). Please request as SPOD and indicate on the request form which parameter is required.			
Semen Parameters	SPOD	Semen ¹	1 day

* If required, comprehensive semen analysis can be reported within 4 hours, with morphology to follow.

** For men who have had a vasectomy, clearance should only be given when there is no evidence of presence of sperm in a single ejaculate when recommendations are met. It is rare that a 'diagnosis' is made without confirmation, therefore patients/clinicians should be able to freely request a second confirmatory sample. Special clearance may be given at the doctor's discretion, when there are <100 000/ml non-motile sperm present after the assessment of two specimens in full accordance with recommendations. Recommendations, as given by the Association of Biomedical Andrologists, the British Andrology Society and the British Association of Urological Surgeons 2016, are as follows:

- 1 Analysis of post vasectomy semen samples should not occur until 12 weeks post-surgery and after a minimum of 20 ejaculates
- 2 Semen samples must be analysed within 4 hours of production, and in cases where sperm is found a repeat analysis must be performed within 1 hour of production
- 3 Semen should be provided in weighed specimen containers provided by TDL Andrology
- 4 Sexual abstinence should be between 2 and 7 days

[†] Sperm antibodies in semen are measured as part of the routine semen analysis.

Sperm swim test

Sperm preparation for overnight survival

Sperm motility and vitality testing for epididymal toxicity

Sperm retrieval procedures (biopsy, PESA, MESA)

Sperm cryopreservation and storage (undertaken by Andrology Solutions – HFEA licensed)

All men who store sperm must be screened for HIV 1&2, Hepatitis B, Hepatitis C and HTLV. Under HFEA regulations, sperm can be stored for an initial period of 10 years with formal consent. All patients are offered counselling prior to sperm cryopreservation.

These arrangements, and details for other specialist semen tests, are available on request. Please contact TDL Andrology on 020 7025 7940 or email sheryl.homa@tdlpathology.com for further information.

Sperm DNA fragmentation

High sperm DNA fragmentation is associated with reduced natural pregnancy rates and assisted conception pregnancy rates as well as live birth rates. In addition, DNA fragmentation leads to higher miscarriage rates as published in the ESHRE Recurrent Pregnancy Loss 2017 Guideline. High levels of DNA fragmentation may be reduced by considering varicocele repair, treatment of underlying infections or inflammation, changes in lifestyle or with antioxidant supplements.

When requesting Sperm DNA Fragmentation there are two options. Please specify whether the request is for sperm DNA fragmentation by **SCSA** or **COMET**.

• Sperm Chromatin Structure Assay (SCSA®) [SEXT]

This test has the ability to measure large numbers of cells (between 5,000 and 10,000 sperm), rapidly in an ejaculate. The SCSA® test monitors the changes in fluorescence of a probe, acridine orange, to detect both single and double DNA strand breaks using flow cytometry. It has been developed using human and animal models over the last 35 years and is one of the most statistically robust tests available for sperm DNA fragmentation. It is a standardised, validated CLIA approved test with high reproducibility and low variability. The test requires a minimum sperm count of approximately 1 million/ml.

• Sperm COMET® Assay [CMET]

When sperm counts are limited, DNA fragmentation can be effectively assessed using the Comet® assay as only ~5,000 sperm are required. The Comet® assay uses electrophoresis to determine abnormal sperm, and can measure both single and double strand breaks. Unlike the SCSA® test, the comet assay may be subject to inter-observer variability and may be less statistically robust as it measures low counts of 50 to 100 sperm cells from each sample.

Sperm Aneuploidy

Chromosomal abnormalities may be somatic cell in origin, in which case they can be detected by a simple blood karyotype analysis. However, most sperm chromosome anomalies arise as a result of errors during meiosis, which cannot be detected by a blood karyotype analysis. These anomalies can only be detected by looking at the sperm chromosomes directly. Studies have shown that sperm with a high rate of aneuploidy have a negative impact on pregnancy rate and are associated with recurrent pregnancy loss.

TDL Andrology

This test uses fluorescent in situ hybridisation (FISH) to label individual chromosomes with specific probes. Hundreds of sperm are assessed from one ejaculate. There are limitations to the test as only 5 probes are currently used routinely for analysis (three of the 22 autosomes: chromosomes 13, 18 and 21, and the sex chromosomes, X and Y), although others are available upon specific request. The results are reported showing incidence of disomy or nullisomy for each of the autosomes and for both sex chromosomes. A sex chromosome ratio is also reported. It is CE marked.

Instructions for collection of Sperm DNA and Aneuploidy specimens

Sperm DNA Fragmentation or Sperm Aneuploidy testing are not part of the Comprehensive Semen Analysis and need to be requested as a separate test, test code SEXT and SPPL, respectively. Semen samples ideally need to be frozen as soon as possible after liquefaction, but not longer than 60 minutes post ejaculation. Samples must be snap-frozen for Sperm DNA Fragmentation and cryopreserved in TYB for Sperm Aneuploidy. If samples are prepared by another laboratory. Two cryovials containing not less than 0.25 mls of semen is required. Frozen samples can be sent to, or collected by TDL, by arrangement, and must be accompanied with relevant patient details, the sperm count and GDPR consent form. A count of a minimum 1 million/ml is required for accurate DNA and aneuploidy reporting.

Oxidative Stress in Semen (ROS + MIOXSYS) and Male infertility

There is now growing evidence to support a link between oxidative stress and male infertility. It is the underlying cause of sperm DNA damage and impairs semen parameters and fertilisation, adversely affects embryo development and is associated with reduced pregnancy rates. It may also increase the risk of miscarriage. High levels of ROS may be reduced by considering varicocele repair, treatment of underlying infections or inflammation, changes in lifestyle or with antioxidant supplements.

TDL provides a comprehensive assessment of oxidative stress by **combined measurement of Reactive Oxygen Species and Redox Potential**. Please request as oxidative stress test (code ROS).

The test includes combined testing for:

- **Chemiluminescence Assay for Reactive Oxygen Species**

Reactive Oxidative stress may be measured by a simple chemiluminescence test in semen, which measures the level of reactive oxygen species.

- **MIOXSYS Electrochemical Assay for Redox Potential**

Oxidative stress may be determined by an electrochemical assay which measures the redox potential in semen. This test measures the overall difference between total oxidants and antioxidants in the system.

References

Homa ST, Vessey W, Perez-Miranda A, Riyait T, Agarwal A (2015). Reactive oxygen species (ROS) in human semen: determination of a reference range. *J Assist Reprod Genet* 32(5):757-64.

Vessey W, Perez-Miranda A, Macfarquhar R, Agarwal A, Homa S. (2014). Reactive oxygen species (ROS) in human semen: validation and qualification of a chemiluminescence assay. *Fertil Steril*. 102:1576-1583.

If you would like to discuss these tests, or any aspect of this service, please contact TDL Andrology on 020 7025 7940 or 020 7307 7373, or email andrology@tdlpathology.com.

Effects of ROS-induced Oxidative Stress on Sperm

- Lipid peroxidation which damages the sperm surface causing an abnormal morphology and impaired motility.
- Damage to proteins on cell surface responsible for cell signalling and may affect enzyme function inside the cell.
- Increased semen viscosity.
- Peroxidation of DNA and subsequent unravelling or fragmentation.
- Possible mutagenic effects.
- Damage to seminiferous epithelium, damage to tubules, testicular atrophy, reduced spermatogenesis.
- Decrease in sperm vitality, motility.
- Impaired fertilization by affecting sperm capacitation and the acrosome reaction.

Causes of Elevated ROS Levels

- Genito-urinary tract infection
- Prostatitis
- Vasectomy reversal
- Varicocele
- Cryptorchidism
- Chronic disease
- Xenobiotics
- Chemical pollutants and occupational hazards
- Heavy metal exposure
- Removal of seminal plasma during sperm preparation for assisted conception
- Drugs – cyclophosphamide, aspirin, paracetamol
- Smoking
- Excessive exercise
- Heat exposure
- Obesity
- Age

Semen samples need specialist handling – for this reason all requests for semen analyses should be made by appointment. Practices or patients should contact TDL Andrology on 020 7025 7940 to make appointments and to confirm instructions for sample collection.

Sexual Health

TEST	CODE	SAMPLE REQ	TAT
7 STI Profile by PCR (7 tests from 1 Sample)	PP12	FCRU/PCR/TPV/Semen	2 days
Chlamydia (PCR swab)	SPCR	PCR	2 days
Chlamydia (Thin Prep)	TPCR	TPV	2 days
Chlamydia (Urine)	CPCR	FCRU	2 days
Chlamydia/Gonorrhoea (PCR Swab)	SCG	PCR	2 days
Chlamydia/Gonorrhoea (Rectal)	RSCG	PCR	2 days
Chlamydia/Gonorrhoea (Thin Prep)	TCG	TPV	5 days
Chlamydia/Gonorrhoea (Throat)	TSCG	PCR	2 days
Chlamydia/Gonorrhoea (Urine)	CCG	FCRU	2 days
Chlamydia/Gonorrhoea/Trichomonas by PCR	CCGT	FCRU/PCR/TPV	2 days
CT/GC/Trichomonas/Mgen (PCR Swab)	SGTM	PCR Swab	2 days
CT/GC/Trichomonas/Mgen (Urine)	CGTM	FCRU	2 days
Early Detection Screen PCR/NAAT	STDx	A 10mls or 2 x 4mls (Vacutainer only)	3 days
Early Detection Screen PCR/NAAT with Syphilis	STXX	B A 10mls or 2 x 4mls	3 days
FASTest Sexual Health Screening Tests			See page 69
Gardnerella vaginalis by PCR	GVPC	FCRU/PCR/TPV	2 days
Gonorrhoea (Culture)	GONN	CS	2-3 days
Gonorrhoea (PCR swab)	SGON	PCR	2 days
Gonorrhoea (Thin Prep)	TGON	TPV	2 days
Gonorrhoea (Urine)	CGON	FCRU	2 days
Haemophilus ducreyi by PCR	DUCR	PCR	7 days
Hepatitis A Profile	HEPA	B	4 hours
Hepatitis B Surface Antigen	AUAG	B	4 hours
Hepatitis C Antibodies	HEPC	B	4 hours
Herpes Simplex I/II by PCR (Swab)	HERS	PCR	5 days
Herpes Simplex I/II by PCR (Urine)	HERD	FCRU/PCR/TPV	5 days
HIV 1 & 2/p24Ag	HDUO	B	4 hours
HIV/HSV/HCV (Early detection by PCR/NAAT) with Syphilis	STXX	B A 10mls or 2 x 4mls	3 days
HIV/HSV/HCV Screen by PCR/NAAT (10 days post exposure)	STDx	A 10mls or 2 x 4mls (Vacutainer only)	3 days
HIV Rapid RNA HIV-1 QUALITATIVE	LHIV	A (Vacutainer only)	4 hours
HIV Rapid RNA HIV-1 QUANTITATIVE	RHIV	A (Vacutainer only)	4 hours
HPV (DNA and reflexed mRNA)	HPVT	TPV	3 days
HPV (HR mRNA types 16, 18 + others)	HPVH	TPV	2-3 days
HPV (Individual low & high risk DNA subtypes)	HP20	TPV/PCR	2-3 days
Lymphogranuloma Venerium (LGV)	LGVP	PCR* ⁴²	1-2 weeks
Macrolide Resistance Test (Mgen)	MGR	FCRU/PCR	1-2 weeks
Mycoplasma genitalium by PCR	MGEN	FCRU/PCR/TPV	2 days
Mycoplasma genitalium/Ureaplasma by PCR	MUPC	FCRU/PCR/TPV	2 days
Rapid Xpert HIV-1 RNA Qualitative – Early Detection from 10 days	LHIV	A (Vacutainer only)	4 hours

* LGV can be added to a positive chlamydia sample using the same swab if requested within 4 days of receipt of result.

Sexual Health

TEST	CODE	SAMPLE REQ	TAT
Rapid Xpert HIV-1 RNS Viral Load – Rapid Testing for HIV-Positive Patient Prognosis and Response To Antiretroviral Therapy	RHIV	A (Vacutainer only)	4 hours
RPR (VDRL)	RPR	B	2 days
STD1 M/F STD Quad	STD1	B FCRU	2 days
STD2 M/F STI Profile Plus (Urine and Serology)	STD2	B FCRU (If culture swabs are needed please request separately)	4 days
STD3 Female STD Quad (PCR Swab and Serology)	STD3	B PCR	2 days
STD4 Female STI Profile Plus (PCR Swab and Serology)	STD4	B PCR (If culture swabs are needed please request separately)	4 days
STD5 Serology only	STD5	B	4 hours
STD6 Serology only without HIV	STD6	B	4 hours
STD8 Vaginitis/BV Profile using Culture & PCR Swab	STD8	PCR/STM	3 days
STD9 Symptomatic lesion sample using PCR Swab from lesion & PCR Swab	STD9	2 x PCR Swab	7 days
STI Profile: MSM1	MSM1	B /FCRU/PCR Swab Throat/PCR Swab Rectal	2 days
STI Profile: MSM2	MSM2	B /FCRU/PCR Swab Throat/PCR Swab Rectal	3 days
Swab for Culture (Any Site)	SWAB	STM [†]	2-4 days
Syphilis by PCR (chancres)	SYPS	PCR	5 days
Syphilis IgG/IgM	SERJ	B	4 hours
TPPA	TPPA	B	2 days
Trichomonas vaginalis by PCR	TVPC	FCRU/PCR/TPV	2 days
Ureaplasma urealyticum by PCR	UGEN	FCRU/PCR/TPV	2 days
Vaginitis/BV Profile using Culture & PCR Swab	STD8	PCR/STM	3 days

RAPID XPRT HIV-1

For some patients earlier diagnosis of HIV infection is important. **Xpert HIV-1 Qual** is a qualitative test that provides on-demand molecular testing for early diagnosis (from 10 days).

FOR PATIENT ON TREATMENT FOR HIV

Xpert HIV-1 Viral Load accommodates on demand testing and measurement of blood plasma HIV-1 RNA concentration (HIV viral load/40 copies/ml) which has been established as the standard of care in assessing HIV-positive patient prognosis and response to antiretroviral therapy. Assessment of viral load levels is a strong predictor of the rate of disease progression and, by itself or in combination with CD4 T-cell counts, has great prognostic value.

- Improve Patient Care: Same day results support better clinical decisions
- Increase Efficiency: Rapid results enable earlier adjustments to appropriate therapy
- Strengthen Communities: Quick decisions can help reduce drug resistance

Sexual Health

Chlamydia

Chlamydia is the most common curable STI diagnosed in the UK. Often asymptomatic, anyone who is sexually active is considered to be at increased risk of chlamydia infection. It is the most commonly recognised, screened and treated of all STI's. **Allow 6 weeks before re-testing to avoid picking up the DNA from a previous infection.**

Gonorrhoea

Gonorrhoea is caused by the bacterium *Neisseria gonorrhea*, which multiplies easily in the mucous membranes of the male and female reproductive tract. It can cause serious and permanent health conditions if not treated. Symptoms of gonorrhoea are usually overt in men with white, yellow, or green discharge from the penis. Gonorrhoea can also infect the throat and rectum – individual PCR swabs from **each site** should be taken to screen for gonorrhoea. Resistance to antibiotics is increasing and treatment is now combined oral and injectable antibiotics. **Partners should be treated at the same time with retesting after two weeks to confirm clearance – test of cure is recommended following treatment for gonococcal infections.**

Mycoplasma Genitalium (M.Gen)

M.gen is an important sexually transmitted pathogen detectable only by NAAT. M.gen lacks a cell wall and has limited treatment options. It spontaneously develops resistance to antimicrobials. BASHH recommends treatment with Resistance Guided Therapy – testing for M.gen with macrolide resistance determination. M.gen cannot be cultured for diagnostic testing. M.gen prevalence is higher than GC, and in some populations can be similar to CT. M.gen risk factors are similar to CT and consider testing M.gen in all males with non-GC urethritis and all individuals with signs or symptoms of PID, cervicitis, endometritis, associated infertility, ano-rectal condition or epididymo-orchitis. Partner testing is advised for current partners only. Rectal infections are common, and appear to be an important reservoir for resistance. BASHH guidance – all patients must return for test of cure at 3-5 weeks.

Macrolide Resistance Testing (M.gen)

Prevalence of M.gen in men and women in the general population is 1-2%. *Mycoplasma genitalium* has been implicated as a cause of acute and chronic non-chlamydial non-gonococcal urethritis in males and post coital bleeding, cervicitis, endometritis and pelvic inflammatory disease in females. It is a sexually transmitted, fastidious microorganism that is extremely difficult to culture – with nucleic acid amplification testing (NAAT urine or swab) being the only method available for routine *M. genitalium* detection. Macrolides are generally considered the first-line treatment for *M. genitalium* infections. However, **resistance to macrolides** seems to be increasing worldwide typically exceeding > 40% in male patients who are detected positive for M.gen at screening.

M.gen can be requested as a single PCR test or with CT/GC, with or without other testing options. Important updates to the UK BASHH *M. genitalium* management guidelines are taking the issue of antimicrobial resistance seriously. The draft guidelines have been posted for consultation and include a grade 1B recommendation to test for antimicrobial resistance, stating the importance of knowing the macrolide resistance status to determine whether azithromycin should be prescribed. The guidelines aim to support laboratories in making a case for increased funding to bring in the necessary testing to manage *M. genitalium* infections and associated antimicrobial resistance.

Ureaplasma

U. Urealyticum and *parvum* are strains of bacteria that can lead to urinary tract infection and pelvic inflammation. Usually asymptomatic, it is part of the normal genital flora of both men and women. It is found in about 70% of sexually active humans. In males with lower sperm quality, ureaplasma infection could lead to a more pronounced decreased in some seminal parameters and compromise sperm motility.

Sexual Health

Trichomoniasis

Trichomoniasis is caused by a tiny parasite called *Trichomonas vaginalis* – and is one of the most common STI's worldwide. Frequency of coinfection with other STI's is well recognised, and notably, infection increases the risk of HIV transmission in both men and women. It is associated with adverse pregnancy outcomes, infertility, and cervical neoplasia. Some women may mistake this infection for a yeast infection or bacterial vaginosis since the symptoms are similar: frothy discharge, strong vaginal odour, pain on intercourse, irritation and itching. Men can get trichomoniasis too, but they don't tend to have symptoms. It seems to be linked to male factor infertility. Partners (male or female) need to be treated to avoid ongoing re-infection. Infected women who are sexually active have a high rate of reinfection, **thus re-screening at 3 month post treatment could be considered.**

Gardnerella vaginalis

'*Gardnerella vaginalis*' is a bacterium rather than a sexually transmitted infection. It is part of the normal vaginal flora but, when the normal balance of bacteria in the vagina is disrupted, it can flourish and overgrow leading to bacterial vaginosis. Does it matter if it not an STI? Yes, because it can be characterised by a fishy smelling, white vaginal discharge, itching, burning, and irritation, and there are some known pregnancy and pelvic inflammatory conditions associated with Gardnerella as well as a higher risk of getting other STI's.

In a patient with signs and symptoms suggestive of bacterial vaginosis detection of Gardnerella vaginalis provides supportive evidence of bacterial vaginosis. It can, however, be detected in asymptomatic individuals and it can also be absent in patients with bacterial vaginosis which has been caused by overgrowth of other similar organisms such as Mobiluncus and Atopobium species. Results should be interpreted in line with patient's clinical symptoms and microscopy.

Herpes/Herpes Simplex Virus I/II

Genital herpes caused by the herpes simplex virus (HSV). The virus lives in the nerves and when active it travels to the surface of the infected area and makes copies of itself – called shedding, because new virus cells can at this time rub off onto another person. The virus travels back down the nerve to a ganglion usually at the base of the spine where it lies dormant for a while. It causes painful blisters on the genitalia and surrounding areas. It can be passed through intimate sexual contact and for this reason is referred to as an STI. Once infected, it remains a chronic long term condition with the virus remaining with recurrent activity with variable frequency. There are two types of herpes simplex virus: Type 1 and Type 2. Both are highly contagious and can be passed easily from one person to another. There is no cure for genital herpes, the symptoms can usually be controlled by antiviral medication. Although using a condom can reduce the risk of herpes transmission, condoms are not 100% effective since herpes can be spread from skin-to-skin.

Lymphogranuloma venereum (LGV)

LGV is a type of chlamydia bacteria that attacks the lymph nodes. It is seen predominantly in gay and bisexual men, and very rarely seen in the UK in heterosexual men and women.

Nearly all LGV infections seen in the UK in recent years have been in the rectum. Within a few weeks of becoming infected, most people get painful inflammation in the rectum with bleeding, pus, constipation or ulcers, sometimes with fever, rash and groin, armpit or neck swelling. Left untreated, LGV can cause lasting damage to the rectum that may require surgery. LGV in the penis might cause a discharge and pain when urinating, with swollen glands in the groin. LGV in the mouth or throat is rare but can cause swollen glands in the neck.

Investigation for possible LGV symptoms is by PCR swab taken from the rectum and penis. If LGV infection is suspected in female patients, cervical and vaginal PCR swabs should be taken. Samples are first tested for chlamydia and if chlamydia is detected, if LGV is suspected, swabs can be further tested, if requested, for LGV as an additional tests, using the same swab samples. Sexual contact partners should also be checked.

FASTest Test Now

Sexual Health Screening – *ahead of expected time*

FAST SSC

Fast Screen *SHORT*

HIV 1&2/p24 Ag
Syphilis IgM/IgG
FAST Urine CT/GC



TAT
4
HOURS

FSSC

B FCRU

FAST USC

Fast Screen with *URINE*

HIV 1&2/p24 Ag
Hep B sAg
Hep C Abs
Syphilis IgG/IgM
FAST Urine CT/GC



TAT
4
HOURS

FUSC

B FCRU

FAST SSS

Fast Screen *SHORT* with *SWAB*

HIV 1&2/p24 Ag
Syphilis IgM/IgG
FAST Swab CT/GC



TAT
4
HOURS

FSSS

B PCR

FAST SSC

Fast Screen with *SWAB*

HIV 1&2/p24 Ag
Hep B sAg
Hep C Abs
Syphilis IgG/IgM
FAST Swab CT/GC



TAT
4
HOURS

FSWS

B PCR



FAST SINGLE TESTS

Sample type

FCT	FAST Chlamydia Urine	FCRU
FGN	FAST Gonorrhoea Urine	FCRU
FCG	FAST CT/GC Urine	FCRU
FSCT	FAST Chlamydia PCR Swab	PCR Swab
FSGN	FAST Gonorrhoea PCR Swab	PCR Swab
FSCG	FAST CT/GC PCR Swab	PCR Swab
FTCG	FAST CT/GC Throat PCR Swab	PCR Swab
FRCG	FAST CT/GC Rectal PCR Swab	PCR Swab

Sexual Health

STI's can be caused by virus, fungus, parasite or bacteria. Anyone who is sexually active may be at risk of acquiring an STI. The risk is higher for those with increased numbers of sexual partners, or who have had sex with someone who has/had many partners, or have had unprotected sex.

STI		INCUBATION PERIOD	SAMPLE SITE
Chlamydia CT	Bacterial	1–3 weeks, up to 6 weeks	Urine Cervix/Vagina Cervix/Vagina
Gonorrhoea GC	Bacterial	2–7 days, up to 1 month	Urine Cervix/Vagina Cervix/Vagina Cervix/Vagina
CT/GC Combined	Bacterial	1–3 weeks, up to 6 weeks	Urine Cervix/Vagina Cervix/Vagina Rectum Throat
Mycoplasma genitalium	Bacterial	Symptoms develop at 1–3 weeks	Urine GU Site Cervix/Vagina
Ureaplasma urealyticum	Bacterial	Symptoms develop at 1–3 weeks	Urine GU Site Cervix/Vagina
Trichomonas vaginalis	Parasitic	4–28 days, many patients are asymptomatic carriers	Urine GU Site Cervix/Vagina
Gardnerella vaginalis	Bacterial	Imbalance of normal flora	Urine GU Site Cervix/Vagina
Bacterial Vaginosis (BV)	Bacterial	Imbalance of normal flora	Cervix/Vagina
Herpes Simplex Viral I/II	Viral	2–14 days, testing is most appropriate for patients with symptomatic lesion(s)	Herpes lesion
Human Papillomavirus	Viral	HPV is the most common sexually transmitted infection – usually asymptomatic	Cervical cells Cells/papilloma from site (throat/penile/anal)
Genital warts	Viral	Weeks/ months after exposure	GU Warts
Syphilis/Herpes	Bacterial/ Viral	Whenever active lesions are present	Symptomatic lesion

Sexual Health

TEST	TEST CODE	SAMPLE TYPE	TAT
Chlamydia	CPCR	First catch Urine	2 days
Chlamydia	SPCR	PCR Swab	2 days
Chlamydia	TPCR	Thin Prep Vial	2 days
Gonorrhoea by PCR	CGON	First Catch Urine	2 days
Gonorrhoea by PCR	SGON	PCR Swab	2 days
Gonorrhoea by PCR	TGON	Thin Prep Vial	2 days
Gonorrhoea by CULTURE	GONN	Black Charcoal swab	2-3 days
CT/GC	CCG	First Catch Urine	2 days
CT/GC	SCG	PCR Swab	2 days
CT/GC	TCG	Thin Prep Vial	5 days
CT/GC	RSCG	PCR Swab	2 days
CT/GC	TSCG	PCR Swab	2 days
Mycoplasma genitalium by PCR	MGEN	First Catch Urine	2 days
Mycoplasma genitalium by PCR	MGEN	PCR Swab	2 days
Mycoplasma genitalium by PCR	MGEN	Thin Prep Vial	2 days
Ureaplasma by PCR	UGEN	First Catch Urine	2 days
Ureaplasma by PCR	UGEN	PCR Swab	2 days
Ureaplasma by PCR	UGEN	Thin Prep Vial	2 days
Trichomonas vaginalis by PCR	TVPC	First Catch Urine	2 days
Trichomonas vaginalis by PCR	TVPC	PCR Swab	2 days
Trichomonas vaginalis by PCR	TVPC	Thin Prep Vial	2 days
Gardnerella vaginalis by PCR	GVPC	First Catch Urine	2 days
Gardnerella vaginalis by PCR	GVPC	PCR Swab	2 days
Gardnerella vaginalis by PCR	GVPC	Thin Prep Vial	2 days
Bacterial Vaginosis (BV) Profile by both MICROSCOPY and PCR	STD8	Both Microscopy & PCR swab	3 days
Herpes by PCR	HERS	PCR Swab	5 days
Herpes by PCR	HERD	First Catch Urine	5 days
HPV DNA/mRNA	HPVT	Thin Prep Vial	3 days
HPV Typed DNA	HP20	PCR Swab	2-3 days
HPV Typed DNA	HP20	Cells / Papilloma	2-3 days
HPV Typed DNA	HPVT	Thin Prep Vial	3 days
HPV Typed DNA	HP20	PCR Swab	2-3 days
HPV Typed DNA	HP20	Cells / Papilloma	2-3 days
Syphilis/Herpes Lesion Profile	STD9	PCR Swab	7 days

Sexual Health

BLOOD		INCUBATION PERIOD	SAMPLE SITE
Syphilis	Bacterial	9–21 days, but up to 90 days	Blood
Herpes Simplex Virus I/II	Viral	IgG 4–6 weeks after exposure IgM 5–35 days after exposure, after which test IgG	Blood Blood
HIV	Viral	Usually 10–90 days, but up to 180 days	Blood Blood
Hep B	Viral	Usually 45–180 days, average of 60–90 days	Blood Blood
Hep C Ab	Viral	Usually 9–180 days, average of 45–65 days	Blood Blood

EARLY DETECTION PROFILES BY PCR	INCUBATION PERIOD	SAMPLE SITE
7 STIs by PCR	One sample for 7 STI Tests	Urine Cervix Vagina
HIV/HBV/HCV	Early Detection Screen by PCR Multiplex (HIV from 10 days)	Blood

Sexual Health

TEST	TEST CODE	SAMPLE TYPE	TAT
Syphilis IgG/ IgM	SERJ	B	4 hours
Herpes IgG (past infection)	HERP	B	2 days
Herpes IgM (current/recent)	HERM	B	2 days
HIV I&II/ p24 antigen (screening from 45 days post exposure (BHIVA))	HDUO	B	4 hours
Hep B surface antigen	AUAG	B	4 hours
Hep C Antibodies	HEPC	B	4 hours

TEST	TEST CODE	SAMPLE TYPE	TAT
Chlamydia	PP12	Thin Prep Vial	2 days
Gonorrhoea		or	
Mycoplasma genitalium	PP12	First Catch Urine	2 days
Macrolide Resistance Test (M.gen)*		or	
Ureaplasma genitalium	PP12	PCR Swab	2 days
Trichomonas vaginalis			
Gardnerella vaginalis			
Herpes Simplex I/II			
<i>*included if POSITIVE M.gen is detected from the same sample</i>			
HIV 1&2 RNA	STDx	A 10mls or 2x4mls	3 days
Hepatitis B (HBV DNA)		(Vacutainer only)	
Hepatitis C (HCV RNA)			

Sexual Health

STD1

M/F STD QUAD
(Urine and Serology)

Serology
HIV 1&2/p24 Antigen
Syphilis IgG/IgM

Urine
Chlamydia
Gonorrhoea

TAT
2
DAYS

STD1

B FCRU

STD3

FEMALE STD QUAD
(PCR swab and Serology)

Serology
HIV 1&2/p24 Antigen
Syphilis IgG/IgM

Vaginal PCR Swab
Chlamydia
Gonorrhoea

TAT
2
DAYS

STD3

B PCR

STD5

SEROLOGY ONLY

HIV 1&2/p24 Antigen
Hepatitis B Surface Antigen
Hep C Abs
Syphilis IgG/IgM

TAT
4
HOURS

STD5

B

STD2

M/F STI PROFILE PLUS
(Urine and Serology)

Serology
HIV 1&2/p24 Antigen
Hep B Surface Antigen
Hep C Abs
Syphilis IgG/IgM

Urine
Chlamydia/Gonorrhoea
Mycoplasma genitalium
Ureaplasma
Trichomonas vaginalis
Gardnerella vaginalis
Herpes Simplex I/II

TAT
4
DAYS

STD2

B FCRU If culture swabs are needed please request separately

STD4

FEMALE STI PROFILE PLUS
(PCR swab and Serology)

Serology
HIV 1&2/p24 Antigen
Hep B Surface Antigen
Hep C Abs
Syphilis IgG/IgM

Vaginal PCR Swab
Chlamydia/Gonorrhoea
Mycoplasma genitalium
Ureaplasma
Trichomonas vaginalis
Gardnerella vaginalis
Herpes Simplex I/II

TAT
4
DAYS

STD4

B PCR If culture swabs are needed please request separately

STD6

SEROLOGY ONLY WITHOUT HIV

Hepatitis B Surface Antigen
Hep C Abs
Syphilis IgG/IgM

TAT
4
HOURS

STD6

B

Sexual Health

STD8 VAGINITIS / BV PROFILE USING CULTURE & PCR SWAB

Candida species
Gardnerella vaginalis by PCR
Trichomonas vaginalis by PCR

TAT
3
DAYS

STD8

PCR STM

STD9 SYMPTOMATIC LESION SAMPLE USING PCR SWAB FROM LESION

Syphilis by PCR
Herpes Simplex I/II by PCR
(from single swab)

TAT
7
DAYS

STD9

PCR PCR

HIV / HBV / HCV SCREEN (HIV1/HIV2/HBV/HCV by PCR/NAAT)

HIV1 and HIV2 (RNA)
Hepatitis B Virus (HBV DNA)
Hepatitis C Virus (HCV RNA)

Samples must be received in the laboratory within 2 days of sample taking

TAT
3
DAYS

STDx

A 10mls or 2x4mls (Vacutainer only)

EARLY DETECTION SCREEN WITH SYPHILIS (HIV1/HIV2/HBV/HCV by PCR/NAAT)

HIV1 and HIV2 (RNA)
Hepatitis B Virus (HBV DNA)
Hepatitis C Virus (HCV RNA)
Syphilis IgG/IgM

Samples must be received in the laboratory within 2 days of sample taking

TAT
3
DAYS

STXX

B **A** 10mls or 2x4mls

CT/GC/TRICHOMONAS/MGEN

Chlamydia
Gonorrhoea
Trichomonas vaginalis
Mycoplasma genitalium
Macrolide Resistance Test (Mgen)*

All tests can be requested individually
**included if POSITIVE M.gen is detected from the same sample.*

TAT
2
DAYS

CGTM (Urine) / SGTm (Swab)

FCRU OR PCR Swab

7 STI PROFILE BY PCR (7 TESTS FROM 1 SAMPLE) (Urine, Swab, Thin Prep or Semen)

Chlamydia trachomatis
N. Gonorrhoea
Mycoplasma genitalium
Macrolide Resistance Test (M.gen)*
Ureaplasma
Trichomonas vaginalis
Gardnerella vaginalis
Herpes Simplex I/II

All tests can be requested individually
**included if POSITIVE M.gen is detected from the same sample.*

TAT
2
DAYS

PP12

FCRU OR PCR Swab OR TPV OR Semen

Sexual Health

STI Profile: MSM1			
HIV 1&2/p24 Ag Syphilis IgG/IgM Urine for CT/GC Throat Swab CT/GC Rectal Swab CT/GC			
			TAT 2 DAYS
MSM1			
B	FCRU	PCR Swab Throat	PCR Swab Rectal

STI Profile: MSM2			
HIV 1&2/p24 Ag Syphilis IgG/IgM 7 STI by PCR Screen Throat Swab CT/GC Rectal Swab CT/GC Macrolide Resistance Test (M.gen)*		Hep B sAg Hep C Abs	
			TAT 3 DAYS
MSM2			
B	FCRU	PCR Swab Throat	PCR Swab Rectal

RAPID XPRT HIV-1

For some patients earlier diagnosis of HIV infection is important. **Xpert HIV-1 Qual** is a qualitative test that provides on-demand molecular testing for early diagnosis (from 10 days).

FOR PATIENT ON TREATMENT FOR HIV

Xpert HIV-1 Viral Load accommodates on demand testing and measurement of blood plasma HIV-1 RNA concentration (HIV viral load/40 copies/ml) which has been established as the standard of care in assessing HIV-positive patient prognosis and response to antiretroviral therapy. Assessment of viral load levels is a strong predictor of the rate of disease progression and, by itself or in combination with CD4 T-cell counts, has great prognostic value.

- Improve Patient Care: Same day results support better clinical decisions
- Increase Efficiency: Rapid results enable earlier adjustments to appropriate therapy
- Strengthen Communities: Quick decisions can help reduce drug resistance

RAPID XPRT HIV-1 RNA QUALITATIVE EARLY DETECTION FROM 10 DAYS	RAPID XPRT HIV-1 RNA VIRAL LOAD RAPID TESTING FOR HIV-POSITIVE PATIENT PROGNOSIS AND RESPONSE TO ANTIRETROVIRAL THERAPY
HIV-1 RNA	HIV-1 RNA VIRAL LOAD (40 copies/ml)
Sample must be received in the laboratory within 24 hours of sample taking	Sample must be received in the laboratory within 24 hours of sample taking
TAT 4 HOURS	TAT 4 HOURS
LHIV	RHIV
A (Vacutainer only)	A (Vacutainer only)

Immunology

TEST	CODE	SAMPLE REQS	TAT
Acute Viral Hepatitis Screen	AHSC	B	4 hours
Adrenal Cortex Antibodies	ACTX	B	2 days
ANCA (Anti-Neutrophil Cytoplasmic Abs)	ANCA	B	2 days
Anti-Actin Antibodies	AAA	B	5 days
Anti-Basal Ganglia Antibodies	ABGA	B	3 weeks
Anti-CCP Antibodies (RF)	CCP	B	2 days
Anti-Liver Cytosol Antibodies	ALCA	B	5 days
Anti-MOG [Myelin Oligodendrocyte Glycoprotein] Antibodies	AMOG	B	3 weeks
Anti-MUSK Antibodies	MUSK	B	2 weeks
Antinuclear Antibodies (titre & pattern)	ANAB	B	2 days
Anti-Phosphatidylserine Antibodies	PHTS	B	5 days
Anti-Phospholipase A2 Receptor	AA2R	B	3 weeks
Anti-Ri Antibodies	RIAB	B	3 days
Anti-SLA (Soluble Liver Antigen) Abs	LSA	B	10 days
Antistaphylolysin Titre (SGOT)	ASTT	B	3 days
Antistreptolysin Titre/ASOT	ASLT	B	2 days
Antisulfatide Antibodies	ASA	B	5 weeks
Aquaporin 4 Antibodies (Neuromyelitis Optica)	AQUA	B	2 weeks
Ascariasis Serology	ASC	B	5 days
Autoantibody Profile I	AUTO	B	2 days
Autoantibody Profile II	ENDO	B	2 days
Avian Precipitins (11 Species)	AVIA	B	5 days
Babesia Antibodies	BABE	B	3 weeks
Beta 2 Glycoprotein 1 Abs	B2GP	B	5 days
Borrelia Antibodies (Lyme Disease) IgG, IgM – see page 88	BORR	B ^{9,14}	2 days
Borrelia Antibodies (Lyme Disease) IgM – see page 88	BORM	B	2 days
Borrelia Confirmation (Immunoblot) – see page 88	BORC	B ^{9,14}	10 days
Brucella Serology	BRUC	B ⁹	2-3 weeks
C1 Esterase Inhibitor	C1EI	B	5 days
C3 Complement	C3	B	4 hours
C3/C4 Complement	COMP	B	4 hours
C4 Complement	C4	B	4 hours
Calprotectin	CALP	RF	5 days
Calprotectin/Elastase Profile	CEP	RF	5 days
Campylobacter Jejuni Antibodies	CJAB	B	5 days
Candida Antibodies	CANA	B	5 days
Candida Antigen	CCAG	B	5 days
Cardiolipin Antibodies (IgG+IgM)	ACAB	B	2 days
Cartilage Antibodies	ACA	B	5 days
CCP Antibodies (RF)	CCP	B	2 days
Centromere Autoantibodies	CENT	B	2 days
CH50 (Classical pathway)	CH50	B (Frozen) ⁴	4 days
Chagas Disease Serology (S.American Trypanosomiasis) T. Cruzi	CHGA	B ^{9,14}	10 days























































Immunology

TEST	CODE	SAMPLE REQ	TAT
Chlamydia Species Specific (MIF) Ab Screen	CHAB	B	2 days
Chronic Fatigue Syndrome Profile	VIP1	A + B ¹⁰	5 days
Coeliac Disease – HLA DQ2/DQ8 Genotype	Q2Q8	A ⁹	10 days
Coeliac/Gluten Profile 2	GSA2	A B	10 days
Coeliac/Gluten Sensitivity Profile	GSA	B	2 days
Colloid Antigen-2 Antibodies	CA2A	B	2 weeks
Cotinine (Serum)	COT	B	4 days
Cotinine (Urine)	COTT	RU	2 days
Cryoglobulins	CRYO	J ⁶	10 days
Diamine Oxidase Activity	DIAM	B	2 weeks
Diphtheria Antibodies	DIPH	B	5 days
DNA (Double Stranded) Antibodies IgG	DNAA	B	2 days
DNA (Single Stranded) Antibodies	DNAS	B	5 days
Echinococcus (Hydatid) Antibodies	EFAT	B ^{9,14}	5 days
Ehrlichiosis Antibodies	EHRL	B ^{9,14}	10 days
Elastase (Faecal)	ELAS	RF	5 days
Elastase/Calprotectin Profile	CEP	RF	5 days
Endomysial Antibodies (IgA)	AEAB	B	2 days
Extractable Nuclear Antibodies (nRNP, Sm, Ro, La, Jo1, Scl70) CENP-B	ENA	B	2 days
Faecal Elastase	ELAS	RF	5 days
Farmers Lung Precipitins	FARM	B	5 days
Fasciola Hepatica Antibodies (Liver Fluke)	FASC	B	2 weeks
Ganglionic Acetylcholine Receptor Antibodies	GACA	B	1 month
Ganglioside GM1, GD1B, GQ1B Abs	GANG	B	5 days
Gastric Parietal Autoantibodies	GASP	B	2 days
Giardia Serology	GIAR	B	5 days
Gliadin Antibodies (IgG) (deamidated)	AGAB	B	2 days
Glomerular Basement Membrane Abs	AGBM	B	2 days
Glutamic Acid Decarboxylase Antibodies (GAD 65)	GAD	B	5 days
Gluten Allergy Profile	GLUT	A B B	10 days
Gluten Sensitivity Evaluation	GSA	B	2 days
Gluten/Coeliac Profile 2	GSA2	A B	10 days
Granulocyte Immunology	GRIM	A A	2 weeks
H. pylori Antibodies (IgG)	HBPA	B	2 days
H. pylori Antigen (Breath)	HBQT	J	5 days
H. pylori Antigen (Stool)	HBAG	RF	3 days
Haemophilus B Influenzae Antibodies	HINF	B	7 days
Histamine (Blood)	HITT	A (Frozen plasma)	5 days
Histamine (Urine)	HITU	RU	5 days
Histamine Releasing Urticaria Test	CURT	B	10-14 days
Histone Antibodies	HISA	B	5 days
Histoplasmosis	HISP	B	10 days

















Immunology

TEST	CODE	SAMPLE REQ	TAT
HLA B27	HLAB	A ⁹	3 days
Human Anti-Mouse Antibodies	HAMA	B (Frozen)	6 weeks
IgE (Total)	IGE	B	1 day
Immune-Complexes	IMCP	B	5 days
Immunoglobulins (IgG, IgM, IgA)	IMM	B	4 hours
Inner Ear Antigen (Ottoblot)	IEA	B	3 weeks
Insulin Antibodies	INAB	B	5 days
Interferon – Alpha	IFA	B (Frozen) ⁹	3 weeks
Interferon – Gamma	IFG	A (Frozen)	3 weeks
Interleukin 1 Beta	ILB	B (Frozen) ^{4,7}	1-2 weeks
Interleukin 2	IL2	B (Frozen) ^{4,7}	1-2 weeks
Interleukin 4	IL4A	B (Frozen) ^{4,7}	1-2 weeks
Interleukin 6	IL6	B (Frozen) ^{4,7}	1-2 weeks
Interleukin 8	IL8	B (Frozen) ^{4,7}	1-2 weeks
Interleukin 10	IL10	B (Frozen) ^{4,7}	1-2 weeks
Interleukin 28b Genotype	IL28	A	2 weeks
Intrinsic Factor Antibodies	IFAB	B	2 days
Islet Cell Antibodies	ICAB	B	2 days
Legionella Antibodies	LEGO	B	2 days
Legionella Urine Antigen	LEGA	RU	1 day
Leptospirosis (Weil's Disease) Abs (IgM)	LEP	B	5 days
Leukotriene E4	LTE4	CU (Frozen)	3 weeks
Listeria Antibody	LIST	B	1 week
Liver Immunoblot	LIVI	B	5 days
Liver Kidney Microsomal Antibodies	LKM	B	2 days
Lupus Anticoagulant and Anticardiolipin Abs	LUPA	B C ^{4,18}	2 days
Lyme Disease (Borrelia Abs) IgG, IgM	BORR	B ^{9,14}	2 days
Lyme Disease (Borrelia Abs) IgM	BORM	B	2 days
Meningococcal Abs	MENI	B	2-4 weeks
Mitochondrial Antibodies	AMIT	B	2 days
Mitochondrial Antibodies M2	MAM2	B	2 days
Myasthenia Gravis Evaluation	MGE	B	5 days
Myelin Associated Glycoprotein Antibodies	MAG	B	5 days
Myelin Basic Protein Antibodies	MBPA	B	2 weeks
Myeloperoxidase Antibodies	MPO	B	2 days
Myocardial Antibodies	MYO	B	1 week
Myositis Panel	MYOS	B	2 days
Neuronal Antibody (Hu, Ri, Yo, Cv2, Ma2)	NEUR	B	10 days
NMDA Receptor Antibodies	NMDA	B	3 weeks
Nucleic Acid Antigen Antibodies	DNA	B	2 days
Oligoclonal Bands	CSFO	CSF + B	5 days
Ovarian Autoantibodies	OVAB	B	2 days
Paragonimus Serology	PRGM	B	2 weeks
Parathyroid Antibodies	PTHA	B	1 week

Immunology





TEST	CODE	SAMPLE REQ	TAT
Pemphigus/Pemphigoid Autoantibodies	SKAB		2 days
Phospholipid Antibodies	PLIP		5 days
Pituitary Antibodies	PITU	 ⁴	1 month
Pneumococcal Antibodies – Serotype Specific	PASS		5 weeks
Pneumococcal Antibody Screen	PNEU		7 days
Proteinase 3 Ab	PR3		2 days
Purkinje Cell Antibody (Hu and Yo)	PURK		10 days
Q Fever (C Burnetti) Antibodies	QFEV	 ⁹	10 days
Rheumatoid Factor (Latex Test)	RF		1 day
Rheumatology Profile 1 (Screen)	RH	 	2 days
Rheumatology Profile 2 (Connective tissue)	RH2	   	3 days
Rheumatology Profile 3 (Rheumatoid/Basic)	RH3	 	2 days
Rheumatology Profile 4 (Systemic Lupus)	RH4	  	2 days
Rheumatology Profile 5 (Mono Arthritis)	RH5	   	3 days
Rheumatology Profile 6 (Rheumatoid Plus)	RH6		2 days
Rheumatology Profile 7 (Sjogren's Syndrome)	RH7		10 days
Rickettsial Species Antibody Profile	RICK		7 days
RPR (VDRL)	RPR		2 days
Saccharomyces Cerevisiae Antibodies	ASCA		2 weeks
Salivary Duct Antibodies	SAB		12 days
Scleroderma Immunoblot	SCLI		5 days
Sjogren's Syndrome	RH7		2 days
Skin (Pemphigus/Pemphigoid) Autoantibodies	SKAB		2 days
Skin Antibodies by Immunofluorescence	STSK		1 month
Sleeping Sickness Serology (African Trypanosomiasis)	TRYP	 ⁹	10 days
Smooth Muscle Antibodies	ASMO		2 days
Sperm Antibodies (Serum)	ASAB		5 days
Steroid Cell Antibody	SCA		2 days
Striated/Skeletal Muscle Antibody	STRA		2 days
Strongyloides Antibodies	STGA		10 days
Syphilis IgG/IgM	SERJ		4 hours
TB Quantiferon®-TB Gold*	TBQ4	Special tubes or  ¹	3 days
Testicular Autoantibodies	TAB		2 days
Tetanus Antibody	TETA		5 days
Thyroid Abs (incl. Thyroglobulin + Thyroid Peroxidase Abs)	THAB		1 day
Thyroid Peroxidase Antibodies/Anti TPO	TPEX		1 day
Tissue Transglutaminase IgA (Coeliac)**	TAA		2 days
Tissue Transglutaminase IgG	TAAG		5 days
Torch Screen	TORC		2 days
Total Immune Function Evaluation	TIE	 +  ^{5,10}	7 days
Total Immunoglobulin E	IGE		1 day
Toxocara Antibodies (IgG)	TFAT	 ⁹	5 days
Toxoplasma Antibodies (IgG+IgM)	TFAM	 ⁹	4 hours







Immunology

TEST	CODE	SAMPLE REQ	TAT
Toxoplasma Antibody Full Evaluation (IgM, Dye Test, IgG Avidity)	TDYE	 ⁹	10 days
Toxoplasma by PCR	TXAG		5 days
TPPA	TPPA		2 days
Trichinella Serology	TRIC		5 days
Trypanosome (Chagas) Antibodies	CHGA	 ^{9,14}	10 days
Tularaemia Antibodies	TULA	 ¹⁴	5 days
TSH-Receptor Antibodies	TSI		4 days
Urinary Methyl Histamine	UHIT	RU (Frozen)	2 weeks
Urticaria Test (Histamine Releasing)	CURT		10-14 days
Vascular Endothelial Growth Factor	VEGF		14 days
VDRL (RPR)	RPR		2 days
Voltage Gated Calcium Channel Antibodies	CCAB		3 weeks
Voltage Gated Potassium Channel Antibodies	VPCA		3 weeks
Whooping Cough (Pertussis) Antibodies	PERS		5 days
Whooping Cough (Pertussis) by PCR	PERP	Prenasal (posterior nasopharynx) swab	5 days
Yellow Fever Antibodies	YELL	 ^{9,14}	10 days
Yersinia Antibodies	YERS		4 days
Zika Abs IgM and IgG	ZKAB		5 days

* Please indicate clearly if samples have/have not been incubated prior to sending to the laboratory. If Lith Hep (green top) tube is used, please request as TBQ4 and ensure sample is received in the laboratory within 16 hours of sample taking.

** If Tissue Transglutaminase (TAA) is regulated and is LOW (<0.1U/ml) total IgA will be reflexed. If total IgA is low (<0.1g/L) deamidated gliadin IgG will be reflexed. If Tissue Transglutaminase (TAA) is HIGH (>10 U/ml), endomysial IgA will be reflexed as confirmatory test.

HLA DQ2/DQ8			
TEST	CODE	SAMPLE REQs	TAT
Coeliac Disease Profile 2	GSA2	 	10 days
Coeliac Disease – HLA DQ2/DQ8 Genotype	Q2Q8	 ⁹	10 days
Coeliac/Gluten Sensitivity Profile	GSA		2 days

GLUTEN SENSITIVITY EVALUATION (COELIAC DISEASE ANTIBODY)	COELIAC DISEASE PROFILE 2	GLUTEN ALLERGY PROFILE
Endomysial IgA Gliadin deamidated IgG Total IgA Tissue Transglutaminase (IgA) <div>TAT 2 DAYS</div>	Endomysial IgA Gliadin deamidated IgG Total IgA Tissue Transglutaminase (IgA) HLA DQ2/DQ8 <div>TAT 10 DAYS</div>	Gluten single IgE Allergen Endomysial Abs IgA Gliadin Abs deamidated IgG Tissue Transglutaminase IgA HLA DQ2/DQ8 Total IgA <div>TAT 10 DAYS</div>
GSA	GSA2	GLUT
	 	  

To determine the new Coeliac Pathway, a TDL audit of more than 12,000 requests for coeliac testing was carried out and results assessed within UKAS current guidelines. The purpose of these new guidelines is to reduce the risk of missing IgA deficient patients. The new pathway covers for this by adding a total IgA to all low **Tissue Transglutaminase (TTG)** IgA results to check for an IgA deficiency. If an IgA deficiency is identified, a reflex deamidated gliadin IgG will be carried out to determine whether the patient is likely to have coeliac disease with an IgG antibody.

The changes are as follows:

- 1 Initial TTG IgA samples are received and tested
- 2 If TTG IgA is LOW <0.1 U/ml reflex testing for Total IgA will be undertaken
- 3 If Total IgA is LOW <0.1 g/L then reflex testing for Gliadin IgG test will be undertaken
- 4 If TTG IgA is HIGH (>= 10 U/ml then reflex testing for Endomesial IgA will be undertaken as a confirmatory test.

Endomysial IgA

- This is no longer available as a stand-alone test. If requested the request will default to TTG IgA.
- However if TTG IgA is positive endomysial IgA will be carried out as a confirmatory test.
This only needs to be done once in the patients history.

Endomysial IgG requests

- No longer available as a single test request.

Immunology

Deamidated gliadin IgA requests

- This is no longer available. If requested the request will default to TTG IgA.

Deamidated gliadin IgG requests

- This can be requested as an individual standalone test as well as being incorporated into the coeliac pathway. This may be useful when testing children's samples.

Appropriate clinical comments will be added to results automatically – as follows:

TTG IgA result U/ml	Total IgA result for new assay g/L	Deamidated gliadin IgG result U/ml	Comment
0.1 to 10	N/A	N/A	Coeliac disease unlikely (please note that if the patient has no dietary gluten results may appear false negative)
≥ 10	N/A	N/A	Suggestive of coeliac disease
< 0.1	≥ 0.1	N/A	Coeliac disease unlikely (please note that if the patient has no dietary gluten, results may appear false negative)
< 0.1	< 0.1	≥ 10	Consistent with coeliac disease in a patient with selective IgA deficiency
< 0.1	< 0.1	< 7	Coeliac disease unlikely (please note that if the patient has no dietary gluten, results may appear false negative)
< 0.1	< 0.1	7-10	Result equivocal suggest referral to a gastroenterologist for consideration of duodenal biopsy

Coeliac Disease (CD) is an immune-mediated disease of the intestines that is triggered by the ingestion of gluten in genetically susceptible individuals. Gluten is the major protein component of wheat, rye, and barley. Genetic predisposition does play a key role in CD, and it is well known that CD is strongly associated with specific HLA class II genes known as HLA-DQ2 and HLA-DQ8. Approximately 95% of CD patients express HLA-DQ2, and the remaining patients are usually HLA-DQ8 positive. The negative predictive value for both tests is higher than 99%. However, the HLA-DQ2 allele is common and is carried by approximately 30% of Caucasian individuals. Thus, HLA-DQ2 or HLA-DQ8 is necessary for disease development but is not sufficient for disease development; its estimated risk effect is only 36-53%.

Note: History taking is important if a patient has been on a gluten-free diet for 6-12 months, approximately 80% will lose their antibody response. After 5 years this increases to >90%.

<div><div>RHEUMATOLOGY PROFILE 1</div><div>FBC ESR Uric Acid RF Anti CCP Antibodies (RF) C Reactive Protein</div><div>TAT 2 DAYS</div><div>RH</div></div> <div>A B</div>	<div><div>RHEUMATOLOGY PROFILE 3 Rheumatoid Disease</div><div>FBC ESR Uric Acid RF Anti CCP Antibodies (RF) Antinuclear Autoantibodies C Reactive Protein</div><div>TAT 2 DAYS</div><div>RH3</div></div> <div>A B</div>	<div><div>RHEUMATOLOGY PROFILE 5 Mono Arthritis</div><div>FBC ESR Uric Acid RF Anti CCP Antibodies (RF) Antinuclear Autoantibodies C Reactive Protein HLA B27</div><div>TAT 3 DAYS</div><div>RH5</div></div> <div>A A B B</div>
<div><div>RHEUMATOLOGY PROFILE 2 General screen for Connective Tissue Disorders</div><div>FBC ESR Uric Acid Antinuclear Autoantibodies Anti-dsDNA IgG Antibodies to Extractable Nuclear Antigens (ENA) <i>Anti nRNP</i> <i>Anti Sm</i> <i>Anti Ro (SS-A)</i> <i>Anti La (SS-B)</i> <i>Anti Jo-1</i> <i>Anti Scl 70</i> <i>Anti CENP</i> RF Anti CCP Antibodies HLA B27 C Reactive Protein CENP-B</div><div>TAT 3 DAYS</div><div>RH2</div></div> <div>A A B B</div>	<div><div>RHEUMATOLOGY PROFILE 4 Systematic Lupus Erythematosus</div><div>FBC ESR Antinuclear Autoantibodies Anti-dsDNA IgG Antibodies to Extractable Nuclear Antigens (ENA) <i>Anti nRNP</i> <i>Anti Sm</i> <i>Anti Ro (SS-A)</i> <i>Anti La (SS-B)</i> <i>Anti Jo-1</i> <i>Anti Scl 70</i> <i>Anti CENP</i> RF Anti CCP Antibodies Anti Cardiolipin Autoantibodies Complement 3,4 C Reactive Protein</div><div>TAT 2 DAYS</div><div>RH4</div></div> <div>A B B</div>	<div><div>RHEUMATOLOGY PROFILE 6 Rheumatoid Factor</div><div>RF Anti CCP Antibodies (RF) C Reactive Protein</div><div>TAT 2 DAYS</div><div>RH6</div></div> <div>B</div>
		<div><div>RHEUMATOLOGY PROFILE 7 Sjogren's Syndrome</div><div>Anti RO (SS-A) Anti La (SS-B) Salivary antibodies (SAB) C Reactive Protein</div><div>TAT 10 DAYS</div><div>RH7</div></div> <div>B</div>

Patients with Irritable Bowel Syndrome (IBS) may benefit by testing for **Calprotectin**, see page 77 for details.

<div><div>AUTOANTIBODY PROFILE I</div><div>Thyroid Peroxidase Antibodies Antinuclear Antibodies Mitochondrial Antibodies Smooth Muscle Antibodies Gastric Parietal Cell Antibodies LKM</div><div>TAT 2 DAYS</div><div>AUTO</div></div> <div>B</div>	<div><div>AUTOANTIBODY PROFILE II</div><div>Thyroid Peroxidase Antibodies Islet Cell Antibodies Adrenal Antibodies Gastric Parietal Cell Antibodies Gonadal (Ovarian/ Testicular) abs</div><div>TAT 2 DAYS</div><div>ENDO</div></div> <div>B</div>	<div><div>CHLAMYDIA SPECIES SPECIFIC (MIF) ANTIBODY SCREEN</div><div>Chlamydia trachomatis (serovar A-K & L1-L3) Chlamydia pneumoniae Chlamydia psittaci</div><div>TAT 2 DAYS</div><div>CHAB</div></div> <div>B</div>
<div><div>FAECAL CALPROTECTIN ELASTASE PROFILE</div><div>Faecal Calprotectin Faecal Elastase</div><div>TAT 5 DAYS</div><div>CEP</div></div> <div>RF</div>	<div><div>CHRONIC FATIGUE SYNDROME PROFILE</div><div>Epstein-Barr Virus Antibody Profile Lymphocyte Subsets (CD4/CD8)* CRP Vitamin D (25 OH)</div><div>TAT 5 DAYS</div><div>VIP1</div></div> <div>A + B¹⁰</div>	

Tropical and travel related immunology

TEST	CODE	SAMPLE REQ	TAT
Amoebic (<i>E. histolytica</i>) Antibodies	AFAT	B	2 days
Amoebic (<i>E. histolytica</i>) PCR	AMAG	RF	2 days
Bilharzia (Schistosome) Antibody Screen	BILH	B ¹⁴	10 days
Bilharzia (Schistosome) Antigen	SHAG	B	15 days
Bilharzia (Urine)	USCH	RU ¹⁴	2 days
Borrelia Antibodies (Lyme Disease) IgG, IgM	BORR	B ^{9,14}	2 days
Borrelia Antibodies (Lyme Disease) IgM	BORM	B	2 days
Borrelia Confirmation (Immunoblot)	BORC	B ^{9,14}	10 days
Cryptosporidium Detection by PCR	CRPA	RF	2 days
Dengue Virus Serology	DENG	B ^{9,14}	5 days
DVT/Pre-travel Screen	DVT1	A A B ⁹	5 days
Echinococcus (Hydatid) Antibodies	EFAT	B ^{9,14}	5 days
Enteric Organism Rapid Detection	EORD	RF	2 days
Filaria (Lymphatic and Non-Lymphatic) Antibodies	FIFA	B ^{9,14}	10 days
Insect/Worm/Ova/Cysts	FLEA	Send Specimen ^{9,14}	5 days
Leishmania Antibodies	LEIS	B	5 days
Malarial Antibodies (<i>Pl. falciparum</i>)	MALA	B ^{9,14}	5 days
Malarial Antibodies (species specific)	MALS	B ^{9,14}	10 days
Post-Travel Screen 1 (Prior to 6 weeks)	PTS	A A B G ¹⁴	10 days
Post-Travel Screen 2 (Prior to 6 weeks)	PTS2	A A B B B G ¹⁴	10 days
Pre-Travel Screen (DVT)	DVT1	A A B ⁹	5 days
Rickettsial Species Antibody Profile	RICK	B	7 days
Schistosome (Bilharzia) Antibodies	BILH	B ¹⁴	10 days
Schistosome Antigen	SHAG	B	15 days
Toxoplasma Antibodies (IgG+IgM)	TFAM	B ⁹	4 hours
Tropical Screen (from 6 weeks post-travel)	TROP	B B ^{9,14}	10 days
Zika Abs IgM and IgG – Antibody detection from 15 days	ZKAB	B	5 days
Zika RNA by PCR in Semen	ZIKS	Semen	5 days
Zika RT PCR – Window of detection from 1-14 days from onset of symptoms	ZIKU	RU	5-7 days
Zika RT PCR – Window of detection from 1-7 days from onset of symptoms	ZIKA	B	5-7 days

Tropical and travel related immunology

TROPICAL SCREEN
(from 6 weeks post-travel)

Amoebic Antibodies
Schistosomal Antibodies (Bilharzia)
Echinococcus Antibodies (Hydatid)
Leishmania Antibodies
Malarial Antibodies (IFA)
Toxoplasma Antibodies IgG
Toxoplasma Antibodies IgM

TAT 10 DAYS

TROP

B B ^{9,14}

POST-TRAVEL SCREEN 1
(Prior to 6 weeks)

Haematology Profile
Biochemistry Profile
Schistosome Abs
Malarial Abs

TAT 10 DAYS

PTS

A A B G ¹⁴

POST-TRAVEL SCREEN 2
(Prior to 6 weeks)

Haematology Profile
Biochemistry Profile
Schistosome Abs
Malarial Abs
Hep A IgM Abs
Hep B s Ag
Hep C Abs
HIV Duo

TAT 10 DAYS

PTS2

A A B B B G ¹⁴

DVT/PRE-TRAVEL SCREEN

FBC
Factor II Prothrombin Gene
Factor V Leiden
Anticardiolipin Antibodies

TAT 5 DAYS

DVT1

A A B ⁹

ENTERIC ORGANISM RAPID DETECTION

Detection of Bacterial, Viral and Parasitic Infection by Multiplex Real-Time PCR

Bacteria and Bacterial Toxins
C. difficile Toxin A/B gene, *Campylobacter* spp., *Enterococcus* spp. (EPEC), *Enteroinvasive E.coli* (EIEC)/*Shigella*, *Enterotoxigenic E.coli* (ETEC), *Enteropathogenic E.coli* (EPEC), *Plesiomonas shigelloides*, *Salmonella*, *Shiga-toxin producing E.coli* (STEC) *stx1/stx2*, *Shiga-toxin producing E.coli* (STEC) *O157:H7*, *Vibrio cholerae*, *Vibrio parahaemolyticus*, *Vibrio vulnificus*, *Yersinia enterocolitica*

Viruses
Adenovirus 40/41, *Astrovirus*, *Norovirus GI*, *Norovirus GII*, *Rotavirus A*, *Sapovirus* (I, II, IV, V)

Parasites
Cyclospora cayentanensis, *Cryptosporidium* spp., *Entamoeba histolytica*, *Gardia lamblia*

This does NOT include stool for m/c/s – this needs to be requested as a separate test. Please provide two samples if this is required.

TAT 2 DAYS

EORD

RF

Tropical and travel related immunology

Borrelia Antibodies (Lyme Disease) *Borrelia burgdorferi*

Presence of antibodies confirms infection with the Lyme Disease spiral bacterium (spirochaete) known as *Borrelia burgdorferi* by a bite from an infected tick. Patients bitten by an infected tick which is not removed within a day or so may develop Lyme disease. An expanding rash would usually appear at the site of the bite within 3 to 30 days in a large proportion of those infected. The rash spreads and often develops a 'bull's-eye' appearance. Many also develop flu-like symptoms with aching joints and muscles. The disease can later affect the nervous system, joints and other body systems.

Borrelia Antibodies IgM (BORM):

detectable after 2-3 weeks
increasing up to 6 weeks.

Borrelia Antibodies IgG/IgM

(BORR): detectable after several weeks increasing to maximum at 4-6 months and may remain at high levels for many years.

Borrelia Confirmation

(Immunoblot) (BORC):

The ELISA test is sensitive but has a well-documented high false positive rate giving positive results in cases of glandular fever, rheumatoid arthritis and other autoimmune conditions. If the ELISA is positive testing by Immunoblot confirms a diagnosis by Lyme disease. IgM and IgG antibodies are tested separately. It is essential that details of the IgG +IgM Elisa are provided for this test.

SPECIAL PATHOLOGY

```
Borrelia ab's Immunoblot      ~

Borrelia antibodies- Immunoblot:
-----
B. burgdorferi IgG/IgM [C6 EIA]      POSITIVE
-----
Borrelia IgG Lineblot [virastripe]
-----
IgG to Borrelia P83 antigen      Negative
IgG to Borrelia P58 antigen      Negative
IgG to Borrelia P43 antigen      Negative
IgG to Borrelia P39 antigen      Negative
IgG to Borrelia P30 antigen      Negative
IgG to Borrelia OspC antigen      POSITIVE
IgG to Borrelia p21 antigen      Negative
IgG to Borrelia Osp17 antigen     Negative
IgG to Borrelia DBPA antigen      Negative
IgG to Borrelia P14 antigen      Negative
IgG to Borrelia VlsE antigen      Negative
IgG to BORRELIA ANTIGENS INTERPRETATION Negative
-----
IgG to Borrelia IgM Lineblot [virastripe]
-----
IgM to P41 antigen               Negative
IgM to P39 antigen               Negative
IgM to Borrelia OspC antigen      POSITIVE
IgM to Borrelia Osp17 antigen     Negative
IgM to Borrelia VlsE antigen      POSITIVE
IgM to BORRELIA ANTIGENS INTERPRETATION POSITIVE
Send Imm Result & Clin detail ~
Report Comments:
-----
The C6 result is very weak but the results could be consistent with recent/current Lyme. Treat erythema migrans on clinical suspicion. If recent infection is suspected, consider sending follow up serology at 2 or more weeks after the original sample, although prompt antibiotic treatment may abrogate the antibody response. If chronic infection was suspected, no further action is needed. If still clinically concerned please contact us to discuss
```


IMMUNE STATUS			
TEST	CODE	SAMPLE REQS	TAT
Hepatitis A Immunity (IgG/IgM)	HAIM	B	4 hours
Hepatitis B Immunity	HBIM	B	4 hours
Measles Antibodies (IgG) Immunity	MEAS	B	1 day
Measles Antibodies (IgM)	MEAM	B ⁹	2 days
Measles, Mumps, Rubella (MMR)	MMR	B	1 day
Mumps Antibodies (IgG)	MUMP	B	1 day
Mumps Antibodies (IgM)	MUMM	B	1 day
Pertussis (Whooping Cough) Antibodies	PERS	B	5 days
Pneumococcal Antibody Screen	PNEU	B	7 days
Polio Virus 1, 2, 3 Antibodies	POLO	B ⁹	15 days
Rabies Antibody	RABI	B	10 days
Rubella Antibody (IgG)	RUBE	B	4 hours
Rubella Antibody (IgM)	RUBM	B	4 hours
Rubella PCR	RUBP	A / Amniotic Fluid	5 days
Tetanus Antibody	TETA	B	5 days
Varicella Zoster Antibodies (IgG)	VZOS	B	1 day
Varicella Zoster Antibodies (IgM)	VZOM	B	1 day

Hepatitis B Immunity/Vaccination

Anti HBs	
less than 10 mIU/ml	Non-immune to Hepatitis B
10–50 mIU/ml	borderline – Booster indicated
50–100 mIU/ml	low level immunity – Booster suggested
100 and over	Immune to Hepatitis B

NEEDLE STICK INJURY PROFILE	
(Donor – Not recipient) Hep B sAg Hep C Abs HIV 1+2 Abs/p24 Antigen Serum saved for 2 years	
	TAT 4 HOURS
NSI	

B B

HEPATITIS VIRAL LOAD SAMPLE INSTRUCTIONS
<p>Whole blood can be stored at 2°C to 30°C and must be centrifuged within 24 hours of specimen collection. Separate the plasma or serum from the pelleted red blood cells following the manufacturer's instructions for the tube used. Plasma or serum can be tested on the Panther system in the primary tube or transferred to a secondary Aptima Specimen Aliquot Tube (SAT) for testing on the Panther system. If not tested immediately, plasma and serum can be stored in accordance with the specifications below. If transferred to the SAT, plasma may be frozen at -20°C or -70°C, and serum may be frozen at -20°C. Do not freeze specimens in EDTA, ACD, or serum primary collection tubes.</p> <p>After centrifugation: In the primary collection tube at 2°C to 8°C for up to 3 days In the Aliquoted Tubes: at 2°C to 8°C for up to 5 days In the Aliquoted Tubes: at -20°C or -70°C for up to 90 days</p>

HEPATITIS TESTING			
TEST	CODE	SAMPLE REQS	TAT
Hepatitis (Acute) Screen	AHSC	B	4 hours
Hepatitis A (IgM)	HAVM	B	4 hours
Hepatitis A Immunity (IgG/IgM)	HAIM	B	4 hours
Hepatitis A Profile	HEPA	B	4 hours
Hepatitis A RNA by PCR	HAVR	A or B	3 weeks
Hepatitis A, B & C Profile	ABC	B	4 hours
Hepatitis B (PCR) Genotype	BGEN	A	7 days
Hepatitis B 'e' Antigen and Antibody	HEPE	B	4 hours
Hepatitis B Core Antibody – IgM	HBCM	B	4 hours
Hepatitis B Core Antibody – Total	HBC	B	4 hours
Hepatitis B DNA (Viral load) – see page 89	DNAB	A	5 days
Hepatitis B Immunity	HBIM	B	4 hours
Hepatitis B Profile	HEPB	B	4 hours
Hepatitis B Resistant Mutation	HBRM	A or B	7 days
Hepatitis B Surface Antigen	AUAG	B	4 hours
Hepatitis C Abs Confirmation (RIBA)	RIBA	B	5 days
Hepatitis C Antibodies	HEPC	B	4 hours
Hepatitis C Genotype	CGEN	A	5 days
Hepatitis C Quantification (Viral Load) – see page 89	QPCR	A or B	5 days
Hepatitis Delta Antibody	HEPD	B	5 days
Hepatitis Delta Antigen	HDAG	B	5 days
Hepatitis Delta RNA	DRNA	A (Frozen plasma)	5 days
Hepatitis E (PCR)	EHEP	A	2 weeks
Hepatitis E IgG/IgM	HBE	B	5 days
Hepatitis G (PCR)	HEPG	A (Frozen plasma)	2 weeks

HEPATITIS B PROFILE

Hep B Surface Antigen
Hep B Surface Antibodies
Hep B Core IgG/IgM

TAT
4
HOURS

HEPB

B

ACUTE VIRAL HEPATITIS SCREEN

Hepatitis A IgM Abs
Hepatitis B Surface Antigen
Hepatitis C Abs

TAT
4
HOURS

AHSC

B

HEPATITIS A, B & C PROFILE

Hepatitis A Profile
Hepatitis B Profile
Hepatitis C Abs
LFT's

TAT
4
HOURS

ABC

B

Virology

All virology samples are processed as per manufacturers sample requirements and guidelines.

Hepatitis virus is named in order of their discovery A, B, C, D, E and G.

Hepatitis A

Hepatitis A is spread through food and water that have been contaminated with the virus derived from human faeces and urine. Hepatitis is an acute infection, not a chronic form of the disease.

HBV Assays

Hepatitis B surface antigen (HBsAg) (AUAG)

A protein on the surface of HBV; it can be detected in high levels in serum during acute or chronic HBV infection. The presence of HBsAg indicates that the person is infectious. The body normally produces antibodies to HBsAg as part of the normal immune response to infection. HBsAg is the antigen used to make Hepatitis B vaccine.

Hepatitis B surface antibody (anti-HBs) (HBIM)

The presence of anti-HBs is generally interpreted as indicating recovery and immunity from HBV infection. Anti-HBs also develops in a person who has been successfully vaccinated against Hepatitis B.

Total Hepatitis B core antibody (anti-HBc) (HBC)

Appears at the onset of symptoms in acute Hepatitis B and persists for life. The presence of anti-HBc indicates previous or ongoing infection with HBV in an undefined time frame.

IgM antibody to Hepatitis B core antigen (IgM anti-HBc) (HBCM)

Positivity indicates recent infection with HBV (≤6 months). Its presence indicates acute infection.

Hepatitis B e antigen and antibody (HEPE)

Hepatitis B e antigen (HBeAg): A secreted product of the nucleocapsid gene of HBV that is found in serum during acute and chronic Hepatitis B. Its presence indicates that the virus is replicating and the infected person has high levels of HBV.

Hepatitis B e antibody (HBeAb or anti-HBe): Produced by the immune system temporarily during acute HBV infection or consistently during or after a burst in viral replication. Spontaneous conversion from e antigen to e antibody (a change known as seroconversion) is a predictor of long-term clearance of HBV in patients undergoing antiviral therapy and indicates lower levels of HBV.

HBV Viral Load (DNAB)

This assay measures the concentration of Hepatitis B viral DNA in patient serum. The test enables the viral load at the beginning of treatment to be established and, thereafter, monitored to indicate treatment success.

HBV Genotyping (BGEN)

Identifies the hepatitis B genotype (A to H) in a patient's serum/plasma. This is critical for determining treatment and monitoring response.

HBV Drug Resistance Detection (HBRM)

Detects hepatitis B virus wild-type and drug-induced mutations, associated with lamivudine, entecavir and tenofovir.

HCV Assays

HCV Antibody (HEPC)

The test indicates exposure to virus but does not necessarily signify current infection. The HCV antibody test may therefore be used to screen patients for possible HCV infection to detect the presence of antibodies to the virus, indicating exposure to HCV. This test cannot tell if the viral infection is active, only that you were exposed to the virus in the past.

HCV Viral Load (QPCR)

Measures the concentration of hepatitis C viral RNA in patient serum. This state-of-the-art assay enables the viral load at the beginning of treatment to be established and, thereafter, monitored to indicate treatment success.

HCV Genotype for Treatment (CGEN)

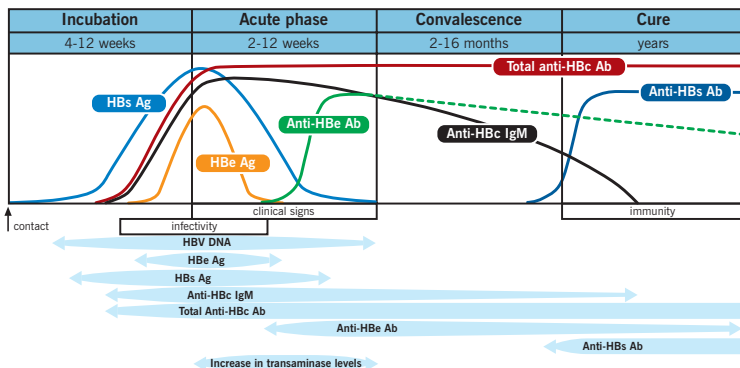
Determines the HCV genotype in a patient's serum. The result is presented as being of either Genotype [1, 5, 6], [4] or [2, 3]. This grouping reflects required treatment duration of the different genotypes.

HCV Drug Resistance

Detects hepatitis C wild-type or drug-induced mutations associated with resistance to HCV drugs including NS5A inhibitors, NS5B inhibitors or NS3 inhibitors.

- **Transmission:** Sexual, parenteral, perinatal, direct contact between individuals.
- **Clinical Signs:** Asymptomatic in 90% of cases.
- **Cure:** 95% of cases (adults).
- **Complications:** Cirrhosis and hepatocellular carcinoma.

- **Development of chronic form:**
Yes (5% of adult cases).
- **Prevention:**
Vaccination ++++; specific IgG.
- **Main Marker:**
HBS Ag, anti HBe IgM, total anti HBe Ab,
Anti-HBs Ab, HBe Ag, Anti-HBe Ab, HBV DNA.

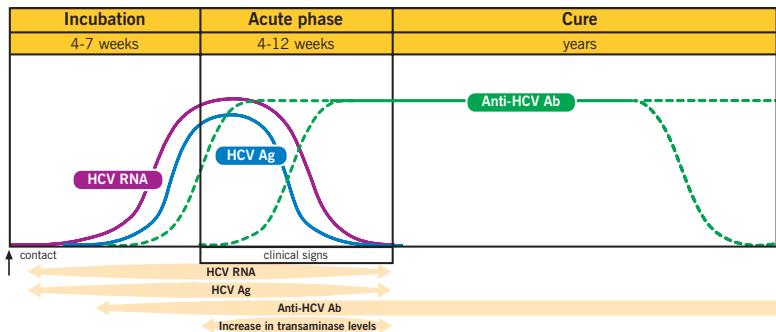
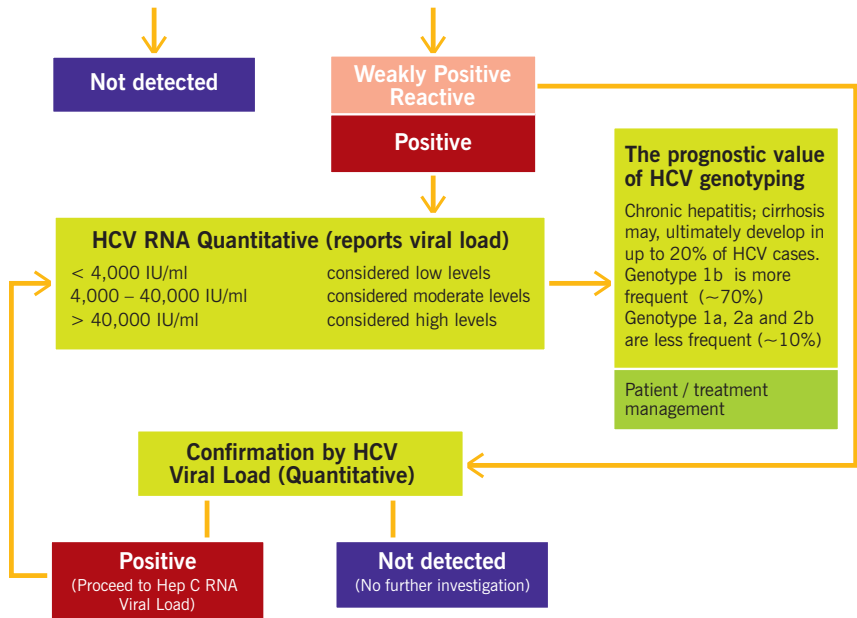


Virology

Hepatitis C Antibodies

HEPATITIS C

- **Transmission:**
Parenteral, nosocomial, sexual.
- **Clinical Signs:**
Asymptomatic in 90% of cases.
- **Cure:**
95% of cases (adults).
- **Complications:**
Cirrhosis and hepatocellular carcinoma.
- **Development of chronic form:**
Yes (80% of adult cases).
- **Prevention:**
Hygiene, no vaccination.
- **Main Marker:**
Anti HCV Ab, HCV RNA.



HIV TESTING

TEST	CODE	SAMPLE REQ	TAT
HIV Screening: HIV1& 2 Abs/p24 Ag (4th Gen)	HDUO	B	4 hours
HIV Screening: HIV1& 2 Abs, p24 (5th Gen)	HIV5	B	4 hours
HIV Confirmation of Positive Screens (Using 3 methodologies)	HIVC	B	1 day
HIV Rapid RNA HIV-1 QUALITATIVE	LHIV	A (Vacutainer only)	4 hours
HIV Rapid RNA HIV-1 QUANTITATIVE	RHIV	A (Vacutainer only)	4 hours
HIV/HBV/HCV Screen (HIV post exposure at 10 days)	STDx	A 10mls or 2x4mls (Vacutainer only)	3 days
HTLV 1& 2 Abs. (Human T Lymphotropic Virus Type I-II)	HTLV	B	8 hours
HTLV by PCR	HTLP	A Whole blood	21 days
HIV 1 Proviral DNA	HIVP	A Whole blood	7 days

TDL TINY™ SELF-COLLECTION HIV TESTS

(please refer to page 146 for information about self-collection tests)

TEST	CODE	SAMPLE REQ	TAT
4th Generation HIV1& 2 Abs/p24 Ag (45 days post-contact)*	THIV	B Tiny™	4 hours
5th Generation HIV1& 2 Abs/p24 Ag (45 days post-contact)*	THV5	B Tiny™	4 hours

*Reactive 4th & 5th Gen HIV Results require confirmation with a follow up venous blood sample.

HIV POSITIVE PATIENT MONITORING

TEST	CODE	SAMPLE REQ	TAT
HIV-1 RNA Viral Load by PCR	HIV1	A A (2x6ml whole blood)	3 days
HIV-2 RNA by PCR	HIV2	A	21 days
HIV Rapid RNA HIV-1 QUANTITATIVE	RHIV	A (Vacutainer only)	4 hours
HIV Therapeutic Drug Monitoring	TDM	J	21 days
CD3/CD4/CD8	LYSS	A ¹⁰	1 day

HIV-1 GENOTYPIC RESISTANCE TESTING

TEST	CODE	SAMPLE REQ	TAT
HIV-1 Genotypic Resistance (RT & Protease)	HIVD	A A (2x6ml whole blood)	21 days
HIV-1 Genotypic Resistance (Integrase)	INTE	A A (2x6ml whole blood)	21 days
HIV-1 Tropism	TRPM	A A (2x6ml whole blood)	28 days
HLA B*57:01	HL57	A ⁹	10 days

HLA-B*57:01 should be tested before starting patients on an Abacavir (ABC) containing regimen to reduce the risk of hypersensitivity reaction. HLA-B*57:01-positive patients should not be prescribed ABC and a positive status should be recorded as an ABC allergy in the patient's medical record.

RAPID XPERT HIV-1 RNA QUALITATIVE
EARLY DETECTION FROM 10 DAYS

HIV-1 RNA

Sample must be received in the laboratory within 24 hours of sample taking

TAT
4
HOURS

LHIV

A (Vacutainer only)

HIV/HBV/HCV SCREEN
(SIMULTANEOUS TESTING FOR
HIV1/HIV2/HBV/HCV BY PCR/NAAT)

Positive findings will be reflexed for individual qualitative confirmatory testing using the Roche Cobas Ampliscreen

HIV1 and HIV2 (RNA)
Hepatitis B Virus (HBV DNA)
Hepatitis C Virus (HCV RNA)

Samples must be received in the laboratory within 2 days of sample taking

TAT
3
DAYS

STDx

A 10mls or 2x4mls (Vacutainer only)

RAPID XPERT HIV-1 RNA VIRAL LOAD
RAPID TESTING FOR HIV-POSITIVE
PATIENT PROGNOSIS AND RESPONSE TO
ANTIRETROVIRAL THERAPY

HIV-1 RNA VIRAL LOAD (40 copies/ml)

Sample must be received in the laboratory within 24 hours of sample taking

TAT
4
HOURS

RHIV

A (Vacutainer only)

**HIV (5TH GENERATION)
Ag-Ab SCREEN**
(Bio-Rad BioPlex 2200)

HIV-1 Abs
HIV-2 Abs
HIV-1 p24 Antigen

TAT
4
HOURS

HIV5

B

TEST	CODE	SAMPLE REQ	TAT
Adenovirus by PCR	ADV	A / PCR / VS / SC	7 days
Arbovirus Antibodies/Abs	ARBO	B ^{9,14}	3 weeks
Atypical Pneumonia Screen	APS	B	2 days
Bancroftia/Oncerciasis/Filarial Antibodies	TFIF	B ¹⁴	2 weeks
BK Polyoma Virus by PCR	BKPV	A / B / RU	5 days
Cat Scratch Fever (Bartonella IgG+IgM)	CAT	B	5 days
CD3/CD4/CD8	LYSS	A ¹⁰	1 day
Chikungunya Virus Abs	CHIK	B ^{9,14}	10 days
CMV DNA (by PCR)	CMVP	A	5 days
CMV DNA by PCR (Semen)	SCVM	Semen	7 days
CMV DNA by PCR (Urine)	CMVU	RU	5 days
CMV Resistance	CMVR	A A (2 x 6mls)	21 days
NEW COVID-19 (SARS-CoV-2) Abbott IgG Antibody	GCOV	SST / Serum B * (Venous)	24 hours
NEW COVID-19 (SARS-CoV-2) RNA by PCR	NCOV	PCR Swab (nasal/ pharyngeal)	48 hours
NEW COVID-19 (SARS-CoV-2) Roche Elecsys Anti-SARS-CoV-2 Total Antibody	TCOV	SST / Serum B ** (Venous and Capillary)	24 hours
Coxsackie Antibodies (IgM)	COXM	B	10 days
CSF Screen by PCR	VPCR	CSF	2 days
Cytomegalovirus (CMV-DNA) Amnio	CMVD	AF	5 days
Cytomegalovirus (IgG/IgM) Antibodies	CMV	B	4 hours
Cytomegalovirus (PCR) Urine	CMVU	RU	5 days
Cytomegalovirus Avidity	CMAV	B	10 days
Cytomegalovirus DNA (PCR)	CMVP	A	5 days
Cytomegalovirus IgM	CMVM	B	4 hours
Dengue Fever PCR	DPCR	A or B ^{9,14}	2 weeks
Epstein-Barr Virus Antibodies IgG/IgM	EBVA	B	2 days
Epstein-Barr Virus PCR	EBVQ	A	5 days
Hantavirus Serology	HANV	B ⁹	10 days
Herpes Simplex I/II Antibody Profile (IgG)	HERP	B	2 days
Herpes Simplex I/II by PCR (Swab)	HERS	PCR	5 days
Herpes Simplex I/II by PCR (Urine)	HERD	FCRU/TPV	5 days
Herpes Simplex I/II IgM	HERM	B	2 days
HIV/HBV/HCV Screen by PCR/NAAT (10 days post exposure)	STDx	A 10mls or 2 x 4mls (Vacutainer only)	3 days
Human Herpes Virus – 6 by PCR	HHV6	A	5 days
Human Herpes Virus – 8 (IgG)	HHV8	B	10 days
Human Herpes Virus – 8 by PCR	HV8D	A	5 days
Human Parvovirus B19 – DNA	PCRP	A	2 weeks
JC Polyoma Virus by PCR	JCPV	A / B / CSF	5 days
Measles Antibodies (IgG) Immunity	MEAS	B	1 day

* Contact the laboratory for patient self-collection sample kits.

** CE marked IVD capillary kits must be used for self-collection samples and can be ordered through TDL Supplies.

TEST	CODE	SAMPLE REQS	TAT
Measles Antibodies (IgM)	MEAM	B ⁹	2 days
Measles PCR	MEAP	Buccal swab	48 hours
MERS Coronavirus Test	MERS	J	1 day
Mumps Antibodies (IgM)	MUMM	B	1 day
Mycoplasma pneumoniae IgM and IgG	MYCO	B	2 days
Mycoplasma species – DNA	MPCR	A	5 days
Needle Stick Injury Profile	NSI	B B	4 hours
Neurological Viral Screen	NVIR	B B	2 days
Parvovirus Antibodies (IgM)	PARV	B	2 days
Parvovirus DNA by PCR	PCRP	A	2 weeks
Parvovirus IgG Antibodies	PARG	B	2 days
Parvovirus IgG/IgM Abs	PARP	B	2 days
Pneumonia (Atypical) Screen	APS	B	2 days
NEW Respiratory Viral RNA Screen by PCR	FLU4	PCR nasopharyngeal	48 hours
Rotavirus in Stool by PCR	ROTA	RF	1 day
Rubella Antibody (IgG)	RUBE	B	4 hours
Rubella Antibody (IgM)	RUBM	B	4 hours
Rubella Avidity	RUAV	B	1 week
NEW SARS-CoV-2 (COVID-19) Abbott IgG Antibody	GCOV	SST / Serum B [*] (Venous)	24 hours
NEW SARS-CoV-2 (COVID-19) RNA by PCR	NCOV	PCR Swab (nasal/ pharyngeal)	48 hours
NEW SARS-Cov-2 (COVID-19) Roche Elecsys Anti-SARS-CoV-2 Total Antibody	TCOV	SST / Serum B ^{**} (Venous and Capillary)	24 hours
Torch Screen	TORC	B	2 days
Varicella Zoster – DNA	VZPC	A	5 days
Varicella Zoster Antibodies (IgG)	VZOS	B	1 day
Varicella Zoster Antibodies (IgM)	VZOM	B	1 day
Viral Antibody Screen	VIRA	B B	2 days
Viral Eye by PCR	VPE	PCR	3 days
Viral Respiratory RNA Screen by PCR	VPR	PCR or as specified on the form	2 days
Viral Skin/Mucosa by PCR	VPSK	PCR	2 days
West Nile Virus Abs	WNV	B	2 weeks
Zika Abs IgM and IgG – Antibody detection from 15 days	ZKAB	B	5 days
Zika RNA by PCR in Semen	ZIKS	Semen	5 days

VIROLOGY BY BLOOD

VIRAL ANTIBODY SCREEN	NEUROLOGICAL VIRAL SCREEN	TORCH SCREEN
Measles IgG Measles IgM Mumps IgG Mumps IgM Mycoplasma pneumonia CMV HSV 1 HSV 2	Measles IgG Measles IgM Mumps IgG Mumps IgM CMV IgG HSV 1 + 2 IgG HSV 1 + 2 IgM VZV IgG	Toxoplasma Antibodies (IgG, IgM) Rubella Antibody (IgG, IgM) CMV Antibody (IgG, IgM) Herpes Antibody (HSV1/HSV2 IgG)
TAT 2 DAYS	TAT 2 DAYS	TAT 2 DAYS
VIRA	NVIR	TORC
B B	B B	B
		ATYPICAL PNEUMONIA SCREEN
		Mycoplasma pneumonia Abs Chlamydia pneumoniae (MIF) Legionella pneumophila (IF)
		TAT 2 DAYS
		APS

VIROLOGY BY PCR

RESPIRATORY VIRAL RNA SCREEN BY PCR	VIRAL SKIN / MUCOSA BY PCR	VIRAL EYE BY PCR
Flu A Flu B Respiratory Syncytal Virus (RSV) COVID-19	<i>If chicken pox or shingles suspected, please indicate clearly on request form</i> Herpes Simplex virus Varicella Zoster virus	Herpes Simplex virus Varicella Zoster virus Adenovirus
NEW 2021 TAT 48 HOURS	TAT 2 DAYS	TAT 3 DAYS
FLU4	VPSK	VPE
PCR nasopharyngeal	PCR	PCR
		CSF SCREEN BY PCR
		Herpes Simplex virus Varicella Zoster virus Enterovirus
		TAT 2 DAYS
		VPCR
		CSF

Tumour markers/sites

TEST	CODE	SAMPLE REQ	TAT
Alpha Feto Protein	AFP	B	4 hours
Beta HCG (Oncology)	HCGQ	B	4 hours
Breast Cancer NGS Panel – full sequencing across 14 genes + deletions/duplications	GENE	A A ^{9,11}	4 weeks
CA 15-3	C153	B	4 hours
CA 19-9	C199	B	4 hours
CA 50	CA50	B	5 days
CA 72-4	C724	B	5 days
CA 125	C125	B	4 hours
Carcino Embryonic Antigen	CEA	B	4 hours
Complex PSA (Prostate Specific Ag)	CPSA	B	3 days
Cyfra 21-1	CY21	B	4 days
Early CDT-Lung	CDTL	B	7 days
HE4 + ROMA (Earlier Detection of Ovarian Tumour)	HE4	B	1 day
Neurone Specific Enolase	NSE	B	5 days
NMP22 (Bladder tumour)	NMP	J ¹	4 days
Osteocalcin	OST	B (Frozen) ⁴	4 days
Prostate Profile (Total & Free PSA)	PR2	B	4 hours
Prostate Specific Antigen (Total)*	PSPA	B	4 hours
Pyruvate Kinase (M2-PK)	M2ST	RF ⁴	5 days
Pyruvate Kinase (M2-PK)	M2PK	A	5 days
S100 Malignant Melanoma	S100	B	4 days
Squamous Cell Carcinoma	SCC	B	4 days
Testicular Tumour Profile	TTP	B	4 hours

* Results that fall between 4.00 ug/L and 10.00 ug/L will automatically reflex to a Free PSA with a calculated ratio. The ratio of Free to Total PSA may help discriminate between prostate cancer and benign prostatic hyperplasia.

TUMOUR MARKERS/SITES	
AFP: Liver, Testes	Cyfra 21-1: Oesophagus, Lung, Bladder
BHCG: Testes	HE4: Ovary
BRCA1/2: Breast	NMP22: Bladder
CA 125: Ovary	NSE: Lung, Brain, Thyroid
CA 15-3: Breast	PSA: Prostate
CA 19-9: Stomach, Colorectal, Gastrointestinal, Pancreas	S100: Melanoma
CA 50: Bladder, Colon	SCC: Oesophagus, Bronchus, Lung, Cervix
CDTL: Lung	
CEA: Stomach, Liver, Breast, Ovary, Gastrointestinal, Lung	

HE4 Earlier Detection of Ovarian Tumour	
HE4/CA 125/ROMA Calculated Algorithm for pre and post menopausal risk of malignant disease	TAT 1 DAY
HE4	

PROSTATE PROFILE Total and Free PSA	
Total PSA Free PSA Calculated Ratio	TAT 4 HOURS
PR2	

Tumour markers/sites

Site	Tumour marker	Sample type	Turnaround time
Oesophagus	CA 19-9	serum	4 hours
	CEA	serum	4 hours
	SCC	serum	4 days

Site	Tumour marker	Sample type	Turnaround time
Bronchial/ Lung	NSE*	serum	5 days
	SCC*	serum	4 days
	CDTL	serum	7 days
	CEA	serum	4 hours
	Cyfra 21-1	serum	4 days

Site	Tumour marker	Sample type	Turnaround time
Bile duct	CA 19-9	serum	4 hours
	CEA	serum	4 hours

Site	Tumour marker	Sample type	Turnaround time
Pancreas	CA 19-9	serum	4 hours
	CEA	serum	4 hours

Site	Tumour marker	Sample type	Turnaround time
Carcinoid	5-HIAA	24 hour urine/acidified	5 days

Site	Tumour marker	Sample type	Turnaround time
Bladder/ Chorion	CEA	serum	4 hours
	CA 50	serum	5 days
	NMP22	urine	4 days

Site	Tumour marker	Sample type	Turnaround time
Cervix/ Uterus	SCC	serum	4 days
	CEA	serum	4 hours

Site	Tumour marker	Sample type	Turnaround time
Prostate	Prostate Profile (Total + Free PSA)	serum	4 hours

Site	Tumour marker	Sample type	Turnaround time
Melanoma	S-100	serum	4 days

Site	Tumour marker	Sample type	Turnaround time
Thyroid	CEA	serum	4 hours
	Thyroglobulin	serum	1 day
	Calcitonin	1ml Frozen serum	

Site	Tumour marker	Sample type	Turnaround time
Breast	Breast Cancer NGS Panel	EDTA	4 weeks
	CA 15-3	serum	4 hours
	CEA	serum	4 hours

Site	Tumour marker	Sample type	Turnaround time
Liver	AFP	serum	4 hours
	CEA	serum	4 hours
	Ferritin	serum	4 hours

Site	Tumour marker	Sample type	Turnaround time
Gastro-intestine	CEA	serum	4 hours
	CA 19-9	serum	4 hours

Site	Tumour marker	Sample type	Turnaround time
Ovary	Ovarian Cancer NGS Panel	EDTA	4 weeks
	CA 125	serum	4 hours
	CA 15-3	serum	4 hours
	HE4	serum	1 day
	AFP	serum	4 hours

Site	Tumour marker	Sample type	Turnaround time
Colon	CEA	serum	4 hours
	CA 19-9	serum	4 hours
	CA 50	serum	5 days

Site	Tumour marker	Sample type	Turnaround time
Testes	AFP	serum	4 hours
	Beta HCG (quantitative)	serum	4 hours

Site	Tumour marker	Sample type	Turnaround time
	Osteocalcin	serum (frozen)	4 days

* NSE: Neurone Specific Enolase
SCC: Squamous Cell Carcinoma

TDL Genetics is a consultant-led service which is able to provide extensive expertise in the testing, diagnosis and genetic counselling of inherited disorders. Genetic tests are performed on DNA



**TDL
GENETICS**

for molecular genetic analysis and on whole chromosomes for cytogenetic analysis. Some tests are part of profiles that can be linked with assays from other TDL disciplines, such as biochemistry and haematology, to give more comprehensive results for the patient.

Genetic tests are available for:

- Prenatal diagnosis and rapid trisomy screening by Amnio-PCR
- Carrier screening
- Newborn chromosome analysis
- Confirmation of symptomatic individuals and pre-symptomatic testing
- Genetic variation that influences risk of disease
- Identity studies (paternity, zygosity, tissue typing)
- Fertility studies
- Products of conception
- Cancer

Genetic testing is sometimes complex and tests will vary in their ability to detect mutations or to detect all patients who have, or will develop, the disease. Some tests are diagnostic for a condition, others are indicative or are associated with an altered risk for a condition. Results can affect the lives of individuals and have implications for their family, for insurance and employment. Where testing will predict the inheritance of a disease in a healthy person, counselling and consent are mandatory. For these tests, please complete the Genetic Request form at the back of the guide (including informed consent). Our service provides result interpretation and risk assessment to patients and their family members. Genetic counselling can be arranged by TDL's Consultant Clinical Geneticist.

To meet the increasing range and complexity of genetic testing we have developed an excellent collaboration with other specialist laboratories.

Tests marked GENE are sent to these laboratories within our network and have a fixed price.

GENE panel composition may change throughout the year to reflect new and improved developments.

Turnaround times may be longer if follow-up studies are required.

Specimen Receipt at The Doctors Laboratory is 24 hours a day. Specifically, TDL Genetics results service is available Monday to Friday 8.30am–5.30pm with the laboratory also open for processing of samples on Saturdays from 9.00am–1.00pm.

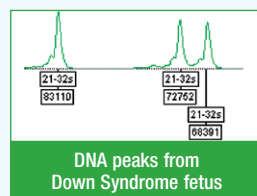
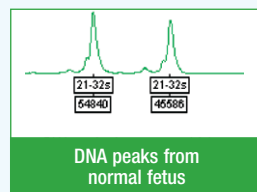
Test codes, sample requirement codes, and turnaround times may be found on the following pages.

All samples must be collected in the specified containers, as shown in the key at the back of this guide.

Samples should be fresh and in good condition (e.g. not clotted if EDTA or heparinised whole blood is required) otherwise testing may be adversely affected and another sample may be required. Small DNA samples are stored routinely for one year, larger DNA samples can be stored by special arrangement.

Instructions for transportation, sample labelling, and the completion of request forms can be found on the reverse of the TDL Genetics Request Form.

The locations of the Laboratory and Patient Reception are indicated on the map on the reverse of each request form. If you do not find the test you require in this directory or need more information and advice please telephone the laboratory on 020 7307 7409.



Sending samples to the laboratory

Transport arrangements

All specimens should be kept at room temperature and despatched to the laboratory as soon as possible, by TDL/international courier, first class post, guaranteed next day delivery or a reliable alternative.

If a delay in sending the sample is unavoidable, please refrigerate overnight – DO NOT FREEZE. Specimens must not be allowed to come in contact with request forms, but should be kept separate by using dual – pocketed plastic bags. Specimens for inland postage must be packed in a rigid crush-proof container according to current Post Office guidelines. IATA guidelines should be followed for international transport (Advice is available from the laboratory).

Labelling of high risk samples

Please note that it is the responsibility of the referring clinician to ensure that high-risk samples are clearly identified to reduce the risk of infection to staff and others.

Patient details on request forms and samples

Request and consent forms are available directly from TDL Genetics.

In order to avoid unnecessary time spent in obtaining details please provide the following information:

Information for request forms:

- Surname, forename (not initials), date of birth and biological sex of patient for postnatal referrals
- Full name (not initials) and location of referring clinician
- Full address of clinician to whom the result should be sent
- Legible clinical summary, including details of any relevant family history
- Address for billing – Doctor, patient or other
- Gestation on prenatal samples
- Hospital or reference number
- Test required

Essential information on sample container label:

- Patients surname and forename (not initials)
- Date of birth
- Hospital number or reference number

Consent forms

Consent forms (at the back of this guide) are available for genetic testing. As genetic testing may have implications for other family members and is regarded as personal data, it is recommended that written consent is obtained wherever possible. In cases with predictive testing for severe disorders, as indicated in the laboratory guide, it is essential that patients should also be offered formal genetic counselling. It is the responsibility of the referring clinician to obtain appropriate consent from the patient.

Unlabelled samples

Unlabelled samples will **ONLY** be processed if the individual who took the sample can confirm the sample is from the patient in question. In the absence of this assurance, the sample will be discarded and a repeat required.

Genetic Testing

THE IMPORTANCE OF CLINICAL DETAILS

Clinical details are very important when providing genetic analysis. The more clinical information that is available (e.g. details of ultrasound information, phenotypic features or family history) the better the service we can provide. Failure to provide this information for cytogenetic studies may result in an inaccurate analysis.

MOLECULAR GENETICS

Clinical details can be extremely important for clinical interpretation of a molecular genetic test.

For example, the clinical comments accompanying a cystic fibrosis screening report will vary depending on whether the patient is a potential gamete donor or a person exhibiting a cystic fibrosis phenotype.

It may also be crucial, where a mutation has already been shown to be segregating in a family, to be provided with information concerning the mutation and a family pedigree to ensure the correct analysis is performed and reliable risk figures calculated.

CYTOGENETICS

Cytogenetic analysis is performed according to the Professional Guidelines for the Association of Clinical Genetic Science and the recommendations provided are dependent on the clinical indications given for each case.

Clinical details inform the investigation at all stages:

- Prior to analysis, clinical details may indicate, for example, that procedures such as chromosome breakage or leukaemic studies are required, which must be referred to the oncogenomic department or specialist centre.
- During analysis they may indicate that extra cells should be screened to investigate the possibility of mosaicism, for example in a diagnosis of suspected Turner syndrome, or that particular chromosomes must be targeted for high-resolution study, for example chromosome 4 in suspected Wolf-Hirschhorn syndrome.
- When the analysis has been completed they may help to provide an accurate interpretation of the findings and in some instances prompt further investigations, for example FISH or molecular genetic studies.

When clinical details are not available a routine analysis will be performed and a conditional report issued.

SAMPLE STABILITY

Molecular Genetic Samples

Whole blood collected in EDTA should be sent to the laboratory between 4°C-28°C within 48 hours.

Long term storage should be at 2-8°C.

Extracted DNA samples should be sent to the laboratory between 4°C-28°C.

Cytogenetic Samples

Cytogenetic studies require living cells, please ensure that samples reach the laboratory as soon as possible. If a delay before dispatch is unavoidable, samples may be stored in a refrigerator (4°C) but they must **not** be frozen.

Samples sent more than 48 hours after sampling, or kept at temperatures below 4°C and greater than 38°C may have inhibited growth.

Information concerning packaging, transportation, and labelling of samples is provided on the reverse of our TDL Genetics Request Form.

Requesting additional tests

Any further tests not requested at the time of sample receipt must be requested within:

- 1 week for tests requiring prenatal culture or cultured cells
- 2 weeks for DNA testing
- 2 weeks for cell culture testing
- 3 months for FISH testing

Samples can be stored for longer periods if specifically requested at the time of sample receipt.

POSTNATAL DIAGNOSIS (BLOOD CULTURE)

Reasons for analysis: Chromosome studies are requested where problems that may have a cytogenetic basis are suspected, e.g. babies with birth defects; children with developmental delay and physical handicaps, or adults with fertility problems. Additionally, prospective gamete donors are screened to detect carriers of balanced chromosome rearrangements.

Sample requirements: Lithium heparin whole blood specimens are required – gently mixed to prevent clotting and must **not** be frozen. See sample stability section for cytogenetic samples. Sample volumes may be reduced for children (2-4ml) and neonates (1-2ml).

Turnaround time: The usual turnaround time is 2-3 weeks however the laboratory will endeavour to respond to urgent requests. Where a major trisomy is suspected, a rapid PCR screen may be performed to provide an urgent provisional result.

Notes

- a) Rarely, blood samples fail to culture (<1%);
- b) The culture may yield chromosomes of insufficient quality. This will be indicated on the report and a repeat study suggested;
- c) The laboratory should be informed if the patient has recently received a blood transfusion.
- d) The laboratory should be informed if the patient has EVER had a bone marrow transplant.
- e) The patient's biological sex should be included on the request form.

PRENATAL DIAGNOSIS

Reasons for analysis: Chromosome studies are requested where pregnancies are identified as being at risk of a cytogenetic abnormality e.g. positive maternal serum screening combined NT test; fetal abnormalities found on ultrasound; or where a parent is a known carrier of a chromosome anomaly, or where a high risk trisomy has been found by NIPT.

Sample requirements:

- a) amniotic fluid – 10ml+ in a plain sterile, leak-proof container. Suitable containers can be provided by the laboratory. The specimen must **not** be frozen. See sample stability section for cytogenetic samples.
- b) chorionic villus – 5mg+ in sterile transport medium. Suitable containers containing medium can be provided by the laboratory. The specimen must **not** be frozen. See sample stability section for cytogenetic samples.
- c) fetal blood – 1-2ml LITHIUM HEPARIN whole blood, gently mixed to prevent clotting. The specimen must **not** be frozen. See sample stability section for cytogenetic samples.

Turnaround time: This is dependent on the rate of cell growth, however, the usual turnaround time is approximately 2 weeks. A number of circumstances now occur more frequently, as invasive prenatal diagnosis becomes less common, that may result in delayed reporting time. These include:

- a) A delay in transportation in order to collect a batch of samples to reduce courier costs. Even when couriered promptly, sample growth may be slower than that seen in samples sent immediately.
- b) Sampling at early or late gestations, for example to confirm non-invasive tests or follow up anomaly scans.
- c) A tendency to take smaller quantities of sample or to take insufficient sample for multiple techniques.
- d) The request for karyotyping as an add-on after an initial PCR test.

Fetal blood results will usually be reported by 10 calendar days. **For all other prenatal tests, please contact the laboratory prior to taking samples.**

Always provide Clinical Details and Family History with requests for Genetic Tests.

Key: See page 21 for sample taking and special handling instructions.

Notes

- a) Maternal contamination, and mosaicism may complicate the analysis and may lead to the suggestion that a second invasive test is performed.
- b) Rarely, cultures fail to grow (overall <1%)
- c) Very small chromosome abnormalities may not be detected (this is why the phrase 'No trisomies or major chromosome abnormalities detected...' is used in our reports).
- d) for TTIs or heavily blood stained amniocentesis samples, please provide a maternal EDTA blood sample for comparison studies.

SOLID TISSUE

Reasons for analysis: Fibroblast cultures may be used in addition to blood cultures, for example where tissue specific mosaicism is suspected, or where blood samples cannot be obtained. POC samples may be requested for early spontaneous miscarriages, stillbirths, or to confirm a prenatal diagnosis.

Sample requirements: All specimens should be placed in a sterile container, preferably containing transport medium. This can be supplied by the laboratory. Sterile normal saline can be used if transport medium is not available. Samples must not be placed in formaldehyde or other preservative and must not be frozen. See sample stability section for cytogenetic samples.

Turnaround time: This is dependent on the rate of cell growth, however, the usual turnaround time is approximately 4 weeks.

Notes

- a) Material from miscarriages has a relatively high culture failure rate (around 20%). Where failure occurs, alternative molecular methods may be attempted, usually a KaryoLite Bacs-on-Beads assay that can detect whole monosomy or trisomy of any chromosome, if possible.
- b) If no villus or fetal parts are identified in supposedly POC material and a normal female chromosome result is found, this may indicate that maternal tissue has been cultured (this will be noted on our report).
- c) Material from miscarriages can be returned for sensitive disposal if requested at the time of receipt. If no special request is made, fetal material will be sent for incineration separate from general clinical waste. Placental and other POC material will be disposed of in general clinical waste for incineration.

FLUORESCENCE IN SITU HYBRIDISATION (FISH)

Where FISH studies for specific microdeletion syndromes are required this must be indicated on the request form.

Note: FISH studies for a rapid pre or postnatal aneuploidy screen have now been superseded in our laboratory by multiplex-PCR technology. Subtelomeric screens are now performed by Array CGH as part of developmental delay investigations. Common microdeletion syndrome testing is now performed by BOBs analysis.

CELL LINE KARYOLOGY

The cytogenetics laboratory can perform cell line karyology on live cultures or fixed cells suspensions (recommended) on a research basis. Please note: a laboratory processing charge of £100+VAT is applicable to those cases wherein a successful analysis cannot be obtained. Please contact the laboratory for further details.

STATEMENT REGARDING MEASUREMENT UNCERTAINTY (MU)

Measurement Uncertainty is determined for each measurement procedure in the examination phase used to report measured quantity values on patients' samples. This is determined during verification of this assay for service introduction; creation of laboratory standard operating procedures (SOP) and interpretation of the results.

Where examinations include a measurement step but do not report a measured quantity value, the laboratory calculates the uncertainty of the measurement step where it has utility in assessing the reliability of the examination procedure or has influence on the reported result.

Estimates of measurement uncertainty are regularly reviewed and are available upon request to laboratory users.










































KEY PERSONNEL

Consultant Clinical Geneticist	Prof. Michael Patton	020 7307 7409	michael.patton@tdlpathology.com
Consultant Clinical Scientist	Elaine Holgado	020 7307 7409	elaine.holgado@tdlpathology.com
Head of Cytogenetics	Rebecca Watts	020 7460 4787	rebecca.watts@hslpathology.com
Senior Cytogeneticist	Kath Masters	020 7307 7409	kath.masters@tdlpathology.com
Cytogenetics Operations Manager	Emma Wilcock	020 7307 7409	emma.wilcock@tdlpathology.com
Postnatal Lab Manager	Allison Daffern	020 7307 7409	allison.daffern@tdlpathology.com
Director of Genetics & Molecular Pathology	Dr Lisa Levett	020 7307 7409	lisa.levett@tdlpathology.com
Head of Genetics & Molecular Pathology	Dr Stuart Liddle	020 7307 7409	stuart.liddle@tdlpathology.com
Operations Manager	Andrew Levett	020 3908 1282	andrew.levett@tdlpathology.com
Molecular Cytogenetics Manager	Alessandra Callegari	020 7307 7409	alessandra.callegari@tdlpathology.com

TEST	CODE	SAMPLE REQ	TAT
1p36 Deletion Syndrome – karyotype + FISH	KARY, FISH	CVS/AF/ H ⁹	12-17 days
21-Hydroxylase Deficiency (Congenital Adrenal Hyperplasia) – 8 mutations screened	GENE	A ^{9,11}	8 weeks
22q11 & 10p14 deletion (Di George Syndrome) – BOBs only	DGB	CVS/AF/ A ⁹	5 days
22q11 & 10p14 deletion (Di George Syndrome) – BOBs (5 days) + karyotype (15 days)	DGB, KARY	CVS/AF/ A H ⁹	5-15 days
Achromatopsia NGS Panel – full sequencing across 6 genes plus BCM genes	GENE	A A ⁹	4 weeks
Aicardi-Goutières Syndrome NGS Panel – full sequencing across 7 genes	GENE	A A ⁹	6 weeks
Alagille Syndrome NGS Panel – full sequencing JAG1 + NOTCH2 genes	GENE	A A ⁹	6 weeks
Alpha Fetoprotein on Amniotic fluid	AFPA	AF ⁹	5-10 days
Alpha Thalassaemia – multiplex PCR for common large deletions	GENE	A ⁹	4 weeks
Alpha-1 Antitrypsin Genotype – PI*M, PI*S, PI*Z	GENE	A ⁹	4 weeks
Alport Syndrome NGS Panel – full sequencing COL4A3 + COL4A4 + COL4A5 + MYH9 genes	GENE	A A ⁹	6 weeks
Amelogenesis/Dentinogenesis Imperfecta NGS Panel – full sequencing across 31 genes	GENE	A A ⁹	6 weeks
AML/ALL Molecular MRD – NPM1, PML-RARA, CBFβ-MYH11, RUNX1-RUNX1T1, ETV6-RUNX1 – <i>Contact Lab for further information</i>	GENE	Bone Marrow / A ⁹	5 days
AmnioBOBs only – rapid aneuploidy diagnosis for all chromosomes + common microdeletion syndromes	ABOB	AF ⁹	5 days
Amniocentesis – rapid BOBs aneuploidy diagnosis for all chromosomes (5 days) + culture (10-15 days) – see profiles	ABK	AF ⁹	5-15 days
Amniocentesis – rapid PCR diagnosis for common aneuploidies (2 days) + culture (10-15 days)	APCC	AF ⁹	2-15 days
Amniocentesis culture (karyotype) only	ACUL	AF ⁹	10-15 days
AmnioPCR only – rapid common aneuploidy diagnosis by QF-PCR	APC	AF ⁹	2 days
Amyotrophic Lateral Sclerosis (Motor Neurone Disease) NGS Panel – full sequencing across 43 genes	GENE	A A ⁹	6 weeks
Androgen Insensitivity – AR gene sequencing	GENE	A ⁹	8 weeks
Aneurysm/Connective Tissue Disorders/Ehlers-Danlos Syndrome NGS Panel – full sequencing across 46 genes + deletions/duplications	GENE	A A ⁹	4 weeks
Angelman Syndrome (Primary Screen) – methylation PCR	PWAM	A ⁹	5 days
Angelman/Rett Syndromes NGS Panel – full sequencing across 30 genes	GENE	A A ⁹	6 weeks
Aniridia, Isolated – PAX6 gene sequencing + deletions/duplications	GENE	A ⁹	8 weeks

Always provide Clinical Details and Family History with requests for Genetic Tests.

Key: See page 21 for sample taking and special handling instructions.

TEST	CODE	SAMPLE REQ	TAT
Anophthalmia/Microphthalmia NGS Panel – full sequencing across 39 genes	GENE	  ⁹	6 weeks
Antithrombin Deficiency – SERPINC1 Gene Variant Analysis (Known Genotype)	ATMA	  (Whole Blood 10ml) ⁴⁰	6 weeks
Antithrombin Deficiency – SERPINC1 Gene Variant Analysis (Unknown Genotype)	ATMA	  (Whole Blood 10ml) ⁴⁰	12 weeks
Aortopathy/Marfan Syndrome/Loeys-Dietz Syndrome NGS Panel – full sequencing across 30 genes	GENE	  ⁹	6 weeks
Apert Syndrome – 2 common FGFR2 mutations	GENE	 ⁹	4 weeks
Apolipoprotein E genotype – E2, E3, E4	APEG	 ⁹	5 days
Array CGH (Comparative Genomic Hybridisation)	CGH	CVS/ AF/   ⁹	10 days
Arrhythmogenic Right Ventricular Cardiomyopathy (ARVC) NGS Panel – sequencing across 46 genes + deletions/duplications	GENE	  ⁹	4 weeks
Ashkenazi Breast Cancer Screen – 3 common mutations	GENE	Requires patient informed consent  ^{9,11}	
Ashkenazi Jewish Carrier Screen – see Pan-ethnic/ Jewish Carrier Profile	ASHJ	 ⁹	4 weeks
Ataxia/Episodic Ataxia Disorders NGS Panel – full sequencing across 152 genes	GENE	  ⁹	6 weeks
Autoinflammation/Periodic Fever NGS Panel – full sequencing across 36 genes	GENE	  ⁹	6 weeks
Azoospermia – karyotype + cystic fibrosis screen + polyT(5T) + Y deletions	GRP	  ⁹	10-15 days
B cell clonality assay (IgH and IgK)	IGHA	 or FFPE	2 weeks
Bardet-Biedl Syndrome NGS Panel – full sequencing across 24 genes	GENE	  ⁹	6 weeks
Batten Disease (Neuronal Ceroid Lipofuscinosis) NGS Panel – full sequencing across 14 genes	GENE	  ⁹	6 weeks
BCR-ABL diagnostic assay	BCRD	 ⁹	2 weeks
BCR/ABL Quantitative – fusion gene sizes p190 + p210 – MUST arrive in the laboratory within 48 hours, before 12pm on Fridays	BCRA	  ⁹	10 days
Becker Muscular Dystrophy – deletions/duplications	DND	 ⁹	10 days
Beckwith-Wiedemann Syndrome – methylation studies on 11p15 imprinting domains KvDMR + H19	GENE	 ⁹	4 weeks
Behcet's Disease – HLA Tissue Typing B*51	B51	 ⁹	10 days
Beta Thalassaemia – beta-globin gene sequencing	GENE	 ⁹	4 weeks
Bleeding and platelet disorders NGS Panel (known familial mutations) – Contact lab	GENE	 	6 weeks
Bleeding and platelet disorders NGS Panel (unknown familial mutations) – Contact lab	GENE	 	12 weeks
Blood PCR for Chromosome 21	BPCR	 ⁹	5 days
Bloom Syndrome – BLM gene sequencing	GENE	 ⁹	4 weeks
BOBs rapid chromosome analysis – see profiles			
BRAF V600E mutation by PCR for Hairy Cell Leukaemia	GENE	Bone Marrow/ 	5 days

Always provide Clinical Details and Family History with requests for Genetic Tests.
Turnaround times are quoted as working days.




























TDL Genetics

TEST	CODE	SAMPLE REQ	TAT
Breast Cancer Ashkenazi Screen – 3 common mutations	GENE	Requires patient informed consent A ^{9,11}	4 weeks
Breast Cancer – BRCA1 + BRCA2 only gene sequencing + deletions/duplications	GENE	A	4 weeks
Breast Cancer NGS Panel – full sequencing across 14 genes + deletions/duplications	GENE	Requires patient informed consent A A ^{9,11}	4 weeks
Brugada Syndrome/Long-QT NGS Panel – full sequencing across 34 genes	GENE	A A ⁹	4 weeks
C-KIT D816V mutation by PCR for Mastocytosis	GENE	Bone Marrow / A	5 days
CADASIL – NOTCH3 gene sequencing	GENE	A ⁹	6 weeks
CAKUT (Congenital Anomalies of Kidney & Urinary Tract) NGS Panel – full sequencing across 38 genes	GENE	A A ⁹	6 weeks
Calreticulin – CALR exon 9 mutation screen	CALR	A ⁹	2 weeks
Cancer, Comprehensive NGS Panel – full sequencing across 123 genes + deletions/duplications	GENE	Requires patient informed consent A A ^{9,11}	4 weeks
Carbohydrate Metabolism Deficiency NGS Panel – full sequencing across 47 genes + deletions/duplications + mitochondrial DNA	GENE	A A ⁹	4 weeks
Cardio-Facio-Cutaneous/Noonan/LEOPARD/Costello Syndromes NGS Panel – full sequencing across 20 genes	GENE	A A ⁹	6 weeks
Cardiomyopathy, Arrhythmogenic Right Ventricular NGS Panel – sequencing across 34 genes + deletions/duplications	GENE	A A ⁹	4 weeks
Cardiomyopathy, Comprehensive NGS Panel – full sequencing across 111 genes + deletions/duplications	GENE	A A ⁹	4 weeks
Cardiomyopathy, Dilated NGS Panel – full sequencing across 78 genes + deletions/duplications	GENE	A A ⁹	4 weeks
Cardiomyopathy, Hypertrophic NGS Panel – full sequencing across 86 genes + deletions/duplications	GENE	A A ⁹	4 weeks
Carrier Screen (Pan-ethnic or Jewish) – see profiles	GENE	A ⁹	4 weeks
Charcot-Marie-Tooth Syndrome NGS Panel – full sequencing across 59 genes	GENE	A A ⁹	6 weeks
Charcot-Marie-Tooth Type 1A – PMP22 duplications	GENE	A ⁹	4 weeks
CHARGE Syndrome – CHD7 gene sequencing	GENE	A ⁹	8 weeks
Chediak-Higashi Syndrome – LYST gene sequencing	GENE	A ⁹	4 weeks
Cholestasis, Intrahepatic NGS Panel – full sequencing across 15 genes	GENE	A A ⁹	6 weeks
Chromosome Analysis (Amniocentesis) – culture only	ACUL	AF ⁹	10-15 days
Chromosome Analysis (Amniocentesis) – rapid BOBs aneuploidy diagnosis for all chromosomes (5 days) + culture (10-15 days) – see profiles	ABK	AF ⁹	5-15 days
Chromosome Analysis (Amniocentesis) – rapid PCR diagnosis for common aneuploidies (2 days) + culture (10-15 days)	APCC	AF ⁹	2-15 days

Always provide Clinical Details and Family History with requests for Genetic Tests.

Key: See page 21 for sample taking and special handling instructions.

TDL Genetics

TEST	CODE	SAMPLE REQ	TAT
Chromosome Analysis (Blood)	KARY	 ⁹	8-18 days
Chromosome Analysis (Chorionic Villus) – rapid BOBs aneuploidy diagnosis for all chromosomes (5 days) + culture (10-15 days) – see profiles	CBK	CVS ⁹	5-15 days
Chromosome Analysis (Chorionic Villus) – rapid PCR diagnosis for common aneuploidies (2 days) + culture (10-15 days)	CVPC	CVS ^{1,9}	2-15 days
Chromosome Analysis (Chorionic Villus) – culture only	CVSC	CVS ^{1,9}	10-15 days
Chromosome Analysis (Products of Conception) – BOBs rapid aneuploidy diagnosis for all chromosomes (5 days) + culture (25 days)	PBK	Placental Sample ^{1,9}	5-25 days
Chromosome Analysis (Products of Conception) – reflex to BOBs testing if culture fails to grow	PROC	Placental Sample ^{1,9}	20-25 days
Chromosome Analysis (Solid Tissue)	PROC	Fetal tissue ^{1,9}	4-5 weeks
Chromosome Analysis (Stem Cells)	STEM/ SUSP	Culture/Fixed cells	Contact lab
Chromosome Y Deletion – AZFa, AZFb, AZFc + SRY	YDEL	 ⁹	5 days
Cockayne Syndrome NGS Panel – full sequencing ERCC6 + ERCC8	GENE	  ⁹	5 weeks
Celiac Disease – HLA DQ2/DQ8 Genotype	Q2Q8	 ⁹	10 days
Colorectal Cancer NGS Panel – full sequencing across 18 genes + deletions/duplications	GENE	Requires patient informed consent   ^{9,11}	4 weeks
Comparative Genomic Hybridisation (Array CGH)	CGH	CVS/AF/   ⁹	10 days
Congenital Absence of Vas Deferens – karyotype + cystic fibrosis screen + polyT(5T) + Y deletions	GRP	  ⁹	10-15 days
Congenital Central Hypoventilation Syndrome (CCHS) – PHOX2B polyalanine repeat analysis	GENE	 ⁹	4 weeks
Congenital Central Hypoventilation Syndrome (CCHS) – full sequencing PHO X2B gene	GENE	 ⁹	4 weeks
Congenital Muscular Dystrophy NGS Panel – full sequencing across 27 genes	GENE	  ⁹	6 weeks
Connective Tissue Disorders/Ehlers-Danlos Syndrome/Aneurysm NGS Panel – full sequencing across 46 genes + deletions/duplications	GENE	  ⁹	5 weeks
Connexin-26 Associated Deafness – full sequencing GJB2 gene (+ GJB6 common deletion)	GENE	 ⁹	8 weeks
Cornelia de Lange Syndrome NGS Panel – full sequencing across 8 genes	GENE	  ⁹	6 weeks
Costello/Noonan/LEOPARD/Cardio-Facio-Cutaneous Syndromes NGS Panel – full sequencing across 20 genes	GENE	  ⁹	6 weeks
Craniosynostosis and related disorders NGS Panel	GENE	 	6 weeks
Cri du Chat Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS/AF/   ⁹	5-15 days
Cri du Chat Syndrome – BOBs only	PBOB	CVS/AF/  ⁹	5 days
CVS PCR for common aneuploidies (2 days) + culture (10-15 days)	CVPC	CVS ⁹	2-15 days
























Always provide Clinical Details and Family History with requests for Genetic Tests.
Turnaround times are quoted as working days.

TDL Genetics




















TEST	CODE	SAMPLE REQ	TAT
CVSBOBs – rapid BOBs aneuploidy diagnosis for all chromosomes (3-5 days) + culture (10-15 days) – see profiles	CBK	CVS ⁹	5-15 days
CVSBOBs only – rapid aneuploidy diagnosis for all chromosomes + common microdeletion syndromes	CBOB	CVS ⁹	5 days
CYP450 2D6 Genotyping	TGEN	A ⁹	10 days
Cystic Fibrosis (139 common mutations) – reflex to Poly T when required	CFS	A ⁹	5 days
Deafness NGS Panel – full sequencing across 179 genes	GENE	A A ⁹	6 weeks
Deafness, Non-Syndromic – GJB2 sequencing + GJB6 common deletion	GENE	A ⁹	8 weeks
Dentinogenesis/Amelogenesis Imperfecta NGS Panel – full sequencing across 31 genes	GENE	A A ⁹	6 weeks
Diabetes Mellitus, MODY NGS Panel – full sequencing across 13 genes	GENE	A A ⁹	6 weeks
Diabetes Mellitus, Neonatal NGS Panel – full sequencing across 26 genes	GENE	A A ⁹	6 weeks
DiGeorge Syndrome (22q11 & 10p14 deletion) – BOBs (5 days) + karyotype (15 days)	DGB, KARY	CVS/AF/A H ⁹	5-15 days
DiGeorge Syndrome (22q11 & 10p14) – BOBs only	DGB	CVS/AF/A ⁹	5 days
Dihydropyrimidine Dehydrogenase deficiency screening (Fluoropyrimidine Toxicity) – 5 mutations	GENE	A ⁹	1-2 weeks
Dilated Cardiomyopathy NGS Panel – full sequencing across 78 genes + deletions/duplications	GENE	A A ⁹	4 weeks
DNA Extraction & Storage – 3 years (longer upon request)	XDNA	A ⁹	20 days
DNA Identity Profile – 15 STR markers	DNAF	A ⁹	10 days
Doyme Honeycomb Retinal Dystrophy – EFEMP1 screening	GENE	A ⁹	4 weeks
Duchenne Muscular Dystrophy – deletions/duplications only	DMD	A ⁹	10 days
Duchenne Muscular Dystrophy – full sequencing DMD1 gene	GENE	A ⁹	6 weeks
DVT/Pre-travel Screen	DVT1	A A B ⁹	5 days
Ehlers-Danlos Syndrome/Aneurysm/Connective Tissue Disorders NGS Panel – full sequencing across 46 genes + deletions/duplications	GENE	A A ⁹	5 weeks
Endometrial Cancer NGS Panel – full sequencing across 10 genes + deletions/duplications	GENE	Requires patient informed consent A A ^{9,11}	4 weeks
Epidermolysis Bullosa, Comprehensive NGS Panel – full sequencing across 13 genes	GENE	A A ⁹	6 weeks
Epidermolysis Bullosa, Simplex Panel – full sequencing of KRT5 + KRT14 genes	GENE	A A ⁹	8 weeks
Epilepsy, Adolescent/Adult Onset Panel – sequencing across 83 genes + deletions/duplications	GENE	A	6 weeks

Always provide Clinical Details and Family History with requests for Genetic Tests.

Key: See page 21 for sample taking and special handling instructions.

TEST	CODE	SAMPLE REQ	TAT
Epilepsy, Childhood Panel – full sequencing across 211 genes + deletions/duplications	GENE		6 weeks
Epilepsy, Comprehensive NGS Panel – full sequencing across 400 genes + deletions/duplications	GENE	 ⁹	6 weeks
Epilepsy, Neonatal Panel – sequencing across 278 genes + deletions/duplications	GENE		6 weeks
Epilepsy, Progressive Myoclonic Panel – sequencing across 18 genes + deletions/duplications	GENE		6 weeks
Exudative Vitreoretinopathy, Familial (FEVR) NGS Panel – full sequencing NDP + FZD4 + LRP5 + TSPAN12 + ZNF408 genes	GENE	 ⁹	4 weeks
Eye Developmental Disease NGS Panel – full sequencing across 59 genes	GENE	 ⁹	4 weeks
Fabry Disease, X-linked – GLA gene sequencing	FABM	 ⁹	4 weeks
Facioscapulohumeral Muscular Dystrophy (FSHD) – D4Z4 repeat deletion – <i>Contact lab prior to sending</i>	GENE	 ⁹	8 weeks
Factor II Prothrombin – G20210A mutation	FX2	 ⁹	5 days
Factor V Leiden – G1691A mutation	FX5	 ⁹	5 days
Factor VII Deficiency – F7 Gene Variant Analysis (Known Genotype)	7MA	 (Whole blood 10ml) ⁴⁰	6 weeks
Factor VII Deficiency – F7 Gene Variant Analysis (Unknown Genotype)	7MA	 (Whole blood 10ml) ⁴⁰	12 weeks
Factor X Deficiency – F10 Gene Variant Analysis (Known Genotype)	10MA	 (Whole blood 10ml) ⁴⁰	6 weeks
Factor X Deficiency – F10 Gene Variant Analysis (Unknown Genotype)	10MA	 (Whole blood 10ml) ⁴⁰	12 weeks
Factor XI Deficiency – F11 Gene Variant Analysis (Known Genotype)	11MA	 (Whole blood 10ml) ⁴⁰	6 weeks
Factor XI Deficiency – F11 Gene Variant Analysis (Unknown Genotype)	11MA	 (Whole blood 10ml) ⁴⁰	12 weeks
Familial Adenomatous Polyposis (FAP) – full sequencing across 18 genes + deletions/duplications	GENE	Requires patient informed consent  ^{9,11}	4 weeks
Familial Exudative Vitreoretinopathy (FEVR) NGS Panel – full sequencing NDP + FZD4 + LRP5 + TSPAN12 + ZNF408 genes	GENE	 ⁹	4 weeks
Familial Hypercholesterolaemia – LDLR + APOB + PCSK9 + LDLRAP1 screening	GENE	 ⁹	4 weeks
Familial Hypocalcaemic Hypercalcaemia (FHH) Panel – full sequencing CASR + AP2S1 + GNA11 genes	GENE	 ⁹	8 weeks
Familial Mediterranean Fever – hotspot sequencing MEFV gene	GENE	 ⁹	4 weeks
Familial Medullary Thyroid Carcinoma – hotspot sequencing RET gene	GENE	Requires patient informed consent  ^{9,11}	8 weeks
Fatty Acid Oxidation Deficiency NGS Panel – full sequencing across 22 genes	GENE	 ⁹	6 weeks





























Always provide Clinical Details and Family History with requests for Genetic Tests.
Turnaround times are quoted as working days.

TEST	CODE	SAMPLE REQ	TAT
NEW Fever (Recurrent) Screening – across 4 genes	GENE		6 weeks
FLT3-ITD and FLT3-TKD screening assay	FLT3		3-5 days
Fragile X Syndrome screen – FMR1 repeat analysis PCR (3 weeks) + Southern Blot (8 weeks) if required	GENE	 ⁹	3-8 weeks
Friedreich Ataxia – frataxin gene repeat analysis	GENE	 ⁹	4 weeks
Gastric Cancer NGS Panel – full sequencing across 15 genes + deletions/duplications	GENE	Requires patient informed consent  ^{9,11}	
Gaucher Disease – 8 common mutations	GENE	 ⁹	4 weeks
Gaucher Disease full gene sequencing	GDMA	 ⁴⁰	4 weeks
Genetic Reproductive Profile (Male) – see profiles	GRP	 ⁹	10-15 days
Gilbert Syndrome – common UGT1A1 repeat variation	GENE	 ⁹	6 weeks
Glaucoma NGS Panel – full gene sequencing across 26 genes	GENE	 ⁹	6 weeks
Glucose-6-Phosphate Dehydrogenase (G6PD) Deficiency – full G6PD gene sequencing	GENE	 ⁹	4 weeks
Glycogen storage disease type 2 (Pompe) mutation analysis	POMP	 ⁹	4 weeks
Haemochromatosis – HFE common mutations C282Y + H63D	HMD	 ⁹	3 days
Haemolytic-Uremic Syndrome NGS Panel – full sequencing across 15 genes	GENE	 ⁹	8 weeks
Haemophilia A Variant Analysis (Known Genotype) – F8 Intron 22 Inversion, F8 Intron 1 Inversion, Sequence analysis of known variants for F8 gene	HACD	 (Whole blood 10ml) ⁴⁰	6 weeks
Haemophilia A Variant Analysis (Unknown Genotype) – F8 Intron 22 Inversion, F8 Intron 1 Inversion, Sequence analysis of unknown variants for F8 gene	GENE	 (Whole blood 10ml) ⁴⁰	12 weeks
Haemophilia A CVS Variant Analysis (Known Genotype) – F8 Intron 22 Inversion, F8 Intron 1 Inversion, Sequence analysis of known variants for F8 gene	8CVS	CVS ⁴⁰	3 days
Haemophilia B Variant Analysis (Known Genotype) – Sequence analysis of known variants for F9	HBCD	 (Whole blood 10ml) ⁴⁰	6 weeks
Haemophilia B Variant Analysis (Unknown Genotype) – Sequence analysis of unknown variants for F9	HBMA	 (Whole blood 10ml) ⁴⁰	12 weeks
Haemophilia B CVS Variant Analysis (Known Genotype) – Sequence analysis of known variants for F9	9CVS	CVS ⁴⁰	3 days
Harmony® Prenatal Test (Non-Invasive Prenatal Testing) – common aneuploidy screening from maternal blood	NIPT	J/Special tubes ¹	3-5 days
Harmony® Prenatal Test (Non-Invasive Prenatal Testing) – common aneuploidy screening from maternal blood including 22q11.2 del	NIPQ	J/Special tubes ¹	3-5 days
Hearing Loss NGS Panel – full sequencing across 179 genes	GENE	 ⁹	6 weeks









































Always provide Clinical Details and Family History with requests for Genetic Tests.

Key: See page 21 for sample taking and special handling instructions.

TDL Genetics

TEST	CODE	SAMPLE REQ	TAT
Hemiplegic Migraine, Familial NGS Panel – full sequencing across 6 genes + mtDNA	GENE		5 weeks
Hereditary Cancer NGS Panel, Comprehensive – full sequencing across 127 genes + deletions/duplications	GENE	Requires patient informed consent 	4 weeks
Hereditary Hemorrhagic Telangiectasia – ACVRL1 + ENG full sequencing + deletions/duplications	GENE		8 weeks
Hereditary Neuropathy NGS Panel – full sequencing across 39 genes	GENE		6 weeks
Hereditary Neuropathy with Liability to Pressure Palsy – PMP22 deletion analysis	GENE		4 weeks
Hereditary Non-Polyposis Colon Cancer (Lynch Syndrome) NGS Panel – full sequencing across 18 genes + deletions/duplications	GENE	Requires patient informed consent 	4 weeks
Hereditary Pancreatitis – PRSS1 hotspot sequencing + deletions/duplications + SPINK1 N34S common mutation	GENE		8 weeks
Hereditary Spastic Paraplegia NGS Panel – full sequencing across 262 genes + deletions/duplications + mitochondrial DNA	GENE		5 weeks
Hermansky-Pudlak Syndrome/Oculocutaneous Albinism/Pigmentation NGS Panel – full sequencing across 30 genes	GENE		4 weeks
HFE gene (Haemochromatosis) – common mutations C282Y + H63D	HMD		3 days
Hirschprung Disease NGS Panel – full sequencing across 6 genes + copy number variant	GENE		4 weeks
HLA Tissue Typing A	HLA		10 days
HLA Tissue Typing A+B	HLBA		10 days
HLA Tissue Typing A+B+C (Class I)	HABC		10 days
HLA Tissue Typing A/B/DRB1/3/4/5	HLAF		10 days
HLA Tissue Typing A/B/DRB1/3/4/5/DQB1	HLF		10 days
HLA Tissue Typing A/B/C/DRB1/3/4/5/DQB1 (Class I & II)	HLFC		10 days
HLA Tissue Typing B	HLB		10 days
HLA Tissue Typing B*27 only	HLAB		3 days
HLA Tissue Typing B*51 (Behcet's Disease)	B51		10 days
HLA Tissue Typing B*57:01 high resolution	HL57		10 days
HLA Tissue Typing C	HLC		10 days
HLA Tissue Typing Coeliac Disease – DQ2/DQ8	Q2Q8		10 days
HLA Tissue Typing DRB1/3/4/5	DRB1		10 days
HLA Tissue Typing DRB1/3/4/5/DQB1 (Class II)	HLDQ		10 days
HLA Tissue Typing Narcolepsy – DQB1*06:02	GENE		4 weeks
Huntington Disease – HD gene repeat analysis PCR	GENE	Requires patient informed consent 	4 weeks
Hyperinsulinism NGS Panel – full sequencing across 8 genes	GENE		8 weeks

Always provide Clinical Details and Family History with requests for Genetic Tests.
Turnaround times are quoted as working days.

TEST	CODE	SAMPLE REQ	TAT
Hyperparathyroidism – CASR sequencing	GENE	 ⁹	8 weeks
Hypertriglyceridemia NGS Panel – full sequencing across 47 genes	GENE	  ⁹	8 weeks
Identity Profile (DNA) – 15 STR markers	DNAF	 ^{9,11}	10 days
IgVH mutation analysis for CLL	IGMU	 ⁹	4 weeks
Incontinentia Pigmenti, X-linked – IKBKG/NEMO common mutation	GENE	 ⁹	4 weeks
Intellectual Disability NGS Panel – full sequencing across 560 genes + deletions/duplications	GENE	  ⁹	6 weeks
Intrahepatic Cholestasis NGS Panel – full sequencing ABCB11 + ABCB4 + ATP8P1	GENE	  ⁹	6 weeks
Iron Overload Profile	IOP	   ⁹	3 days
JAK 2 – exon 12 sequencing (rare mutations) – MUST arrive in the laboratory within 48 hours, before 12pm on Fridays	GENE	 ⁹	4 weeks
JAK2 V617F genotyping assay	JAK2	 ⁹	2 weeks
Jervell and Lange-Nielsen Syndrome – full sequencing KCNE1 + KCNQ1 genes	GENE	  ⁹	6 weeks
Jewish/Pan-ethnic carrier screening – see profiles	ASHJ	 ⁹	4 weeks
Joubert/Meckel-Gruber Syndrome NGS Panel – full sequencing across 39 genes	GENE	  ⁹	6 weeks
Kallmann Syndrome NGS Panel – full sequencing across 19 genes	GENE	  ⁹	6 weeks
Karyotype – See Chromosome Analysis			
Kennedy Disease (Spinal Bulbar Muscular Atrophy) – AR repeat expansion	GENE	 ⁹	6 weeks
Ketolysis Disorders NGS Panel – full sequencing across 7 genes	GENE	  ⁹	6 weeks
Kidney/Urinary Tract Cancer NGS Panel – full sequencing across 27 genes + deletions/duplications	GENE	Requires patient informed consent   ^{9,11}	4 weeks
Krabbe Disease – GALC sequencing + 502T/del common deletion	GENE	 ⁹	6 weeks
Lactose Intolerance Gene	LACG	 ⁹	2 weeks
Langer-Giedion Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS/AF/   ⁹	5-15 days
Langer-Giedion Syndrome – BOBs only	PBOB	CVS/AF/  ⁹	5 days
Leber's Congenital Amaurosis NGS Panel – full sequencing across 32 genes	GENE	  ⁹	6 weeks
Lebers Hereditary Optic Neuropathy – m.3460G>A + m.11778G>A + m.14484T>C common mutations	GENE	 ⁹	8 weeks
Leigh Syndrome NGS Panel – full sequencing across 78 genes + deletions/duplications + mitochondrial DNA	GENE	  ⁹	4 weeks
LEOPARD/Noonan/Cardio-Facio-Cutaneous/Costello Syndromes NGS Panel – full sequencing across 20 genes	GENE	  ⁹	6 weeks
Leukaemia Fusion Gene Screening Assay (Q30)	LMPX	 ⁹	2 weeks

Always provide Clinical Details and Family History with requests for Genetic Tests.






























Key: See page 21 for sample taking and special handling instructions.

TDL Genetics

TEST	CODE	SAMPLE REQ	TAT
Li-Fraumeni Syndrome (p53-related cancer predisposition) – TP53 sequencing + MLPA	GENE	Requires patient informed consent A ^{9,11}	6 weeks
Limb-Girdle Muscular Dystrophy (LGMD) NGS Panel – full sequencing across 34 genes	GENE	A A ⁹	6 weeks
Lissencephaly NGS Panel – full sequencing across 14 genes	GENE	A A ⁹	8 weeks
Loeys-Dietz Syndrome/Marfan Syndrome/Aortopathy NGS Panel – full sequencing across 30 genes	GENE	A A ⁹	8 weeks
Long-QT Syndrome/Brugada Syndrome – full sequencing across 34 genes	GENE	A A ⁹	4 weeks
Lowie (Oculocerebrorenal) Syndrome – OCRL sequencing + large deletions	GENE	A ⁹	8 weeks
Lung Disorders NGS Panel – full sequencing across 51 genes	GENE	A A ⁹	6 weeks
Lynch Syndrome (HNPCC) NGS Panel – full sequencing across 18 genes + deletions/duplications	GENE	Requires patient informed consent A A ^{9,11}	4 weeks
Lysosomal Disorders NGS Panel – full sequencing across 106 genes	GENE	A A ⁹	6 weeks
Male Genetic Reproductive Profile – see profiles	GRP	A H ⁹	10-15 days
Marfan Syndrome – FBN1 sequencing + deletions/duplications	GENE	A ⁹	5 weeks
Marfan Syndrome/Loeys-Dietz Syndrome/Aortopathy NGS Panel – full sequencing across 30 genes	GENE	A A ⁹	6 weeks
Maturity-Onset Diabetes of the Young (MODY) NGS Panel – full sequencing across 14 genes	GENE	A A ⁹	6 weeks
Meckel-Gruber/Joubert Syndrome NGS Panel – full sequencing across 39 genes	GENE	A A ⁹	6 weeks
Medium-Chain Acyl-CoA Dehydrogenase Deficiency – ACADM sequencing	GENE	A ⁹	4 weeks
Melanoma NGS Panel – full sequencing across 14 genes + deletions/duplications	GENE	Requires patient informed consent A A ^{9,11}	4 weeks
Microdeletion (common) Syndromes – BOBs only	PBOB	CVS/AF/A ⁹	5 days
Microphthalmia/Anophthalmia/Coloboma NGS Panel – full sequencing across 39 genes	GENE	A A ⁹	6 weeks
Miller-Dieker Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS/AF/A H ⁹	5-15 days
Miller-Dieker Syndrome – BOBs only	PBOB	CVS/AF/A ⁹	5 days
Mitochondrial genome – full mitochondrial DNA sequencing + deletions	GENE	A ⁹	5 weeks
Mitochondrial genome sequencing	GENE	A ⁹	5 weeks
Motor Neurone Disease (Amyotrophic Lateral Sclerosis) NGS Panel – full sequencing across 43 genes	GENE	A A ⁹	6 weeks
MPL exon 10 analysis	MPL	A	2 weeks
MTHFR – common C677T + A1298C mutations	MTHF	A ⁹	5 days

































Always provide Clinical Details and Family History with requests for Genetic Tests.
Turnaround times are quoted as working days.

TDL Genetics





















TEST	CODE	SAMPLE REQ	TAT
Mucopolysaccharidosis NGS Panel – full sequencing across 11 genes	GENE	  ⁹	8 weeks
Multiple Endocrine Neoplasia Type 1 – full MEN1 sequencing	GENE	Requires patient informed consent  ^{9,11}	8 weeks
Multiple Endocrine Neoplasia Type 2 – RET gene hotspot sequencing	GENE	Requires patient informed consent  ^{9,11}	8 weeks
Myotonic Dystrophy Type 1 – DMPK repeat PCR	GENE	 ⁹	4 weeks
Myotonic Dystrophy Type 2 (PRMM) – ZNF9 repeat PCR	GENE	 ⁹	4 weeks
Narcolepsy (HLA DQB1*06:02)	GENE	 ⁹	4 weeks
Nephrotic Syndrome, Steroid-Resistant NGS Panel – full sequencing across 14 genes	GENE	  ⁹	6 weeks
Nervous System/Brain Cancer NGS Panel – full sequencing across 27 genes + deletions/duplications	GENE	Requires patient informed consent   ^{9,11}	4 weeks
Neurofibromatosis Type 1 – NF1 + SPRED1 sequencing + deletions/duplications – <i>Contact lab prior to sending</i>	GENE	Requires patient informed consent   ^{9,11}	8 weeks
Neurofibromatosis Type 2 (Bilateral Acoustic) – NF2 sequencing + deletions/duplications	GENE	 ⁹	8 weeks
Neuronal Ceroid Lipofuscinosis (Batten Disease) NGS Panel – full sequencing across 14 genes	GENE	  ⁹	6 weeks
Non-Invasive Prenatal Testing – common aneuploidy screening from maternal blood	NIPT	J/Special tubes ¹	3-5 days
Non-Invasive Prenatal Testing – common aneuploidy screening from maternal blood including 22q11.2 del	NIPQ	J/Special tubes ¹	3-5 days
Noonan Syndrome Prenatal Screening – PTPN11 exons 3 & 8 only	GENE	CVS/AF	2 weeks
Noonan/LEOPARD/Cardio-Facio-Cutaneous/Costello Syndromes NGS Panel – full sequencing across 20 genes	GENE	  ⁹	6 weeks
NPM1 mutascreen assay	NPM1	 ⁹	2 weeks
Nystagmus, X-linked Infantile – FRMD7 gene sequencing	GENE	 ⁹	4 weeks
Oculocutaneous Albinism/Hermansky-Pudlak Syndrome/Pigmentation NGS Panel – full sequencing across 30 genes	GENE	  ⁹	4 weeks
Oculopharyngeal Muscular Dystrophy – PABPN1 repeat analysis	GENE	 ⁹	4 weeks
Optic Atrophy NGS Panel – full sequencing OPA1 + OPA3 genes	GENE	  ⁹	4 weeks
Osteogenesis Imperfecta NGS Panel – full sequencing across 14 genes	GENE	  ⁹	5 weeks
Ovarian Cancer NGS Panel – full sequencing across 16 genes + deletions/duplications	GENE	Requires patient informed consent   ^{9,11}	4 weeks

Always provide Clinical Details and Family History with requests for Genetic Tests.

Key: See page 21 for sample taking and special handling instructions.

TEST	CODE	SAMPLE REQ	TAT
p53-related cancer predisposition (Li-Fraumeni Syndrome) – TP53 sequencing + MLPA	GENE	Requires patient informed consent  ^{9,11}	6 weeks
Pan-Ethnic/Jewish Carrier Screening – see profiles	GENE	 ⁹	4 weeks
Pancreatic Cancer NGS Panel – full sequencing across 22 genes + deletions/duplications	GENE	Requires patient informed consent   ^{9,11}	4 weeks
Pancreatitis (Hereditary) – PRSS1 hotspot sequencing + deletions/duplications + SPINK1 N34S common mutation	GENE	 ⁹	8 weeks
Paraganglioma/Pheochromocytoma NGS Panel – full sequencing across 11 genes + deletions/duplications	GENE	Requires patient informed consent   ^{9,11}	4 weeks
Paternity Testing (postnatal and prenatal) – sample required from each person being tested (3 people)	PATT	 / AF/CVS ^{9,11,12} Contact lab	5 days
Pelizaeus-Merzbacher Disease – PLP1 sequencing + deletions/duplications	GENE	 ⁹	8 weeks
Pendred Syndrome – SLC26A4 gene sequencing	GENE	 ⁹	4 weeks
Periodic Fever/Autoinflammation NGS Panel – full sequencing across 36 genes	GENE	  ⁹	6 weeks
Peutz-Jegher Syndrome – STK11 sequencing + deletions/duplications	GENE	 ⁹	8 weeks
Phelan-McDermid Syndrome – karyotype + FISH	KARY, FISH	CVS/AF/  ⁹	12-17 days
Pheochromocytoma/Paraganglioma NGS Panel – full sequencing across 11 genes + deletions/duplications	GENE	Requires patient informed consent   ^{9,11}	4 weeks
Pigmentation/Oculocutaneous Albinism/ Hermansky-Pudlak Syndrome NGS Panel – full sequencing across 30 genes	GENE	  ⁹	4 weeks
POLG-Related Disorders – full POLG sequencing + copy number variant	GENE	 ⁹	5 weeks
Polycystic Kidney/NGS Panel – full sequencing across 7 genes	GENE	  ⁹	6 weeks
Pontocerebellar Hypoplasia NGS Panel – full sequencing across 9 genes	GENE	  ⁹	6 weeks
Postnatal array CGH	CGH	  ⁹	10 days
Prader-Willi Syndrome (Primary Screen) – methylation PCR	PWAM	 ⁹	5 days
Prenatal array CGH	CGH	Amniotic fluid or CVS ⁹	10 days
Prenatal Diagnosis (BOBs + Culture)	ABK or CBK	AF/CVS ⁹	3-5 days, 15 days
Prenatal Diagnosis for haemoglobinopathies	PND	CVS/Amniocentesis/ fetal blood	3 days
Pre-Travel Screen (DVT)	DVT1	    ⁹	5 days
Primary Ciliary Dyskinesia (PCD) NGS Panel – full sequencing of 38 genes	GENE	  ⁹	6 weeks








































TDL Genetics

TEST	CODE	SAMPLE REQ	TAT
Primary Hyperoxaluria Panel – full sequencing across 3 genes + CNV	GENE		6 weeks
Products of Conception – rapid BOBs aneuploidy diagnosis for all chromosomes (5 days) + culture (25 days)	PBK	Placental Sample ^{1,9}	5-25 days
Products of Conception BOBs only – rapid aneuploidy diagnosis for all chromosomes	KBOB	Placental Sample or Solid Tissue ^{1,9}	3-6 days
Prostate Cancer NGS Panel – full sequencing across 12 genes + deletions/duplications	GENE	Requires patient informed consent  ^{9,11}	4 weeks
Protein C Deficiency – PROC Gene Variant Analysis (Known Genotype)	PCMA	 (Whole blood 10ml) ⁴⁰	6 weeks
Protein C Deficiency – PROC Gene Variant Analysis (Unknown Genotype)	PCMA	 (Whole blood 10ml) ⁴⁰	12 weeks
Pseudoachondroplasia (Multiple Epiphyseal Dysplasia) – COMP hotspot sequencing	GENE	 ⁹	8 weeks
PTEN-related disorders (including Bannayan-Riley-Ruvalcaba, Cowden & Proteus Syndromes) – sequencing + deletions/duplications	GENE	 ^{9,11}	8 weeks
QF-PCR rapid common aneuploidy screen	APC	AF/A ⁹	1-2 days
NEW Recurrent Fever Screening – across 4 genes	GENE		6 weeks
Recurrent Miscarriage Profile (female) – see profiles	RMP	 ^{9,18}	10-15 days
Renal Cysts and Diabetes (RCAD) – HNF-1 β sequencing + deletions/duplications	GENE	 ⁹	8 weeks
Renal/Urinary Tract Cancer NGS Panel – full sequencing across 28 genes + deletions/duplications	GENE	Requires patient informed consent  ^{9,11}	4 weeks
Retinal Dystrophy/NGS Panel – full sequencing across 537 genes	GENE	 ⁹	5 weeks
Retinoblastoma – RB1 sequencing + deletions/duplications	GENE	Requires patient informed consent  ^{9,11}	8 weeks
Rett Syndrome (MECP2 gene only) – full sequencing + deletions/duplications	GENE	Requires patient informed consent  ^{9,11}	8 weeks
Rett/Angelman Syndromes NGS Panel – full sequencing across 30 genes	GENE	 ⁹	6 weeks
Sarcoma NGS Panel – full sequencing across 26 genes + deletions/duplications	GENE	Requires patient informed consent  ^{9,11}	4 weeks
Short-Chain Acyl-CoA Dehydrogenase Deficiency – ACADS sequencing	GENE	 ⁹	5 weeks
Short Stature – SHOX mutation screening + deletions/duplications	GENE	 ⁹	8 weeks
Silver-Russell Syndrome – methylation studies on 11p15 imprinting domains KvDMR + H19	GENE	 ⁹	4 weeks
Skeletal Dysplasia NGS Panel – full sequencing across 179 genes	GENE	 ⁹	6 weeks
Smith-Lemli-Opitz Syndrome – DHCR7 sequencing	GENE	 ⁹	8 weeks

Always provide Clinical Details and Family History with requests for Genetic Tests.











Key: See page 21 for sample taking and special handling instructions.

TDL Genetics

TEST	CODE	SAMPLE REQ	TAT
Smith-Magenis Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS/AF/   ⁹	5-15 days
Smith-Magenis Syndrome – BoBs only	PBOB	CVS/AF/  ⁹	5 days
Sotos Syndrome (Cerebral Gigantism) – NSD1 sequencing + deletions/duplications	GENE	 ⁹	5 weeks
Spastic Paraplegia NGS Panel – full sequencing across 262 genes + deletions/duplications + mitochondrial DNA	GENE	  ⁹	5 weeks
Spinal Bulbar Muscular Atrophy (Kennedy Disease) – AR repeat analysis	GENE	 ⁹	6 weeks
Spinal Muscular Atrophy – SMN1 deletions/duplications	SMA	 ⁹	10 days
Spinocerebellar Ataxia – multiplex SCA1+2+3+6+7+17 common repeat expansions	GENE	 ⁹	4 weeks
SRY (Sex-determining Region Y)	SRY	 ⁹	2 days
Stargardt/Macular Dystrophy NGS Panel – full sequencing across 13 genes	GENE	  ⁹	4 weeks
Stickler Syndrome NGS Panel – full sequencing across 11 genes	GENE	  ⁹	6 weeks
Systemic mastocytosis – C-Kit common mutation (KIT D816V)	GENE	Bone Marrow /  ⁹	5 days
T cell clonality assay (TCR beta and TCR gamma)	TCRA	 or FFPE	2 weeks
Tay Sachs Screen – 5 common mutations. See also Pan-Ethnic/Jewish Carrier Profile	GENE	 ⁹	5 weeks
Thrombotic Risk Profile – see profiles	PROP	      ¹⁸	5 days
Thyroid Cancer NGS Panel – full sequencing across 7 genes + deletions/duplications	GENE	Requires patient informed consent   ^{9,11}	4 weeks
Torsion Dystonia (DYT1) – TOR1A common mutation c.904-906delGAG	GENE	 ⁹	5 weeks
Treacher-Collins Syndrome NGS Panel – full sequencing POLR1C + POLR1D + TCOF1	GENE	  ⁹	6 weeks
Tuberous Sclerosis – full TSC1 + TSC2 gene sequencing	GENE	  ⁹	5 weeks
Uni Parental Disomy (UPD) – parents and child – <i>Specify chromosome</i>	Specify type	 ^{9,12}	5 days
Urinary Tract/Renal Cancer NGS Panel – full sequencing across 28 genes + deletions/duplications	GENE	Requires patient informed consent   ^{9,11}	4 weeks
Usher Syndrome NGS Panel – full sequencing across 19 genes	GENE	  ⁹	6 weeks
Very Long-Chain Acyl-CoA Dehydrogenase Deficiency – ACADVL sequencing	GENE	 ⁹	6 weeks
Von Hippel-Lindau Syndrome – VHL sequencing + deletions/duplications	GENE	 ⁹	8 weeks
Von Willebrands Disease – Type 2 (Ex28) Variant Analysis (VWF) (Known Genotype)	VW2A	  ⁴⁰ (Whole blood 10ml)	6 weeks

Always provide Clinical Details and Family History with requests for Genetic Tests.
Turnaround times are quoted as working days.

TDL Genetics

TEST	CODE	SAMPLE REQ	TAT
Von Willebrands Disease – Type 2 (Ex28) Variant Analysis (VWF) (Unknown Genotype)	VW2A	 (Whole blood 10ml) ⁴⁰	12 weeks
Von Willebrands Disease – Type 2 VWD Variant Analysis (VWF) (Known Genotype)	2AVW	 (Whole blood 10ml) ⁴⁰	6 weeks
Von Willebrands Disease – Type 2 VWD Variant Analysis (VWF) (Unknown Genotype)	2AVW	 (Whole blood 10ml) ⁴⁰	12 weeks
Von Willebrands Disease – Type 2N Variant Analysis (VWF) (Known Genotype)	VW2N	 (Whole blood 10ml) ⁴⁰	6 weeks
Von Willebrands Disease – Type 2N Variant Analysis (VWF) (Unknown Genotype)	VW2N	 (Whole blood 10ml) ⁴⁰	12 weeks
Wolf-Hirschhorn Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS/AF/  ⁹	5-15 days
Wolf-Hirschhorn Syndrome – BOBs only	PBOB	CVS/AF/  ⁹	5 days
Y chromosome microdeletions – AZFa + AZFb + AZFc + SRY	YDEL	 ⁹	5 days
Zellweger Syndrome NGS Panel – full sequencing across 12 genes	GENE	 ⁹	6 weeks
Zygosity testing – comparative DNA profile	DNAC	 (From each twin and both parents) ⁹	5 days

Always provide Clinical Details and Family History with requests for Genetic Tests.

Key: See page 21 for sample taking and special handling instructions.

TDL Genetics

ARRAY CGH TESTING

Chromosome abnormalities can be associated with developmental delay, autism spectrum disorder, learning difficulties, dysmorphic features and other congenital abnormalities.

Array CGH can detect smaller genetic changes than is possible by conventional karyotyping, and can provide accurate information on the size and possible consequences of the gains (duplications) or losses (deletions) identified. Multiple studies have shown that Array CGH, when applied to appropriate patients, will detect up to three times more pathogenic chromosome imbalances than karyotyping due to its greater precision and sensitivity.

Array CGH testing is now considered to be the front line test for patients presenting with developmental delay (motor or growth), autism spectrum disorder, moderate to severe learning difficulties, dysmorphic features, with or without congenital abnormalities. Consortiums in the USA and many EU countries have adopted Array CGH as the front line test in this patient cohort.

Array CGH is now more frequently used for prenatal studies as an adjunct or replacement for conventional cytogenetic studies, particularly where structural fetal abnormalities are seen at ultrasound scan but also at a patient's or doctor's request. The technique may also be utilised as a follow up test to characterise anomalies detected by a previous study (e.g. an apparently balanced de novo rearrangement or marker chromosome).

When to use Array CGH

In postnatal cases, patients should present with one or more of the following:

- Mental retardation
- Autism/autism spectrum disorder
- Congenital malformations
- Developmental delay
- Dysmorphic features

In prenatal cases, patients may present with:

- Abnormalities or increased nuchal translucency on ultrasound scan which may be associated with a chromosome imbalance.

Approximately 10-20% of results identify extra or missing DNA which may or may not be relevant to the clinical phenotype, and will require further family studies to assist with interpretation.

What can Array CGH detect?

Deletions and duplications with greater sensitivity than conventional karyotyping.

What does Array CGH not detect?

- Balanced chromosome rearrangements such as translocations or inversions.
The chromosome location of duplications (this would require additional FISH testing).
- Low frequency mosaicism (<30% abnormal cells), some types of polyploidy like triploidy, Uniparental disomy (UPD) and Fragile X syndrome, imprinting defects, genetic diseases caused by point mutations or multifactorial inheritance.

Further information is provided by the UNIQUE website at www.rarechromo.org

TEST	CODE	SAMPLE REQs	TAT
Postnatal array CGH	CGH	  ⁹	10 days

Blood from both parents may also be provided in case of follow up studies at no extra charge.

TEST	CODE	SAMPLE REQs	TAT
Prenatal array CGH	CGH	Amniotic fluid or CVS ⁹	10 days

EDTA and heparin blood from both parents should be provided at the time of prenatal sampling, if possible, in case of follow up studies at no extra charge.

Always provide Clinical Details and Family History with requests for Genetic Tests.
Turnaround times are quoted as working days.

NON-INVASIVE PRENATAL TESTING (NIPT)

The Harmony test is a cell-free DNA-based prenatal blood screen. It is being used in more than 100 countries around the world, and has been used to guide clinical care in over 1.4 million pregnancies. The test can be used in singleton, twin, and egg-donor pregnancies and has been validated for use in pregnant women aged 18 to 48. It can be administered as early as 10 weeks gestation.

The test can screen for:

- Trisomies 21, 18, and 13
- Sex chromosome aneuploidy
- Monosomy X
- Fetal sex
- 22q11.2 deletion

Patient information

Non-invasive prenatal testing (NIPT) analyses cell-free DNA circulating in a pregnant mother's blood. It is used screen for Down syndrome (trisomy 21) and other common chromosomal conditions (trisomies 18 and 13). Options are also available to screen for X and Y chromosome conditions or for a deletion in chromosome 22q11.2.

About the test

DNA from the fetus circulates in the mother's blood. Cell-free DNA (cfDNA) results from the natural breakdown of fetal cells (presumed to be mostly placental) and clears from the maternal system within hours of giving birth.

During a pregnancy, cfDNA can be tested to give the most accurate screening approach in estimating the risk of a fetus having a common chromosome condition sometimes called a trisomy. This occurs when there are three copies of a particular chromosome instead of the expected two. The test looks to detect the following conditions:

- **Trisomy 21** is the most common trisomy at the time of birth. Also called Down syndrome, it is associated with moderate to severe intellectual disabilities and may also lead to digestive disease, congenital heart defects and other malformations.

- **Trisomy 18** (Edwards syndrome) and **Trisomy 13** (Patau syndrome) are associated with a high rate of miscarriage. These babies are born with severe brain abnormalities and often have congenital heart defects as well as other birth defects. Most affected individuals die before or soon after birth, and very few survive beyond the first year of life.
- **Sex chromosome conditions** occur when there is a missing, extra, or incomplete copy of the X or Y chromosomes. The Harmony test with sex chromosome aneuploidy panel option can assess risk for XXX, XYY, XXY (Klinefelter syndrome), and a missing X chromosome in a girl (Turner syndrome).

Options are also available to look for Turner syndrome only (and not the other sex chromosome conditions), and/or to look for a deletion in chromosome 22q11.2. If the mother is interested in having this optional testing, she should talk with her healthcare provider to determine if it is right for her. This option is not available for twin pregnancies.

Risk

The testing is non-invasive: it involves taking a blood sample from the mother. The pregnancy is not put at risk of miscarriage, or from other adverse outcomes that are associated with invasive testing procedures such as amniocentesis.

Accuracy

A 'high probability' result is indicative of a high probability for a trisomy. In singleton pregnancies, the test identifies more than 99% of fetuses with trisomy 21, 97% of fetuses with trisomy 18, 94% of fetuses with trisomy 13, and 96% of fetuses with Turner syndrome. X and Y analysis provides >99% accuracy for fetal sex. Accuracy for detecting other sex chromosome anomalies varies by condition.

After the test, less than 1% of women need to have a CVS or an amniocentesis procedure.

The Harmony test is considered a prenatal screening test, not a diagnostic test. So if the test results show there is a high risk of the fetus having trisomy 21, 18, 13 or a sex chromosome condition, it does not mean that the fetus definitely has one of these conditions – although it is highly likely. For this reason, in the event of a 'high risk' (or positive) result, follow-up testing by an invasive procedure is recommended.

TDL Genetics

In the same way, if the test results show a 'low probability' of the fetus having trisomy 21, 18, 13 or a sex chromosome condition, it is unlikely that the fetus has one of these conditions. However, there is a very small risk that not all trisomic fetuses will be detected.

Who can have this test?

The Harmony test can be ordered by healthcare professionals for women with pregnancies of at least 10 weeks' gestational age. This test can be requested for any singleton or twin pregnancy, including those conceived naturally or by IVF using the patient's own egg or a donor egg. Note that, in twin pregnancies, sex chromosome (X and Y) analysis can determine fetal sex but not sex chromosome conditions. The Harmony test also does not assess risk for mosaicism, partial trisomies or translocations.

Results will be ready in approximately 3-5 days. Women still can have their 12-week scan for a detailed examination of the fetal anatomy, including measurement of nuchal translucency, nasal bone and other important factors. In this visit, patients can discuss the DNA and ultrasound results with their obstetricians. On the basis of the NIPT result and the ultrasound findings, a patient can decide whether or not she wants to have an invasive procedure (for example, CVS or amniocentesis).

Repeat samples

There needs to be enough fetal DNA in the maternal blood to be able to provide a result. If there is insufficient fetal DNA in the sample (which occurs in 3% of cases), another blood sample from the mother may be required. This will be processed in the laboratory at no extra charge.

What is the process?

Once the mother has taken an independent personal decision that she wants to have the NIPT performed, she will be asked to sign a consent form and her blood sample can be taken from a vein in her arm.



Who carries out the analysis of the test?

Her sample and completed request form need to be sent to TDL Genetics, where the Harmony test is performed on the DNA extracted from her blood sample.

Will the mother need to have any other tests?

The Harmony test does not provide information on mosaicism, partial trisomies or translocations, or other rare chromosomal abnormalities. If the ultrasound scan shows a high nuchal translucency or other major physical defects such as brain abnormalities, heart abnormalities, the risk for some rare chromosomal defects may be high. In such cases, the mother may choose to have a CVS or an amniocentesis.

The non-invasive prenatal test does not provide information on other physical defects such as spina bifida, or information on fetal growth. It is therefore advisable that the mother has all the usual ultrasound scans during her pregnancy.

Sample stability

Samples must be taken in special tubes provided by the laboratory. These samples must not be refrigerated, but stored at ambient temperature protected by the gel packs provided. The lab must receive the samples within 7 days to allow testing to proceed.

TEST	CODE	SAMPLE REQs	TAT
Non-Invasive Prenatal Testing – common aneuploidy screening from maternal blood	NIPT	Two 10ml tubes of maternal blood – special tubes provided by the laboratory	3-5 days
Non-Invasive Prenatal Testing – common aneuploidy screening from maternal blood including 22q11.2 del	NIPQ	Two 10ml tubes of maternal blood – special tubes provided by the laboratory	3-5 days

Always provide Clinical Details and Family History with requests for Genetic Tests.

Key: See page 21 for sample taking and special handling instructions.

22Q DELETION SCREENING

TDL Genetics will include 22q11.2 deletion, if requested as an additional option in the Harmony prenatal test menu. 22q11.2 deletion is the underlying cause of conditions described as DiGeorge syndrome and velocardiofacial syndrome (VCFS).

Why is 22q11.2 being included in the Harmony test (and not other microdeletion syndromes)?

- The 22q11.2 deletion has been carefully chosen as the only clinically relevant microdeletion syndrome to include with NIPT.
- 22q11.2 deletion is the most common chromosomal microdeletion, occurring in up to 1 in 1000 pregnancies.
- Other microdeletion syndromes have a much lower incidence and would increase the false positive rate of the test.

What is the performance of the 22q.11.2 addition?

- Inclusion of 22q11.2 deletion is aimed at a screening population, the test has been shown to identify 75% of pregnancies with a 22q11.2 deletion. Therefore, pregnancies with a known higher risk of 22q11.2 deletion, whether ascertained through ultrasound scan or family history should consider invasive diagnostic testing as this test will not identify 1 in 4 (25%) of cases.
- There is a false-positive rate of up to 0.5% associated with the 22q11.2 part of the Harmony test. This means that in 200 women with a pregnancy unaffected by 22q11.2 deletion 199 will receive a low probability result and 1 will receive a high probability result.

What is the benefit of finding out that a pregnancy has a high probability of a 22q11.2 deletion?


- Early screening and diagnosis of 22q11.2 deletions affects pregnancy management.
- Following confirmatory diagnosis of 22q.11.2 deletion the following may be recommended:
 - Level II ultrasound with fetal echocardiogram to evaluate for anomalies such as congenital heart defect and cleft palate.
 - Screening for and coordinated management of associated conditions.
 - Delivery at a tertiary care centre.

How do I request the 22q11.2 additional test option?

- Our updated request forms include the option of selecting 22q11.2 deletion. Tick this box if this is required.
- The 22q11.2 deletion cannot be requested in twin pregnancies or in pregnancies where the mother has a 22q11.2 duplication or deletion.
- There is an additional charge for 22q11.2 deletion.
- When discussing the informed consent for the Harmony test with your patient you must ensure they have read all the information on the reverse of the request form including the additional section headed 'What are the limitations of the Harmony prenatal test for 22q11.2?'

If 22q11.2 deletion is detected, we will undertake a confirmatory aCGH (microarray) on a CVS or Amnio, if undertaken, at no additional charge.

If you would like any further information about the 22q11.2 test please contact us at TDL Genetics by phone 020 7307 7409 or email harmony@tdlgenetics.com



harmony®

harmony®

THE RELIABILITY YOU WANT, AND THE ACCURACY YOU NEED.

harmony®

harmony®

Always provide Clinical Details and Family History with requests for Genetic Tests.

Key: See page 21 for sample taking and special handling instructions.

MALE GENETIC REPRODUCTIVE PROFILE

Chromosome Analysis
Y-Chromosome Microdeletions
Cystic Fibrosis Carrier Screen
(139 common mutations)
PolyT (5T,7T,9T) if clinically indicated

TAT
10-15
DAYS

GRP

A H⁹

THROMBOTIC RISK PROFILE

FBC
Coagulation Profile
Antithrombin III
Factor V Leiden
Common Mutation
Factor II Prothrombin
Common Mutation
MTHFR Common Variants
Lupus Anticoagulant
Protein C
Free Protein S Ag
Anticardiolipin Abs

TAT
5
DAYS

PROP

A A B C C C¹⁸

PRE-TRAVEL (DVT) SCREEN

FBC
Anticardiolipin Antibodies
Factor II Prothrombin Mutation
(G20210A)
Factor V Leiden Mutation
(G1691A)

TAT
5
DAYS

DVT1

A A B⁹

PAN-ETHNIC CARRIER SCREEN

2000+ Common Mutations
across 250+ Diseases*

includes 20+ X-linked Diseases
and 60+ Jewish Panel Diseases

TAT
4
WEEKS

GENE

A⁹

JEWISH CARRIER SCREEN

60+ Jewish Panel Diseases*

uses the same technology as the
Pan-Ethnic Carrier Screen, but
filters results to only report on
mutations commonly seen in
the Jewish Population

TAT
4
WEEKS

ASHJ

A⁹

* Disease list available from
the Laboratory

IRON OVERLOAD PROFILE

Iron
Total Iron Binding Capacity
Ferritin
Haemochromatosis
C282Y, H63D

TAT
3
DAYS

IOP

A A B⁹

RECURRENT MISCARRIAGE PROFILE (FEMALE)

FBC
Coagulation Profile
Antithrombin III
Factor V Leiden
Common Mutation
Factor II Prothrombin
Common Mutation
MTHFR Common Variants
Fibrinogen
Lupus Anticoagulant
Protein C
Free Protein S Ag
Anticardiolipin Abs
Chromosome Analysis

Please request Partner's
Chromosome Analysis using
a separate request form.

TAT
10-15
DAYS

RMP

A A B C C C H^{9,18}

PRENATAL DIAGNOSIS (BOBS + CULTURE)

Rapid Aneuploidy Diagnosis for
all Chromosomes + Common
Microdeletion Syndromes
by BOBs Analysis

TAT
3-5
DAYS

Chromosome Analysis
(Karyotype)

TAT
15
DAYS

ABK or CBK

AF/CVS⁹

PRODUCTS OF CONCEPTION (BOBS + CULTURE)

Rapid Aneuploidy Diagnosis for
all Chromosomes
by BOBs Analysis

TAT
3-5
DAYS

Chromosome Analysis
(Karyotype)

TAT
25
DAYS

PBK

Placental sample^{1,9}

In-vivo tests

These tests, ideally, must be carried out by appointment. Please telephone 020 7307 7383 for details, information for patient preparation, and appointment times. Sample taking fees for Extended tests are charged at £98.00 per visit.

EXTENDED TESTING

- 50g liquid glucose is consumed for the glucose challenge test/Mini-GTT.
- 75g liquid glucose is consumed for all other glucose tests.
- Each sample tube must be labelled with time of collection.

GLUCOSE TOLERANCE TESTS

TEST	CODE	SAMPLE REQs	COLLECTION TIME (MINUTES POST-GLUCOSE)	TAT
Glucose Challenge Test/Mini-GTT	RBGM	G	1 at 60 mins (50gm glucose)	1 day
Glucose Tolerance Test/OGTT	GTT	3x G 3x RU	1 each at 0, 60 and 120 mins (75gm glucose load)	1 day
Glucose Tolerance with Insulin	GTTI	3x B 3x G 3x RU	1 each at 0, 60 and 120 mins	1 day
Glucose Tolerance with Growth Hormone	GTT+GHDF	3x B ³⁵ 3x G 3x RU	1 each at 0, 60 and 120 mins	1 day
Glucose Tolerance Test (Short)	GTTS	2x G 2x RU	1 each at 0 and 120 mins	1 day
Glucose Tolerance Test (Extended)	GTTE	5x G 5x RU	1 each at 0, 30, 60, 90 and 120 mins	1 day
Glucose Tolerance Test (Extended Plus)	GTTX	7x G 7x RU	1 each at 0, 30, 60, 90, 120, 150 and 180 mins	1 day

EXTENDED TESTS

TEST	CODE	SAMPLE REQs	COLLECTION TIME (MINUTES POST-GLUCOSE)	TAT
Lactose Tolerance Test	LTT	By appointment only	Contact 020 7025 7997 (Phlebotomy)	1 day
Synacthen Stimulation Test	SYNA	By appointment only	Contact 020 7025 7997 (Phlebotomy)	1 day

ANTIBIOTIC ASSAYS

TEST	CODE	SAMPLE REQs	TAT
Amikacin Level (State dose)	AMIK	B ⁴	4 hours
Gentamicin Assay	GENT	B ⁴	4 hours
Metronidazole Level	METR	B ⁴	7 days
Teicoplanin Assay	TEIC	B	5 days
Tobramycin Assay (Provide Clinical Details)	TOBR	B	3 days
Vancomycin Hydrochloride	VANC	B	4 hours

Therapeutic drug assays

There are three different collection times for Therapeutic Drug Monitoring:

TROUGH LEVEL

Blood should be collected just before the next dose. Trough Levels check that the appropriate drug concentration is being maintained.

PEAK LEVELS

Sample collection time is dependent on specific drug type and method of administration. Peak levels check that the drug level is not in the toxic range.

SUSPECTED TOXICITY

Blood can be collected any time.
















All collections should have the following noted on the request form:

- Dosage schedule including the amount and frequency and time of the last dose
- Time of specimen collection
- Clinical status of patient (e.g. routine, suspected toxicity)
- Name(s) of other drugs being taken by the patient

TEST	CODE	SAMPLE REQ	TAT
Amitriptyline	AMTR	A ⁴	5 days
Anafranil (Clomipramine)	CHLO	A	7 days
Carbamazepine (Tegretol)	CARB	B	4 hours
Clobazam	CLOB	A	5 days
Clomipramine (Anafranil)	CHLO	A	7 days
Clonazepam	CLON	A	7 days
Diazepam (Valium)	DIAZ	A	7 days
Digoxin	DIGO	B	4 hours
Epanutin (Phenytoin)	PHEN	B	4 hours
Erythropoietin	ERY	B	4 days
Ethosuximide	ETHO	A	7 days
FK506 (Tacrolimus/Prograf)	FK5	A ⁴	1-2 days
Flecainide (Tambocor)	FLEC	A	5 days
Fluoxetine (Prozac)	PROZ	A ⁴	5 days
Gabapentin	GABA	B ⁴	5 days
Imipramine	IMIP	A ⁴	4 days
Lamotrigine	LAMO	B ⁴	5 days
Levetiracetam (Keppra)	LEVE	B ⁴	3 days
Lithium (take 12 hours after dose)	LITH	B	4 hours
Lorazepam	LORA	A ⁴	10 days
Methotrexate	METX	B	2 days
Mycophenolic Acid (Cellcept)	MYCP	A	5 days
Mysoline (Primidone)	PRIM	B ⁴	3 days
Olanzapine	OLAN	A ⁴	5 days
Paracetamol	PARA	B	4 hours
Phenobarbitone	PHB	B	4 hours
Phenytoin (Epanutin)	PHEN	B	4 hours
Primidone (Mysoline)	PRIM	B ⁴	3 days

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Therapeutic drug assays

TEST	CODE	SAMPLE REQ	TAT
Propanalol	PRO	 ⁴	7 days
Risperidone	RISP	 ⁴	7 days
Sinequan (Doxepin)	DOXE	 ⁴	10 days
Sirolimus	SIRO	 ⁴	3 days
Streptomycin Levels	STRM	 ⁴	5 days
Sulpiride	SULP	 ⁴	4 days
Tacrolimus/Prograf (FK506)	FK5	 ⁴	1-2 days
Tegretol (Carbamazepine)	CARB	 ⁴	4 hours
Temazepam	TEMA	 ⁴	4 days
Theophylline	THEO	 ⁴	4 hours
Topiramate (Topamax)	TOPI	 ⁴	4 days
Trimipramine	TRIM	 ⁴	5 days
Valium (Diazepam)	DIAZ	 ⁴	7 days
Valproic Acid (Epilim)	VALP	 ⁴	4 hours
Vigabatrin (Sabril)	VIGA	 ⁴	10 days

Allergy

Allergy, Asthma and Autoimmune diseases are increasing around the world, especially in industrialised countries and affect all ages. Since every country has their own dietary habits there are noteworthy differences in the allergens causing food allergy.



UK PROFILE

Total IgE plus:

Food Mix inc.

Cod, Cow's Milk, Egg White,
Soya Bean, Peanut, Wheat

Grass Mix inc.

Cocksfoot, Meadow Fescue,
Meadow, Rye, Timothy

Fish: Cod

Cat Dander
Cladosporium Herbarum
Dog Dander
House Dust Mite
Latex

TAT
2
DAYS



ALUK

B

MEDITERRANEAN PROFILE

Total IgE plus:

A. alternata
Cat Epithelium and Dander
Cow's Milk
Egg White
House Dust Mite
(Dermatophagoides
pteronysinus and
Dermatophagoides farinae)
Olive
Peanut
Rye-grass
Timothy Grass

TAT
2
DAYS



ALMD

B

MIDDLE EAST PROFILE

Total IgE plus:

Food Mix inc.

Cod, Cow's Milk, Egg White,
Soya Bean, Peanut, Wheat

Fish: Cod

Dust Mix inc.

House Dust Mite,
Dermatophagoides
pteronysinus,
Dermatophagoides farinae,
Blatella germanica

TAT
2
DAYS



ALME

B

Allergy

TEST	CODE	SAMPLE REQs	TAT
Allergy – Individual Allergens See list on page 137	ALLE	B	2 days
Total IgE	IGE	B	1 day
Allergy Profile (Mediterranean)	ALMD	B	2 days
Allergy Profile (Middle East)	ALME	B	2 days
Allergy Profile (UK)	ALUK	B	2 days
Allergy Profile 1 (Food & Inhalants)	1A	B B	2 days
Allergy Profile 2 (Inhalants)	2A	B	2 days
Allergy Profile 3 (Food)	3A	B	2 days
Allergy Profile 4 (Nuts & Seeds)	4A	B	2 days
Allergy Profile 5 (Children's Panel)	5A	B	2 days
Allergy Profile 6 (Shellfish)	6A	B	2 days
Allergy Profile 7 (Finfish)	7A	B	2 days
Allergy Profile 8 (Cereal – singles)	8A	B	2 days
Allergy Profile 9 (Antibiotics)	9A	B	2 days
Allergy Profile 10 (Insects)	10A	B	2 days
Allergy Profile 11 (Combined Shellfish/Finfish)	11A	B	2 days
Allergy Profile 12 (Milk & Milk Proteins)	12A	B	2 days
Allergy Profile 13 (Stone fruit/Rosaceae family)	13A	B	2 days
Eczema Provoking Profile	ALEC	B	2 days
Gluten Allergy Profile	GLUT	A B B	10 days
Rhinitis Provoking Profile	ALRN	B	2 days
Tryptase	STRY	B	2 days
Allergen Component Profiles See page 141			
Histamine Releasing Urticaria Test	CURT	B	10-14 days
ISAC Panel	ISAC	B	3 days
Prealbumin	PALB	B	3 days

ECZEMA PROVOKING PROFILE (9 Allergens)

Total IgE with individual IgE allergens for:

- Milk
- Peanut
- Soya Bean
- Wheat
- Cat Dander
- Egg White
- Egg Yolk
- Fish Mix
- Hazelnut
- House Dust Mite

TAT
2
DAYS

ALEC

B

RHINITIS PROVOKING PROFILE (10 Allergens)

Total IgE with individual IgE allergens for:

- Milk
- Nettle
- Peanut
- Timothy Grass
- Birch
- Cat Dander
- Dog Dander
- Egg White
- Egg Yolk
- House Dust Mite

TAT
2
DAYS

ALRN

B

GLUTEN ALLERGY PROFILE

Gluten single IgE Allergen
Endomysial Antibodies IgA
Deamidated Gliadin IgG
Antibodies
Tissue Transglutaminase IgA
HLA DQ2/DQ8
Total IgA

TAT
10
DAYS

GLUT

A B B

Allergy

IgE ALLERGY PROFILE 1 (Food and inhalants)

Total IgE with individual
IgE allergens for:

Grass Mix, inc.

Cocksfoot
Meadow Fescue
Meadow
Rye
Timothy

Weed Mix, inc.

Common Ragweed
Giant Ragweed
Western Ragweed

Dust Mix, inc.

Blatella germanica
Dermatophagoides
pteronyssinus
Dermatophagoides
farinae
Hollister-Stier Labs

Mould Mix, inc.

A. alternata
Aspergillus fumigatus
Candida albicans
Cladosporium herbarum
Helminthosporium
halodes
Penicillium notatum

Tree Mix, inc.

Box Elder
Common Silverbirch
Hazel
Oak
London Plane
Maple
Sycamore

Single Allergens (19)

Beef
Bermuda Grass
Cat Dander
Clam
Common Silver Birch
Cow's Milk
Crab
Dog Dander
Egg White
Egg Yolk
Fish (Cod)
Hazel Nut
Horse Dander
Latex
Nettle
Peanut
Shrimp/Prawn
Soya Bean
Wheat

**TAT
2
DAYS**

1A

B B

IgE ALLERGY PROFILE 2 (Inhalants)

Total IgE with individual
IgE allergens for:

Alternaria
Aspergillus
Birch Pollen
Cat Dander
Cladosporium

Common Ragweed
Derma farinae
Dog Dander
House Dust Mite
Horse Dander
Timothy Grass

**TAT
2
DAYS**

2A

B

IgE ALLERGY PROFILE 3 (Food)

Total IgE with individual
IgE allergens for:

Codfish
Cow's Milk
Egg White

Egg Yolk
Kiwi
Peanut
Sesame
Soya
Wheat

**TAT
2
DAYS**

3A

B

IgE ALLERGY PROFILE 4 (Nuts and Seeds)

Total IgE with individual
IgE allergens for:

Almond
Brazil Nut
Cashew
Hazel Nut
Macadamia Nut
Peanut

Pecan
Pine Nut
Pistachio
Poppy Seed
Pumpkin Seed
Sesame Seed
Sunflower Seed
Walnut

**TAT
2
DAYS**

4A

B

IgE ALLERGY PROFILE 5 (Children's Panel)

Total IgE with individual
IgE allergens for:

Cat Dander
Cow's Milk
Egg White
Egg Yolk

Mite, Pteronyssinus
Peanut
Soya Bean
Timothy Grass
Wheat Flour

**TAT
2
DAYS**

5A

B

IMMUNOCAP ISAC PANEL

Simultaneous measurement in a single test of specific
antibodies to more than one hundred allergen
components from more than 50 preselected
allergen sources.

**TAT
3
DAYS**

ISAC

B

Allergy

IgE ALLERGY PROFILE 6 (Shellfish)		
Total IgE with individual IgE allergens for:	Lobster	TAT 2 DAYS
	Octopus	
	Prawns/Shrimp	
	Scallop	
	Squid	
Clam		
Crab		
Crawfish/Crayfish		
6A		

B

IgE ALLERGY PROFILE 7 (Finfish)		
Total IgE with individual IgE allergens for:	Sardine/Pilchard	TAT 2 DAYS
	Salmon	
	Sole	
	Swordfish	
	Tuna	
Codfish		
Mackerel		
Plaice		
7A		

B

IgE ALLERGY PROFILE 8 (Cereal – singles)	
Total IgE with individual IgE allergens for: Barley Oat Rye Wheat	TAT 2 DAYS
	8A

B

IgE ALLERGY PROFILE 9 (Antibiotics)		
Total IgE with individual IgE allergens for:		TAT 2 DAYS
	Cefaclor	
	Pen G	
	Pen V	
9A		

B

IgE ALLERGY PROFILE 10 (Insects)		
Total IgE with individual IgE allergens for: Common Wasp – Yellow Jacket Bee	Paper Wasp	TAT 2 DAYS
	Yellow Hornet	
	White Faced Hornet	
10A		

B

IgE ALLERGY PROFILE 11 (Combined Shellfish/Finfish)		
Total IgE with individual IgE allergens for:	Salmon	TAT 2 DAYS
	Scallop	
	Squid	
	Tuna	
	Cod	
Prawn/Shrimp		
11A		

B

IgE ALLERGY PROFILE 12 (Milk & Milk Proteins)		
Total IgE with individual IgE allergens for:	Cow's Milk	TAT 2 DAYS
	Goat's Milk	
	Mare's Milk	
	Sheep's Milk	
	Whey (cow and ewe)	
Alpha-lactalbumin – milk proteins		
Beta-lactoglobulin – milk proteins		
Casein – milk proteins		
12A		

B

IgE ALLERGY PROFILE 13 (Stone Fruit, Rosaceae family)		
Total IgE with individual IgE allergens for:	Cherry	TAT 2 DAYS
	Peach	
	Pear	
	Plum	
	Raspberry	
Almond		
Apple		
Apricot		
13A		

B

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Allergy

Allergens, when requested individually are priced as single tests, sample 1 x **B** (up to 5 allergens).
Protein allergens are manufactured by Thermofisher (Phadia) and are IgE specific.

GRASS POLLENS

Bahia grass **g17**
Barley **g201**
Bermuda grass **g2**
Brome grass **g11**
Canary grass **g71**
Cocksfoot **g3**
Common reed **g7**
Cultivated oat **g14**
Cultivated rye **g12**
Cultivated wheat **g15**
Johnson grass **g10**
Maize, Corn **g202**
Meadow fescue **g4**
Meadow foxtail **g16**
Meadow grass,
 Kentucky blue **g8**
Redtop, Bentgrass **g9**
Rye-grass **g5**
Sweet vernal grass **g1**
Timothy grass **g6**
Velvet grass **g13**
Wild rye grass **g70**

WEED POLLENS

Alfalfa **w45**
Camomile **w206**
Careless weed **w82**
Cocklebur **w13**
Common pigweed **w14**
Common ragweed **w1**
Dandelion **w8**
Dog fennel **w46**
False ragweed **w4**
Firebush (Kochia) **w17**
Giant ragweed **w3**
Goldenrod **w12**
Goosefoot,
 Lamb's quarters **w10**
Japanese Hop **w22**
Lupin **w207**
Marguerite, Ox-eye daisy **w7**
Mugwort **w6**
Nettle **w20**
Parietaria officinalis **w19**
Parietaria judaica **w21**

Plantain (English), Ribwort **w9**
Rape **w203**
Rough marshelder **w16**
Saltwort (prickly),
 Russian thistle **w11**
Scale, Lenscale **w15**
Sheep sorrel **w18**
Sunflower **w204**
Wall pellitory **w19**
Wall pellitory **w21**
Western ragweed **w2**
Wormwood **w5**
Yellow dock **w23**

TREE POLLENS

Acacia **t19**
American beech **t5**
Australian pine **t73**
Bald cypress **t37**
Bayberry **t56**
Box-elder **t1**
Cedar **t212**
Cedar elm **t45**
Chestnut **t206**
Common silver birch **t3**
Cottonwood **t14**
Cypress **t222**
Date **t214**
Douglas fir **t207**
Elder **t205**
Elm **t8**
Eucalyptus, Gum-tree **t18**
European ash **t25**
Grey alder **t2**
Hackberry **t44**
Hazel **t4**
Horn beam **t209**
Horse chestnut **t203**
Italian/Mediterranean/
 Funeral cypress **t23**
Japanese cedar **t17**
Linden **t208**
Maple leaf sycamore,
 London plane **t11**
Melaleuca, Cajeput-tree **t21**
Mesquite **t20**

Mountain juniper **t6**
Mulberry **t70**
Oak **t7**
Oil Palm **t223**
Olive **t9**
Paloverde **t219**
Pecan, Hickory **t22**
Peppertree **t217**
Pine **t213**
Privet **t210**
Queen palm **t72**
Red cedar **t57**
Red mulberry **t71**
Scotch broom **t55**
Spruce **t201**
Sweet gum **t211**
Walnut **t10**
White ash **t15**
White hickory **t41**
White pine **t16**
Willow **t12**
Virginia live oak **t218**

MICROORGANISMS

Acremonium kiliense **m202**
Alternaria alternata **m6**
Aspergillus flavus **m228**
Aspergillus fumigatus **m3**
Aspergillus niger **m207**
Aspergillus terreus **m36**
Aureobasidium pullulans **m12**
Botrytis cinerea **m7**
Candida albicans **m5**
Chaetomium globosum **m208**
Cladosporium herbarum **m2**
Curvularia lunata **m16**
Epicoccum purpurascens **m14**
Fusarium proliferatum
 (*F. moniliforme*) **m9**
Setomelanomma rostrata
 (*Helminthosporium halodes*) **m8**
Malassezia spp. **m227**
Mucor racemosus **m4**
Penicillium chrysogenum
 (*P. notatum*) **m1**
Penicillium glabrum **m209**

Allergy

Phoma betae m13
Rhizopus nigricans m11
Staphylococcal enterotoxin A m80
Staphylococcal enterotoxin B m81
Staphylococcal enterotoxin C m223
Staphylococcal enterotoxin TSST m226
Stemphylium herbarum
(*S. botryosum*) m10
Tilletia tritici m201
Trichoderma viride m15
Trichophyton mentagrophytes var.
goetzii m210
Trichophyton mentagrophytes var.
interdigitale m211
Trichophyton rubrum m205
Ulocladium chartarum m204

EPIDERMALS AND

ANIMAL PROTEINS

Budgerigar droppings e77
Budgerigar feathers e78
Camel dander u328
Canary bird droppings e200
Canary bird feathers e201
Cat dander e1
Chicken droppings e218
Chicken feathers e85
Chicken, serum proteins e219
Chinchilla epithelium e208
Cow dander e4
Deer epithelium e216
Dog dander e5
Duck feathers e86
Ferret epithelium e217
Finch feathers e214
Fox epithelium e210
Gerbil epithelium e209
Goat epithelium e80
Goose feathers e70
Guinea pig epithelium e6
Hamster epithelium e84
Horse dander e3
Mink epithelium e203
Mouse epithelium e71
Mouse epithelium,
serum proteins and urine
proteins e88
Mouse serum proteins e76

Mouse urine proteins e72
Parakeet droppings e197
Parakeet serum e198
Parrot feathers e213
Pigeon droppings e7
Pigeon feathers e215
Rabbit epithelium e82
Rabbit, serum proteins e206
Rabbit, urine proteins e211
Rat epithelium e73
Rat epithelium, serum proteins
and urine proteins e87
Rat serum proteins e75
Rat urine proteins e74
Reindeer epithelium e202
Sheep epithelium e81
Swine epithelium e83
Turkey feathers e89

MITES

Acarus siro (Storage mite) d70
Blomia tropicalis
(House dust mite) d201
Dermatophagoides farinae
(House dust mite) d2
Dermatophagoides microceras
(House dust mite) d3
Dermatophagoides pteronyssinus
(House dust mite) d1
Euroglyphus maynei
(House dust mite) d74
Glycyphagus domesticus
(Storage mite) d73
Lepidoglyphus destructor
(Storage mite) d71
Tyrophagus putrescentiae
(Storage mite) d72

ALLERGEN COMPONENTS

See page 141 for Component
Testing and Component
Allergen Profiles

HOUSE DUST

Greer Labs., Inc. h1
Hollister-Stier Labs. h2

INSECTS

Berlin beetle i76
Blood worm i73
Cockroach, American i206
Cockroach, German i6
Cockroach, Oriental i207
Fire ant i70
Grain weevil i202
Green nimitti i72
Horse fly i204
Mediterranean flour moth i203
Mosquito i71
Moth i8

VENOMS

Bumblebee i205
Common wasp (Yellow jacket i3
European Paper Wasp i77
European hornet i75
Honey bee i1
Paper wasp i4
White-faced hornet i2
Yellow hornet i5

DRUGS

Amoxicilloyl c6
Ampicilloyl c5
Cefaclor c7
Chlorhexidine c8
Gelatin bovine c74
Insulin human c73
Penicilloyl G c1
Penicilloyl V c2
Pholcodine c261
Morphine c260
Suxamethonium
(succinylcholine) c202

OCCUPATIONAL

Bougainvillea k214
Cotton seed k83
Ethylene oxide k78
Ficus k81
Formaldehyde/Formalin k80
Green coffee bean k70
Hexahydrophthalic anhydrid k209
Isocyanate HDI (Hexamethylene
diisocyanate) k77
Isocyanate MDI (Diphenylmethane
diisocyanate) k76

Allergy

Iso cyanate TDI (Toluene diisocyanate) **k75**

Ispaghula **k72**

Latex **k82**

Methyltetrahydroptalic anhydrid **k211**

Phthalic anhydride **k79**

Silk **k74**

Silk waste **k73**

Sunflower seed **k84**

Trimellitic anhydride, TMA **k86**

PARASITES

Anisakis **p4**

Ascaris **p1**

Echinococcus **p2**

MISCELLANEOUS

Cotton, crude fibers **o1**

Mealworm **o211**

MUXF3 CCD, Bromelain **o214**

Seminal fluid **o70**

Streptavidin **o212**

FOODS – FRUITS & VEGETABLES

Apple **f49**

Apricot **f237**

Asparagus **f261**

Aubergine, eggplant **f262**

Avocado **f96**

Bamboo shoot **f51**

Banana **f92**

Beetroot **f319**

Blackberry **f211**

Blueberry **f288**

Broccoli **f260**

Brussel sprouts **f217**

Cabbage **f216**

Carrot **f31**

Cauliflower **f291**

Celery **f85**

Cherry **f242**

Cucumber **f244**

Date **f289**

Fennel, fresh **f276**

Fig **f328**

Garlic **f47**

Grape **f259**

Grapefruit **f209**

Guava **f292**

Jack fruit **f318**

Jujube **f336**

Kiwi **f84**

Lemon **f208**

Lettuce **f215**

Lime **f306**

Mandarin (tangerine, clementine, satsumas) **f302**

Mango **f91**

Melon **f87**

Olive (black, fresh) **f342**

Onion **f48**

Orange **f33**

Papaya **f293**

Passion fruit **f294**

Peach **f95**

Pear **f94**

Persimon (kaki fruit, sharon) **f301**

Pineapple **f210**

Plum **f255**

Potato **f35**

Pumpkin **f225**

Raspberry **f343**

Red currant **f322**

Spinach **f214**

Strawberry **f44**

Sweet potato **f54**

Tomato **f25**

Watermelon **f329**

FOODS – SEED, LEGUMES & NUTS

Almond **f20**

Barley **f6**

Blue vetch **f310**

Brazil nut **f18**

Buckwheat **f11**

Cashew nut **f202**

Chick pea **f309**

Coconut **f36**

Common millet **f55**

Fenugreek **f305**

Foxtail millet **f56**

Gluten **f79**

Green bean **f315**

Hazel nut **f17**

Lentil **f235**

Lima bean **f182**

Linseed **f333**

Lupin seed **f335**

Macadamia nut **f345**

Maize, Corn **f8**

Oat **f7**

Pea **f12**

Peanut **f13**

Pecan nut **f201**

Pine nut, pignoles **f253**

Pistachio **f203**

Poppy seed **f224**

Pumpkin seed **f226**

Quinoa **f347**

Rape seed **f316**

Red kidney bean **f287**

Rice **f9**

Rye **f5**

Sesame seed **f10**

Soybean **f14**

Spelt wheat **f124**

Sugar-beet seed **f227**

Sweet chestnut **f299**

Walnut **f256**

Wheat **f4**

White bean **f15**

FOODS – SPICES

Allspice **f339**

Anise **f271**

Basil **f269**

Bay leaf **f278**

Black pepper **f280**

Caraway **f265**

Cardamon **f267**

Chilipepper **f279**

Clove **f268**

Coriander **f317**

Curry (Santa Maria) **f281**

Dill **f277**

Ginger **f270**

Green pepper (unripe seed) **f263**

Lovage **f275**

Mace **f266**

Marjoram **f274**

Mint **f332**

Mustard **f89**

Oregano **f283**

Paprika, Sweet pepper **f218**

Parsley **f86**

Tarragon **f272**

Thyme **f273**

Vanilla **f234**

Allergy

FOODS – FISH, SHELLFISH & MOLLUSCS

Abalone f346
Anchovy f313
Blue mussel f37
Cat fish f369
Chub mackerel f50
Clam f207
Crab f23
Crayfish f320
Eel f264
Fish (cod) f3
Grouper f410
Gulf flounder f147
Haddock f42
Hake f307
Halibut f303
Herring f205
Jack mackerel, Scad f60
Langust (spiny lobster) f304
Lobster f80
Mackerel f206
Megrim f311
Octopus f59
Orange roughy f412
Oyster f290
Pacific squid f58
Plaice f254
Pollock f413
Red snapper f381
Salmon f41
Sardine (Pilchard) f308
Sardine, Japanese Pilchard f61
Scallop f338
Shrimp f24
Snail f314
Sole f337
Squid f258
Swordfish f312
Tilapia f414
Trout f204
Tuna f40
Walleye pike f415
Whitefish (Inconnu) f384

FOODS – EGG & FOWL

Chicken f83
Egg f245
Egg white f1
Egg yolk f75
Turkey meat f284

FOODS – MEAT

Beef f27
Elk/moose meat f285
Mutton f88
Pork f26
Rabbit f213

FOODS – MILK

Cheese, cheddar type f81
Cheese, mold type f82
Cow's whey f236
Goat milk f300
Mare's milk f286
Milk f2
Milk, boiled f231
Sheep milk f325
Sheep whey f326

FOODS – ADDITIVES

Carob (E410) f296
Guar, guar gum (E412) f246
Gum arabic (E414) f297
Tragacanth (E413) f298
Cochineal extract
(Carmine red) (E120) f340

FOODS – MISCELLANEOUS

Cacao f93
Coffee f221
Honey f247
Hop (fruit cone) f324
Malt f90
Mushroom (champignon) f212
Tea f222
Yeast f45

Allergy

COMPONENT TESTING

Using ImmunoCAP Allergen Components can help refine the understanding of sensitisation, by assessing a person's sensitisation pattern at the molecular level. When used in conjunction with traditional extract-based IgE testing, these provide information at the individual component level.

For more information, please contact the Immunology Department on 020 7025 7917.

TEST	CODE	SAMPLE REQ	TAT
Alpha Gal Components (related to red meat)	ZZ37	B	2 days
Alternaria Components	ZZ1	B	2 days
Apple Components	ZZ36	B	2 days
Aspergillus Components	ZZ2	B	2 days
Birch Components	ZZ3	B	2 days
Brazil Components	ZZ4	B	2 days
Cashew Components	ZZ35	B	2 days
Cat Components	ZZ5	B	2 days
Celery Components	ZZ6	B	2 days
Cow's Milk Components	ZZ7	B	2 days
Dog Components	ZZ8	B	2 days
Egg Components	ZZ9	B	2 days
Fish Components	ZZ10	B	2 days
Hazelnut Components	ZZ11	B	2 days
House Dust Mite Components	ZZ12	B	2 days
Kiwi Components	ZZ32	B	2 days
Latex Components	ZZ13	B	2 days
Olive Components	ZZ14	B	2 days
Peach Components	ZZ15	B	2 days
Peanut Components	ZZ16	B	2 days
Shrimp Components	ZZ17	B	2 days
Soybean Components	ZZ18	B	2 days
Timothy Grass Components	ZZ19	B	2 days
Venom Components	ZZ33	B	2 days
Wall Pellitory Components	ZZ20	B	2 days
Walnut Components	ZZ34	B	2 days
Wheat Components	ZZ21	B	2 days
PR-10 Proteins	ZZ22	B	2 days
Lipid Transfer Proteins	ZZ23	B	2 days
Profilins	ZZ24	B	2 days
Polcalcins	ZZ25	B	2 days
Seed Storage Proteins	ZZ26	B	2 days
Glycan Determinants	ZZ27	B	2 days
Lipocalins	ZZ28	B	2 days
Parvalbumins	ZZ29	B	2 days
Serum Albumins	ZZ30	B	2 days
Tropomyosins	ZZ31	B	2 days

* Please quote the ZZ Code when requesting Allergen Component Profiles.





































Key: See page 21 for sample taking and special handling instructions.

Vitamins, Nutrition and Lifestyle























VITAMIN B PROFILE Vitamin B1 Vitamin B2 Vitamin B3 Vitamin B6 Vitamin B9 (red cell) Vitamin B12 (Active) TAT 5 DAYS VBP	VITAMIN PROFILE 1 Vitamin A Beta Carotene Vitamin B1 Vitamin B2 Vitamin B6 Vitamin C (Frozen) Vitamin E TAT 5 DAYS VITS	MINERAL SCREEN Calcium Magnesium Zinc Iron Copper Chromium Manganese TAT 5 DAYS MINE
A A B	A B B ⁷	B K
SPORTS/PERFORMANCE PROFILE FBC/ESR Biochemistry Profile HDL/LDL Ferritin C-Reactive Protein Omega 3/Omega 6 Mineral Screen Vitamin B9 (Red Cell Folate) Vitamin B12 (Active) TAT 5 DAYS SPOR	VITAMIN PROFILE 2 Vitamin A Beta Carotene Vitamin B1 Vitamin B2 Vitamin B3 Vitamin B6 Vitamin B9 (Red Cell Folate) Vitamin B12 (Active) Vitamin C (Frozen) Vitamin D (25-OH) Vitamin E TAT 5 DAYS VIT2	MINERAL SCREEN – WHOLE BLOOD Whole Blood Potassium Whole Blood Magnesium Whole Blood Calcium Whole Blood Manganese Whole Blood Zinc Whole Blood Copper Whole Blood Selenium Whole Blood Chromium TAT 5 DAYS RMIN
A A A B B B B G K ⁴	A A A B B ^{7,13}	H H

Patients taking supplements may be advised to stop medication prior to testing.














Vitamins, Nutrition and Lifestyle

TEST	CODE	SAMPLE REQ	TAT
Ceruloplasmin	CERU		1 day
Copper (Serum)	COPP		5 days
Essential Fatty Acid Profile (Red Cell)	EFAR	 ⁴	10 days
Folate (Red Cell)	RBCF		2 days
Glutathione (Red Cell)	GLUR	 ⁵	5 days
Glutathione Peroxidase	GLPX		5 days
Lutein	LUTE	 ¹³	2 weeks
Lycopene	LYCO		2 weeks
Magnesium (Whole blood)	RCMG	 or 	4 days
Mineral Screen	MINE	 	5 days
Mineral Screen (Whole blood)	RMIN	 	5 days
Mineral Screen and Industrial Heavy Metal Screen (Trace Metals)	TRAC	   	7-10 days
Omega 3/Omega 6 (see page 145)	OMG3	 ⁴	4 days
Selenium (Serum)	SELE		4 days
Selenium (Whole Blood)	SELR	 or 	4 days
Sports/Performance Profile	SPOR	         ⁴	5 days
Xylose Tolerance Test	XTT	 ¹	7 days
Zinc (Serum/Plasma)	ZINC		1 day
Zinc (Urine)	URZN		5 days
Zinc (Whole Blood)	RBCZ	 or 	5 days

This provides valuable diagnostic information, which can be assimilated with other diagnostic markers in the assessment of nutritional status, and compares favourably to semi-quantitative functional assays. For fertility and lifestyle refer to page 56.

TEST	CODE	SAMPLE REQ	TAT
1,25 Vitamin D	D3		5-8 days
Beta Carotene	CARO		5 days
Biotin	BIOS		1 week
Carotenes	CARO	 ¹³	5 days
Vitamin A (Retinol)	VITA		5 days
Vitamin B (Functional)	FUNC	  or  ¹³	5 days
Vitamin B Profile	VBP	  	5 days
Vitamin B1 (Thiamine)	VIT1		5 days
Vitamin B2 (Riboflavin)	VIB2		5 days
Vitamin B3 (Nicotinamide)	VIB3		5 days
Vitamin B5 (Pantothenic Acid)	VB5S		5 days
Vitamin B6 (Pyridoxine)	VITB		5 days
Vitamin B8 (Biotin)	BIOS		5 days
Vitamin B9 (Folic acid) – Red cell	RBCF		2 days
Vitamin B9 (Folic acid) – Serum	FOLA		1 day
Vitamin B12 (Active)	B12		1 day
Vitamin B12 (Active)/Red Cell Folate	B12F	 	2 days

Vitamins, Nutrition and Lifestyle

TEST	CODE	SAMPLE REQS	TAT
Vitamin C (Active)	VITC	 (Frozen) ⁷	5 days
Vitamin D (1, 25 Dihydroxy)	D3		5-8 days
Vitamin D (25-OH)	VITD		4 hours
Vitamin E (Alpha Tocopherol)	VITE		5 days
Vitamin K (Nutritional)	VKN	 ¹³	5 days
Vitamin Profile 1	VITS	   ⁷	5 days
Vitamin Profile 2	VIT2	     ^{7,13}	5 days

Omega3/6

Essential Red Cell Fatty Acids Omega-3/Omega-6

Omega-3 is the name given to a family of polyunsaturated fatty acids, which the body needs but cannot manufacture itself. Omega-3 fats are used as the building blocks for fat derived hormones such as prostaglandins and leukotrienes. The hormones with an Omega-3 base tend to reduce inflammation, while those that have an Omega-6 base increase inflammation. In the cell membrane the competition between these two essential fats has a direct bearing on the type of local hormone produced and the level of inflammation in the cell.

The Omega-6 to Omega-3 ratio in the cell membranes is key to the development of inflammatory disorders such as rheumatoid arthritis and heart disease. Diets low in oily fish and high in grains will promote inflammation and affect good health. The ratio of Omega-6 to Omega-3 in the West is around 15 to 1, fifteen times more Omega-6 on the cell membrane promoting inflammation. Having twice as much Omega-6 is considered by most experts to be the optimal amount but a ratio of 2:1 is not easy to produce by diet alone. Many people are aware of the health benefits of Omega-3 but the supplementation to achieve optimal health is erratic. Being able to test for Essential Red Cell Fatty Acids (Omega-6/Omega-3 ratio) identifies a person's current status and is sufficiently specific to allow an accurate supplementation recommendation to be made.

Results show the Omega Ratio with a clear recommendation for the required level of Omega Supplementation (if any) to achieve optimal levels.

Results show the ratio of Omega 3 to Omega 6, against an optimal ratio and provide a supplementation recommendation to achieve this optimal ratio.

TEST	CODE	SAMPLE REQS	TAT
Omega 3/Omega 6	OMG3	 ⁴	4 days

TDL Tinies™ & Self-collection samples

TDL TINIES™ (tinies@tdlpathology.com)

This list of tests covers some of the range that can be offered to patients for self-collection, using TDL TINIES™ and Royal Mail postal packs. Orders for TDL TINIES™ (packs with instructions) can be made up by TDL, by arrangement, and sent individually to patients, or supplied directly to doctors or healthcare companies. This is not a patient self-referral service and it is not point of care testing. All testing is undertaken in the laboratory and results are always returned directly to the healthcare company or doctor, **not to the patient**.

TDL TINY™ samples can be combined with other self-collected samples types (urine, stool, swabs, HPV).

In the case of positive Sexual Health, results will be reported with the recommendation for a venous sample to undertake confirmatory sample.

The sample volume from one TINY sample, when filled to the upper fill line, is **600 microlitres**. These, on receipt in the laboratory, are centrifuged and provide a volume of 300 microlitres of serum/plasma (depending on the tube type used). Different tests require varying amounts of sample, and this, together with analyser dead volumes, means that although certain tests can be carried out from TINY tubes, many tests simply cannot be achieved from these smaller sample volumes.

TDL TINY™ microtainers are manufactured by BD Diagnostics. They are designed for samples collection from skin puncture. BD Microtainers come with a variety of additives for various tests, have visible fill lines, and are colour coded as for standard BD Vacutainer tubes. Tubes and Lancets are CE marked. TDL TINY™ packs are made up by TDL and contain everything needed for a patient to self-collect their blood sample.

Recommendation: most people are not experienced at self-collection of their own blood. Whilst it is certainly possible to do a number of tests from one TINY and it is possible to collect for two or three microtainers – the most successful outcomes are collected by patients who read the instructions given in each pack, and who collect enough sample for one microtainer. Instructions for sample collection are enclosed in each pack. A completed **request form** must be enclosed with the returned sample. Results will always be sent to the requesting doctor/healthcare organisation.

There is a TDL TINY™ video to assist patients with sample collection.

Visit <http://www.tdlpathology.com/test-information/test-service-updates/tdl-tinies>

This can be personalised with logo and details.

For information and packs, please contact Annette Wilkinson 020 7307 7343 or email tinies@tdlpathology.com.

Tests that can be self-collected using TDL TINIES™

HAEMATOLOGY		
TEST	CODE	SAMPLE REQ
Full Blood Count	FBC	A
HbA1c	GHB	A

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

TDL Tinies™ & Self-collection samples

BIOCHEMISTRY

TEST	CODE	SAMPLE REQ
Amylase	AMY	B
Calcium	CA	B
Calcium + Vitamin D	CALD	B
Carbohydrate Deficient Transferrin	CDT	B
C Reactive Protein	CRP	B
C Reactive Protein (High Sensitivity)	HCRP	B
Ferritin	FERR	B
HbA1c	GHB	A
Iron Status Profile (FE/TIBC/FERR)	ISP	B
Liver Function Tests	LFT	B
Lipid Profile	LIPP	B
Lp-PLA2 (PLAC) Test	PLA2	B
Uric Acid	UA	B
Vitamin B12 (Active)	B12	B
Vitamin D (25-OH)	VITD	B

ENDOCRINOLOGY

TEST	CODE	SAMPLE REQ
AFP	AFP	B
Antimullerian Hormone	AMH	B
Beta HCG (Quantitative)	QHCG	B
Cortisol	CORT	B
DHEA Sulphate	DHEA	B
Female Hormone (LH/FSH/PROL/OEST)	FIP	B
FSH	FSH	B
HRT Profile 1 (FSH/OEST/PROG)	HRT	B
Oestradiol	OEST	B
Progesterone	PROG	B
Prolactin	PROL	B
SHBG	SHBG	B
Testosterone	TEST	B
Thyroid Profile 1 (Free T4/TSH)	TF	B
Thyroid Profile 3 (Free T3/Free T4/TSH)	TF3	B

IMMUNOLOGY

TEST	CODE	SAMPLE REQ
Borrelia Antibodies (IgG/IgM)	BORR	B
Borrelia Antibodies (IgM)	BORM	B
Endomysial Antibodies IgA	AEAB	B
Gliadin Antibodies (IgG)	AGAB	B
H. pylori Antibodies (IgG)	HBPA	B
Tissue Transglutaminase IgA	TAA	B

TDL Tinies™ & Self-collection samples

VIROLOGY / SEXUAL HEALTH

TEST	CODE	SAMPLE REQ
COVID-19 Roche Total Antibody IgG/IgM (SARS-CoV-2)	TCOV	CE marked self-collection kit*
Hepatitis B Surface Antigen	THBA	B
Hepatitis B Immunity (IgG)	THBI	B
Hepatitis C Antibodies	THCV	B
HIV1&2 Abs/p24 Ag	THIV	B
HPV mRNA (All High Risk Subtypes)	HPVY	Self-collection kit
HPV Individually Typed High Risk DNA Subtypes	HPVZ	Self-collection kit
Syphilis IgG/IgM	TSYP	B

*See details below – CE marked self-collection kits for COVID must be used.

TUMOUR MARKERS

TEST	CODE	SAMPLE REQ
AFP	AFP	B
Beta HCG(Oncology)	HCGQ	B
CA 15-3	C153	B
CA 19-9	C199	B
CA 125	C125	B
CEA	CEA	B
HE4 + ROMA	HE4	B
Prostate Specific Antigen	PSPA	B

LIFESTYLE

TEST	CODE	SAMPLE REQ
Omega 3/Omega 6	OMG3	A
Vitamin B9 (Folic Acid) Red Cell	RBCF	A
Vitamin B9 (Folic Acid) Serum	FOLA	B
Vitamin B12 (Active)	B12	B
Vitamin D (25-OH)	VITD	B

COVID-19 (SARS-CoV-2) Roche Elecsys Anti-SARS-CoV-2 Total Antibody NEW

Roche Elecsys Anti-SARS-CoV-2 reports both IgG and IgM as a TOTAL antibody result. The Roche Antibody test is CE marked for **capillary** samples, and one of the PHE selected antibody tests.

Test Code: TCOV

Sample Type	SST/Serum B Capillary (>14 days after onset of symptoms)
Performance	Specificity 100%, Sensitivity 97.4%
Analysers	Roche e801
Turnaround time	24 hours from receipt of sample

Self-collection capillary samples must be taken using CE marked IVD for COVID Postal kits NEW

The kits include a Royal Mail Tracked 24 return label. Contact TCOV@tdlpathology.com for details.

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

TDL Tinies™ & Self-collection samples

STEP 1

TDL TINY™ SAMPLE COLLECTION



THE DOCTORS
LABORATORY

Sample collection instructions

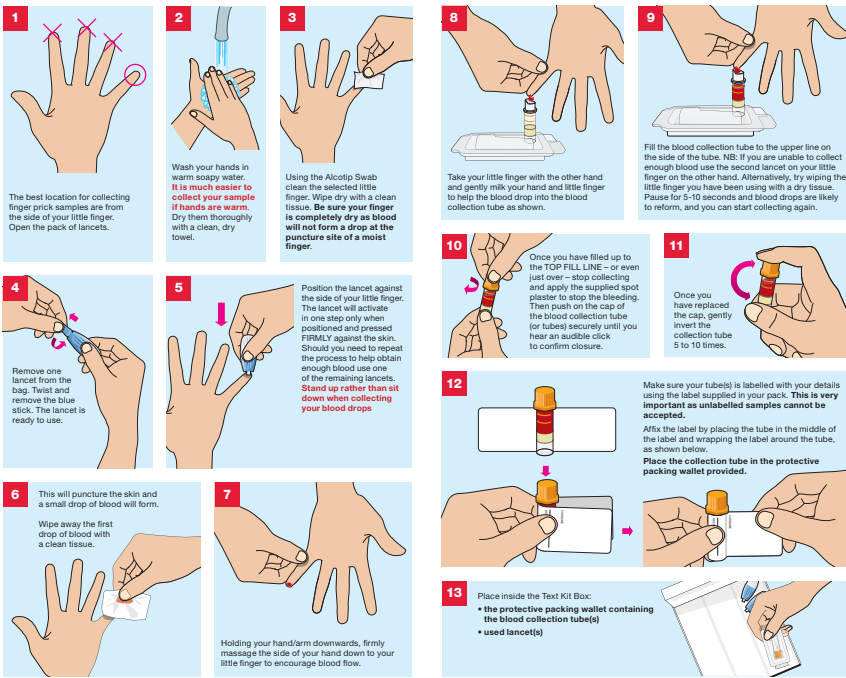
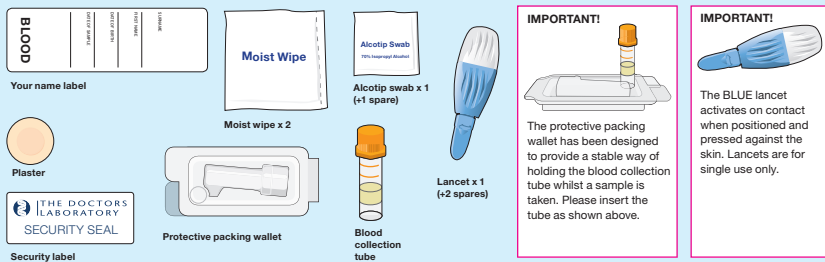
Please read these instructions first, slowly and carefully, the whole way through before attempting to collect your sample.

Clearly complete the Name Label using a ball point pen with:

• Your Surname • Your Date of Birth • Your First name • Date of Blood Collection

Do not affix the label to the blood collection tube until after collecting your sample. This is important as you will not be able to see how much blood you have collected if the label covers the sides of the tube. Sample self-collection is carried out at an individual's own risk.

Your sample collection pack contents: Step 1



TDL Tinies™ & Self-collection samples

Sample collection instructions

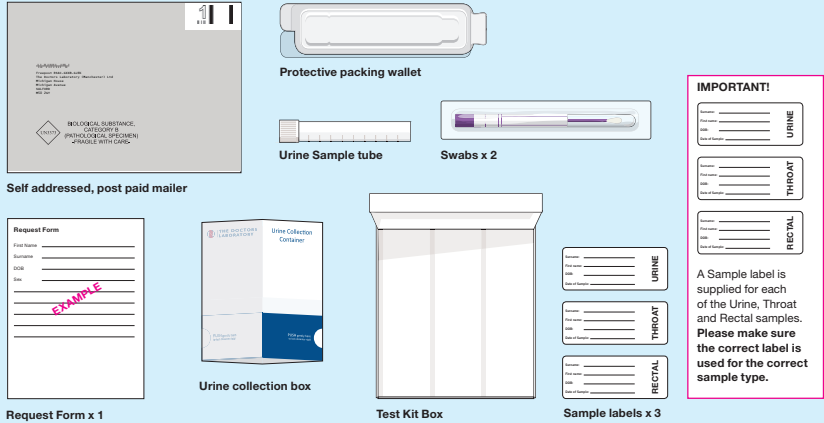
Please read these instructions first, slowly and carefully, the whole way through before attempting to collect your sample.

Clearly complete the Specimen bottle and Swab labels using a ball point pen with:

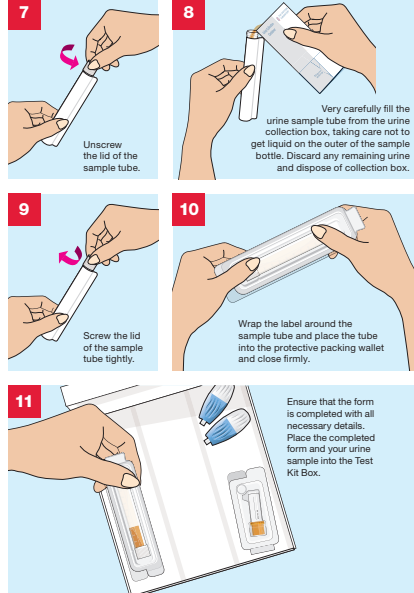
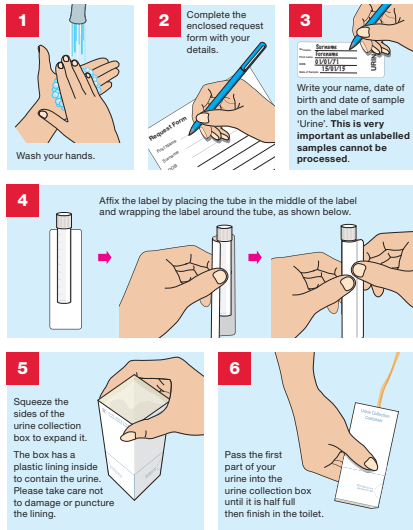
- Patient Surname
- Patient First name
- Patient Date of Birth

Sample self-collection is carried out at an individual's own risk.

Your sample collection pack contents: Steps 2-4



STEP 2 URINE SAMPLE COLLECTION



Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

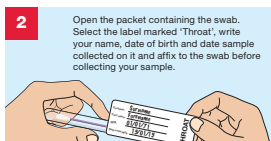
TDL Tinies™ & Self-collection samples

STEP 3

THROAT SWAB SAMPLE COLLECTION



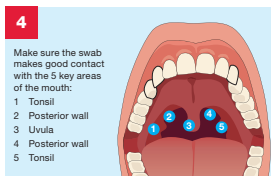
Wash your hands.



Open the packet containing the swab. Select the label marked 'Throat', write your name, date of birth and date sample collected on it and affix to the swab before collecting your sample.

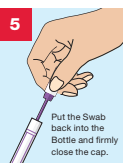


3



Make sure the swab makes good contact with the 5 key areas of the mouth:

- 1 Tonsil
- 2 Posterior wall
- 3 Uvula
- 4 Posterior wall
- 5 Tonsil



5



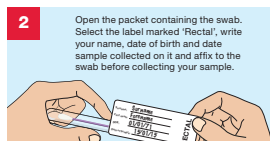
Place your sample into the Test Kit Box.

STEP 4

RECTAL SWAB SAMPLE COLLECTION



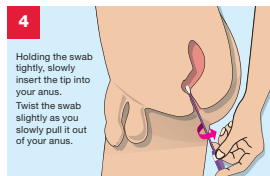
Wash your hands.



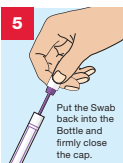
Open the packet containing the swab. Select the label marked 'Rectal', write your name, date of birth and date sample collected on it and affix to the swab before collecting your sample.



3



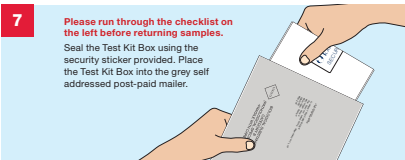
4



5



Place your sample into the Test Kit Box.



7

Please run through the checklist on the left before returning samples. Seal the Test Kit Box using the security sticker provided. Place the Test Kit Box into the grey self addressed post-paid mailer.

STEP 5

CHECKLIST

Before you return your samples please tick off the contents of the grey self addressed post-paid mailer.

- ☐ Completed Request form
- ☐ Blood collection tube(s) in the protective packing wallet
- ☐ Used lancets
- ☐ Urine sample in the outer transport bottle
- ☐ Throat swab in the swab bottle
- ☐ Rectal swab in its swab bottle
- ☐ Place all of the samples into the Test Kit Box and seal with the security sticker

You are now ready to seal the grey self addressed post-paid mailer. Please post your samples to **The Doctors Laboratory** as soon as possible from **ANY** Royal Mail post box in the UK. No stamp is required within the UK.

If you need assistance please contact **The Doctors Laboratory** on 020 7307 7373 or email samples@tdlpathology.com.

The Doctors Laboratory, The Halo Building, 1 Mabledon Place, London WC1H 9AX
Tel: 020 7307 7373 Fax: 020 7307 7374 E-mail: tdl@tdlpathology.com
Website: www.tdlpathology.com

© The Doctors Laboratory, 2019 TAP00002/13-05-19/V2

Screening for Drugs of Abuse/Alcohol

TEST	CODE	SAMPLE REQ	TAT
Alcohol Profile	AP	A B B G	5-7 days
Alcohol Profile 2	ALCP	A A B B G RU	5-7 days
Amphetamines – Blood	AMPB	B B	5 days
Cannabinoids (Urine) Screen	CANN	RU	1 day
Cocaine (Urine) Screen	UCOC	RU	1 day
Drugs of Abuse From Blood	DOAP	B	5 days
Drugs of Abuse Profile – Random Urine Sample/No Chain of Custody Plus Alcohol	DOA3	RU	2 days (5 days with LCMS/MS confirmation)
Drugs of Abuse Profile – Random Urine Sample/ No Chain of Custody	DOA	RU	2 days (5 days with LCMS/MS confirmation)
Drugs of Abuse Profile – With Chain of Custody	DOAL	RU/CoC Collection Containers ^{1,2}	2 days (5 days with LCMS/MS confirmation)
Drugs of Abuse Profile – Without Chain of Custody	DOAN	RU ²	2 days (5 days with LCMS/MS confirmation)
Ketamine Screen	KETA	RU	7-10 days
LSD	LSD	RU	5 days
Opiate Screen (Urine)	UOPI	RU	2 days
PEth (Phosphatidylethanol)	PETH	A ³⁸	5-7 days
Urine EtG (Ethyl glucuronide)	ETG	RU	1 week

Chain of custody refers to the system of controls governing the entire urine collection, processing and storage of sample to ensure that a particular urine specimen originated from a particular individual and that the reported results relate, beyond doubt, to that specimen. Chain of custody requires attention to detail so that it is possible to prove that there has been no opportunity for the sample to be accidentally or maliciously adulterated. Sample collection should be undertaken by collectors who are well versed in the protocols of chain of custody.

Samples submitted for analysis will undergo initial screening. Urinary creatinine is routinely measured during testing to verify the validity of the sample submitted. Creatinine levels below normal occur when the urine has been diluted, either directly or by drinking large amounts of water before providing the urine sample. Chain of custody containers, forms, seals and barcodes are provided by TDL on request. All Chain of Custody, and non-chain, samples with positive findings will proceed to identification/confirmation by Gas Chromatography/Mass Spectrometry.

Screening for Drugs of Abuse/Alcohol

DRUGS OF ABUSE SCREENING

DRUGS OF ABUSE PROFILE – WITH CHAIN OF CUSTODY			
Alcohol	LSD		
Amphetamines	MDMA		
Barbiturates	Methadone		
Benzodiazepine	Methaqualone		
Cannabinoids	Morphine – opiate		
Cocaine	Phencyclidine		
Codeine – opiate	Propoxyphene		
Dihydrocodeine – opiate			
Ketamine		TAT 2 DAYS	TAT 5 DAYS WITH LCMS/MS CONFIRMATION
DOAL			

RU/CoC collection containers^{1,2} * See page 153

DRUGS OF ABUSE PROFILE – WITHOUT CHAIN OF CUSTODY			
As above but with NO Chain of Custody			
		TAT 2 DAYS	TAT 5 DAYS WITH LCMS/MS CONFIRMATION
DOAN			

RU²

ALCOHOL PROFILE			
LFT	Alcohol Level		
CDT	MCV		
PEth			TAT 5-7 DAYS
AP			

A B B G

DRUGS OF ABUSE PROFILE – RANDOM URINE SAMPLE/NO CHAIN OF CUSTODY			
Amphetamines	MDMA		
Barbiturates	Methadone		
Benzodiazepine	Morphine – opiate		
Cannabinoids			
Cocaine			
Codeine – opiate			
Dihydrocodeine – opiate		TAT 2 DAYS	TAT 5 DAYS WITH LCMS/MS CONFIRMATION
DOA			
plus Alcohol DOA3			

RU






















DRUGS OF ABUSE FROM BLOOD – WITHOUT CHAIN OF CUSTODY			
Amphetamines	Opiates		
Barbiturates	Cocaine		
Tricyclic Antidepressants			
Benzodiazepine			
Cannabinoids			TAT 5 DAYS
DOAP			

B

ALCOHOL PROFILE 2			
LFT	Alcohol Level		
CDT	MCV		
PEth			TAT 5-7 DAYS
Urine Ethyl Glucuronaride (EtG)			
ALCP			

A A B B G RU

Occupational health

OCCUPATIONAL HEALTH – TRACE METALS IN BLOOD			
TEST	CODE	SAMPLE REQs	TAT
Aluminium	ALUM		7 days
Arsenic	ARS	 or 	5 days
Cadmium	CADM	 or 	5 days
Chromium	CHRO		5 days
Cobalt (Serum)	COBB		5 days
Copper (Serum)	COPP		5 days
Lead	LEAD		5 days
Lead Profile (Hb, ZPP, Lead)	LEAZ	 ¹³	3-5 days
Magnesium (Serum)	MG		4 hours
Manganese (Serum)	MANG		5 days
Mercury	MERC	 or 	5 days
Nickel	NICK		5 days
Silver	SILV		5 days
Trace Metal (Blood) Profile	TRAC	   	7-10 days
Zinc (Serum/Plasma)	ZINC		1 day

TRACE METAL (BLOOD) PROFILE						
Aluminium	Iron	Zinc	Copper	Mercury	Chromium	TAT 7-10 DAYS
Manganese	Calcium	Magnesium	Cadmium	Lead		
						TRAC

Occupational health

OCCUPATIONAL HEALTH – TRACE METALS IN URINE

TEST	CODE	SAMPLE REQ	TAT
Aluminium	ALUU	RU	1-2 weeks
Arsenic	ARSE	RU ³⁰	5 days
Cadmium	URCD	RU ³⁰	5 days
Chromium	URCR	RU ³⁰	10 days
Cobalt	COBA	RU ³⁰	5 days
Copper	URCU	CU	5 days
Lead	URPB	RU	5 days
Magnesium	URMG	PU	1 day
Mercury	URHG	RU ¹	5 days
Nickel	NICU	RU	10 days
Silver	USIL	RU	5 days
Zinc	URZN	CU	5 days

OCCUPATIONAL HEALTH – TESTS FOR SPECIFIC EXPOSURE

TEST	CODE	SAMPLE REQ	TAT
2-Butanone GC	BUTA	RU	7 days
2-Furoic Acid	2FA	RU	10 days
Acetone – Blood	ACTB	A or H	2 weeks
Acetone – Urine	ACTU	RU	5 days
Alcohol Profile	AP	A B B G	5-7 days
Alcohol Profile 2	ALCP	A A B B G RU	5-7 days
Benzene	BENZ	J ^{1,6}	3 days
Beta 2 Microglobulin (Serum)	B2MG	B	2 days
Beta 2 Microglobulin (Urine)	UB2M	RU	3 days
Bromide	BROM	B	3 days
Cholinesterase (Blood)	CHRC	H	5 days
Cholinesterase (Serum/Pseudo)	CHPS	B	4 hours
Cotinine (Saliva)	SCOT	Saliva Kit ¹	1-2 weeks
Doxepin Level (Sinequan)	DOXE	A	10 days
MBOCA in Urine	MBOC	RU	10 days
Molybdenum (Serum)	MOLY	B	5 days
Pethidine – Urine	UPET	RU	4 weeks
Thallium (Blood)	THAL	A / H	1 week
Thallium (Urine)	URTH	RU	1 week
Toluene (Blood)	TOL	J	10 days
Toluene (Urine)	UTOL	RU	10 days
Trichloroacetic Acid (Urine)	UTCA	RU	5 days
Xanthine – Blood	XANB	A	2 weeks
Xylene – Urine	UXYL	RU ³⁰	2 weeks
Zinc Protoporphyrin	ZNPR	A ¹³	5 days

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Cervical Screening

The Cytology Laboratory provides a rapid service for liquid based cervical samples. Urine cytology is performed in house while other non-gynaecological cytology samples are referred to a UKAS accredited laboratory for reporting.

Human papilloma virus (HPV), Chlamydia and Gonorrhoea testing is carried out routinely from ThinPrep vials and can be requested at the time the cervical sample is taken.

Laboratory hours

The laboratory department is open between 9.00am and 6.00pm.
Out of hours results available on 020 7307 7373.

Urgent samples

It is helpful if requests for urgent samples can be discussed with the Cytology Manager. Please telephone 020 7307 7323.

Use of service/Information required

Request forms must include **3 identifiers** (this can be patient's full name = 2, date of birth, hospital number or reference number) and need to accompany each sample.

Appropriate clinical information providing previous treatment/histological diagnosis is essential to ensure correct management recommendations can be given in the patient report. Tick boxes are provided to assist you.

The specimen container must be clearly labelled with patient details. Forms and samples which are mismatched will result in the sample being returned to the sender for correction and will delay the report turn around time.

Clinical advice

The Consultant Cytopathologists and the Advanced Practitioner work together to provide clinical and technical advice, including recommendations for follow-up, HPV testing and management of complex cases. To contact the department directly, please telephone 020 7307 7323.

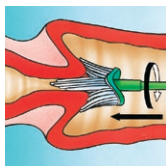
Cervical Screening



RECORD...

...the patient's 3 identifiers to include date of birth on the vial.

...the patient information and medical history on the cytology requisition form.



OBTAIN...

...an adequate sample from the cervix using a Cervex Brush (broom-like device). Insert the central bristles of the brush into the endocervical canal deep enough to allow the shorter bristles to fully contact the ectocervix. Push gently and rotate the brush in a clockwise direction five times.



RINSE...

...the Cervex Brush immediately into the PreservCyt Solution vial by pushing it into the bottom of the vial 10 times, forcing the bristles apart. As a final step, swirl the brush vigorously to further release material. Visually inspect the Cervex Brush to ensure that no material remains attached. Discard the brush.

**Do not leave the head of the Cervex Brush in the vial.
Check the vial is in date before use.**



TIGHTEN...

...the cap so that the black torque line on the cap passes the black torque line on the vial. Do not over-tighten.



PLACE...

...the vial and request form in a specimen bag for transportation to TDL.

Cervical Screening

ThinPrep® PAP Test Cervex Brush Protocol

PREPARE ALL EQUIPMENT BEFORE STARTING THE PROCEDURE

- Note expiry date on sample collection vial. Do not use expired vials.
- Ensure the entire plastic seal is removed from the lid of the vial and discarded.
- Complete patient details on both the request form and the vial.
Specimens may be returned or discarded if details are missing from the vial.
- Remove the lid from the vial before taking the sample.
- **Use of lubricant is NOT recommended.**

DO

- If excessive mucus is present, this should be gently removed before sampling.
- Use either the Cervex Brush (broom-like device) on its own or a Plastic spatula and endocervical brush combination.
- The Cervex Brush should be rotated 5 times in a clockwise direction. The Plastic spatula should be rotated through 360 degrees and the endocervical brush rotated through one quarter to one half turn.
- Immediately rinse the collected material into the vial.
- Replace the lid and tighten so that the black torque line on the cap passes the black torque line on the vial to avoid leakage.
- Keep the unlabelled portion of the sample vial free of labels so that the contents can be seen.
- If barcoded labels are used these must be applied horizontally around the vial.
- Samples should be sent to the laboratory without delay.

DON'T

- DO NOT leave the head of the Cervex Brush in the vial.
- DO NOT routinely clean the cervix or take a cervical swab before taking a cervical sample.
- An endocervical brush should never be used in isolation.
- DO NOT under any circumstances use a wooden spatula.
- DO NOT leave the collection device sitting in the vial whilst dealing with the patient.
- DO NOT over-tighten the lid on the vial.
- DO NOT place multiple labels on the outside of the vial.
- DO NOT apply barcoded labels vertically on the vial.
- DO NOT use expired vials.
- DO NOT delay the sending of vials to the laboratory. The sample needs to be processed within 3 weeks of collection.
- DO NOT use excessive lubricant – please AVOID if possible.

Cervical Screening

Gynaecological Samples

The Cytology department processes cervical samples directly referred from all sectors of practice – Health Screening, Occupational Health, GP's, Consultants, Colposcopy Units, Clinics, Hospitals and other Laboratories.

Liquid Based Cytology (LBC) is processed using the Hologic ThinPrep system.

The Doctors Laboratory uses the Hologic Imaging system as an enhanced Quality Control.

Information for Sample Takers is available by contacting the department. **Important: the head of the cervical broom must NOT be left in the vial.** The use of lubricant interferes with LBC sampling and may result in an inadequate sample. Use of lubricant is NOT recommended as it can affect the processing quality of the sample. Supplies of Thin prep vials are available from TDL.

STI Screening from Hologic Thin Prep Vial (HPV – see page 162)

Tests are priced individually. Please request tests individually. Thin Prep Vials are kept for 21 days after receipt of sample. Requests for additional tests from the vial already received in the laboratory can be made by contacting the Cytology Department.

Infection by PCR (singles)

TEST	CODE	SAMPLE REQS	TAT
Chlamydia trachomatis	TPCR	TPV	2 days
N. gonorrhoea	TGON	TPV	2 days
Chlamydia/Gonorrhoea	TCG	TPV	5 days
Mycoplasma genitalium	MGEN	TPV	2 days
Ureaplasma urealyticum	UGEN	TPV	2 days
Trichomonas vaginalis	TVPC	TPV	2 days
Gardnerella vaginalis	GVPC	TPV	2 days
Herpes Simplex I/II	HERD	TPV	5 days

7 STI PROFILE BY PCR FROM THIN PREP VIAL

Chlamydia trachomatis	All tests can be requested individually <i>*included if POSITIVE M.gen is detected from the same sample.</i>
N. gonorrhoea	
Mycoplasma genitalium	
Macrolide Resistance Test (M.gen)*	
Ureaplasma	
Trichomonas vaginalis	
Gardnerella vaginalis	
Herpes Simplex I/II	

TAT
**2
DAYS**

PP12

TPV

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Cervical Screening

Human papillomavirus (HPV) is a common virus transmitted through sexual contact. High Risk sub-types of HPV (HR-HPV) are linked to the development of abnormal cells and can cause cervical cancer. HPV is a necessary cause of invasive cervical cancer. Evidence shows HPV testing is a more effective way to identify women at risk of cervical cancer than by testing microscopically for abnormal cells from a PAP smear.

HR-HPV testing has been used in the UK since 2011 to identify women with low grade cytology abnormalities and as a follow up test of cure in women who have received treatment. In 2017 the UK NHSCSP recommended that **testing for HPV should replace cytology as the first (primary test) in cervical screening**. Primary HR-HPV testing has higher sensitivity for high grade CIN than primary cytology. HR-HPV testing also has a lower false negative rate than cytology. Primary HR-HPV testing was fully implemented in the UK during 2020. Sample taking remains unchanged: HR-HPV testing is carried out from Thin Prep samples. Cytology will be undertaken as a triage if HPV is DETECTED.

WHAT DOES THIS CHANGE MEAN?

It means that HPV testing is the **FIRST LINE TEST**. It will be carried out as a single test, with a single result reported as DETECTED/NOT DETECTED.

- If HR-HPV is **NEGATIVE (NOT DETECTED)** – this means no further testing is needed for your patient: she returns to Routine Recall
- If HR-HPV is **POSITIVE (DETECTED)** – this means that **CYTOLOGY** will be processed from the same Thin Prep Vial. **A further specimen is not required.**
- **If the CYTOLOGY result from this sample is HR-HPV NOT DETECTED** – the patient Recall will be determined by the screening history and will either be a repeat HR-HPV test in 12 months' time or, if HR-HPV remains persistent, a referral to colposcopy will be recommended.
- **If the CYTOLOGY result from this sample is ABNORMAL** the recommendation is to refer this patient for COLPOSCOPY.

<https://www.gov.uk/government/publications/cervical-screening-primary-hpv-screening-implementation/cervical-screening-implementation-guide-for-primary-hpv-screening>

Since 1st January 2019 all TDL requests for HPV have been processed as follows:

- **If HPV is requested as a single test, and the result is NEGATIVE/NOT DETECTED, cervical cytology (PAPT) will only be processed if specifically requested. The PAPT would be charged as an additional test.**
- **If HPV result is DETECTED, cervical cytology (PAPT) will be processed, even if not requested. The PAPT cervical sample will NOT be charged additionally.**
- **If cervical cytology (PAPT) is requested, HPV will always be processed with the PAPT. The PAPT will be charged.**

Cervical Screening

UNDERSTANDING THE SIGNIFICANCE OF HPV TESTING

The benefit of a negative HPV result is its negative predictive value – meaning that a negative HPV result indicates that a patient is at very low risk of developing cervical disease. The negative predictive value of both DNA and mRNA testing is the same. DNA tests detect presence of virus only. A mRNA test detects the presence of viral oncogenic expression.

Requests for Cervical Cytology (PAPT) only will no longer be processed without HPV. HPV testing will be charged.

Requests for PAPT

TEST	CODE	SAMPLE REQ	TAT
Cervical Cytology	PAPT will include HPVH	TPV	2-3 days

If PAPT is requested as a single test, HR-HPV will be undertaken additionally, and a combined report will be issued. **PAPT and HPVH will be charged.**

Requests for PAPT with selected HPV (HPVH or HP20 or HPV2)

TEST	CODE	SAMPLE REQ	TAT
PAPT and HPVH	PAPT + HPVH	TPV	2-3 days

If PAPT and HPVH are requested together, results will be given as a combined report, **PAPT and selected HPVH test will be charged.**

Requests for HPV as the PRIMARY TEST will reflex to PAPT if HPV is DETECTED/POSITIVE. PAPT will NOT be charged.

TEST	CODE	SAMPLE REQ	TAT
HPV mRNA (All High Risk Subtypes)	HPVH	TPV	2-3 days

If HPV is DETECTED/POSITIVE, cervical cytology (PAPT) will be processed **without charge**. The PAPT will be processed from the same vial.

Requests for HP20 as a single test

TEST	CODE	SAMPLE REQ	TAT
HPV Typed DNA	HP20	TPV/PCR Swab	2-3 days

HPV low and high risk DNA subtypes will be reported individually (5 low and 14 high risk).

If HPV is DETECTED/POSITIVE, cervical cytology (PAPT) will be processed **without charge**. The PAPT will be processed from the same vial.

Requests for HPV2 as a single test

TEST	CODE	SAMPLE REQ	TAT
HPV Typed DNA	HPV2	TPV	3 days

If one or more of DNA types 16, 18, 31, 33, 45 are DETECTED/POSITIVE, reflex testing for expression of E6/E7 oncoproteins will be undertaken and cervical cytology (PAPT) will be processed **without charge**. The PAPT will be processed from the same vial.

HPV/PAPT Combined Report

Where HPV result is reported with Cervical Cytology, a recommendation for patient management will be given, based on the combined findings.

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Self-collection HPV samples

TDL Self-Collection HPV Test

Human Papillomavirus (HPV) is the primary cause of nearly all cervical cancer. In most cases, the HPV virus is harmless and causes no symptoms. Most women who acquire HPV are able to clear the infection through their own immune systems. Persistent presence of high-risk types of HPV can cause cervical lesions which over time may develop into cancer if untreated. Testing for HPV determines the presence, or absence, of HPV and will determine whether the HPV type present is high risk for CIN and cervical cancer.

The **Self Collection HPV Test** provides women with the option to self-collect a vaginal specimen that is then sent to the laboratory for testing. There is well documented high level of concordance between the HPV DNA results from self-collected and clinician-collected specimens.

The **Self-Collection HPV Test** is validated, using a CE marked sample collection device for vaginal cell collection. This sample is then sent to the laboratory for processing for 14 high risk HPV DNA subtypes. A negative result means that these high-risk subtypes HPV were not detected and the patient is at extremely low risk of developing high-grade cervical disease/CIN2+ before their next routine visit.

A positive HPV result might indicate an increased risk of developing CIN/cervical cancer, and the report from the laboratory will provide a clear recommendation for follow-up/colposcopy.

The value of HPV DNA testing in cervical cancer screening and disease detection has been proven over and over again. Self-collection of specimens for HPV testing is not intended to replace existing patient management pathways but allows for:

- Those who wish to test following a change of sexual partner
- Option for identifying individual high risk DNA subtypes
- Personal preference to self-collect vaginal samples
- An acceptable option for women who avoid having regular cervical smears
- Self-collection for HPV increases acceptability and coverage rate of cervical cancer prevention

Results will always be sent to the requesting clinician, clinic or healthcare organisation.

HPVY Self-Collected HPV DNA with reporting of high risk subtypes (16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, 66, 68).

HPVZ Self-Collected HPV DNA with **individual** reporting of all subtypes 16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, 66, 68.

For more information, or to order Self-Collection HPV Test Packs, please contact Annette Wilkinson on 020 7307 7373 or annette.wilkinson@tdlpathology.com

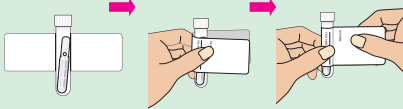
TEST	CODE	SAMPLE REQ	TAT
HPV mRNA (All High Risk Subtypes)	HPVY	Self-collection kit	3 days
HPV Individually Typed High Risk DNA Subtypes	HPVZ	Self-collection kit	10 days

Self-collection HPV samples

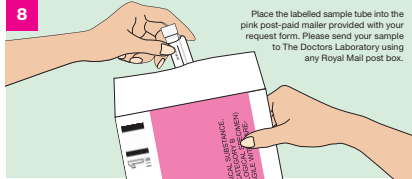
7

Make sure your sample is labelled with your details using the label supplied in your pack. This is very important as unlabelled samples cannot be processed.

Affix the label by placing the sample tube in the middle of the label and wrapping the label around the shaft as shown below.



8



The Doctors Laboratory
The Halo Building, 1 Mableton Place, London WC1H 9AX
Tel: 020 7307 7373 Fax: 020 7307 7374 E-mail: tdl@tdlpathology.com
Website: www.tdlpathology.com

© The Doctors Laboratory, 2017

1602860/19-09-09-04

Ovintip® is a registered trademark of Aprecise AB, Sweden.



**THE DOCTORS
LABORATORY**

SELF-COLLECTION HPV SAMPLE

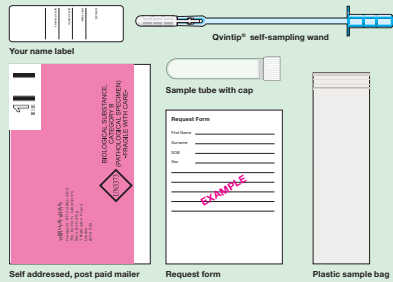
Sample Collection Instructions

Please read these instructions the whole way through before attempting to collect your sample.

Clearly complete the Name Label using a ball point pen with:

- Your surname
- Your first name
- Your date of birth
- Date of sample collection

Your sample collection pack contents



Self-sampling step-by-step

Before use, check that the product is intact (blue and white self-sampling wand, sample tube with cap, pink post-paid mailer). The self-sampling wand should be handled with care and only according to these instructions. Hold the wand straight when taking it in and out of your vagina. You can take your test in a standing or lying position. Don't collect a sample during your period. Sampling can be carried out during the first three months of pregnancy.

General Information

An infection with human papilloma virus (HPV) could potentially lead to cervical cancer. Your sample will be tested for prevalence of high-risk HPV. Your request will be handled confidentially. The results of the analysis will be posted to you.

Negative results

If the results are negative and the test shows no high-risk HPV, it means there is currently very little risk of cervical cancer. Please note that you might be infected at a later stage. HPV is sexually transmitted.

Positive results

If the results are positive, it means you have an infection with high risk subtypes. Please contact your gynaecologist for follow-up counselling. Women with persistent infection run an increased risk of cell changes which may lead to cervical cancer. Detecting an infection at an early stage allows for treatment.

PLEASE NOTE - FOR EASY SELF SAMPLING

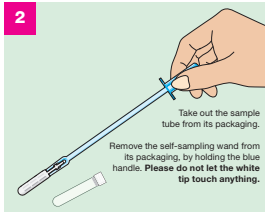
- Please note that the self-sampling wand is intended for **single use only**. The self-sampling wand should be handled with care and **only according to these instructions**.
- The white tip of the wand **must not be bent or removed** before self-sampling.
- To ensure correct results, the sample must **immediately be sent in by post** after taking.

1



Wash your hands in warm soapy water.

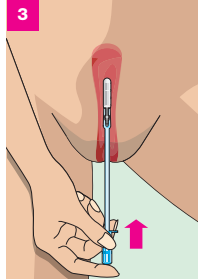
2



Take out the sample tube from its packaging.

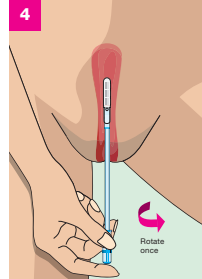
Remove the self-sampling wand from its packaging, by holding the blue handle. **Please do not let the white tip touch anything.**

3



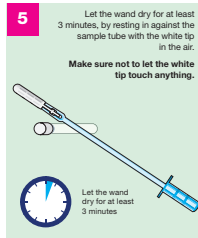
Keep the wand straight while collecting your sample. Insert the wand into your vagina (see diagram).

4



Rotate the self-sampling wand once. Take the wand out (keeping it straight).

5



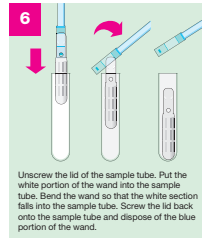
Let the wand dry for at least 3 minutes, by resting it against the sample tube with the white tip in the air.

Make sure not to let the white tip touch anything.



Let the wand dry for at least 3 minutes

6



Unscrew the lid of the sample tube. Put the white portion of the wand into the sample tube. Bend the wand so that the white section falls into the sample tube. Screw the lid back onto the sample tube and dispose of the blue portion of the wand.

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Non-Gynae Cytology

Non-Gynaecological Cytology

Urines

To prevent cell degeneration it is advisable to collect urine samples in a sample pot containing preservative (available from TDL Supplies). Use of preservative will ensure the cellular material is preserved up to 48 hours.

Ideally 10 mls (excluding preservative) from a freshly fully voided urine (when the bladder is emptied) mid-morning sample should be submitted for cytological assessment. If microbiology or chemistry investigations are also required, **please submit separate urine samples** and mark the vials accordingly. A mid-stream urine sample is NOT recommended for cytological assessment as it could lead to a low cellular yield. If a delay of greater than 24 hours in reaching the laboratory is anticipated samples should be refrigerated at 4°C.

Sputum

Sputum should be collected on at least three occasions if underlying lung carcinoma is suspected. A single sputum is sufficient for microbiological assessment. Sputum should be sent to the laboratory immediately following production, or stored in a universal container containing cytolyt cell fixative if there is a likely delay. Please note that this is only acceptable if sputum is only for Cytology. Microbiology cannot be performed on fixed material. Early morning sputum is ideal, but contamination with food, toothpaste and tobacco should be avoided.

Fluids

All available material should be submitted in a sterile container without fixative as quickly as possible. If any delay is anticipated, the material should be submitted in cytolyt fixative.

Cerebrospinal fluid (CSF)

Ideally CSF should be submitted fresh or as an air dried cytospin slide, unstained and in a plastic transport slide box. A minimum of 3mls should be submitted either in fresh form or spun on multiple slides for cytopathologists' review and opinion. Please contact TDL Cytology for advice if required on 020 7307 7323 / 7373.

URINE/SPUTUM/FLUID			
TEST	CODE	SAMPLE REQs	TAT
Fluid Cytology	CATF	Fluid ⁴	3 days
Urine Cytology (Urine cytology containers available from TDL Supplies)	URCY	Urine (30mls) ²¹	2 days

Histopathology

CATEGORY	CODE	TISSUE SAMPLE
Breast	HIS1	Breast Capsule
Breast	HIS4	Breast Reduction (Bilateral)
Breast	HIS3	Breast Reduction (Unilateral)
Breast	HIS2	Breast Tissue
Breast	HIS2	Cavity Shavings
Breast	HIS1	Core Biopsy (1 Specimen)
Breast	HIS2	Core Biopsy (2 Specimens)
Breast	HIS3	Core Biopsy (3 Specimens)
Breast	HIS4	Core Biopsy (4 Specimens)
Breast	HIS3	Lumpectomy
Breast	HIS5	Mastectomy (simple)/Wide Local Excision (WLE)
Breast	HIS5+HIS4	Mastectomy + axillary clearance
Breast	HIS4	Microdochectomy
Breast	HIS2	Nipple
Breast	HIS5	Sentinal Nodes
Cardiac	HIS3	Aorta
Cardiac	HIS2	Cardiac Biopsy
Cardiac	HIS3	Cardiac Tumour Excision
Cardiac	HIS2	Heart Valves
Cardiac	HIS2	Mediastinal Tissue
Cardiac	HIS2	Pericardium
Cardiac	HIS2	Temporal Artery Biopsy
Endocrine	HIS5	Adrenal
Endocrine	HIS4	Parathyroid
Endocrine	HIS4	Thyroid (Lobe)
Endocrine	HIS5	Thyroid (Total)
ENT – Biopsy	HIS2	Bronchial Biopsy
ENT – Biopsy	HIS1	Cholesteatoma
ENT – Biopsy	HIS1	Dental Cyst
ENT – Biopsy	HIS1	Ear Canal Biopsy
ENT – Biopsy	HIS1	Ear Polyp
ENT – Biopsy	HIS1	Epiglottis
ENT – Biopsy	HIS1	Gingival Tissue
ENT – Biopsy	HIS1	Laryngeal Biopsy
ENT – Biopsy	HIS2	Laryngeal Nodule (Bilateral)
ENT – Biopsy	HIS1	Laryngeal Nodule (Unilateral)
ENT – Biopsy	HIS2	Mandible Biopsy
ENT – Biopsy	HIS2	Maxillary Mucosa
ENT – Biopsy	HIS2	Mucocele
ENT – Biopsy	HIS1	Nasal Biopsy
ENT – Biopsy	HIS1	Nasal Polyps
ENT – Biopsy	HIS1	Oral Biopsy
ENT – Biopsy	HIS1	Palatal Biopsy

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Histopathology

CATEGORY	CODE	TISSUE SAMPLE
ENT – Biopsy	HIS1	Pharyngeal Biopsy
ENT – Biopsy	HIS2	Pleural Biopsy
ENT – Biopsy	HIS1	Thyroid Biopsy
ENT – Biopsy	HIS1	Tongue Biopsy
ENT – Biopsy	HIS1	Tonsil (1 Specimen)
ENT – Biopsy	HIS2	Tonsil Biopsy
ENT – Biopsy	HIS2	Tonsils (2 Specimens)
ENT – Biopsy	HIS2	Uvelectomy
ENT – Biopsy	HIS1	Vocal chords
ENT – Resections	HIS5+HIS2	Glossectomy
ENT – Resections	HIS5	Laryngectomy
ENT – Resections	HIS5+HIS2	Maxillectomy
ENT – Resections	HIS5+HIS2	Neck Dissection
ENT – Resections	HIS5+HIS5	Neck Dissection (Bilateral)
ENT – Resections	HIS4	Parotidectomy
ENT – Resections	HIS4	Partial Thyroidectomy
ENT – Resections	HIS5+HIS5	Pharyngectomy
ENT – Resections	HIS5+HIS2	Rhinectomy
ENT – Resections	HIS3	Submandibular Gland – Excision
ENT – Resections	HIS2	Thyroglossal Cyst
GI Endoscopic – Biopsy	HIS1	Bile duct biopsy
GI Endoscopic – Biopsy	HIS1	Colonic Polyp
GI Endoscopic – Biopsy	HIS1	Endoscopic Biopsy (1 specimen)
GI Endoscopic – Biopsy	2H1	Endoscopic Biopsy (2 specimens)
GI Endoscopic – Biopsy	3H1	Endoscopic Biopsy (3 specimens)
GI Endoscopic – Biopsy	4H1	Endoscopic Biopsy (4 specimens)
GI Endoscopic – Biopsy	5H1	Endoscopic Biopsy (5 specimens)
GI Endoscopic – Biopsy	6H1	Endoscopic Biopsy (6 specimens)
GI Endoscopic – Biopsy	7H1	Endoscopic Biopsy (7 specimens)
GI Endoscopic – Biopsy	8H1	Endoscopic Biopsy (8 specimens)
GI Endoscopic – Biopsy	9H1	Endoscopic Biopsy (9 specimens)
GI Endoscopic – Biopsy	10H1	Endoscopic Biopsy (10-15 specimens)
GI Endoscopic – Biopsy	HIS5	Liver Biopsy – Medical
GI Endoscopic – Biopsy	HIS3	Liver Biopsy – Tumour
GI Endoscopic – Biopsy	HIS3	Omental Biopsy
GI Endoscopic – Biopsy	HIS1	Pancreatic Biopsy
GI Endoscopic – Biopsy	HIS1	Perianal Biopsy
GI-Resection – Small	HIS215	Anal Fistula
GI-Resection – Small	HIS2	Appendix
GI-Resection – Small	HIS3	Endo Mucosal Resection (EMR/ESD)
GI-Resection – Small	HIS2	Gallbladder
GI-Resection – Small	HIS2	Haemorrhoidectomy
GI-Resection – Small	HIS2	Hernia Sac
GI-Resection – Small	HIS3	Meckel's Diverticulum

Histopathology

CATEGORY	CODE	TISSUE SAMPLE
GI-Resection – Small	HIS2	Mesentery
GI-Resection – Small	HIS2	Perianal Biopsy / Warts
GI-Resection – Small	HIS2	Pilonidal Sinus
GI-Resection – Small	HIS2	Polypectomy
GI-Resection – Small	HIS2	Umbilical Lesion
GI Resection – Large	HIS5	Biliary Resection
GI Resection – Large	HIS5+HIS2	Colon
GI Resection – Large	HIS5	Distal Pancreatectomy
GI Resection – Large	HIS5+HIS2	Gastrectomy
GI Resection – Large	HIS5	Gastric Wedge Resection
GI Resection – Large	HIS5	Ileoanal Pouch Resection
GI Resection – Large	HIS4	Ileostomy
GI Resection – Large	HIS3	Ileum
GI Resection – Large	HIS5+HIS2	Large Bowel Resection – Benign/ Malignant
GI Resection – Large	HIS4	Liver Wedge Resection
GI Resection – Large	HIS5+HIS2	Oesophagectomy
GI Resection – Large	HIS5	Partial Hepatectomy
GI Resection – Large	HIS5	Small Bowel Resection – Benign/ Malignant
GI Resection – Large	HIS5+HIS5	Whipple's Procedure / Pancreatoduodenectomy
Gynaecology	HIS2	Cervical Biopsy
Gynaecology	HIS1	Cervical Polyp
Gynaecology	HIS4	Cervix
Gynaecology	HIS1	Curettings – Endocervical
Gynaecology	HIS1	Curettings – Endometrial
Gynaecology	HIS2	Endometrial Biopsy
Gynaecology	HIS1	Endometrial Pipelle
Gynaecology	HIS1	Endometrial Polyp
Gynaecology	HIS2	Fallopian Tube
Gynaecology	HIS3	Fibroids
Gynaecology	HIS2	Fimbrial Cyst
Gynaecology	HIS4	LLETZ and/or Cone Biopsy
Gynaecology	HIS2	Mastoid
Gynaecology	HIS2	Ovarian Biopsy
Gynaecology	HIS2	Ovarian Cyst
Gynaecology	HIS1	Ovarian Pipelle
Gynaecology	HIS5	Ovaries (Bilateral)
Gynaecology	HIS3	Ovary (Unilateral)
Gynaecology	HIS4	Ovary and Tube (Unilateral)
Gynaecology	HIS5	Ovary and Tube (Bilateral)
Gynaecology	HIS2	Pelvic Mass
Gynaecology	HIS1	Peritoneal Biopsy
Gynaecology	HIS5	Placenta
Gynaecology	HIS2	Pouch of Douglas
Gynaecology	HIS1	Products of Conception

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Histopathology

CATEGORY	CODE	TISSUE SAMPLE
Gynaecology	HIS2	Uterine Polyp
Gynaecology	HIS4	Uterus
Gynaecology	HIS5	Uterus and Cervix
Gynaecology	HIS5	Uterus, Tubes And Ovaries
Gynaecology	HIS1	Vulval Biopsy
Haemato-Oncology	HIS5	Bone Marrow
Haemato-Oncology	HIS2	Lymph Node
Haemato-Oncology	HIS3	Lymph Node (Lymphoma)
Haemato-Oncology	HIS3	Lymph Node (Metastatic Disease)
Haemato-Oncology	HIS5	Spleen
Haemato-Oncology	HIS5	Thymus
Lung – Biopsy	HIS3	Lung Biopsy
Lung – Resections	HIS3	Lung Lesion Small Wedge Resection
Lung – Resections	HIS5+HIS5	Lung Resection
Lung – Resections	HIS5	Lung Tumour Resection +/- Nodes
Neurosurgery	HIS3	Brain Biopsy
Neurosurgery	HIS3	Brain Resection
Neurosurgery	HIS5+HIS5	Muscle Biopsy
Neurosurgery	HIS3	Pituitary Gland – Resection
Neurosurgery	HIS3	Spinal Tumour Biopsy
Neurosurgery	HIS3	Spinal Tumour Resection
Neurosurgery	HIS4	Vertebrea
Ophthalmic	HIS1	Conjunctival Biopsy
Ophthalmic	HIS1	Cornea
Ophthalmic	HIS4	Globe/Removal of Eye
Ophthalmic	HIS2	Lacrimal Gland Biopsy/Excision
Ophthalmic	HIS1	Orbit Contents Of Eye
Orthopaedic	HIS1	Bone Biopsy
Orthopaedic	HIS2	Bone Currettings
Orthopaedic	HIS2	Bursa
Orthopaedic	HIS2	Duputrenes Contracture
Orthopaedic	HIS3	Femoral Head Resection
Orthopaedic	HIS1	Ganglion Cyst
Orthopaedic	HIS3	Joint Resurfacing/Redo Prosthesis Capsule
Orthopaedic	HIS1	Neuroma
Orthopaedic	HIS2	Synovial Biopsy
Orthopaedic	HIS3	Tendon
Skin and Soft Tissue	HIS2	Abscess
Skin and Soft Tissue	HIS3	Alopecia Biopsies
Skin and Soft Tissue	HIS1	Cyst Excision
Skin and Soft Tissue	HIS1	Fossa
Skin and Soft Tissue	HIS1	Granuloma
Skin and Soft Tissue	HIS3	Lipoma
Skin and Soft Tissue	HIS2	Skin Excision BCC/SCC


Histopathology

CATEGORY	CODE	TISSUE SAMPLE
Skin and Soft Tissue	HIS1	Nail
Skin and Soft Tissue	HIS1	Pilonidal Sinus
Skin and Soft Tissue	HIS5	Sentinel Nodes In Skin Cancer (Melanoma)
Skin and Soft Tissue	1SK	Skin Biopsy (1 specimen)
Skin and Soft Tissue	2SK	Skin Biopsy (2 specimens)
Skin and Soft Tissue	3SK	Skin Biopsy (3 specimens)
Skin and Soft Tissue	4SK	Skin Biopsy (4 specimens)
Skin and Soft Tissue	5SK	Skin Biopsy (5 specimens)
Skin and Soft Tissue	6SK	Skin Biopsy (6 specimens)
Skin and Soft Tissue	7SK	Skin Biopsy (7 specimens)
Skin and Soft Tissue	8SK	Skin Biopsy (8 specimens)
Skin and Soft Tissue	9SK	Skin Biopsy (9 specimens)
Skin and Soft Tissue	10SK	Skin Biopsy (10 specimens)
Skin and Soft Tissue	11SK	Skin Biopsy (11-15 specimens)
Skin and Soft Tissue	HIS3	Soft Tissue Tumour Biopsy
Skin and Soft Tissue	HIS3	Soft Tissue Tumour Resection
Urology – Biopsy	HIS1	Bladder Biopsy
Urology – Biopsy	HIS1	Core Biopsy (Urology)
Urology – Biopsy	HIS2	Hydrocele
Urology – Biopsy	HIS2	Penile Biopsy
Urology – Biopsy	HIS1	Prostate biopsy
Urology – Biopsy	2H1	Prostate biopsies x 2
Urology – Biopsy	3H1	Prostate biopsies x 3
Urology – Biopsy	4H1	Prostate biopsies x 4
Urology – Biopsy	5H1	Prostate biopsies x 5
Urology – Biopsy	6H1	Prostate biopsies x 6
Urology – Biopsy	7H1	Prostate biopsies x 7
Urology – Biopsy	8H1	Prostate biopsies x 8
Urology – Biopsy	9H1	Prostate biopsies x 9
Urology – Biopsy	10H1	Prostate biopsies x 10-12
Urology – Biopsy	HIS5	Testicular Biopsy (Bilateral)
Urology – Biopsy	HIS4	Testicular Biopsy (Unilateral)
Urology – Biopsy	HIS1	Urethral Biopsy
Urology – Biopsy	HIS2	Vasectomy
Urology – Resection	HIS5+HIS5	Cystoprostatectomy
Urology – Resection	HIS3	Epididymis
Urology – Resection	HIS1	Foreskin / Circumcision
Urology – Resection	HIS5	Nephrectomy / Kidney
Urology – Resection	HIS5+HIS5	Prostatectomy
Urology – Resection	HIS5+HIS5	Radical Cystectomy
Urology – Resection	HIS3	Testis
Urology – Resection	HIS3 – HIS5+	TURBT (dependent on number of blocks)
Urology – Resection	HIS3 – HIS5	TURP (dependent on number of blocks)

Special instructions for samples

- 1 Contact the laboratory for special sample tubes/containers/instructions.
- 2 Confirmation of not negative drug screens by LCMS/MS may take up to 5 days.
- 3 Clinical history essential and protect from light.
- 4 Send to the laboratory without delay.
- 5 Do not send sample to the laboratory between Friday noon and Monday morning.
- 6 Contact the Referrals Department before taking and sending sample to the laboratory.
- 7 Sample should be separated and frozen if sending overnight.
- 8 DRP Form required. DRP Form can be found at the back of the guide.
- 9 Clinical history must be provided.
- 10 Contact the laboratory for special stability tubes for lymphocyte subsets – or take an EDTA sample and ensure same day delivery to the laboratory, Monday to Friday noon (do not send sample between Friday noon and Monday morning).
- 11 Patient consent required. Consent Form can be found at the back of this guide.
- 12 Please provide one sample for each person being tested.
- 13 Protect from light.
- 14 Provide details of travel history.
- 15 Ammonia
Sample: EDTA plasma only. Full tubes and tightly stoppered. On ice, centrifuged and analysed 20-30 mins post venepuncture (or plasma can be frozen). If haemolysed gives falsely high results.
Patient: Fasting. Avoid smoking.
- 16 Lactate
Sample: Fluoride oxalate plasma only.
On ice and separate from cells 15-30 mins, analyse promptly. Handle with care as sweat contains large amounts of lactate. No tourniquet.
Patient: Rest 30 mins prior to test.
- 17 Homocysteine
Should be spun and separated with 1 hour of venepuncture.
- 18 Citrate Samples
Samples should be double spun and separated and frozen within 4-8 hours of sample taking, if a delay is expected with transportation to the laboratory, samples must be transported as frozen.
- 19 Must include patient's age, height and weight.
- 20 Sample types: FCRU or PCR swab or TPV or Semen.
- 21 Urine cytology container, ideally first catch, mid-morning specimen.
- 22 Must be fresh.
- 30 Collect sample at end of exposure.
- 33 Sample must be labelled by hand with first name, family name, gender and date of birth detailed on sample and form. Do not use labels other than the tube label.
- 34 Samples must arrive in the laboratory on the same day of sample taking or contact the laboratory.
- 35 Patient should be fasting and resting for 30 mins before sample taking. Samples need handling urgently.
- 36 Renin: Sample collected either upright/active or resting/supine (3 hours lying).
- 37 Provide sample time and date of collection.
- 38 EDTA sample should not be separated: send whole blood.
- 39 Urgent samples have a 3 day TAT if genotype is required for prenatal diagnosis or two weeks TAT if urgent for other factors.
- 40 Informed Consent is required for these tests.
- 41 Recommendation for patient to attend Patient Reception for sample taking.
- 42 LGV can be added to a positive chlamydia sample using the same swab if requested within 4 days of receipt of result.

Example of profile panel information

Profile name	PRE-TRAVEL SCREEN (DVT)	
Profile content	FBC Factor II Prothrombin Gene Factor V Leiden Anticardiolipin Antibodies	
Turnaround time		TAT 5 DAYS
Sample requirements	DVT1 	Code Reference to sample taking and special handling instructions (see above)

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
1,25 Vitamin D	D3	B	5-8 days	144
2-Butanone GC	BUTA	RU	7 days	156
2-Furoic Acid	2FA	RU	10 days	156
4th Generation HIV1& 2 Abs/p24 Ag (45 days post-contact)*	THIV	B Tiny™	4 hours	94
5 HIAA	RU5H	PU ¹	5 days	27
5' Nucleotidase	5NT	B	5 days	27
5th Generation HIV1& 2 Abs/p24 Ag (45 days post-contact)*	THV5	B Tiny™	4 hours	94
6-Thioguanine Nucleotides	TGN	A A	2 weeks	27
7 STI Profile by PCR (7 tests from 1 Sample)	PP12	FCRU/PCR/TPV/Semen	2 days	65,75
11 Deoxycorticosterone	DEOX	B	10 days	49
11 Deoxycortisol	11DC	B (Frozen)	10 days	49
16S rRNA Bacterial Gene	16S	J	1 week	40
17 Hydroxyprogesterone	17OH	B	5 days	49
18S rRNA Fungal Gene	18S	J	1 week	40
21 Hydroxylase Ab's	21HA	B (Frozen)	10 days	27
Acetone – Blood	ACTB	A or H	2 weeks	156
Acetone – Urine	ACTU	RU	5 days	156
Acetylcholine Receptor Autoantibodies	ACRA	B ⁴	5 days	27
Acetylcholinesterase Isoenzymes	ACEI	AF	7 days	27
Acid Phosphatase – Total	APT	B	5 days	27
ACTH (Adreno Corticotrophic Hormone)	ACTH	A (Plasma Frozen) ⁴¹	1 day	49
Activated Protein C Resistance	APCR	C (Frozen) ^{4,18}	3 days	37
Acute Viral Hepatitis Screen	AHSC	B	4 hours	77
ADAMTS-13 Activity	CP13	C (Frozen)	3 days	37
ADAMTS-13 Antibody	A13A	C (Frozen)	1 month	37
Adenosine Deaminase	AD	A / B / Fluid	3 weeks	27
Adenovirus by PCR	ADV	A / PCR / VS / SC	7 days	96
Adiponectin	ADIP	B	2 weeks	27
Adrenal Cortex Antibodies	ACTX	B	2 days	77
Albumin	ALB	B	4 hours	27
Alcohol (Medical)	ALCO	G ¹	4 hours	27
Alcohol (Urine)	UALC	RU	4 hours	27
Alcohol Profile	AP	A B B G	5-7 days	153-154,156
Alcohol Profile 2	ALCP	A A B B G RU	5-7 days	153-154,156
Aldolase	ALDO	B	5 days	27
Aldosterone	ALDN	B	5 days	49
Aldosterone (Urine)	UALD	PU	5 days	49
Alk Phosphatase Isoenzymes	APIE	B	5 days	27
Alkaline Phosphatase	ALP	B	4 hours	27
Allergen Component Profiles				141
Allergy – Individual Allergens See list on page 137	ALLE	B	2 days	134

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
Allergy Profile (Mediterranean)	ALMD	B	2 days	133-134
Allergy Profile (Middle East)	ALME	B	2 days	133-134
Allergy Profile (UK)	ALUK	B	2 days	133-134
Allergy Profile 1 (Food & Inhalants)	1A	B B	2 days	134-135
Allergy Profile 2 (Inhalants)	2A	B	2 days	134-135
Allergy Profile 3 (Food)	3A	B	2 days	134-135
Allergy Profile 4 (Nuts & Seeds)	4A	B	2 days	134-135
Allergy Profile 5 (Children's Panel)	5A	B	2 days	134-135
Allergy Profile 6 (Shellfish)	6A	B	2 days	134, 136
Allergy Profile 7 (Finfish)	7A	B	2 days	134, 136
Allergy Profile 8 (Cereal – singles)	8A	B	2 days	134, 136
Allergy Profile 9 (Antibiotics)	9A	B	2 days	134, 136
Allergy Profile 10 (Insects)	10A	B	2 days	134, 136
Allergy Profile 11 (Combined Shellfish/Finfish)	11A	B	2 days	134, 136
Allergy Profile 12 (Milk & Milk Proteins)	12A	B	2 days	134, 136
Allergy Profile 13 (Stone fruit/Rosaceae family)	13A	B	2 days	134, 136
Alpha 1 Antitrypsin (Serum)	A1AT	B	1 day	27
Alpha 1 Antitrypsin (Stool)	A1AF	RF	10 days	27
Alpha 1 Antitrypsin Genotype – Pi*M, Pi*S, Pi*Z	GENE	A ⁹	4 weeks	27, 107
Alpha 1 Glycoprotein	OROS	B (Frozen)	5 days	27
Alpha 1 Microglobulin	A1MG	RU ^{1,22}	10 days	27
Alpha 2 Macroglobulins	A2MG	B	5 days	27
Alpha Feto Protein	AFP	B	4 hours	49, 99
Alpha Feto Protein (Maternal)	AFPM	B	4 hours	27
Alpha Gal Components (related to red meat)	ZZ37	B	2 days	141
ALT (Alanine Aminotransferase) (SGPT)	ALT	B	4 hours	27
Alternaria Components	ZZ1	B	2 days	141
Aluminium (Blood)	ALUM	K	7 days	27, 155
Aluminium (Urine)	ALUU	RU	1-2 weeks	156
Amenorrhoea Profile	AMEN	B	4 hours	49, 55
Amikacin Level (State dose)	AMIK	B ⁴	4 hours	129
Amino Acid (Serum/Plasma)	AMIN	B	7 days	27
Amino Acid Quantitative (Urine)	UAAQ	RU	7 days	27
Amino-Laevalinic Acid (Urine)	RUAL	100mls PU	5 days	27
Amitriptyline	AMTR	A ⁴	5 days	130
AML/ALL Molecular MRD – NPM1, PML-RARA, CBFB-MYH11, RUNX1-RUNX1T1, ETV6-RUNX1 – <i>Contact Lab for further information</i>	GENE	Bone Marrow / A	5 days	107
Ammonia	AMMO	A (Frozen) ¹⁵	4 hours	27
Amniocentesis – rapid BOBs aneuploidy diagnosis for all chromosomes (5 days) + culture (10-15 days)	ABK	AF ⁹	5-15 days	107
Amniocentesis – rapid PCR diagnosis for common aneuploidies (2 days) + culture (10-15 days)	APCC	AF ⁹	2-15 days	107
Amoebic (E. histolytica) Antibodies	AFAT	B	2 days	86









































Key: See page 21 for sample taking and special handling instructions.

Alphabetical test index







































TEST	CODE	SAMPLE REQ	TAT	PAGE
Amoebic (E. histolytica) PCR	AMAG	RF	2 days	86
Amphetamines – Blood	AMPB	B B	5 days	153
Amylase	AMY	B	4 hours	27
Amylase (Urine)	UAMY	CU	4 hours	27
Amylase Isoenzymes	AMYI	B	5 days	27
Amyloidosis (Amyloid A Protein)	SAA	B	5 days	27
Anaemia Profile	ANAE	A A B	2 days	36, 39
Anafranil (Clomipramine)	CHLO	A	7 days	130
ANCA (Anti-Neutrophil Cytoplasmic Abs)	ANCA	B	2 days	77
Andropause Profile	ANDP	B B	8 hours	49, 54
Androstanediolglucoronide	ANDG	B	3 weeks	27
Androstenedione	ANDR	B (Frozen)	4 days	49
Angiotensin II	ANG2	A (Frozen)	2 weeks	27
Angiotensin Converting Enzyme	ACE	B	4 hours	27
Angiotensin Converting Enzyme – CSF	ACEF	CSF (Frozen)	2 weeks	27
Antenatal Profile	ANTE	A A ³³ B B B G	3 days	36, 39
Anti-Actin Antibodies	AAA	B	5 days	77
Anti-Basal Ganglia Antibodies	ABGA	B	3 weeks	77
Anti-CCP Antibodies (RF)	CCP	B	2 days	77
Anti-Liver Cytosol Antibodies	ALCA	B	5 days	77
Anti-MOG [Myelin Oligodendrocyte Glycoprotein] Antibodies	AMOG	B	3 weeks	77
Anti-MUSK Antibodies	MUSK	B	2 weeks	77
Anti-Phosphatidylserine Antibodies	PHTS	B	5 days	77
Anti-Phospholipase A2 Receptor	AA2R	B	3 weeks	77
Anti-Ri Antibodies	RIAB	B	3 days	77
Anti-SLA (Soluble Liver Antigen) Abs	LSA	B	10 days	77
Anti-Xa Apixaban monitoring	APIX	C (Frozen)*	3 days	37
Anti-Xa Fondaparinux monitoring	FOND	C (Frozen)*	3 days	37
Anti-Xa- LMWH monitoring	LMWX	C (Frozen)*	3 days	37
Anti-Xa- Rivaroxaban monitoring	RIVA	C (Frozen)*	3 days	37
Antidiuretic Hormone	ADH	A A (Plasma Frozen) ⁴	10 days	49
Antimony (Urine)	ANTI	RU ³⁰	10 days	27
Antimullerian Hormone (AMH Plus)	AMH	B	4 hours	27, 49, 54
Antinuclear Antibodies (titre & pattern)	ANAB	B	2 days	77
Antistaphylolysin Titre (SGOT)	ASTT	B	3 days	77
Antistreptolysin Titre/ASOT	ASLT	B	2 days	77
Antisulfatide Antibodies	ASA	B	5 weeks	77
Antithrombin III	A111	C (Frozen) ^{4,9,18}	3 days	37
AP50 Alternative Hemolytic Complement	AP50	B (Frozen)	2 weeks	27
Apolipoprotein A1	APOA	B	3 days	27
Apolipoprotein B	APOB	B	3 days	27
Apolipoprotein C	APOC	B	3 months	27
Apolipoprotein E (12 hours fasting)	APOE	B (fasting)	5 days	27

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.














































Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
Apolipoprotein E genotype – E2, E3, E4	APEG	 ⁹	5 days	108
Apple Components	ZZ36		2 days	141
APTT/KCCT	KCCT	 ¹⁸	4 hours	36
Aquaporin 4 Antibodies (Neuromyelitis Optica)	AQUA		2 weeks	77
Arbovirus Antibodies/Abs	ARBO	 ^{9,14}	3 weeks	96
Array CGH (Comparative Genomic Hybridisation)	CGH	CVS/AF/   ⁹	10 days	108
Arsenic (Blood)	ARS	 or 	5 days	28,155
Arsenic (Urine)	ARSE	RU ³⁰	5 days	28,156
Arylsulphatase A	ARYL	 ^{5,6}	8 weeks	28
Ascariasis Serology	ASC		5 days	77
Ashkenazi Jewish Carrier Screen	ASHJ	 ⁹	4 weeks	108,115, 123,128
Aspartate Transaminase (AST) (SGOT)	AST		4 hours	28
Aspergillus Components	ZZ2		2 days	141
Aspergillus Precipitins	ASPP		5 days	40
Atypical Antibody Screen (handwritten tube label)	AASC	 ^{22,33}	2 days	36
Atypical Pneumonia Screen	APS		2 days	96,98
Autoantibody Profile I	AUTO		2 days	77,85
Autoantibody Profile II	ENDO		2 days	77,85
Avian Precipitins (11 Species)	AVIA		5 days	77
Azoospermia – karyotype + cystic fibrosis screen + polyT(5T) + Y deletions	GRP	  ⁹	10-15 days	108
Babesia Antibodies	BABE		3 weeks	77
Bancroftia/Oncerciasis/Filarial Antibodies	TFIF	 ¹⁴	2 weeks	96
BCR/ABL Quantitative – fusion gene sizes p190 + p210 – <i>MUST arrive in the laboratory within 48 hours, before 12pm on Fridays</i>	BCRA	  ⁹	10 days	108
Becker Muscular Dystrophy – deletions/duplications	DND	 ⁹	10 days	108
Behcet's Disease – HLA Tissue Typing B*51	B51	 ⁹	10 days	108
Bence-Jones Protein	RBJP	1 x 30mls (RU)	5 days	28
Benzene	BENZ	J ^{1,6}	3 days	156
Beta 2 Glycoprotein 1 Abs	B2GP		5 days	77
Beta 2 Microglobulin (Serum)	B2MG		2 days	28,156
Beta 2 Microglobulin (Urine)	UB2M	RU	3 days	28,156
Beta Carotene	CARO		5 days	144
Beta D Glucan	XBDG		3 days	40
Beta HCG (Oncology)	HCGQ		4 hours	99
Beta HCG (Quantitative)	QHCG		4 hours	49
Beta-Glucuronidase (Sly Disease)	BGLU	  ^{9,4}	8 weeks	28
Bicarbonate	HCO3		4 hours	28
Bile Acids – Serum	BILE		4 hours	28
Bilharzia (Schistosome) Antibody Screen	BILH	 ¹⁴	10 days	86
Bilharzia (Schistosome) Antigen	SHAG		15 days	86



















Alphabetical test index

	TEST	CODE	SAMPLE REQ	TAT	PAGE
	Bilharzia (Urine)	USCH	RU ¹⁴	2 days	86
	Bilirubin (Direct/Indirect)	DBIL		4 hours	28
	Bilirubin (Total)	BILI		4 hours	28
	Bilirubin (Urine)	UBIL	RU	1 day	28
	Biotin	BIOS		1 week	144
	Biotinidase	BIOT	 (Frozen plasma) ⁴	3 weeks	28
	Birch Components	ZZ3		2 days	141
	Bismuth	BISM		5 days	28
	BK Polyoma Virus by PCR	BKPV	 /  /RU	5 days	96
NEW	Bleeding and platelet disorders NGS Panel (known familial mutations) – Contact lab	GENE	 	6 weeks	108
NEW	Bleeding and platelet disorders NGS Panel (unknown familial mutations) – Contact lab	GENE	 	12 weeks	108
	Blood Culture	BCUL	2 x BC ⁴	6 days +	40
	Blood Film Examination	FILM		1 day	36
	Blood Group	ABO	 ^{22,33}	2 days	36
	BNP (NT-pro BNP)	BNP		4 hours	28, 49
	Bone Alkaline Phosphatase	BALP	 (Frozen)	2 weeks	28
	Bone Marrow (Aspirate)	BMAS	J ¹	14 days	38
	Bone Marrow (Trephine Biopsy)	BMI	J ¹	3 days	38
	Bone Screen	BONE	 CU	4 hours	28, 35
	Bone Screen (Bloods only)	BON2		4 hours	28, 35
	Borrelia Antibodies (Lyme Disease) IgG, IgM	BORR	 ^{9,14}	2 days	77, 86
	Borrelia Antibodies (Lyme Disease) IgM	BORM		2 days	77, 86
	Borrelia Confirmation (Immunoblot)	BORC	 ^{9,14}	10 days	77, 86
	BRAF V600E mutation by PCR for Hairy Cell Leukaemia	GENE	Bone Marrow/ 	5 days	108
	Brazil Components	ZZ4		2 days	141
	Breast Cancer – BRCA1 + BRCA2 only gene sequencing + deletions/duplications	GENE		4 weeks	109
	Breast Cancer NGS Panel – full sequencing across 14 genes + deletions/duplications. Requires patient informed consent	GENE	  ^{9,11}	4 weeks	99, 109
	Bromide	BROM		3 days	156
	Brucella Serology	BRUC	 ⁹	2-3 weeks	77
	BUN (Blood Urea Nitrogen)	BUN		4 hours	28
	C-KIT D816V mutation by PCR for Mastocytosis	GENE	Bone Marrow/ 	5 days	109
	C Peptide	CPEP		3 days	49
	C Reactive Protein	CRP		4 hours	28
	C Reactive Protein (High Sensitivity)	HCRP		4 hours	28
	C1 Esterase Inhibitor	C1EI		5 days	77
	C1 Esterase: Function & Total	FC1E	  (Plasma Frozen) ^{4,18}	10 days	28
	C1q Binding Immune Complex	IMCP		5 days	28
	C3 Complement	C3		4 hours	77

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
C3/C4 Complement	COMP		4 hours	77
C4 Complement	C4		4 hours	77
CA 15-3	C153		4 hours	99
CA 19-9	C199		4 hours	99
CA 50	CA50		5 days	99
CA 72-4	C724		5 days	99
CA 125	C125		4 hours	99
Cadmium (Blood)	CADM	 or 	5 days	28, 155
Cadmium (Urine)	URCD	RU ³⁰	5 days	28, 156
Calcitonin	CATO	 (Frozen) ⁴	1 day	49
Calcium	CA		4 hours	28
Calcium (24 hour Urine)	UCA	PU	4 hours	28
Calcium/Creatinine Ratio	CACR	RU 	4 hours	28
Calprotectin	CALP	RF	5 days	77
Calprotectin/Elastase Profile	CEP	RF	5 days	77, 85
Campylobacter Jejuni Antibodies	CJAB		5 days	77
Candida Antibodies	CANA		5 days	77
Candida Antigen	CCAG		5 days	77
Cannabinoids (Urine) Screen	CANN	RU	1 day	153
Carbamazepine (Tegretol)	CARB		4 hours	130
Carbapenemase producing organism screen	MDR	STM (rectal)	4-5 days ±	40
Carbohydrate Deficient Glycoprotein	CDG		2 weeks	28
Carbohydrate Deficient Transferrin (CDT)	CDT	 ⁴	3 days	28
Carboxyhaemoglobin	CBHB		1 week	36
Carcino Embryonic Antigen	CEA		4 hours	99
Cardiac Enzymes (not chest pain)	CENZ		4 hours	28
Cardiolipin Antibodies (IgG+IgM)	ACAB		2 days	77
Cardiovascular Risk Profile 1	PP10	  	3 days	28, 35
Cardiovascular Risk Profile 2	PP11	    ³⁴	3 days	28, 35
Carnitine – Free & Total	CARN	  (Frozen Plasma)	10 days	28
Carotenes	CAR0	 ¹³	5 days	144
Cartilage Antibodies	ACA		5 days	77
Cashew Components	ZZ35		2 days	141
Cat Components	ZZ5		2 days	141
Cat Scratch Fever (Bartonella IgG+IgM)	CAT		5 days	96
Catecholamines (Plasma)	CATE	  (Plasma Frozen) ⁴	5 days	49
Catecholamines (Urine)	UCAT	PU ¹	5 days	49
CCP Antibodies (RF)	CCP		2 days	77
CD3/CD4/CD8	LYSS	 ¹⁰	1 day	38, 94, 96
CD16	CD16	 ⁴	1 day	38
CD19 B Cells	CD19	 ⁴	1 day	38
CD20	CD20	 ¹⁰	2 days	38
CD25	CD25	 ¹⁰	2 days	38
CD56	CD56	 ⁴	1 day	38

Alphabetical test index




























TEST	CODE	SAMPLE REQ	TAT	PAGE
CD57	CD57		1 day	38
Celery Components	ZZ6		2 days	141
Centromere Autoantibodies	CENT		2 days	77
Ceruloplasmin	CERU		1 day	28,144
Cervical Cytology	PAPT will include HPV	TPV	2-3 days	162
CH50 (Classical pathway)	CH50	 (Frozen) ⁴	4 days	77
Chagas Disease Serology (S.American Trypanosomiasis) T. Cruzi	CHGA	 ^{9,14}	10 days	77
Chest Pain Profile	CPP		STAT	28, 35
Chikungunya Virus Abs	CHIK	 ^{9,14}	10 days	96
Chlamydia (PCR swab)	SPCR	PCR	2 days	65
Chlamydia (Thin Prep)	TPCR	TPV	2 days	65,160
Chlamydia (Urine)	CPCR	FCRU	2 days	65
Chlamydia Species Specific (MIF) Ab Screen	CHAB		2 days	78,85
Chlamydia/Gonorrhoea (PCR Swab)	SCG	PCR	2 days	65
Chlamydia/Gonorrhoea (Rectal)	RSCG	PCR	2 days	65
Chlamydia/Gonorrhoea (Thin Prep)	TCG	TPV	5 days	65,160
Chlamydia/Gonorrhoea (Throat)	TSCG	PCR	2 days	65
Chlamydia/Gonorrhoea (Urine)	CCG	FCRU	2 days	65
Chlamydia/Gonorrhoea/Trichomonas by PCR	CCGT	FCRU / PCR / TPV	2 days	65
Chloride	CL		4 hours	28
Cholesterol	CHO		4 hours	28
Cholesterol (Familial Hypercholesterolaemia)	GENE	  ⁹	4 weeks	28,112
Cholinesterase (Blood)	CHRC		5 days	28,156
Cholinesterase (Serum/Pseudo)	CHPS		4 hours	28,156
Chromium (Blood)	CHRO		5 days	28,155
Chromium (Urine)	URCR	RU ³⁰	10 days	29,156
Chromogranin A	CGA		5 days	29
Chromogranin A & B	MTAB	J ¹	3 weeks	29
Chromosome Analysis (Amniocentesis) – culture only	ACUL	AF ⁹	10-15 days	109
Chromosome Analysis (Amniocentesis) – rapid BOBs aneuploidy diagnosis for all chromosomes (5 days) + culture (10-15 days)	ABK	AF ⁹	5-15 days	109
Chromosome Analysis (Amniocentesis) – rapid PCR diagnosis for common aneuploidies (2 days) + culture (10-15 days)	APCC	AF ⁹	2-15 days	109
Chromosome Analysis (Blood)	KARY	 ⁹	8-18 days	110
Chromosome Analysis (Chorionic Villus) – rapid BOBs aneuploidy diagnosis for all chromosomes (5 days) + culture (10-15 days)	CBK	CVS ⁹	5-15 days	110

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
Chromosome Analysis (Chorionic Villus) – rapid PCR diagnosis for common aneuploidies (2 days) + culture (10-15 days)	CVPC	CVS ^{1,9}	2-15 days	110
Chromosome Analysis (Chorionic Villus) – culture only	CVSC	CVS ^{1,9}	10-15 days	110
Chromosome Analysis (Products of Conception) – BOBs rapid aneuploidy diagnosis for all chromosomes (5 days) + culture (25 days)	PBK	Placental Sample ^{1,9}	5-25 days	110
Chromosome Analysis (Products of Conception) – reflex to BOBs testing if culture fails to grow	PROC	Placental Sample ^{1,9}	20-25 days	110
Chromosome Analysis (Solid Tissue)	PROC	Fetal tissue ^{1,9}	4-5 weeks	110
Chromosome Analysis (Stem Cells)	STEM/ SUSP	Culture / Fixed cells	Contact lab	110
Chronic Fatigue Syndrome Profile	VIP1	A + B ¹⁰	5 days	78, 85
Citrate (Blood)	CITR	B	5 days	29
Citrate (Urine)	UCIT	CU (Frozen)	5 days	29
CK (MB Fraction)	CKMB	B	4 hours	29
CK Isoenzymes	CKIE	B	5 days	29
Clobazam	CLOB	A	5 days	130
Clomipramine (Anafranil)	CHLO	A	7 days	130
Clonazepam	CLON	A	7 days	130
Clostridium Difficile Toxin by PCR	CLOS	RF*	2 days	40
CMV DNA (by PCR)	CMVP	A	5 days	96
CMV DNA by PCR (Semen)	SCVM	Semen	7 days	96
CMV DNA by PCR (Urine)	CMVU	RU	5 days	96
CMV Resistance	CMVR	A A (2 x 6mls)	21 days	96
Coagulation Profile 1	CLPF	C ¹⁸	4 hours	36, 39
Coagulation Profile 2	CLOT	A C ¹⁸	4 hours	36, 39
Cobalt (Blood)	COB	A	5 days	29
Cobalt (Serum)	COBB	B	5 days	29, 155
Cobalt (Urine)	COBA	RU ³⁰	5 days	29, 156
Cocaine (Urine) Screen	UCOC	RU	1 day	153
Coeliac Disease – HLA DQ2/DQ8 Genotype	Q2Q8	A ⁹	10 days	78, 82, 110
Coeliac/Gluten Profile 2	GSA2	A B	10 days	78, 82
Coeliac/Gluten Sensitivity Profile	GSA	B	2 days	78, 82
Coenzyme Q10	CQ10	B	2 weeks	29
Cold Agglutinin	CAGG	J ¹	5 days	29
Collagen (Type I, II, IV) Antibodies	COAB	B	10 days	29
Collagen Type 1 Cross-Linked N-Telopeptide – NTX	NTX	2nd EMU	2 weeks	29
Colloid Antigen-2 Antibodies	CA2A	B	2 weeks	78
Colorectal Cancer NGS Panel – full sequencing across 18 genes + deletions/duplications. Requires patient informed consent	GENE	A A ^{9,11}	4 weeks	110
Comparative Genomic Hybridisation (Array CGH)	CGH	CVS/AF/A H ⁹	10 days	110

Alphabetical test index


























TEST	CODE	SAMPLE REQ	TAT	PAGE
Complement C1q	C1Q		5 days	29
Complement C2	C2		10 days	29
Complement C5	C5A		2 weeks	29
Complement C6	C6	 (Frozen)*	5 weeks	29
Complement C7	C7	 (Frozen)*	5 weeks	29
Complement C8	C8	 (Frozen)*	5 weeks	29
Complement C9	C9	 (Frozen)*	5 weeks	29
Complement Factor H	FACH		3 weeks	29
Complex PSA (Prostate Specific Ag)	CPSA		3 days	99
Congenital Absence of Vas Deferens – karyotype + cystic fibrosis screen + polyT(5T) + Y deletions	GRP	  ⁹	10-15 days	110
Coombs (Direct Antiglobulin Test)	COOM		2 days	38
Copper (Serum)	COPP		5 days	29,144,155
Copper (Urine)	URCU	CU	5 days	29,156
Cortisol	CORT		4 hours	49
Cortisol (Urine)	UCOR	CU	5 days	49
Cortisol Binding Globulin	CBG	 (Frozen)	1 month	29
Cotinine (Saliva)	SCOT	Saliva Kit¹	1-2 weeks	156
Cotinine (Serum)	COT		4 days	78
Cotinine (Urine)	COTT	RU	2 days	78
NEW COVID-19 (SARS-CoV-2) Abbott IgG Antibody	GCOV	SST/Serum * (Venous)	24 hours	96
NEW COVID-19 (SARS-CoV-2) RNA by PCR	NCOV	PCR Swab (nasal/pharyngeal)	48 hours	96
NEW COVID-19 (SARS-CoV-2) Roche Elecsys Anti-SARS-CoV-2 Total Antibody	TCOV	SST/Serum ** (Venous and Capillary)	24 hours	96
Cow's Milk Components	ZZ7		2 days	141
Coxsackie Antibodies (IgM)	COXM		10 days	96
Creatine Kinase (CK, CPK)	CKNA		4 hours	29
Creatinine	CREA		4 hours	29
Creatinine (Urine)	UCR	CU	4 hours	29
Creatinine Clearance	CRCL	 CU	4 hours	29
Cri du Chat Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS/AF/  ⁹	5-15 days	110
Cri du Chat Syndrome – BOBs only	PBOB	CVS/AF/  ⁹	5 days	110
Crosslaps (Serum DPD)	SDPD	 (Freeze within 24 hours)	4 days	29
Cryoglobulins	CRYO	J⁶	10 days	78
Cryptococcal Antigen	CRYC	Serum or CSF	1 day	40
Cryptosporidium	CRPO	RF	2 days	40
Cryptosporidium Detection by PCR	CRPA	RF	2 days	86
CSF for Microscopy and Culture	CSF	CSF	1-3 days	40
CSF Screen by PCR	VPCR	CSF	2 days	96,98
CT/GC/Trichomonas/Mgen (PCR Swab)	SGTM	PCR Swab	2 days	65,75
CT/GC/Trichomonas/Mgen (Urine)	CGTM	FCRU	2 days	65,75
Culture (Any site)	CULT		up to 5 days	40

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
CVS PCR for common aneuploidies (2 days) + culture (10-15 days)	CVPC	CVS ⁹	2-15 days	110
CVSBOBs – rapid BOBs aneuploidy diagnosis for all chromosomes (3-5 days) + culture (10-15 days)	CBK	CVS ⁹	5-15 days	111
CVSBOBs only – rapid aneuploidy diagnosis for all chromosomes + common microdeletion syndromes	CBOB	CVS ⁹	5 days	111
Cyclic Amp (Urine)	CAMP	CU (Frozen)	5 days	29
Cyclosporin (Monoclonal)	CYCL	A	1 day	29
Cyfra 21-1	CY21	B	4 days	99
CYP450 2D6 Genotyping	TGEN	A ⁹	10 days	111
Cystatin C	CYCC	B	5 days	29
Cystic Fibrosis (139 common mutations) – reflex to Poly T when required	CFS	A ⁹	5 days	111
Cystine – Quantitative (Beta-CTX)	QCYS	PU	5 days	29
Cytomegalovirus (CMV-DNA) Amnio	CMVD	AF	5 days	96
Cytomegalovirus (IgG/IgM) Antibodies	CMV	B	4 hours	96
Cytomegalovirus (PCR) Urine	CMVU	RU	5 days	96
Cytomegalovirus Avidity	CMAV	B	10 days	96
Cytomegalovirus DNA (PCR)	CMVP	A	5 days	96
Cytomegalovirus IgM	CMVM	B	4 hours	96
D-Dimers (Fibrinogen Degradation Products)	DDIT	C ⁴	4 hours	36
Dengue Fever PCR	DPCR	A or B ^{9,14}	2 weeks	96
Dengue Virus Serology	DENG	B ^{9,14}	5 days	86
Deoxyypyridinoline (DPD) – Serum	SDPD	B (Freeze within 24 hours)	4 days	29
Deoxyypyridinoline (DPD) – Urine	DPD	EMU	4 days	29
DHEA	DHEX	B	7-10 days	49
DHEA – Urine (Dehydroepiandrosterone)	UDHE	CU	3 weeks	49
DHEA Sulphate	DHEA	B	4 hours	49
Diabetic Profile 1	DIAB	A G	8 hours	29, 35
Diabetic Profile 2	DIA2	A G RU	2 days	29, 35
Diamine Oxidase Activity	DIAM	B	2 weeks	78
Diazepam (Valium)	DIAZ	A	7 days	130
DiGeorge Syndrome (22q11 & 10p14 deletion) – BOBs (5 days) + karyotype (15 days)	DGB, KARY	CVS/AF/A H ⁹	5-15 days	111
DiGeorge Syndrome (22q11 & 10p14) – BOBs only	DGB	CVS/AF/A ⁹	5 days	111
Digoxin	DIGO	B	4 hours	130
Dihydrotestosterone	DHT	B B	7 days	49
Diphtheria Antibodies	DIPH	B	5 days	78
DL1-DL12 Screening Profiles				22-23
DNA (Double Stranded) Antibodies IgG	DNAA	B	2 days	78
DNA (Single Stranded) Antibodies	DNAS	B	5 days	78

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
DNA Extraction & Storage – 3 years (longer upon request)	XDNA	 9	20 days	111
DNA Identity Profile – 15 STR markers	DNAF	 9	10 days	111
Dog Components	ZZ8		2 days	141
Down Syndrome Risk Bloods only (Risk to be calculated by clinician)	HCGF/ PAPA		4 hours	49
Down Syndrome Risk Profile (2nd trimester) Quad	DRP	 DRP form ^{7,8}	2 days	49
Down Syndrome Risk Profile with risk calculation first trimester	DRP	 DRP form + image of scan ^{7,8}	2 days	49
Doxepin Level (Sinequan)	DOXE		10 days	156
Drugs of Abuse From Blood	DOAP		5 days	153-154
Drugs of Abuse Profile – Random Urine Sample/ No Chain of Custody	DOA	RU	2 days (5 days with LCMS/MS confirmation)	153-154
Drugs of Abuse Profile – Random Urine Sample/No Chain of Custody Plus Alcohol	DOA3	RU	2 days (5 days with LCMS/MS confirmation)	153-154
Drugs of Abuse Profile – With Chain of Custody	DOAL	RU/CoC Collection Containers ^{1,2}	2 days (5 days with LCMS/MS confirmation)	153-154
Drugs of Abuse Profile – Without Chain of Custody	DOAN	RU ²	2 days (5 days with LCMS/MS confirmation)	153-154
Duchenne Muscular Dystrophy – deletions/duplications only	DMD	 9	10 days	111
DVT/Pre-travel Screen	DVT1	   9	5 days	36, 39, 86-87, 111, 128
Early CDT-Lung	CDTL		7 days	99
Early Detection Screen PCR/NAAT	STDx	 10mls or 2 x 4mls (Vacutainer only)	3 days	65, 75
Early Detection Screen PCR/NAAT with Syphilis	STXX	  10mls or 2 x 4mls	3 days	65, 75
Echinococcus (Hydatid) Antibodies	EFAT	 9,14	5 days	78, 86
Eczema Provoking Profile	ALEC		2 days	134
Egg Components	ZZ9		2 days	141
Ehlers-Danlos Syndrome/Aneurysm/Connective Tissue Disorders NGS Panel – full sequencing across 46 genes + deletions/duplications	GENE	  9	5 weeks	111
Ehrlichiosis Antibodies	EHRL	 9,14	10 days	78
Elastase (Faecal)	ELAS	RF	5 days	78
Elastase/Calprotectin Profile	CEP	RF	5 days	78, 85
Electrolytes	ELEC		4 hours	29
Electrolytes (Urine)	UELE	CU	4 hours	29
ELF/Enhanced Liver Fibrosis	ELF		5-7 days	29
Endometrial Biopsy Immune Profiling	23RF	J (Contact Referrals)	2 weeks	52
Endomysial Antibodies (IgA)	AEAB		2 days	78

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
Enteric Organism Rapid Detection	EORD	RF	2 days	86-87
Eosinophil Cationic Protein	ECP	B	7 days	30
Epanutin (Phenytoin)	PHEN	B	4 hours	130
Epstein-Barr Virus Antibodies IgG/IgM	EBVA	B	2 days	96
Epstein-Barr Virus PCR	EBVQ	A	5 days	96
Erectile Dysfunction Profile	IMPO	A B B G	3 days	49, 54
Erythropoietin	ERY	B	4 days	38, 130
ESR	ESR	A	4 hours	36
Essential Fatty Acid Profile (Red Cell)	EFAR	A ⁴	10 days	144
Ethosuximide	ETHO	A	7 days	130
Extractable Nuclear Antibodies (nRNP, Sm, Ro, La, Jo1, Scl70) CENP-B	ENA	B	2 days	78
Factor II Assay	FAC2	C (Frozen) ^{9,18}	5 days	37
Factor II Prothrombin – G20210A mutation	FX2	A ⁹	5 days	112
Factor V Assay	FAC5	C (Frozen) ^{9,18}	5 days	37
Factor V Leiden – G1691A mutation	FX5	A ⁹	5 days	112
Factor VII Assay	FAC7	C (Frozen) ^{9,18}	5 days	37
Factor VIII Assay	FAC8	C (Frozen) ^{9,18}	5 days	37
Factor VIII Inhibiting Antibody	F8IA	C C ¹⁸	2 weeks	37
Factor IX Assay	F1X	C (Frozen) ^{9,18}	5 days	37
Factor IX Inhibiting Antibody	F9IA	C C ¹⁸	2 weeks	37
Factor X Assay	FX	C (Frozen) ^{9,18}	5 days	37
Factor XI Assay	FX1	C (Frozen) ^{9,18}	5 days	37
Factor XII Assay	FX11	C (Frozen) ^{9,18}	5 days	37
Factor XIII Assay	FA13	C (Frozen) ^{9,18}	5 days	37
Faecal Elastase	ELAS	RF	5 days	78
Faecal Fat (1 Day Collection)	TFFA	LF ⁶	5 days	30
Faecal Fat (3 day)	FFAT	LF ⁶	5 days	30
Faecal Lactoferrin	FLAC	RF	5 days	30
Faecal Occult Blood/FOB (immunochemical/FIT)	QFIT	QFIT	1 day	40
Faecal Sugar Chromatography	FCRO	RF (Frozen)	3 weeks	30
Faecal Urobilinogen	FURO	RF	5 days	30
Familial Hypercholesterolaemia – LDLR + APOB + PCSK9 + LDLRAP1 screening	GENE	A A ⁹	4 weeks	112
Farmers Lung Precipitins	FARM	B	5 days	78
Fasciola Hepatica Antibodies (Liver Fluke)	FASC	B	2 weeks	78
FASTest Sexual Health Screening Tests				69
Fat Globules in Faeces	FGLO	RF	1 week	30
Female Hormone Profile	FIP	B	4 hours	49, 54
Ferritin	FERR	B	4 hours	30
Fibrinogen	FIB	C ^{4,18}	4 hours	36
Fibrotest (Liver Fibrosis)	FIBT	B	2 weeks	30
Filaria (Lymphatic and Non-Lymphatic) Antibodies	FIFA	B ^{9,14}	10 days	86

Alphabetical test index








































TEST	CODE	SAMPLE REQ	TAT	PAGE
First Trimester Antenatal Screen (Risk to be calculated by requesting clinician)	HCGF/ PAPA	B	4 hours	49, 55
Fish Components	ZZ10	B	2 days	141
FK506 (Tacrolimus/Prograf)	FK5	A ⁴	1-2 days	130
Flecainide (Tambocor)	FLEC	A	5 days	130
Fluid Culture	FLUD	SC	2-7 days	40
Fluid Cytology	CATF	Fluid ⁴	3 days	165
Fluid for Crystals	FLU2	SC	1 day	40
Fluoride (Urine)	UFL	RU	5 days	30
Fluoxetine (Prozac)	PROZ	A ⁴	5 days	130
Folate (Red Cell)	RBCF	A	2 days	30, 144
Folate (Serum)	FOLA	B	1 day	30
Fragile X Syndrome screen – FMR1 repeat analysis PCR (3 weeks) + Southern Blot (8 weeks) if required	GENE	A A A ⁹	3-8 weeks	113
Free Cortisol (Urine)	UCOR	CU	5 days	49
Free Fatty Acids	FFA	B (Frozen) ¹	10 days	30
Free T3	FT3	B	4 hours	49
Free T4	FT4	B	4 hours	49
Fructosamine	FRUC	B	1 day	30
Fructose – Plasma	FRU	G ⁷ (Frozen)	5 days	30
FSH	FSH	B	4 hours	49
Full Blood Count	FBC	A	4 hours	36
Fungal ID + Sens	FUID	Fungal sample/STM	14 days	40
G6PD	G6PD	A	3 days	38
Gabapentin	GABA	B ⁴	5 days	130
Galactomanan (Aspergillus Antigen)	SGAL	B	2 weeks	40
Galactose-1-Phosphate Uridyltransferase	GAL1	H ^{5,6}	2 weeks	30
Galactosidase – Alpha*	GALA	J *	6 weeks	30
Gall Stone Analysis	RSTA	STONE	10 days	30
Gamma GT	GGT	B	4 hours	30
Ganglionic Acetylcholine Receptor Antibodies	GACA	B	1 month	78
Ganglioside GM1, GD1B, GQ1B Abs	GANG	B	5 days	78
Gardnerella vaginalis by PCR	GVPC	FCRU/PCR/TPV	2 days	65, 160
Gastric Parietal Autoantibodies	GASP	B	2 days	78
Gastrin	GAST	B (Frozen)	5 days	30
Genetic Reproductive Profile (Male)	GRP	A H ⁹	10-15 days	113, 116, 128
GENETICS: TDL Genetics – see pages 101-128				101-128
Gentamicin Assay	GENT	B ⁴	4 hours	129
Giardia Serology	GIAR	B	5 days	78
Gliadin Antibodies (IgG) (deamidated)	AGAB	B	2 days	78
Globulin	GLOB	B	4 hours	30
Glomerular Basement Membrane Abs	AGBM	B	2 days	78
Glucagon	GLUG	J ¹	10 days	30

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index














































TEST	CODE	SAMPLE REQ	TAT	PAGE
Glucose	RBG	G	4 hours	30
Glucose Challenge Test/Mini-GTT	RBGM	G	1 day	129
Glucose Tolerance Test (Short)	GTTS	2x G 2xRU	1 day	129
Glucose Tolerance Test (Extended Plus)	GTTX	7x G 7xRU	1 day	129
Glucose Tolerance Test (Extended)	GTTE	5x G 5xRU	1 day	129
Glucose Tolerance with Growth Hormone	GTT + GHDF	3x B ³⁵ 3x G 3xRU	1 day	129
Glucose Tolerance with Insulin	GTTI	3x B 3x G 3xRU	1 day	129
Glucose Tolerance Test/OGTT	GTT	3x G 3xRU	1 day	129
Glutamic Acid Decarboxylase Antibodies (GAD 65)	GAD	B	5 days	78
Glutathione (Red Cell)	GLUR	H ⁵	5 days	144
Glutathione Peroxidase	GLPX	H	5 days	144
Gluten Allergy Profile	GLUT	A B B	10 days	78, 82, 134
Gluten Sensitivity Evaluation	GSA	B	2 days	78
Gluten/Coeliac Profile 2	GSA2	A B	10 days	78
Glycan Determinants	ZZ27	B	2 days	141
Gonorrhoea (Culture)	GONN	CS	2-3 days	40, 65
Gonorrhoea (PCR swab)	SGON	PCR	2 days	65
Gonorrhoea (Thin Prep)	TGON	TPV	2 days	65
Gonorrhoea (Urine)	CGON	FCRU	2 days	65
Granulocyte Immunology	GRIM	A A	2 weeks	78
Group B Strep	GBSX	2 x STM	3-4 days	40
Growth Hormone (Fasting)	GH	B ^{7,35}	4 hours	49
Gut Hormone Profile	GUTP	A A (Frozen within 15 minutes) ⁴¹	3 weeks	49
H. pylori Antibodies (IgG)	HBPA	B	2 days	78
H. pylori Antigen (Breath)	HBQT	J	5 days	78
H. pylori Antigen (Stool)	HBAG	RF	3 days	78
H. pylori Culture	HPCU	J	3 weeks	40
Haematology Profile	PP3	A	4 hours	36, 39
Haemochromatosis – HFE common mutations C282Y + H63D	HMD	A ⁹	3 days	30, 113
Haemoglobin	HB	A	4 hours	36
Haemoglobin Electrophoresis	HBEL	A	4 days	38
Haemophilus B Influenzae Antibodies	HINF	B	7 days	78
Haemophilus ducreyi by PCR	DUCR	PCR	7 days	65
Haemosiderin (Urine)	HSID	EMU	2 weeks	30
Hams Test for PNH (CD59)	HAMS	J ^{34,5}	5 days	38
Hantavirus Serology	HANV	B ⁹	10 days	96
Haptoglobin	HAPT	B	5 days	30
Harmony® Prenatal Test (Non-Invasive Prenatal Testing) – common aneuploidy screening from maternal blood	NIPT	J/Special tubes ¹	3-5 days	113

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
Harmony® Prenatal Test (Non-Invasive Prenatal Testing) – common aneuploidy screening from maternal blood including 22q11.2 del	NIPQ	J/Special tubes ¹	3-5 days	113
Hazelnut Components	ZZ11		2 days	141
HbA1c	GHB		6 hours	30
HDL Cholesterol	HDL		4 hours	30
HDL2 & HDL3 Fractions	HDLF		3 weeks	30
HE4 + ROMA (Earlier Detection of Ovarian Tumour)	HE4		1 day	99
Hepatitis (Acute) Screen	AHSC		4 hours	90
Hepatitis A (IgM)	HAVM		4 hours	90
Hepatitis A Immunity (IgG/IgM)	HAIM		4 hours	89-90
Hepatitis A Profile	HEPA		4 hours	65,90
Hepatitis A RNA by PCR	HAVR	 or 	3 weeks	90
Hepatitis A, B & C Profile	ABC		4 hours	90
Hepatitis B 'e' Antigen and Antibody	HEPE		4 hours	90
Hepatitis B (PCR) Genotype	BGEN		7 days	90
Hepatitis B Core Antibody – IgM	HBCM		4 hours	90
Hepatitis B Core Antibody – Total	HBC		4 hours	90
Hepatitis B DNA (Viral load)	DNAB		5 days	90
Hepatitis B Immunity	HBIM		4 hours	89-90
Hepatitis B Profile	HEPB		4 hours	90
Hepatitis B Resistant Mutation	HBRM	 or 	7 days	90
Hepatitis B Surface Antigen	AUAG		4 hours	65,90
Hepatitis C Abs Confirmation (RIBA)	RIBA		5 days	90
Hepatitis C Antibodies	HEPC		4 hours	65,90
Hepatitis C Genotype	CGEN		5 days	90
Hepatitis C Quantification (Viral Load)	QPCR	 or 	5 days	90
Hepatitis Delta Antibody	HEPD		5 days	90
Hepatitis Delta Antigen	HDAG		5 days	90
Hepatitis Delta RNA	DRNA	 (Frozen plasma)	5 days	90
Hepatitis E (PCR)	EHEP		2 weeks	90
Hepatitis E IgG/IgM	HBE		5 days	90
Hepatitis G (PCR)	HEPG	 (Frozen plasma)	2 weeks	90
Herpes Simplex I/II Antibody Profile (IgG)	HERP		2 days	96
Herpes Simplex I/II by PCR (Swab)	HERS	PCR	5 days	65,96
Herpes Simplex I/II by PCR (Urine)	HERD	FCRU/PCR/TPV	5 days	65,96,160
Herpes Simplex I/II IgM	HERM		2 days	96
HFE gene (Haemochromatosis) – common mutations C282Y + H63D	HMD	 ^a	3 days	38,114
Hirsutism Profile	HIRP		4 hours	49,55
Histamine (Blood)	HITT	 (Frozen plasma)	5 days	78
Histamine (Urine)	HITU	RU	5 days	78
Histamine Releasing Urticaria Test	CURT		10-14 days	78,134

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
Histone Antibodies	HISA		5 days	78
Histopathology				166-170
Histoplasmosis	HISP		10 days	78
HIV 1 & 2/p24Ag	HDUO		4 hours	65
HIV 1 Proviral DNA	HIVP	 Whole blood	7 days	94
HIV Confirmation of Positive Screens (Using 3 methodologies)	HIVC		1 day	94
HIV Rapid RNA HIV-1 QUALITATIVE	LHIV	 (Vacutainer only)	4 hours	65, 76, 94-95
HIV Rapid RNA HIV-1 QUANTITATIVE	RHIV	 (Vacutainer only)	4 hours	65, 76, 94-95
HIV Screening: HIV1& 2 Abs, p24 (5th Gen)	HIV5		4 hours	94-95
HIV Screening: HIV1& 2 Abs/p24 Ag (4th Gen)	HDUO		4 hours	94
HIV Therapeutic Drug Monitoring	TDM	J	21 days	94
HIV-1 Genotypic Resistance (Integrase)	INTE	  (2x6ml whole blood)	21 days	94
HIV-1 Genotypic Resistance (RT & Protease)	HIVD	  (2x6ml whole blood)	21 days	94
HIV-1 RNA Viral Load by PCR	HIV1	  (2x6ml whole blood)	3 days	94
HIV-1 Tropism	TRPM	  (2x6ml whole blood)	28 days	94
HIV-2 RNA by PCR	HIV2		21 days	94
HIV/HBV/HCV (Early detection by PCR/NAAT) with Syphilis	STXX	  10mls or 2 x 4mls	3 days	65, 75
HIV/HBV/HCV Screen by PCR/NAAT (10 days post exposure)	STDx	 10mls or 2 x 4mls (Vacutainer only)	3 days	65, 75, 94-96
HLA B*57:01	HL57	 ⁹	10 days	94
HLA B27	HLAB	 ⁹	3 days	79
HLA DQ Alpha Antigens	10RF	 	2 weeks	52
HLA DQ Beta Antigens	11RF	 	2 weeks	52
HLA DR Antigens	9RF	 	2 weeks	52
HLA Tissue Typing A	HLA	 ⁹	10 days	114
HLA Tissue Typing A+B	HLBA	 ⁹	10 days	114
HLA Tissue Typing A+B+C (Class I)	HABC	 ⁹	10 days	114
HLA Tissue Typing A/B/DRB1/3/4/5	HLAF	 ⁹	10 days	114
HLA Tissue Typing A/B/DRB1/3/4/5/DQB1	HLF	 ⁹	10 days	114
HLA Tissue Typing A/B/C/DRB1/3/4/5/DQB1 (Class I & II)	HLFC	 ⁹	10 days	114
HLA Tissue Typing B	HLB	 ⁹	10 days	114
HLA Tissue Typing B*27 only	HLAB	 ⁹	3 days	114
HLA Tissue Typing B*51 (Behcet's Disease)	B51	 ⁹	10 days	114
HLA Tissue Typing B*57:01 high resolution	HL57	 ⁹	10 days	114
HLA Tissue Typing C	HLC	 ⁹	10 days	114
HLA Tissue Typing Coeliac Disease – DQ2/DQ8	Q2Q8	 ⁹	10 days	114
HLA Tissue Typing DRB1/3/4/5	DRB1	 ⁹	10 days	114
HLA Tissue Typing DRB1/3/4/5/DQB1 (Class II)	HLDQ	 ⁹	10 days	114
HLA Tissue Typing Narcolepsy – DQB1*06:02	GENE	 ⁹	4 weeks	114
Homocysteine (Quantitative)	HOMO	 ¹⁷	1 day	30










































Key: See page 21 for sample taking and special handling instructions.

Alphabetical test index










































TEST	CODE	SAMPLE REQ	TAT	PAGE
Homocysteine (Urine)	HCYS	CU	2 weeks	30
Homovanillic Acid (HVA)	HVA	PU	5 days	30
House Dust Mite Components	ZZ12	B	2 days	141
HPV (DNA and reflexed mRNA)	HPVT	TPV	3 days	65,162
HPV (HR mRNA types 16, 18 + others)	HPVH	TPV	2-3 days	65,162
HPV (Individual low & high risk DNA subtypes)	HP20	TPV/PCR	2-3 days	65,162
HPV Individually Typed High Risk DNA Subtypes	HPVZ	Self-collection kit	10 days	163
HPV mRNA (All High Risk Subtypes)	HPVY	Self-collection kit	3 days	163
HRT Profile 1	HRT	B	4 hours	49,55
HRT Profile 2	HRT2	B G	4 hours	49,55
HTLV 1& 2 Abs. (Human T Lymphotropic Virus Type I-II)	HTLV	B	8 hours	94
HTLV by PCR	HTLP	A Whole blood	21 days	94
Hughes Syndrome	LUPA	B C 4,18	2 days	37
Human Anti-Mouse Antibodies	HAMA	B (Frozen)	6 weeks	79
Human Herpes Virus – 6 by PCR	HHV6	A	5 days	96
Human Herpes Virus – 8 (IgG)	HHV8	B	10 days	96
Human Herpes Virus – 8 by PCR	HHV8D	A	5 days	96
Human Parvovirus B19 – DNA	PCRPP	A	2 weeks	96
HVS	HVS	STM ^{†††}	2-4 days	40
Hyaluronic Acid	AHT	B	1 week	30
Hydroxybutyrate Dehydrogenase	HBD	B (Frozen)	1 week	30
Hydroxyprenalene	UHYD	CU	2 weeks	30
Identity Profile (DNA) – 15 STR markers	DNAF	A 9,11	10 days	115
IgE (Total)	IGE	B	1 day	79
IGF-1 (Somatomedin)	SOMA	B (Frozen) ⁴	1 day	50
IGF-BP3	IGF3	B (Frozen) ⁴	5 days	50
IgG Subclasses	IGSC	B	4 days	30
Imipramine	IMIP	A ⁴	4 days	130
Immune Function Evaluation (Total)	TIE	A + B 5,10	7 days	36
Immune-Complexes	IMCP	B	5 days	79
Immunoglobulin A	IGA	B	4 hours	30
Immunoglobulin D	IGD	B	5 days	30
Immunoglobulin E – Total	IGE	B	1 day	30
Immunoglobulin G	IGG	B	4 hours	31
Immunoglobulin M	IGM	B	4 hours	31
Immunoglobulins (IgG, IgM, IgA)	IMM	B	4 hours	31,79
Impotence Profile	IMPO	A B B B G	3 days	50,54
Inhibin A	INIA	B	1 month	50
Inhibin B	INIB	B (Day 3 of cycle, frozen)	5 days	50
Inner Ear Antigen (Ottoblot)	IEA	B	3 weeks	79
INR	PTIM	C 18	4 hours	36
Insect/Worm/Ova/Cysts	FLEA	Send Specimen ^{9,14}	5 days	86

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
Insulin	INSU		4 hours	50
Insulin Antibodies	INAB		5 days	79
Insulin Resistance (Fasting)	FIRI	 	4 hours	50
Insulin-Like Growth Factor 2	IGF2	 ⁶	1 month	31
Interferon – Alpha	IFA	 (Frozen) ⁹	3 weeks	79
Interferon – Gamma	IFG	 (Frozen)	3 weeks	79
Interleukin 1 Beta	ILB	 (Frozen) ^{4,7}	1-2 weeks	79
Interleukin 2	IL2	 (Frozen) ^{4,7}	1-2 weeks	79
Interleukin 4	IL4A	 (Frozen) ^{4,7}	1-2 weeks	79
Interleukin 6	IL6	 (Frozen) ^{4,7}	1-2 weeks	79
Interleukin 8	IL8	 (Frozen) ^{4,7}	1-2 weeks	79
Interleukin 10	IL10	 (Frozen) ^{4,7}	1-2 weeks	79
Interleukin 28b Genotype	IL28		2 weeks	79
Intrinsic Factor Antibodies	IFAB		2 days	79
Iodide – Urine	UIOD	RU	1 week	31
Iodine – Serum	IODI		1 week	31
Ionised Calcium	ICPA		5 days	31
Iron (TIBC included)	FE		4 hours	31
Iron Overload Profile	IOP	   ⁹	3 days	31, 34, 115, 128
Iron Status Profile	ISP		4 hours	31, 34
ISAC Panel	ISAC		3 days	134-135
Islet Cell Antibodies	ICAB		2 days	79
IUCD for Culture	IUCD	Send Device	11-12 days	40
JAK2 V617F genotyping assay	JAK2		2 weeks	115
JC Polyoma Virus by PCR	JCPV	  / CSF	5 days	96
Jewish/Pan-ethnic carrier screening	ASHJ	 ⁹	4 weeks	108, 115, 123, 128
Ketamine Screen	KETA	RU	7-10 days	153
KIR (Killer-like Immunoglobulin-like Receptors) Genotyping	17RF	  	2-3 weeks	52
Kiwi Components	ZZ32		2 days	141
Lactate (Plasma)	LACT	 ¹⁶	1 day	31
Lactate Dehydrogenase (LDH)	LDH		4 hours	31
Lactate Pyruvate Ratio	LPR	J ¹	4-6 weeks	31
Lactose Intolerance Gene	LACG		2 weeks	115
Lactose Tolerance Test	LTT	By appointment only	1 day	129
Lamotrigine	LAMO	 ⁴	5 days	130
Langer-Giedion Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS/AF/  H ⁹	5-15 days	115
Langer-Giedion Syndrome – BOBs only	PBOB	CVS/AF/  ⁹	5 days	115
Latex Components	ZZ13		2 days	141
LDH Isoenzymes	ISOL		5 days	31
LDL7 Subfractions	LDL7		10 days	31

Alphabetical test index






























TEST	CODE	SAMPLE REQ	TAT	PAGE
Lead (Blood)	LEAD		5 days	31, 155
Lead (Urine)	URPB	RU	5 days	31, 156
Lead Profile (Hb, ZPP, Lead)	LEAZ	 ¹³	3-5 days	155
Legionella Antibodies	LEGO		2 days	79
Legionella Urine Antigen	LEGA	RU	1 day	40, 79
Leishmania Antibodies	LEIS		5 days	86
Leptin	LEPT	 ¹⁹	5 days	31
Leptospirosis (Weil's Disease) Abs (IgM)	LEP		5 days	79
Leucine Amino Peptidase	LAP		5 days	31
Leucocyte Antibody Detection Panel FEMALE	8RF		1 week	52
Leucocyte Antibody Detection Panel MALE	7RF	   ^{3,4,6}	1 week	52
Leukaemia Immunophenotyping	LYPT	 ^{4,5}	5 days	38
Leukotriene E4	LTE4	CU (Frozen)	3 weeks	79
Levetiracetam (Keppra)	LEVE	 ⁴	3 days	130
Lipase	LIPA		4 hours	31
Lipid Profile	LIPP		4 hours	31, 34
Lipid Transfer Proteins	ZZ23		2 days	141
Lipocalins	ZZ28		2 days	141
Lipoprotein (a)	LPOA		4 hours	31
Lipoprotein Electrophoresis	LEL		5 days	31
Listeria Antibody	LIST		1 week	79
Lithium (take 12 hours after dose)	LITH		4 hours	31, 130
Liver Fibrosis (Enhanced Liver Fibrosis ELF)	ELF		5-7 days	31
Liver Fibrosis Fibrotest	FIBT		2 weeks	31
Liver Function Tests	LFT		4 hours	31, 34
Liver Immunoblot	LIVI		5 days	79
Liver Kidney Microsomal Antibodies	LKM		2 days	79
Lorazepam	LORA	 ⁴	10 days	130
Lp-PLA2 (PLAC) Test	PLA2		2 days	31
LSD	LSD	RU	5 days	153
Lupus Anticoagulant and Anticardiolipin Abs	LUPA	  ^{4,18}	2 days	37, 79
Lupus Anticoagulant only	LUPC	 ¹⁸	2 days	37
Lutein	LUTE	 ¹³	2 weeks	144
Luteinising Hormone (LH)	LH		4 hours	50
Lycopene	LYCO		2 weeks	144
Lyme Disease (Borrelia Abs) IgG, IgM	BORR	 ^{9,14}	2 days	79
Lyme Disease (Borrelia Abs) IgM	BORM		2 days	79
Lymphocyte Subsets (CD3/CD4/CD8)	LYSS	 ¹⁰	1 day	36
Lymphogranuloma Venerium (LGV)	LGVP	PCR* ⁴²	1-2 weeks	65
Lysosomal Enzyme Screen	LE	  ⁶	2 months	31
Lysozyme	LYSO		5 days	31
Macrolide Resistance Test (Mgen)	MGR	FCRU/PCR	1-2 weeks	65
Macroprolactin	PRLD		4 days	50

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
Magnesium (Serum)	MG	B	4 hours	31,155
Magnesium (Urine)	URMG	PU	1 day	31,156
Magnesium (Whole blood)	RCMG	A or H	4 days	144
Malarial Antibodies (Pl. falciparum)	MALA	B ^{9,14}	5 days	86
Malarial Antibodies (species specific)	MALS	B ^{9,14}	10 days	86
Malarial Parasites	MALP	A ^{4,9,14}	STAT	36
Male Genetic Reproductive Profile	GRP	A H ⁹	10-15 days	113, 116, 128
Male Hormone Profile	MIPR	B	4 hours	50, 54
Manganese (Serum)	MANG	B	5 days	31, 155
Mannose Binding Lectin	MBL	B	3 weeks	31
MBOCA in Urine	MBOC	RU	10 days	156
Mean Cell Volume (MCV)	MCV	A	4 hours	36
Measles Antibodies (IgG) Immunity	MEAS	B	1 day	89, 96
Measles Antibodies (IgM)	MEAM	B ⁹	2 days	89, 97
Measles PCR	MEAP	Buccal swab	48 hours	97
Measles, Mumps, Rubella (MMR)	MMR	B	1 day	89
Melanin	MELA	RU ¹³	5 days	50
Melatonin (Serum)	MEL	B (Frozen)	5 days	50
Melatonin (Urine)	UMEL	CU ¹³	2 weeks	50
Meningococcal Abs	MENI	B	2-4 weeks	79
Menopause Profile	MENO	B	4 hours	50, 55
Mercury (Blood)	MERC	A or H	5 days	31, 155
Mercury (Urine)	URHG	RU ¹	5 days	31, 156
MERS Coronavirus Test	MERS	J	1 day	97
Metabolic Syndrome Profile	METS	A B B G	9 days	50, 55
Metanephrines (Plasma)	PMET	A (Frozen plasma)	7 days	50
Metanephrines (Urine)	UMEX	PU ¹	5 days	50
Methaemoglobin	METH	A	3 days	31
Methaqualone	METQ	RU	5 days	31
Methotrexate	METX	B	2 days	130
Methylmalonic Acid – Serum	MMAS	B	5 days	31
Methylmalonic Acid – Urine	MMA	CU	2 weeks	31
Metronidazole Level	METR	B ⁴	7 days	129
Microalbumin (Urine)	UMA	RU	4 hours	31
Microdeletion (common) Syndromes – BOBs only	PBOB	CVS/AF/ A ⁹	5 days	116
Microfilaria Blood Film	MICF	A	STAT	36
Miller-Dieker Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS/AF/ A H ⁹	5-15 days	116
Miller-Dieker Syndrome – BOBs only	PBOB	CVS/AF/ A ⁹	5 days	116
Mineral Screen	MINE	B K	5 days	143-144
Mineral Screen (Whole blood)	RMIN	H H	5 days	143-144
Mineral Screen and Industrial Heavy Metal Screen (Trace Metals)	TRAC	A B H K	7-10 days	144

Alphabetical test index










































TEST	CODE	SAMPLE REQ	TAT	PAGE
Miscarriage/Thrombotic Risk Profile	PROP	 18	5 days	37, 39, 120, 128
Mitochondrial Antibodies	AMIT		2 days	79
Mitochondrial Antibodies M2	MAM2		2 days	79
Molybdenum (Serum)	MOLY		5 days	156
MRSA (Rapid PCR) one swab per site	MRSA	Blue Micro Swab	4 hours	40
MRSA Culture one swab per site	MRSW	Blue Micro Swab	2 days	40
Mucopolysaccharides	MPS	RU (Frozen)	3 weeks	31
Mumps Antibodies (IgG)	MUMP		1 day	89
Mumps Antibodies (IgM)	MUMM		1 day	89, 97
Myasthenia Gravis Evaluation	MGE		5 days	79
Mycology/Skin Scrapings by PCR	DERM	Submit Sample	3-7 days	40
Mycophenolic Acid (Cellcept)	MYCP		5 days	130
Mycoplasma genitalium by PCR	MGEN	FCRU / PCR / TPV	2 days	65, 160
Mycoplasma genitalium/Ureaplasma by PCR	MUPC	FCRU / PCR / TPV	2 days	65
Mycoplasma pneumoniae IgM and IgG	MYCO		2 days	97
Mycoplasma species – DNA	MPCR		5 days	97
Myelin Associated Glycoprotein Antibodies	MAG		5 days	79
Myelin Basic Protein Antibodies	MBPA		2 weeks	79
Myeloma Screen	MYEL	 RU	5 days	31, 34
Myeloperoxidase Antibodies	MPO		2 days	79
Myocardial Antibodies	MYO		1 week	79
Myoglobin (Serum)	SMYO		4 hours	32
Myoglobin (Urine)	UMYO	RU	5-10 days	32
Myositis Panel	MYOS		2 days	79
Mysoline (Primidone)	PRIM	 4	3 days	130
N. gonorrhoea	TGON	TPV	2 days	160
Nail Clippings	DERM	Nail clippings	3-7 days	40
Natural Killer Profile 2	NKP2		2 days	36, 39
Needle Stick Injury Profile	NSI		4 hours	89, 97
Neurological Viral Screen	NVIR		2 days	97-98
Neuronal Antibody (Hu, Ri, Yo, Cv2, Ma2)	NEUR		10 days	79
Neurone Specific Enolase	NSE		5 days	99
Newborn Screening Panel	GUTH	J ¹	2 weeks	32
Nickel (Serum)	NICK		5 days	32, 155
Nickel (Urine)	NICU	RU	10 days	32, 156
NK (CD69) and NK Cytotoxicity	69C	 *	Send Mon-Thurs only	53
NK (CD69) Cell Assay	CD69	 *	Send Mon-Thurs only	53
NK Assay Follow-Up Panel	5RF		1 week	52
NK Assay Panel + Intralipids	16RF		1 week	52
NK Assay/Cytotoxicity Panel	4RF		1 week	52

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index






























TEST	CODE	SAMPLE REQ	TAT	PAGE
NK Cytotoxicity Assay	HSNK	H H H*	Send Mon-Thurs only	53
NK Cytotoxicity with suppression with steroid, IVIg and intralipin, and NK (CD69) cell assay	69CI	H H H*	Send Mon-Thurs only	53
NK Cytotoxicity with suppression, steroid, IVIg & Intralipin	NKCY	H H H*	Send Mon-Thurs only	53
NMDA Receptor Antibodies	NMDA	B	3 weeks	79
NMP22 (Bladder tumour)	NMP	J ¹	4 days	32,99
Non-Invasive Prenatal Testing – common aneuploidy screening from maternal blood	NIPT	J/Special tubes ¹	3-5 days	117,125
Non-Invasive Prenatal Testing – common aneuploidy screening from maternal blood including 22q11.2 del	NIPQ	J/Special tubes ¹	3-5 days	117,125
Nucleic Acid Antigen Antibodies	DNA	B	2 days	79
Oestradiol (E2)	OEST	B	4 hours	50
Oestriol (Estriol)	E3	B B	4 days	50
Oestrone	E1	B B	4 days	50
Olanzapine	OLAN	A ⁴	5 days	130
Oligoclonal Bands	CSFO	CSF + B	5 days	79
Oligosaccharides	UOLI	RU	6 weeks	32
Olive Components	ZZ14	B	2 days	141
Omega 3/Omega 6	OMG3	A ⁴	4 days	144-145
Opiate Screen (Urine)	UOPI	RU	2 days	153
Orosomucoid (A1AG – Alpha 1 Glycoprotein)	OROS	B (Frozen)	5 days	32
Osmolality (Serum)	OSMO	B	1 day	32
Osmolality (Urine)	ROSM	RU	1 day	32
Osteocalcin	OST	B (Frozen) ⁴	4 days	50,99
Osteoporosis Screen	OPS	B B	4 days	32, 35
Ovarian Autoantibodies	OVAB	B	2 days	79
Oxalate (Plasma)	POXA	A (Frozen)	7 days	32
Oxalate (Urine)	UOXA	PU	5 days	32
Oxidative Stress in Semen (ROS + MIOXSYS)	SROS	Semen ¹	1 day	61
P2Y12 Receptor Platelet Function Analysis (Clopidogrel Resistance)	P2Y	C (Whole blood)**	1 day	37
PAI1 4G/5G Polymorphism	PAIP	A	10 days	36
Pan-Ethnic/Jewish Carrier Screening	GENE	A ⁹	4 weeks	118, 128
Pancreatic Peptide	PP	J	4 weeks	32
PAPT and HPV	PAPT + HPV	TPV	2-3 days	162
Paracetamol	PARA	B	4 hours	130
Paragomius Serology	PRGM	B	2 weeks	79
Parathyroid Antibodies	PTHA	B	1 week	79
Parathyroid Hormone (Whole)	PTHI	B ⁴	1 day	50
Parathyroid Related Peptide	PTRP	2ml A Plasma frozen (Freeze immediately) ¹	2 weeks	32

Alphabetical test index



















































TEST	CODE	SAMPLE REQ	TAT	PAGE
Parvalbumins	ZZ29		2 days	141
Parvovirus Antibodies (IgM)	PARV		2 days	97
Parvovirus DNA by PCR	PCRP		2 weeks	97
Parvovirus IgG Antibodies	PARG		2 days	97
Parvovirus IgG/IgM Abs	PARP		2 days	97
Paternity Testing (postnatal and prenatal) – sample required from each person being tested (3 people)	PATT	 / AF/ CVS ^{9,11,12} Contact lab	5 days	118
Paul Bunnell (Monospot)	PAUL	 or 	8 hours	36
Peach Components	ZZ15		2 days	141
Peanut Components	ZZ16		2 days	141
Pemphigus/Pemphigoid Autoantibodies	SKAB		2 days	80
Pertussis (Whooping Cough) Antibodies	PERS		5 days	89
PEth (Phosphatidylethanol)	PETH	 ³⁸	5-7 days	32, 153
Pethidine – Urine	UPET	RU	4 weeks	156
Phelan-McDermid Syndrome – karyotype + FISH	KARY, FISH	CVS/ AF/  ⁹	12-17 days	118
Phencyclidine (PCP)	DUST	RU	5 days	32
Phenobarbitone	PHB		4 hours	130
Phenytoin (Epanutin)	PHEN		4 hours	130
Phosphate	PHOS		4 hours	32
Phosphate (24 hour Urine)	UPH	PU	4 hours	32
Phospholipid Antibodies	PLIP		5 days	80
Pituitary Antibodies	PITU	 ⁴	1 month	80
Pituitary Function Profile	PITF	 	1 day	50, 55
PLAC Test (Lp-PLA2)	PLA2		2 days	32
Plasminogen	PLAS	 (Frozen plasma) ⁴	5 days	32
Plasminogen Activator Inhibitor – 1	PAI1	 (Frozen plasma)	2 weeks	32
Platelet Aggregation Studies	PLAG	J ^{5,6}	3 days	37
Pleural Fluid for Culture	FLUP	SC	7 days	40
Pneumococcal Antibodies – Serotype Specific	PASS		5 weeks	80
Pneumococcal Antibody Screen	PNEU		7 days	80, 89
Pneumococcal Antigen	PNAG	RU	1 day	40
Pneumocystis Jiroveci (PCP) Examination	PCYS	BAL ^{††}	2-3 days	40
Pneumonia (Atypical) Screen	APS		2 days	97-98
Polcalcins	ZZ25		2 days	141
Polio Virus 1, 2, 3 Antibodies	POLO	 ⁹	15 days	89
Polycystic Ovary Syndrome Profile	PCOP	      ⁷	5 days	50, 55
Polycystic Ovary Syndrome SHORT	PCOS	 	4 hours	50, 55
Porphyrin (Blood)	PORP	 ³	15 days	32
Porphyrins (Faeces)	FPOR	RF ³	3 weeks	32
Porphyrins Full Screen (Total: Urine, Stool, Blood)	PORS	 RU, RF ³	3 weeks	32
Porphyrins Screen (Urine)	RPOR	RU ³	3 weeks	32
Postnatal array CGH	CGH	  ⁹	10 days	118

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
Post-Travel Screen 1 (Prior to 6 weeks)	PTS	 ¹⁴	10 days	86-87
Post-Travel Screen 2 (Prior to 6 weeks)	PTS2	 ¹⁴	10 days	86-87
Potassium	K		4 hours	32
PR-10 Proteins	ZZ22		2 days	141
Prader-Willi Syndrome (Primary Screen) – methylation PCR	PWAM	 ⁹	5 days	118
Pre-Travel Screen (DVT)	DVT1	 ⁹	5 days	36, 39, 86-87, 118, 128
Prealbumin	PALB		3 days	134
Pregnancy (Serum) [Quantitative]	QHCG		4 hours	32, 50
Pregnancy Test (Urine)	PREG	RU	4 hours	32
Pregnanetriol (Urine)	UPTR	CU (Frozen)	5 days	50
Pregnenolone	PREN		15 days	50
Prenatal array CGH	CGH	Amniotic fluid or CVS ⁹	10 days	118
Primidone (Mysoline)	PRIM	 ⁴	3 days	130
Procalcitonin	PCAL	 (Frozen) ^{4,7}	1 day	32
Procollagen 1 Peptide N-Terminal (NTX)	P1NP		5 days	32
Procollagen III Peptide	PRCO		5 days	32
Products of Conception – rapid BOBs aneuploidy diagnosis for all chromosomes (5 days) + culture (25 days)	PBK	Placental Sample ^{1,9}	5-25 days	119, 128
Products of Conception BOBs only – rapid aneuploidy diagnosis for all chromosomes	KBOB	Placental Sample or Solid Tissue ^{1,9}	3-6 days	119
Profilins	ZZ24		2 days	141
Progesterone	PROG		4 hours	50
Proinsulin	PROI	 (Frozen plasma) ⁴	5 days	50
Prolactin	PROL		4 hours	50
Prolactin (Macro)	PRLD		4 days	50
Propanalol	PRO	 ⁴	7 days	131
Propoxyphene	DPRO	RU	5 days	32
Prostate Profile (Total & Free PSA)	PR2		4 hours	99
Prostate Specific Antigen (Total)*	PSPA		4 hours	99
Prostatic Acid Phosphatase	PACP	 (Frozen)	3 days	32
Protein (Urine)	UPRT	CU	4 hours	32
Protein 14.3.3 (Creutzfeldt-Jakob Disease)	CJD	CSF (Frozen)	5 weeks	32
Protein C	PRC	 (Frozen) ^{4,9,18}	3 days	37
Protein Electrophoresis incl. immunogloblin	PRTE		2-4 days	32
Protein S Activity	PS1	 (Frozen)	5 days	37
Protein S Free Ag	FPRS	 (Frozen) ^{4,9,18}	3 days	37
Protein Total (Blood)	PROT		4 hours	32
Protein/Creatinine Ratio (Urine)	UCPR	RU	4 hours	32
Proteinase 3 Ab	PR3		2 days	80
Prothrombin Time	PTIM	 ¹⁸	4 hours	36

Alphabetical test index



































TEST	CODE	SAMPLE REQ	TAT	PAGE
Prothrombin Time + Dose	PT+D	 ¹⁸	4 hours	36
Purkinje Cell Antibody (Hu and Yo)	PURK		10 days	80
Pyruvate Kinase (M2-PK)	M2ST	RF ⁴	5 days	99
Pyruvate Kinase (M2-PK)	M2PK		5 days	99
QF-PCR rapid common aneuploidy screen	APC	AF/A ⁹	1-2 days	119
Q Fever (C Burnetti) Antibodies	QFEV		10 days	80
Rabies Antibody	RABI		10 days	89
Rapid Strep (incl. m/c/s)	RAPS	STM ^{**}	1-3 days ^{**}	40
Rapid Xpert HIV-1 RNA Qualitative – Early Detection from 10 days	LHIV	 (Vacutainer only)	4 hours	65, 76
Rapid Xpert HIV-1 RNS Viral Load – Rapid Testing for HIV-Positive Patient Prognosis and Response To Antiretroviral Therapy	RHIV	 (Vacutainer only)	4 hours	66, 76
NEW Recurrent Fever Screening – across 4 genes	GENE	 	6 weeks	119
Recurrent Miscarriage Profile (female)	RMP	       ^{9,18}	10-15 days	119, 128
Renal Calculi Screen (Metabolic)	RSPR	J ⁶	5 days	32
Renal Stone Analysis	RSTA	STONE	10 days	32
Renin	RENI	 (Frozen plasma) ³⁶	5 days	50
Reproductive Immunophenotype Panel	3RF	  	1 week	52
NEW Respiratory Viral RNA Screen by PCR	FLU4	PCR nasopharyngeal	48 hours	97-98
Reticulocyte Count	RETC		4 hours	36
Retinol Binding Protein	RBP		3 days	32
Retrograde Ejaculation	RTR0	Contact Lab	2 days	61
Reverse T3	RT3	 ^{7,37}	10 days	50
Rheumatoid Factor (Latex Test)	RF		1 day	80
Rheumatology Profile 1 (Screen)	RH	 	2 days	80, 84
Rheumatology Profile 2 (Connective tissue)	RH2	   	3 days	80, 84
Rheumatology Profile 3 (Rheumatoid/Basic)	RH3	 	2 days	80, 84
Rheumatology Profile 4 (Systemic Lupus)	RH4	  	2 days	80, 84
Rheumatology Profile 5 (Mono Arthritis)	RH5	   	3 days	80, 84
Rheumatology Profile 6 (Rheumatoid Plus)	RH6		2 days	80, 84
Rheumatology Profile 7 (Sjogren's Syndrome)	RH7		10 days	80, 84
Rhinitis Provoking Profile	ALRN		2 days	134
Rickettsial Species Antibody Profile	RICK		7 days	80, 86
Risperidone	RISP	 ⁴	7 days	131
Rotavirus in Stool by PCR	ROTA	RF	1 day	97
RPR (VDRL)	RPR		2 days	66, 80
Rubella Antibody (IgG)	RUBE		4 hours	89, 97
Rubella Antibody (IgM)	RUBM		4 hours	89, 97
Rubella Avidity	RUAV		1 week	97
Rubella PCR	RUBP	 /Amniotic Fluid	5 days	89
S100 Malignant Melanoma	S100		4 days	99

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
Saccharomyces Cerevisiae Antibodies	ASCA	B	2 weeks	80
Salicylates	SALI	B	4 hours	32
Salivary Duct Antibodies	SAB	B	12 days	80
NEW SARS-CoV-2 (COVID-19) Abbott IgG Antibody	GCOV	SST/Serum B * (Venous)	24 hours	97
NEW SARS-CoV-2 (COVID-19) RNA by PCR	NCOV	PCR Swab (nasal/pharyngeal)	48 hours	97
NEW SARS-CoV-2 (COVID-19) Roche Elecsys Anti-SARS-CoV-2 Total Antibody	TCOV	SST/Serum B ** (Venous and Capillary)	24 hours	97
Schistosoma (Urine)	USCH	Mid-morning terminal urine	8 hours	41
Schistosoma (Bilharzia) Antibodies	BILH	B ¹⁴	10 days	86
Schistosome Antigen	SHAG	B	15 days	86
Scleroderma Immunoblot	SCLI	B	5 days	80
Screening Profile 1 – Biochemistry	PP1	B G	4 hours	22
Screening Profile 2 – Haematology/Biochemistry	PP2	A B G	4 hours	22
Screening Profile 3 – Haematology	PP3	A	4 hours	22, 36, 39
Screening Profile 4 – Haematology/Biochemistry (Short)	PP4	A B G	4 hours	22
Screening Profile 5 – Haematology/Biochemistry (Postal)	PP5	A B G	4 hours	22
Screening Profile 6 – Well Person	PP6	A B G	4 hours	22
Screening Profile 7 – Well Man	PP7	A B G	4 hours	23
Screening Profile 8 – Well Person	PP8	A B G	2 days	23
Screening Profile 9F – Senior Female	PP9F	A B B G RU QFIT ⁴	2 days	23
Screening Profile 9M – Senior Male	PP9M	A B B G RU QFIT ⁴	2 days	23
Screening Profile 10 – Cardiovascular Risk 1	PP10	B B	3 days	23, 28, 35
Screening Profile 11 – Cardiovascular Risk 2	PP11	B B B C ³⁴	3 days	23, 28, 35
Screening Profile 12 – Sexual Health Screen	PP12	FCRU/PCR/TPV/Semen	2 days	23, 65, 75, 160
Seed Storage Proteins	ZZ26	B	2 days	141
Selenium (Serum)	SELE	B	4 days	33, 144
Selenium (Whole Blood)	SELR	A or H	4 days	33, 144
Sellotape Test	SELL	Send Sample***	1 day	41
Semen Analysis, Comprehensive*	SPER	Semen ¹	2 days*	61
Semen Analysis, Post-Vasectomy**	PVAS	Semen ¹	2 days	61
Semen Analysis, Vasectomy Reversal*	SPER	Semen ¹	2 days*	61
Semen Culture	SPCU	Semen	2-4 days	41, 61
Semen Fructose	SPCF	Semen	2 days	61
Semen Leucocytes	PMNS	Semen	2 days	61
Semen Parameters	SPOD	Semen ¹	1 day	61
Semen Zinc	SPCZ	Semen	up to 10 days	61
Serotonin	SERT	H (Frozen whole blood) ¹	10 days	50
Serotonin (Urine)	USER	PU 50mls (Frozen) ¹	5 days	50
Serum Albumins	ZZ30	B	2 days	141
Serum Free Light Chains	SLC	B	1 week	33

Alphabetical test index




















































TEST	CODE	SAMPLE REQ	TAT	PAGE
Sex Hormone Binding Globulin	SHBG		4 hours	50
Shrimp Components	ZZ17		2 days	141
Sickle Solubility	SSOL		4 days	38
Silver (Blood)	SILV		5 days	33,155
Silver (Urine)	USIL	RU	5 days	33,156
Sinequan (Doxepin)	DOXE		10 days	131
Sirolimus	SIRO		3 days	131
Sjogren's Syndrome	RH7		2 days	80,84
Skin (Pemphigus/Pemphigoid) Autoantibodies	SKAB		2 days	80
Skin Antibodies by Immunofluorescence	STSK		1 month	80
Skin Scrapings/Mycology by PCR	DERM	Send Sample	3-7 days	41
Sleeping Sickness Serology (African Trypanosomiasis)	TRYP	 ⁹	10 days	80
Smith-Magenis Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS/AF/   ⁹	5-15 days	120
Smith-Magenis Syndrome – BoBs only	PBOB	CVS/AF/  ⁹	5 days	120
Smooth Muscle Antibodies	ASMO		2 days	80
Sodium	NA		4 hours	33
Somatomedin (IGF-1)	SOMA	 (Frozen) ⁴	1 day	50
Soybean Components	ZZ18		2 days	141
Specific Gravity (Urine)	USG	RU	24 hours	41
Sperm Aneuploidy	SPPL	Semen ¹	4 weeks	61
Sperm Antibodies (Serum)	ASAB		5 days	61,80
Sperm Antibodies/MAR Test (Semen) ₁	ASPA	Semen	1 day	61
Sperm Comet [®]	CMET	Semen	1-2 weeks	61
Sperm Count (Post-Vasectomy)	PVAS	Semen ¹	2 days	61
Sperm DNA Fragmentation (SCSA)	SEXT	Semen ¹	1-2 weeks	61
Sperm Morphology (Kruger strict criteria)	MRPH	Semen ¹	2 days	61
Spinal Muscular Atrophy – SMN1 deletions/duplications	SMA	 ⁹	10 days	120
Sports/Performance Profile	SPOR	         ⁴	5 days	143-144
Sputum for Routine Culture	SPU1	SC	2-4 days	41
Sputum for TB Culture (AFB)	SPU2	SC	up to 8 weeks	41
Squamous Cell Carcinoma	SCC		4 days	99
SRY (Sex-determining Region Y)	SRY	 ⁹	2 days	120
STD1 M/F STD Quad	STD1	 FCRU	2 days	66,74
STD2 M/F STI Profile Plus (Urine and Serology)	STD2	 FCRU (If culture swabs are needed please request separately)	4 days	66,74
STD3 Female STD Quad (PCR Swab and Serology)	STD3	 PCR	2 days	66,74
STD4 Female STI Profile Plus (PCR Swab and Serology)	STD4	 PCR (If culture swabs are needed please request separately)	4 days	66,74

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index


































TEST	CODE	SAMPLE REQ	TAT	PAGE
STD5 Serology only	STD5	B	4 hours	66,74
STD6 Serology only without HIV	STD6	B	4 hours	66,74
STD8 Vaginitis/BV Profile using Culture & PCR Swab	STD8	PCR/STM	3 days	66,75
STD9 Symptomatic lesion sample using PCR Swab from lesion & PCR Swab	STD9	2 x PCR Swab	7 days	66,75
Steroid Cell Antibody	SCA	B	2 days	80
STI Profile: MSM1	MSM1	B /FCRU/PCR Swab Throat/PCR Swab Rectal	2 days	66,76
STI Profile: MSM2	MSM2	B /FCRU/PCR Swab Throat/PCR Swab Rectal	3 days	66,76
Stool for OCP and Culture	PENT	RF	2-3 days	41
Stool for OVA Cysts & Parasites by PCR	OCP	RF	1 day	41
Stool Reducing Substances	STRS	RF ⁷	5 days	41
Streptomycin Levels	STRM	F	5 days	131
Striated/Skeletal Muscle Antibody	STRA	B	2 days	80
Strongyloides Antibodies	STGA	B	10 days	80
Sulpiride	SULP	B ⁴	4 days	131
Superoxide Dismutase Inhibitor	SODI	A/H	5 days	33
Suppression with steroid, IVIg and intralipin, NK (CD69) cell assay, TH1/TH2 cytokines	NCIT	H H H [*]	Send Mon-Thurs only	53
Swab (Ear)	EARS	STM	2-4 days (Culture) 8-9 days (Fungal) – same swab	41
Swab for Culture (Any Site)	SWAB	STM [†]	2-4 days	41,66
Synacthen Stimulation Test	SYNA	By appointment only	1 day	129
Synovial Fluid (for microscopy and culture)	FLU2	SC ^{†††}	14 days	41
Syphilis by PCR (chancres)	SYPS	PCR	5 days	66
Syphilis IgG/IgM	SERJ	B	4 hours	66, 80
T Regulatory Cells	25RF	H	3 days	52
T3	T3	B	4 hours	50
T3 (Reverse)	RT3	B ^{7,37}	10 days	50
Tacrolimus/Prograf (FK506)	FK5	A ⁴	1-2 days	131
Taipan Snake Venom Time	TTVT	C ¹⁸	1 week	37
TB (pleural fluid)	TBCU	SC	up to 8 weeks	41
TB Culture	SPU2	SC	up to 8 weeks	41
TB Culture (Urine)	TBUR	3 x EMU	up to 8 weeks	41
TB Quantiferon®-TB Gold*	TBQ4	Special tubes or H ¹	3 days	80
TB Slopes – Confirmation and Sensitivity	TBSL	TB slope (LJ medium-green) ⁶	up to 8 weeks	41
TDL Tinies & Self-collection samples				146-151
Tegretol (Carbamazepine)	CARB	B	4 hours	131
Teicoplanin Assay	TEIC	B	5 days	129

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
Temazepam	TEMA	 ⁴	4 days	131
Testicular Autoantibodies	TAB		2 days	80
Testicular Tumour Profile	TTP		4 hours	99
Testosterone	TEST		4 hours	50
Testosterone (Bioavailable)	BTES		5 days	50
Testosterone (Free)	FTES		3 days	50
Tetanus Antibody	TETA		5 days	80,89
TH1/TH2 Cytokine Profile	1TH2	   *	Send Mon-Thurs only	53
TH1/TH2 Cytokine Ratio	6RF	   ⁵	1 week	52
TH1/TH2 Intracellular Cytokine Ratios with IVIG, Prednisolone	20RF	   ⁵	1 week	52
TH1/TH2 Intracellular Cytokine Ratios with Prednisolone	22RF	   ⁵	1 week	52
TH1/TH2 Intracellular Cytokine Ratios with IVIG	21RF	   ⁵	1 week	52
Thalassaemia Screen	HBEL		4 days	38
Thallium (Blood)	THAL	 / 	1 week	156
Thallium (Urine)	URTH	RU	1 week	156
Theophylline	THEO		4 hours	131
Thiopurine Methyl Transferase	TPMT	 ⁵	5 days	33
Thrombin Time	THRO	 ¹⁸	4 hours	36
Thrombotic Risk Profile	PROP	      ¹⁸	5 days	37, 39, 120, 128
Thyroglobulin Abs	TGAB		1 day	51
Thyroglobulin Assay	TGA		1 day	51
Thyroid Abs (incl. Thyroglobulin + Thyroid Peroxidase Abs)	THAB		1 day	51,80
Thyroid Peroxidase Antibodies/Anti TPO	TPEX		1 day	51,80
Thyroid Profile 1	TF		4 hours	51,54
Thyroid Profile 2	TF2		2 days	51,54
Thyroid Profile 3	TF3		4 hours	51,54
Thyroxine (T4)	T4		4 hours	51
Thyroxine Binding Globulin	TBG	 (Frozen)	10 days	51
Timothy Grass Components	ZZ19		2 days	141
Tissue for culture	TISS	Tissue sample	up to 14 days	41
Tissue Polypeptide Antigen	TPA		1 week	33
Tissue Transglutaminase IgA (Coeliac)**	TAA		2 days	80
Tissue Transglutaminase IgG	TAG		5 days	80
Tobramycin Assay (Provide Clinical Details)	TOBR		3 days	129
Toluene (Blood)	TOL	J	10 days	156
Toluene (Urine)	UTOL	RU	10 days	156
Topiramate (Topamax)	TOPI	 ⁴	4 days	131
Torch Screen	TORC		2 days	80, 97-98
Total Acid Phosphatase	APT		5 days	33

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index


























TEST	CODE	SAMPLE REQ	TAT	PAGE
Total Bile Acid/Bile Salts	BILS		1 week	33
Total IgE	IGE		1 day	33, 80, 136
Total Immune Function Evaluation	TIE	 +  ^{5,10}	7 days	80
Toxocara Antibodies (IgG)	TFAT	 ⁹	5 days	80
Toxoplasma Antibodies (IgG+IgM)	TFAM	 ⁹	4 hours	80, 86
Toxoplasma Antibody Full Evaluation (IgM, Dye Test, IgG Avidity)	TDYE	 ⁹	10 days	81
Toxoplasma by PCR	TXAG		5 days	81
TPPA	TPPA		2 days	66, 81
Trace Metal (Blood) Profile	TRAC	   	7-10 days	155
Transferrin	TRAN		1 day	33
Transferrin Electrophoresis	TREL		2 weeks	33
Trichinella Serology	TRIC		5 days	81
Trichloroacetic Acid (Urine)	UTCA	RU	5 days	156
Trichomonas vaginalis by PCR	TVPC	FCRU/PCR/TPV	2 days	66, 160
Triglycerides	TRI		4 hours	33
Trimethylaminuria (Fish Odour Syndrome)	FOS	PU	6 weeks	33
Trimipramine	TRIM		5 days	131
Tropical Screen (from 6 weeks post-travel)	TROP	  ^{9,14}	10 days	86-87
Tropomyosins	ZZ31		2 days	141
Troponin T (High sensitive)	TROT		4 hours	33
Trypanosome (Chagas) Antibodies	CHGA	 ^{9,14}	10 days	81
Tryptase	STRY		2 days	33, 134
TSH	TSH		4 hours	51
TSH-Receptor Antibodies	TSI		4 days	51, 81
Tularaemia Antibodies	TULA	 ¹⁴	5 days	81
Tumour Necrosis Factor – Alpha	TNF	 (Frozen) ⁴	2 weeks	33
Uni Parental Disomy (UPD) – parents and child – <i>Specify chromosome</i>	Specify type	 ^{9,12}	5 days	120
Urate (Uric acid)	UA		4 hours	33
Urea	UREA		4 hours	33
Urea (Urine)	UURE	CU	4 hours	33
Urea and Electrolytes	U/E		4 hours	33, 34
Urea Electrolytes (Urine)	UELE	CU	4 hours	33
Ureaplasma urealyticum by PCR	UGEN	FCRU/PCR/TPV	2 days	66, 160
Uric Acid (Serum)	UA		4 hours	33
Uric Acid (Urine)	UURI	CU	4 hours	33
Urinary Methyl Histamine	UHIT	RU (Frozen)	2 weeks	81
Urine (Microscopy Only)	UMIC	RU	1 day	41
Urine Cytology (Urine cytology containers available from TDL Supplies)	URCY	Urine (30mls) ²¹	2 days	165
Urine EtG (Ethyl glucuronide)	ETG	RU	1 week	153
Urine for Microscopy and Culture	UCEM	MSU ^{††††}	1-2 days	41
Urine Free Light Chains	UFLC	RU	1 week	33

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
Urine Organic Acids	UORG	RU (Frozen)	3 weeks	33
Urine Steroid Screen (Steroid Hormones)	USTE	CU or RU ⁹	2 weeks	33
Urine Sugar Chromatography	UCRO	RU (Frozen)	3 weeks	33
Urobilinogen (Urine)	UURO	RU	1 day	33
Urticaria Test (Histamine Releasing)	CURT	B	10-14 days	81
Vaginitis/BV Profile using Culture & PCR Swab	STD8	PCR/STM	3 days	66, 75
Valium (Diazepam)	DIAZ	A	7 days	131
Valproic Acid (Epilem)	VALP	B	4 hours	131
Vancomycin Hydrochloride	VANC	B	4 hours	129
Varicella Zoster – DNA	VZPC	A	5 days	97
Varicella Zoster Antibodies (IgG)	VZOS	B	1 day	89, 97
Varicella Zoster Antibodies (IgM)	VZOM	B	1 day	89, 97
Vascular Endothelial Growth Factor	VEGF	B	14 days	81
VDRL (RPR)	RPR	B	2 days	81
Venom Components	ZZ33	B	2 days	141
Very Long Chain Fatty Acids	VLCF	A or H (Frozen) ⁹	4-6 weeks	33
Vigabatrin (Sabril)	VIGA	A	10 days	131
Viral Antibody Screen	VIRA	B B	2 days	97-98
Viral Eye by PCR	VPE	PCR	3 days	97-98
Viral Respiratory RNA Screen by PCR	VPR	PCR or as specified on the form	2 days	97
Viral Skin/Mucosa by PCR	VPSK	PCR	2 days	97-98
Viscosity (Plasma)	VISC	A ⁴	3 days	37
Vitamin A (Retinol)	VITA	B	5 days	144
Vitamin B (Functional)	FUNC	A A or H ¹³	5 days	144
Vitamin B Profile	VBP	A A B	5 days	143-144
Vitamin B1 (Thiamine)	VIT1	A	5 days	144
Vitamin B2 (Riboflavin)	VIB2	A	5 days	144
Vitamin B3 (Nicotinamide)	VIB3	B	5 days	144
Vitamin B5 (Pantothenic Acid)	VB5S	B	5 days	144
Vitamin B6 (Pyridoxine)	VITB	A	5 days	144
Vitamin B8 (Biotin)	BIOS	B	5 days	144
Vitamin B9 (Folic acid) – Red cell	RBCF	A	2 days	144
Vitamin B9 (Folic acid) – Serum	FOLA	B	1 day	144
Vitamin B12 (Active)	B12	B	1 day	33, 144
Vitamin B12 (Active)/ Red Cell Folate	B12F	A B	2 days	33, 144
Vitamin B12 (Total)	TB12	B	1 day	33
Vitamin C (Active)	VITC	B (Frozen) ⁷	5 days	145
Vitamin D (1, 25 Dihydroxy)	D3	B	5-8 days	145
Vitamin D (25-OH)	VITD	B	4 hours	33, 145
Vitamin E (Alpha Tocopherol)	VITE	B	5 days	145
Vitamin K (Nutritional)	VKN	B ¹³	5 days	145
Vitamin K (With PIVKA II)	VITK	B ¹³	10 days	36

Ensure all specimens and forms are labelled with given Forename, Surname, DOB, Date and Time of collection. Turnaround times are quoted as working days.

Alphabetical test index

TEST	CODE	SAMPLE REQ	TAT	PAGE
Vitamin Profile 1	VITS	 7	5 days	143,145
Vitamin Profile 2	VIT2	 7,13	5 days	143,145
VLDL Cholesterol	VLDL	 13	1 week	33
VMA	UVMA	PU ¹	5 days	33
Voltage Gated Calcium Channel Antibodies	CCAB		3 weeks	81
Voltage Gated Potassium Channel Antibodies	VPCA		3 weeks	81
Von Willebrand Profile	FVWF	 4,12	5 days	37,39
Von Willebrands Multimers	VWM	 18	3 months	37
Wall Pellitory Components	ZZ20		2 days	141
Walnut Components	ZZ34		2 days	141
West Nile Virus Abs	WNV		2 weeks	97
Wheat Components	ZZ21		2 days	141
Whooping Cough (Pertussis) Antibodies	PERS		5 days	81
Whooping Cough (Pertussis) by PCR	PERP	Prenasal (posterior nasopharynx) swab	5 days	81
Wolf-Hirschhorn Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS / AF /  ⁹	5-15 days	121
Wolf-Hirschhorn Syndrome – BOBs only	PBOB	CVS / AF /  ⁹	5 days	121
Xanthine – Blood	XANB		2 weeks	156
Xylene – Urine	UXYL	RU ³⁰	2 weeks	156
Xylose Tolerance Test	XTT	J ¹	7 days	144
Y chromosome microdeletions – AZFa + AZFb + AZFc + SRY	YDEL	 ⁹	5 days	121
Yellow Fever Antibodies	YELL	 ^{9,14}	10 days	81
Yersinia Antibodies	YERS		4 days	81
Zika Abs IgM and IgG – Antibody detection from 15 days	ZKAB		5 days	81, 86, 97
Zika RNA by PCR in Semen	ZIKS	Semen	5 days	86, 97
Zika RT PCR – Window of detection from 1-14 days from onset of symptoms	ZIKU	RU	5-7 days	86
Zika RT PCR – Window of detection from 1-7 days from onset of symptoms	ZIKA		5-7 days	86
Zinc (Serum/Plasma)	ZINC		1 day	144,155
Zinc (Urine)	URZN	CU	5 days	144,156
Zinc (Whole Blood)	RBCZ	 or 	5 days	144
Zinc Protoporphyrin	ZNPR	 ¹³	5 days	156
Zygosity testing – comparative DNA profile	DNAC	 (From each twin and both parents) ⁹	5 days	121

TDL Referral Laboratories

For certain specialist tests TDL has developed a selected network of TDL Group and Reference Laboratories. These Group or specialist laboratories can be identified by a code assigned to reports. The quality of these laboratories is recognised by UKAS, or similar accrediting bodies for the laboratories outside the UK.

Addenbrooke's Hospital – BGU and Immunology	Health & Safety Laboratory
Affinity Biomarker Labs	HFL Sport Science (LGC Group)
Alder Hey Children's NHS Foundation Trust – Biochemistry Department	Homerton University Hospital – Department of Clinical Biochemistry
Analytical Services International Ltd, St George's University of London – Forensic Toxicology Service	Igenomix UK
Animal and Plant Health Agency – Veterinary labs	Imperial College Healthcare NHS Trust – Charing Cross Hospital, Chemical Pathology Department
Antenatal Screening Service, Wolfson Institute of Preventive Medicine, Barts and The London School of Medicine and Dentistry	Imperial College Healthcare NHS Trust – Charing Cross Hospital, Infection and Immunity Department
Bio Predictive	Imperial College Healthcare NHS Trust – Charing Cross Hospital, Medical Oncology
Biodesix, Inc.	Imperial College Healthcare NHS Trust – Hammersmith Hospital, Molecular Endocrinology
Biolab Medical Unit	Imperial College Healthcare NHS Trust, St Mary's Hospital – Virology Department
Bioscientia	Independent Histopathology Services
Birmingham Children's Hospital NHS Foundation Trust – Clinical Chemistry	Institute of Aquaculture – University of Stirling
Brucella Reference Unit – Liverpool Clinical Laboratories, Royal Liverpool and Broadgreen Hospital	Institute of Neurology – Neurogenetics Unit
Cambridge Clinical Laboratory	Instituto Bernabeu Biotech
Cambridge Life Sciences	King's College Hospital – HMDC Laboratory for Molecular Haemato-Oncology
Cambridge Nutritional Science Ltd	Labor Augsburg MVZ GmbH
Cardiff and Vale University Health Board – The Analytical Toxicology Department	Latis Scientific
Cerba	London School of Hygiene & Tropical Medicine – Diagnostic Parasitology Lab
Chelsea and Westminster Hospital NHS Foundation Trust	Matrix Diagnostics
CNC Forensic Toxicology Service LTD	Mayo Clinic Laboratories
Douglass Hanly Moir Pathology	Meningococcal reference unit (Men RU)
Epsom and St Helier University Hospital NHS Trust – Biochemistry Department	Manchester – Manchester Royal Infirmary
Epsom and St Helier University Hospital NHS Trust – Immunology Department	Micropathology Ltd
Eurofins – Biomnis, France	National Blood Service – Colindale, Red Cell Immuno Haematology Department
Great Ormond Street Hospital – Department of Chemical Pathology	NHS Blood and Transplant – Birmingham
Great Ormond Street Hospital – Enzyme Unit, Chemical Pathology	NHS Blood and Transplant – H & I Laboratory
Great Ormond Street Hospital – Immunology Department	NHS Blood and Transplant – Tooting
Great Ormond Street Hospital – Neurometabolic Unit	Norfolk and Norwich University Hospital NHS Foundation Trust – SAS Metabolic Bone Laboratory
Guildford RSCH Trace Element Laboratory, SAS Trace Element Centre	Oxford University Hospital NHS Foundation Trust – Churchill Hospital
HCA Healthcare UK – HCA Laboratories	Pathcare

TDL Referral Laboratories

PHE – Bacteriology Reference Department (BRD), Colindale

PHE – Virus Reference Department (VRD) – Colindale

PHE Mycology Reference Laboratory –

PHE South West Laboratory, Southmead Hospital, Bristol

PHE National Mycobacterium Reference Service
National Infection Service, Colindale

PHE Rare and imported pathogens laboratory –
Porton Down

Queens University Hospital, Belfast –
Institute of Clinical Science

Radboud University Nijmegen Medical Center

Randox Health – London

Reflab – Copenhagen

Rosalind Franklin University

Royal Berkshire Hospital NHS Foundation Trust –
Clinical Biochemistry

Royal Surrey County Hospital – SAS Peptide Hormone Section

Sandwell and West Birmingham NHS Trust –
City Hospital Birmingham, Clinical Biochemistry Department

SCSA Diagnostics

Sheffield Children's NHS Trust – Clinical Chemistry

Sheffield Teaching Hospital NSH Foundation Trust –
Protein Reference Laboratory Unit and Immunology
Department

Southmead Hospital – Antimicrobial Reference Laboratory,
Bristol

St George's University Hospital NHS Foundation Trust –
Cell Marker Department

SYNLAB Budapest Diagnostic Center,
Genoid Molecular Diagnostic Laboratory

SYNLAB Laboratory Service – Abergavenny

The European Laboratory of Nutrients

The Leeds Teaching Hospital NHS Trust –
Endocrinology Laboratory (including SAS Steroid Centre),
Department of Specialist Laboratory Medicine,
ST James University Hospital

The Leeds Teaching Hospitals NHS Trust –
Mycology Reference Centre

The Newcastle upon Tyne Hospitals – Royal Victoria Infirmary

The Royal Marsden Hospital
– Department of Haematology/Oncology

Toxoplasma Reference Unit, Public Health Wales
Microbiology ABM, Singleton Hospital – Swansea

Trace Laboratories Ltd

UCL Great Ormond Street Institute of Child Health

UCL Queen Square Institute of Neurology –
Department of Neuroimmunology

University Hospital Birmingham NHS Foundation Trust –
Heartlands Hospital

University Hospital of Wales – Immunology Department

Viapath – Guy's Hospital, Biochemistry Genetics Laboratory

Viapath – King's College Hospital, Clinical Biochemistry

Viapath – St Thomas' Hospital Haemophilia Centre

Viapath – St Thomas' Hospital Immunohistology

Viapath – St Thomas' Hospital Purine Research Laboratory

GROUP LABORATORIES

Royal Free London NHS Foundation Trust – Haemostasis

University College London Hospitals NHS Foundation Trust
(UCLH) – Cytology

University College London Hospitals NHS Foundation Trust
(UCLH) – Hospital for Tropical disease

University College London Hospitals NHS Foundation Trust
(UCLH) – Molecular Virology

University College London Hospitals NHS Foundation Trust
(UCLH) – Special Chemistry

TDL Genetics Referral Laboratories

All Wales Medical Genetics Service	Progenika Biopharma Grifols
Anthony Nolan, Histocompatibility and Immunogenetics	Protein Reference Unit & Immunology Department – Sheffield Protein Unit
Asper Biotech	Purine Research Laboratory – St Thomas' Hospital
Bioscientia GmbH	Royal Marsden – Haemato-Oncology Unit
Bristol Genetics Laboratory (North Bristol NHS Trust)	Sheffield Diagnostic Genetics Service
CentoGene	SIHMDS – Cytogenetics Laboratory, Great Ormond Street Hospital
DiaGenom GmbH	South East Scotland Genetics Service (NHS Lothian)
Douglass Hanly Moir Pathology	South West Thames Regional Genetics Service
East Scotland Regional Genetics Service (NHS Tayside)	SYNLAB Budapest Diag Center
Exeter Clinical Laboratory – Department of Molecular Genetics	The Leeds Genetics Laboratory
Fulgent Diagnostics	Viapath Analytics LLP
Institute of Neurology, Queen's Square	Wessex Region Genetics Service
International Blood Group Reference Laboratory	West Midlands Regional Genetics Laboratory
London South East Genetics Service	West of Scotland Genetic Service (NHS Greater Glasgow and Clyde)
Medical Genetics Laboratory – Central Manchester University Hospitals NHS Foundation Trust	
Medical Neurogenetics Laboratory LLC	
Micropathology Ltd	
Molecular Genetics Laboratory – Liverpool's Women NHS Foundation Trust	
Molecular Vision Laboratory	
Newcastle Mitochondrial NGC Diagnostic Service	
North East Thames Regional Genetic Service	
North West London Pathology	
North West Thames Regional Genetic Service	
Northern Genetics Service	
Oxford Genetics Laboratory – Oxford University Hospitals	
Prevention Genetics	

Terms & conditions of business from 1st Jan 2021

The definitions which apply to these Terms and Conditions are set out in clause 18.

1 THE SERVICES

- 1.1 These Terms and Conditions will apply to any services that TDL provides to the Client, unless those services are the subject of a separate written agreement signed by TDL and the Client. These Terms and Conditions apply to the exclusion of any other terms presented by the Client or implied by custom or course of dealing.
- 1.2 By submitting a request for any services described in the Laboratory Guide or in any other proposal provided by TDL (an '**Order**'), the Client offers to purchase those services on these Terms and Conditions. A contract between TDL and the Client for the provision of services incorporating these Terms and Conditions (an '**Agreement**') takes effect when TDL confirms acceptance of the Client's Order in writing, logs the relevant Pathology Request in its laboratory information management system, or begins performing the Services (whichever occurs first). Any request for add-on Tests (as described in the Laboratory Guide) constitutes a request for further Services under that Agreement, which TDL may accept or decline.
- 1.3 TDL will provide the Services under the Agreement:
 - 1.3.1 in accordance with Good Industry Practice;
 - 1.3.2 in accordance with the UKAS medical laboratory accreditation standard (ISO 15189); and
 - 1.3.3 using suitably skilled and experienced staff.
- 1.4 TDL will use reasonable efforts to achieve the Test turn-around times quoted in the Laboratory Guide, but does not warrant that it will achieve those times in the case of any particular Sample.
- 1.5 The Laboratory Guide sets out Sample rejection criteria. If the Sample meets those criteria, or if TDL considers that the Sample is otherwise unsuitable for Testing or TDL is unable to conduct the Testing then TDL may decline to carry out the Testing under the Agreement and will be entitled to dispose of the Sample.
- 1.6 As part of its Services TDL will, on request, arrange for collection of Samples from locations within the M25 motorway. Such collection service is included within the price of the Test unless otherwise specified by TDL. Collection of Samples from locations outside the M25 is by special arrangement, and may incur an additional charge. Where collection by TDL has not been requested and agreed, the Client will be responsible, at its own cost, for the transport of Samples to TDL. Where TDL arranges collection of Samples it will use reasonable efforts to achieve the timescales it quotes for collection, but does not warrant that it will achieve those timescales in the case of any particular collection.

- 1.7 TDL may destroy or dispose of a Sample after completing the Testing or on termination of the Agreement, unless otherwise agreed in writing with the Client.

2 PRICE AND PAYMENT TERMS

- 2.1 The fees payable by the Client for the Services will be the most recent price confirmed by TDL to the Client in writing or by telephone prior to the Client submitting its Order. If TDL has not confirmed the price for the Services, the price will be that indicated in the Laboratory Guide.
- 2.2 As at the date of these Terms and Conditions many of TDL's services are VAT exempt. All of TDL's prices are stated exclusive of VAT and where VAT is chargeable on the Services the Client will pay it at the applicable rate.
- 2.3 Invoices are normally issued on a monthly basis, but TDL reserves the right to issue them more frequently. The client will pay TDL's invoices under the Agreement within 30 days of the date of the invoice, without any deduction or set off. At TDL's option interest may be charged on late payment at the statutory rate prescribed from time to time by regulations under the Late Payments of Commercial Debts (Interest) Act 1998. Invoices paid from outside the UK must be paid by either direct bank transfer or by cheque drawn on a UK branch. All payments will be made in pounds sterling.
- 2.4 Without affecting any of its other rights, TDL may suspend provision of the Services if the Client fails to pay an invoice due to TDL.

3 CONFIDENTIALITY

- 3.1 TDL agrees that it will hold and maintain the confidence of:
 - 3.1.1 all information of a confidential nature which is received by TDL from the Client or its patients in connection with the Services; and
 - 3.1.2 all Test results, invoices and other information of a confidential nature issued by TDL to the Client or its patients in connection with the Services, and, save with the Client's consent or as otherwise permitted under this Agreement, will not disclose such information other than to its professional staff, independent consultants and/or persons to whom it has delegated the performance of the Services and who require the information for such purpose. Where TDL has been provided with the details of a patient's private medical insurance in connection with the Services, TDL will be entitled to assume (and the Client so warrants) that both the Client and the patient consent to the disclosure of information relating to that patient to the insurer concerned.

Terms & conditions of business from 1st Jan 2021

- 3.2 The restrictions in clause 3.1 will not apply to information which: (i) was in TDL's possession prior to disclosure by the Client; or (ii) is now or hereafter comes into the public domain other than by default of TDL; or (iii) was lawfully received by TDL from a third party acting in good faith having a right of further disclosure; or (iv) is required by law to be disclosed by TDL; or (v) which is required by a regulatory or accreditation body to be disclosed to it for the purpose of regulating or accrediting the TDL Group.

4 CLIENT RESPONSIBILITIES

- 4.1 Except where TDL obtains the Sample directly from the patient during a home visit or at TDL's patient reception facility, the Client will ensure that the Sample is obtained from the patient, packaged, and labelled in accordance with Applicable Law and good clinical practice.
- 4.2 Except where TDL agrees to arrange transport of the Sample to TDL's laboratory, the Client will ensure that the Sample is transported to TDL's laboratory in accordance with Applicable Law and good clinical practice.
- 4.3 The Client will ensure that all necessary consents and permissions are obtained and all necessary information provided to the patient, which is required under Applicable Law or good clinical practice in order to permit the Testing, the performance of and any other Services, and the use of the Protected Data as contemplated in the Agreement.
- 4.4 The Client will provide TDL with any information reasonably necessary for performing the Services, including by ensuring that the Pathology Request contains sufficient information regarding the Sample, the relevant patient, and the persons to whom the Test results are to be reported, and will ensure that any information the Client provides to TDL in connection with the Services is accurate and complete.

5 LIABILITY

- 5.1 Nothing in the Agreement will limit or exclude liability for death or personal injury caused by negligence or any other liability that cannot be limited or excluded under Applicable Law.
- 5.2 In these Terms and Conditions 'liability' means any liability whether in contract, tort (including negligence), misrepresentation, breach of statutory duty or otherwise, which arises in connection with the Services or under or in connection with any Agreement.
- 5.3 The liability of TDL and the Client will each be limited to £2,000,000 in total. This limit applies per Agreement and in aggregate for all Agreements made in a calendar year.

- 5.4 Neither TDL nor the Client will have any liability for:

- 5.4.1 loss of profit or revenue;
- 5.4.2 loss of anticipated savings;
- 5.4.3 loss of reputation or goodwill; or
- 5.4.4 indirect, special or consequential loss.

- 5.5 TDL will have no liability for any delay or failure in performance of the Services arising from the Client's delay or failure in performing its obligations under clause 4 (Client Responsibilities).

- 5.6 All of the warranties which TDL gives in relation to the Services are expressly set out in these Terms and Conditions. All other warranties, whether implied or express, are excluded from the Agreement where it is lawful to exclude them.

- 5.7 In this clause 5 references to TDL include the members of TDL's Group, and for the purpose of the limit in clause 5.3 the liabilities of TDL and the TDL Group Members will be counted in aggregate. The members of TDL's Group may enforce this clause 5.

6 FORCE MAJEURE

If the performance of any obligation under the Agreement (except for an obligation to pay) is prevented, restricted or interfered with by reason of circumstances beyond the reasonable control of that party obliged to perform it (a 'Force Majeure Event'), the party so affected will be excused from any resulting failure or delay in performance, and the time for performance will be extended by an amount of time equal to the duration of the Force Majeure Event. The party so affected will use reasonable endeavours to mitigate the effect of the Force Majeure Event on its performance of its obligations. If the Force Majeure Event delays or prevents performance of a party's obligations for more than three months, either party may terminate the agreement on written notice to the other.

7 DATA PROCESSOR AND DATA CONTROLLER

- 7.1 When TDL processes Protected Data on behalf of the Client in providing the Services the parties agree that the Client will be the data controller and TDL will be the data processor. The Annex to these Terms and Conditions sets out when TDL processes Protected Data on behalf of the Client. Clause 16 describes the circumstances where TDL will use Protected Data on its own behalf as data controller.
- 7.2 When TDL processes Protected Data as the data processor, clauses 8 to 15 will apply in relation to the Protected Data. Where TDL processes Protected Data as data controller, clause 16 will apply instead.

Terms & conditions of business from 1st Jan 2021

- 7.3 The Client will comply with the Data Protection Laws in relation to the Protected Data, and ensure that all instructions given by it to TDL in respect of Protected Data will at all times be in accordance with Data Protection Laws.

8 DATA PROCESSING INSTRUCTIONS

- 8.1 When TDL processes Protected Data as the data processor, TDL will comply with the obligations of data processors under Data Protection Laws.
- 8.2 Unless required to do otherwise by Applicable Law, TDL will (and will take steps to ensure each person acting under its authority will) process the Protected Data only in accordance with the Client's documented instructions as set out in the Order, pursuant to the Terms & Conditions, and in the Annex (the '**Processing Instructions**').
- 8.3 If Applicable Law requires TDL to process Protected Data other than in accordance with the Processing Instructions, TDL will notify the Client of any such requirement before processing the Protected Data (unless Applicable Law prohibits TDL from doing so).
- 8.4 TDL will promptly inform the Client if TDL becomes aware of a Processing Instruction that, in TDL's opinion, infringes Data Protection Laws. TDL will have no liability for any processing in accordance with those Processing Instructions after giving the notice. TDL's obligations under this clause 8.4 do not limit the Client's obligations under clause 7.3.

9 DATA SECURITY MEASURES

In relation to the processing of the Protected Data, TDL will implement and maintain, at its cost and expense, appropriate technical and organisational measures to ensure for the Protected Data a level of security appropriate to the risks presented by the processing, taking into account the state of the art, the cost of implementation and the nature, scope, context and purpose of the processing of the Protected Data as well as the risk of varying likelihood and severity of the rights and freedoms of natural persons.

10 USING STAFF AND OTHER PROCESSORS

- 10.1 TDL will not engage any data processor to process the Protected Data on the Client's behalf (a '**Sub-Processor**') without the Client's authorisation of that specific Sub-Processor. The Client will not unreasonably withhold, condition or delay such consent. By accepting these Terms and Conditions the Client authorises the appointment of the Authorised Sub-Processors.
- 10.2 TDL will ensure that each Sub-Processor is appointed under a written contract containing materially the same obligations as clauses 8 to 15.

- 10.3 TDL will ensure that all persons authorised to process Protected Data are subject to a binding obligation to keep the Protected Data confidential (except where disclosure is required in accordance with Applicable Law, in which case TDL will, where practicable and not prohibited by Applicable Law, notify the Client of any such requirement before such disclosure).

11 ASSISTANCE WITH THE CLIENT'S COMPLIANCE AND DATA SUBJECT RIGHTS

- 11.1 Taking into account the nature of the processing, TDL will implement and maintain reasonable measures to assist the Client to respond to the Data Subject Requests relating to the Protected Data that TDL processes on the Client's behalf. TDL will refer such Data Subject Requests it receives to the Client promptly, and in any event within five Business Days of receipt of the request.
- 11.2 TDL will provide such assistance as the Client reasonably requires (taking into account the nature of processing and the information available to TDL) to the Client in ensuring compliance with the Client's obligations under Data Protection Laws with respect to: (i) security of processing, (ii) data protection impact assessments (as such term is defined in Data Protection Laws), (iii) prior consultation with the relevant regulator regarding high risk processing, (iv) and notifications to the regulator and/or communications to data subjects by the Client in response to any Personal Data Breach. The Client will pay TDL's charges for providing the assistance in this clause 11, such charges to be calculated on a time and materials basis at TDL's applicable daily or hourly rates in force from time to time.

12 INTERNATIONAL DATA TRANSFERS

The Client agrees that TDL may transfer Protected Data to countries outside the United Kingdom for the purpose of providing the Services, provided all transfers by TDL of Protected Data to such recipients are in accordance with such safeguards or other mechanism(s) for transfers of personal data as may be permitted under Data Protection Laws from time to time. The Client agrees that TDL may implement such safeguards by entering into standard data protection clauses authorised under the Data Protection Laws, which TDL may do as agent on behalf of the Client. The provisions of clauses 8 to 15 (inclusive) will constitute the Client's instructions with respect to transfers in accordance with clause 8.2.

Terms & conditions of business from 1st Jan 2021

13 RECORDS, INFORMATION AND AUDIT

- 13.1 TDL will maintain, in accordance with Data Protection Laws binding on TDL, written records of all categories of processing activities carried out on behalf of the Client.
- 13.2 TDL will, in accordance with Data Protection Laws, make available to the Client such information as is reasonably necessary to demonstrate TDL's compliance with its obligations as a data processor under these Terms and Conditions and the Data Protection Laws, and allow for and contribute to audits, including inspections, by the Client (or another auditor mandated by the Client) for this purpose, subject to the Client:
 - 13.2.1 giving TDL reasonable prior notice of such information request, audit and/or inspection being required by the Client;
 - 13.2.2 ensuring that all information obtained or generated by the Client or its auditor(s) in connection with such information requests, inspections and audits is kept strictly confidential (save for disclosure to the relevant regulator or as otherwise required by Applicable Law);
 - 13.2.3 ensuring that such audit or inspection is undertaken during normal business hours, with minimal disruption to TDL's business, the Sub-Processors' business and the business of other customers of TDL.

14 BREACH NOTIFICATION

TDL will, without undue delay notify the Client of the Personal Data Breach involving the Protected Data, and provide the Client with details of the Personal Data Breach.

15 DELETION OR RETURN OF PROTECTED DATA AND COPIES

TDL will, at the Client's written request, either delete or return all of the Protected Data to the Client in such form as the Client reasonably requests within a reasonable time after the end of the provision of the relevant Services related to processing, and delete existing copies (unless storage of any data is required by Applicable Law and, if so, TDL will inform the Client of any such requirement). Where TDL will process that Protected Data as data controller under clause 16, TDL may retain the Protected Data.

16 PROTECTED DATA THAT TDL PROCESSES AS A DATA CONTROLLER

- 16.1 TDL may process Protected Data as data controller in the circumstances and for the purposes set out in TDL's Privacy Notice. In particular TDL may:

- 16.1.1 retain and submit Protected Data to a Health Authority in the United Kingdom for the purposes of a Public Health Programme operated by that Health Authority, or to regulator for the purpose of complying with regulatory obligations; and
- 16.1.2 retain and process Protected Data in its laboratory records in order to meet the requirements of the UKAS medical laboratory accreditation standard (ISO 15189) and implement the guidelines of the Royal College of Pathologists for the retention and storage of pathological records and specimens.
- 16.3 When TDL processes Protected Data to provide Harmony® Non-Invasive Prenatal Tests, TDL does so as a data controller.
- 16.4 When TDL processes personal data on its own behalf as data controller, it will do so in accordance with the obligations of data controllers under Data Protection Laws and with the applicable terms of the Agreement.

17 GENERAL

- 17.1 Dispute resolution
 - 17.1.1 If any dispute arises relating to this Agreement or any breach or alleged breach of this Agreement, the parties will make a good faith effort to resolve such dispute without recourse to legal proceedings. If, notwithstanding such good faith efforts, the dispute is not resolved either party may submit the dispute to the jurisdiction of the English Courts.
 - 17.1.2 Except to the extent clearly prevented by the area of dispute, the parties will continue to perform their respective obligations under this Agreement while such dispute is being resolved.
- 17.2 Variation
 - 17.2.1 TDL may amend these Terms and Conditions by updating the Laboratory Guide and providing the Client with a copy of the update or publishing it on TDL's website. Such amendments will only apply to an Order submitted after the date of the update, and the Client will be deemed to accept those amendments by submitting an Order after that date.
 - 17.2.2 Except as set out in clause 17.2.1, any amendments to this Agreement will not be effective unless in writing and signed by an authorised signatory on behalf of each of the parties. The terms of this Agreement may be varied by agreement of the parties but without the consent of any third party whether or not the rights of such third party are affected by such variation. The Client will not unreasonably withhold, delay or condition its agreement to any variation to this Agreement requested by TDL in order to ensure the Services and TDL (and each Sub-Processor) can comply with any change in Applicable Laws.

Terms & conditions of business from 1st Jan 2021

17.3 Rights and waiver

All rights granted to either of the parties will be cumulative and not exhaustive of any rights and remedies provided by law. The failure of either party to enforce (or delay in enforcing) at any time for any period any one or more of the terms of this Agreement will not be a waiver of such term or of the right of such party at any time subsequently to enforce all the terms of this Agreement.

17.4 Severability

If any provision of this Agreement is or becomes invalid, illegal or unenforceable in any respect under any law, the validity, legality and enforceability of the remaining provisions will not be in any way affected.

17.5 Assignment

TDL may assign or sub-contract the performance of this Agreement (in whole or in part) or any one or more of the Tests to be performed hereunder to suitably accredited laboratories including those listed in the Laboratory Guide. The Client may not assign this Agreement or any of its rights or obligations hereunder without the prior approval of TDL.

17.6 Relationship of the parties

It is acknowledged and agreed that TDL and the Client are independent contractors and nothing in this Agreement will create or be construed as creating a partnership or (except as provided in clause 12 and the Annex) a relationship of agent and principal between the parties. The Client acknowledges and agrees that, in requesting Services from TDL, it is not acting as agent for any patient or patients to which the Services relate.

17.7 Notices

All notices given under this Agreement will be in writing and will be delivered by hand or sent by prepaid first class post or by prepaid first class recorded delivery or by email transmission. All notices will be delivered at or sent, in the case of TDL, to The Halo Building, 1 Mabledon Place, London WC1H 9AX, email notices@tdlpathology.com and, in the case of the Client to the address and/or email address set out in the Order (or such other address as that party will notify in writing to the other for this purpose). A notice sent by post will be deemed to be served at 9.00 am on the second business day following the date of posting; a notice sent by email transmission will (provided the sender receives no error message indicating that delivery has been unsuccessful) be deemed to have been served at the time it is transmitted if transmitted within business hours (9.00 am to 6.00 pm) on a business day or, if transmitted outside such business hours on a business day or on a day which is not a business day as soon thereafter as such business hours commence.

17.8 Entire agreement

The Agreement is set out in the Order and these Terms and Conditions, which together set out the entire contract between the Client and TDL relating to their subject matter. In the event of a conflict between the Order and these Terms and Conditions, the Terms and Conditions will take priority. Each party acknowledges that it has not entered into the Agreement in reliance on, and will have no remedies in respect of, any representation or warranty that is not expressly set out in the Agreement except in the case of fraudulent misrepresentation.

17.9 Third parties

The Agreement is not intended to create any rights for, nor be enforceable by, any third party except as set out in clause 5.

17.10 Governing law

The Agreement and any dispute arising out of or in connection with it (including non-contractual disputes and claims) will be governed by and construed in accordance with English law and each of the parties submits to the exclusive jurisdiction of the English Courts.

18 INTERPRETATION

18.1 In these Terms and Conditions and the Annex:-

'Agreement' has the meaning given in clause 1.2;

'Annex' means the annex to the Terms and Conditions;

'Applicable Law' means the laws, regulations, judgments, binding on the relevant party, as amended from time to time;

'Authorised Sub-Processors' means:

a) Health Service Laboratories LLP and any other member of the TDL Group which provides the applicable Test or Service;

b) accredited specialist centres for onward referral of esoteric assays as identified in the TDL Laboratory Guide;

c) persons who provide information technology services that TDL uses in the course of providing the Services.; and

d) any Sub-Processor referred to in the Annex;

'Client' means the person or organisation requesting Services from TDL and for whom TDL has agreed to provide the Services;

'controller', 'data subject', 'personal data', 'process' and 'processor' have the meanings given to those terms in Data Protection Laws;

Terms & conditions of business from 1st Jan 2021

'Data Protection Laws' means the UK GDPR, the Data Protection Act 2018, and any other Applicable Law having effect in the United Kingdom concerning privacy or the use of personal data;

'data subject' and 'personal data' have the meaning given to those terms in Data Protection Laws;

'Data Subject Request' means a request made by a data subject to exercise any rights of data subjects under Data Protection Laws;

'Good Industry Practice' means the standard of skill and care reasonably to be expected from a professional provider of the Services;

'Group' in respect of any undertaking, means such undertaking and its group undertakings ('undertaking' and 'group undertaking' having the meanings given in the Companies Act 2006);

'Health Authority' means (i) a department of the UK government or of a devolved administration, (ii) an executive agency of such department, or (iii) a body exercising statutory functions in relation to public health in the UK or any part of the UK;

'Laboratory Guide' means TDL's Laboratory Guide current at the time the Client submits the Order, as supplied to the Client or, if not so supplied, available on request from TDL, including any updates or supplements issued by TDL;

'Order' has the meaning given in clause 1.2;

'Pathology Request' means an Order requesting Testing;

'Personal Data' has the meaning given to that term in Data Protection Laws;

'Personal Data Breach' means any breach of security leading to the accidental or unlawful destruction, loss, alteration, unauthorised disclosure of, or access to, any Protected Data;

'Privacy Notice' means TDL's detailed Privacy Notice available at tdlpathology.com;

'processing' has the meanings given to that term in Data Protection Laws (and related terms such as process have corresponding meanings);

'Processing Instructions' has the meaning given to that term in paragraph 8.2;

'Protected Data' means personal data provided to TDL by the Client or a third party on the instructions of the Client, or collected or generated by TDL in the course of the Services;

'Public Health Programme' means a programme administered by a Health Authority to monitor or analyse health data for the purpose of public health or for statistical, scientific or research purposes in the public interest;

'Sample' means a sample provided by the Client to TDL for Testing;

'Services' means the services to be provided under the Agreement;

'Sub-Processor' has the meaning given in clause 10.1;

'TDL' means The Doctors Laboratory Limited or such other member of the TDL Group as has agreed to provide the Services;

'TDL Group' means The Doctors Laboratory Limited and its Group and Health Service Laboratories LLP and its Group;

'Test' means a laboratory test to be carried out by TDL on a Sample, and 'Testing' means the process of conducting that Test and reporting the results;

'UKAS' means the United Kingdom Accreditation Service, or any successor to it.

- 18.2 References to the singular include the plural and vice versa.
- 18.3 Clause headings and paragraph headings are for ease of reference only and are not part of these Terms and Conditions for the purpose of construction.
- 18.4 References to paragraphs are to paragraphs of the Annex.
- 18.5 Words following the terms 'including', 'include', 'in particular', 'for example' or any similar expression shall be construed as illustrative and shall not limit the sense of the words, preceding those terms.
- 18.6 The Annex is incorporated into these Terms and Conditions.

Terms & conditions of business from 1st Jan 2021

ANNEX

1 Subject matter and nature of processing

- 1.1 TDL processes Protected Data as data processor on behalf of the Client:
 - 1.1.1 in the case of Testing, when TDL receives a Pathology Request and Sample and processes the corresponding Protected Data to carry out the Test and report the Test results in accordance with the Client's documented instructions;
 - 1.1.2 when TDL carries out the Client's 'fee to patient' instructions, as described below; and
 - 1.1.3 in the case of any other Services, when TDL is required to process the Protected Data on the Client's behalf to fulfil the Client's instructions.
- 1.2 The subject matter and nature of TDL's processing of the Protected Data are:
 - 1.1.1 pathology samples and test results for the purpose of providing clinical pathology services;
 - 1.1.2 information about clinicians who order pathology tests, for the purposes of reporting the test results to the Client;
 - 1.1.3 information about a patient's health insurance for the purposes of administering payment for the Services; and
 - 1.1.4 billing information for a patient where the Client has asked TDL to direct TDL's invoice to the patient.

2 Duration of processing

The duration of the processing is the time necessary to carry out the Services.

3 Types of personal data

- 3.1 The Protected Data comprise the following types of personal data:
 - 3.1.1 Name
 - 3.1.2 Gender
 - 3.1.3 Date of birth
 - 3.1.4 Address
 - 3.1.5 Identity numbers assigned by TDL or the Client
 - 3.1.6 Types of pathology tests conducted
 - 3.1.7 Results of pathology tests
 - 3.1.8 Health insurance policy details
 - 3.1.9 Billing information
 - 3.1.10 The types of data referred to in the TDL Laboratory Guide

4 Categories of data subjects

The Protected Data concerns patients in respect of whom TDL conducts pathology tests, and clinicians who request pathology tests.

5 Reporting pathology test results

- 5.1 TDL will report Test results using the method selected by the Client from the range of options offered by TDL or, if no method is selected by the Client, using a method selected by TDL from that range of options.
- 5.2 TDL will report the Test results using the contact details supplied to TDL in the relevant section of the Pathology Request. The Client will be responsible for ensuring that those contact details are correct.
- 5.3 Where TDL supplies Test results electronically it will ensure that the results are supplied in the format selected by the Client (from the range of options offered by TDL) and are supplied to the address indicated when the Client selects electronic results reporting. The Client will be responsible for ensuring that the selected format is compatible with the Client's information systems and for making the results available to the users of those systems.

6 Fee to patient

Where the Client selects the 'fee to patient' option in a Pathology Request Form, the Client instructs TDL to seek payment from the patient of the fees owed by the Client in respect of that test. The Client confirms that the patient has agreed with the Client to pay those fees to TDL for the Client. The Client instructs TDL to recover the fees by invoicing the patient using the personal data provided by the Client. The Client instructs TDL on the Client's behalf to appoint debt collectors to recover the fees from the patient if the patient does not pay the invoice by the date payment falls due. The Client authorises TDL to appoint those debt collectors as Sub-Processors in accordance with clauses 8 to 15.



THE DOCTORS
LABORATORY

Antenatal Screening Service for Down's, Edwards & Patau Syndromes and Open Neural Tube Defects

First ☐ Second ☐ Trimester (please tick as required)

Weeks 11-13 ☐ Weeks 14-21 (16 ideal) ☐ Name of Requesting Doctor: _____

MATERNAL SCREENING FOR DOWN'S SYNDROME AND NEURAL TUBE DEFECTS

If you have a query with completing this form, please telephone the Referrals Dept at The Doctors Laboratory on 020 7307 7373

PATIENT DETAILS

Surname:	<input type="text"/>	Hospital No.:	<input type="text"/>
Forename:	<input type="text"/>	Date of birth:	<input type="text" value="DD"/> <input type="text" value="MM"/> <input type="text" value="YY"/>
NHS No.:	<input type="text"/>	Post code:	<input type="text"/>

CLINICAL DETAILS (To be completed by Midwife or Doctor)

First day of Last Menstrual Period (LMP)	<input type="text" value="DD"/> <input type="text" value="MM"/> <input type="text" value="YY"/>	Does the patient have Insulin dependent diabetes? (no=0, yes=1)	<input type="checkbox"/>
Vaginal bleed in the last 7 days? (no=0, yes=1) If yes please see overleaf	<input type="checkbox"/>	Is this an IVF pregnancy? (no=0, yes=1)	<input type="checkbox"/>
Maternal weight (kgs)	<input type="text"/>	If yes egg collection date:	<input type="text" value="DD"/> <input type="text" value="MM"/> <input type="text" value="YY"/>
Height (cms)	<input type="text"/>	embryo transfer date	<input type="text" value="DD"/> <input type="text" value="MM"/> <input type="text" value="YY"/>
Previous Neural Tube Defect pregnancies (none=0, one=1, two or more=2)	<input type="checkbox"/>	If egg(s) donated enter the donor's DOB	<input type="text" value="DD"/> <input type="text" value="MM"/> <input type="text" value="YY"/>
Previous Down's Syndrome pregnancies (none=0, non-inherited=1, inherited translocation=2, type not known=3)	<input type="checkbox"/>	If unknown, enter donor age	<input type="text"/>
If the patient had a previous pregnancy with Down's syndrome how old was she at the time?	<input type="text"/>	Does the patient smoke? (no=0, yes=1, given up during pregnancy=2, e-cigarettes=3, patches=4)	<input type="checkbox"/>
Previous other chromosomal pregnancy (no=0, yes=1). If yes, please specify abnormality and year diagnosed:	<input type="checkbox"/>	If yes, number of cigarettes per day	<input type="text"/>
Family origin: (Black Caribbean/African=1, White European=2 Indian/Pakistani/Bangladeshi/Sri Lankan=4, Chinese/Japanese/SE Asian=5, Other=6). If other, please specify:	<input type="checkbox"/>	Did the patient take a daily supplement containing Folic Acid? (no=0, before becoming pregnant=1, once she knew she was pregnant=2)	<input type="checkbox"/>
		Has the patient had pre-eclampsia in a previous pregnancy? (no=0, yes=1)	<input type="checkbox"/>
		If the patient has had an amniocentesis performed prior to this test please see overleaf.	

ULTRASOUND SCAN

Date of scan	<input type="text" value="DD"/> <input type="text" value="MM"/> <input type="text" value="YY"/>	FETUS 1	FETUS 2
Hospital where scanned	_____	Nuchal translucency (NT) (mm):	<input type="text"/>
Number of fetuses	<input type="checkbox"/>	Crown rump length (CRL) (mm):	<input type="text"/>
If twins are they monochorionic or dichorionic? (MC=1, DC=2)	<input type="checkbox"/>	Head circumference (HC) (mm):	<input type="text"/>
Name of Sonographer	_____	Gestational age at time of scan	<input type="text"/> weeks <input type="text"/> days
Sonographer ID Code	<input type="text"/>	EDD	<input type="text" value="DD"/> <input type="text" value="MM"/> <input type="text" value="YY"/>

Date of serum sample	<input type="text" value="DD"/> <input type="text" value="MM"/> <input type="text" value="YY"/>	Time taken	_____	Sample taken by	_____
Was the DNA sample taken at the same time (no=0, yes=1)	<input type="checkbox"/>	If no, please complete below:			
Date of DNA sample	<input type="text" value="DD"/> <input type="text" value="MM"/> <input type="text" value="YY"/>	Time taken	_____	Sample taken by	_____

ADDRESS TO WHICH REPORT SHOULD BE SENT

Telephone No. _____ Fax No. _____

Leukaemic studies request

(Cytogenetics/Molecular Genetics)



THE DOCTORS
LABORATORY

Lab No: _____

Priority Code: _____

Surname:

First Name:

Hospital No.:

Date of Birth:

Consultant: _____

Gender: ☐ Male ☐ Female

Sample Type: _____

Sample WBC ($\times 10^9/l$): _____

Sample Date: _____

Sample Vol. (ml): _____

Date Received:

Time Received: _____

Sample Comments: _____

Amount Sample/Culture: _____ Check: _____

Referral centre/hospital: _____

Full postal address: _____

Tel No.: _____

Fax No.: _____

Referral reason/Clinical details: _____

Disease stage: _____

Treatment stage: _____

Karyotype analysis required? ☐ Yes ☐ No

FISH required? ☐ Yes ☐ No

Probes: _____

RT-PCR Required? ☐ Yes ☐ No

Gene Fusion: _____

SAMPLE REQUIREMENTS

In preservative-free heparin and RPMI medium

Preferred volume

Peripheral Blood

Adult: 10mls

Child: 2-5mls

Bone Marrow

Adult: 5-10ml

Child: 2-5mls

Optimal time in transit

Peripheral Blood: 48hrs

Bone Marrow: 24hrs

☐ Fee to be paid by Patient/Other. **PLEASE PROVIDE ADDRESS DETAILS**

☐ Fee to be paid by
Doctor/Clinic as above

Insurance Co. _____ Membership No. _____

Patient address _____

Postcode _____ Contact telephone number _____

Genetic Request



THE DOCTORS
LABORATORY

In order to provide an efficient service for Genetic Requests, please complete the following:

PATIENT DETAILS

Surname: _____

First Name: _____

Date of Birth: _____ Gender: ☐ M ☐ F

Patient Number: _____

Ethnic Origin: _____

Gestation (if applicable): _____ weeks

REFERRING DOCTOR

Name: _____

Address: _____

Telephone: _____

Fax: _____

TEST REQUEST

Disease Name: _____

Gene(s) to be Analysed: _____

Test for: ☐ Diagnosis ☐ Carrier Screening ☐ Known Family Mutation

Clinical Symptoms: _____

Family History: _____

Please state any Family Gene Mutation(s) if known: _____

Please also provide copies of any relevant genetic or pathology (ie. haematology) reports.

INFORMED CONSENT

PATIENT OR GUARDIAN

Please cross-out where applicable:

I consent /do not consent to be tested for the genetic test(s), which have been explained to me

I consent /do not consent for the results of this test to be available to assist in testing other family members

I consent /do not consent for DNA from this sample to be stored

I consent /do not consent for DNA to be used anonymously for relevant research

Signed: _____

Date: ____/____/____

DOCTOR/GENETIC COUNSELLOR

I have explained the purpose of obtaining a blood or tissue sample for genetic testing.

Signed: _____

Date: ____/____/____

This consent form is for use with diagnostic testing. It is important to think through the implications of genetic testing for other family members. We strongly recommend genetic counselling for predictive testing in disorders such as Huntington's Disease or inherited cancers. Please contact our Consultant if you have queries about consent or counselling issues.

☐ Fee to be paid by Patient/Other. **PLEASE PROVIDE ADDRESS DETAILS**

Insurance Co. _____ Membership No. _____

Patient address _____

_____ Postcode _____ Contact telephone number _____

☐ Fee to be paid by
Doctor/Clinic as above

TAP4157/05-11-19/V2

Supplies re-order form

Tel: 020 7307 7373 Fax:020 7307 7340

E-mail:supplies@tdlpathology.com



THE DOCTORS
LABORATORY

Doctor/Practice: _____

Address: _____

DATE OF ORDER

--	--	--	--	--	--

IF URGENT BY

--	--	--	--	--	--

Requested by: _____ Tel: _____

VACUTAINER TUBES No. Required

<input type="checkbox"/> EDTA 4ml Lavender	[]
<input type="checkbox"/> EDTA 10ml Lavender (For STDx)	[]
<input type="checkbox"/> SST/Serum 5ml Gold	[]
<input type="checkbox"/> Fluoride Ox./Glucose 4ml Grey	[]
<input type="checkbox"/> Lithium Heparin 6ml Green	[]
<input type="checkbox"/> No Additive Red 6ml	[]
<input type="checkbox"/> Sod. Heparin 6ml Dark Blue	[]
<input type="checkbox"/> Citrate 4.5ml Light Blue	[]

VACUTAINER NEEDLES No. Required

<input type="checkbox"/> 21g Green	[]
<input type="checkbox"/> 21g Butterfly Green	[]
<input type="checkbox"/> 22g Black	[]
<input type="checkbox"/> 23g Butterfly Blue	[]
<input type="checkbox"/> VACUTAINER BARREL WHITE	[]

SYRINGES (20)

☐ 10ml ☐ 20ml

HELICOBACTER PYLORI No. Required

☐ Breath/Blow Bags []

URINE/STOOL CONTAINERS No. Required

<input type="checkbox"/> Urine/Universal Container pots 30ml	[]
<input type="checkbox"/> Urine/Universal Container pots 60ml	[]
<input type="checkbox"/> 24 hour Urine Containers	[]
<input type="checkbox"/> Stool Pot	[]
<input type="checkbox"/> FOB Pot	[]

REQUEST FORMS

☐ Singles ☐ Duplicates

PERSONALISED BARCODED FORMS

☐ Singles ☐ Duplicates

SAMPLE BAGS No. Required

<input type="checkbox"/> Clear Small	[]
<input type="checkbox"/> Clear Large	[]
<input type="checkbox"/> Red (Urgent)	[]
<input type="checkbox"/> Large Sample Practice Packing Bag	[]

SWABS, GYNAE & NON-GYNAE CYTOLOGY

No. Required

<input type="checkbox"/> Speculum (10) S <input type="checkbox"/> M <input type="checkbox"/> L <input type="checkbox"/>	
<input type="checkbox"/> Thin Prep Vial + Thin Prep Brush	[]
<input type="checkbox"/> Microbiology CULTURE Swabs BLUE	[]
<input type="checkbox"/> ENT/Urethral CULTURE Swabs ORANGE	[]
<input type="checkbox"/> PCR Swabs (chlamydia, herpes, etc) BLUE	[]
<input type="checkbox"/> PCR Swabs (chlamydia, herpes, etc) PINK	[]
<input type="checkbox"/> Histology Pots 60ml	[]
<input type="checkbox"/> Virology Swabs GREEN	[]
<input type="checkbox"/> Blood Culture Bottles	[]

OTHERS – PLEASE SPECIFY

POSTAL PACKS (All postal packs are made up with Royal Mail Track 24 return postal envelopes and labels.)

No. Required

<input type="checkbox"/> Haem/Bio (Lavender/Gold/Grey vacutainer)	[]
<input type="checkbox"/> Single SST vacutainer	[]
<input type="checkbox"/> 30ml MSU/DOA (Non Chain of Custody)	[]
<input type="checkbox"/> COVID-19 Antibody (blood) kit for self-collection	[]
<input type="checkbox"/> COVID-19 PCR swab kit	[]
<input type="checkbox"/> DOA (with Chain of Custody)	[]
<input type="checkbox"/> FOB pack to QFIT pack	[]
<input type="checkbox"/> Group B Strep (GBS) kit	[]
<input type="checkbox"/> HPV Swab kit for self-collection	[]
<input type="checkbox"/> Stool (now brown not blue)	[]
<input type="checkbox"/> Thin Prep Vial postal pack	[]

**PATIENT RECEPTION AT:
THE DOCTORS LABORATORY**
76 Wimpole Street, London W1G 9RQ
Monday to Friday 7.00am – 7.00pm
Saturday 7.00am – 1.00pm
Main Tel: 020 7307 7373
Patient Reception Fax: 020 7307 7373
***Out of hours samples may
be dropped at 76 Wimpole St***


Fax

Additional copy of results to:

SURNAME																DOB	/	/	When completing this form please provide at least three unique identifiers for your patient.
FORENAME											TITLE					M/F			

Please Tick		Home Visit	Patient Ref/ID No.	PROFILES AND TESTS <i>Please specify</i>									
(Biochemistry)	DL1	<input type="checkbox"/>											
(Biochemistry/HDL)	DL1L	<input type="checkbox"/>											
(Haem/Bio)	DL2	<input type="checkbox"/>											
(Haem/Bio/HDL)	DL2L	<input type="checkbox"/>											
(Haematology)	DL3	<input type="checkbox"/>											
(Haem/Bio (short))	DL4	<input type="checkbox"/>											
(Haem/Bio/HDL)	DL4L	<input type="checkbox"/>											
(Postal Haem/Bio)	DL5	<input type="checkbox"/>											
(Postal Haem/Bio/HDL)	DL5L	<input type="checkbox"/>											
Well Person Screen (DL2/T4/TSH/Ferritin)	DL6	<input type="checkbox"/>											
Well Person Screen (DL2L/T4/TSH/Ferritin)	DL6L	<input type="checkbox"/>											
Well Man Screen (DL6/PSA/Ferritin)	DL7	<input type="checkbox"/>											
Well Man Screen (DL6L/PSA/Ferritin)	DL7L	<input type="checkbox"/>											
Well Person Screen (DL6/VITD/Ferritin)	DL8	<input type="checkbox"/>											
Well Person Screen (DL6/HDL/VITD/Ferritin)	DL8L	<input type="checkbox"/>											
Senior Male Profile 60+	DL9M	<input type="checkbox"/>											
Senior Female Profile 60+	DL9F	<input type="checkbox"/>											
Cardiovascular Risk Evaluation Profile	DL10	<input type="checkbox"/>											
Cardiovascular Risk Plus Profile	DL11	<input type="checkbox"/>											
Sexual Health 7 STI screen by PCR	DL12	<input type="checkbox"/>											

<input style="width: 20px; height: 20px; margin-right: 5px;" type="checkbox"/> Fee to be paid by Patient/Other. PLEASE PROVIDE ADDRESS DETAILS Insurance Co. _____ Membership No. _____ Patient address _____ _____ _____ Postcode _____ Contact telephone number _____	<input style="width: 20px; height: 20px; margin-right: 5px;" type="checkbox"/> Fee to be paid by Doctor/Clinic as above Signature _____ Date sample taken _____ Time sample taken _____
--	--

For Practice Use Only:						For Laboratory Use Only:						For Patient Service's Use Only:				 THE DOCTORS LABORATORY
EDTA	SST	GREY	MSU	OTHERS	INITIALS	EDTA	SST	GREY	MSU	OTHERS	INITIALS	TIME IN R	TIME IN Ph	TIME OUT Ph	TAKEN BY INITIALS	

Vacutainer	Anticoagulant	Capacity	SAMPLE TYPES
Lavender	EDTA	4ml/10ml*	A
Gold	SST/Gel	5ml	B
Light Blue	Citrate	4.5ml	C
Red	None	6ml	F
Grey	Fluoride oxalate	2ml, 4ml	G
Green	Lithium heparin	6ml	H
Dark Blue	Sodium heparin	7ml	K

* 10ml EDTA tubes are used for specific PCR assays

Blood culture bottle: contact laboratory	BC
Contact laboratory for advice on sample taking	J
Test by appointment	X
Random Faeces	RF
Faecal Collection	LF
Random Urine	RU
Mid Stream Urine	MSU
First Catch Random Urine (for DL12/Chlamydia, etc.)	FCRU
30ml aliquot from a 24 hour urine collection – state total volume	CU
30ml aliquot from a 24 hour urine collection with 10ml of 0.1N Hydrochloric Acid added – state total volume	PU
Early Morning Urine (1st sample of the day)	EMU
60ml container (sterile)	SC
Cytoc Thin Prep Vial	TPV
Orange/Blue swab for culture – swab in transport medium/Blue microswab	STM
Black Charcoal swab	CS
Green Viral swab	VS
PCR swab for Chlamydia/PCR Infection Screening	PCR
Tap/bottled water mouth wash – 20mls	MW
Ammotic fluid (5mls PCR – 10mls Karyotype)	AF
Chorionic Villus (medium provided by laboratory)	CVS
Urine cytology container	UCYT

The Doctors Laboratory
The Halo Building, 1 Mabledon Place, London WC1H 9AX
Tel: 020 7307 7373 Fax: 020 7307 7374 E-mail: tdl@tdlpathology.com
Web: www.tdlpathology.com



Please recycle responsibly. Printed on FSC accredited Novatech Matt.