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The TDL 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 of the tests and services you need, please contact the laboratory on **020 7307 7373** for advice and information.

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

#### TDL services include:

- Comprehensive, multidisciplinary pathology services
- Specialist diagnostic analysis for other laboratories
- Pathology partnerships with NHS Trusts
- Support for CRO and pharmaceutical companies

# Sonic Healthcare core values

Sonic Healthcare's core values were created by our staff more than 20 years ago, and act as guiding principles for how we conduct ourselves as an organisation.

Our core values set the standard for the collegiate and supportive way in which we behave towards one another, as well as the professionalism with which we conduct ourselves in our day-to-day duties. Individually, our core values articulate our commitment to medical excellence. Collectively, they empower our people to deliver exceptional medical services to doctors and patients.

Since their inception, Sonic Healthcare's core values have been embraced by Sonic Healthcare staff around the world as a unifying code of conduct.

#### **Commit to service excellence**

 To willingly serve all those with whom we deal, with unsurpassed excellence.

# Treat each other with respect and honesty

To grow a workplace where trust, team spirit and equity are an integral part of everything we do.

# Demonstrate responsibility and accountability

 To set an example, to take ownership of each situation to the best of our ability and to seek help when needed.

# Be enthusiastic about continuous improvement

 To never be complacent, to recognise limitations and opportunities for ourselves and processes and to learn through these.

# **Maintain confidentiality**

 To keep all information pertaining to patients, as well as professional and commercial issues, in strict confidence.



# **Complaints policy/procedure**

It is the aim of the company to maintain its core values. Two of these core values are to commit to service excellence, and to be enthusiastic about continuous improvement.

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

The initial complaint will be acknowledged within 3 working days and the investigation, and any follow up actions will be completed within 30 days.

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 and subsequent corrective and preventative actions will be introduced where needed.

Internally, any complaints received will be shared and discussed at Executive Director level where appropriate, as it is the intention of TDL to provide unsurpassed excellence of service.

The Doctors Laboratory
The Halo Building, 1 Mabledon Place
London, WC1H 9AX, UK

Tel: +44 (0)20 7307 7373 – 24 hour telephone (Main switchboard/All services)

Email: tdl@tdlpathology.com

Laboratory times: 24 hours

Samples can be delivered at any time to this location.

Patients' samples cannot be taken at The Halo Building. This service is undertaken at 76 Wimpole Street, London W1G 9RT



To download a location map or to get directions visit:

www.tdlpathology.com/ about-us/locations/

SCAN ME

# TDL Manchester Regents Place, 4 Windsor Street Salford, M5 4HB, UK

Tel: +44 (0)161 332 7181

Email: tdlmanchester@tdlpathology.com

Laboratory times: 24 hours

Samples can be delivered at any time to this location.

Patients' samples cannot be taken at TDL Manchester.

#### **TDL Manchester Couriers**

Direct Tel: +44 (0)161 332 7187 Email: couriersman@tdlpathology.com



# Patient Reception/ Phlebotomy Services

Patient Reception provides a sample collection service for patients attending at the request of their doctor/clinic.

Patients, of all ages, are welcome to attend Patient Reception, 76 Wimpole Street, London W1G 9RT for their samples to be taken. Patients need to be referred by their clinic or doctor and are required to bring a request form or letter of referral.

Appointments are only necessary if a patient needs specialised investigations or care. Instructions can be telephoned or emailed ahead of the patient's attendance, if this is more convenient.

Sample-taking is undertaken by qualified phlebotomy staff for which a standard sample-taking fee of  $\mathfrak{L}60.00$  is charged to patients. Doctors and clinics are charged  $\mathfrak{L}35.00$  for each patient. Sample-taking services for Extended Tests and Drugs of Abuse with Chain of Custody, and semen analysis are routinely available.

Cervical cytology, HVS and cervical swabs are not taken at Patient Reception.

Patient Reception sample-taking services are not available in Manchester.

TDL Patient Reception 76 Wimpole Street, London, W1G 9RT, UK

Tel: +44 (0)20 7307 7383

Email: patientreception@tdlpathology.com

Out of hours samples can be dropped off at this location. **Phlebotomy Services are only available at this location**. Patients' samples cannot be taken at the main laboratory.

#### Opening times

Monday to Friday 7.00am – 7.00pm

Saturday 7.00am - 1.00pm

Closed Sunday and public holidays.



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To download a location map or to get directions visit:

www.tdlpathology.com/ patients/patient-reception/



# TDL Collect: specimen collection services by courier

TDL Collect provides a dedicated medical sample collection service (vans by arrangement) on a scheduled or ad hoc basis.

No charge is made for collections from practices within the M25. Courier collections from private addresses are not undertaken.

The courier collection service for Inner London postcodes operates on a 24/7 basis, as shown. Postcodes extending beyond to the M25 operate from 9.00am to 8.00pm. Outside the M25, and throughout the UK, sample collections are by arrangement and may incur courier charges.

TDL Collect Online Courier Booking is a time-saving option for arranging couriers for sample collection: www.tdlpathology.com/services/tdl-collect/

Please contact **couriers@tdlpathology.com** for your practice's secure login and password.

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

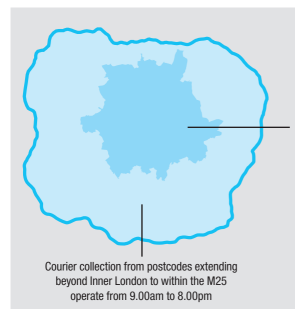
TDL's couriers cannot transport samples containing Hazard Group 4 Pathogens such as Ebola Fever or Haemorrhagic Fever.



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Use the TDL Collect Online Courier Booking service to arrange a courier for sample collection:

www.tdlpathology.com/ services/tdl-collect/



Courier collection from Inner London postcodes (see below) operates 24/7: E1, E2, E3, E4, E5, E6, E7, E8, E9, E10, E11,

E12, E13, E14, E15, E16, E17, E18, E20 EC1, EC2, EC3, EC4

N1, N2, N3, N4, N5, N6, N7, N8, N9, N10, N11, N12, N13, N14, N15, N16, N17, N18, N19, N20, N21, N22

NW1, NW2, NW3, NW4, NW5, NW6, NW7, NW8, NW9, NW10, NW11

SE1, SE2, SE3, SE4, SE5, SE6, SE7, SE8, SE9, SE10, SE11, SE12, SE13, SE14, SE15, SE16, SE17, SE18, SE19, SE20, SE21, SE22, SE23, SE24, SE25, SE26, SE27, SE28

SW1, SW2, SW3, SW4, SW5, SW6, SW7, SW8, SW9, SW10, SW11, SW12, SW13, SW14, SW15, SW16, SW17, SW18, SW19, SW20

W1, W2, W3, W4, W5, W6, W7, W8, W9, W10, W11, W12, W13, W14

WC1, WC2

# **Semen Analysis**

Semen samples need specialist and immediate 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 7025 7940** to make appointments and confirm instructions for sample collection. There is an attendance fee of £50.00.

- Patients must abstain from ejaculation for at least 2 days but not longer than 5 days before the test. Instructions will be given to patients at the time of arranging their appointment.
- Semen samples should be produced at The Doctors Laboratory, 76 Wimpole Street, unless there are exceptional circumstances. If there are exceptional circumstances please contact **TDL Andrology** on **020 7025 7940** for special arrangements and instructions. Refer to Andrology, see page 67.

Semen Analysis services are not provided in Manchester.



SCAN ME

To make an appointment for Semen Analysis online please visit:

www.tdlpathology.com/ andrologybooking

# **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 matched, fully labelled, and include three unique patient identifiers and other relevant Information.

- 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
- Known cases of Hazard Group 4 pathogens such as Ebola or Viral Haemorrhagic Fever must NOT be sent to the laboratory. If there is doubt about a patient's symptoms and presentation please contact the Imported Fever Service on 0844 778 8990 for advice before sending samples to TDL or any 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.



SCAN ME

Download TDL Request Forms from:

www.tdlpathology.com/ tests/request-forms/

# **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 on **020 7307 7373** or by email to **addons@tdlpathology.com**. Please specify the details of the test(s) to be added.

If requests for **Add ons** are made by email, the **patient's details** and **Laboratory Number** need to be referenced.

### **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, London. 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 £150.00 to patients within the M25, and from £250.00 for children when two nurses need to attend. 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**.

# Sample packing

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

Samples need to be collected and packed into appropriate sample containers provided by the laboratory in order to maintain integrity. Attention needs to be given to temperature, special transport containers and time limitations. Each testing has a different sample requirement, which should be referenced prior to sample taking.

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
- International air transport regulations
- 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 (for example a substance that causes viral haemorrhagic fevers) 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, UN3373.

# **Packaging requirements**

There are specific labelling and triple packaging requirements for Category B samples such that it meets packaging instruction P650:

- Primary receptacle tube or vial containing the sample which is placed in the secondary packaging.
- Secondary packaging for example, a protective packaging case or ziplock bag with absorbent material.
- The outer packaging intended to protect the entire contents.

There may also be additional postal envelopes to place the entire package in for postal return. The external surface of the package must be labelled with UN3373 and clearly state BIOLOGICAL SUBSTANCE CATEGORY B.

There are additional packaging requirements for frozen samples requiring shipment using BioFreeze bottles or Dry Ice.

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

# **Postal pathology**

Postal pathology services should be considered by all practices in the UK who need a rapid delivery service to the laboratory as it is a quick and efficient method of sample return, which causes little to no disruption to the patient. Royal Mail require that ALL pathology postal packs are sent using Tracked 24 returns. This provides a particularly suitable method of transport for any healthcare organisation. Royal Mail postal pathology with Tracked 24 returns provides:

- Simple and convenient sample handling throughout the UK for most tests. It is not suitable for samples that need to be received within 24 hours of sample taking (e.g. coagulation, Quantiferon TBQ).
- Scope for large and small numbers of samples.
- Next morning delivery.
- Allows patients and practices to track samples to the Distribution Office through the Royal Mail system.
- Samples can be posted from any Royal Mail post box.
- There is a charge of £3.74 for each Royal Mail Tracked 24 pack. This charge will be itemised in monthly invoices to the practice or patient, as requested.

# **TDL** website

The TDL website gives updated details of our tests — sample types, turnaround times and special instructions. The 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 can be requested by emailing **refranges@tdlpathology.com**. Full details of our tests and profiles are also available in the TDL TestGuide app.



Visit the TDL website at:

www.tdlpathology.com

SCAN ME



# **DX System**

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

TDL Supplies Department provides all appropriate sample collection consumables required for sample collection. Orders will be dispatched on the same or next day and can be made by email to **supplies@tdlpathology.com**. A Supplies Order Form is available from the TDL website.



Download TDL Request Forms from:

www.tdlpathology.com/ tests/request-forms/

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

Security of information in TDL systems and processes is managed by our Information Security Management System, which is certified to the latest International Standard for Information Security ISO/IEC 27001:2013.

#### TDL eViewPlus

Provides the most accurate requesting option for clinics who don't have a practice management system. As well as producing QR coded forms to accompany samples to the laboratory, registered users of this secure Login/Password protected system can request self-collection kits to be sent directly to their patients.

eViewPlus users can also view their results online, with cumulative reporting, anytime, anywhere.

For information about eViewPlus please contact **eviewplus@tdlpathology.com**.

# **Printed Copy**

Printed results will only be sent, as standard, if requested.

# **Emailed results incorporating your logo**

If a practice or company receives results by email, and would like these to be personalised with the practice's logo, please email your company details and logo in GIF format to logo@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 if invoices are sent to patients. All normal credit, debit or charge cards are accepted and payment can be made by following the telephone payment instructions given with each invoice.

The Terms and Conditions of Business appearing on pages 222-231 of this Laboratory Guide shall apply to the services we provide to you, unless otherwise agreed.

# Protection of personally identifiable information

The General Data Protection (GDPR) and UK Data Protection Act 2018 came in to force in 2018 and have had 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.

# **TDL TestGuide app**

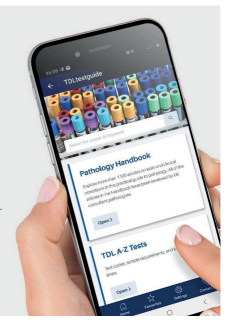
Available for iOS and Android, the TDL TestGuide app offers:

- Full details of TDL's tests and profiles
- The TDL Pathology Handbook, which provides information on more than 1000 pathology topics, reflecting our deep collective knowledge across all areas of pathology

The app can be downloaded from the Apple App Store or Google Play Store. To register for the app, you will just need your TDL Source Code and an email address.

Please contact **testguide@tdlpathology.com** if you need help with finding your Source Code.

Feedback for the TestGuide app is always welcome; please send suggestions and comments to **tdl@tdlpathology.com**.

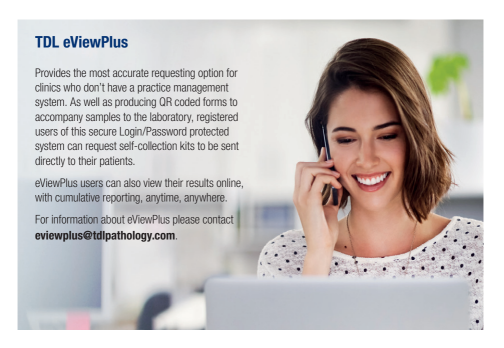


At TDL, these requirements have been implemented within the context of a mature ISO 27001 Information Security Management System – the globally accepted standard by which information is secured.

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.

Customers can find out more about how TDL protects their data by reading the TDL Privacy Notice at www.tdlpathology.com/about-us/corporate-information/tdl-group-privacy-notice.



# **Key contacts**

# 24 HOUR TELEPHONE (MAIN SWITCHBOARD/ALL SERVICES): 020 7307 7373

#### CE<sub>0</sub>

#### **David Byrne**

david.byrne@tdlpathology.com

# **Group Commercial Director**

#### **Brian Madden**

brian.madden@tdlpathology.com

#### **Chief Medical Officer**

#### **Dr Rachael Liebmann OBE**

rachael.liebmann@tdlpathology.com

#### **Group Laboratory Director**

#### **Tim Herriman**

tim.herriman@tdlpathologv.com

#### Director of Sales / Service

#### **Annette Wilkinson**

annette.wilkinson@tdlpathology.com

# Director of Genetics & Molecular Pathology

#### Molecular rati

Dr Lisa Levett

lisa.levett@tdlpathologv.com

#### **Chief Information Officer (IT)**

#### John Matthews

john.matthews@tdlpathology.com

# **Director of Group Laboratory Operations**

#### Lisa Manze

lisa.manze@tdlpathology.com

# **Heads of Support Departments**

# **Director of Laboratory Compliance**

#### **Cyril Taylor**

Cyril.taylor@tdlpathology.com

#### **Director of Governance**

#### **Emer Nestor**

emer.nestor@tdlpathology.com

#### **Credit Control Manager**

#### William Howard

william.howard@tdlpathology.com

#### Logistics / Couriers

#### Steve Kettle

steve.kettle@tdlpathology.com

#### **Patient Reception**

#### Becca Gallagher

Becca.Gallagher@tdlpathology.com patient.reception@tdlpathology.com

#### **Call and Service Centre**

#### Chris Tanalega

chris.tanalega@tdlpathology.com

#### IT Operations / Customer Service

#### Rochelle Fakhri

rochelle.fakhri@tdlpathology.com

#### **Sample Reception**

#### **Amy Scott**

amy.scott@tdlpathology.com

#### **Referrals Department**

#### **Maulik Trivedi**

maulik.trivedi@tdlpathology.com

# **Heads of Laboratory Departments** (London)

# Haem/Bio/Automated Pathology

#### Naina Chavda

naina.chavda@hslpathology.com

# Microbiology/Infection Sciences

#### **Alan Spratt**

alan.spratt@tdlpathology.com

# **Andrology**

# **Wayne Vessey**

wayne.vessey@tdlpathology.com

#### **Cervical Screening**

#### **Julie Smith**

Julie.smith@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

# **TDL Manchester**

# **Operational Site Lead**

#### **Diane Camliyer**

diane.camliyer@tdlpathology.com

# Deputy Site Lead

#### **Andy Leeson**

andy.leeson@tdlpathology.com

# **SRA and Kit Distribution Manager**

# **Georgina Taylor**

georgina.taylor@tdlpathology.com

# **Quality Manager**

#### **Carol Tonge**

carol.tonge@tdlpathology.com

# **Courier Control**

#### **Marc Rennard**

marc.rennard@tdlpathology.com

# 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. The United Kingdom Accreditation Service (UKAS) provides accreditation to the internationally recognised ISO 15189 Medical Laboratories: Requirements for Quality and Competence standard. 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.

Quality Assurance is administered by TDL's Quality Management Group (QMG), who also adhere to regulatory and accreditation requirements.

#### **BIOCHEMISTRY**

#### UKNEQAS, WEQAS, RIQAS, BIORAD, LABQuality for

ACE

AFP/CEA & HCG

Antibiotics (Gentamicin, Vancomycin and Amikacin)

Anti-Hbs Detection

Ammonia

Autoimmune (RF and TPO)

B2 Microalobulin

Cardiac Markers

Clinical Chemistry

CMV laG/laM

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

HTI V

IGF-1

Infectious Immunology

Lipase

Lipid Investigations

NT-Pro BNP

Paediatric Bilirubins

Parasitology

Peptide Hormones

PSA. Free PSA

PTH. ACTH and hCT

QFIT

Rubella IgG Serology

Salicylate and Paracetamol

Specific Proteins

Steroid Hormones

Syphilis Serology

Thyroglobulin Surveys

Thyroid Hormones

Total IgE

**Tumour Markers** 

Toxoplasma IgM Serology

Toxoplasma IgG Serology

Trace Elements

Urine Chemistry

Vitamin D (25 OH)

#### **HAEMATOLOGY**

#### **UKNEQAS**

Automated Differential Leucocyte Count

Blood Film Morphology

Blood Transfusion Laboratory Practice Scheme (BTLP)

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

#### **Special Coagulation**

Anti-Xa assays

ADAMTS-13 activity

ADAMTS-13 antibody

Heparin/Platelet Factor 4

Induced Antibodies

Lupus anticoagulant:

DRVVT assay

Taipan Venom Time

Plasma viscosities

Platelet function analysis (RCPA)

Von Willebrand (vWD) screen

#### **GENETICS AND MOLECULAR VIROLOGY**

#### **Molecular Genetics and Cytogenetics**

# GENQA, ISFG, EMON, UKNEQAS, ECAT, LABQuality for

Acquired array (CLL/MDS)

Acute Lymphoblastic Leukaemia (ALL)

- G banding and FISH

BCR ABL1 and AML Translocation Identification

BCR ABL1 Kinase Domain Variant

BCR ABL1 Major Quantification

BCR ABL1 Minor Quantification

BoBs Rapid Aneuploidy detection

BRAF p.Val600Glu (V600E) Mutation

Status for Hairy Cell Leukaemia

Chlamydia & Gonorrhoea detection by PCR

Chronic Lymphocytic Leukaemia (CLL)

Constitutional Clinical Cytogenetics (Rounds for Amniocentesis, CVS, Solid Tissue, Blood, Array CGH)

Cystic Fibrosis

Duchenne/Becker Muscular Dystrophy

FLT3 Mutation Status

Haematological Technical FISH

Hereditary Haemochromotosis

(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 2C19 genotyping

Human Papillomavirus DNA

IG/TCR Clonality Status

IGHV for CLL

Inborn Errors of Metabolism

JAK2 p.Val617Phe (V617F) Mutation Status

KIT p.Asp816Val (D816V) Mutation

Status for Mast Cell Disease

Lymphoid Gene Panels

Lymphoma

Lymphoplasmacytic Lymphoma /

Waldenstrom Macroglobulinaemia

Measurable Residual Disease for

AML by Molecular Methods

Myeloid (AML/MDS/CML) - G-banding and FISH

Myeloid Gene Panels

Myeloma – sample FISH set up and analysis plus online

Myeloproliferative Neoplasms Diagnostic Testing

NGS AML gene panel

NGS Myeloid Target Panel

NIPT for aneuploidies and sexing

NMP1 Mutation Status

Paediatric Acute Leukaemia Translocations

Paternity Testing

Prader-Willi and Angelman Syndromes

QF-PCR Aneuploidy Detection

Sexually Transmitted Diseases

(CT/NG/MGEN/TV/UU/UP)

Spinal Muscular Atrophy

Thrombophilia (Factor II, V, MTHFR)

TP53 for CLL

Y Microdeletion PCR Assay

#### **Molecular Virology**

#### QCMD, INSTAND for

Adenovirus DNA Viral load and Qualitative PCR

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

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 and Qualitative PCR

HIV-1 DNA Genome Detection

HIV-1 Tropism Genome Detection

HSV 1&2 DNA

HSV 1&2 DNA HSV Drug Resistance

HIV-2 Viral Load

Human Herpes virus 6 DNA

Human Herpes Virus 8 Viral load and Qualitative PCR

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 Variants of Concern (VOC) sequencing

Syphilis PCR

Transplantation Virus Panel

**VZV DNA** 

#### **MICROBIOLOGY**

### **Laboratory Quality Scheme**

Aspergillus PCR

Blood culture and gram stain

Candida PCR

Helicobacter pylori antigen from faeces

Mycoplasma PCR

Polarising crystal microscopy from synovial fluid

Streptococcus pyogenes (Group A)

detection in pharyngeal samples

Surveillance for multi drug resistant bacteria

#### **UKNEQAS**

Antifungal assays

Antifungal susceptibilities

Clostridium difficile detection and toxin testing

Cryptococcal antigen

Faecal parasites

Fungal culture

Fungal biomarkers

General bacteriology

Genital pathogens

MRSA screening

Microbial susceptibilities

Mycobacterial microscopy

Mycobacterial culture and molecular detection

Urinary antigen

#### **WEQAS POCT**

Urinalysis

#### **QCMD**

Atypical pneumoniae PCR

Dermatophyte PCR

PCP PCR

#### **IMMUNOLOGY**

### UKNEQAS - General Immunology

Allergen Component Testing

Allergen Specific IgE Antibodies

Anti-Phospholipid Antibodies (B2GP)

Autoimmune Serology ANCA/GBM Antibodies

**Bullous Dermatosis Antibodies** 

Coeliac Disease (Endomysium, Tissue transglutaminase)

Covid 19 Antibodies

Diabetic Marker (Islet Cell Antibodies)

Faecal Markers (Calprotectin)

General Autoimmune Serology

Hepatitis E (IgG and IgM)

IGRA (Interferon gamma release assay)

Intrinsic Factor Antibodies

Lyme (IgG + IgM)

Myositis Associated Antibodies

Nuclear and Related Antigens

Specific Microbial Antibodies

Syphilis (THPA and RPR)

Tryptase

#### **UKNEQAS – Infectious Immunology**

Anti-Hbs Detection

CMV lgG/lgM

Diagnostic Serology Hepatitis

Helicobacter pylori antigen from faeces

Hepatitis B Serology

Hepatitis C Serology

HIV Serology/POCT

HTLV

Measles and Mumps Serology

Parasite Serology

Parvovirus and Rubella Serology

Syphilis Serology

Toxoplasma IgM Serology

Toxoplasma IgG Serology

#### **RCPAQAP Scheme**

Chlamydia Serology

Legionella (IgG) Serology

Striated Muscle Antibodies

#### **INSTAND Scheme**

Adrenal Antibodies

**HDV Serology and Functional Complement** 

Hepatitis E Serology

#### **CSCQ Scheme**

Lyme Borrelia Serology

#### **Laboratory Quality Scheme**

Antistreptolysin O Titre

Cytomegalovirus Antibodies

**EBV Serology** 

Euroimmun ifQ-Lubeck (Liver)

Autoimmune Disease Scheme

Helicobacter Pylori IgG Antibodies

Herpes Simplex 1 & 2 Antibodies

Measles Serology

Mumps Serology

Mycoplasma Serology

RNA Polymerase III

VZV Serology

#### **ENDOCRINOLOGY**

#### **UKNEQAS**

AFP/CEA

Allergens Scheme

Peptide Schemes 1 to 4

Prostate Specific Antigen

PTH

SHBG

Specific IgE/Total IgE

Steroid Hormones

Thyroid Scheme

Tumour Markers

#### **CERVICAL SCREENING**

#### **NHS England**

Gynaecological Cytopathology EQA Scheme (GEQA)

National EQA Scheme for the Preparation and Staining of Cervical Liquid Based Cytology Samples (TEQA)

#### **HOLOGIC EQA scheme for**

ThinPrep Stain

#### **UKNEQAS** for Microbiology

Molecular Detection of HPV

#### **DIAGNOSTIC CYTOLOGY**

#### UKNEQAS for CPT

Stained Non-Gynaecological Cytology Module.

All non-gynaecological (diagnostic cytology), including Urine Cytology, are referred to a UKAS accredited laboratory for reporting.

#### **ANDROLOGY**

#### UKNEQAS

Semen Analysis Scheme

#### INFORMATION SECURITY

Accredited by British Standards Institute ISO/IEC 27001:2013

#### **Links to the UKAS Schedules of Accreditation**

**HSL Blood Sciences (8169)** 

https://www.ukas.com/wp-content/uploads/schedule\_uploads/00007/8169-Medical-Single.pdf

**HSL Infection Sciences (8860)** 

https://www.ukas.com/wp-content/uploads/schedule\_uploads/00007/8860-Medical-Single.pdf

HSL Molecular Pathology and Genetics (8059)

https://www.ukas.com/wp-content/uploads/schedule\_uploads/00007/8059-Medical-Single.pdf

TDL Manchester (8812)

https://www.ukas.com/wp-content/uploads/schedule\_uploads/00007/8812-Medical-Multiple.pdf

TDL Andrology (10199)

https://www.ukas.com/wp-content/uploads/schedule\_uploads/00007/10199-Medical-Single.pdf

**HSL Cervical Screening (8511)** 

https://www.ukas.com/wp-content/uploads/schedule\_uploads/00007/8511-Medical-Single.pdf

# **Measurement Uncertainty**

Medical laboratories are responsible for ensuring that test results are fit for clinical application by defining analytical performance goals, selecting and qualifying appropriate measurement procedures. All measurement results have some inaccuracies due to analytical bias and imprecision; therefore a measurement result is an estimate at the time of undertaking such measurements. To properly use such results, medical laboratories and their clinical users need some knowledge of the accuracy of such estimates and the uncertainty it may have on the interpretation of patient results.

This estimate of such uncertainties is referred to as Measurement Uncertainty (MU) which incorporates the cumulative range of factors involved in the examination procedure, which may potentially influence the overall test result and thus the interpretation of patient results.

The complete result of a measurement is a value, a unit and an estimate of uncertainty.

Medical laboratories consider the impact of such uncertainties on the interpretation of patient results and ensure uncertainties are minimised through control measures such as standardised procedures, Internal Quality Control monitoring and trending and routine evaluation of MU. Evaluating measurement uncertainty is an ISO 15189:2022 accreditation requirement.

It should be noted that all assays within the TDL/ HSL group of laboratories use standard operating procedures followed by trained and competency assessed scientists.

The MU is initially estimated for each assay during the qualification of the selected examination procedure and is evaluated against the pre-defined maximum allowable measurement uncertainty. Each MU is then re-estimated at regular intervals with additional data and reviewed against the pre-defined maximum allowable measurement uncertainty to ensure uncertainty values and therefore errors and inaccuracies are minimised.

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 such as clear sample labelling. In these cases, the potential risk to the patient management is that the laboratory may need to reject the samples, and not carry out processing. Sometimes the laboratory can rectify a situation where a sample falls short of the sample acceptance criteria though in this case the risk to the patient management may be a breach of stated turnaround time and a delay to provision of the result. In order to reduce the risk of sample rejection or delay to provision of results, please ensure all sample taking criteria are met.

# **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 uricyte container).
- Insufficient sample received.
- No sample received.
- Labelling or form issues (mislabelled/ 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.
- 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.
- Discontinued tests.

# **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.
- Cervical Cytology cannot accept samples containing a sample broom.

- Cervical Cytology can only accept samples received in a Hologic ThinPrep Vial.
- Cervical cytology cannot accept samples received in an expired ThinPrep Vial.
- Urine cytology cannot accept delayed samples unless they have been refrigerated.

Samples deemed to be unrepeatable (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).

# **Consultant advice and opinion**

Each department in the laboratory is consultant led. The TDL Consultants listed below have defined advice or professional support, TDL consultants can be contacted via the laboratory.

# **TDL Lead Consultants**

#### **Chief Medical Officer**

#### Dr Rachael Liebmann OBE

BSc Hons, MB, BCh, BAO, FRCPath, FAcadMed, SFFMLM, FIBMS (Hon)

# **Allergy and Immunology**

#### **Dr Scott Pereira**

MA, MB, B Chir, PhD, FRCPath

# **Prof. Suranjith Seneviratne**

DPhil (Oxon), FRCP, FRCPath

# **Andrology**

# **Dr Sheryl Homa**

PhD. ARCS. FIBMS

# **Biochemistry**

### Dr Frank Geoghegan

**FRCPath** 

#### **Blood Transfusion**

# **Dr Vivienne Andrews**

**FRCPath** 

# **Cervical Cytology**

#### **Dr Mary Falzon**

MRCS, LRCP, FRCPath

#### Dr Geraldine Soosav

MB, BS, FRCPath

# Diagnostic (Non-Cervical) Cytology

#### Dr Mohamed Elshiekh

MBBCh, MSc, FRCPath

# **Dr Miguel Perez Machado**

**FRCPath** 

#### Genetics

#### **Prof. Michael Patton**

FRCP. FRCPCH

#### **Haematology**

#### **Prof. Adrian Bloor**

MA. PhD. FRCP. FRCPath

# Histopathology

#### **Dr Rachael Liebmann OBE**

BSc Hons, MB, BCh, BAO, FRCPath, FAcadMed, SFFMLM, FIBMS (Hon)

# **Medical Microbiology**

#### **Dr Robin Smith**

**FRCPath** 

#### **Parasitology**

#### Dr Laura Nabarro

FRCPath

#### **Point of Care Testing**

#### **Dr Gilbert Wieringa**

MSc. FRCPath, EuSpLM

#### **Virology**

#### **Dr Mark Atkins**

BSc (Hons), MSc, MBBS, FRCPath

# **TDL Consultants**

# **Allergy and Immunology**

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#### **Andrology**

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# **Dr Bryan Woodward**

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# Biochemistry and Point of Care Testing

# **Dr Frank Geoghegan**

**FRCPath** 

# **Dr Gilbert Wieringa**

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### **Dr Bernie Croal**

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#### **Dr Denise Darby**

MRCP, FRCPath

# **Dr Paul Holloway**

**FRCPath** 

# Mr Ed Kearney

MSc, MCB, FRCPath

# Prof. Carel le Roux

FRCPath

#### Dr Moya O'Doherty

MB ChB, MRCPUK, DRCOG, MRCGP, DFFP, FRCPath

Dr Colleen Ross FRCPath

**Dr Hussam Rostom** BA BMBCh, FHEA, RCPath

**Dr Rajeev Srivastava**MBBS, MS, FRCS, FRCPath, EuSpl.m

**Dr Michael Thomas** BSc, PhD, MSc, FRCPath, EuSpLM, SRCS, CSci

**Prof. Pankaj Vadgama** MB, BS, FRCPath

**Dr Royce Vincent** MB, BS, FRCPath

Mr Craig Webster FRCPath

Dr Rachel Webster
PhD. FRCPath

# **Blood Transfusion** and Haematology

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**Prof. Adrian Bloor**MA, PhD, FRCP, FRCPath

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**Dr Gillian Evans** MB. ChB. FRCPath

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**Dr Keith Gomez** FRCPath

**Dr Will Lester**BSc Hons, FRCP, FRCPath, PhD

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**Dr Chris McNamara** MB. BS. FRCPath

**Prof. Atul Mehta** B Chir. FRCPath

**Dr Ryan Mullally** MRCP, FRCPath

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MRCP FRCPath, PGCert, MedEd

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**Dr Philip Robson** FRCPath

Dr Sajitha Sachchithanantham ERCPath

Prof. Marie Scully MRCP. FRCPath

**Dr Mallika Sekhar** FRCPath

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# **Cervical Cytology**

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**Dr Miguel Perez Machado** FRCPath

**Genetics:** Molecular/Cytogenetics

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**FRCPath** 

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#### **Dr Olivier Giger**

MD. PhD

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#### Dr Delaram Kermani

MD. FRCPath

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#### Dr Kav Lawson

MB BS BSc (Hons), FRCPath

# Dr Tanya Levine

FRCPath

# Prof. Teresa Marafioti

MD. FRCPath

#### Dr Adriana Martinez

**FRCPath** 

#### Dr Miriam Mitchison

# **Dr David Moore**

MD, FRCPath

#### Dr Morgan Moorghen

MB ChB, MRCPath, MD, FRCPath

#### Dr Ezra Nigar

RCPath, DipRCPATH

### Dr Laiia Panchal

**MBBS** 

# **Dr Miguel Perez-Machado**

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# **Dr Sabine Pomplun**

MRCPath, FRCPath

#### Dr Nidhi Prasad

MD. FRCPath

#### **Dr Carmel Rvan**

**FRCPath** 

#### **Dr Anna Silvanto**

**FRCPath** 

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#### Dr Madhuri Warren

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# Dr Gillian Williams

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#### **Dr Alison Winstanley**

#### Dr Martin Young

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# **Medical Microbiology**

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**FRCPath** 

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#### Dr Sophie Collier

FRCPath

#### Dr Vanya Gant

**FRCPath** 

#### **Prof. Brian Jones**

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#### Dr Natasha Karunaharan

MRCP. FRCPath

### **Dr Jonathon Lambourne**

MB. BS. FRCPath

#### Dr Alistair Leanord

**FRCPath** 

### **Dr Damien Mack**

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# Dr Stephen Mepham

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# Dr Rajesh Rajendran

MB, BS, FRCPath

# **Dr Antonia Scobie**

MB BS, BSc MSc, DTM&H, MRCP(ID), FRCPath

# **Dr Simon Warren**

FRCPath

# **Dr Emmanuel Wey**

MB, BS, FRCPath

# **Parasitology**

#### **Dr Laura Nabarro**

**FRCPath** 

#### Dr Gauri Godbole

**FRCPath** 

# Virology

# **Dr Mark Atkins**

BSc (Hons), MSc, MBBS, FRCPath

#### **Dr Colin Graham Fink**

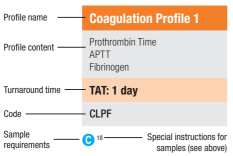
MB. ChB. PhD. FRCPath

# **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 same day.
- 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 the 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.

# **Profile panel information**



16 Lactate

Sample: Fluoride oxalate plasma only.
On ice and separate from cells within 15 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 within 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.
- 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). EDTA Plasma must be frozen within 2 hours.
- 37 Provide sample time and date of collection.
- 38 EDTA sample should not be separated: send whole blood.
- 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.
- 43 Please contact lisa.levett@tdlpathology.com for details for referring samples to the laboratory for sequencing testing.

# **TDL Screening Profiles DL1-DL12**

#### **DL1 Biochemistry Profile**

#### CHANGE

Urea and Electrolytes: Sodium, Potassium, Chloride, Bicarbonate, Urea. Creatinine, eGFR

#### **Liver Function Tests:**

Bilirubin, Alkaline Phosphatase. AST, ALT, Gamma GT, Protein Total, Albumin, Globulin

#### Bone Markers:

Calcium, Phosphate, Uric Acid. Magnesium

Glucose Triglycerides

Cholesterol Iron (TIBC included)

#### TAT: 1 day

#### DL<sub>1</sub>

#### DL1L

plus HDL, LDL (Calculated) and Non-HDL Cholesterol



# **DL5 Biochemistry** & Haematology **Postal Profile**

#### **CHANGE**

#### AS DL4

DL5/DL5L do not include ESR and Phosphate as these results may be more affected by overnight transit times.

#### TAT: 1 day

#### DL5

#### DL5L

plus HDL, LDL (Calculated) and Non-HDL Cholesterol



# **DL2 Biochemistry** (24 Parameters) & **Haematology Profile**

#### CHANGE

#### **HAEMATOLOGY**

Full Blood Count (FBC), ESR

#### **BIOCHEMISTRY**

Urea and Electrolytes: Sodium, Potassium. Chloride. Bicarbonate. Urea, Creatinine, eGFR

#### **Liver Function Tests:**

Bilirubin, Alkaline Phosphatase. AST, ALT, Gamma GT, Protein Total, Albumin, Globulin

#### **Bone Markers:**

Calcium, Phosphate, Uric Acid. Magnesium

Iron (TIBC included)

Glucose

Triglycerides Cholesterol

#### TAT: 1 day

#### DL<sub>2</sub>

#### DL2L

plus HDL, LDL (Calculated) and Non-HDL Cholesterol



# **DL6 General Well Person Profile**

#### CHANGE

DL2 HbA1c Free T4 / TSH Ferritin

#### TAT: 1 day

#### DL<sub>6</sub>

#### DL6L

plus HDL, LDL (Calculated) and Non-HDI Cholesterol



# **DL3 Haematology Profile**

Full Blood Count (FBC)

#### TAT: 1 day

#### DL3



# **DL4 Biochemistry** (16 Parameters) & Haematology Profile

#### CHANGE

#### **HAEMATOLOGY**

Full Blood Count (FBC), ESR

#### BIOCHEMISTRY

#### Renal Function:

Urea, Creatinine, eGFR

#### **Liver Function Tests:**

Bilirubin, Alkaline Phosphatase, AST, ALT, Gamma GT, Protein Total, Albumin, Globulin

#### **Bone Markers:**

Calcium. Phosphate. Uric Acid. Magnesium

Glucose Triglycerides Cholesterol

# TAT: 1 day

#### DL4

#### DL4L

plus HDL, LDL (Calculated) and Non-HDL Cholesterol





### **TDL Screening Profiles DL1-DL12**

#### **DL7 Well Man Profile**

CHANGE Ferritin
DL2 HbA1c
Free T4 / TSH Prostate Profile

TAT: 1 day

DL7

DL7L

plus HDL, LDL (Calculated) and Non-HDL Cholesterol



# DL9M Senior Male Profile 60+

CHANGE DL2

HDL/LDL Cholesterol

Free T4 / TSH HbA1c

Prostate Profile

C Reactive Protein (CRP)

C Reactive Protein (High Sensitivity) Ferritin

MSU

Vitamin D (25-0H) Lp-PLA2 (PLAC) Test

TAT: 2 days

DL9M



#### **DL8 Well Person Profile**

**CHANGE** Ferritin DL2 HbA1c

Free T4 / TSH Vitamin D (25-OH)

TAT: 1 day

DL8

DL8L

plus HDL, LDL (Calculated) and Non-HDL Cholesterol



# DL10 Cardiovascular Risk Profile 1

Lipid Profile (Cholesterol, Triglycerides, HDL Cholesterol, LDL Cholesterol, Non-HDL Cholesterol) Apolipoprotein A1 Apolipoprotein B Lipoprotein (a) C Reactive Protein (High Sensitivity) Lp-PLA2 (PLAC) Test

TAT: 3 days

DL10



# DL9F Senior Female Profile 60+

CHANGE

DL2

HDL/LDL Cholesterol (Calculated)

Free T4 / TSH

HbA1c

C Reactive Protein (CRP)

C Reactive Protein (High Sensitivity) Ferritin

MSU

Vitamin D (25-OH)

HE4

Lp-PLA2 (PLAC) Test

TAT: 2 days

DL9F

ABBGRU4

# DL11 Cardiovascular Risk Profile 2

Lipid Profile (Cholesterol, Triglycerides, HDL Cholesterol, LDL Cholesterol (Calculated), Non-HDL Cholesterol) Apolipoprotein A1

Apolipoprotein B Lipoprotein (a)

Fibrinogen

C Reactive Protein (High Sensitivity) Lp-PLA2 (PLAC) Test

Homocysteine (Quantitative)

TAT: 3 days

DL11

**B B B C** 34

# **DL12 7 STI Profile by PCR (7 PCR Tests from 1 Sample)**

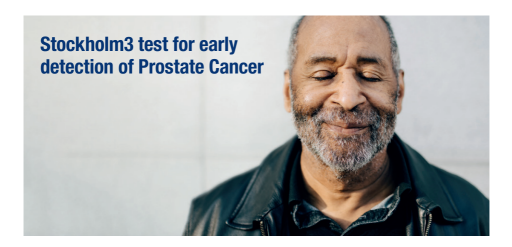
Chlamydia trachomatis Mycoplasma genitalium Trichomonas vaginalis Herpes Simplex I/II Neisseria gonorrhoea Ureaplasma species Gardnerella vaginali

All tests can be requested individually

TAT: 2 days

DL<sub>12</sub>

FCRU / PCR Swab / TPV



Stockholm3 is a blood test that helps to predict risk of clinically significant prostate cancer in men aged 45–74 years with a PSA level greater than 1.5 ng/ml where no previous diagnosis of prostate cancer has been made. Stockholm3 combines genetic markers, proteins and clinical data in an algorithm to help identify clinically significant prostate cancer. It allows for screening in primary or secondary care settings and is equivalent across diverse ethnicities. Apart from a strong health economic case, the result from a Stockholm3 test can be used to reduce unnecessary imaging or invasive diagnostic procedures.

#### **Key characteristics of Stockholm3**

- Increased early detection increased sensitivity
- Increased specificity reduces over testing, unnecessary biopsies by 50% and treatment
- Higher accuracy compared to PSA, PSAD and prostate cancer risk calculator
- Can distinguish between aggressive and benign tumours in a way that PSA testing cannot
- Validated in combination with MRI and in multiple ethnicities
- Shown to detect clinically significant prostate cancers in PSA levels of 1.5–2.9 ng/ml
- Reduces healthcare costs

# Stockholm3: diagnostic patient pathway, including STKR (reflex testing to STK3 from PSA with results of >1.5)

Gender	Male
Intended age	45–74 years, not had prostate cancer, PSA > 1.5 ng/ml
Clinical data required	Age, family history of prostate cancer, previous biopsies, use of 5-alpha reductase inhibitors (Avodart [Dutasteride] or Proscar [Finasteride].
Test code STK3	2 x EDTA tubes must be received within 24–36 hours of sample taking. TAT up to 2 weeks
Test code STKR	PSA levels of >1.5 combined with reflex testing to STK3. 1 x SST, 2 x EDTA tubes must be received within 24–36 hours of sample taking. TAT up to 2 weeks.

A Stockholm3 risk score of >11 is considered to be an indicator of clinically significant prostate cancer risk and referral to a urologist for further investigation is recommended.

For further information about the test please contact stockholm3@tdlpathology.com. Don't post samples to TDL as the timing for receipt of samples within 24–36 hours is important.

See page 106 for test information.

TEST	CODE	SAMPLE REQS	TAT
5 HIAA	RU5H	PU (collect on acid) <sup>1</sup>	5 days
5' Nucleotidase	5NT	В	5 days
6-Thioguanine Nucleotides	TGN	AA	2 weeks
21 Hydroxylase Ab's	21HA	(Frozen)	10 days
Acetylcholine Receptor Autoantibodies	ACRA	<b>B</b> 4	5 days
Acid Phosphatase – Total	APT	B	5 days
Adenosine Deaminase	AD	A / B / Fluid	3 weeks
Adiponectin	ADIP	B	2 weeks
Albumin	ALB	B	1 day
Alcohol (Medical) [Do not use alcohol swab prior to sample taking]	ALCO	<b>G</b> 1	1 day
Alcohol (Urine)	UALC	RU	1 day
Aldolase	ALD0	<b>B</b>	5 days
Alkaline Phosphatase	ALP	<b>B</b>	1 day
Alkaline Phosphatase Isoenzymes	APIE	<b>B</b>	5 days
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	<b>A</b> 9	3 weeks
Requires patient informed consent.	0000	O (5 )	F.1
Alpha-1-Glycoprotein	OROS	(Frozen)	5 days
Alpha-1-Microglobulin	A1MG	RU 1,22	10 days
Alpha-2-Macroglobulins	A2MG	<u>B</u>	5 days
Alpha-Fetoprotein	AFP	B	1 day
ALT (Alanine Aminotransferase) (SGPT)	ALT	B	1 day
Aluminium (Blood)	ALUM	<b>(</b> )	7 days
Amino Acid (EDTA Plasma)	AMIN	(Frozen EDTA Plasma)	7 days
Amino Acid Quantitative (Urine)	UAAQ	RU (Frozen)	7 days
Aminolevulinic Acid (Urine)	RUAL	100mls <b>PU</b>	5 days
Ammonia	AMMO	(Frozen) <sup>15</sup>	1 day
Amylase (Self-collect)	AMY	(TDL Tiny)	1 day
Amylase (Urine)	UAMY	CU	1 day
Amylase (Venous)	AMY	B	1 day
Amylase Isoenzymes	AMYI	B	5 days
Amyloidosis (Amyloid A Protein)	SAA	В	5 days
Androstanediol Glucuronide	ANDG	В	3 weeks
		-	

TEST	CODE	SAMPLE REQS	TAT
Angiotensin II	ANG2	(Frozen plasma)	2 weeks
Angiotensin Converting Enzyme	ACE	В	1 day
Angiotensin Converting Enzyme – CSF	ACEF	CSF (Frozen)	2 weeks
Antimony (Urine)	ANTI	RU 30	10 days
Antimullerian Hormone (AMH) (Self-collect)	AMH	B (TDL Tiny) or (TDL Tiny)	1 day
Antimullerian Hormone (AMH) (Venous) 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 (Venous).	АМН	<b>B</b>	1 day
AP50 Alternative Hemolytic Complement	AP50	(Frozen)	2 weeks
Apolipoprotein A1 (Self-collect) NEW	APOA	(TDL Tiny)	3 days
Apolipoprotein A1 (Venous)	APOA	B	3 days
Apolipoprotein B (Self-collect) NEW	APOB	(TDL Tiny)	3 days
Apolipoprotein B (Venous)	AP0B	B	3 days
Apolipoprotein C	APOC	B	3 months
Apolipoprotein E (12 hours fasting)	AP0E	(fasting)	5 days
Arsenic (Blood)	ARS	(A) or (1)	5 days
Arsenic (Urine)	ARSE	RU 30	5 days
Arylsulphatase A	ARYL	5,6	8 weeks
Aspartate Transaminase (AST) (SGOT)	AST	В	1 day
Bence-Jones Protein	RBJP	RU or CU	5 days
Beta 2 Microglobulin (Serum)	B2MG	В	2 days
Beta 2 Microglobulin (Urine)	UB2M	RU	3 days
Beta-Glucuronidase (Sly Disease)	BGLU	<b>(1)</b> (1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	8 weeks
Bicarbonate	HCO3	В	1 day
Bile Acids – Serum	BILE	В	1 day
Bilirubin (Direct)	DBIL	В	1 day
Bilirubin (Indirect)	IBIL	В	1 day
Bilirubin (Total)	BILI	В	1 day
Biotinidase	BIOT	(Frozen plasma) <sup>4</sup>	3 weeks
Bismuth	BISM	В	5 days
BNP (NT-pro BNP)	BNP	В	1 day
Bone Alkaline Phosphatase	BALP	(Frozen)	2 weeks
Bone Screen	BONE	<b>₿</b> CU	1 day
Bone Screen (Bloods only)	BON2	В	1 day
BUN (Blood Urea Nitrogen) (Calculated)	BUN	3	1 day

TEST	CODE	SAMPLE REQS	TAT
C Reactive Protein (Self-collect)	CRP	(TDL Tiny)	1 day
C Reactive Protein (Venous)	CRP	В	1 day
C Reactive Protein (High Sensitivity) (Self-collect)	HCRP	(TDL Tiny)	1 day
C Reactive Protein (High Sensitivity) (Venous)	HCRP	B	1 day
C1 Esterase: Function & Total	FC1E	(Plasma Frozen) <sup>4,18</sup>	10 days
C1q Binding Immune Complex	IMCP	B	5 days
Cadmium (Blood)	CADM	A or (1)	5 days
Cadmium (Urine)	URCD	RU <sup>30</sup>	5 days
Caeruloplasmin	CERU	В	1 day
Calcium (24 hour Urine)	UCA	PU or acid urine	1 day
Calcium (Venous)	CA	B	1 day
Calcium + Vitamin D (Venous)	CALD	В	1 day
Calcium/Creatinine Ratio	CACR	CU B	1 day
Calprotectin	CALP	<b>QFIT</b> sample collection device	5 days
Calprotectin (Self-collect)	CALP	<b>QFIT</b> sample collection device	5 days
Calprotectin/QFIT Profile (Combined) (QFIT)	QCAL	QFIT	5 days
Calprotectin/QFIT Profile (Combined) (Self-collect)	QCAL	<b>QFIT</b> sample collection device	5 days
Carbohydrate Deficient Glycoprotein	CDG	<b>B</b>	2 weeks
Carbohydrate Deficient Transferrin (CDT) (Self-collect)	CDT	(TDL Tiny)	3 days
Carbohydrate Deficient Transferrin (CDT) (Venous)	CDT	B	3 days
Cardiovascular Risk Profile 1	PP10	88	3 days
Cardiovascular Risk Profile 2	PP11	<b>BBB C</b> 34	3 days
Chest Pain Profile	CPP	<b>B</b>	STAT
Chloride	CL	<b>B</b>	1 day
Cholesterol	СНО	В	1 day
Cholesterol (Familial Hypercholesterolaemia) Requires patient informed consent.	GENE	<b>A A</b> 9	7 weeks
Cholinesterase (Serum/Pseudo)	CHPS	B	1 day
Chromium (Blood)	CHRO	A / Trace metal / (1)	5 days
Chromium (Urine)	URCR	RU <sup>30</sup>	4 weeks

TEST	CODE	SAMPLE REQS	TAT
Chromogranin A	CGA	B	1 week
Chromogranin A & B	MTAB	(Frozen plasma)	3 weeks
Citrate (Blood)	CITR	<b>B</b>	5 days
Citrate (Urine)	UCIT	CU (Frozen)	5 days
CK (MB Fraction)	СКМВ	B	1 day
CK Isoenzymes	CKIE	B	5 days
Cobalt (Blood)	COB	A	5 days
Cobalt (Urine)	COBA	RU <sup>30</sup>	5 days
Coenzyme Q10	CQ10	B	2 weeks
Cold Agglutinin	CAGG	<b>J</b> <sup>1</sup>	5 days
Complement C1q	C1Q	B	5 days
Complement C2	C2	B	10 days
Complement C3	C3	B	1 day
Complement C4	C4	B	1 day
Complement C5	C5A	B	2 weeks
Complement C6	C6	(Frozen)*	5 weeks
* Separate and freeze within 2 hours after collection.		O.E. 14	
<b>Complement C7</b> * Separate and freeze within 2 hours after collection.	C7	(Frozen)*	5 weeks
Complement C8	C8	B (Frozen)*	5 weeks
* Separate and freeze within 2 hours after collection.			
Complement C9	C9	B (Frozen)*	5 weeks
* Separate and freeze within 2 hours after collection.			
Complement Factor H	FACH	B	3 weeks
Copper (Serum)	COPP	⊕ or	5 days
Copper (Urine)	URCU	CU	5 days
Cortisol Binding Globulin	CBG	(Frozen)	1 month
Cotinine (Urine)	COTT	RU	2 days
Creatine Kinase (CK, CPK)	CKNA	В	1 day
Creatinine (including eGFR) (Self-collect)	CREA	(TDL Tiny)	1 day
Creatinine (including eGFR) (Venous)	CREA	B	1 day
Creatinine (Urine)	UCR	CU	1 day
Creatinine Clearance	CRCL	<b>₿</b> CU	1 day
Crosslaps (Serum DPD)	SDPD	(Freeze within 24 hours)	4 days
Cryoglobulins	CRY0	<b>J</b> 6	10 days
Cyclosporin	CYCL	A	1 day

TEST	CODE	SAMPLE REQS	TAT
Cystatin C	CYCC	B	5 days
Cystine – Quantitative (Beta-CTX)	QCYS	PU	5 days
Deoxypyridinoline (DPD) – Serum	SDPD	(Freeze within 24 hours)	4 days
Deoxypyridinoline (DPD) – Urine	DPD	EMU	4 days
Diabetic Profile 1 Please clearly state fasting or non-fasting status.	DIAB	<b>A G</b>	1 day
Diabetic Profile 2 Please clearly state fasting or non-fasting status.	DIA2	A G RU	2 days
Diamine Oxidase Activity	DIAM	B	2 weeks
Elastase (RF)	ELAS	RF	5 days
Elastase (Self-collect)	ELAS	Stool/faecal container	5 days
Electrolytes	ELEC	В	1 day
Electrolytes (Urine)	UELE	CU	1 day
ELF/Enhanced Liver Fibrosis	ELF	В	5 days
Eosinophil Cationic Protein	ECP	В	7 days
Erythropoietin	ERY	B	4 days
Faecal Fat (1 day collection)	TFFA	<b>LF</b> <sup>6</sup>	5 days
Faecal Fat (3 day)	FFAT	<b>LF</b> <sup>6</sup>	5 days
Faecal Lactoferrin	FLAC	RF	5 days
Faecal Sugar Chromatography	FCR0	RF (Frozen)	3 weeks
Ferritin (Self-collect)	FERR	(TDL Tiny)	1 day
Ferritin (Venous)			
Fibrotest (Liver Fibrosis)	FERR	B	1 day
,	FERR FIBT	<b>B</b>	1 day 2 weeks
Fluoride (Urine)			
- '	FIBT	<b>B</b>	2 weeks
Fluoride (Urine)	FIBT UFL	B RU	2 weeks 5 days
Fluoride (Urine) Folate (Red Cell)	FIBT UFL RBCF	© RU	2 weeks 5 days 2 days
Fluoride (Urine) Folate (Red Cell) Folate (Serum)	FIBT UFL RBCF FOLA	B RU	2 weeks 5 days 2 days 1 day
Fluoride (Urine) Folate (Red Cell) Folate (Serum) Free Fatty Acids	FIBT UFL RBCF FOLA FFA	B (Frozen) <sup>1</sup>	2 weeks 5 days 2 days 1 day 10 days
Fluoride (Urine) Folate (Red Cell) Folate (Serum) Free Fatty Acids Fructosamine	FIBT UFL RBCF FOLA FFA FRUC	(a) (Frozen) 1 (b) (Frozen) 2	2 weeks 5 days 2 days 1 day 10 days 1 day
Fluoride (Urine) Folate (Red Cell) Folate (Serum) Free Fatty Acids Fructosamine Galactose-1-Phosphate Uridyltransferase Galactosidase - Alpha* *Sample must reach TDL Referrals Dept. urgently, to be tested within 24 hours of collection. Monday—	FIBT UFL RBCF FOLA FFA FRUC GAL1	(a) (Frozen) <sup>1</sup> (b) (5,6	2 weeks 5 days 2 days 1 day 10 days 1 day 2 weeks
Fluoride (Urine) Folate (Red Cell) Folate (Serum) Free Fatty Acids Fructosamine Galactose-1-Phosphate Uridyltransferase Galactosidase - Alpha* *Sample must reach TDL Referrals Dept. urgently, to be tested within 24 hours of collection. Monday—Thursday only. Referrals to send Immediately.	FIBT UFL RBCF FOLA FFA FRUC GAL1 GALA	B RU  A B (Frozen) <sup>1</sup> B 5.6  J*	2 weeks 5 days 2 days 1 day 10 days 1 day 2 weeks 6 weeks
Fluoride (Urine)  Folate (Red Cell)  Folate (Serum)  Free Fatty Acids  Fructosamine  Galactose-1-Phosphate Uridyltransferase  Galactosidase – Alpha*  *Sample must reach TDL Referrals Dept. urgently, to be tested within 24 hours of collection. Monday—Thursday only. Referrals to send Immediately.  Gall Stone Analysis	FIBT UFL RBCF FOLA FFA FRUC GAL1 GALA RSTA	(a) (Frozen) <sup>1</sup> (b) (5.6 (c) J*	2 weeks 5 days 2 days 1 day 10 days 1 day 2 weeks 6 weeks

TEST	CODE	SAMPLE REQS	TAT
Glucagon	GLUG	(Plasma)	10 days
Glucose	RBG	G	1 day
Please clearly state fasting or non-fasting status.	LIMP	•	0.4-
Haemochromatosis – HFE common variants C282Y + H63D	HMD	<b>A</b> 9	3 days
Haemosiderin (Urine)	HSID	EMU	2 weeks
Haptoglobin	HAPT	В	5 days
HbA1c (Self-collect)	GHB	(TDL Tiny)	1 day
HbA1c (Venous)	GHB	A	1 day
HDL Cholesterol	HDL	B	1 day
Homocysteine (Quantitative)	НОМО	B 17 or (A) (Plasma)	1 day
Homocysteine (Urine)	HCYS	CU	2 weeks
Homovanillic Acid (HVA)	HVA	PU	5 days
Hyaluronic Acid	AHT	B	1 week
Hydroxybutyrate Dehydrogenase	HBD	(Frozen)	1 week
Hydroxyprolene	UHYD	CU	2 weeks
IgG Subclasses	IGSC	В	5 days
Immunoglobulin A	IGA	B	1 day
Immunoglobulin D	IGD	B	5 days
Immunoglobulin E – Total	IGE	В	1 day
Immunoglobulin G	IGG	В	1 day
Immunoglobulin M	IGM	B	1 day
Immunoglobulins (IgG, IgM, IgA)	IMM	В	1 day
Insulin-Like Growth Factor 2	IGF2	<b>B</b> 6	1 month
lodide – Urine	UIOD	RU	1 week
lodine – Serum	IODI	В	1 week
Ionised Calcium	ICPA	B	5 days
Iron (TIBC included) (Venous)	FE	В	1 day
Iron Overload Profile	IOP	<b>A B</b> 9	3 days
Iron Status Profile (Venous)	ISP	В	1 day
Lactate (Plasma)	LACT	<b>G</b> 16	1 day
Lactate Dehydrogenase (LDH)	LDH	B	1 day
Lactate Pyruvate Ratio	LPR	J <sup>1</sup>	4-6 weeks
Lactose Tolerance Test Collection timings and sample requirements: Contact 0207 307 7383 (Phlebotomy)	LTT	By appointment only	1 day
LDL7 Subfractions	LDL7	В	10 days
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TEST	CODE	SAMPLE REQS	TAT
Lead (Blood)	LEAD	A	5 days
Lead (Urine)	URPB	RU	5 days
Leptin	LEPT	(height & weight required) 19	5 days
Lipase (Self-collect)	LIPA	(TDL Tiny)	1 day
Lipase (Venous)	LIPA	B	1 day
Lipid Profile (Self-collect)	LIPP	(TDL Tiny)	1 day
Lipid Profile (Venous)	LIPP	В	1 day
Lipoprotein (a) (Self-collect)	LP0A	(TDL Tiny)	1 day
Lipoprotein (a) (Venous)	LP0A	B	1 day
Lipoprotein Electrophoresis	LEL	B	5 days
Lithium (take 12 hours after dose)	LITH	B	1 day
Liver Fibrosis (Enhanced Liver Fibrosis ELF)	ELF	B	5 days
Liver Fibrosis Fibrotest	FIBT	В	2 weeks
Liver Function Tests (Excluding AST) (Self-collect)	TLFT	(TDL Tiny) and (TDL Tiny)	1 day
Liver Function Tests (Venous)	LFT	<b>B</b>	1 day
Lp-PLA2 (PLAC) Test	PLA2	B	2 days
Lysosomal Enzyme Screen	LE	J <sup>1</sup>	2 months
Lysozyme	LYS0	В	5 days
Magnesium (Serum)	MG	В	1 day
Magnesium (Urine)	URMG	PU	1 day
Manganese (Serum)	MANG	В	5 days
Mercury (Blood)	MERC	A or (1)	5 days
Mercury (Urine)	URHG	RU <sup>1</sup>	5 days
Methaqualone	METQ	RU	5 days
Methylmalonic Acid – Serum	MMAS	В	5 days
Methylmalonic Acid – Urine	MMA	CU	2 weeks
Mucopolysaccharides	MPS	RU (Frozen)	3 weeks
Myeloma Screen CHANGE Please clearly state fasting or non-fasting status.	MYEL	ABG	5 days
Myoglobin (Serum)	SMY0	В	1 day
Myoglobin (Urine)	UMY0	RU	5-10 days
Newborn Screening Panel	GUTH	<b>J</b> <sup>1</sup>	2 weeks
Nickel (Serum)	NICK	В	5 days
Nickel (Urine)	NICU	RU	4 weeks

TEST	CODE	SAMPLE REQS	TAT
Oligosaccharides	UOLI	RU	6 weeks
Orosomucoid (A1AG – Alpha 1 Glycoprotein)	OROS	(Frozen)	5 days
Osmolality (Serum)	OSM0	В	1 day
Osmolality (Urine)	ROSM	RU	1 day
Osteoporosis Screen	0PS	BB	4 days
Oxalate (Plasma)	POXA	(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) <sup>1</sup>	2 weeks
PEth (Phosphatidylethanol) (Self-collect)	PETH	(TDL Tiny)	5-7 days
PEth (Phosphatidylethanol) (Venous)	PETH	A	5-7 days
Phencyclidine (PCP)	DUST	RU	5 days
Phosphate	PH0S	В	1 day
Phosphate (24 hour Urine)	UPH	PU	1 day
PLAC Test (Lp-PLA2) (Self-collect)	PLA2	(TDL Tiny)	2 days
PLAC Test (Lp-PLA2) (Venous)	PLA2	В	2 days
Plasminogen	PLAS	(Frozen plasma) <sup>4</sup>	5 days
Plasminogen Activator Inhibitor – 1	PAI1	(Frozen plasma)	2 weeks
Porphyrin (Blood)	PORP	<b>A</b> 3	15 days
Porphyrin (Stool)	FP0R	RF <sup>3</sup>	3 weeks
Porphyrin (Urine)	RPOR	RU <sup>3</sup>	3 weeks
Porphyrin Full Screen (Total: Urine, Stool, Blood)	PORS	A RU, RF <sup>3</sup>	3 weeks
Potassium	K	В	1 day
Pregnancy (Serum) [Quantitative]	QHCG	В	1 day
Pregnancy Test (Urine)	PREG	RU	1 day
Procalcitonin	PCAL	(Frozen) <sup>4,7</sup>	1 day
Procollagen 1 Peptide N-Terminal (NTX)	P1NP	В	5 days
Procollagen 3 Peptide	PRC0	В	5 days
Propoxyphene	DPR0	RU	5 days
Prostatic Acid Phosphatase	PACP	(Frozen)	3 days
Protein (Urine)	UPRT	CU	1 day
Protein 14.3.3 (Creutzfeldt–Jakob Disease)	CJD	J	5 weeks
Protein Electrophoresis incl. immunoglobulin	PRTE	В	5 days

Protein Total (Blood) PROT  I day  Protein/Creatinine Ratio (Urine) PROT Protein/Creatinine Ratio (Urine) Protein/Creatinine Ratio (Urine)  QFIT Protein/Calprotectin Profile (Combined) (QFIT) QCAL QFIT sample collection device QUANTITIAL (QFIT) QFIT QFIT QFIT QFIT QFIT QFIT QFIT QFIT	TEST	CODE	SAMPLE REQS	TAT
QFIT/Calprotectin Profile (Combined) (QFIT)       QCAL       QFIT       5 days         QFIT/Calprotectin Profile (Combined) (Self-collect)       QCAL       QFIT ample collection device       5 days         QFIT QFIT       QFIT       1 day         Immunochemical Test (QFIT)         QFIT Sample collection device         QFIT Sample collection device         Real Calculi Screen (Metabolic)       RSPR       J6       5 days         Renal Stone Analysis       RSTA       STONE       10 days         Retinol Binding Protein       RBP       3 days         Salicylates       SALI       3       1 day         Selenium (Serum) (Self-collect)       SELE       3 (TDL Tiny)       4 days         Selenium (Serum) (Venous)       SELE       3 (TDL Tiny)       4 days         Serum Free Light Chains       SLC       3       5 days         Silver (Blood)       SILV       3       5 days         Silver (Urine)       USIL       RU       5 days         Sodium       NA       3       1 day         Superoxide Dismutase       SODI       A       5	Protein Total (Blood)	PR0T	В	1 day
QFIT/Calprotectin Profile (Combined) (Self-collect)       QCAL collection device       QFIT sample collection device       5 days         Quantitative Faecal Immunochemical Test (QFIT)       QFIT QFIT ample collection device       1 day         Quantitative Faecal Immunochemical Test (QFIT) (Self-collect)       QFIT QFIT sample collection device       1 day         Renal Calculi Screen (Metabolic)       RSPR J <sup>6</sup> 5 days         Renal Stone Analysis       RSTA STONE       10 days         Retinol Binding Protein       RBP 3 3 days         Salicylates       SALI 3 1 day         Selenium (Serum) (Self-collect)       SELE 3 (TDL Tiny)       4 days         Selenium (Serum) (Venous)       SELE 3 (TDL Tiny)       4 days         Serum Free Light Chains       SLC 3 5 days         Silver (Blood)       SILV 3 5 days         Silver (Urine)       USIL RU 5 days         Sodium       NA 3 1 day         Superoxide Dismutase       SODI 4 1 day         Thiopurine Methyl Transferase       TPMT 4 5 5 days         Tissue Polypeptide Antigen       TPA 3 1 week	Protein/Creatinine Ratio (Urine)	UCPR	RU	1 day
(Combined) (Self-collect)       collection device         Quantitative Faecal Immunochemical Test (QFIT)       QFIT       QFIT ample collection device       1 day         Quantitative Faecal Immunochemical Test (QFIT) (Self-collect)       QFIT QFIT sample collection device       1 day         Renal Calculi Screen (Metabolic)       RSPR       J6       5 days         Renal Stone Analysis       RSTA       STONE       10 days         Retinol Binding Protein       RBP       3 days         Salicylates       SALI       1 day         Selenium (Serum) (Self-collect)       SELE       3 (TDL Tiny)       4 days         Selenium (Serum) (Venous)       SELE       1 days         Serum Free Light Chains       SLC       3 days         Silver (Blood)       SILV       5 days         Silver (Urine)       USIL       RU       5 days         Sodium       NA       1 day         Superoxide Dismutase       SODI       4 days         Thiopurine Methyl Transferase       TPMT       5 days         Tissue Polypeptide Antigen       TPA       1 week	QFIT/Calprotectin Profile (Combined) (QFIT)	QCAL	QFIT	5 days
Immunochemical Test (QFIT)   QFIT   QFIT sample collection device   1 day	•	QCAL		5 days
Test (QFIT) (Self-collect)  Renal Calculi Screen (Metabolic)  RSPR  J  STA  STONE  10 days  Retinol Binding Protein  RBP  Salicylates  SALI  SELE  (TDL Tiny)  4 days  Selenium (Serum) (Venous)  SELE  SELE  (SOCIONA  SELE	****	QFIT	QFIT	1 day
Renal Stone Analysis Retinol Binding Protein RBP 3 days Salicylates SALI 3 tday Selenium (Serum) (Self-collect) SELE 3 (TDL Tiny) 4 days Selenium (Serum) (Venous) SELE 3 tdays Serum Free Light Chains SLC 3 tdays Silver (Blood) SILV 5 days Silver (Urine) USIL RU 5 days Sodium NA 3 tday Superoxide Dismutase SODI A/A 5 tdays Thiopurine Methyl Transferase TPMT A 5 tdays Tissue Polypeptide Antigen TPA 3 tweek	· · · · · · · · · · · · · · · · · · ·	QFIT		1 day
Retinol Binding Protein  RBP  Salicylates  SALI  SALI  SELE  SITDL Tiny)  4 days  Selenium (Serum) (Venous)  SELE  SELE	Renal Calculi Screen (Metabolic)	RSPR	<b>J</b> 6	5 days
Salicylates SALI Selenium (Serum) (Self-collect) SELE SITDL Tiny) 4 days Selenium (Serum) (Venous) SELE SITDL Tiny) 4 days Selenium (Serum) (Venous) SELE SITURE (SITURE) SUC SILV SILV SILV SILV SILV SILV SILV SILV	Renal Stone Analysis	RSTA	STONE	10 days
Selenium (Serum) (Self-collect)       SELE       ③ (TDL Tiny)       4 days         Selenium (Serum) (Venous)       SELE       ③ 4 days         Serum Free Light Chains       SLC       ⑤ 5 days         Silver (Blood)       SILV       ⑥ 5 days         Silver (Urine)       USIL       RU       5 days         Sodium       NA       ⑥ 1 day         Superoxide Dismutase       SODI       ⑥ / ⑥       5 days         Thiopurine Methyl Transferase       TPMT       ⑥ 5 days         Tissue Polypeptide Antigen       TPA       ⑥ 1 week	Retinol Binding Protein	RBP	B	3 days
Selenium (Serum) (Venous)     SELE     3     4 days       Serum Free Light Chains     SLC     3     5 days       Silver (Blood)     SILV     3     5 days       Silver (Urine)     USIL     RU     5 days       Sodium     NA     3     1 day       Superoxide Dismutase     SODI     3 / 1     5 days       Thiopurine Methyl Transferase     TPMT     4 s     5 days       Tissue Polypeptide Antigen     TPA     3     1 week	Salicylates	SALI	В	1 day
Serum Free Light Chains  SLC  Silver (Blood)  SILV  Silver (Urine)  USIL  RU  5 days  Sodium  NA  1 day  Superoxide Dismutase  SODI  A/1  Thiopurine Methyl Transferase  TPMT  TPA  1 week	Selenium (Serum) (Self-collect)	SELE	(TDL Tiny)	4 days
Silver (Blood)       SILV       5 days         Silver (Urine)       USIL       RU       5 days         Sodium       NA       3       1 day         Superoxide Dismutase       SODI       4 / 1       5 days         Thiopurine Methyl Transferase       TPMT       4 s       5 days         Tissue Polypeptide Antigen       TPA       3       1 week	Selenium (Serum) (Venous)	SELE	B	4 days
Silver (Urine) USIL RU 5 days Sodium NA 3 1 day Superoxide Dismutase SODI 4 1 day  Thiopurine Methyl Transferase TPMT 4 5 5 days Tissue Polypeptide Antigen TPA 3 1 week	Serum Free Light Chains	SLC	В	5 days
Sodium NA 3 1 day Superoxide Dismutase SODI 4 5 days Thiopurine Methyl Transferase TPMT 4 5 5 days Tissue Polypeptide Antigen TPA 3 1 week	Silver (Blood)	SILV	В	5 days
Superoxide Dismutase  SODI  A/B  5 days  Thiopurine Methyl Transferase  TPMT  A 5  5 days  Tissue Polypeptide Antigen  TPA  1 week	Silver (Urine)	USIL	RU	5 days
Thiopurine Methyl Transferase TPMT (A) 5 days  Tissue Polypeptide Antigen TPA (3) 1 week	Sodium	NA	В	1 day
Tissue Polypeptide Antigen TPA 3 1 week	Superoxide Dismutase	SODI	<b>A</b> / <b>(1)</b>	5 days
	Thiopurine Methyl Transferase	TPMT	<b>A</b> 5	5 days
Total Acid Phosphatase APT 3 days	Tissue Polypeptide Antigen	TPA	В	1 week
	Total Acid Phosphatase	APT	В	5 days
Total Bile Acid/Bile Salts BILS 1 week	Total Bile Acid/Bile Salts	BILS	В	1 week
Total IgE IGE 1 day	Total IgE	IGE	В	1 day
Transferrin TRAN (3) 1 day	Transferrin	TRAN	В	1 day
Transferrin Electrophoresis TREL 3 weeks	Transferrin Electrophoresis	TREL	В	2 weeks
Triglycerides TRI (3) 1 day		TRI	В	1 day
Trimethylaminuria (Fish Odour Syndrome) FOS J 6 weeks		F0S		6 weeks
Troponin I (High sensitive) TROC (3) 1 day	Troponin I (High sensitive)	TROC		1 day
Troponin T (High sensitive) TROT (3) 1 day	Troponin T (High sensitive)	TROT		1 day
Tryptase STRY 3 days	Tryptase	STRY	В	2 days
Tumour Necrosis Factor – Alpha TNF (3 (Frozen) <sup>4</sup> 2 weeks	Tumour Necrosis Factor – Alpha	TNF	B (Frozen)⁴	2 weeks
Urate (Uric acid) UA (3) 1 day		UA	В	1 day
Urea (Self-collect) UREA (3 (TDL Tiny) 1 day	Urea (Self-collect)	UREA	(TDL Tiny)	1 day
Urea (Venous) UREA (3) 1 day	Urea (Venous)	UREA	<b>B</b>	1 day

TEST	CODE	SAMPLE REQS	TAT
Urea (Urine)	UURE	CU	1 day
Urea and Electrolytes	U/E	B	1 day
Urea/Creatinine/eGFR (Self-collect)	TCU	(TDL Tiny)	1 day
Uric Acid (Serum)	UA	B	1 day
Uric Acid (Urine)	UURI	CU	1 day
Urinary Bladder Cancer Antigen  ** It is recommended to collect mid-stream urine. Do not first morning urine. Collection of urine specimen before surgical intervention or treatment or 1–2 weeks after speak and the collected with an instrument e.g. catheter.	any	<b>RU</b> (Freeze within 48 hours)**	5 days
Urine Microalbumin/Creatinine Ratio	UMA	RU	1 day
Urine Organic Acids	UORG	RU (Frozen)	3 weeks
Urine Steroid Screen (Steroid Hormones)	USTE	CU <sup>9</sup>	2 weeks
Urine Sugar Chromatography	UCR0	RU (Frozen)	3 weeks
Very Long Chain Fatty Acids	VLCF	A or (Frozen) 9	4-6 weeks
Vitamin B12 (Active) (Self-collect)	B12	(TDL Tiny)	1 day
Vitamin B12 (Active) (Venous)	B12	B	1 day
Vitamin B12 (Active)/Red Cell Folate	B12F	AB	2 days
Vitamin B12 (Total)	TB12	B	1 day
Vitamin D (25-OH) (Self-collect)	VITD	(TDL Tiny)	1 day
Vitamin D (25-OH) (Venous)	VITD	B	1 day
VLDL Cholesterol	VLDL	<b>B</b> 13	1 week
VMA	UVMA	<b>PU</b> <sup>1</sup>	5 days

#### **Bone Screen**

Alkaline Phosphatase Protein Total (Blood) Albumin Globulin Calcium

TAT: 1 day

**BONE** 

CU

# Bone Screen (Bloods only)

Urea and Electrolytes Liver Function Tests (LFTs) Calcium Phosphate Vitamin D (25-0H)

TAT: 1 day

BON2

В

### Cardiovascular Risk Profile 1

Lipid Profile (Cholesterol, Triglycerides, HDL Cholesterol, LDL Cholesterol (Calculated), Non-HDL Cholesterol) Apolipoprotein A1 Apolipoprotein B Lipoprotein (a) C Reactive Protein (High Sensitivity) Lp-PLA2 (PLAC) Test

TAT: 3 days

PP10



#### Cardiovascular **Risk Profile 2**

Lipid Profile (Cholesterol, Trialycerides. HDL Cholesterol. LDL Cholesterol (Calculated), Non-HDL Cholesterol) Apolipoprotein A1 Apolipoprotein B Lipoprotein (a) Fibrinogen C Reactive Protein (High Sensitivity) Lp-PLA2 (PLAC) Test

TAT: 3 days

PP11





Homocysteine (Quantitative)

#### **Chest Pain Profile**

Myoglobin (Serum) CK (MB Fraction) Troponin T (High sensitive)

TAT: STAT

**CPP** 

B

# Calprotectin/QFIT **Profile (Combined)**

Calprotectin

Quantitative Faecal Immunochemical Test (QFIT)

#### If CALP < 50ug/g then the below comment will be appended:

Calprotectin: < 50 ug/g- Not indicative of GI inflammation. Consider IBS, or quiescent IBD if this is a known patient.

#### If CALP = 50 ug/g or higher,then the below comment will be appended:

Calprotectin: 50-150 ug/g repeat calprotectin in 2 weeks (Also consider other potential causes (infection, NSAIDS, GI malignancy) depending on the magnitude of the result and clinical context.)

Repeated Calprotectin result: 100-250 ug/g routine referral to gastroenterology.

Calprotectin: >250 ug/g urgent referral to gastroenterology.

TAT: 5 days

QCAL

**QFIT** 

#### **Diabetic Profile 1**

Glucose HbA1c

Please clearly state fasting or non-fasting status

TAT: 1 day

DIAB



#### Diabetic Profile 2

Glucose HbA1c

Microalbumin

Please clearly state fasting or non-fasting status

TAT: 2 days

DIA2



#### Iron Overload Profile

Iron (TIBC included) Ferritin

Transferrin Saturation

Haemochromatosis C282Y, H63D

TAT: 3 days

IOP



#### **Iron Status Profile** (Venous)

Iron (TIBC included) Ferritin Transferrin Saturation

TAT: 1 day

**ISP** 



#### **Lipid Profile (Venous)**

Triglycerides Cholesterol

HDL Cholesterol

LDL Cholesterol (Calculated) Non-HDL Cholesterol

TAT: 1 day

LIPP



# **Liver Function Tests (Venous)**

ALT (Alanine Aminotransferase) (SGPT)

Aspartate Transaminase (AST) (SGOT)

Bilirubin (Total)

Total Protein

Alkaline Phosphatase Albumin

Globulin

Gamma GT

#### TAT: 1 day

LFT



#### **Myeloma Screen**

#### CHANGE

Full Blood Count (FBC)

**ESR** Biochemistry Profile

Protein Electrophoresis

Immunoglobulins (IgA, IgG, IgM) Serum Free Light Chains

Please clearly state fasting or non-fasting status

TAT: 5 days

MYFL





#### **Osteoporosis Screen**

Alkaline Phosphatase

Calcium

Albumin

Phosphate

Crosslaps (Serum DPD)

Vitamin D (25 OH)

#### TAT: 4 days

0PS



### **Porphyrin Full Screen** (Total: Blood, Stool, Urine)

Porphyrin Blood Porphyrin Stool Porphyrin Urine

#### TAT: 3 weeks

**PORS** 



#### **Urea and Electrolytes**

Sodium

Potassium

Chloride Bicarbonate

Urea

Creatinine

#### TAT: 1 day

U/E



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	AAB	2 days
Antenatal Profile	ANTE	<b>A A</b> <sup>33</sup> <b>B B G</b>	3 days
APTT/KCCT	KCCT	<b>(</b> ) 18	1 day
Atypical Antibody Screen (handwritten tube label)	AASC	<b>A</b> 22,33	2 days
Blood Film Examination	FILM	A	1 day
Blood Group † † 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.	AB0	<b>A</b> 22,33	2 days
Carboxyhaemoglobin	СВНВ	A	1 week
Coagulation Profile 1	CLPF	<b>(</b> ) 18	1 day
Coagulation Profile 2	CLOT	<b>A C</b> 18	1 day
D-Dimers (Fibrinogen Degradation Products)	DDIT	<b>6</b> 4	1 day
DVT/Pre-travel Screen	DVT1	<b>A A B</b> <sup>9</sup>	5 days
ESR	ESR	A	1 day
Fibrinogen	FIB	C 4,18	1 day
Full Blood Count	FBC	A	1 day
Full Blood Count* (Haemoglobin, White Cell Count, Red Cell Count, Platelets) (Self-collect) NEW *Mix sample on collection.	TFBC	(TDL Tiny)	1 day
Haematology Profile	PP3	A	1 day
Haemoglobin	НВ	A	1 day
Immune Function Evaluation (Total)	TIE	A + B 5,10	7 days
INR	PTIM	<b>(</b> ) 18	1 day
Lymphocyte Subsets (CD3/CD4/CD8)	LYSS	A	1 day
Malarial Parasites	MALP	<b>A</b> 4,9,14	STAT
Malarial Parasites (visa, non-urgent)	MP48	A	2 days
Mean Cell Volume (MCV)	MCV	A	1 day
Microfilaria Blood Film	MICF	A	STAT
Natural Killer Profile 2	NKP2	<b>A</b> 10	2 days
PAI-1 4G/5G Polymorphism	PAIP	A	10 days
Paul Bunnell (Monospot)	PAUL	A or B	1 day

TEST	CODE	SAMPLE REQS	TAT	
Pre-Travel Screen (DVT)	DVT1	<b>A A B</b> 9	5 days	
Prothrombin Time	PTIM	<b>C</b> 18	1 day	
Reticulocyte Count	RETC	A	1 day	
Thrombin Time	THR0	<b>C</b> 18	1 day	

# **Special Haemostasis**

TEST	CODE	SAMPLE REQS	TAT
Activated Protein C Resistance	APCR	C (Frozen) <sup>4,18</sup>	3 days
ADAMTS-13 Activity	CP13	C (Frozen) <sup>4,18</sup>	3 days
ADAMTS-13 Antibody	A13A	C (Frozen) <sup>9,18</sup>	2 weeks
Anti-Xa Apixaban Monitoring * Please state drug and time of dose on request.	APIX	C (Frozen)*18	3 days
Anti-Xa Edoxaban Monitoring *Please state drug and time of dose on request.	ED0X	C (Frozen)*18	3 days
Anti-Xa Fondapariux Monitoring  * Please state drug and time of dose on request.	FOND	C Frozen)*18	3 days
Anti-Xa LMWH Monitoring * Please state drug and time of dose on request.	LMWX	(Frozen)*18	3 days
Anti-Xa Rivaroxaban Monitoring * Please state drug and time of dose on request.	RIVA	C (Frozen)*18	3 days
Antithrombin III Activity	A111	(Frozen) <sup>4,9,18</sup>	3 days
Factor II Assay	FAC2	C (Frozen) <sup>9,18</sup>	5 days
Factor V Assay	FAC5	C (Frozen)9,18	5 days
Factor VII Assay	FAC7	C (Frozen) <sup>9,18</sup>	5 days
Factor VIII Assay	FAC8	C (Frozen) <sup>9,18</sup>	5 days
Factor VIII Inhibiting Antibody	F8IA	<b>()</b> () 18	2 weeks
Factor IX Assay	F1X	C (Frozen) <sup>9,18</sup>	5 days
Factor IX Inhibiting Antibody	F9IA	<b>()</b> () 18	2 weeks
Factor X Assay	FX	C (Frozen)9,18	5 days
Factor XI Assay	FX1	© (Frozen) <sup>9,18</sup>	5 days
Factor XII Assay	FX11	C (Frozen) <sup>9,18</sup>	5 days
Factor XIII Assay	FA13	C (Frozen)9,18	5 days
FXIII A Subunit	F13S	(Frozen) <sup>9,18</sup>	14 days
Hughes Syndrome	LUPA	<b>B C 4</b> ,18	2 days
Lupus Anticoagulant and Anticardiolipin Abs	LUPA	<b>B C C</b> 4,18	2 days
Lupus Anticoagulant only	LUPC	<b>© ©</b> 9,18	2 days

TEST	CODE	SAMPLE REQS	TAT
Miscarriage/Thrombotic Risk Profile	PROP	<b>AABCC</b> <sup>18</sup>	5 days
P2Y12 Receptor Platelet Function	P2Y	<b>J</b> 1	1 day
Analysis (Clopidogrel Resistance)			
** Samples not processed at Halo, contact haemostasis UCLH.Haemostasis@tdlpathology.com prior to sample to			
Platelet Aggregation Studies	PLAG	J** <sup>9</sup>	3 days
**Samples not processed at Halo, contact haemostasis Haemostasis@tdlpathology.com. Please contact Haemostaboratory prior to arranging patient appointment. Platel aggregation studies are not processed without prior arra	stasis et		
Platelet Function Test Screen - PFA-100/200	PFAT	<b>J</b> ** <sup>1</sup>	1 day
** Samples not processed at Halo, please contact haem on UCLH.Haemostasis@tdlpathology.com prior to sampl			
Protein C Activity	PRC	(Frozen) <sup>4,9,18</sup>	3 days
Protein S Activity	PS1	(Frozen) <sup>4,9,18</sup>	5 days
Protein S Free Ag	FPRS	(Frozen) <sup>4,9,18</sup>	3 days
Taipan Snake Venom Time	TTVT	<b>() ()</b> 9,18	2-3 weeks
Thrombotic Risk Profile	PROP	<b>AABCC</b> <sup>18</sup>	5 days
Viscosity (Plasma) *EDTA plasma must be separated within 24 hours of collection and sent at room temperature.	VISC	<b>A</b> *4	3 days
Von Willebrand Profile	FVWF	<b>C C C</b> 4,9,12	5 days
Von Willebrands Multimers	VWM	<b>()</b> () () 18	3 months

# **Special Haematology**

TEST	CODE	SAMPLE REQS	TAT
Coombs (Direct Antiglobulin Test)	COOM	A	2 days
Eosin-5 Maleimide Dye binding test for Hereditary spherocytosis (EMA)*	EMA	A	2 days
*Sample to be received by laboratory within 24 hours of being taken and the test is done Tuesday to Thursday (test must be performed within 48 hours of sample being taken.			
G6PD	G6PD	A	4 days
Haemoglobin Electrophoresis	HBEL	A	4 days
HFE gene (Haemochromatosis) – common variants C282Y + H63D	HMD	<b>A</b> 9	3 days
Thalassaemia Screen	HBEL	A	4 days

# **Flow Cytometry**

TEST	CODE	SAMPLE REQS	TAT
Bone Marrow (Aspirate)	BMAS	<b>J</b> <sup>1</sup>	14 days
Bone Marrow (Trephine Biopsy)	BMI	J <sup>1</sup>	3 days
CD3/CD4/CD8	LYSS	<b>A</b> 10	1 day
CD16	CD16	<b>A</b> 4	1 day
CD19 B Cells	CD19	<b>A</b> 4	1 day
CD20	CD20	<b>A</b> 10	2 days
CD25	CD25	<b>A</b> 10	2 days
CD56	CD56	<b>A</b> 4	1 day
CD57	CD57	A	1 day
Leukaemia Immunophenotyping	LYPT	<b>A</b> 4,5	5 days

#### **Anaemia Profile**

Full Blood Count (FBC) ESR Iron (TIBC included) Ferritin Vitamin B12 (Active) Folate (RBC)

#### TAT: 2 days

ANAE



#### **Antenatal Profile**

Full Blood Count (FBC) Blood Group and Rh Type Atypical Antibody Screen Haemoglobin electrophoresis Syphilis IgG/IgM Glucose Free T4 / TSH Rubella Antibody (IaG) Toxoplasma Antibodies (IgG, IgM) Cytomegalovirus (lgG/lgM) Antibodies Hepatitis B Surface Antigen Hepatitis C Antibodies Varicella zoster Antibodies (IqG) HIV 1 & 2 Abs Please ensure the blood group (EDTA) tube label is handwritten. Do not

affix a secondary label.

TAT: 3 days

ANTE

**AA** 33 **BBBG** 

#### **Coagulation Profile 1**

Prothrombin Time APTT/KCCT Fibrinogen

TAT: 1 day

**CLPF** 



### **Coagulation Profile 2**

Full Blood Count (FBC)
Prothrombin Time
APTT/KCCT
Fibrinogen

TAT: 1 day

CLOT



#### **DVT/Pre-travel Screen**

Full Blood Count (FBC)
Factor II Prothrombin –
G20210A Variant
Factor V Leiden – G1691A Variant
Cardiolipin Antibodies (IgG+IgM)

TAT: 5 days

DVT1



#### **Haematology Profile**

Full Blood Count (FBC) FSR

TAT: 1 day

PP3



# Miscarriage/ Thrombotic Risk Profile

Full Blood Count (FBC)
Coagulation Profile 1
Antithrombin III
Factor V Leiden — G1691A Variant
Factor II Prothrombin —
G20210A Variant
MTHFR — common C677T
+ A1298C variants
Lupus Anticoagulant
Protein C
Free Protein S Aq

Cardiolipin Antibodies (IgG+IgM)

TAT: 5 days

PROP.



# Natural Killer Profile 2

CD3 CD4 CD8 CD16/CD56

CD16/CD36

TAT: 2 days

NKP2



#### e 2

Von Willebrand Factor Von Willebrand Activity (GPIbM assay) Factor VIII Assay

Von Willebrand Profile

TAT: 5 days

**FVWF** 



# **Pre-Travel Screen (DVT)**

Full Blood Count (FBC)
Factor II Prothrombin —
G20210A Variant
Factor V Leiden — G1691A Variant
Cardiolipin Antibodies (IgG+IgM)

TAT: 5 days

DVT1



#### **Thrombotic Risk Profile**

Full Blood Count (FBC)

Coagulation Profile 1
Antithrombin III
Factor V Leiden — G1691A Variant
Factor II Prothrombin —
G20210A Variant
MTHFR — common C677T
+ A1298C variants
Lupus Anticoagulant
Protein C
Protein S Free Ag
Cardiolipin Antibodies (IgG+IgM)

TAT: 5 days

PR<sub>OP</sub>





TEST	CODE	SAMPLE REQS	TAT
16S rRNA Bacterial Gene	16S	J	1 week
18S rRNA Fungal Gene	18S	J	1 week
Beta D Glucan	XBDG	В	3 days
Blood Culture#	BCUL	2 x <b>BC</b> <sup>4</sup>	6 days +

- # Please contact 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.

Candida (Culture for ID Only)	CANC	STM/CS	2-4 days
Candida (Culture for ID + Sensitivities)	CAND	STM/CS	2-4 days
Candida auris Screen	CANS	STM/CS	2-4 days
Carbapenemase producing organism screen  ‡ Presumptive positive isolates will be sent to the UKHSA reference laboratory for confirmation.	MDR	STM (rectal)	4-5 days *
Clostridium Difficile Toxin by PCR	CLOS	RF*	2 days
* Not performed on formed stool specimens.			
Cryptococcal Antigen	CRYC	Serum or CSF	1 day
Cryptosporidium	OCP	RF	2 days
CSF for Microscopy and Culture	CSF	1.5ml <b>CSF</b>	1-3 days
Fluid Culture	FLUD	SC	2-7 days
Specify sample site and sterility of sample collection.			
Fungal investigations (non- superficial extended culture)	FUN	All specimens other than Skin, Hair and Nails	3-21 days
Please send in a dermaPak where possible.			
Fungal investigations (superficial/ dermatophyte PCR test)	DERM	Skin, Hair, Nails	3-7 days
Galactomanan (Aspergillus Antigen)	SGAL	B	2 weeks
Gonorrhoea Culture – Cervix  † ‡ † The optimal sample type from the female genita an endocervical swab. Gonorrhoea does not survive w the endocervical epithelium; a negative gonorrhoea co from a vaginal swab is not reliable for excluding infec	vell outside ulture result	CS***	3-5 days
Gonorrhoea Culture – Rectal	GONR	CS	3-5 days
Gonorrhoea Culture – Throat	GONT	CS	3-5 days
Gonorrhoea Culture – Urethral	GONU	CS	3-5 days
Gonorrhoea Culture – Other site	GONO	CS	3-5 days
Group B Strep – Vaginal and Rectal (Self-collect)	GBSX	Blue gel Amies swab x2	3-5 days

TEST	CODE	SAMPLE REQS	TAT
Group B Strep – Vaginal and Rectal (STM)	GBSX	2 x <b>STM</b>	3-5 days
H. pylori Antigen – Stool (RF)	HBAG	RF	3 days
H. pylori Antigen – Stool (Self-collect)	HBAG	Stool/faecal container	3 days
H. pylori Culture	HPCU	J	1 month
Histoplasma Antigen	HANT	RU	3 days
Fresh urine less than 24 hours, directly to Mycology.			
HVS	HVS	STM/CS	2-4 days
IUCD for Culture	IUCD	Send Device	11-12 days
Legionella Urine Antigen	LEGA	Urine with boric acid	1 day
MRSA (Rapid PCR) one swab per site	MRSA	Blue liquid Amies swab	1 day
MRSA (Rapid PCR) one swab per site x 2	MRS2	Blue liquid Amies swab x 2	1 day
MRSA Culture one swab per site	MRSW	Blue liquid Amies swab	2 days
MRSA Culture one swab per site x 2	MRW2	Blue liquid Amies swab x 2	2 days
MRSA Culture (Self-collect) – Nose/Groin	MRW2	Purple liquid Amies swab x2	2 days
MRSA PCR (Self-collect) – Nose/Groin	MRS2	Purple liquid Amies swab x2	1 day
Pleural Fluid for Culture	FLUP	SC	7 days
Pneumococcal Antigen	PNAG	Urine with boric acid	1 day
Pneumocystis jiroveci (PJP) PCR	MPCP	SC BAL#	2-3 days
‡ ‡ BAL: Induced sputum or bronchoalveolar larage.			
Rapid Strep PCR (incl. m/c/s)  ** Do not use a black swab for RAPS. Use Blue only. I PCR is reported within 4 hours with full culture to foll		Blue liquid Amies swab**	1-3 days**
Schistosoma (Urine)	USCH	Mid-morning terminal urine following exercise 14	1-2 days
Sellotape Test	SELL	Send Sample***	1 day
*** Use clear Sellotape only and attach to slide.			
Semen Culture	SPCU	Semen	2-4 days
Skin Scrapings/Mycology by PCR	DERM	Send Sample	3-7 days
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 by PCR	PENT	<b>RF</b> <sup>††</sup>	2-3 days
† † 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]. Unless relevant clinical details are provided MOCP will not be done.			

TEST	CODE	SAMPLE REQS	TAT
Stool for OVA Cysts & Parasites by Microscopy	MOCP	RF	2 days
Stool Reducing Substances	STRS	RF <sup>7</sup>	2-3 weeks
Swab (Cervical)	CERS	STM / CS	2-4 days
Swab (Ear) Includes fungal culture over 7 days.	EARS	STM	2-4 days (Culture) 8-9 days (Fungal) – same swab
Swab (Eye)	EYES	STM	2-4 days
Swab (Nasal)	NASS	STM	2-4 days
Swab (Oral)	ORSW	STM/CS	2-4 days
Swab (Penile)	PENS	STM/CS	2-4 days
Swab (Rectal)	RECG	STM/CS	2-4 days
Swab (Skin) Specify sample site swab taken from.	SKIS	STM	2-4 days
Swab (Throat)	THRS	STM	2-4 days
Swab (Urethral)	URES	STM/CS	2-4 days
Swab (Vaginal)	VAGS	STM/CS	2-4 days
Swab (Vulval)	VULV	STM/CS	2-4 days
Swab (Wound) Specify sample site.	WOUS	STM	2-4 days
Synovial Fluid (for microscopy, crystals and culture)	FLU2	SC <sup>†††</sup>	14 days
† † † If prosthetic joint is present please state in clini to ensure that enrichment culture is prolonged for 14			
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 PCR (PCR detection of Mycobacterium tuberculosis complex and mutations for Rifampicin resistance)  This test is automatically added if the direct auramine	TBPC	All samples except blood cultures and urine, as clinically requested.	1 day
is positive on AFB culture requests and no known pre			
TB Slopes – Confirmation and Sensitivity	TBSL	<b>TB slope</b> (LJ medium-green) <sup>6</sup>	up to 8 weeks
Tissue for culture Specify sample site.	TISS	Tissue sample	up to 14 days
Urine (Microscopy Only)	UMIC	RU	1 day
Urine Chemistry and Microscopy (Self-collect)	UMIC	Urine (Universal). Mid stream.	1-2 days

TEST	CODE	SAMPLE REQS	TAT	
Urine Chemistry, Microscopy and Culture (Self-collect)	UCEM	Urine (Universal & Boric). Mid stream.	1-2 days	
Urine for Extended Culture	UCXD	MSU ††††	up to 7 days	
† † † † Optimal sample type for urine culture is a clean catch urine sent in a sterile pot containing preservative. Testing must be done within 24 hou	boric acid	ity.		
Urine for Microscopy and Culture	UCEM	MSU ††††	1-2 days	
† † † † Optimal sample type for urine culture is a catch urine sent in a sterile pot containing boric a				

# 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 103 cfu/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 LIKAS-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 (< 103 cfu/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.

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.

# **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.

# **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.



# **Swabs: Types and Codes**

**Patient Request Forms** and **Swabs** must 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.

#### **Culture Swabs**

SITE	CODE	SAMPLE TYPE
Candida only swab	CANC	Black or Blue Micro Swab
Cervical swab	CERS	Black or Blue Micro Swab
Ear swab	EARS	Blue or Orange Micro Swab
Eye swab	EYES	Blue or Orange Micro Swab
Gonorrhoea	GONN	Black Charcoal Swab (specify site)
High vaginal swab	HVS	Black or Blue Micro Swab
Nasal swab	NASS	Blue or Orange Micro Swab
Oral swab	ORSW	Black or Blue Micro Swab
Penile swab	PENS	Black or Orange Micro Swab
Rectal swab	RECG	Black or Blue Micro Swab
Skin swab	SKIS	Blue Micro Swab
Throat swab	THRS	Blue Micro Swab
Urethral swab	URES	Orange Micro Swab or Orange Charcoal Swab
Vaginal swab	VAGS	Black or Blue Micro Swab
Vulval swab	VULV	Black or Blue Micro Swab
Wound swab	WOUS	Black or Blue Micro Swab (specify site)

#### **MRSA by Culture**

Order code: MW171

MRW2	<b>Blue</b> Micro Swab x 2 – state sites
MRSW	<b>Blue</b> Micro Swab x 1 – state site
CODE	SAMPLE TYPE

Order code: MW170
Dark Blue Microbiology Swab (Culture)

Black Charcoal Microbiology Swab (Culture)

# **Rapid MRSA by PCR**

CODE	SAMPLE TYPE
MRSA	Blue Micro Swab x 1 – state site
MRS2	<b>Blue</b> Micro Swab x 2 – state sites

Order code: **MW172C**Orange Wire Shaft Microbiology Swab (Charcoal

Order code: MW172P

Orange Wire Shaft Microbiology Swab (Culture)

#### **PCR Swabs**

CODE	SAMPLE TYPE
SPCR	PCR Swab: Aptima Swab
SCG	PCR Swab: Aptima Swab
RSCG	PCR Swab: Aptima Swab
TSCG	PCR Swab: Aptima Swab
SCGT	PCR Swab: Aptima Swab
SGTM	PCR Swab: Aptima Swab
GVPC	PCR Swab: Aptima Swab
SGON	PCR Swab: Aptima Swab
DUCR	PCR Swab: Aptima or Green Swab
HERS	PCR Swab: Aptima or Green Swab
HP20	PCR Swab / Qvintip (HPV kit)
LGVP	PCR Swab: Aptima Swab
MGR	PCR Swab: Aptima Swab
MPXV	PCR Swab: Aptima Swab
MGEN	PCR Swab: Aptima Swab
MUPC	PCR Swab: Aptima Swab
SYPS	PCR Swab: Aptima or Green Swab
TVPC	PCR Swab: Aptima Swab
UGEN	PCR Swab: Aptima Swab
STD8	PCR Swab: Aptima Swab and Blue or Orange Micro Swab
VPR	PCR Swab: Aptima or Green Swab
NCOV	PCR Swab: Aptima or Green Swab
PERP	Orange PCR Swab / Prenasal (posterior nasopharynx) Swab
	SPCR SCG RSCG TSCG SCGT SGTM GVPC SGON DUCR HERS HP20 LGVP MGR MPXV MGEN MUPC SYPS TVPC UGEN STD8 VPR NCOV

Order code: MW142

Orange Aluminium Wire Dry Microbiology Swab (PCR)

Order code: MW951S

Green Sigma Virocult Swab and Vial (PCR)

Order code: PRD-03546

Aptima Multitest Swab Specimen Collection Kit

# 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	Skin, Hair and Nail.	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> <li>Dermatophyte PCR has replaced microscopy for Skin, Hair and Nail (72 Hour TAT).</li> <li>Non-dermatophyte culture will take 7 days.</li> <li>Microscopy is carried out to confirm significance of rare fungi.</li> <li>Pseudomonas investigation in Nail specimens is available on request.</li> </ul>	<ul> <li>Non-sterile specimen fungal cultures are performed on Sabouraud's agar plates for 7 days with no microscopy.</li> <li>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.</li> </ul>

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

#### **Stool for OCP**

Sample type		Comments
Stool	Please request as <b>MOCP</b> Requests for MOCP 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 ty	уре	Comments
Stool Please	Please request as <b>CLOS</b>	Two tier PCR and Toxin c.diff
	Serosep Enteric Bio PCR	screening based on PHE guidance.
	Alere Techlab EIA (Toxin)	

**Enteric Organism Rapid Detection – see Tropical Immunology page 93** 

CODE	SAMPLE REOS	TAT
DEOX	B	10 days
11DC	B (Frozen)	10 days
170H	B	5 days
ACTH	A (EDTA on ice, Plasma, spun and frozen within 2 hours) <sup>41</sup>	1 day
ALDN	A or B	5 days
UALD	PU	5 days
AFP	B	1 day
TAME ole	(TDL Tiny)	1 day
AMEN	B	1 day
ANDP	88	1 day
ANDR	B	5 days
ADH	A (Plasma Frozen) <sup>4</sup>	10 days
AMH	(TDL Tiny) or (TDL Tiny)	1 day
AMH	<b>B</b>	1 day
BNP	В	1 day
CPEP	B	3 days
CAT0	⊕ (Frozen) <sup>4</sup>	1 day
CATE	A (Plasma Frozen) <sup>4</sup>	5 days
UCAT	PU (collect on acid) <sup>1</sup>	5 days
CORT	(TDL Tiny)	1 day
UCOR	CU	5 days
CORT	B	1 day
DHEX	B	7-10 days
UDHE	CU	3 weeks
DHEA	(TDL Tiny) or (TDL Tiny)	1 day
DHEA	B	1 day
DHT	BB	7 days
HCGF/ PAPA	<b>B</b>	1 day
DRP	<b>□</b> DRP form <sup>7,8</sup>	5 days
	11DC 17OH ACTH ALDN UALD AFP TAME AMEN ANDP ANDR ANDR AMH AMH CPEP CATO CATE UCAT UCOR CORT UCOR CORT UCOR UCOR CORT UCOR COR COR COR COR COR COR COR COR COR	DEOX : (Frozen)  11DC : (Frozen)  17OH : (EDTA on ice, Plasma, spun and frozen within 2 hours) (Frozen)  ALDN

TEST	CODE	SAMPLE REQS	TAT
Down Syndrome Risk Profile with risk calculation first trimester	DRP	B DRP form + image of scan <sup>7,8</sup>	5 days
<b>Erectile Dysfunction Profile</b> Please clearly state fasting or non-fasting status.	IMP0	<b>ABBG</b>	3 days
Fasting Insulin Resistance Index (FIRI) Both samples taken at the same time and must be fast Please indicate fasting status clearly on request form.	FIRI ing.	<b>3 6</b>	1 day
Female Hormone Profile (LH, FSH, PROL, TOES) (Self-collect) Avoid taking samples from any area an HRT cream is applied.	TFIP	(TDL Tiny) and (3) (TDL Tiny)	1 day
Female Hormone Profile (Venous)	FIP	<b>B</b>	1 day
First Trimester Maternal Screen (PAPP-A/Free Beta-hCG) (Risk to be calculated by requesting clinician)	FTMS	<b>B</b>	1 day
Free T3 (Self-collect)	FT3	(TDL Tiny)	1 day
Free T3 (Venous)	FT3	В	1 day
Free T4 (Self-collect)	FT4	(TDL Tiny)	1 day
Free T4 (Venous)	FT4	В	1 day
FSH (Self-collect)	FSH	(TDL Tiny) or (7DL Tiny)	1 day
FSH (Venous)	FSH	В	1 day
Growth Hormone (Fasting)	GH	<b>3</b> 7,35	1 day
Gut Hormone Profile	GUTP	(Frozen within 15 minutes) <sup>41</sup>	3 weeks
HCG (Quantitative) (Self-collect)	QHCG	(TDL Tiny)	1 day
HCG (Quantitative) (Venous)	QHCG	<b>B</b>	1 day
Hirsutism Profile	HIRP	<b>B</b>	1 day
HRT Profile 1	HRT	<b>B</b>	1 day
HRT Profile 2 Please clearly state fasting or non-fasting status.	HRT2	<b>B G</b>	1 day
IGF-1 (Somatomedin)	SOMA	(Frozen) <sup>4,7</sup>	1 day
IGF-BP3	IGF3	B (Frozen)⁴	5 days
Impotence Profile	IMP0	ABBG	3 days
Inhibin A	INIA	В	1 month
Inhibin B	INIB	(Day 3 of cycle, frozen)	5 days
Insulin	INSU	<b>B</b> 4,7	1 day
Luteinising Hormone (LH) (Self-collect)	LH	(TDL Tiny) or (TDL Tiny)	1 day
Luteinising Hormone (LH) (Venous)	LH	3	1 day
Macroprolactin	PRLD	В	4 days

TEST Date Harrison Burgle	CODE	SAMPLE REQS	TAT
Male Hormone Profile	MIPR	В	1 day
Melatonin (Serum)	MEL	(Frozen)	5 days
Melatonin (Urine)	UMEL	CU <sup>13</sup>	2 weeks
Menopausal Profile (FSH, LH, TOES, TSH, FT4) (Self-collect) Avoid taking samples from any area an HRT cream is applied.	TMEN	(TDL Tiny) and (3) (TDL Tiny)	1 day
Menopause Profile (Venous)	MENO	<b>B</b>	1 day
Metabolic Syndrome Profile Please clearly state fasting or non-fasting status.	METS	<b>ABB</b> G	9 days
Metanephrines (Plasma) Must be frozen within 2 hours.	PMET	(Frozen plasma, must be frozen within 2 hours)	7 days
Metanephrines (Urine)	UMEX	PU (collect on acid) <sup>1</sup>	5 days
Oestradiol-17-Beta (Self-collect)	T0ES	(TDL Tiny)	1 day
Oestradiol-17-Beta (Venous)	0EST	<b>B</b>	1 day
Oestriol (Estriol)	E3	BB	4 days
Oestrone	E1	BB	4 days
Osteocalcin	0ST	B (Frozen)⁴	4 days
Parathyroid Hormone (Whole) Requires its own EDTA tube, if other tests require EDTA an extra EDTA sample should be taken for PTHI.	PTHI	<b>A</b> 4	1 day
Pituitary Function Profile	PITF	<b>BB</b> <sup>7</sup>	1 day
Polycystic Ovary Syndrome Profile Please clearly state fasting or non-fasting status.	PCOP	<b>ABBB G</b> <sup>7</sup>	5 days
Polycystic Ovary Syndrome SHORT Please clearly state fasting or non-fasting status.	PCOS	<b>A B G</b>	1 day
Pregnancy (Serum) [Quantitative]	QHCG	B	1 day
Pregnenolone	PREN	<b>B</b>	15 days
Progesterone (Self-collect) Avoid taking samples from any area an HRT cream is applied.	PROG	(TDL Tiny) or (TDL Tiny)	1 day
Progesterone (Venous)	PROG	<b>B</b>	1 day
Proinsulin	PROI	(Frozen plasma) <sup>4</sup>	5 days
Prolactin (Macro)	PRLD	В	4 days
Prolactin (Self-collect)	PROL	(TDL Tiny) or (TDL Tiny)	1 day
Prolactin (Venous)	PROL	B	1 day
Renin	RENI	A (Frozen plasma) <sup>36</sup>	5 days
Reverse T3	RT3	<b>B</b> 7,37	15 days
Serotonin	SERT	(Frozen whole blood) <sup>1</sup>	10 days
-		,	

TEST	CODE	SAMPLE REQS	TAT
Serotonin (Urine)	USER	PU 50mls (Frozen) <sup>1</sup>	5 days
Sex Hormone Binding Globulin (Self-collect)	SHBG	(TDL Tiny) or (TDL Tiny)	1 day
Sex Hormone Binding Globulin (Venous)	SHBG	B	1 day
Somatomedin (IGF-1)	SOMA	(Frozen) <sup>4,7</sup>	1 day
Т3	T3	B	1 day
T3 (Reverse)	RT3	B 7,37	15 days
<b>Testosterone (Self-collect)</b> Avoid taking samples from any area an HRT cream is applied.	TEST	(TDL Tiny) or (TDL Tiny)	1 day
Testosterone (Total), LC MS Mass Spec NEW	MSTT	B	5-7 days
Testosterone (Venous)	TEST	<b>B</b>	1 day
<b>Testosterone (Free) (Self-collect)</b> Avoid taking samples from any area an HRT cream is applied.	FTES	(TDL Tiny) or (TDL Tiny)	3 days
Testosterone (Free) (Venous)	FTES	<b>B</b>	3 days
Thyroglobulin Abs	TGAB	<b>B</b>	1 day
Thyroglobulin Assay	TGA	B	1 day
Thyroid Abs (Thyroglobulin + Thyroid Peroxidase Abs) (Self-collect)	THAB	(TDL Tiny)	2 days
Thyroid Abs (Thyroglobulin + Thyroid Peroxidase Abs) (Venous)	THAB	<b>B</b>	1 day
Thyroid Peroxidase Antibodies/Anti TPO	TPEX	В	1 day
Thyroid Profile 1 (FT4/TSH) (Self-collect)	TF	(TDL Tiny)	1 day
Thyroid Profile 1 (FT4/TSH) (Venous)	TF	В	1 day
Thyroid Profile 2 (Venous)	TF2	В	2 days
Thyroid Profile 3 (FT3/FT4/TSH) (Self-collect)	TF3	(TDL Tiny)	1 day
Thyroid Profile 3 (FT3/FT4/TSH) (Venous)	TF3	В	1 day
Thyroxine (T4)	T4	В	1 day
Thyroxine Binding Globulin	TBG	(Frozen)	10 days
Total Testosterone, LC MS Mass Spec NEW	MSTT	В	5-7 days
TSH (Self-collect)	TSH	(TDL Tiny)	1 day
TSH (Venous)	TSH	B	1 day
TSH-Receptor Antibodies	TSI	В	4 days
Ziwig Endotest® For information about this test and to order kits please contact endotest@tdlpathology.com. The quality of the sample collection is important. Samples can be collecte the clinic or at home following instructions provided.		Endotest saliva collection kit	25 days

# Reproductive Immunology at Rosalind Franklin Laboratory, Chicago, USA

TEST	CODE	SAMPLE REQS	TAT
Reproductive Immunophenotype Panel	3RF	000	1 week
NK Assay/Cytotoxicity Panel	4RF	000	1 week
NK Assay Follow-Up Panel	5RF	000	1 week
TH1/TH2 Cytokine Ratio	6RF	<b>OOO</b> <sup>5</sup>	1 week
Leucocyte Antibody Detection Panel MALE	7RF	<b>(1)</b> (1) (6,34)	1 week
Leucocyte Antibody Detection Panel FEMALE	8RF	B	1 week
HLA DR Antigens	9RF	AA	2 weeks
HLA DQ Alpha Antigens	10RF	AA	2 weeks
HLA DQ Beta Antigens	11RF	AA	2 weeks
HLA A, B, C	14RF	AA	2 weeks
NK Assay Panel + Intralipids	16RF	000	1 week
KIR (Killer-like Immunoglobulin-like Receptors) Genotyping	17RF	AAA	2-3 weeks
TH1/TH2 Intracellular Cytokine Ratios with IVIG, Prednisolone	20RF	<b>000</b> <sup>5</sup>	1 week
TH1/TH2 Intracellular Cytokine Ratios with IVIG	21RF	<b>000</b> <sup>5</sup>	1 week
TH1/TH2 Intracellular Cytokine Ratios with Prednisolone	22RF	<b>000</b> <sup>5</sup>	1 week
T Regulatory Cells	25RF	0	3 days
HLA-C	26RF	AA	2 weeks
Decidualization Score (DS)	DSRF	J (Contact Referrals)	2-3 weeks
PAI-1 4G/5G Polymorphism	PAIP	A	10 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 from St Helier**

TEST	CODE	SAMPLE REQS	TAT
NK (CD69) Cell Assay	CD69	<b>(</b> )*	Send Mon-Thurs only
NK (CD69) and NK Cytotoxicity	69C	<b>000</b> *	Send Mon-Thurs only
NK Cytotoxicity Assay	HSNK	<b>000</b> *	Send Mon-Thurs only
NK Cytotoxicity with suppression with steroid, IVIg and intralipin, and NK (CD69) cell assay	69CI	<b>000</b> *	Send Mon-Thurs only
NK Cytotoxicity with suppression, steroid, IVIg & Intralipin	NKCY	<b>000</b> *	Send Mon-Thurs only
Suppression with steroid, IVIg and intralipin, NK (CD69) cell assay, TH1/TH2 cytokines	NCIT	<b>0000</b> *	Send Mon-Thurs only
TH1/TH2 Cytokine Profile	1TH2	000*	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.

# **Amenorrhoea Profile (Venous)**

Luteinising Hormone (LH) **FSH** Prolactin Testosterone Oestradiol-17-Beta Sex Hormone Binding Globulin (SHBG) Free Androgen Index

#### TAT: 1 day

#### **AMEN**



#### **Andropause Profile**

**DHEA Sulphate** FSH Testosterone Free Androgen Index Luteinising Hormone (LH) Sex Hormone Binding Globulin (SHBG)

#### TAT: 1 day

ANDP



# **Female Hormone Profile (Venous)**

Luteinising Hormone (LH) FSH Prolactin Oestradiol-17-Beta

TAT: 1 day

FIP



#### **Erectile Dysfunction Profile**

Lipid Profile Glucose HbA1c Free T4 / TSH Prolactin **Total Testosterone** Free Testosterone PSA

Sex Hormone Binding Globulin (SHBG) Free Androgen Index

Please clearly state fasting or non-fasting status.

#### TAT: 3 days

IMP0





# **First Trimester Maternal** Screen (PAPP-A/ Free Beta-hCG) (Risk to be calculated by requesting clinician)

Free B-hCG PAPP-A

Free B-hCG and PAPP-A in serum and sonographic determination of nuchal translucency (NT) are markers of choice to identify women at increased risk of Down Syndrome during the first trimester (week 11-13) of pregnancy.

#### TAT: 1 day

#### **FTMS**



#### **Hirsutism Profile**

DHEA sulphate FSH Luteinising Hormone (LH) Sex Hormone Binding Globulin (SHBG) Testosterone

#### TAT: 1 day

# HIRP



#### **HRT Profile 1**

**FSH** Oestradiol-17-Beta Progesterone

# TAT: 1 day

HRT

**B** 

#### **HRT Profile 2**

Lipid Profile (Cholesterol. Triglycerides, HDL Cholesterol, LDL Cholesterol (Calculated), Non-HDL Cholesterol)

Glucose Free T4 **TSH FSH** 

Oestradiol-17-Beta

Please clearly state fasting or non-fasting status.

#### TAT: 1 day

HRT2



# Impotence Profile

Lipid Profile Glucose HbA1c **TSH** Prolactin Total Testosterone

Free Testosterone

PSA

Sex Hormone Binding Globulin (SHBG)

Free Androgen Index

Please clearly state fasting or non-fasting status.

#### TAT: 3 days

IMP0



#### **Male Hormone Profile**

FSH

Luteinising Hormone (LH)

Testosterone

Free Androgen Index Prolactin

Sex Hormone Binding Globulin (SHBG)

TAT: 1 day

MIPR

B

### **Menopause Profile** (Venous)

FSH

Luteinising Hormone (LH) Oestradiol-17-Beta

Free T4 **TSH** 

TAT: 1 day

**MENO** 

B

# **Metabolic Syndrome Profile**

Lipid Profile Glucose HbA1c

Insulin

C Reactive Protein (High Sensitivity) Adiponectin

Please clearly state fasting or non-fasting status.

TAT: 9 days

METS





#### **Pituitary Function Profile**

**TSH FSH** 

Luteinising Hormone (LH)

Prolactin

Growth Hormone (Fasting)

Cortisol

Please provide details of time of day sample is taken. Patient should be resting for 30 mins before sample taking.

#### TAT: 1 day

PITF



# **Polycystic Ovary** Syndrome SHORT

Testosterone

Sex Hormone Binding Globulin (SHBG)

Free Androgen Index

Luteinising Hormone (LH)

Glucose Insulin

Lipid Profile

Free T4 / TSH

HbA1c

Please clearly state fasting or non-fasting status.

#### TAT: 1 day

**PCOS** 



# **Polycystic Ovary Syndrome Profile**

Testosterone

TSH

Glucose

HbA1c

**FSH** 

**DHEA Sulphate** 

Insulin

Luteinising Hormone (LH)

17 Hydroxyprogesterone

Lipid Profile

Prolactin

Cortisol

Antimullerian Hormone Androstenedione

Sex Hormone Binding

Globulin (SHBG)

A fasting 9.00am sample is recommended. Please clearly state fasting or non-fasting status.

#### TAT: 5 days

**PCOP** 







# **Thyroid Profile 1** (FT4/TSH) (Venous)

Free T4 TSH

#### TAT: 1 day

TF

# **Thyroid Profile 2** (Venous)

**TSH** 

Free T3 Free T4

Thyroglobulin Abs Thyroid Peroxidase

TAT: 2 days

TF2



# **Thyroid Profile 3** (FT3/FT4/TSH) (Venous)

Free T3 Free T4 TSH

TAT: 1 day

TF3



# Ziwig Endotest®

Ziwig Endotest® is a new non-invasive diagnostic test for reliable and rapid diagnosis for all types of endometriosis. The test relies on Next Generation Sequencing of micro RNA present in saliva and on the use of Artificial Intelligence to process the very large volume of data generated. Ziwig Endotest® is an in vitro diagnostic test for diagnosis of endometriosis on salivary samples with sensitivity 97.4% and specificity 93.7%.

About 10% of all women are affected by endometriosis with average times for diagnosis of around 9 years. Patients not infrequently see up to 10 doctors before being diagnosed with endometriosis (MRI, pelvic ultrasound, laparoscopy). Ziwig Endotest® is able to detect all types of endometriosis, mild and advanced. Endometriosis is not infrequently mistaken for other conditions that can cause pelvic pain, such as pelvic inflammatory disease (PID) or ovarian cysts. The effects of endometriosis range from asymptomatic, often identified during investigations for infertility, to chronic or progressively severe symptomatic related conditions. Once diagnosed, optimised clinical management of endometriosis would apply.

Laparoscopy is the gold standard for diagnosis of endometriosis but is relatively expensive, invasive, and requires an anaesthetic. The saliva sample required for the test is straight forward to collect, non-invasive and provides a definitive diagnosis even in the most complex cases. It has been validated by one of the largest clinical studies in the world.

Ziwig Endotest® provides a Bioinformatics Approach to microRNA sequencing analysis, from saliva.

For access to scientific publications visit: ziwig.com/en/our-publications/

- Clear Positive/Negative result
- Definitive diagnosis for all forms of endometriosis
- Non-invasive, saliva sample
- Cost contained single test outcome

For information about this test and to order kits please contact endotest@tdlpathology.com

The quality of the saliva sample collection is important. Samples can be collected under supervision of a referring clinician or self-collected at home following the instruction video:

www.tdlpathology.com/ziwig-endotest



# LC MS Mass Spec Total Testosterone NEW

Testosterone must be the world's most discussed hormone, but general understanding about testing is much less well known. Immunoassay methods are widely used to measure testosterone, providing a reliable method of measurement in many use cases. However, whilst this is accepted for standard testing, this methodology carries a possibility of analytical interference and cross-reactivity with structurally similar biological compounds, which can lead to a false, overestimation of circulating testosterone concentration.

Measurement of total testosterone by tandem mass spectrometry (LC-MS/MS) is a methodology with greater specificity, providing a more accurate measurement of testosterone, important when assessing lower levels of testosterone found in females, children and hypogonadal males.

The British Menopause Society recommends LC-MS/MS for the measurement of total testosterone levels, both to exclude high baseline concentrations before treatment is commenced and to ensure that levels remain within the female physiological range when monitoring supplementation (https://thebms.org.uk/wp-content/uploads/2022/12/08-BMS-TfC-Testosterone-replacement-in-menopause-DEC2022-A.pdf)

See page 60 for test information.

Female Reference		0.7–2.8 nmol/L (normal)	
Ranges		0-0.7 (low)	
Male Reference Ranges	15 years	0.9-2.7 nmol/L	
	15-49 years	9.2-55.8 nmol/L	
	50+ years	6.3-26.5 nmol/L	

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.

### **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, generally no less than 2 days and no longer than 7 days before the test is acceptable. 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. For other semen tests like ROS and DNA fragmentation the abstinence period is minimum 2 days but no more than 3 days. 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.

TDL Andrology provides reference values to those given in the most recent WHO guidelines (2021). WHO 2021 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 online at **www.tdlpathology.com/andrologybooking** or by calling **020 7025 7940**. There is an attendance fee of £50.00 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 including clinical interpretation by the consultant please contact TDL Andrology on **020 7025 7940** or email **andrology@tdlpathology.com** for further information



Book an appointment online:

www.tdlpathology.com/andrologybooking

SCAN MF

TEST	CODE	SAMPLE REQS	TAT
Individual Semen Parameters***  *** Semen parameters may be requested individually (e.g. count only, vitality only, motility etc.). Please request as SPOD and indicate on the request form which parameter is required.	SPOD	Semen <sup>1</sup>	1 day
Oxidative Stress in Semen (ROS + MIOXSYS)	SROS	Semen 1	1 day
Retrograde Ejaculation	RTR0	Contact lab	2 days
Semen Analysis, Comprehensive* * If required, comprehensive semen analysis can be reported within 4 hours, with morphology to follow.	SPER	Semen <sup>1</sup>	2 days*
Semen Analysis, Post-Vasectomy**	PVAS	Semen 1	2 days

<sup>\*\*</sup> 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:

- · Analysis of post vasectomy semen samples should not occur until 12 weeks post-surgery and after a minimum of 20 ejaculates
- 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
- . Semen should be provided in weighed specimen containers provided by TDL Andrology
- . Sexual abstinence should be between 2 and 7 days.

Semen Analysis, Vasectomy Reversal* * If required, comprehensive semen analysis can be reported within 4 hours, with morphology to follow.	SPER	Semen 1	2 days*
Semen Culture	SPCU	Semen	2-4 days
Semen Fructose (Qualitative assessment)	SPCF	Semen	2 days
Semen Leucocytes	PMNS	Semen	2 days
Semen Zinc	SPCZ	Semen	up to 10 days
Sperm Aneuploidy	SPPL	Semen 1	4 weeks
Sperm Antibodies (Serum)	ASAB	В	5 days
Sperm Antibodies/MAR Test (Semen) <sup>†</sup> † Sperm antibodies in semen are measured as part of the routine semen analysis.	ASPA	Semen	1 day
Sperm Comet®	CMET	Semen 1	1-2 weeks
Sperm Comet® Exact Focus	CMT2	Semen 1	1-2 weeks
Sperm Comet® Extend	CMT3	Semen 1	1-2 weeks
Sperm Comet® Extend Focus	CMT4	Semen 1	1-2 weeks
Sperm Count (Post-Vasectomy)	PVAS	Semen 1	2 days
Sperm DNA Fragmentation (SCSA type test)	SEXT	Semen 1	1-2 weeks
Sperm Morphology (Kruger strict criteria)	MRPH	Semen 1	2 days

#### By special arrangement

- 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 (SDF) is associated with reduced natural pregnancy rates and assisted conception pregnancy rates as well as live birth rates. Sperm DNA fragmentation also 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. Sperm DNA can be damaged when sperm are being made in the testes or as they mature before ejaculation. This damage breaks the DNA into fragments, so sperm DNA tests are also known as sperm DNA fragmentation tests. Men with high levels of sperm DNA damage are less likely to get their partner pregnant and have increased risk of miscarriage. Even if semen analysis results are 'normal', the sperm DNA could be damaged and therefore poor quality.

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

# Sperm Chromatin Structure Assay (SCSA type test) [SEXT]

This test has the ability to measure large numbers of cells (between 5,000 and 15,000 sperm), rapidly in an ejaculate. The SCSA test monitors the changes

in fluorescence of a probe using Acridine Orange Flowcytometry test, to detect both single and double DNA strand breaks. 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 100,000/ml.

# Sperm COMET® Assay [CMET/CMET2/CMET3/CMET4]

Exact® tests, powered by SpermComet technology measure sperm DNA damage.

The COMET® assay sperm DNA fragmentation (SDF) tests from Examen® provide one of the highest performance levels among commercially available SDF tests. The COMET® tests directly measure total single- and double-strand DNA breaks (CMET/CMT2), and double-strand breaks only (CMT3/CMT4). Use Examen's COMET® tests for

- couples struggling to achieve pregnancy (idiopathic infertility), especially where a semen analysis was 'normal', or where there have been repeated failed IVF cycles (CMET/CMT2)
- couples struggling to keep the pregnancy/ suffering multiple recurrent pregnancy loss, especially if multiple early losses <14 weeks, and female aged mid-thirties and above (CMT3/CMT4)

It has the advantage of requiring only 5000 sperm so it can be used for men with low sperm counts and for surgically retrieved sperm samples. There are 4 types of COMET tests available at different costs. The clinician must select the desired type prior to sending the patient as well as ensuring completion of the Examen request form.

# **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.

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 fragmentation 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, CMET1-CMET4 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 andthe sperm count. For CMET test please ensure a completed Examen request form accompanies the TDL request form.

A count of a minimum 0.1 million/ml is required for accurate DNA fragmentation 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.

#### **TDL Andrology**

• 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.

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.

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. Appointments can also be booked via **www.tdlpathology.com/andrologybooking** 

#### 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

#### References

Vassiliou A, Martin CH, Homa ST, Stone J, Dawkins A, Genkova MN, Skyla Dela Roca H, Parikh S, Patel J, Yap T, Killeen AP. Redox potential in human semen: Validation and qualification of the MiOXsys assay. Andrologia. 2021 Mar;53(2):e13938. doi: 10.1111/and.13938. Epub 2020 Dec 30. PMID: 33377541.

TEST	CODE	SAMPLE REQS	TAT
7 STI Profile by PCR (7 tests from 1 Sample)	DL12	FCRU / PCR Swab / TPV	2 days
7 STI Profile by PCR (7 tests from 1 Sample) (Self-collect)	DL12	Aptima urine or multisite swab	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 (PCR)	RSCG	PCR	2 days
Chlamydia/Gonorrhoea – Rectal (Self-collect)	RSCG	Aptima multisite swab	2 days
Chlamydia/Gonorrhoea – Thin Prep	TCG	TPV	2 days
Chlamydia/Gonorrhoea – Throat (PCR)	TSCG	PCR	2 days
Chlamydia/Gonorrhoea – Throat (Self-collect)	TSCG	Aptima multisite swab	2 days
Chlamydia/Gonorrhoea – Urine (FCRU)	CCG	FCRU	2 days
Chlamydia/Gonorrhoea – Urine (Self-collect)	CCG	Aptima urine	2 days
Chlamydia/Gonorrhoea – Vaginal (Self-collect)	SCG	Aptima multisite swab	2 days
Chlamydia/Gonorrhoea/Trichomonas – PCR Swab	SCGT	PCR	2 days
Chlamydia/Gonorrhoea/Trichomonas – Thin Prep	TCGT	TPV	2 days
Chlamydia/Gonorrhoea/Trichomonas – Urine	CCGT	FCRU	2 days
			<u> </u>
CT/GC/Trichomonas/Mgen – PCR Swab	SGTM	PCR Swab	2 days
CT/GC/Trichomonas/Mgen – PCR Swab CT/GC/Trichomonas/Mgen – Thin Prep	SGTM TGTM	PCR Swab TPV	2 days 2 days
CT/GC/Trichomonas/Mgen – Thin Prep	TGTM	TPV	2 days
CT/GC/Trichomonas/Mgen – Thin Prep CT/GC/Trichomonas/Mgen – Urine	TGTM CGTM	TPV FCRU	2 days 2 days
CT/GC/Trichomonas/Mgen – Thin Prep CT/GC/Trichomonas/Mgen – Urine Gardnerella vaginalis by PCR	TGTM CGTM GVPC	TPV FCRU FCRU / PCR / TPV	2 days 2 days 2 days
CT/GC/Trichomonas/Mgen – Thin Prep CT/GC/Trichomonas/Mgen – Urine Gardnerella vaginalis by PCR Gonorrhoea – PCR swab	TGTM CGTM GVPC SGON	TPV FCRU FCRU/PCR/TPV PCR	2 days 2 days 2 days 2 days
CT/GC/Trichomonas/Mgen – Thin Prep CT/GC/Trichomonas/Mgen – Urine Gardnerella vaginalis by PCR Gonorrhoea – PCR swab Gonorrhoea – Thin Prep	TGTM CGTM GVPC SGON TGON CGON GONC	TPV FCRU FCRU/PCR/TPV PCR TPV	2 days 2 days 2 days 2 days 2 days 2 days
CT/GC/Trichomonas/Mgen – Thin Prep CT/GC/Trichomonas/Mgen – Urine Gardnerella vaginalis by PCR Gonorrhoea – PCR swab Gonorrhoea – Thin Prep Gonorrhoea – Urine Gonorrhoea Culture – Cervix ‡ ‡ † The optimal sample type from the female genital tract is an endocervical swab. Gonorrhoea does not survive well outs the endocervical epithelium; a negative gonorrhoea culture re	TGTM CGTM GVPC SGON TGON CGON GONC	TPV FCRU FCRU/PCR/TPV PCR TPV FCRU	2 days
CT/GC/Trichomonas/Mgen – Thin Prep CT/GC/Trichomonas/Mgen – Urine Gardnerella vaginalis by PCR Gonorrhoea – PCR swab Gonorrhoea – Thin Prep Gonorrhoea – Urine Gonorrhoea Culture – Cervix ‡ ‡ † The optimal sample type from the female genital tract is an endocervical swab. Gonorrhoea does not survive well outs the endocervical epithelium; a negative gonorrhoea culture refrom a vaginal swab is not reliable for excluding infection.	TGTM CGTM GVPC SGON TGON CGON GONC S Sidde sult	TPV FCRU FCRU/PCR/TPV PCR TPV FCRU CS <sup>‡‡‡</sup>	2 days 3-5 days
CT/GC/Trichomonas/Mgen – Thin Prep CT/GC/Trichomonas/Mgen – Urine Gardnerella vaginalis by PCR Gonorrhoea – PCR swab Gonorrhoea – Thin Prep Gonorrhoea – Urine Gonorrhoea Culture – Cervix ‡‡ † The optimal sample type from the female genital tract is an endocervical swab. Gonorrhoea does not survive well outs the endocervical epithelium; a negative gonorrhoea culture refrom a vaginal swab is not reliable for excluding infection. Gonorrhoea Culture – Rectal	TGTM CGTM GVPC SGON TGON CGON GONC S side sult	TPV FCRU FCRU/PCR/TPV PCR TPV FCRU CS***	2 days 2 days 2 days 2 days 2 days 2 days 3-5 days
CT/GC/Trichomonas/Mgen – Thin Prep CT/GC/Trichomonas/Mgen – Urine Gardnerella vaginalis by PCR Gonorrhoea – PCR swab Gonorrhoea – Thin Prep Gonorrhoea – Urine Gonorrhoea Culture – Cervix ‡ ‡ † The optimal sample type from the female genital tract is an endocervical swab. Gonorrhoea does not survive well outs the endocervical epithelium; a negative gonorrhoea culture refrom a vaginal swab is not reliable for excluding infection. Gonorrhoea Culture – Rectal Gonorrhoea Culture – Throat	TGTM CGTM GVPC SGON TGON CGON GONC Sidde sult GONR GONT	TPV FCRU FCRU/PCR/TPV PCR TPV FCRU CS***	2 days 3-5 days 3-5 days 3-5 days
CT/GC/Trichomonas/Mgen - Thin Prep CT/GC/Trichomonas/Mgen - Urine Gardnerella vaginalis by PCR Gonorrhoea - PCR swab Gonorrhoea - Thin Prep Gonorrhoea - Urine Gonorrhoea Culture - Cervix ‡‡ † The optimal sample type from the female genital tract is an endocervical swab. Gonorrhoea does not survive well outs the endocervical epithelium; a negative gonorrhoea culture refrom a vaginal swab is not reliable for excluding infection. Gonorrhoea Culture - Rectal Gonorrhoea Culture - Throat Gonorrhoea Culture - Urethral	TGTM CGTM GVPC SGON TGON CGON GONC s ide sult GONR GONT GONU	TPV FCRU FCRU/PCR/TPV PCR TPV FCRU CS***  CS CS	2 days 2 days 2 days 2 days 2 days 2 days 3 days
CT/GC/Trichomonas/Mgen – Thin Prep CT/GC/Trichomonas/Mgen – Urine Gardnerella vaginalis by PCR Gonorrhoea – PCR swab Gonorrhoea – Thin Prep Gonorrhoea – Urine Gonorrhoea Culture – Cervix ‡ ‡ † The optimal sample type from the female genital tract is an endocervical swab. Gonorrhoea does not survive well outs the endocervical epithelium; a negative gonorrhoea culture refrom a vaginal swab is not reliable for excluding infection. Gonorrhoea Culture – Rectal Gonorrhoea Culture – Throat Gonorrhoea Culture – Urethral Gonorrhoea Culture – Other site	TGTM CGTM GVPC SGON TGON CGON GONC Sidde sult GONR GONT GONU GONU	TPV FCRU FCRU/PCR/TPV PCR TPV FCRU CS***  CS CS CS CS	2 days 2 days 2 days 2 days 2 days 2 days 3 days 2 days 3 days
CT/GC/Trichomonas/Mgen – Thin Prep CT/GC/Trichomonas/Mgen – Urine Gardnerella vaginalis by PCR Gonorrhoea – PCR swab Gonorrhoea – Thin Prep Gonorrhoea – Urine Gonorrhoea Culture – Cervix ‡ ‡ † The optimal sample type from the female genital tract is an endocervical swab. Gonorrhoea does not survive well outs the endocervical epithelium; a negative gonorrhoea culture refrom a vaginal swab is not reliable for excluding infection. Gonorrhoea Culture – Rectal Gonorrhoea Culture – Throat Gonorrhoea Culture – Urethral Gonorrhoea Culture – Other site Haemophilus ducreyi by PCR	TGTM CGTM GVPC SGON TGON CGON GONC Side GONR GONT GONU GONU	TPV FCRU FCRU/PCR/TPV PCR TPV FCRU CS****  CS CS CS CS CS CS CS PCR	2 days 2 days 2 days 2 days 2 days 2 days 3 days

TEST	CODE	SAMPLE REQS	TAT
Hepatitis C Antibodies (Self-collect)	THCV	(TDL Tiny)	1 day
Hepatitis C Antibodies (Venous)	HEPC	B	1 day
Hepatitis C Antigen (Early detection) (Self-collect)	TCAG	(TDL Tiny)	1 day
Hepatitis C Antigen (Early detection) (Venous)	HCAG	B	1 day
Herpes Simplex (HSV) 1 & 2 – Genital lesion (Self-collect)	HERS	Aptima multisite swab	5 days
Herpes Simplex (HSV) 1 & 2 – Oral lesion (Self-collect)	HERS	Aptima multisite swab	5 days
Herpes Simplex (HSV) 1 & 2 (PCR) (Oral or Genital)	HERS	PCR	5 days
Herpes Simplex I/II by PCR (Urine)	HERD	FCRU	5 days
HIV 1 & 2 Abs/p24Ag (Self-collect)	THIV	(TDL Tiny)	1 day
HIV 1 & 2/p24Ag (Venous)	HDU0	B	1 day
HIV/HBV/HCV (Early detection by PCR/NAAT) with Syphilis	STXX	B A 2 x 6mls or 2 x 4mls	3 days
HIV/HBV/HCV Screen by PCR/ NAAT (10 days post exposure)	STDX	A 2 x 6mls or 2 x 4mls (Vacutainer only)	3 days
HIV Rapid RNA HIV-1 QUALITATIVE	LHIV	(Vacutainer only)	1 day
HIV Rapid RNA HIV-1 QUANTITATIVE	RHIV	(Vacutainer only)	1 day
HPV (19 high risk DNA subtypes, reported as types 16, 18 or Others) (Self-collect)	HPVY	Qvintip vaginal swab	3 days
HPV (Individually typed high risk DNA subtypes) (Self-collect)	HPVZ	Qvintip vaginal swab	3 days
HPV (28 individually typed low risk (LR) & high risk (HR) DNA subtypes and reflexed mRNA for types 16, 18, 31, 33 and 45)	HPVT	TPV	5 days
HPV (28 individually typed LR & HR DNA subtypes)	HP20	TPV	3 days
HPV (A group of 14 HR mRNA types)	HPVH	TPV	3 days
Lymphogranuloma Venerium (LGV)  – Rectal (Self-collect)*  * This test can be configured to be automatically reflexed as required.	LGVP	Aptima multisite swab	1-2 weeks
Lymphogranuloma Venerium (LGV) (PCR)  * LGV can be added to a positive chlamydia sample using the same swab if requested within 4 days of receipt of result (PC)		PCR*42	1-2 weeks
Macrolide Resistance Test (Mgen)	MGR	FCRU / PCR	1-2 weeks
MPOX (Virus) – Lesion (Self-collect)	MPXV	Aptima multisite swab	2 days
Mycoplasma genitalium by PCR	MGEN	FCRU / PCR / TPV	2 days

TEST	CODE	SAMPLE REQS	TAT
Mycoplasma genitalium by PCR – Urine and Vaginal (Self-collect)	MGEN	Aptima urine or multisite swab	2 days
Mycoplasma genitalium Resistance – Urine or Vaginal (Self-collect)	MGR	Aptima urine or multisite swab	1-2 weeks
Mycoplasma genitalium/Ureaplasma by PCR	MUPC	FCRU / PCR / TPV	2 days
Rapid Xpert HIV-1 RNA Qualitative – Early Detection from 10 days	LHIV	(Vacutainer only)	1 day
Rapid Xpert HIV-1 RNS Viral Load – Rapid Testing for HIV-Positive Patient Prognosis and Response To Antiretroviral Therapy	RHIV	(Vacutainer only)	1 day
RPR (Syphilis)	RPR	B	2 days
STD1 M/F STD Quad (Urine and Serology)	STD1	□ FCRU	2 days
STD2 M/F STI Profile Plus (Urine and Serology)	STD2	E FCRU (If culture swabs are needed please request separately)	4 days
STD3 Female STD Quad (PCR Swab and Serology)	STD3	□ PCR	2 days
STD4 Female STI Profile Plus (PCR Swab and Serology)	STD4	<b>E) PCR</b> (If culture swabs are needed please request separately)	4 days
STD5 Serology only	STD5	B	1 day
STD6 Serology only without HIV	STD6	В	1 day
STD8 Vaginitis/BV Profile using Culture & PCR Swab	STD8	PCR and STM	3 days
STD9 Symptomatic lesion sample using PCR Swab from lesion	STD9	PCR Swab	7 days
STI Profile by PCR (7 tests from 1 sample) (Self-collect)	DL12	Aptima urine or multisite swab	2 days
STI Profile: MSM1 (Blood + Urine/ Throat/Rectal Swabs) (Self-collect)	MSM1	(TDL Tiny) / Aptima Urine / Aptima multisite swab x 2	2 days
STI Profile: MSM1 (Venous)	MSM1	3 / FCRU / PCR Swab Throat / PCR Swab Rectal	2 days
STI Profile: MSM2 (Blood + Urine/ Throat/Rectal Swabs) (Self-collect)	MSM2	(TDL Tiny) / Aptima urine / Aptima multisite swab x 2	3 days
STI Profile: MSM2 (Venous)	MSM2	3 / FCRU / PCR Swab Throat / PCR Swab Rectal	3 days
Syphilis by PCR (chancre)	SYPS	PCR	5 days
Syphilis IgG/IgM (Self-collect)	TSYP	(TDL Tiny)	1 day

TEST	CODE	SAMPLE REQS	TAT
Syphilis IgG/IgM (Venous)	SERJ	B	1 day
ТРНА	TPPA	B	2 days
Trichomonas vaginalis (PCR)	TVPC	FCRU / PCR / TPV	2 days
Trichomonas vaginalis (TV) – Urine or Vaginal (Self-collect)	TVPC	Aptima urine or multisite swab	2 days
Triple Swab Female STI Profile (Vaginal/ Throat/Rectal Swabs) (PCR)	3SWA	PCR swab x 3 (label by site)	2 days
Triple Swab Female STI Profile (Vaginal/ Throat/Rectal Swabs) (Self-collect)	3SWA	Aptima multisite swab x 3 (label by site)	2 days
Ureaplasma urealyticum by PCR	UGEN	FCRU / PCR / TPV	2 days
Vaginitis/BV Profile (Culture & PCR)	STD8	PCR and STM	3 days
Vaginitis/BV Profile using Culture & PCR Swab (Self-collect)	STD8	Aptima multisite swab and Blue gel Amies swab	3-5 days

## STD1 M/F STD Quad (Urine and Serology)

#### **SEROLOGY**

HIV 1&2/p24 Antigen Syphilis IgG/IgM

#### URINE

Chlamydia Gonorrhoea

#### TAT: 2 days

STD1

**□** FCRU

# STD4 Female STI Profile Plus (PCR Swab and Serology)

#### SEROLOGY

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

#### **VAGINAL PCR SWAB**

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

#### TAT: 4 days

STD4

PCR (If culture swabs are needed please request separately)

## STD2 M/F STI Profile Plus (Urine and Serology)

#### **SEROLOGY**

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

#### URINE

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

#### TAT: 4 days

STD2

**E FCRU** (If culture swabs are needed please request separately)

### **STD5 Serology only**

HIV 1&2/p24 Antigen Hepatitis B Surface Antigen Hepatitis C Antibodies Hepatitis C Antigen (Early detection) Syphilis IgG/IgM

#### TAT: 1 day

STD5



## STD3 Female STD Quad (PCR Swab and Serology)

#### **SEROLOGY**

HIV 1&2/p24 Antigen Syphilis IqG/IqM

#### **VAGINAL PCR SWAB**

Chlamydia Gonorrhoea

TAT: 2 days

STD3

PCR

## STD6 Serology only without HIV

Hepatitis B Surface Antigen Hepatitis C Antibodies Hepatitis C Antigen (Early detection) Syphilis IgG/IgM

#### TAT: 1 day

STD6



### STD8 Vaginitis/BV Profile using Culture & PCR Swab

Candida (Culture)
Gardnerella vaginalis by PCR
Mycoplasma genitalium
Trichomonas vaginalis by PCR
Ureaplasma urealyticum

#### TAT: 3 days

STD8

PCR and STM

### STD9 Symptomatic lesion sample using PCR Swab from lesion

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

TAT: 7 days

STD9

**PCR Swab** 

## 7 STI Profile by PCR (7 tests from 1 Sample)

Chlamydia trachomatis Neisseria gonorrhoea Mycoplasma genitalium Ureaplasma species Trichomonas vaginalis Gardnerella vaginalis Herpes Simplex I/II

All tests can be requested individually

TAT: 2 days

DL12

FCRU / PCR / TPV

### CT/GC/Trichomonas/ Mgen – PCR Swab

Chlamydia Gonorrhoea Trichomonas vaginalis Mycoplasma genitalium

All tests can be requested individually

TAT: 2 days

**SGTM** 

**PCR Swab** 

### CT/GC/Trichomonas/ Mgen – Thin Prep

Chlamydia Gonorrhoea Trichomonas vaginalis Mycoplasma genitalium

All tests can be requested individually

TAT: 2 days

**TGTM** 

TPV

### CT/GC/Trichomonas/ Mgen – Urine

Chlamydia Gonorrhoea Trichomonas vaginalis Mycoplasma genitalium

All tests can be requested individually

TAT: 2 days

**CGTM** 

**FCRU** 

## HIV/HBV/HCV Screen by PCR/NAAT (10 days post exposure)

Positive findings will be reflexed for confirmatory testing

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

STDX provides diagnostic confirmatory testing only when used in addition to serology for Ag/Ab HIV-1&2, HBV, HCV

TAT: 3 days

STDX

(Vacutainer only)

## HIV/HBV/HCV (Early detection by PCR/ NAAT) with Syphilis

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

STXX provides diagnostic confirmatory testing only when used in addition to serology for Ag/Ab HIV-1&2, HBV, HCV

TAT: 3 days

STXX

## HIV Rapid RNA HIV-1 QUALITATIVE

Early detection from 10 days HIV-1 RNA

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

#### TAT: 1 day

LHIV

(Vacutainer only)

## HIV Rapid RNA HIV-1 QUANTITATIVE

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: 1 day

RHIV

(Vacutainer only)

## STI Profile: MSM1 (Venous)

HIV 1&2/p24 Ag Syphilis IgG/IgM Chlamydia/Gonorrhoea – Urine Chlamydia/Gonorrhoea – Throat Chlamydia/Gonorrhoea – Rectal

#### TAT: 2 days

MSM<sub>1</sub>

(3) FCRU / PCR Swab
Throat / PCR Swab Rectal

## STI Profile: MSM2 (Venous)

HIV 1&2/p24 Ag
Syphilis IgG/IgM
Hepatitis B Surface Antigen
Hepatitis C Antibodies
7 STI by PCR Screen
Chlamydia/Gonorrhoea – Throat
Chlamydia/Gonorrhoea – Rectal
Macrolide Resistance Test (M.gen)

#### TAT: 3 days

MSM2

B / FCRU / PCR Swab Throat / PCR Swab Rectal

## Triple Swab Female STI Profile (Vaginal/Throat/ Rectal Swabs) (PCR)

Chlamydia/Gonorrhoea – Vaginal Chlamydia/Gonorrhoea – Throat Chlamydia/Gonorrhoea – Rectal

#### TAT: 2 days

3SWA

PCR Swab x 3 (label by site)

STI	INCUBATION PERIOD	SAMPLE SITE	TEST	TEST CODE	SAMPLE TYPE	TAT
<b>Chlamydia CT</b> (Bacterial)	1–3 weeks, up to 6 weeks	Urine Cervix/Vagina Cervix/Vagina	Chlamydia Chlamydia Chlamydia	CPCR SPCR TPCR	First catch Urine PCR Swab Thin Prep Vial	2 days 2 days 2 days
Gonorrhoea GC (Bacterial)	2-7 days, up to 1 month	Urine Cervix/Vagina Cervix/Vagina Cervix Rectal Throat Urethral Other site	Gonorrhoea by PCR Gonorrhoea by PCR Gonorrhoea by PCR Gonorrhoea by Culture Gonorrhoea by Culture Gonorrhoea by Culture Gonorrhoea by Culture	CGON SGON TGON GONC GONT GONU	First Catch Urine PCR Swab Thin Prep Vial Black Charcoal swab	2 days 2 days 2 days 3-5 days 3-5 days 3-5 days 3-5 days
CT/GC Combined (Bacterial)	up to 6 weeks,	Urine Cervix/Vagina Cervix/Vagina Rectum Throat	CT/GC CT/GC CT/GC CT/GC CT/GC	SCG SCG TCG RSCG TSCG	First Catch Urine PCR Swab Thin Prep Vial PCR Swab PCR Swab	2 days 2 days 2 days 2 days 2 days
Mycoplasma genitalium (Bacterial)	Symptoms develop at 1-3 weeks	Urine GU Site Cervix/Vagina	Mycoplasma genitalium by PCR Mycoplasma genitalium by PCR Mycoplasma genitalium by PCR	MGEN MGEN MGEN	First Catch Urine PCR Swab Thin Prep Vial	2 days 2 days 2 days
Ureaplasma urealyticum (Bacterial)	Symptoms develop at 1-3 weeks	Urine GU Site Cervix/Vagina	Ureaplasma by PCR Ureaplasma by PCR Ureaplasma by PCR	UGEN UGEN UGEN	First Catch Urine PCR Swab Thin Prep Vial	2 days 2 days 2 days
Trichomonas vaginalis (Parasitic)	4–28 days, many patients are asymptomatic carriers	Urine GU Site Cervix/Vagina	Trichomonas vaginalis by PCR Trichomonas vaginalis by PCR Trichomonas vaginalis by PCR	TVPC TVPC TVPC	First Catch Urine PCR Swab Thin Prep Vial	2 days 2 days 2 days
<b>Gardnerella vaginalis</b> (Bacterial)	Imbalance of normal flora	Urine GU Site Cervix/Vagina	Gardnerella vaginalis by PCR Gardnerella vaginalis by PCR Gardnerella vaginalis by PCR	GVPC GVPC GVPC	First Catch Urine PCR Swab Thin Prep Vial	2 days 2 days 2 days

STI	INCUBATION PERIOD	SAMPLE SITE	TEST	TEST CODE	SAMPLE TYPE	TAT
Bacterial Vaginosis (BV) (Bacterial)	Imbalance of normal flora	Cervix/Vagina	Bacterial Vaginosis (BV) Profile by both MICROSCOPY and PCR	STD8	Both Microscopy & PCR swab	3 days
Herpes Simplex Viral I/II (Viral)	2–14 days, testing is most appropriate for patients with symptomatic lesion(s)	Herpes lesion	Herpes by PCR Herpes by PCR	HERS HERD	PCR Swab First Catch Urine	5 days 5 days
Human Papillomavirus (Viral)	HPV is the most common sexually transmitted infection – usually asymptomatic	Cervical cells Cells/ papilloma from site (anal)	HPV (28 individually typed LR & HR DNA subtypes)	HP20	Thin Prep Vial Brush sampler	3 days
Genital warts (Viral)	Weeks/months after exposure	GU Warts	HPV (28 individually typed LR & HR DNA subtypes)	HP20	Thin Prep Vial Brush sampler Cells/Papilloma	3 days
Syphilis/Herpes (Bacterial/Viral)	Whenever active lesions are present	Symptomatic lesion	Syphilis/Herpes Lesion Profile	STD9	PCR Swab	7 days
Syphilis (Bacterial)	9–21 days, but up to 90 days	Blood	Syphilis IgG/IgM	SERJ	8	1 day
Herpes Simplex Virus I/II (Viral)	lgG 4—6 weeks after exposure, IgM 5—35 days after exposure, after which test IgG	Blood	Herpes IgG (past infection Herpes IgM (current/recent)	HERP HERM	<b>0 0</b>	2 days 2 days
HIV (Viral)	Usually 10 – 90 days, but up to 180 days	Blood	HIV I&II / p24 antigen (screening from 45 days post exposure (BHIVA))	нрио	<b>©</b>	1 day

STI	INCUBATION SAMPLE PERIOD SITE	PLE	TEST	TEST CODE	SAMPLE	TAT
Hepatitis B (Viral)	Usually 45–180 Blood days, average of 60–90 days	5	Hepatitis B surface antigen	AUAG	<b>a</b>	1 day
Hepatitis C Ab (Viral)	Usually 9–180 Blood days, average of 45–65 days	q	Hepatitis C Antibodies	НЕРС	<b>@</b>	1 day
EARLY DETECTION PROFILES BY PCR	INCUBATION PERIOD	SAMPLE SITE	TEST	TEST CODE	SAMPLE TYPE	TAT
7 STIs by PCR	One sample for 7 STI Tests	Urine Cervix Vagina	Chlamydia Gonorrhoea Mycoplasma genitalium Ureaplasma genitalium Trichomonas vaginalis Gardnerella vaginalis Herpes Simplex I/II	PP12	Thin Prep Vial or First Catch Urine or PCR Swab or Aptima urine or multisite swab	2 days
ни/нви/нси	Early Detection Screen by PCR Multiplex	Blood	HIV 1&2 RNA Hepatitis B (HBV DNA) Hepatitis C (HCV RNA)	STDX	2 x 6mls or 2x4mls 3 days (Vacutainer only)	3 days

#### **TDL** website

The TDL website gives updated details of our tests — sample types, turnaround times and special instructions. The 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 can be requested by emailing **refranges@tdlpathology.com**. Full details of our tests and profiles are also available in the TDL TestGuide app.



Visit the TDL website at:

www.tdlpathology.com

SCAN ME



TEST	CODE	SAMPLE REQS	TAT
Acute Viral Hepatitis Screen	AHSC	B	1 day
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	CCP	B	2 days
Anti-Liver Cytosol Antibodies	ALCA	B	5 days
Anti-MOG [Myelin Oligodendrocyte Glycoprotein] Antibodies	AMOG	В	3 weeks
Anti-MUSK Antibodies	MUSK	B	2 weeks
Anti-Nuclear Antibodies (titre & pattern)	ANAB	B	2 days
Anti-Phosphatidylserine Antibodies	PHTS	B	5 days
Anti-Phospholipase A2 Receptor	AA2R	B	6 weeks
Anti-SLA (Soluble Liver Antigen) Abs	LSA	B	5 days
Anti-Staphylolysin Titre (SGOT)	ASTT	B	3 days
Anti-Streptolysin Titre/ASOT	ASLT	B	2 days
Anti-Sulfatide Antibodies	ASA	B	5 weeks
Aquaporin 4 Antibodies (Neuromyelitis Optica)	AQUA	B	2 weeks
Ascariasis Serology	ASC	B	5 days
Aspergillus Precipitins	ASPP	B	5 days
Autoantibody Profile I	AUT0	B	2 days
Autoantibody Profile II	ENDO	B	3 days
Avian Precipitins (11 Species)	AVIA	B	5 days
Babesia PCR	PCRB	A	7 days
Beta 2 Glycoprotein 1 Abs	B2GP	B	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
Brucella Serology	BRUC	<b>B</b> 9	2-3 weeks
C1 Esterase Inhibitor	C1EI	В	5 days
C3 Complement	C3	B	1 day
C3/C4 Complement	COMP	В	1 day
C4 Complement	C4	В	1 day
Campylobacter Jejuni Antibodies	CJAB	В	5 days
Candida Antibodies	CANA	В	5 days
Cardiolipin Antibodies (IgG+IgM)	ACAB	B	2 days

TEST	CODE	SAMPLE REQS	TAT
CCP Antibodies (RF)	CCP	B	2 days
CH50 (Classical pathway)	CH50	(Frozen)⁴	4 days
Chagas Disease Serology (S.American Trypanosomiasis) T. Cruzi	CHGA	<b>B</b> 9,14	10 days
Chlamydia Species Specific (MIF) Ab Screen	CHAB	В	3 days
Chronic Fatigue Syndrome Profile	VIP1	A + B 10	5 days
Coeliac Disease – HLA DQ2/DQ8 Genotype	Q2Q8	<b>A</b> 9	10 days
Coeliac/Gluten Genetic Profile 2 See page 91	GSA2	<b>A</b> B	10 days
Coeliac/Gluten Sensitivity Profile See page 91	GSA	<b>B</b>	2 days
Colloid Antigen-2 Antibodies	CA2A	B	2 weeks
Cotinine (Serum)	COT	B	4 days
COVID-19 (SPIKE) Antibodies (Self-collect)	SCOV	(TDL Tiny)	1 day
COVID-19 (SPIKE) Antibodies (Venous)	SCOV	SST/Serum (B) (Venous)	1 day
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>B</b> 9,14	5 days
Ehrlichiosis Antibodies	EHRL	<b>B</b> 9,14	10 days
Extractable Nuclear Antibodies (nRNP, Sm, Ro, La, Jo1, ScI70) CENP-B	ENA	В	2 days
Farmers Lung Precipitins	FARM	B	5 days
Fasciola Hepatica Antibodies (Liver Fluke)	FASC	B	2 weeks
Ganglionic Acetylcholine Receptor Antibodies	GACA	B	9 weeks
Ganglioside GM1, GD1B, GQ1B Abs	GANG	B	5 days
Gastric Parietal Autoantibodies	GASP	B	2 days
Gliadin Antibodies (IgG) (deamidated) (Self-collect)	AGAB	(TDL Tiny)	2 days
Gliadin Antibodies (IgG) (deamidated) (Venous)	AGAB	B	2 days
Glomerular Basement Membrane Abs	AGBM	B	2 days
Glutamic Acid Decarboxylase Antibodies (GAD 65)	GAD	В	5 days
Gluten Sensitivity Evaluation	GSA	B	2 days
Gluten Sensitivity Profile	GLUT	ABB	10 days
Gluten/Coeliac Genetic Profile 2	GSA2	AB	10 days
Granulocyte Immunology	GRIM	(or 2 x 6ml)	2 weeks

TEST	CODE	SAMPLE REQS	TAT
H. pylori Antibodies (IgG)	НВРА	B	2 days
Haemophilus B Influenzae Antibodies	HINF	B	5 days
Histamine (Blood)	HITT	(Frozen plasma)	5 days
Histamine (Urine)	HITU	RU	5 days
Histamine Releasing Urticaria Test	CURT	B	3 weeks
Histone Antibodies	HISA	B	5 days
Histoplasmosis	HISP	B	10 days
HLA B27	HLAB	<b>A</b> 9	3 days
IgE (Total)	IGE	B	1 day
Immune-Complexes	IMCP	B	5 days
Immunofluorescence in Skin Biopsies	IHCS	Skin sample in Michels solution	2 weeks
Immunoglobulins (IgG, IgM, IgA)	IMM	B	1 day
Insulin Antibodies	INAB	B	5 days
Interleukin 1 Beta	ILB	(Frozen) <sup>4,7</sup>	1-2 weeks
Interleukin 2	IL2	(Frozen) <sup>4,7</sup>	1-2 weeks
Interleukin 4	IL4A	(Frozen) <sup>4,7</sup>	1-2 weeks
Interleukin 6	IL6	(Frozen) <sup>4,7</sup>	1-2 weeks
Interleukin 8	IL8	(Frozen) <sup>4,7</sup>	1-2 weeks
Interleukin 10	IL10	(Frozen) <sup>4,7</sup>	1-2 weeks
Interleukin 28b Genotype	IL28	A	2 weeks
Intrinsic Factor Antibodies	IFAB	В	2 days
Islet Cell Antibodies	ICAB	В	3 days
Legionella Antibodies	LEG0	B	3 days
Legionella Urine Antigen	LEGA	Urine with boric acid	1 day
Leptospirosis (Weil's Disease) Abs (IgM)	LEP	В	5 days
Leukotriene E4	LTE4	CU (Frozen)	3 weeks
Liver Immunoblot	LIVI	В	3 days
Liver Kidney Microsomal Antibodies	LKM	В	2 days
Lupus Anticoagulant and Anticardiolipin Abs	LUPA	B C C 4,18	2 days
Lyme Disease (Borrelia Abs) IgG, IgM	BORR	B 9,14	2 days
Lyme Disease (Borrelia Abs) IgM	BORM	В	2 days
Meningococcal Serology (only serogroup C)	MENI	В	6 weeks
Mitochondrial Antibodies	AMIT	В	3 days
Mitochondrial Antibodies M2	MTM2	В	2 days
Myasthenia Gravis Evaluation	MGE	В	5 days

TEST	CODE	SAMPLE REQS	TAT
Myelin Associated Glycoprotein Antibodies	MAG	B	5 days
Myelin Basic Protein Antibodies	MBPA	B	2 weeks
Myeloperoxidase Antibodies	MP0	B	2 days
Myocardial Antibodies	MYO	B	1 week
Myositis Panel	MYOS	B	3 days
Neuronal Antibody (Hu, Ri, Yo, Cv2, Ma2)	NEUR	B	10 days
NMDA Receptor Antibodies	NMDA	В	3 weeks
Nucleic Acid Antigen Antibodies	DNA	В	2 days
Oligoclonal Bands	CSF0	2ml CSF + 🕃	5 days
Ovarian Autoantibodies	OVAB	В	3 days
Paragomius Serology	PRGM	B	2 weeks
Parathyroid Antibodies	PTHA	В	3 weeks
Pemphigus/Pemphigoid Autoantibodies	SKAB	В	2 days
Pertussis (Whooping Cough) Antibodies	PERS	В	5 days
Pertussis (Whooping Cough) by PCR	PERP	Pernasal or dry swab	2-3 days
Pituitary Antibodies	PITU	<b>B</b> 4	1 month
Pneumococcal Antibodies – Serotype Specific	PASS	<b>B</b>	5 weeks
Pneumococcal Antibody Screen	PNEU	<b>3</b>	5 days
Proteinase 3 Ab	PR3	B	2 days
Purkinje Cell Antibody (Hu and Yo)	PURK	В	10 days
Q Fever (C Burnetti) Antibodies	QFEV	<b>B</b> 9	10 days
Rheumatoid Factor (Latex Test)	RF	В	3 days
Rheumatology Profile 1 (Screen)	RH	<b>A</b> B	2 days
Rheumatology Profile 2 (Connective tissue)	RH2	AABB	3 days
Rheumatology Profile 3 (Rheumatoid/Basic)	RH3	<b>A</b> B	2 days
Rheumatology Profile 4 (Systemic Lupus)	RH4	<b>A</b> BB	2 days
Rheumatology Profile 5 (Mono Arthritis)	RH5	<b>AABB</b>	3 days
Rheumatology Profile 6 (Rheumatoid Plus)	RH6	B	3 days
Rheumatology Profile 7 (Sjogren's Syndrome)	RH7	B	15 days
Rickettsial Species Antibody Profile	RICK	B	7 days
RNA Polymerase Antibodies	RNAP	<b>B</b>	3 days
RPR (Syphilis)	RPR	<b>B</b>	2 days
Saccharomyces Cerevisiae Antibodies	ASCA	<b>3</b>	2 weeks
Salivary Duct Antibodies	SAB	<b>B</b>	15 days

TEST	CODE	SAMPLE REQS	TAT
Scleroderma Immunoblot	SCLI	B	3 days
Sjogren's Syndrome	RH7	B	15 days
Skin (Pemphigus/Pemphigoid) Autoantibodies	SKAB	B	2 days
Skin Antibodies by Immunofluorescence	STSK	B	1 month
Sleeping Sickness Serology (African Trypanosomiasis)	TRYP	<b>B</b> 9	10 days
Smooth Muscle Antibodies	ASM0	B	2 days
Sperm Antibodies (Serum)	ASAB	B	5 days
Steroid Cell Antibody	SCA	B	2 days
Striated/Skeletal Muscle Antibody	STRA	В	3 days
Strongyloides Antibodies	STGA	B	10 days
Syphilis IgG/IgM (Self-collect)	TSYP	(TDL Tiny)	1 day
Syphilis IgG/IgM (Venous)	SERJ	B	1 day
TB Quantiferon®-TB Gold*  * 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.	TBQ4	Special tubes or 1	3 days
Tetanus Antibody	TETA	B	5 days
Thyroid Abs (Thyroglobulin + Thyroid Peroxidase Abs) (Self-collect)	THAB	(TDL Tiny)	2 days
Thyroid Abs (Thyroglobulin + Thyroid Peroxidase Abs) (Venous)	THAB	В	1 day
Thyroid Peroxidase Antibodies/Anti TP0	TPEX	B	1 day
Tissue Transglutaminase IgA (Coeliac) (Self-collect)	TAA	(TDL Tiny)	2 days
Tissue Transglutaminase IgA (Coeliac) (Venous)**	TAA	В	2 days
Tissue Transglutaminase IgG	TAAG	B	5 days
Total Immune Function Evaluation	TIE	A + B 5,10	7 days
Total Immunoglobulin E	IGE	B	1 day
Toxocara Antibodies (IgG)	TFAT	<b>B</b> 9	5 days
Toxoplasma Antibodies (IgG, IgM)	TFAM	<b>B</b> 9	1 day
Toxoplasma Antibody Full Evaluation (IgM, Dye Test, IgG Avidity)	TDYE	<b>B</b> 9	10 days
Toxoplasma by PCR	TXAG	A	5 days
ТРНА	TPPA	В	2 days
Trichinella Serology	TRIC	В	5 days

TEST	CODE	SAMPLE REQS	TAT
Trypanosome (Chagas) Antibodies	CHGA	<b>B</b> 9,14	10 days
TSH-Receptor Antibodies	TSI	B	4 days
Tularaemia Antibodies	TULA	<b>B</b> 14	5 days
Urinary Methyl Histamine	UHIT	RU (Frozen)	2 weeks
Urticaria Test (Histamine Releasing)	CURT	B	3 weeks
Vascular Endothelial Growth Factor	VEGF	B	14 days
Voltage Gated Calcium Channel Antibodies	CCAB	B	3 weeks
Voltage Gated Potassium Channel Antibodies	VPCA	B	3 weeks
Whooping Cough (Pertussis) Antibodies	PERS	B	5 days
Whooping Cough (Pertussis) by PCR	PERP	Pernasal or dry swab	2-3 days
Yellow Fever Antibodies	YELL	<b>B</b> 9,14	10 days
Yersinia Antibodies	YERS	B	4 days
Zika Abs IgM and IgG – Antibody detection from 15 days	ZKAB	B	Up to 14 days
Zika RNA by PCR in Semen	ZIKS	Semen	Up to 14 days
Zika RT PCR – Window of detection from 1-7 days from onset of symptoms	ZIKA	В	Up to 14 days
Zika RT PCR – Window of detection from 1-14 days from onset of symptoms	ZIKU	RU	Up to 14 days

### Acute Viral Hepatitis Screen

Hepatitis A (IgM) Hepatitis B Surface Antigen Hepatitis C Antibodies

TAT: 1 day

AHSC

₿

#### **Autoantibody Profile I**

Thyroid Abs (Thyroglobulin +
Thyroid Peroxidase Abs)
Anti-Nuclear Antibodies
Mitochondrial Antibodies
Smooth Muscle Antibodies
Gastric Parietal Autoantibodies
Liver Kidney Microsomal Antibodies

TAT: 2 days

AUTO

₿

## **Autoantibody Profile II**

Thyroid Abs (Thyroglobulin + Thyroid Peroxidase Abs) Islet Cell Antibodies Adrenal Cortex Antibodies Gastric Parietal Autoantibodies Gonadal (Ovarian) Autoantibodies

TAT: 3 days

**ENDO** 

₿

### Chlamydia Species Specific (MIF) Ab Screen

Chlamydia trachomatis (serovar A-K & L1-L3) Chlamydia pneumoniae Chlamydia psittaci

TAT: 3 days

CHAB



## Chronic Fatigue Syndrome Profile

Epstein-Barr Virus Antibodies IgG/IgM Lymphocyte Subsets (CD3/CD4/CD8) C Reactive Protein (CRP) Vitamin D (25-OH)

TAT: 5 days

VIP1



## Coeliac/Gluten Genetic Profile 2

Gliadin Antibodies (IqG)

(deamidated)
HLA Tissue Typing Coeliac
Disease – DQ2/DQ8
Immunoglobulin A
Tissue Transglutaminase
IgA (Coeliac)

TAT: 10 days

GSA2



### Coeliac/Gluten Sensitivity Profile

Gliadin Antibodies (IgG) (deamidated) Immunoglobulin A Tissue Transglutaminase (IgA)

TAT: 2 days

GSA



#### **Gluten Sensitivity Profile**

Gluten Single IgE Allergen Gliadin Antibodies (IgG) (deamidated) HLA Tissue Typing Coeliac Disease – DQ2/DQ8 Immunoglobulin A Tissue Transglutaminase IgA (Coeliac)

TAT: 10 days

GLUT



### **Rheumatology Profile 1** (Screen)

Full Blood Count (FBC) **ESR** Uric Acid (Serum) Rheumatoid Factor (Latex Test) Anti-CCP Antibodies

C Reactive Protein (CRP)

TAT: 2 days

RH



### **Rheumatology Profile 2** (Connective tissue)

Full Blood Count (FBC) **FSR** Uric Acid (Serum) Anti-Nuclear Antibodies DNA (Double Stranded) Antibodies IaG

Antibodies to Extractable Nuclear Antigens (ENA):

Anti-Sm Anti-Ro (SS-A) Anti-La (SS-B) Anti-Jo-1

Anti-nRNP

Anti-Scl 70 Anti-CFNP

Rheumatoid Factor (Latex Test)

Anti-CCP Antibodies HI A B27

C Reactive Protein (CRP)

TAT: 3 days

CENP-B

RH2



### **Rheumatology Profile 3** (Rheumatoid/Basic)

Full Blood Count (FBC) **FSR** Uric Acid (Serum) Rheumatoid Factor (Latex Test) Anti-CCP Antibodies Anti-Nuclear Antibodies C Reactive Protein (CRP)

TAT: 2 days

RH3



### **Rheumatology Profile 4** (Systemic Lupus)

Full Blood Count (FBC) Anti-Nuclear Antibodies DNA (Double Stranded) Antibodies IaG Antibodies to Extractable Nuclear Antigens (ENA): Anti-nRNP Anti-Sm Anti-Ro (SS-A)

Anti-La (SS-B) Anti-Jo-1 Anti-Scl 70 Anti-CFNP

Rheumatoid Factor (Latex Test) Anti-CCP Antibodies Anti-Cardiolipin Autoantibodies Complement 3/4

C Reactive Protein (CRP)

TAT: 2 days

RH4



## **Rheumatology Profile 5** (Mono Arthritis)

Full Blood Count (FBC) **FSR** Uric Acid (Serum) Rheumatoid Factor (Latex Test) Anti-CCP Antibodies Anti-Nuclear Antibodies C Reactive Protein (CRP) HLA B27

TAT: 3 days

RH5





## **Rheumatology Profile 6** (Rheumatoid Plus)

Rheumatoid Factor (Latex Test) Anti-CCP Antibodies C Reactive Protein (CRP)

TAT: 3 days

RH6



### **Rheumatology Profile 7** (Sjogren's Syndrome)

Anti-Ro (SS-A) Anti-La (SS-B) Salivary Antibodies (SAB) C Reactive Protein (CRP)

TAT: 15 days

RH7



## **Coeliac Disease (CD)**

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%.

## **Coeliac pathway**

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 this by adding a total IgA to all low Tissue Transglutaminase (TGG) 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:

- Initial TTG IgA samples are received and tested
- If TTG IgA is LOW <0.2 U/ml reflex testing for Total IgA will be undertaken
- If Total IgA is LOW <0.1 g/L then reflex testing for Gliadin IgG test will be undertaken
- If TTG IgA is HIGH >/= 10 U/ml then reflex testing for Endomysial IgA will be undertaken as a confirmatory test for first time positive samples.

#### **Endomysial IgA**

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.

#### 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 – see table.

TTG IgA result U/ml	Total IgA result for new assay g/L	Deamidated gliadin IgG result U/ml	Comment
0.2 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
<0.2	>/= 0.1	N/A	Coeliac disease unlikely (please note that if the patient has no dietary gluten, results may appear false negative)
<0.2	<0.1	>/=10	Consistent with coeliac disease in a patient with selective IgA deficiency
<0.2	<0.1	<7	Coeliac disease unlikely (please note that if the patient has no dietary gluten, results may appear false negative)
<0.2	<0.1	7-10	Result equivocal suggest referral to a gastroenterologist for consideration of duodenal biopsy

## **Tropical and Travel-Related Immunology**

TEST	CODE	SAMPLE REQS	TAT
Amoebic (E. histolytica) Antibodies	AFAT	В	1 week
Amoebic (E. histolytica) PCR	AMAG	RF	2 days
Bancroftia/Oncerciasis/Filarial Antibodies	TFIF	B 14	2 weeks
Bilharzia (Schistosome) Antibody Screen	BILH	B 14	10 days
Bilharzia (Urine)	USCH	Mid-morning terminal urine following exercise 14	1-2 days
Borrelia Antibodies (Lyme Disease) IgG, IgM	BORR	<b>B</b> 9,14	2 days
Borrelia Antibodies (Lyme Disease) IgM	BORM	В	2 days
Borrelia Confirmation (Immunoblot)	BORC	<b>B</b> 9,14	10 days
Cryptosporidium Detection by PCR	CRPA	RF	2 days
Dengue Virus Serology	DENG	<b>B</b> 9,14	5 days
DVT/Pre-travel Screen	DVT1	<b>A A B</b> <sup>9</sup>	5 days
Echinococcus (Hydatid) Antibodies	EFAT	B 9,14	5 days
Enteric Organism Rapid Detection (RF)	EORD	RF	2 days
Enteric Organism Rapid Detection (Self-collect)	EORD	Stool/faecal container	2 days
Filaria (Lymphatic and Non- Lymphatic) Antibodies	FIFA	<b>B</b> 9,14	10 days
Gastrointestinal Pathogen Profile by PCR (Self-collect)	EORD	Stool/faecal container	2 days
Insect/Worm/Ova/Cysts	FLEA	Send Specimen <sup>9,14</sup>	5 days
Leishmania Antibodies	LEIS	B	5 days
Malarial Antibodies (Pl. falciparum)	MALA	B 9,14	5 days
Malarial Antibodies (species specific)	MALS	B 9,14	10 days
Post-Travel Screen 1 (Up to 6 weeks post travel)	PTS	<b>A B C</b> 14	10 days
Post-Travel Screen 2 (6 weeks after travel)	PTS2	<b>A B B B G</b> 14	10 days
Pre-Travel Screen (DVT)	DVT1	<b>A A B</b> 9	5 days
Rickettsial Species Antibody Profile	RICK	B	7 days
Schistosome (Bilharzia) Antibodies	BILH	B 14	10 days
Toxoplasma Antibodies (IgG, IgM)	TFAM	<b>B</b> 9	1 day
Tropical Screen (from 6 weeks post-travel)	TR0P	<b>B B</b> 9,14	10 days
Zika Abs IgM and IgG – Antibody detection from 15 days	ZKAB	В	Up to 14 days
Zika RNA by PCR in Semen	ZIKS	Semen	Up to 14 days
Zika RT PCR – Window of detection from 1-7 days from onset of symptoms	ZIKA	В	Up to 14 days
Zika RT PCR – Window of detection from 1-14 days from onset of symptoms	ZIKU	RU	Up to 14 days

#### **Tropical and Travel-Related Immunology**

### Post-Travel Screen 1 (Up to 6 weeks post travel)

Haematology Profile Biochemistry Profile Schistosome Abs Malarial Abs

#### TAT: 10 days

PTS



## Post-Travel Screen 2 (6 weeks after travel)

Haematology Profile Biochemistry Profile Schistosome Abs Malarial Abs Hepatitis A IgM Abs Hepatitis B Surface Antigen Hepatitis C Antibodies HIV Duo

#### TAT: 10 days

PTS2



#### **DVT/Pre-travel Screen**

Full Blood Count (FBC)
Factor II Prothrombin –
G20210A Variant
Factor V Leiden – G1691A Variant
Cardiolipin Antibodies (IgG+IgM)

#### TAT: 5 days

DVT1



### Tropical Screen (from 6 weeks post-travel)

Amoebic (E. histolytica) Antibodies Schistosome (Bilharzia) Antibodies Echinococcus (Hydatid) Antibodies Leishmania Antibodies Malarial Parasites Toxoplasma Antibodies (IgG, IgM)

#### TAT: 10 days

TR<sub>0</sub>P



## Enteric Organism Rapid Detection (RF)

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

#### **Bacteria and Bacterial Toxins**

C. difficile Toxin A/B gene,
Campylobacter spp.,
Enteroaggregative E.coli (EAEC),
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) 0157: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 cayetanensis, 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

## **Immune status**

TEST	CODE	SAMPLE REQS	TAT
Hepatitis A Immunity (IgG/IgM)	HAIM	B	1 day
Hepatitis B Immunity (IgG) (Self-collect)	THBI	(TDL Tiny)	1 day
Hepatitis B Immunity (IgG) (Venous)	HBIM	<b>B</b>	1 day
Measles Antibodies (IgG) Immunity	MEAS	B	1 day
Measles Antibodies (IgM)	MEAM	<b>B</b> 9	2 days
Measles, Mumps, Rubella (MMR)	MMR	<b>B</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	5 days
Rabies Antibody	RABI	B	20 days
Rubella Antibody (IgG)	RUBE	B	1 day
Rubella Antibody (IgM)	RUBM	B	1 day
Rubella PCR	RUBP	🔼 / Amniotic Fluid	5 days
Tetanus Antibody	TETA	B	5 days
Varicella zoster Antibodies (IgG)	VZ0S	<b>B</b>	1 day
Varicella zoster Antibodies (IgM)	VZOM	<b>B</b>	1 day

## **Hepatitis testing**

TEST	CODE	SAMPLE REQS	TAT
Hepatitis (Acute) Screen	AHSC	В	1 day
Hepatitis A (IgM)	HAVM	<b>B</b>	1 day
Hepatitis A Immunity (IgG/IgM)	HAIM	<b>B</b>	1 day
Hepatitis A Profile	HEPA	<b>3</b>	1 day
Hepatitis A RNA by PCR	HAVR	(A) or (B)	3 weeks
Hepatitis A, B & C Profile	ABC	88	1 day
Hepatitis B (PCR) Genotype	BGEN	(A) or (B)	7 days
Hepatitis B 'e' Antigen and Antibody	HEPE	<b>3</b>	1 day
Hepatitis B Core Antibody – IgM	НВСМ	<b>B</b>	1 day
Hepatitis B Core Antibody – Total	НВС	<b>B</b>	1 day
Hepatitis B DNA (Viral load)	DNAB	(A) or (B)	5 days
Hepatitis B Immunity (IgG) (Self-collect)	THBI	(TDL Tiny)	1 day
Hepatitis B Immunity (IgG) (Venous)	HBIM	<b>3</b>	1 day

TEST	CODE	SAMPLE REQS	TAT
Hepatitis B Profile	HEPB	В	1 day
Hepatitis B Resistant Mutation	HBRM	A or B	7 days
Hepatitis B Surface Antigen (Self-collect)	THBA	(TDL Tiny)	1 day
Hepatitis B Surface Antigen (Venous)	AUAG	B	1 day
Hepatitis C Abs Confirmation (RIBA)	RIBA	B	5 days
Hepatitis C Antibodies (Self-collect)	THCV	(TDL Tiny)	1 day
Hepatitis C Antibodies (Venous)	HEPC	B	1 day
Hepatitis C Antigen (Early detection) (Self-collect)	TCAG	B (TDL Tiny)	1 day
Hepatitis C Antigen (Early detection) (Venous)	HCAG	В	1 day
Hepatitis C Genotype	CGEN	A or B	5 days
Hepatitis C Quantification (Viral Load)	QPCR	A or B	5 days
Hepatitis Delta Antibody	HEPD	В	5 days
Hepatitis Delta Antigen	HDAG	В	5 days
Hepatitis Delta RNA	DRNA	A	5 days
Hepatitis E IgG/IgM	HBE	В	5 days
Hepatitis E RNA (PCR)	EHEP	A	2 weeks
Hepatitis G (PCR)	HEPG	(Frozen plasma)	2 weeks

## Hepatitis viral load sample instructions

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.

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.

<b>Hepatitis B I</b> I	mmunity/
<b>Vaccination</b>	Anti-HBs

less than 10 mIU/mI	Non-immune to Hepatitis B
10-50 mIU/mI	Borderline – booster indicated
50-100 mIU/mI	Low level immunity – booster suggested
100 mIU/mI and over	Immune to Hepatitis B

## **HAV, HBV and HCV assays**

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 A 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 Antigen (HCAG)**

HVC Antigen is detectable well before the occurrence of antibodies against HCV. When virus is present, but antibodies are not detectable, a negative antibody test does not rule out HCV infection. Active HCV infection, either acute or chronic is characterised by the presence of HCV Antigen. This is analogous to HepB sAg (AUAG) in active HBV Infection.

#### **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.

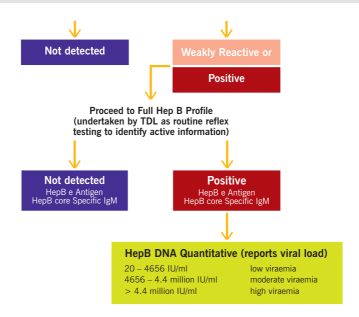
#### **HCV Drug Resistance**

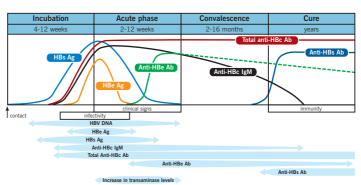
Detects hepatitis C wild-type or drug-induced mutations associated with resistance to HCV drugs including NS5A inhibitors, NS5B inhibitors or NS3 inhibitors.

## **Hepatitis B Surface Antigen**

#### **Hepatitis B**

- 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 HBc IgM, total anti HBc Ab, Anti-HBs Ab, HBe Ag, Anti-HBe Ab, HBV DNA.



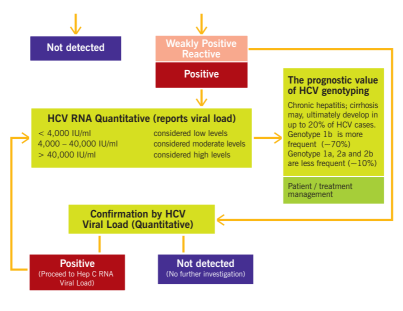


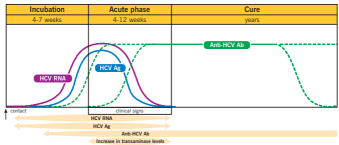
## **Hepatitis C Antibodies**

#### **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 REQS	TAT
HIV-1 Proviral DNA	HIVP	A	7 days
HIV Confirmation of Positive Screens (3 methodologies)	HIVC	3	1 day
HIV/HBV/HCV Screen by PCR/ NAAT (10 days post exposure)	STDX	(Vacutainer only)	3 days
HIV Rapid RNA HIV-1 QUALITATIVE	LHIV	(Vacutainer only)	1 day
HIV Rapid RNA HIV-1 QUANTITATIVE	RHIV	(Vacutainer only)	1 day
HIV Screening: HIV1 & 2 Abs/p24 Ag (4th Gen)	HDU0	<b>B</b>	1 day
HTLV 1 & 2 Abs. (Human T Lymphotropic Virus Type I-II)	HTLV	3	1 day
HTLV by PCR	HTLP	A	21 days

## **HIV** positive patient monitoring

TEST	CODE	SAMPLE REQS	TAT
HIV-1 RNA Viral Load by PCR	HIV1	(2 x 6ml)	3 days
HIV-2 RNA by PCR	HIV2	A	10 days
HIV Rapid RNA HIV-1 QUANTITATIVE	RHIV	(Vacutainer only)	1 day
HIV Therapeutic Drug Monitoring	TDM	<b>J</b> <sup>1</sup>	21 days
Lymphocyte Subsets (CD3/CD4/CD8)	LYSS	A	1 day

## **HIV-1** genotypic resistance testing

TEST	CODE	SAMPLE REQS	TAT
HIV-1 Genotypic Resistance (Integrase)	INTE	<b>A</b> ( 2 x 6ml)	21 days
HIV-1 Genotypic Resistance (RT & Protease)	HIVD	(2 x 6ml)	21 days
HIV-1 Tropism	TRPM	(2 x 6ml)	28 days
HLA B*57:01	HL57	<b>A</b> 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.

## **Virology - General**

TEST	CODE	SAMPLE REQS	TAT
Adenovirus by PCR	ADV	A / PCR / VS	7 days
Arbovirus Antibodies/Abs	ARB0	<b>B</b> 9,14	3 weeks
Atypical Pneumonia Screen	APS	В	3 days
BK Polyoma Virus by PCR	BKPV	(A)/RU	5 days
Cat Scratch Fever (Bartonella IgG)	CAT	В	5 days
CD3/CD4/CD8	LYSS	A	1 day
Chikungunya Virus Abs	CHIK	B 9,14	10 days
CMV IgM Antibodies	CMVM	(Plasma) or (Serum)	1 day
COVID-19 (SARS-CoV-2) (PCR) Contact Laboratory.	NCOV	PCR Swab (nasal/ pharyngeal)	1 day
COVID-19 (SARS-CoV-2) RNA by PCR (Self-collect) Contact Laboratory.	NCOV	Aptima multisite swab of nose/throat	1 day
CSF Screen by PCR	VPCR	CSF	2 days
Cytomegalovirus (CMV-DNA) Amnio	CMVD	AF	5 days
Cytomegalovirus (IgG/IgM) Antibodies	CMV	В	1 day
Cytomegalovirus (PCR) Semen	SCVM	Semen	7 days
Cytomegalovirus (PCR) Urine	CMVU	RU	5 days
Cytomegalovirus Avidity	CMAV	В	10 days
Cytomegalovirus DNA (PCR)	CMVP	A	5 days
Cytomegalovirus Resistance	CMVR	(6mls)	21 days
Dengue Fever PCR	DPCR	A or B 9,14	2 weeks
Epstein-Barr Virus Antibodies IgG/IgM	EBVA	В	2 days
Epstein-Barr Virus PCR	EBVQ	A	5 days
Hantavirus Serology	HANV	B 9	10 days
Herpes Simplex (HSV) 1 & 2 – Genital lesion (Self-collect)	HERS	Aptima multisite swab	5 days
Herpes Simplex (HSV) 1 & 2 – Oral lesion (Self-collect)	HERS	Aptima multisite swab	5 days
Herpes Simplex (HSV) 1 & 2 (PCR) (Oral or Genital)	HERS	PCR	5 days
Herpes Simplex I/II Antibody Profile (IgG)	HERP	<b>3</b>	2 days
Herpes Simplex I/II by PCR (Urine)	HERD	FCRU	5 days
Herpes Simplex I/II IgM	HERM	<b>B</b>	2 days
HIV/HBV/HCV Screen by PCR/ NAAT (10 days post exposure)	STDX	2 x 6mls or 2 x 4mls (Vacutainer only)	3 days

TEST	CODE	SAMPLE REQS	TAT
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) / CSF	5 days
Measles Antibodies (IgG) Immunity	MEAS	В	1 day
Measles Antibodies (IgM)	MEAM	<b>B</b> 9	2 days
Measles PCR	MEAP	Buccal swab	48 hours
MERS Coronavirus Test	MERS	J	1 day
Mumps Antibodies (IgG)	MUMP	В	1 day
Mumps Antibodies (IgM)	MUMM	В	1 day
Mycoplasma pneumoniae IgM and IgG	MYCO	В	2 days
Mycoplasma species – DNA	MPCR	A	5 days
Needle Stick Injury Profile	NSI	BB	1 day
Neurological Viral Screen	NVIR	BB	2 days
Parvovirus Antibodies (IgG)	PARG	В	2 days
Parvovirus Antibodies (IgM)	PARV	В	2 days
Parvovirus IgG/IgM Abs	PARP	В	2 days
Pneumonia (Atypical) Screen	APS	В	3 days
Respiratory PCR Panel (COVID-19, Flu A/B and RSV) (PCR)	FLU4	<b>PCR</b> nasopharyngeal swab	2 days
Respiratory PCR Panel (COVID-19, Flu A/B and RSV) (Self-collect)	FLU4	Aptima multisite swab of nose/throat	2 days
Rotavirus in Stool by PCR	ROTA	RF	1 day
Rubella Antibody (IgG)	RUBE	В	1 day
Rubella Antibody (IgM)	RUBM	В	1 day
Rubella Avidity	RUAV	В	1 week
Torch Screen	TORC	В	2 days
Varicella zoster – DNA	VZPC	A	5 days
Varicella zoster Antibodies (IgG)	VZOS	В	1 day
Varicella zoster Antibodies (IgM)	VZOM	В	1 day
Viral Antibody Screen	VIRA	BB	2 days
Viral Eye by PCR	VPE	PCR	3 days
Viral Respiratory RNA Screen by PCR	VPR	<b>PCR</b> or as specified on the form	2 days
Viral Respiratory RNA Screen by PCR (Self-collect)	VPR	Aptima multisite swab of nose/throat	2 days

TEST	CODE	SAMPLE REQS	TAT
Viral Skin/Mucosa by PCR	VPSK	PCR	5 days
West Nile Virus Abs	WNV	B	2 weeks
Zika Abs IgM and IgG – Antibody detection from 15 days	ZKAB	В	Up to 14 days
Zika RNA by PCR in Semen	ZIKS	Semen	Up to 14 days

#### Atypical Pneumonia Screen

Mycoplasma pneumonia Abs Chlamydia pneumoniae (MIF) Legionella pneumophila (IF)

TAT: 3 days

APS

₿

# Respiratory PCR Panel (COVID-19, Flu A/B and RSV) (PCR)

Flu A Flu B Respiratory Syncytal Virus (RSV) COVID-19

TAT: 2 days

FLU4

PCR nasopharyngeal swab

## **CSF Screen by PCR**

Herpes Simplex virus Varicella zoster virus Enterovirus Parechovirus

TAT: 2 days

**VPCR** 

**CSF** 

### **Hepatitis (Acute) Screen**

Hepatitis A (IgM) Hepatitis B Surface Antigen Hepatitis C Antibodies

TAT: 1 day

**AHSC** 

**3** 

### **Hepatitis A, B & C Profile**

Hepatitis A Profile Hepatitis B Profile Hepatitis C Antibodies Liver Function Tests (LFT)

TAT: 1 day

ABC

BB

#### **Hepatitis B Profile**

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

TAT: 1 day

**HEPB** 

ß

## HIV/HBV/HCV Screen by PCR/NAAT (10 days post exposure)

Positive findings will be reflexed for confirmatory testing

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

STDX provides diagnostic confirmatory testing only when used in addition to serology for Ag/Ab HIV-1&2, HBV, HCV

TAT: 3 days

STDX

A 2 x 6mls or 2 x 4mls (Vacutainer only)

## HIV Rapid RNA HIV-1 OUALITATIVE

Early detection from 10 days HIV-1 RNA

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

TAT: 1 day

LHIV

(Vacutainer only)

## HIV Rapid RNA HIV-1 OUANTITATIVE

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: 1 day

RHIV

(Vacutainer only)

#### Needle Stick Injury Profile

(Donor – Not recipient)
Hepatitis B Surface Antigen
Hepatitis C Antibodies
HIV 1 & 2/p24Ag
Serum saved for 2 years

TAT: 1 day

NSI

**B B** 

## Neurological Viral Screen

Measles IgG Measles IgM Mumps IgG Mumps IgM CMV IgG HSV 1+2 IgG HSV 1+2 IgM VZV IgG

TAT: 2 days

**NVIR** 



#### **Torch Screen**

Toxoplasma Antibodies (IgG, IgM) Rubella Antibody (IgG, IgM) CMV Antibody (IgG, IgM) HSV Antibody (HSV1/HSV2 IgG)

TAT: 2 days

TORC



## **Viral Antibody Screen**

Measles IgG Measles IgM Mumps IgG Mumps IgM Mycoplasma pneumonia CMV HSV 1 HSV 2

TAT: 2 days

VIRA



## Viral Eye by PCR

Herpes Simplex virus Varicella zoster virus Adenovirus

TAT: 3 days

**VPE** 

PCR

#### Viral Respiratory RNA Screen by PCR

Throat swabs, nasopharyngeal aspirates

Adenovirus Parainfluenza 1,2,3,4 Influenza A and B

Coronavirus (seasonal) SARS-CoV-2 (COVID-19) Parechovirus

Rhinovirus

Enterovirus

Respiratory Syncytial virus A and B Human metapneumovirus

TAT: 2 days

**VPR** 

PCR or as specified on the form

## Viral Skin/ Mucosa by PCR

If chicken pox or shingles suspected, please indicate clearly on request form

Herpes Simplex virus Varicella zoster virus

TAT: 5 days

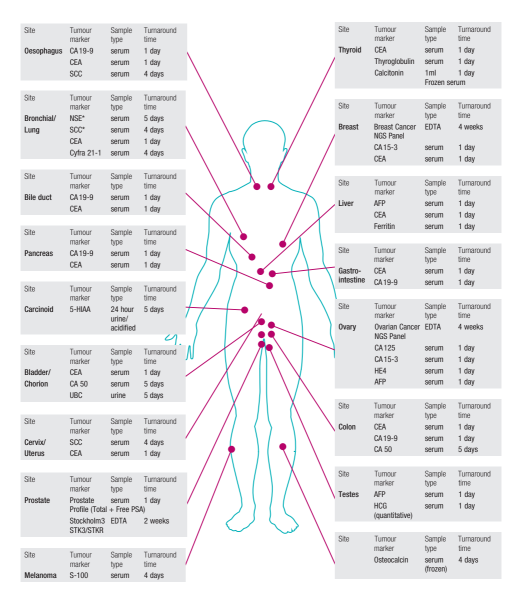
**VPSK** 

**PCR** 

## **Tumour Markers/Sites**

TEST	CODE	SAMPLE REQS	TAT
Alpha-Fetoprotein	AFP	B	1 day
Breast Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
CA 15-3	C153	В	1 day
CA 19-9	C199	B	1 day
CA 50	CA50	B	5 days
CA 72-4	C724	B	5 days
CA 125 (Self-collect)	C125	(TDL Tiny)	1 day
CA 125 (Venous)	C125	B	1 day
Carcino Embryonic Antigen	CEA	B	1 day
Complex PSA (Prostate Specific Ag)	CPSA	B	3 days
Cyfra 21-1	CY21	B	4 days
HCG (Oncology)	HCGQ	B	1 day
HE4 + ROMA (Earlier Detection of Ovarian Tumour)	HE4	В	1 day
Neurone Specific Enolase	NSE	B	5 days
Osteocalcin	0ST	○ (Frozen) <sup>4</sup>	4 days
Prostate Profile (Total & Free PSA)	PR2	B	1 day
Prostate Specific Antigen (Total) (Self-collect)	PSPA	(TDL Tiny)	1 day
Prostate Specific Antigen (Total) (Venous)*  * 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.	PSPA	<b>B</b>	1 day
Pyruvate Kinase (M2-PK)	M2ST	RF <sup>4</sup>	5 days
Pyruvate Kinase (M2-PK)	M2PK	(Frozen plasma) <sup>7</sup>	5 days
S100 Malignant Melanoma	S100	B	4 days
Squamous Cell Carcinoma	SCC	B	4 days
Stockholm3 NEW Samples must be received by TDL within 24 hours of sample taking.	STK3	AA	2 weeks
Stockholm3 Reflex NEW Sample must be received by TDL within 24 hours of sample taking.	STKR	AAB	2 weeks
Testicular Tumour Profile (LDH, AFP, HCQG)	TTP	В	1 day
Urinary Bladder Cancer Antigen	UBC	RU (Freeze within	5 days
** It is recommended to collect mid-stream urine. Do not use first morning urine. Collection of urine specimen before any surgical intervention or treatment or 1–2 weeks after specim shall not be collected with an instrument e.g. catheter.		48 hours)**	

#### **Tumour Markers/Sites**



\* NSE: Neurone Specific Enolase SCC: Squamous Cell Carcinoma

#### **Tumour Markers/Sites**

## HE4 + ROMA (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 & Free PSA)

Prostate Specific Antigen (Total) Free PSA Calculated Ratio

The ratio of Free to Total PSA may help discriminate between prostate cancer and benign prostatic hyperplasia.

#### TAT: 1 day

PR2

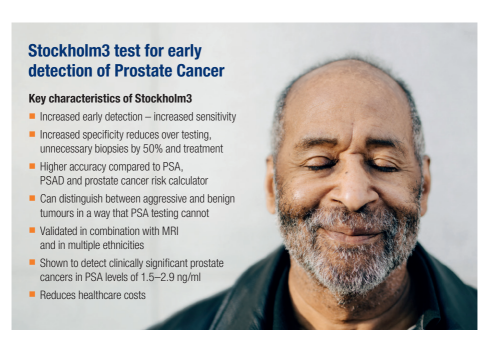
#### **Testicular Tumour Profile**

Lactate Dehydrogenase (LDH) Alpha-Fetoprotein HCG (Oncology)

## TAT: 1 day

TTP





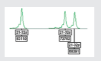
TDL Genetics is a consultant-led service which is able to provide extensive expertise in the testing, diagnosis and genetic counselling of inherited disorders. Genetic tests are performed on DNA for molecular genetic analysis and on whole chromosomes for cytogenetic analysis. Some tests are part of profiles 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 QF-PCR (Amnio-PCR and CVS-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

QF-PCR: DNA peaks from an unaffected fetus

QF-PCR: DNA peaks from a fetus with Down Syndrome



Genetic testing is sometimes complex and tests will vary in their ability to detect variants 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 (including informed consent). Our service provides result interpretation and risk assessment. Genetic counselling can be arranged by TDL's Consultant Clinical Geneticist.



Download TDL Request Forms from:

www.tdlpathology.com/ tests/request-forms/

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. The Non-invasive prenatal testing (NIPT) laboratory is open Monday to Saturday.

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. For NIPT sample stability see page 134, do not refrigerate or freeze NIPT bloods.

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 birth 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 (reason for referral), 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

- Patient's surname and forename (not initials)
- Date of birth
- Hospital number or reference number

#### **Consent forms**

Consent forms (see tdlpathology.com/tests/ request-forms) 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 variant has already been shown to be segregating in a family, to be provided with information concerning the variant 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 Genomic 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 22q11.2 in suspected DiGeorge 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, delayed puberty, 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

- Rarely, blood samples fail to culture (<1%);</p>
- The culture may yield chromosomes of insufficient quality. This will be indicated on the report and a repeat study suggested;
- The laboratory should be informed if the patient has recently received a blood transfusion.
- The laboratory should be informed if the patient has EVER had a bone marrow transplant.
- The patient's birth sex should be included on the request form.
- For fetal blood samples a maternal blood sample must also be provided to confirm fetal origin.

## **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:

- 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.
- 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.

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. For fetal blood samples a maternal blood sample must also be provided to confirm fetal origin.

**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 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.
- Sampling at early or late gestations, for example to confirm non-invasive tests or follow up anomaly scans.
- A tendency to take smaller quantities of sample or to take insufficient sample for multiple techniques.
- 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.

#### Notes

- Maternal contamination, and mosaicism may complicate the analysis and may lead to the suggestion that a second invasive test is performed.
- Rarely, cultures fail to grow (overall <1%)</p>
- Chromosome abnormalities smaller than resolution of G-band analysis may not be detected (higher resolution techniques such as Array-CGH may be more appropriate in certain circumstances).
- For TTTs 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.



For more information on Products of conception for genetic investigations

SCAN ME

www.tdlpathology.com/poc

**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

- 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.
- 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).

- If a request is made for remaining pregnancy loss tissue to be returned to the patient or hospital for burial or cremation, we will return the sample as soon as possible once adequate tissues have been used for testing. Please ensure that this is communicated to the lab using a hospital consent form, noted on the referral form or by email. Patients can arrange to collect remaining tissues from TDL Patient Reception.
- The lab will send all remaining tissue for samples without specific consent, for sensitive incineration. Please note that there is no distinction made between fetal and other pregnancy tissues for this process and there will be no ashes afterwards. The lab keeps detailed records of all pregnancy tissue sent for incineration and a Certificate of Destruction is available if required.

# 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.

# 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**

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TEST	CODE	SAMPLE REQS	TAT
1p36 Deletion Syndrome – karyotype + CGH	KARY, FISH	CVS / AF / (1) 9	12-17 days
21-Hydroxylase Deficiency (Congenital Adrenal Hyperplasia CYP21A2) Requires patient informed consent.	GENE	9,11	5 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) (1) 9	5-15 days
Achromatopsia NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	5 weeks
Adenomatous Polyposis NGS Panel	GENE	<b>A</b> 9	4 weeks
Aicardi-Goutières Syndrome NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	5 weeks
Alagille Syndrome NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
Alpha Thalassaemia – alpha globin gene sequencing + deletions/duplications	GENE	A	4 weeks
Alpha Thalassaemia – multiplex PCR for common large deletions Requires patient informed consent.	GENE	<b>A</b> 9	3 weeks
Alpha-1-Antitrypsin Genotype – PI*M, PI*S, PI*Z Requires patient informed consent.	GENE	<b>A</b> 9	3 weeks
Alport Syndrome NGS Panel – full sequencing with deletions and duplications Requires patient informed consent.	GENE	<b>A A</b> 9	5 weeks
Alzheimer and Dementia NGS Panel Requires patient informed consent.	GENE	<b>A</b> 9,11	6 weeks
AML/ALL Molecular MRD – NPM1, PML-RARA, CBFB-MYH11, RUNX1-RUNX1T1, ETV6-RUNX1 Requires patient informed consent. Must arrive in the laborate 48 hours, before 12pm on Fridays. Contact lab for further info	,	Bone Marrow / 🛕 9	5 days
AmnioBOBs only – rapid aneuploidy diagnosis for all chromosomes + common microdeletion syndromes	ABOB	AF <sup>9</sup>	5 days
Amniocentesis – rapid BOBs aneuploidy diagnosis for all chromosomes (5 days) + culture (10-15 days)	ABK	AF <sup>9</sup>	5-15 days
Amniocentesis – rapid PCR diagnosis for common aneuploidies (2 days) + culture (10-15 days)	APCC	AF <sup>9</sup>	2-15 days
Amniocentesis culture (karyotype) only	ACUL	AF 9	10-15 days
AmnioPCR only – rapid common	APC	AF 9	2 days

TEST	CODE	SAMPLE REQS	TAT
Amyotrophic Lateral Sclerosis (Motor Neurone Disease) NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	5 weeks
Androgen Insensitivity – AR gene sequencing Requires patient informed consent.	GENE	<b>A</b> 9	5 weeks
Angelman Syndrome (Primary Screen)  – methylation test	PWAM	<b>A</b> 9	10 days
Angelman/Rett Syndromes NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	5 weeks
Aniridia, Isolated – PAX6 gene sequencing + deletions/duplications Requires patient informed consent.	GENE	<b>A</b> 9	5 weeks
Anophthalmia/Microphthalmia/ Coloboma NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Apolipoprotein E genotype – E2/E3/E4	APEG	<b>A</b> 9,11	2 weeks
Array-CGH (Comparative Genomic Hybridisation) SNP array	CGH	CVS / AF / (A) (1) 9	10 days
Ashkenazi Breast Cancer Screen  – common variants Requires patient informed consent.	GENE	<b>A</b> 9,11	4 weeks
Ashkenazi Jewish Carrier Screen Requires patient informed consent. See Carrier Screen on page 133 for details	GENE	<b>A</b> 9	4 weeks
Ataxia NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
Autoinflammation/Periodic Fever NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Azoospermia – karyotype + Y deletions + cystic fibrosis screen (+ polyT(5T) when clinically relevant)	GRP	<b>A (1)</b> 9	10-15 days
B cell Clonality Assay (IgH and IgK)	IGHA	(A) or FFPE	2 weeks
Bardet-Biedl Syndrome NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Batten Disease (Neuronal Ceroid Lipofuscinosis) NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
BCR-ABL Diagnostic Assay	BCRD	A	2 weeks
BCR/ABL Quantitative – fusion gene sizes p190 + p210 MUST arrive in the laboratory within 48 hours, before 12pm on Fridays.	BCRQ	<b>A A</b> <sup>9</sup>	10 days
Becker/Duchenne Muscular Dystrophy – deletions/duplications	DMD1	<b>A</b> 9	10 days

TEST	CODE	SAMPLE REQS	TAT
Beckwith-Wiedemann Syndrome  – methylation studies on 11p15 imprinting domains KvDMR + H19 Requires patient informed consent.	GENE	<b>A</b> 9,11	6 weeks
Behcet's Disease – HLA Tissue Typing B*51	B51	<b>A</b> 9	10 days
Beta Thalassaemia – beta-globin gene sequencing + deletions/duplications Requires patient informed consent.	GENE	<b>A</b> 9	4 weeks
Bleeding and Platelet Gene Panel Requires patient informed consent. Contact lab.	R90U	<b>A A</b> <sup>9</sup>	12 weeks
Blood PCR for Chromosome 13, 18, 21 and sex chromosomes	BPCR	Δ	5 days
Breast Cancer – BRCA1 + BRCA2 genes only Requires patient informed consent.	GENE	<b>A</b> 9,11	4 weeks
Breast Cancer Ashkenazi Screen  – common variants Requires patient informed consent.	GENE	<b>A</b> 9,11	4 weeks
Breast Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
Brugada Syndrome/Long QT Syndrome NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	4-6 weeks
C-KIT D816V variant – Mastocytosis Requires patient informed consent.	GENE	Bone Marrow / (A) 9	4 weeks
CADASIL – NOTCH3 gene sequencing Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
CAKUT (Congenital Anomalies of Kidney & Urinary Tract) NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
Cancer, Comprehensive NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	5 weeks
Cardiomyopathy, Dilated NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Cardiomyopathy, Hypertrophic NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Cardiovascular, Comprehensive NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Carrier Screen (Ashkenazi Jewish) Requires patient informed consent.	GENE	<b>A</b> 9	4 weeks
Carrier Screen (Ashkenazi Jewish)  — Partnered Report  Requires patient informed consent. Please contact the lab for special requirements before sending.	GENE	<b>A</b> 9	4 weeks
Carrier Screen (Pan-Ethnic) Requires patient informed consent.	GENE	<b>A</b> 9	4 weeks

Carrier Screen (Pan-Ethnic) – Partnered Report	_		
Requires patient informed consent. Please contact the lab for special requirements before sending.	GENE	<b>A</b> 9	4 weeks
Charcot-Marie-Tooth Syndrome NGS Panel Requires patient informed consent. Contact lab prior to sendi neurologist or clinical geneticist required with genetic conser		A A <sup>9</sup> from clinical	6 weeks
Charcot-Marie-Tooth Type 1A  – PMP22 duplications	GENE	<b>A</b> 9	6 weeks
Requires patient informed consent. Contact lab prior to sending. Referral from clinical neurologist or clinical geneticist required with genetic consent form.			
<b>CHARGE Syndrome – CHD7 gene sequencing</b> Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Cholestasis NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Chromosome Analysis (Amniocentesis)  – culture only	ACUL	<b>AF</b> <sup>9</sup>	10-15 days
Chromosome Analysis (Amniocentesis)  – rapid BOBs aneuploidy diagnosis for all chromosomes (5 days) + culture (10-15 days)	ABK	<b>AF</b> <sup>9</sup>	5-15 days
Chromosome Analysis (Amniocentesis) – rapid PCR diagnosis for common aneuploidies (2 days) + culture (10-15 days)	APCC	<b>AF</b> <sup>9</sup>	2-15 days
Chromosome Analysis (Blood)	KARY	<b>(1)</b> 9	2-3 weeks
Chromosome Analysis (Chorionic Villus)  - rapid BOBs aneuploidy diagnosis for all chromosomes (5 days) + culture (10-15 days)	CBK	CVS <sup>9</sup>	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)	PROC	Placental Sample 1,9	20-25 days
Chromosome Analysis (Products of Conception)  – BOBs rapid aneuploidy diagnosis for all chromosomes (10 days) + culture (25 days)	PBK	Placental Sample 1,9	10-25 days
Chromosome Analysis (Solid Tissue)	PROC	Fetal tissue 1,9	4-5 weeks
Chromosome Y Deletion – AZFa, AZFb, AZFc + SRY	YDEL	<b>A</b> 9	5 days
Coeliac Disease – HLA DQ2/DQ8 Genotype	Q2Q8	<b>A</b> 9	10 days
Colorectal Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
Comparative Genomic Hybridisation (CGH) SNP array	CGH	CVS / AF / (A) (1) 9	10 days

TEST	CODE	SAMPLE REQS	TAT
Congenital Absence of Vas Deferens – karyotype + Y deletions + cystic fibrosis screen (+ polyT(5T) when clinically relevant)	GRP	<b>A</b> ( ) 9	10-15 days
Congenital Adrenal Hyperplasia NGS Panel	GENE	<b>A</b> 9	6 weeks
Congenital Myopathy NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
<b>Connective Tissue Disorders NGS Panel</b> Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Cornelia de Lange Syndrome NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
<b>Craniosynostosis NGS Panel</b> Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Cri du Chat Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, Kary	CVS / AF / (A) (1) 9	5-15 days
Cri du Chat Syndrome – BOBs only	PB0B	CVS / AF / (A) 9	5 days
CVS PCR for common aneuploidies (2 days) + culture (10-15 days)	CVPC	CVS 1,9	2-15 days
CVSBOBs – rapid BOBs aneuploidy diagnosis for all chromosomes (5 days) + culture (10-15 days)	CBK	CVS <sup>9</sup>	5-15 days
CVSBOBs only – rapid aneuploidy diagnosis for all chromosomes + common microdeletion syndromes	CBOB	CVS 9	5 days
Cystic Fibrosis (139 common variants)  – reflex to Poly T when required  Please provide relevant clinical and family history.	CFS	<b>A</b> 9	5-7 days
Cytochrome P450 2C19	2C19	<b>A</b> 9	10 days
Diabetes – Obesity NGS Panel Requires patient informed consent.	GENE	A	6 weeks
DiGeorge Syndrome (22q11 & 10p14 deletion) – BOBs (5 days) + karyotype (15 days)	DGB, KARY	CVS / AF / (A) (1) 9	5-15 days
DiGeorge Syndrome (22q11 & 10p14)  – BOBs only	DGB	CVS / AF / (A) 9	5 days
Dihydropyrimidine Dehydrogenase deficiency screening (Fluoropyrimidine Toxicity)	5FU	<b>A</b> 9	1-2 weeks
Dilated Cardiomyopathy NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
DNA Extraction & Storage – 3 years (longer upon request)	XDNA	<b>A</b> 9	20 days
DNA Identity Profile – 15 STR markers	DNAF	<b>A</b> 9,11	10 days
Duchenne Muscular Dystrophy – deletions/duplications only	DMD1	<b>A</b> 9	10 days

TEST	CODE	SAMPLE REQS	TAT
Duchenne Muscular Dystrophy – full sequencing DMD1 gene Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
DVT/Pre-travel Screen	DVT1	<b>A A B</b> <sup>9</sup>	5 days
Ehlers-Danlos Syndrome NGS Panel	GENE	<b>A</b> 9,11	6 weeks
Endometrial Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
Epidermolysis Bullosa NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
Epilepsy, Adolescent/Adult Onset Panel Requires patient informed consent.	GENE	Δ	6 weeks
<b>Epilepsy, Comprehensive NGS Panel</b> Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
Fabry Disease, X-linked – GLA gene sequencing	GENE	<b>A</b> 9	4 weeks
Facioscapulohumeral Muscular Dystropy (FSHD) – D4Z4 repeat deletion Requires patient informed consent. Contact lab prior to sending. Referrals only from consultant neurologist or clinical geneticist. Genetic consent form required.	GENE	<b>A A</b> <sup>9</sup>	9 weeks
Factor II Prothrombin – G20210A Variant	FX2	<b>A</b> 9	5 days
Factor V Leiden – G1691A Variant	FX5	<b>A</b> 9	5 days
Familial Hypercholesterolaemia NGS panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Familial Hypocalciuric Hypercalcaemia (FHH) Panel Requires patient informed consent.	GENE	<b>AA</b> <sup>9</sup>	6-7 weeks
Familial Mediterranean fever MEFV gene sequencing Requires patient informed consent.	GENE	<b>A</b> 9	5 weeks
Familial Medullary Thyroid Carcinoma  – hotspot sequencing RET gene Requires patient informed consent.	GENE	<b>A</b> 9,11	6-7 weeks
Fatty Acid Oxidation Deficiency NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
FLT3-ITD and FLT3-TKD screening assay	FLT3	A	24 hours
Fragile X Syndrome screen – FMR1 repeat analysis PCR Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	5 weeks
Friedreich Ataxia – frataxin gene repeat analysis Requires patient informed consent.	GENE	<b>A</b> 9	5 weeks
Gaucher Disease full gene sequencing	GENE	<b>A</b> 9	6 weeks
Genetic Reproductive Profile (Male)	GRP	<b>A</b> (1) 9	10-15 days

TEST	CODE	SAMPLE REQS	TAT
Gilbert Syndrome – common UGT1A1 repeat variation Requires patient informed consent.	GENE	<b>A</b> 9	2-3 weeks
Glucose-6-Phosphate Dehydrogenase (G6PD) Deficiency – full G6PD gene sequencing Requires patient informed consent.	GENE	<b>A</b> 9	3-4 weeks
Glycogen storage disease type 2 (Pompe) variant analysis	POMP	A	4 weeks
Haemochromatosis – HFE common variants C282Y + H63D	HMD	<b>A</b> 9	3 days
Haemophilia A (Factor VIII deficiency) – CVS	8CVS	CVS 40	3 days
Haemophilia B (Factor IX deficiency) – CVS	9CVS	CVS 40	3 days
Hearing Loss NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Hereditary Colorectal Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
Hereditary Comprehensive Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	5 weeks
Hereditary Neuropathy with Liability to Pressure Palsy – PMP22 deletion analysis Requires patient informed consent. Contact lab prior to sending. Referrals only from consultant neurologist or clinical geneticist. Genetic consent form required.	GENE	<b>A</b> 9	6 weeks
Hereditary Spastic Paraplegia Comprehensive NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	5 weeks
HFE gene (Haemochromatosis) – common variants C282Y + H63D	HMD	<b>A</b> 9	3 days
Hirschprung Disease NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
HLA Tissue Typing A	HLA	<b>A</b> 9	10 days
HLA Tissue Typing A+B	HLBA	<b>A</b> 9	10 days
HLA Tissue Typing A+B+C (Class I)	HABC	<b>A</b> 9	10 days
HLA Tissue Typing A/B/DRB1/3/4/5	HLAF	<b>A</b> 9	10 days
HLA Tissue Typing A/B/DRB1/3/4/5/DQB1	HLF	<b>A</b> 9	10 days
HLA Tissue Typing A/B/C/ DRB1/3/4/5/DQB1 (Class I & II)	HLFC	<b>A</b> 9	10 days
HLA Tissue Typing B	HLB	<b>A</b> 9	10 days
HLA Tissue Typing B*27 only	HLAB	<b>A</b> 9	3 days
HLA Tissue Typing B*51 (Behcet's Disease)	B51	<b>A</b> 9	10 days
HLA Tissue Typing B*57:01 high resolution	HL57	<b>A</b> 9	10 days

TEST	CODE	SAMPLE REQS	TAT
HLA Tissue Typing C	HLC	<b>A</b> 9	10 days
HLA Tissue Typing Coeliac Disease – DQ2/DQ8	Q2Q8	<b>A</b> 9	10 days
HLA Tissue Typing DRB1/3/4/5	DRB1	<b>A</b> 9	10 days
HLA Tissue Typing DRB1/3/4/5/DQB1 (Class II)	HLDQ	<b>A</b> 9	10 days
HLA Tissue Typing Narcolepsy – DQB1*06:02 Requires patient informed consent.	GENE	<b>A</b> 9	3 weeks
Huntington Disease —  HD gene repeat analysis PCR  Requires patient informed consent. Contact lab prior to sending. Referrals only from consultant neurologist or clinical geneticist. Genetic consent form required.	GENE	<b>A A</b> 9,11	5 weeks
Hyperinsulinism NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Hyperparathyroidism – CASR sequencing Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
dentity Profile (DNA) – 15 STR markers	DNAF	A 9,11	10 days
IDH1/2 Screening Assay Requires patient informed consent.	GENE	<b>A</b>	48 hours
gVH variant analysis for CLL	IGMU	A	4 weeks
Inherited bleeding and platelet disorders (R90) Clinical synopsis, factor levels, bleeding history, family history and informed consent required. Please contact the laboratory for the request and consent forms, or for further guidance.		88	12 weeks
Intellectual Disability NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
ron Overload Profile	IOP	<b>A B</b> 9	3 days
Joubert/Meckel-Gruber Syndrome NGS Panel Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Kallmann Syndrome NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
Kennedy Disease (Spinal Bulbar Muscular Atrophy) – AR repeat expansion Requires patient informed consent.	GENE	<b>A</b> 9	5 weeks
Kidney/Urinary Tract Comprehensive Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
Lactose Intolerance Gene	LACG	A	2 weeks
Langer-Giedion Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS / AF / (A) (1) 9	5-15 days
Langer-Giedion Syndrome – BOBs only	PB0B	CVS / AF / (A) 9	5 days
Leber's Hereditary Optic Neuropathy – m.3460G>A + m.11778G>A + m.14484T>C common variants Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks

TEST	CODE	SAMPLE REQS	TAT
Leukaemia (Rapid Acute) DNA and RNA NGS Panel Requires patient informed consent.	ALRP	(3mL minimum) or bone marrow aspirate sample	3 days
Leukaemia Fusion Gene Screening Assay (Q30)	LMPX	A	24 hours
Leukaemia/Lymphoma RNA Sequencing (Fusion Gene and SNV/Indel) Panel Requires patient informed consent.	PHFP	A	2 weeks
Li-Fraumeni Syndrome (p53-related cancer predisposition) – TP53 sequencing + MLPA Requires patient informed consent.	GENE	<b>A</b> 9,11	6 weeks
Limb-Girdle Muscular Dystrophy (LGMD) NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Lissencephaly NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Long QT Syndrome/Brugada Syndrome NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	4-6 weeks
<b>Lung Disorders NGS Panel</b> Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
<b>Lynch Syndrome (HNPCC) NGS Panel</b> Requires patient informed consent.	GENE	<b>A</b> 9	4 weeks
Lysosomal Storage Disorders NGS Panel – full gene sequencing Requires patient informed consent.	LSDS	<b>A A</b> <sup>9</sup>	4-6 weeks
Male Genetic Reproductive Profile	GRP	<b>A (1)</b> 9	10-15 days
Marfan Syndrome – FBN1 sequencing + deletions/duplications Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Marfan Syndrome and Thoracic Aortic Aneurysm and Dissection NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Maturity-Onset Diabetes of the Young (MODY) Diabetes NGS Panel Requires patient informed consent.	GENE	<b>A</b> 9	12 weeks
Meckel-Gruber/Joubert Syndrome NGS Panel Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Medium-Chain Acyl-CoA Dehydrogenase Deficiency – ACADM sequencing Requires patient informed consent.	GENE	<b>A</b> 9	5 weeks
Melanoma Comprehensive Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
Microdeletion (common) Syndromes – BOBs only	PB0B	CVS / AF / (A) 9	5 days

TEST	CODE	SAMPLE REQS	TAT
Microphthalmia/Anophthalmia/ Coloboma NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Miller-Dieker Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS / AF / (A) (1) 9	5-15 days
Miller-Dieker Syndrome – BOBs only	PB0B	CVS / AF / (A) 9	5 days
Mitochondrial Genome Sequencing Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Motor Neurone Disease (Amylotrophic Lateral Sclerosis) NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	5 weeks
MTHFR – common C677T + A1298C variants	MTHF	<b>A</b> 9	5 days
Mucopolysaccharidosis NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
Multiple Endocrine Neoplasia Type 1  – full MEN1 sequencing Requires patient informed consent.	GENE	<b>A</b> 9,11	6-7 weeks
Multiple Endocrine Neoplasia Type 2  — RET gene hotspot sequencing Requires patient informed consent.	GENE	<b>A</b> 9,11	6-7 weeks
Myeloid Gene Panel Requires patient informed consent.	MVPS	(3mL minimum) or bone marrow aspirate sample	2 weeks
Myeloproliferative Neoplasm NGS Screening Panel Requires patient informed consent.	MPNS	(3mL minimum) or bone marrow aspirate sample	1 week
Myotonic Dystrophy Type 1 – DMPK repeat PCR Requires patient informed consent.	GENE	<b>A</b> 9	5 weeks
Myotonic Dystrophy Type 2 (PROMM) – ZNF9 repeat PCR Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Narcolepsy (HLA DQB1*06:02) Requires patient informed consent.	GENE	<b>A</b> 9	3 weeks
Nephrotic Syndrome, Steroid- Resistant NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Nervous System/Brain Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
Neurofibromatosis Type 1 – NF1 + SPRED1 sequencing + deletions/duplications Requires patient informed consent. Contact lab prior to sending.	GENE	<b>A A</b> 9,11	8 weeks
Neuronal Ceroid Lipofuscinosis (Batten Disease) NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks

TEST	CODE	SAMPLE REQS	TAT
Non-Invasive Prenatal Testing (NIPT) – common aneuploidy screening from maternal blood	NIPT	J / Special tube <sup>1</sup>	2-4 days
<b>Noonan Syndrome and RASopathies NGS Panel</b> Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
Nystagmus, X-linked Infantile – FRMD7 gene sequencing Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Osteogenesis Imperfecta NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Ovarian Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
p53-related cancer predisposition (Li-Fraumeni Syndrome) – TP53 sequencing + MLPA Requires patient informed consent.	GENE	<b>A</b> 9,11	6 weeks
Pancreatic Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
Paraganglioma/Pheochromocytoma NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
Parkinson-Alzheimer-Dementia NGS Panel Requires patient informed consent.	GENE	<b>A</b> 9,11	6 weeks
Paternity Testing (postnatal and prenatal) – sample required from each person being tested (3 people)	PATT	A / AF / CVS 1,12 Contact Genetics lab	5 days
Pelizaeus-Merzbacher Disease – PLP1 sequencing + deletions/duplications Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
<b>Pendred Syndrome – SLC26A4 gene sequencing</b> Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Periodic Fever/Autoinflammation NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
Peutz-Jegher Syndrome – STK11 sequencing + deletions/duplications Requires patient informed consent.	GENE	<b>A</b> 9	5 weeks
Phelan-McDermid Syndrome – karyotype + FISH	KARY, FISH	CVS / AF / (1) 9	12-17 days
Pheochromocytoma/Paraganglioma NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	5 weeks
POLG-Related Disorders – full POLG sequencing + deletions and duplications Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Polycystic Kidney NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
Pontocerebellar Hypoplasia NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks

TEST	CODE	SAMPLE REQS	TAT
Postnatal SNP Array CGH	CGH	<b>A (1)</b> 9	10 days
Prader-Willi Syndrome (Primary Screen)  – methylation test	PWAM	<b>A</b> 9	10 days
Prenatal Diagnosis (BOBs + Culture)	ABK or	AF / CVS 9	3-5 days,
	CBK		15 days
Prenatal SNP Array CGH	CGH	Amniotic fluid, CVS or POC <sup>9</sup>	10 days
Pre-Travel Screen (DVT)	DVT1	<b>A B</b> 9	5 days
Primary Ciliary Dyskinesia (PCD) NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Primary Hyperoxaluria NGS Panel Requires patient informed consent.	GENE	A	7 weeks
Products of Conception – rapid BOBs aneuploidy diagnosis for all chromosomes (10 days) + culture (25 days)	PBK	Placental Sample 1,9	10-25 days
Products of Conception (BOBs + Culture)	PBK	Placental Sample 1,9	10-25 days
Products of Conception BOBs only – rapid aneuploidy diagnosis for all chromosomes	KB0B	Placental Sample or Solid Tissue 1,9	10 days
Prostate Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
QF-PCR rapid common aneuploidy screen	APC	<b>AF</b> / <b>A</b> <sup>9</sup>	2 days
Recurrent Miscarriage Profile (female)	RMP	<b>A A B C C C C H</b> 9,18	10-15 days
Renal Cysts and Diabetes (RCAD)  – HNF-1β sequencing exons 1-9 and dosage analysis by MLPA Requires patient informed consent.	GENE	<b>A</b> 9	8 weeks
Renal/Urinary Tract Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
Retinoblastoma – RB1 sequencing + deletions/duplications Requires patient informed consent.	GENE	<b>A A</b> 9,11	6 weeks
Rett Syndrome (MECP2 gene only) – full sequencing + deletions/duplications Requires patient informed consent.	GENE	<b>A</b> 9,11	6 weeks
Rett/Angelman Syndromes NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	5 weeks
Short-Chain Acyl-CoA Dehydrogenase Deficiency – ACADS sequencing Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Short Stature – SHOX variant screening + deletions/duplications Requires patient informed consent.	GENE	<b>A</b> 9	8 weeks

TEST	CODE	SAMPLE REQS	TAT
Silver-Russell Syndrome – methylation studies on 11p15 imprinting domains KvDMR + H19 Requires patient informed consent.	GENE	<b>A</b> 9	7 weeks
Skeletal Dysplasia NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Smith-Lemli-Opitz Syndrome  – DHCR7 sequencing Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Smith-Magenis Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS / AF / (A) (1) 9	5-15 days
Smith-Magenis Syndrome – BoBs only	PB0B	CVS / AF / (A) 9	5 days
Sotos Syndrome (Cerebral Gigantism) – NSD1 sequencing + deletions/duplications Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Spastic Paraplegia NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Spinal Bulbar Muscular Atrophy (Kennedy Disease) – AR repeat analysis Requires patient informed consent.	GENE	<b>A</b> 9	5 weeks
Spinal Muscular Atrophy – SMN1 deletions/duplications	SMA	<b>A</b> 9	10 days
Spinocerebellar Ataxia – multiplex SCA1+2+3+6+7+8+10+12 +17 common repeat expansions Requires patient informed consent.	GENE	<b>A</b> 9	5 weeks
Spinocerebellar Ataxia NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks
SRY (Sex-determining Region Y)	SRY	<b>A</b> 9	2 days
Systemic Mastocystosis – C-Kit common variants (KIT D816V) Requires patient informed consent.	GENE	<b>A</b> 9	4 weeks
T cell Clonality Assay (TCR beta and TCR gamma)	TCRA	A or FFPE	2 weeks
Tay Sachs Screen Requires patient informed consent. See also Carrier Screen (Ashkenazi Jewish/Pan-Ethnic).	GENE	<b>A</b> 9	4 weeks
Thrombophilia with a likely monogenic cause (R97) Clinical synopsis, levels of relevant proteins, thrombosis hist family history and informed consent required. Please contac laboratory for the request and consent forms, or for further the content is the content forms.	t the	<b>AA</b>	12 weeks
Thrombotic Risk Profile	PR0P	<b>A A B O O O</b> 18	5 days
Thyroid Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks

TEST	CODE	SAMPLE REQS	TAT
Treacher Collins Syndrome and Related Disorders NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9	6 weeks
Tuberous Sclerosis (TSC1 + TSC2) Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	7 weeks
Urinary Tract/Renal Cancer NGS Panel Requires patient informed consent.	GENE	<b>A A</b> 9,11	4 weeks
Usher Syndrome NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Very Long-Chain Acyl-CoA Dehydrogenase Deficiency – ACADVL sequencing Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Von Hippel-Lindau Syndrome – VHL sequencing + deletions/duplications Requires patient informed consent.	GENE	<b>A</b> 9	6 weeks
Whole Genome Sequencing (solo/duo/trio) For whole exome sequencing please contact the lab for further information.	GENE	<b>A</b> 9,11	5-8 weeks
Wolf-Hirschhorn Syndrome – BOBs (5 days) + karyotype (15 days)	PBOB, KARY	CVS / AF / (A) (1) 9	5-15 days
Wolf-Hirschhorn Syndrome – BOBs only	PB0B	CVS / AF / (A) 9	5 days
Y chromosome microdeletions – AZFa + AZFb + AZFc + SRY	YDEL	<b>A</b> 9	5 days
Zellweger Syndrome NGS Panel Requires patient informed consent.	GENE	<b>A A</b> <sup>9</sup>	6 weeks
Ziwig Endotest® For information about this test and to order kits please contact endotest@tdlpathology.com. The quality of the saliva sample collection is important. Samples can be collected in the clinic or at home following instructions provided.	ENDT	Endotest saliva collection kit	25 days
Zygosity Testing – comparative DNA profile	DNAC	(From each twin and both parents) <sup>9</sup>	5 days

# Carrier Screen (Ashkenazi Jewish)

This test is optimised for individuals and couples of Ashkenazi Jewish ancestry.\*\*

Uses the same technology as the Pan-Ethnic Carrier Screen.

\*\*Male patients will not be screened for X-linked conditions (e.g., FMR1, etc.). Requires patient informed consent.

#### TAT: 4 weeks

#### **GENE**



# Carrier Screen (Pan-Ethnic)

Targets 400+ Autosomal Recessive and X-linked Inherited Disorders\*\*

\*\* Male patients will not be screened for X-linked conditions (e.g., FMR1, etc.). Requires patient informed consent.

#### TAT: 4 weeks

#### **GENE**



#### **DVT/Pre-travel Screen**

Full Blood Count (FBC)
Factor II Prothrombin –
G20210A Variant
Factor V Leiden – G1691A Variant
Cardiolipin Antibodies (IgG+IgM)

#### TAT: 5 days

DVT1



#### **Iron Overload Profile**

Iron (TIBC included)
Ferritin
Transferrin Saturation
Haemochromatosis C282Y, H63D

#### TAT: 3 days

#### 10P



## Male Genetic Reproductive Profile

Chromosome Analysis (Blood) Cystic Fibrosis (139 common variants) – reflex to Poly T when required

Y chromosome microdeletions
- AZFa + AZFb + AZFc + SRY

#### **TAT: 10-15 days**

#### GRP



# Products of Conception (BOBs + Culture)

Rapid Aneuploidy Diagnosis for all Chromosomes by BOBs Analysis (10 days) Chromosome Analysis (Karyotype) (25 days)

#### **TAT: 10-25 days**

#### **PBK**

Placental Sample 1,9

# Recurrent Miscarriage Profile (female)

Full Blood Count (FBC)
Coagulation Profile
Antithrombin III
Factor V Leiden Common Variant
Factor II Prothrombin
Common Variant
MTHFR Common Variants
Fibrinogen
Lupus Anticoagulant
Protein C
Free Protein S Ag
Anticardiolipin Abs
Chromosome Analysis
Please request Partner's
Chromosome Analysis using

#### **TAT: 10-15 days**

a separate request form.

#### **RMP**



#### **Thrombotic Risk Profile**

Full Blood Count (FBC)
Coagulation Profile 1
Antithrombin III
Factor V Leiden — G1691A Variant
Factor II Prothrombin —
G20210A Variant
MTHFR — common C677T
+ A1298C variants
Lupus Anticoagulant
Protein C
Protein S Free Ag
Cardiolipin Antibodies (IqG+IqM)

#### TAT: 5 days

#### PR0P



## Leukaemia (Rapid Acute) DNA and RNA NGS Panel / Myeloproliferative Neoplasm NGS Screening Panel

This NGS assay allows for rapid generation of comprehensive profile of variants (both DNA and RNA) from a single NGS run. This assay can profile both DNA and RNA targets including DNA mutations and translocations detected from RNA targets and allows for simultaneous interrogation of 45 DNA target genes and 30 RNA fusion driver genes. The broad fusion panel enables sequencing of over 700 unique fusion transcripts.

The panel covers relevant targets for acute myeloid leukaemia, myelodyplastic syndromes and myeoproliferative neoplasms, including CML, CMML and JMML. Among the targets are Calreticulin (CALR), JAK2, MPL and NPM1, which were previously offered as individual tests.

Requires patient informed consent.

### TAT: 3 days / 1 week

#### ALRP (DNA & RNA)

#### MPNS (DNA)

(3mL minimum) or bone marrow aspirate sample

## Leukaemia/Lymphoma RNA Sequencing (Fusion Gene and SNV/Indel) Panel

The Leukaemia / Lymphoma RNA Sequencing panel is an Anchored Multiplex PCR (AMP<sup>TM</sup>)- based next-generation sequencing (NGS) panel to detect and identify fusions, point mutations and expression levels from ribonucleic acid (RNA) input. The panel encompasses targets in over 199 genes relating to lymphoid and myeloid malignancies. By using gene-specific primers to amplify into molecular barcodes ligated onto the cDNA fragment ends, both known and novel fusions can be identified. Requires patient informed consent.

#### TAT: 2 weeks

#### PHFP



## Lysosomal Storage Disorders NGS Panel – full gene sequencing

This is a 55 gene custom NGS panel which can be used to detect both pathogenic SNP/ Indels and copy number variants (including whole exon insertions / deletions) which cause the various Lysosomal storage disorders.

All known lysosomal storage diseases are covered on this panel including:

Fabry disease, Gaucher disease, Pompe disease, metachromatic leukodystrophy, all the different mucopolysaccharidoses, fucosidosis, Krabbe disease, Tay-Sachs disease, Sandhoff disease, Danon disease, lysosomal acid lipase deficiency, Niemann-Pick disease types A, B and C, lipfuscinoses, prosaposin deficiency and Salla disease.

Requires patient informed consent.

#### TAT: 4-6 weeks

#### LSDS



## **Myeloid Gene Panel**

This is a 75 gene targeted NGS panel for acute myeloid leukaemia, myeloproliferative neoplasms, myelodysplastic syndromes, and also contains a number of targets which are useful for lymphoid malignancies (ALL and lymphoma). It uses Anchored Multiplex PCR (AMPTM) chemistry which enables deep strand-specific amplification of molecular barcoded DNA fragments for sequencing. Requires patient informed consent.

#### TAT: 2 weeks

#### **MVPS**

(3mL minimum) or bone marrow aspirate sample

# **SNP 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 karvotyping due to its greater precision and sensitivity. SNP (Single Nucleotide Polymorphism) arrays enable low-level mosaicism visualisation, loss of heterozygosity (LOH) and UPD detection, copy number change confirmation, triploidy detection and parent-of-origin analysis.

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).

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

## When to use SNP Array?

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

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

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 SNP Array detect?

Deletions and duplications with greater sensitivity than conventional karyotyping and loss of heterozygosity (LOH).

## What does SNP Array not detect?

- Balanced chromosome rearrangements such as translocations or inversions.
   The chromosome location of duplications (this would require additional FISH testing).
- Fragile X syndrome, genetic diseases caused by point mutations or multifactorial inheritance.

Prenatal SNP Array CGH	CGH	Amniotic fluid, CVS or POC 9	10 days	
TEST	CODE	SAMPLE REQS	TAT	
Blood from both parents may	also be prov	ided in case of follow up studies.		
Postnatal SNP Array CGH	CGH	<b>A (1)</b> 9	10 days	
TEST	CODE	SAMPLE REQS	TAT	

EDTA and heparin blood from both parents should be provided at the time of prenatal sampling, if possible, in case of follow up studies.

# Pan-ethnic carrier screening

The Fulgent Beacon carrier panel is a comprehensive genetic screen for people of all ethnic backgrounds. The panel analyses more than 400 genes, in which mutations may cause over 440 different recessive disorders. Testing includes Cystic Fibrosis, Sickle Cell Disease, Thalassemia and Spinal Muscular Atrophy. These conditions vary in morbidity, mortality and treatment.

The Beacon carrier screen can also be filtered to report only on diseases common to the Jewish population – such as Bloom Syndrome, Canavan Disease, Gaucher Syndrome and Tay-Sachs Disease.

#### Indications for use

- Pre-pregnancy screening for couples that wish to check if they are silent carriers for a disease that would have serious implications for the future health of any children.
- For patients who are concerned about a family history of a particular disease, where common mutation detections are very high – such as Tav-Sachs Disease.

The report comes with a synopsis of any diseases for which a mutation was found, including prognosis, treatment and mode of inheritance. It includes a risk assessment and recommendations for further testing.

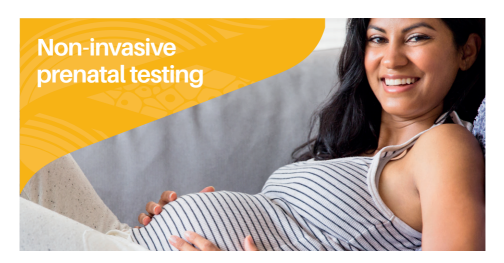
A full list of diseases covered by this test is available from the laboratory.

Male patients will not be screened for X-linked conditions. If an X-linked condition is suspected in a male patient please contact the laboratory or a genetics specialist about diagnostic testing for that particular condition.

### **Limitations**

A normal result does not rule out the possibility that the patient carries a rare mutation not detectible by this particular assay. For this reason, this test is also not appropriate to use as a direct prenatal screen (both parents must be confirmed carriers for a particular disease before we can offer prenatal diagnosis). Screening is not designed to detect somatic mutations.

TEST	CODE	SAMPLE REQS	TAT
Carrier Screen (Ashkenazi Jewish) Requires patient informed consent.	GENE	<b>A</b> 9	4 weeks
Carrier Screen (Ashkenazi Jewish)  — Partnered Report  Requires patient informed consent. Please contact the lab for special requirements before sending.	GENE	<b>A</b> 9	4 weeks
Carrier Screen (Pan-Ethnic) Requires patient informed consent.	GENE	<b>A</b> 9	4 weeks
Carrier Screen (Pan-Ethnic) – Partnered Report Requires patient informed consent. Please contact the lab for special requirements before sending.	GENE	<b>A</b> 9	4 weeks



# **Non-invasive prenatal testing (NIPT)**

Non-invasive prenatal testing (NIPT) screens for the presence of specific chromosome disorders in the developing fetus. The test analyses fragments of cell-free DNA in maternal plasma that have been released from both maternal and placental cells.

By analysing the proportions of cell-free DNA fragments derived from different chromosomes or chromosome regions, NIPT can screen for the presence or absence of specific chromosome disorders

NIPT is more accurate than first trimester maternal serum screening and ultrasound in identifying pregnancies with or without these disorders.

TDL Genetics uses the NIPT assay VeriSeq NIPT Solution v2, which is manufactured by Illumina and is processed at our laboratory in London.

# Targeted screening for specific common chromosome disorders

Our NIPT assay is designed to screen for:

 Trisomy 21 (Down syndrome), which is associated with moderate to severe intellectual disability, congenital heart defects and other malformations;

- Trisomy 18 (Edwards syndrome) and trisomy 13 (Patau syndrome), which are associated with severe brain and cardiac malformations. There is a high risk of stillbirth or death during infancy; and
- Sex chromosome aneuploidy (abnormalities in the number of X or Y chromosomes), which can be associated with malformations and infertility, Turner syndrome (45,X) and Klinefelter syndrome (47,XXY). Triple X syndrome and XYY syndrome can also be detected. This screen is optional (no additional cost).

In addition, NIPT can also assess fetal sex. This is optional (no additional cost).

NIPT does not screen for non-chromosome disorders, familial mutations, malformations, fetal growth or fetal viability.

## **Accuracy of NIPT**

NIPT provides fewer false-positive and false-negative results than combined first trimester screening for trisomy 21, 18 and 13.

It is important to note that NIPT is a screening test and does not provide a definitive genetic diagnosis, as NIPT cannot differentiate potential chromosome differences between the placenta and fetus.

A definitive genetic diagnosis of the fetus requires cytogenetic analysis of either amniotic fluid or chorionic villus sampling (CVS).

## When to perform NIPT

NIPT should not be performed before a gestational age of 10 weeks. However, it is suitable at any time after that, preferably while there is sufficient time for further investigation or decision-making (should this be required). An ultrasound scan is required prior to NIPT to confirm dates and fetal viability, and to check for twins. Performing first trimester screening before NIPT may provide supplementary information regarding the status of the fetus.

## Who is eligible for NIPT?

Eligible patients:

- Women who are at least ten weeks pregnant
- Women with singleton or twin pregnancies
- Women with IVF pregnancies and non-IVF pregnancies

NIPT is not suitable for patients with:

- Recent maternal blood transfusion (within the last 4 months)
- Maternal mosaicism
- Maternal prior organ transplant/stem cell transplant
- Maternal copy number variations
- Chromosomal copy number variations
- Fetoplacental mosaicism/confined placental mosaicism
- Maternal autoimmune disease excluding IVIg treatments
- Maternal neoplasms (benign and malignant)
- Pregnancies with fetal demise/vanishing twin

Patients with a twin pregnancy are not eligible for the sex chromosome aneuploidy component of the screen.

## Reporting results

Results will be ready within 2–4 business days upon receipt of sample in the laboratory.

TDL first checks that there is sufficient cell-free fetal DNA in the maternal sample and quality data to provide an accurate assessment. A re-collection may be recommended if the sample is not suitable or an assessment may not be feasible.

The report then summarises the screening assessment for each disorder specified by the requesting doctor (see example below).

#### **Example report**

Chromosome	Result	Recommendation
Trisomy 21	HIGH Probability	Genetic counselling and additional testing
Trisomy 18	Low probability <1:10,000	Review result with patient
Trisomy 13	Low probability <1:10,000	Review result with patient
Sex chromosome aneuploidy	Not requested	
Fetal sex	Male	Review result with patient

A high probability NIPT result should always be confirmed by amniocentesis or CVS before making any decision regarding subsequent management of the pregnancy.

#### **Limitations of NIPT**

The VeriSeq NIPT Solution v2 is not validated for use in pregnancies with more than two fetuses, fetal demise, mosaicism, partial chromosome aneuploidy, triploidy, translocations, maternal aneuploidy, transplant or malignancy. VeriSeq NIPT Solution v2 does not detect neural tube defects. Certain rare biological conditions may also affect the accuracy of the test.

For twin pregnancies, HIGH PROBABILITY test results apply to at least one fetus; male test results apply to one or both fetuses; female test results apply to both fetuses. Due to the limitations of the test, inaccurate results are possible.

A LOW PROBABILITY result does not guarantee that a fetus is unaffected by a chromosomal or genetic condition. Some non-aneuploid fetuses may have HIGH PROBABILITY results. In cases of HIGH PROBABILITY results and/or other clinical indications of a chromosomal condition, confirmatory testing is necessary for diagnosis.

## If an assessment cannot be provided

On rare occasions, NIPT is unable to provide an assessment of the probability of specific chromosome disorders. This usually reflects the complex biology of genetics and pregnancy, and is not due to a failing in the laboratory.

If NIPT cannot provide a specific assessment after a repeat blood draw, it is not worth repeating the NIPT (unless advised by the laboratory). A decision about other tests (maternal serum screening, detailed ultrasound, amniocentesis or CVS) should be based on the doctor's assessment of all risk factors identified, and may require specialist consultation.

#### **Further information**

- TDL Genetics website: www.tdlpathology.com/tdlgenetics
- Borth H, et al. Analysis of cell-free DNA in a consecutive series of 13,607 routine cases for the detection of fetal chromosomal aneuploidies in a single center in Germany. Arch Gynecol Obstet. 2021 Jun;303(6):1407-1414.



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Find out more about NIPT on the TDL website:

www.tdlpathology.com/noninvasive-prenatal-testing/

2-4 days

TEST	CODE	SAMPLE REQS	TAT

NIPT

Non-Invasive Prenatal Testing (NIPT)

- common aneuploidy screening
from maternal blood

J / Special tube<sup>1</sup>

# **In-vivo** Tests

All *in-vivo* tests (except Glucose Challenge Test/Mini-GTT) require an appointment. Please email **phlebotomy@tdlpathology.com** or call **020 7307 7373** for details, information for patient preparation, and appointment times. Sample taking fees for Extended tests are charged at £90.00 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	TAT	COLLECTION TIME (MINUTES POST-GLUCOSE)
Glucose Challenge Test/Mini-GTT	RBGM	G	1 day	1 at 60 mins (50gm glucose)
Glucose Tolerance Test (Extended Plus)	GTTX	7 x <b>(</b> , 7 x <b>RU</b>	1 day	1 each at 0, 30, 60, 90, 120, 150 and 180 mins.
Glucose Tolerance Test (Extended)	GTTE	5 x 🕒 , 5 x RU	1 day	1 each at 0, 30, 60, 90 and 120 mins.
Glucose Tolerance Test (Short)	GTTS	2 x 🕝 , 2 x RU	1 day	1 each at 0 and 120 mins.
Glucose Tolerance Test/OGTT	GTT	3 x <b>G</b> , 3 x <b>RU</b>	1 day	1 each at 0, 60 and 120 mins (75gm glucose load)
Glucose Tolerance with Growth Hormone	GTT + GHDF	3 x B <sup>35</sup> , 3 x <b>G</b> , 3 x <b>RU</b>	1 day	1 each at 0, 60 and 120 mins.
Glucose Tolerance with Insulin	GTTI	3 x B , 3 x 🕝 , 3 x RU	1 day	1 each at 0, 60 and 120 mins

## **Extended tests**

TEST	CODE	SAMPLE REQS	TAT	COLLECTION TIME (MINUTES POST-GLUCOSE)
Lactose Tolerance Test	LTT	By appointment only	1 day	Contact 0207 307 7383 (Phlebotomy)
Synacthen Stimulation Test	SYNA	By appointment only	1 day	

# **Antibiotic assays**

CODE	SAMPLE REQS	TAT
AMIK	<b>B</b> 4	1 day
GENT	<b>B</b> 4	1 day
METR	<b>B</b> 4	10 days
TEIC	В	5 days
TOBR	0	3 days
VANC	B	1 day
	AMIK GENT METR TEIC TOBR	AMIK 3 4  GENT 3 4  METR 3 4  TEIC 3  TOBR 1

# **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 REQS	TAT
Amitriptyline	AMTR	<b>A</b> 4	5 days
Anafranil (Clomipramine)	CHLO	A	7 days
Carbamazepine (Tegretol)	CARB	B	1 day
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	1 day
Epanutin (Phenytoin)	PHEN	B	1 day
Erythropoietin	ERY	B	4 days
Ethosuximide	ETH0	A	7 days
FK506 (Tacrolimus/Prograf)	FK5	<b>A</b> 4	1-2 days
Flecainide (Tambocor)	FLEC	A	5 days
Fluoxetine (Prozac)	PROZ	<b>A</b> 4	5 days
Gabapentin	GABA	<b>B</b> 4	5 days
Imipramine	IMIP	<b>A</b> 4	4 days
Lamotrigine	LAM0	<b>B</b> 4	5 days
Levetiracetam (Keppra)	LEVE	<b>B</b> 4	3 days
Lithium (take 12 hours after dose)	LITH	B	1 day
Lorazepam	LORA	<b>A</b> 4	10 days
Methotrexate	METX	B	2 days
Mycophenolic Acid (Cellcept)	MYCP	A	5 days
Mysoline (Primidone)	PRIM	<b>B</b> 4	3 days

# **Therapeutic Drug Assays**

TEST	CODE	SAMPLE REQS	TAT
Olanzapine	OLAN	<b>A</b> 4	5 days
Paracetamol	PARA	B	1 day
Phenobarbitone	PHB	B	1 day
Phenytoin (Epanutin)	PHEN	B	1 day
Primidone (Mysoline)	PRIM	<b>B</b> 4	3 days
Propanalol	PR0	<b>B</b> 4	7 days
Risperidone	RISP	<b>A</b> 4	7 days
Sinequan (Doxepin)	DOXE	A or 🕕	10 days
Sirolimus	SIR0	A	3 days
Streptomycin Levels	STRM	<b>(</b>	5 days
Sulpiride	SULP	<b>B</b> 4	4 days
Tacrolimus/Prograf (FK506)	FK5	<b>A</b> 4	1-2 days
Tegretol (Carbamazepine)	CARB	B	1 day
Temazepam	TEMA	<b>B</b> 4	4 days
Theophylline	THE0	B	1 day
Topiramate (Topamax)	TOPI	<b>B</b> 4	4 days
Trimipramine	TRIM	A	5 days
Valium (Diazepam)	DIAZ	A	7 days
Valproic Acid (Epilim)	VALP	B	1 day
Vigabatrin (Sabril)	VIGA	A	10 days

# **Allergy**

For a list of individual allergens see page 147.

TEST	CODE	SAMPLE REQS	TAT
Allergy – Individual Allergens	ALLE	B	2 days
Allergy – 5 x Single Individual Allergens	5AL	B	2 days
Allergy – 10 x Single Individual Allergens	10AL	B	2 days
Allergy Profile 1 (Food & Inhalants)	1A	BB	2 days
Allergy Profile 2 (UK Aero Allergen)	2A	BB	2 days
Allergy Profile 3 (Food)	3A	BB	2 days
Allergy Profile 4 (Nuts & Seeds)	4A	BB	2 days
Allergy Profile 5 (Children's Panel)	5A	BB	2 days
Allergy Profile 6 (Shellfish)	6A	BB	2 days
Allergy Profile 7 (Finfish)	7A	BB	2 days
Allergy Profile 8 (Cereal – singles)	8A	BB	2 days
Allergy Profile 9 (Antibiotics)	9A	BB	2 days
Allergy Profile 10 (Insects)	10A	BB	2 days
Allergy Profile 11 (Combined Shellfish/Finfish)	11A	BB	2 days
Allergy Profile 12 (Milk & Milk Proteins)	12A	BB	2 days
Allergy Profile 13 (Stone fruit/Rosaceae family)	13A	BB	2 days
ALEX <sup>2</sup> Allergy Test (Self-collect)	ALEX	(TDL Tiny)	3-4 days
ALEX <sup>2</sup> Allergy Test (Venous)	ALEX	(Serum)	3-4 days
ISAC Panel (Self-collect)	ISAC	(TDL Tiny)	3 days
ISAC Panel (Venous)	ISAC	В	3 days
Allergic Rhinitis/Asthma Profile	ALRN	BB	2 days
Atopic Dermatitis/Eczema Profile (14 allergens)	ALEC	BB	2 days
Gluten Sensitivity Profile	GLUT	ABB	10 days
Histamine Releasing Urticaria Test	CURT	В	3 weeks
Prealbumin	PALB	B	3 days
Total IgE	IGE	B	1 day
Tryptase	STRY	В	2 days

## **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**.



SCAN MF

Find out more details about component testing:

www.tdlpathology.com/ specialties/allergy/ allergy-component-testing/

	CODE	SAMPLE REQS	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
Glycan Determinants	ZZ27	B	2 days
Hazelnut Components	ZZ11	B	2 days
Horse Components	ZZ38	B	2 days
House Dust Mite Components	ZZ12	B	2 days
Kiwi Components	ZZ32	B	2 days
Latex Components	ZZ13	B	2 days
Lipid Transfer Proteins	ZZ23	B	2 days
Lipocalins	ZZ28	B	2 days
Olive Components	ZZ14	B	2 days
Parvalbumins	ZZ29	B	2 days
Peach Components	ZZ15	B	2 days
Peanut Components	ZZ16	B	2 days

CODE	SAMPLE REQS	TAT
ZZ25	B	2 days
ZZ22	B	2 days
ZZ24	В	2 days
ZZ26	B	2 days
ZZ30	B	2 days
ZZ39	В	2 days
ZZ17	B	2 days
ZZ18	B	2 days
ZZ19	В	2 days
ZZ31	B	2 days
ZZ33	B	2 days
ZZ20	В	2 days
ZZ34	В	2 days
ZZ21	B	2 days
	ZZ25 ZZ22 ZZ24 ZZ26 ZZ30 ZZ39 ZZ17 ZZ18 ZZ19 ZZ31 ZZ33 ZZ33 ZZ34	ZZ25 3 ZZ22 3 ZZ24 3 ZZ26 3 ZZ30 3 ZZ39 3 ZZ17 3 ZZ18 3 ZZ19 3 ZZ18 3 ZZ19 3 ZZ31 3 ZZ33 3 ZZ33 3 ZZ33 3

## **Allergy Profile 1 (Food & 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), Hazelnut, Horse dander, Latex, Nettle, Peanut, Shrimp/Prawn, Soya bean, Wheat

#### TAT: 2 days

1A



## Allergy Profile 2 (UK Aero Allergen)

Total IgE with individual IgE allergens for:

Alternaria Derma farinae
Aspergillus Dog dander
Birch pollen House dust mite
Cat dander Horse dander
Cladosporium Timothy grass
Common ragweed

TAT: 2 days

2A

BB

### **Allergy Profile 3 (Food)**

Total IgE with individual IgE allergens for:

Codfish Egg yolk Sesame
Cow's milk Kiwi Soya
Egg white Peanut Wheat

TAT: 2 days

3A



## Allergy Profile 4 (Nuts & Seeds)

Total IgE with individual IgE allergens for:

Almond Peanut Pumpkin seed
Brazil nut Pecan Sesame seed
Cashew Pine nut Sunflower seed
Hazelnut Pistachio Walnut
Macadamia nut Poppy seed

TAT: 2 days

4A



## Allergy Profile 5 (Children's Panel)

**Total IgE** with individual IgE allergens for:

Cat dander Egg white Soya bean
Cod Egg yolk Timothy grass
Cow's milk Hazelnut Wheat flour
Dog dander Peanut
Dust mite Silver birch

TAT: 2 days

5A



#### **Alleray**

## **Allergy Profile 6 (Shellfish)**

Total IgE with individual IgE allergens for:

Clam Lobster Scallop Crab Octopus Sauid

Crawfish/Crayfish Prawns/Shrimp

TAT: 2 days

6A

**B B** 

### Allergy Profile 7 (Finfish)

Total IgE with individual IgE allergens for:

Codfish Sardine/Pilchard Swordfish Mackerel Salmon Tuna Plaice Sole

TAT: 2 days

7A



## Allergy Profile 8 (Cereal - singles)

Total IgE with individual IgE allergens for:

Barley Rve Wheat 0at

TAT: 2 days

8A



### **Allergy Profile 9 (Antibiotics)**

Total IgE with individual IgE allergens for:

Amoxicillovl Pen G Pen V AmpicillovI Cefaclor

TAT: 2 days

9A

**BB** 

## **Allergy Profile 10 (Insects)**

Total IgE with individual IgE allergens for:

Common wasp -Paper wasp yellow jacket Yellow hornet Ree White faced hornet

TAT: 2 days

10A

BB

## **Allergy Profile 11** (Combined Shellfish/Finfish)

Total IgE with individual IgE allergens for:

Scallop Squid Prawn/Shrimp Salmon Tuna

TAT: 2 days

11A

BB

## Allergy Profile 12 (Milk & Milk Proteins)

Total IgE with individual IgE allergens for:

Alpha-lactalbumin -Goat's milk milk proteins Mare's milk Beta-lactoglobulin -Sheep's milk milk proteins Whey (cow and ewe) Casein - milk proteins

Cow's milk

TAT: 2 days

12A

BB

# Allergy Profile 13 (Stone fruit/Rosaceae family)

Total IgE with individual IgE allergens for:

Almond Pear
Apple Plum
Apricot Raspberry
Cherry Strawberry

Peach

TAT: 2 days

13A

**B B** 

## **ALEX<sup>2</sup> Allergy Test (Venous)**

ALEX® Allergy Explorer rapidly tests for up to 300 allergens simultaneously and providing a comprehensive analysis, from a single sample (0.5µl). The panel of allergens includes pollen, mites, cat and dog fur, insect venoms, moulds and yeasts, food and latex, supplemented with total IgE.

TAT: 3-4 days

ALEX

(Serum)

## **ISAC Panel (Venous)**

Simultaneous measurement from single sample (0.5µl) of specific antibodies to more than one hundred allergen components from more than 48 preselected allergen sources.

TAT: 3 days

**ISAC** 

B

### Allergic Rhinitis/Asthma Profile

Total IgE with individual IgE allergens for:

Cat dander Aspergillius fumigatus
Dog dander Cladosporium herbarum
Common silver birch Mugwort
Timothy grass London plane

Dust mite - Peanut
Dermatophagoides Egg white
pteronyssinus Cow's milk
Alternaria alternata

TAT: 2 days

ALRN

 $\mathbf{B}\mathbf{B}$ 

# Atopic Dermatitis/Eczema Profile (14 allergens)

TOTAL IGE with individual IgE allergens for:

Cod fish Apple Cow's milk Dust mite -Egg white dermatophagoides Soyabean pteronyssinus Peanut Cat dander Hazelnut Dog dander Shrimp Timothy grass Common silver birch Wheat

TAT: 2 days

ALEC



### **Gluten Sensitivity Profile**

Gluten Single IgE Allergen Immunoglobulin A
Gliadin Antibodies (IgG) Tissue Transglutaminase
(deamidated) IgA (Coeliac)

HLA Tissue Typing Coeliac
Disease – D02/D08

TAT: 10 days

**GLUT** 



## **Individual allergens**

Allergens, when requested individually are priced as single tests, sample 1 x 3 (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 q12

Cultivated wheat q15

Johnson grass g10

Maize, Corn q202

Meadow fescue q4

Meadow foxtail q16

Meadow grass, Kentucky blue g8

Redtop, Bentgrass q9

Rye-grass q5

Sweet vernal grass q1

Timothy grass q6

Velvet grass g13

Wild rye grass q70

**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

Elder t205

Flm 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

0ak t7

Oil Palm t223

Olive t9

Pecan, Hickory t22

Peppertree t217

Pine t213

Privet t210

Privet t21

Queen palm t72

Red cedar t57

Red mulberry t71

Scotch broom t55

Spruce t201

Sweet aum 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

Asperaillus niaer 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

Setomelanomma rostrata

(Helminthosporium halodes) m8

Malassezia spp. m227

Mucor racemosus m4

Penicillium chrysogenum (P. notatum) m1 Penicillium glabrum m209 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) m<sub>10</sub>

Tilletia tritici m201 Trichoderma viride m15 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, serum proteins e219 Chinchilla epithelium e208

Cow dander e4 Dog dander e5

Chicken feathers e85

Duck feathers e86 Ferret epithelium e217

Finch feathers e214 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

Parrot feathers e213

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

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

Lepidoalyphus destructor (Storage mite) d71

Tvrophagus putrescentiae (Storage mite) d72

#### ALLERGEN COMPONENTS

See page 142 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 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

Amoxicillovl c6 Ampicilloyl c5

Cefaclor c7

Chlorhexidine c8

Gelatin bovine c74

Insulin human c73

PenicillovI G c1

Penicilloyl V c2

Pholcodine c261

Morphine c260

Suxamethonium (succinylcholine)

c202

#### OCCUPATIONAL

Bougainvillea k214

Cotton seed k83

Ethylene oxide k78

Figus k81

Formaldehyde/Formalin k80

Hexahydrophtalic anhydrid k209

Isocyanate HDI (Hexamethylene

diisocyanate) k77

Isocyanate MDI (Diphenylmethane

diisocvanate) k76

Isocyanate TDI (Toluene diisocyanate) k75

Ispaghula k72

Latex k82

Methyltetrahydrophtalic anhydrid k211

Phthalic anhydride k79

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

Grane f259

Grapefruit f209

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

Spinach f214

Strawberry f44

Sweet potato f54

Tomato f25

Watermelon f329

FOODS - SEED, LEGUMES & NUTS

Almond f20

Barley f6

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

iviacauaiiiia iiut

Maize, Corn f8

0at 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

Rve f5

Sesame seed f10

Sovbean f14

Spelt wheat f124

Sugar-beet seed f227

Sweet chestnut f299

Walnut f256

Wheat f4

White bean f15

FOODS - SPICES

Anise f271

Basil f269

Bay leaf f278

Black pepper f280

Caraway f265

Chilipepper f279

Clove f268

Coriander f317

Coriandei

Dill f277

Ginger f270

Green pepper (unripe seed) f263

Lovage f275

Mace f266

Marioram f274

Mint f332

Mustard f89

Oregano f283

Paprika. Sweet pepper f218

Parslev f86

Tarragon f272

Thyme f273

Vanilla f234

FOODS - FISH, SHELLFISH

& MOLLUSCS

Abalone f346 Anchovy f313

Blue mussel f37

Cat fish f369

Chub mackerel f50 Clam f207

Crab f23

Cravfish f320

Fish (cod) f3

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

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

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 volk f75

Turkey meat f284

#### FOODS - MEAT

Beef f27

Mutton f88

Pork f26

Rabbit f213

#### FOODS - MILK

Casein f78

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

Cochineal extract (Carmine red) (E120)

f340

#### **FOODS - MISCELLANEOUS**

Cacao f93

Coffee f221

Malt f90

Mushroom (champignon) f212

Tea f222

Yeast f45

## **Vitamins**

TEST	CODE	SAMPLE REQS	TAT
1,25 Vitamin D	D3	<b>B</b> *	5-8 days
*Serum sample stable for 3 days ambient.			
Beta Carotene	CAR0	В	5 days
Biotin	BIOS	<b>B</b> 7	5 days
Carotenes	CARO	В	5 days
Vitamin A (Retinol)	VITA	B	5 days
Vitamin B (Functional)	FUNC	A A or H	5 days
Vitamin B Profile	VBP	AAB	5 days
Vitamin B1 (Thiamine)	VIT1	A	5 days
Vitamin B2 (Riboflavin)	VIB2	A	5 days
Vitamin B3 (Nicotinamide)	VIB3	B	5 days
Vitamin B5 (Pantothenic Acid)	VB5S	В	5 days
Vitamin B6 (Pyridoxine)	VITB	A	5 days
Vitamin B7 (Biotin)	BIOS	<b>B</b> 7	5 days
Vitamin B9 (Folic acid) – Red cell	RBCF	A	2 days
Vitamin B9 (Folic acid) – Serum	F0LA	B	1 day
Vitamin B12 (Active) (Self-collect)	B12	(TDL Tiny)	1 day
Vitamin B12 (Active) (Venous)	B12	B	1 day
Vitamin B12 (Active)/Red Cell Folate	B12F	AB	2 days
Vitamin C (Active) *Serum should be separated and frozen within 3 hours of venepuncture.	VITC	(spun and frozen within 3 hours)*	5 days
Vitamin D (1, 25 Dihydroxy) *Serum sample stable for 3 days ambient.	D3	<b>B</b> *	5-8 days
Vitamin D (25-OH) (Self-collect)	VITD	(TDL Tiny)	1 day
Vitamin D (25-OH) (Venous)	VITD	<b>B</b>	1 day
Vitamin E (Alpha Tocopherol)	VITE	B	5 days
Vitamin K (Nutritional)  * Sample should be light protected after collection, spun/separated and frozen within 24 hours of collection.	VKN	Serum (SST or B ) *	5 days
Vitamin Profile 1	VITS	<b>A B B</b> <sup>7</sup>	5 days
Vitamin Profile 2	VIT2	<b>A A A</b> 38 <b>B B</b> 7,13	5 days

## **Nutrition and lifestyle**

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.

TEST	CODE	SAMPLE REQS	TAT
Caeruloplasmin	CERU	<b>B</b>	1 day
Copper (Serum)	COPP	3 or 🚯	5 days
Essential Fatty Acid Profile (Red Cell)	EFAR	<b>A</b> 4	10 days
Folate (Red Cell)	RBCF	A	2 days
Magnesium (Whole blood)	RCMG	A or (1)	4 days
Mineral Screen	MINE	<b>B (3</b>	5 days
Mineral Screen (Whole blood)	RMIN	00	5 days
Mineral Screen and Industrial Heavy Metal Screen (Trace Metals)	TRAC		7-10 days
Omega 3/Omega 6 (Self-collect)	OMG3	(TDL Tiny)	5 days
Omega 3/Omega 6 (Venous)	OMG3	<b>A</b> 4	5 days
Selenium (Serum) (Self-collect)	SELE	(TDL Tiny)	4 days
Selenium (Serum) (Venous)	SELE	B	4 days
Sports/Performance Profile	SPOR		5 days
Zinc (Serum)	ZINC	ß	2 days
Zinc (Urine)	URZN	CU	5 days
Zinc (Whole Blood)	RBCZ	A or (1)	5 days

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

#### **Mineral Screen**

Calcium Magnesium

Zinc

Iron

Copper Chromium

Manganese

TAT: 5 days

MINE



## **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



## Mineral Screen and **Industrial Heavy Metal** Screen (Trace Metals)

Aluminium Manganese

Iron

Calcium

7inc

Magnesium

Copper

Cadmium

Mercury

Lead

Chromium

**TAT: 7-10 days** 

TRAC



## **Sports/Performance Profile**

Full Blood Count (FBC)

Biochemistry Profile

HDL/LDL Cholesterol (Calculated)

Ferritin

C Reactive Protein (CRP)

Omega 3/Omega 6

Mineral Screen

Vitamin B9 (Folic acid) - Red cell Vitamin B12 (Active)

TAT: 5 days

SP0R









#### Vitamin B Profile

Vitamin B1 (Thiamine)

Vitamin B2 (Riboflavin) Vitamin B3 (Nicotinamide)

Vitamin B6 (Pyridoxine)

Vitamin B9 (Folic acid) - Red cell Vitamin B12 (Active)

TAT: 5 days

**VBP** 



#### Vitamin Profile 1

Vitamin A (Retinol)

Beta Carotene

Vitamin B1 (Thiamine)

Vitamin B2 (Riboflavin)

Vitamin B6 (Pyridoxine)

Vitamin D (25-OH)

Vitamin E (Alpha

Tocopherol)

TAT: 5 days

VITS



#### Vitamin Profile 2

Vitamin A (Retinol)

Beta Carotene

Vitamin B1 (Thiamine)

Vitamin B2 (Riboflavin)

Vitamin B3 (Nicotinamide)

Vitamin B6 (Pyridoxine)

Vitamin B9 (Folic acid) - Red cell

Vitamin B12 (Active)

Vitamin D (25-OH)

Vitamin E (Alpha Tocopherol)

TAT: 5 days

VIT2



# 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.

TEST	CODE	SAMPLE REQS	TAT	
Omega 3/Omega 6 (Self-collect)	OMG3	(TDL Tiny)	5 days	
Omega 3/Omega 6 (Venous)	OMG3	<b>A</b> 4	5 days	



The TDL Self-Collect range of testing has been gathering increased and important attention for healthcare services. Self-collection is being adopted across different target areas of healthcare: sexual health screening, wellness testing, genetic conditions, lifestyle review, pre-operative work ups, etc. Self-collection sampling allows patients to collect samples at home, and together with Royal Mail Tracked postal returns, facilitates safe and effective delivery of samples throughout the UK to the laboratory.

As part of the ongoing development of the TDL self-collection service, the sample collection kits ensure that TDL is aligned to regulatory requirement around ISO:13485 kit manufacture and UKCA marking across the UK. This requirement for UKCA marked kits also addresses the need for clinically approved stability and comparative performance. The interest being generated for this service ensures best attention, continued development, with regular review of the repertoire, and where possible more tests will be added to the available Self-Collection list.

The TDL Self-Collect capillary blood and sample kits include a helpful range of diagnostic and screening tests. These sample collection kits are not home test kits that provide an immediate result for the patient. Samples collected at home are all returned to the laboratory for testing, using Royal Mail Tracked 24 postal services. Results are returned directly to the healthcare organisation, doctor or healthcare

professional — not to the patient. Self-Collect kits need to be UKCA marked (or dual marked with UKCA/CE) to represent the product claims that kits are being used for the collection of samples in a non-clinic setting.

The Self-Collect kit itself allows for combinations of sample types (urine, stool, swabs) — and the range of UKCA marked kits are listed on page 157. Instructions for sample collection are enclosed in each collection kit. The best results are obtained by patients who closely follow the instructions that are provided, and by collecting enough blood drops to sufficiently fill the microtainer tube(s) in their kit. A request form or specially provided tube label must be returned with the collected sample. It is exciting that the scope of this service, together with its performance and quality standards, will be revised, developed, and updated on a regular basis.

For more details relating to this service, please email **UKCAkits@tdlpathology.com** 



Find out more information about the TDL Self-Collect kits:

www.tdlpathology.com/ self-collect-kits/

#### Quality is key

- Components: verified for the specific intended use of the kit and linked to the accredited tests performed in the laboratory.
- Instructions: monitored for ease of use, version controlled, with regular feedback for ongoing improvement.
- Quality: Management of technical files, regulatory submissions and manufactured to the required ISO:13485 medical device manufacturing standards.
- Supply: Assembled within the UK. Both individual kit fulfilment services and larger size kit orders are available.
- **Security**: Test kits are security sealed.
- Accompanying information: Request forms cannot be inserted into the sealed kits.
   An accompanying envelope (TDL will provide) or other clearly visible method must be sent with each kit to clearly display the request form.
- Laboratory testing: Verified diagnostic tests performed in an ISO:15189 accredited clinical laboratory

We recommend that all healthcare organisations and healthcare professionals using our TDL Tiny™ and TDL Self-Collect kits are up to date with latest diagnostic testing guidelines and relevant updates, including but not limited to those published by:

- UKHSA Standards for Microbiology Investigations (SMI)
- British Association of Sexual Health and HIV Guidelines (BASHH)
- Royal College of Obstetrics and Gynaecology Guidelines (RCOG)
- NICE Evidence-based recommendations on faecal immunochemical tests (DG30)
- British Society of Haematology Evidence Based Guidance (BSH)
- Association of Clinical Biochemistry (ACB)

## **TDL's range of kits**

Respi	ratory virus PCR	
KIT CODE	KIT TYPE	SAMPLE TYPE
KT293	Respiratory Virus Swab Collection Kit (2mL)	Oropharyngeal and Nasal swab
Capill	ary blood	
KIT CODE	KIT TYPE	SAMPLE TYPE
KT353	Capillary Blood Collection Kit (SST)	Capillary blood (SST)
KT354	Capillary Blood Collection Kit (EDTA)	Capillary blood (EDTA)
KT466	Capillary Blood Collection Kit (Plain Red)	Capillary blood (Plain)
KT384	Capillary Blood Collection Kit (SST x2)	Capillary blood (SST x 2)
KT355	Capillary Blood Collection Kit (SST and EDTA)	Capillary blood (SST and EDTA)
KT423	Capillary Blood Collection Kit (SST x2 and EDTA)	Capillary blood (SST x 2 and EDTA)
KT445	Capillary Blood Collection Kit (Plain Red and SST)	Capillary blood (Plain and SST)
Sexua	il health	
KIT CODE	KITTYPE	SAMPLE TYPE
KT356	Sexual Health Collection Kit (Urine)	Aptima urine
KT357	Sexual Health Collection Kit (Vaginal)	Aptima multisite swab
KT358	Sexual Health Collection Kit (Blood and Vaginal)	Capillary blood and Aptima multisite swab
KT359	Sexual Health Collection Kit (Throat and Rectal)	Aptima multisite swab x2
KT360	Sexual Health Collection Kit (Blood and Urine)	Capillary blood and Aptima urine
KT361	Sexual Health Collection Kit (Blood, Urine, Throat and Rectal – MSM)	Capillary blood, Aptima urine and Aptima multisite swab x 2
KT424	Sexual Health Collection Kit (Blood, Vaginal, Throat and Rectal)	Capillary blood and Aptima multisite swab x3
KT404	Sexual Health Collection Kit (Oral lesion)	Oral swab
KT405	Sexual Health Collection Kit (Genital lesion)	Genital swab
KT421	Sexual Health Collection Kit (Urine, Throat and Rectal – MSM)	Aptima urine and multisite swab x 2
KT425	Sexual Health Collection Kit (Throat)	Aptima multisite swab
KT426	Sexual Health Collection Kit (Rectal)	Aptima multisite swab
KT428	Sexual Health Collection Kit (Vaginal, Throat and Rectal)	Aptima multisite swab x3

KT429	Sexual Health Collection Kit	Capillary blood, Aptima urine
	(Blood, Urine, Vaginal, Throat and Rectal)	and Aptima multisite swab x3

Microbial/Viral screening						
KIT CODE	KIT TYPE	SAMPLE TYPE				
KT364	HPV Swab Collection Kit	Qvintip swab				
KT365	MRSA Collection Kit (Nose and Groin)	Purple liquid Amies swab x 2				
KT422	MRSA Collection Kit (Nose, Groin and Axilla)	Purple liquid Amies swab x3				
KT366	GBS Collection Kit (Vaginal and Rectal)	Blue gel Amies swab x2				
KT441	Vaginitis Collection Kit (Vaginal – Culture and PCR)	Aptima multisite swab and Blue gel Amies swab				
KT385	Urinalysis Collection Kit (Chemistry and Microscopy)	Urine (Universal)				
KT386	Urinalysis Collection Kit (Chemistry, Microscopy and Culture)	Urine (Universal and Boric)				

Gastrointestinal						
KIT CODE	KIT TYPE	SAMPLE TYPE				
KT362	QFIT Collection Kit	QFIT sample collection device				
KT363	Faecal Collection Kit	Stool/faecal container				
KT430	Faecal Collection Kit (QFIT and Stool x2)	QFIT sample collection device and stool/faecal container x2				

Please post self-collected samples on the same day they are taken, avoid posting over weekends and bank holidays.

TEST	CODE	SAMPLE REQS	TAT	KIT CODE
7 STI Profile by PCR (7 tests from 1 Sample)	DL12	Aptima urine or multisite swab	2 days	KT356 or KT357
ALEX <sup>2</sup> Allergy Test 300 allergen panel, from single sample (0.5µl), supply Total IgE, which includes pollens, mites, cat and fur, insect venoms, moulds, yeasts, food and latex.		(TDL Tiny)	3-4 days	KT353
Amenorrhoea Profile (LH, FSH, PROL, TEST, TOES, SHBG, FAI) CHANGE LH, FSH, PROL, TEST, TOES and SHBG now also avindividually from a (5) (TDL Tiny) sample. Avoid taki samples from any area an HRT cream is applied.		(TDL Tiny)	1 day	KT466
Amylase	AMY	(TDL Tiny)	1 day	KT353
Antimullerian Hormone (AMH)	АМН	B (TDL Tiny) or (TDL Tiny)	1 day	KT353 or KT466

TEST	CODE	SAMPLE REQS	TAT	KIT CODE
Apolipoprotein A1 NEW	AP0A	(TDL Tiny)	3 days	KT353
Apolipoprotein B NEW	AP0B	(TDL Tiny)	3 days	KT353
C Reactive Protein	CRP	(TDL Tiny)	1 day	KT353
C Reactive Protein (High Sensitivity)	HCRP	(TDL Tiny)	1 day	KT353
CA 125	C125	(TDL Tiny)	1 day	KT353
Calprotectin	CALP	<b>QFIT</b> sample collection device	5 days	KT362 or KT430
Calprotectin/QFIT Profile (Combined)	QCAL	<b>QFIT</b> sample collection device	5 days	KT362 or KT430
Carbohydrate Deficient Transferrin (CDT)	CDT	(TDL Tiny)	3 days	KT353
Chlamydia/Gonorrhoea – Rectal	RSCG	Aptima multisite swab	2 days	KT426
Chlamydia/Gonorrhoea – Throat	TSCG	Aptima multisite swab	2 days	KT425
Chlamydia/Gonorrhoea – Urine	CCG	Aptima urine	2 days	KT356
Chlamydia/Gonorrhoea – Vaginal	SCG	Aptima multisite swab	2 days	KT357
Cortisol	CORT	(TDL Tiny)	1 day	KT353
COVID-19 (SARS-CoV-2) RNA by PCR Contact Laboratory.	NCOV	Aptima multisite swab of nose/throat	1 day	KT293
COVID-19 (SPIKE) Antibodies	SCOV	(TDL Tiny)	1 day	KT353
Creatinine (including eGFR)	CREA	(TDL Tiny)	1 day	KT353
DHEA Sulphate	DHEA	(TDL Tiny) or (TDL Tiny)	1 day	KT353 or KT466
DL12 7 STI Profile by PCR (7 PCR tests from 1 Sample)	DL12	Aptima urine or multisite swab	2 days	KT356 or KT357
Elastase	ELAS	Stool/faecal container	5 days	KT363
Enteric Organism Rapid Detection	EORD	Stool/faecal container	2 days	KT363 or KT430
Female Hormone Profile (LH, FSH, PROL, TOES) Avoid taking samples from any area an HRT cream is	TFIP applied.	(TDL Tiny) and (TDL Tiny)	1 day	KT445
Ferritin	FERR	(TDL Tiny)	1 day	KT353
Free T3	FT3	(TDL Tiny)	1 day	KT353
Free T4	FT4	(TDL Tiny)	1 day	KT353
FSH	FSH	(TDL Tiny) or (TDL Tiny)	1 day	KT353 or KT466
Full Blood Count* (Haemoglobin, White Cell Count, Red Cell Count, Platelets) NEW *Mix sample on collection.	TFBC	(TDL Tiny)	1 day	KT354
Gastrointestinal Pathogen Profile by PCR	EORD	Stool/faecal container	2 days	KT363 or KT430

TEST	CODE	SAMPLE REQS	TAT	KIT CODE
Gliadin Antibodies (IgG) (deamidated)	AGAB	(TDL Tiny)	2 days	KT353
Group B Strep – Vaginal and Rectal	GBSX	Blue gel Amies swab x2	3-5 days	KT366
H. pylori Antigen – Stool	HBAG	Stool/faecal container	3 days	KT363 or KT430
HbA1c	GHB	(TDL Tiny)	1 day	KT354
HCG (Quantitative)	QHCG	(TDL Tiny)	1 day	KT353
Hepatitis B Immunity (IgG)	THBI	(TDL Tiny)	1 day	KT353
Hepatitis B Surface Antigen	THBA	(TDL Tiny)	1 day	KT353
Hepatitis C Antibodies	THCV	(TDL Tiny)	1 day	KT353
Hepatitis C Antigen (Early detection)	TCAG	(TDL Tiny)	1 day	KT353
Herpes Simplex (HSV) 1 & 2 – Genital lesion	HERS	Aptima multisite swab	5 days	KT405
Herpes Simplex (HSV) 1 & 2 – Oral lesion	HERS	Aptima multisite swab	5 days	KT404
HIV 1 & 2 Abs/p24Ag	THIV	(TDL Tiny)	1 day	KT353
HPV (19 high risk DNA subtypes, reported as types 16, 18 or Others)	HPVY	Qvintip vaginal swab	3 days	KT364
HPV (Individually typed high risk DNA subtypes)	HPVZ	Qvintip vaginal swab	3 days	KT364
ISAC Panel 112 allergen panel from single sample (0.5µl) which components from 48 preselected allergen sources	ISAC includes	(TDL Tiny)	3 days	KT353
112 allergen panel from single sample (0.5µl) which		3 (TDL Tiny)	3 days 1 day	KT353
112 allergen panel from single sample (0.5µl) which components from 48 preselected allergen sources	includes			
112 allergen panel from single sample (0.5µl) which components from 48 preselected allergen sources  Lipase	includes	(TDL Tiny)	1 day	KT353
112 allergen panel from single sample (0.5µl) which components from 48 preselected allergen sources  Lipase  Lipid Profile	LIPA LIPP	(TDL Tiny) (TDL Tiny)	1 day 1 day	KT353 KT353
112 allergen panel from single sample (0.5µl) which components from 48 preselected allergen sources  Lipase  Lipid Profile  Lipoprotein (a)	LIPA LIPP LPOA	3 (TDL Tiny) 3 (TDL Tiny) (3 (TDL Tiny) (7 (TDL Tiny) and	1 day 1 day 1 day	KT353 KT353 KT353
112 allergen panel from single sample (0.5µl) which components from 48 preselected allergen sources  Lipase  Lipid Profile  Lipoprotein (a)  Liver Function Tests (Excluding AST)  Luteinising Hormone (LH)  Lymphogranuloma Venerium (LGV)  - Rectal  * This test can be configured to be	LIPA LIPP LPOA TLFT	3 (TDL Tiny) 3 (TDL Tiny) 5 (TDL Tiny) 7 (TDL Tiny) and 7 (TDL Tiny) 8 (TDL Tiny) or	1 day 1 day 1 day 1 day 1 day	KT353 KT353 KT353 KT445 KT353 or KT466
112 allergen panel from single sample (0.5µl) which components from 48 preselected allergen sources  Lipase  Lipid Profile  Lipoprotein (a)  Liver Function Tests (Excluding AST)  Luteinising Hormone (LH)  Lymphogranuloma Venerium (LGV)  Rectal	LIPA LIPP LPOA TLFT LH LGVP	3 (TDL Tiny) 3 (TDL Tiny) 5 (TDL Tiny) 7 (TDL Tiny) and 7 (TDL Tiny) and 7 (TDL Tiny) or 7 (TDL Tiny)	1 day 1 day 1 day 1 day 1 day	KT353 KT353 KT353 KT445 KT353 or KT466
112 allergen panel from single sample (0.5µl) which components from 48 preselected allergen sources  Lipase  Lipid Profile  Lipoprotein (a)  Liver Function Tests (Excluding AST)  Luteinising Hormone (LH)  Lymphogranuloma Venerium (LGV)  Rectal  * This test can be configured to be automatically reflexed as required.  Menopausal Profile (FSH, LH, TOES, TSH, FT4)	LIPA LIPP LPOA TLFT LH LGVP	3 (TDL Tiny) 3 (TDL Tiny) 5 (TDL Tiny) 6 (TDL Tiny) and 7 (TDL Tiny) or 7 (TDL Tiny)  Aptima multisite swab	1 day	KT353 KT353 KT353 KT445 KT353 or KT466 KT426
112 allergen panel from single sample (0.5µl) which components from 48 preselected allergen sources  Lipase  Lipid Profile  Lipoprotein (a)  Liver Function Tests (Excluding AST)  Luteinising Hormone (LH)  Lymphogranuloma Venerium (LGV)  — Rectal  * This test can be configured to be automatically reflexed as required.  Menopausal Profile (FSH, LH, TOES, TSH, FT4)  Avoid taking samples from any area an HRT cream	LIPA LIPP LPOA TLFT LH LGVP TMEN as applied. MPXV	3 (TDL Tiny) 3 (TDL Tiny) 6 (TDL Tiny) 7 (TDL Tiny) and 7 (TDL Tiny) or 7 (TDL Tiny) Aptima multisite swab 7 (TDL Tiny) and 8 (TDL Tiny)	1 day	KT353 KT353 KT353 KT445  KT353 or KT466 KT426
112 allergen panel from single sample (0.5µl) which components from 48 preselected allergen sources  Lipase  Lipid Profile  Lipoprotein (a)  Liver Function Tests (Excluding AST)  Luteinising Hormone (LH)  Lymphogranuloma Venerium (LGV)  - Rectal  * This test can be configured to be automatically reflexed as required.  Menopausal Profile (FSH, LH, TOES, TSH, FT4)  Avoid taking samples from any area an HRT cream  MPOX (Virus) – Lesion	LIPA LIPP LPOA TLFT LH LGVP TMEN as applied. MPXV	3 (TDL Tiny) 3 (TDL Tiny) (TDL Tiny) (TDL Tiny) and (TDL Tiny) (TDL Tiny) (TDL Tiny) (TDL Tiny) Aptima multisite swab  4 (TDL Tiny) Aptima multisite swab  Purple liquid	1 day 2 days	KT353 KT353 KT353 KT445  KT353 or KT466 KT426  KT445

TEST	CODE	SAMPLE REQS	TAT	KIT CODE
Mycoplasma genitalium by PCR – Urine and Vaginal	MGEN	Aptima urine or multisite swab	2 days	KT356 or KT357
Mycoplasma genitalium Resistance – Urine or Vaginal	MGR	Aptima urine or multisite swab	1-2 weeks	KT356 or KT357
Oestradiol-17-Beta	T0ES	(TDL Tiny)	1 day	KT466
Omega 3/Omega 6	OMG3	(TDL Tiny)	5 days	KT354
PEth (Phosphatidylethanol)	PETH	(TDL Tiny)	5-7 days	KT354
PLAC Test (Lp-PLA2)	PLA2	(TDL Tiny)	2 days	KT353
<b>Progesterone</b> Avoid taking samples from any area an HRT cream is	PROG s applied.	(TDL Tiny) or (TDL Tiny)	1 day	KT353 or KT466
Prolactin	PROL	(TDL Tiny) or (TDL Tiny)	1 day	KT353 or KT466
Prostate Specific Antigen (Total)	PSPA	(TDL Tiny)	1 day	KT353
QFIT/Calprotectin Profile (Combined)	QCAL	<b>QFIT</b> sample collection device	5 days	KT362 or KT430
Quantitative Faecal Immunochemical Test (QFIT)	QFIT	<b>QFIT</b> sample collection device	1 day	KT362 or KT430
Respiratory PCR Panel (COVID-19, Flu A/B and RSV)	FLU4	Aptima multisite swab of nose/throat	2 days	KT293
Selenium (Serum)	SELE	(TDL Tiny)	4 days	KT353
Sex Hormone Binding Globulin	SHBG	(TDL Tiny) or (TDL Tiny)	1 day	KT353 or KT466
STI Profile by PCR (7 tests from 1 sample)	DL12	Aptima urine or multisite swab	2 days	KT356 or KT357
STI Profile: MSM1 (Blood + Urine/ Throat/Rectal Swabs)	MSM1	(TDL Tiny) / Aptima Urine / Aptima multisite swab x 2	2 days	KT361
STI Profile: MSM2 (Blood + Urine/ Throat/Rectal Swabs)	MSM2	(TDL Tiny) / Aptima urine / Aptima multisite swab x 2	3 days	KT361
Syphilis IgG/IgM	TSYP	(TDL Tiny)	1 day	KT353
<b>Testosterone</b> Avoid taking samples from any area an HRT cream is	TEST s applied.	(TDL Tiny) or (TDL Tiny)	1 day	KT353 or KT466
<b>Testosterone (Free)</b> Avoid taking samples from any area an HRT cream is	FTES s applied.	B (TDL Tiny) or (TDL Tiny)	3 days	KT353 or KT466
Thyroid Abs (Thyroglobulin + Thyroid Peroxidase Abs)	THAB	B (TDL Tiny)	2 days	KT353
Thyroid Profile 1 (FT4/TSH)	TF	(TDL Tiny)	1 day	KT353
Thyroid Profile 3 (FT3/FT4/TSH)	TF3	(TDL Tiny)	1 day	KT353
Tissue Transglutaminase IgA (Coeliac)	TAA	(TDL Tiny)	2 days	KT353

TEST	CODE	SAMPLE REQS	TAT	KIT CODE
Trichomonas vaginalis (TV) – Urine or Vaginal	TVPC	Aptima urine or multisite swab	2 days	KT356 or KT357
Triple Swab Female STI Profile (Vaginal/Throat/Rectal Swabs)	3SWA	Aptima multisite swab x 3 (label by site)	2 days	KT428
TSH	TSH	(TDL Tiny)	1 day	KT353
Urea	UREA	(TDL Tiny)	1 day	KT353
Urea/Creatinine/eGFR	TCU	(TDL Tiny)	1 day	KT353
Urine Chemistry and Microscopy	UMIC	Urine (Universal). Mid stream.	1-2 days	KT385
Urine Chemistry, Microscopy and Culture	UCEM	Urine (Universal & Boric). Mid stream.	1-2 days	KT386
Vaginitis/BV Profile using Culture & PCR Swab	STD8	Aptima multisite swab and Blue gel Amies swab	3-5 days	KT441
Viral Respiratory RNA Screen by PCR	VPR	Aptima multisite swab of nose/throat	2 days	KT293
Vitamin B12 (Active)	B12	(TDL Tiny)	1 day	KT353
Vitamin D (25-OH)	VITD	(TDL Tiny)	1 day	KT353

## **Screening for Drugs of Abuse/Alcohol**

TEST	CODE	SAMPLE REQS	TAT
Alcohol Profile	AP	<b>ABBG</b>	5-7 days
Alcohol Profile 2	ALCP	A A B B G RU	5-7 days
Amphetamines – Blood	AMPB	BB	5 days
Cannabinoids (Urine) Screen	CANN	RU	1 day
Cocaine (Urine) Screen	UCOC	RU	1 day
Drugs of Abuse from Blood without Chain of Custody	DOAP	B	5 days
Drugs of Abuse Profile – Random Urine Sample/No Chain of Custody	DOA	RU	2 days (5 days with LC-MS/ MS confirmation)
Drugs of Abuse Profile – Random Urine Sample/No Chain of Custody Plus Alcohol	DOA3	RU	2 days (5 days with LC-MS/ MS confirmation)
Drugs of Abuse Profile — With Chain of Custody* *Appointment required at 76 Wimpole Street and Photo ID to be shown.	DOAL	RU/CoC Collection Containers 1,2	2 days (5 days with LC-MS/ MS confirmation)
Drugs of Abuse Profile – Without Chain of Custody	DOAN	RU <sup>2</sup>	2 days (5 days with LC-MS/ MS confirmation)
Ketamine Screen	KETA	RU	7-10 days
LSD	LSD	RU	5 days
Opiate Screen (Urine)	UOPI	RU	2 days
PEth (Phosphatidylethanol) (Self-collect)	PETH	(TDL Tiny)	5-7 days
PEth (Phosphatidylethanol) (Venous)	PETH	A	5-7 days
Urine EtG (Ethyl glucuronide)	ETG	RU	1 week

#### **Alcohol Profile**

Liver Function Tests (LFT)
Alcohol Level
PEth (Phosphatidylethanol)
Carbohydrate Deficient
Transferrin (CDT)
Mean Cell Volume (MCV)

#### TAT: 5-7 days

ΑP



#### **Alcohol Profile 2**

Liver Function Tests (LFT)
Alcohol Level
PEth (Phosphatidylethanol)
Carbohydrate Deficient
Transferrin (CDT)
Mean Cell Volume (MCV)
Urine Ethyl Gluconaride (EtG)

#### TAT: 5-7 days

**ALCP** 



## Drugs of Abuse from Blood without Chain of Custody

Amphetamines Barbiturates Tricyclic Antidepressants Benzodiazepine Cannabinoids Opiates Cocaine

#### TAT: 5 days

D0AP



#### **Screening for Drugs of Abuse/Alcohol**

## Drugs of Abuse Profile – Random Urine Sample/ No Chain of Custody

Amphetamines Barbiturates

Benzodiazepine Cannabinoids

Cocaine

Codeine – opiate

Dihydrocodeine - opiate

MDMA

Methadone

Morphine - opiate

## TAT: 2 days (5 days with LC-MS/MS confirmation)

DOA

RU

## Drugs of Abuse Profile – Random Urine Sample/No Chain of Custody Plus Alcohol

Alcohol

Amphetamines

Barbiturates

Benzodiazepine

Cannabinoids

Cocaine

Codeine - opiate

Dihydrocodeine – opiate

 $\mathsf{MDMA}$ 

Methadone

Morphine - opiate

## TAT: 2 days (5 days with LC-MS/MS confirmation)

DOA3

RU

# Drugs of Abuse Profile – With Chain of Custody\*

Alcohol

**Amphetamines** 

Barbiturates

Benzodiazepine Cannabinoids

Cocaine

Codeine - opiate

Dihydrocodeine - opiate

Ketamine

LSD

MDMA Methadone

Methagualone

Morphine - opiate

Phencyclidine

Propoxyphene
\*Appointment required at

76 Wimpole Street] and Photo ID to be shown

## TAT: 2 days (5 days with LC-MS/MS confirmation)

DOAL.

RU/CoC Collection Containers 1,2

## Drugs of Abuse Profile – Without Chain of Custody

Alcohol

Amphetamines

Barbiturates

Benzodiazepine Cannabinoids

Cocaine

Codeine – opiate

Dihydrocodeine – opiate

Ketamine

LSD MDMA

Methadone

Methagualone

Morphine – opiate

Phencyclidine Propoxyphene

TAT: 2 days (5 days with LC-MS/MS confirmation)

DOAN

RU<sup>2</sup>

## **Occupational Health**

## **Trace metals in blood**

TEST	CODE	SAMPLE REQS	TAT
Aluminium (Blood)	ALUM	<b>(</b>	7 days
Arsenic (Blood)	ARS	A or (1)	5 days
Cadmium (Blood)	CADM	(A) or (1)	5 days
Chromium (Blood)	CHR0	🛕 / Trace metal / 🕕	5 days
Copper (Serum)	COPP	B or <	5 days
Lead (Blood)	LEAD	A	5 days
Magnesium (Serum)	MG	B	1 day
Manganese (Serum)	MANG	B	5 days
Mercury (Blood)	MERC	A or (1)	5 days
Nickel (Serum)	NICK	B	5 days
Silver (Blood)	SILV	B	5 days
Trace Metal (Blood) Profile	TRAC	<b>ABB</b>	7-10 days
Zinc (Serum)	ZINC	<b>®</b>	2 days

## **Trace metals in blood**

## **Occupational Health**

## **Tests for specific exposure**

TEST	CODE	SAMPLE REQS	TAT
2-Butanone GC	BUTA	RU	7 days
Acetone – Blood	ACTB	A or (1)	2 weeks
Acetone – Urine	ACTU	RU	5 days
Alcohol Profile	AP	ABBG	5-7 days
Alcohol Profile 2	ALCP	A A B B G RU	5-7 days
Benzene	BENZ	J <sup>1,6</sup>	3 days
Beta 2 Microglobulin (Serum)	B2MG	B	2 days
Beta 2 Microglobulin (Urine)	UB2M	RU	3 days
Bromide	BROM	В	3 days
Cholinesterase (Serum/Pseudo)	CHPS	B	1 day
Doxepin Level (Sinequan)	DOXE	A or (1)	10 days
MBOCA in Urine	MBOC	RU	10 days
Molybdenum (Serum)	MOLY	В	5 days
Thallium (Blood)	THAL	<b>A</b> / <b>(1)</b>	1 week
Thallium (Urine)	URTH	RU	1 week
Toluene (Blood)	T0L	J (Contact Referrals)	10 days
Toluene (Urine)	UT0L	RU	10 days
Trichloracetic Acid (Urine)	UTCA	RU	5 days
Xanthine – Blood	XANB	A	2 weeks
Xylene – Urine	UXYL	RU <sup>30</sup>	2 weeks

#### **Alcohol Profile**

Liver Function Tests (LFT)
Alcohol Level
PEth (Phosphatidylethanol)
Carbohydrate Deficient
Transferrin (CDT)
Mean Cell Volume (MCV)

TAT: 5-7 days

ΑP



#### **Alcohol Profile 2**

Liver Function Tests (LFT)
Alcohol Level
PEth (Phosphatidylethanol)
Carbohydrate Deficient
Transferrin (CDT)
Mean Cell Volume (MCV)
Urine Ethyl Gluconaride (EtG)

TAT: 5-7 days

ALCP



# Trace Metal (Blood) Profile

Aluminium Manganese Iron Calcium Zinc Magnesium Copper Cadmium Mercury Lead Chromium

**TAT: 7-10 days** 

TRAC



The cervical cytology laboratory provides a rapid service for liquid based cervical samples.

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 8.00am to 6.00pm. Out-of-hours results are available on 020 7307 7373.

### **Urgent samples**

It is helpful if requests for urgent samples can be discussed with the Senior Management Team prior to sending. Please contact the laboratory on cytology.reporting@tdlpathology.com in the first instance.

## Use of service/Information required

Request forms must include 3 patient identifiers (this can be patient's full name, date of birth, hospital number or reference number). Samples will not be processed without a request form. TDL Request Forms do not include the information required for NHS requests for cervical cytology and should not be used for NHS requests. For further information on NHS requests please contact hsl.csl.queries@nhs.net.

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.

Sample takers are requested to check the expiry date of the vial prior to taking the sample. The laboratory are unable to process samples if the expiry date has passed.

#### Clinical advice

The Consultant Cytopathologists and the Consultant Biomedical Scientist work together to provide clinical and technical advice, including recommendations for follow-up, HPV testing and management of complex cases. TDL will provide recommendation for patient management, but not undertake to provide a direct referral. No result will be entered onto the NHS CSP database and will therefore not be part of an individual's NHS screening record. Failsafe and management of the patient and their follow up, including referral for colposcopy where indicated, would need to be arranged by their referring clinician. To contact the department directly, please call 020 7307 7387 or email cytology.reporting@tdlpathology.com.



#### RECORD...

...the patient's 3 identifiers to include date of birth on the vial, and the patient information and medical history on the cytology requisition form. TDL Request Forms do not include the information required for NHS requests and should not be used for NHS requests.



#### 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.

## 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.
   Samples received in an expired vial will not be processed.
- 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 Jubricant is not recommended.

#### D<sub>0</sub>

- If excessive mucus is present, this should be gently removed before sampling.
- Use either the Cervex Brush (broom-like device) on its own or in combination with an endocervical brush.
- 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 6 weeks of collection.
- DO NOT use excessive lubricant
   please avoid if possible.

## **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.

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 ThinPrep vials are available by contacting the Laboratory Service Centre on 020 7307 7373.

# STI Screening from Hologic Thin Prep Vial

Tests are priced individually. Please request tests individually. Requests for additional test can be made by contacting the laboratory by telephone on **020 7307 7373** or by email to **addons@tdlpathology.com**.

## **Infection by PCR (single tests)**

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

## Multiple tests from a single sample

TEST	CODE	SAMPLE REQS	TAT
7 STI Profile by PCR (7 tests from 1 Sample) Chlamydia trachomatis, Neisseria gonorrhoea, Mycoplasma genitalium, Ureaplasma species, Trichomonas vaginalis, Gardnerella vaginalis, Herpes Simplex I/II	PP12	TPV	2 days

## **Human papillomavirus (HPV)**

Human papillomavirus (HPV) is a common virus which infects the skin and may be transmitted through sexual contact. There are over 200 types of HPV which are split into two groups depending on whether they are linked to an increased risk of cervical cancer. These are described as high risk HPV subtypes or low risk HPV subtypes.

Infection with a high-risk HPV (hrHPV) has been established as a necessary cause of cervical cancer. This has led in recent years to the inclusion of hrHPV testing as an adjunct to cervical cytology in organised cervical screening programmes.

Compared to cervical cytology, hrHPV testing has been shown to reduce the risk of developing cervical cancer through increased sensitivity. The high negative predictive value of hrHPV testing and lower false negative rate provides assurance to woman and clinicians that the risk of developing cervical cancers between screening tests, is rare.

### 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 and they may return to Routine Recall.
- If HR-HPV is POSITIVE (Detected) this means that CYTOLOGY will be processed from the same ThinPrep Vial.

A further specimen is not required.

Patient recall (management) will be determined by the individual screening history and current test results.

All TDL requests for HPV are processed as follows:

- If HPV is requested as a single test and the result is Negative/Not Detected, cervical cytology (PAPT) would only be processed if specifically requested. Should HPV and PAPT be undertaken, there would be a charge for both the HPV and the PAPT.
- If the HPV result is HR-HPV Detected, cervical cytology (PAPT) will be processed, even if the PAPT has not been requested. The PAPT will not be charged.

# Understanding the significance of HPV testing

The benefit of a negative HPV result is its negative predictive value (NPP). A negative HPV result indicates that a patient is a very low risk of developing cervical disease. However neither HPV testing not negative cervical cytology are able to reduce the risk to zero. The negative predictive value of both DNA and mRNA testing is the same.

Requests for Cervical Cytology as a single test are no longer processed without testing for HPV. In these circumstances, the HPV test will be charged in addition to the Cervical Cytology. Requests for HPV as the PRIMARY TEST will reflex to Cervical Cytology if HR-HPV is Detected/Positive at no additional charge.

## **Requests for HPV Primary Screening as a single test**

TEST	CODE	SAMPLE REQS	TAT
HPV (A group of 14 HR mRNA types)	HPVH	TPV	3 days

If HR-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 REQS	TAT
HPV (28 individually typed LR & HR DNA subtypes)	HP20	TPV	3 days

HPV low and high risk DNA subtypes will be reported individually (9 low and 19 high risk). If High Risk DNA subtypes are positive then Cervical Cytology (PAPT) using the same vial will be processed **without charge**.

### Requests for HPVT as a single test

TEST	CODE	SAMPLE REQS	TAT
HPV (28 individually typed low risk (LR) & high risk (HR) DNA subtypes and reflexed mRNA for types 16, 18, 31, 33 and 45)	HPVT	TPV	5 days

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

## **Requests for Cervical Cytology (PAPT)**

TEST	CODE	SAMPLE REQS	TAT
Cervical Cytology	PAPT	TPV	6 days (combined report)
Cervical Cytology + HPVH	PAPT + HPVH	TPV	6 days (combined report)

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 as two separate tests**.

## Requests for Cervical Cytology (PAPT) with selected HPV (HPVH or HP20 or HPVT)

TEST	CODE	SAMPLE REQS	TAT
Cervical Cytology + HPVH	PAPT + HPVH	TPV	6 days (combined report)
Cervical Cytology + HP20	PAPT + HP20	TPV	6 days (combined report)
Cervical Cytology + HPVT	PAPT + HPVT	TPV	6 days (combined report)

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

### **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 19 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 incorporating a collective of high risk subtypes.

#### **HPVZ**

Self-Collected HPV DNA with **individual** reporting of all High Risk subtypes (16, 18, 31, 33, 45, 35, 39, 51, 52, 56, 58, 59, 66, 68, 26, 53, 69, 73, 82).

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 REQS	TAT
HPV (19 high risk DNA subtypes, reported as types 16, 18 or Others) (Self-collect)	HPVY	Qvintip vaginal swab	3 days
HPV (Individually typed high risk DNA subtypes) (Self-collect)	HPVZ	Qvintip vaginal swab	3 days

## **Non-Gynae Cytology**

## **Non-Gynaecological Cytology**

## **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.

#### 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.

#### **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 is 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.

TEST	CODE	SAMPLE REQS	TAT
Fluid Cytology	CATF	Fluid <sup>4</sup>	3 days
Urine Cytology (Urine cytology containers available from TDL Supplies)	URCY	Urine (30mls) <sup>21</sup>	2 days

TDL's Histopathology service supports a full range of pathology sub-specialities.

To prevent tissue degeneration, it is advisable to collect histopathology samples in sample pot(s) containing preservative, usually formalin, to at least ten times the volume of the tissue sample (available from TDL Supplies). Use of preservative will ensure that the tissue architecture and microscopic appearances of specimens are preserved.

Patient demographics, together with clinical and sample details need to be provided with the specimens. Testicular investigations for reproductive investigations are best submitted fixed in Bouins solution. Requests for products of conception require the patient's signed consent/instruction regarding sensitive disposal when the histopathology is complete. Please contact **020 7307 7380** or **020 7307 7373** for information or any query relating to histopathology.

All specimens are initially stained with H&E. However special stains and immunohistochemistry (IHC) may be recommended if additional information is needed to provide a more detailed analysis. The choice of stain depends on the findings on initial assessment, the clinical context and the preference of the pathologist within their specialist expertise. IHC may be added when routine or regular histological testing is not sufficient to form a diagnosis. There are additional charges for special stains and immunohistochemistry, as below.

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	Mastecomy (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

CATEGORY	CODE	TISSUE SAMPLE
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	Gingivial 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
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

CATEGORY	CODE	TISSUE SAMPLE
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	HIS2	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
GI-Resection – Small	HIS2	Mesentery
GI-Resection – Small	HIS2	Perianal Biopsy / Warts
GI-Resection – Small	HIS2	Pilonidal Sinus

CATEGORY	CODE	TISSUE SAMPLE
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	lleostomy
GI Resection – Large	HIS3	lleum
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 / Pancreatectoduodenectomy
Gynaecology	HIS2	Cervical Biopsy
Gynaecology	HIS1	Cervical Polyp
Gynaecology	HIS4	Cervix
Gynaecology	HIS1	Curettings – Endocervical
Gynaecology	HIS1	Curettings – Endometial
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)

## Histopathology

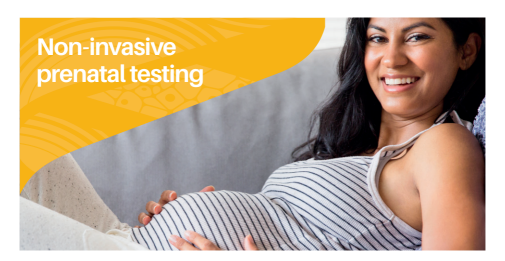
Gynaecology         HIS2         Pelvic Mass           Gynaecology         HIS5         Placenta           Gynaecology         HIS5         Placenta           Gynaecology         HIS2         Pouch of Douglas           Gynaecology         HIS1         Products of Conception           Gynaecology         HIS4         Uterus           Gynaecology         HIS5         Uterus and Cervix           Gynaecology         HIS5         Uterus, Tubes and Ovaries           Gynaecology         HIS5         Dene Marrow           Haemato-Oncology         HIS2         Lymph Node           Haemato-Oncology         HIS3         Lymph Node (Metastatic Disease)           Haemato-Oncology         HIS5         Spleen           Haemato-Oncology         HIS5         Thymus           Lung - Resections         HIS5         Lung Resection           Lung - Re	CATEGORY	CODE	TISSUE SAMPLE
Gynaecology HIS5 Placenta  Gynaecology HIS2 Pouch of Douglas  Gynaecology HIS1 Products of Conception  Gynaecology HIS2 Uterine Polyp  Gynaecology HIS3 Uterus and Cervix  Gynaecology HIS5 Uterus, Tubes and Ovaries  Gynaecology 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 Spleen  Haemato-Oncology HIS5 Thymus  Lung - Resections  HIS3 Lung Biopsy  Lung - Resections  HIS3 Lung Lesion Small Wedge Resection  Lung - Resections HIS3 Lung Tumour Resection +/- Nodes  Neurosurgery HIS3 Brain Biopsy  Neurosurgery HIS3 Brain Resection  Neurosurgery HIS3 Spiral Tumour Biopsy  Neurosurgery HIS3 Spiral Tumour Resection  Neurosurgery HIS3 Spiral Tumour Biopsy  Neurosurgery HIS3 Spiral Tumour Resection  Neurosurgery HIS4 Vertebrea  Opthalmic HIS1 Conjunctival Biopsy / Excision  Opthalmic HIS1 Cornea  Opthalmic HIS1 Orbit Contents of Eye  Orthopaedic HIS2 Bone Currettings  Orthopaedic HIS2 Bone Currettings  Orthopaedic HIS2 Bone Currettings	Gynaecology	HIS2	Pelvic Mass
Gynaecology HIS1 Products of Conception  Gynaecology HIS2 Uterine Polyp  Gynaecology HIS4 Uterus  Gynaecology HIS5 Uterus and Cervix  Gynaecology HIS5 Uterus and Cervix  Gynaecology HIS5 Uterus and Ovaries  Gynaecology HIS5 Uterus, Tubes and Ovaries  Gynaecology HIS1 Vulval Biopsy  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 Spleen  Haemato-Oncology HIS5 Thymus  Lung - Biopsy HIS3 Lung Biopsy  Lung - Resections HIS3 Lung Lung Resection  Lung - Resections HIS5 Lung Turmour Resection  Lung - Resections HIS5 Lung Turmour Resection  Neurosurgery HIS3 Brain Resection  Neurosurgery HIS3 Brain Resection  Neurosurgery HIS3 Spinal Turmour Biopsy  Neurosurgery HIS3 Spinal Turmour Resection  Neurosurgery HIS4 Vertebrea  Opthalmic HIS1 Conjunctival Biopsy  Opthalmic HIS1 Cornea  Opthalmic HIS1 Cornea  Opthalmic HIS1 Orbit Contents of Eye  Orthopaedic HIS1 Bone Biopsy  Orthopaedic HIS2 Bone Currettings  Orthopaedic Orthopaedic HIS2 Borea	Gynaecology	HIS1	Peritoneal Biopsy
Gynaecology HIS1 Products of Conception  Gynaecology HIS2 Uterine Potyp  Gynaecology HIS4 Uterus  Gynaecology HIS5 Uterus and Cervix  Gynaecology HIS5 Uterus, Tubes and Ovaries  Gynaecology HIS1 Vulval Biopsy Haemato-Oncology HIS2 Lymph Node Haemato-Oncology HIS3 Lymph Node (Lymphoma) Haemato-Oncology HIS3 Lymph Node (Metastatic Disease) Haemato-Oncology HIS3 Lymph Node (Metastatic Disease) Haemato-Oncology HIS5 Spleen Haemato-Oncology HIS5 Spleen Haemato-Oncology HIS5 Spleen  Haemato-Oncology HIS5 Spleen  Haemato-Oncology HIS5 Spleen  Haemato-Oncology HIS5 Lung Biopsy Lung - Resections HIS3 Lung Lesion Small Wedge Resection  Lung - Resections HIS5 Lung Tumour Resection +/- Nodes  Neurosurgery HIS3 Brain Biopsy Neurosurgery HIS3 Brain Resection  Neurosurgery HIS3 Pituitary Gland - Resection  Neurosurgery HIS3 Spinal Tumour Biopsy Opthalmic HIS1 Conjunctival Biopsy Opthalmic HIS1 Corporation Opthalmic HIS1 Corporation Opthalmic HIS2 Lacrimal Gland Biopsy / Excision Opthalmic HIS1 Orbit Contents of Eye Orthopaedic HIS1 Bone Biopsy Orthopaedic HIS2 Bone Currettings Orthopaedic Orthopaedic HIS2 Bone Currettings	Gynaecology	HIS5	Placenta
Gynaecology Gynaecology HIS4 Uterus Gynaecology HIS5 Uterus and Cervix Gynaecology HIS5 Uterus, Tubes and Ovaries Gynaecology HIS1 Vulval Biopsy 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 Spleen Haemato-Oncology HIS5 Thymus Lung Biopsy Lung Biopsy Lung Resections HIS5 Lung Resection Lung - Resections HIS5 Lung Tumour Resection +/- Nodes Neurosurgery HIS3 Brain Resection Neurosurgery HIS3 Prituitary Gland - Resection Neurosurgery HIS3 Spinal Tumour Resection Neurosurgery HIS4 Vertebrea Opthalmic HIS1 Conjunctival Biopsy Opthalmic HIS1 Conjunctival Biopsy / Excision Opthalmic HIS1 Orbit Contents of Eye Orthopaedic HIS1 Bone Biopsy Orthopaedic HIS2 Bone Currettings Orthopaedic HIS2 Bone Currettings	Gynaecology	HIS2	Pouch of Douglas
Gynaecology HIS5 Uterus and Cervix  Gynaecology HIS5 Uterus, Tubes and Ovaries  Gynaecology HIS1 Vulval Biopsy 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 Spleen Haemato-Oncology HIS5 Thymus Lung - Biopsy HIS3 Lung Biopsy Lung - Resections HIS3 Lung Lesion Small Wedge Resection Lung - Resections HIS5 Lung Tumour Resection +/- Nodes Neurosurgery HIS3 Brain Biopsy Neurosurgery HIS3 Brain Resection Neurosurgery HIS3 Pituitary Gland - Resection Neurosurgery HIS3 Spinal Tumour Resection Neurosurgery HIS3 Spinal Tumour Resection Neurosurgery HIS3 Spinal Tumour Resection Neurosurgery HIS4 Vertebrea Opthalmic HIS1 Cornea Opthalmic HIS1 Orbit Contents of Eye Orthopaedic HIS1 Bone Biopsy Orthopaedic HIS2 Bursa	Gynaecology	HIS1	Products of Conception
Gynaecology HIS5 Uterus and Cervix  Gynaecology HIS5 Uterus, Tubes and Ovaries  Gynaecology HIS1 Vulval Biopsy 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 Spleen Haemato-Oncology HIS5 Thymus Lung - Biopsy HIS3 Lung Biopsy Lung - Resections HIS3 Lung Lesion Small Wedge Resection Lung - Resections HIS5 Lung Tumour Resection +/- Nodes Neurosurgery HIS3 Brain Biopsy Neurosurgery HIS3 Brain Biopsy Neurosurgery HIS3 Pituitary Gland - Resection Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS3 Spinal Tumour Resection Neurosurgery HIS4 Vertebrea Opthalmic HIS1 Cornea Opthalmic HIS1 Orbit Contents of Eye Orthopaedic HIS1 Bone Biopsy Orthopaedic HIS2 Bursa	Gynaecology	HIS2	Uterine Polyp
Gynaecology HIS5 Uterus, Tubes and Ovaries Gynaecology HIS1 Vulval Biopsy 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 Lung - Biopsy Lung - Resections HIS3 Lung Biopsy Lung - Resections HIS5 Lung Tumour Resection Lung - Resections HIS5 Brain Biopsy Neurosurgery HIS3 Brain Resection Neurosurgery HIS3 Pituitary Gland - Resection Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS3 Spinal Tumour Resection Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS4 Vertebrea Opthalmic HIS1 Cornea Opthalmic HIS4 Globe / Removal of Eye Opthalmic HIS1 Orbit Contents of Eye Orthopaedic HIS2 Bursa Orthopaedic HIS2 Bursa	Gynaecology	HIS4	Uterus
Gynaecology HIS1 Vulval Biopsy 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 Lung Tumour Resection +/- Nodes Neurosurgery HIS3 Brain Biopsy Neurosurgery HIS3 Brain Resection Neurosurgery HIS3 Pituitary Gland - Resection Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS4 Vertebrea Opthalmic HIS1 Cornea Opthalmic HIS4 Globe / Removal of Eye Opthalmic HIS5 Bone Biopsy Orthopaedic HIS2 Bursa Orthopaedic HIS2 Bursa	Gynaecology	HIS5	Uterus and Cervix
Haemato-Oncology HIS5 Bone Marrow Haemato-Oncology HIS2 Lymph Node Haemato-Oncology HIS3 Lymph Node (Lymphoma) Haemato-Oncology HIS3 Lymph Node (Lymphoma) Haemato-Oncology HIS5 Spleen Haemato-Oncology HIS5 Thymus Lung - Biopsy HIS3 Lung Biopsy Lung - Resections HIS3 Lung Lesion Small Wedge Resection Lung - Resections HIS5 Lung Tumour Resection Lung - Resections HIS5 Lung Tumour Resection +/- Nodes Neurosurgery HIS3 Brain Biopsy Neurosurgery HIS3 Brain Resection Neurosurgery HIS3 Pituitary Gland - Resection Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS3 Spinal Tumour Resection Neurosurgery HIS3 Spinal Tumour Resection Neurosurgery HIS4 Vertebrea Opthalmic HIS1 Conjunctival Biopsy Opthalmic HIS4 Globe / Removal of Eye Opthalmic HIS5 Orbit Contents of Eye Orthopaedic HIS2 Bone Currettings Orthopaedic HIS2 Bone Currettings Orthopaedic HIS2 Bursa	Gynaecology	HIS5	Uterus, Tubes and Ovaries
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-HIS5 Lung Tumour Resection +/- Nodes Neurosurgery HIS3 Brain Biopsy Neurosurgery HIS3 Brain Resection Neurosurgery HIS3 Pituitary Gland - Resection Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS3 Spinal Tumour Resection Neurosurgery HIS3 Spinal Tumour Resection Neurosurgery HIS4 Vertebrea Opthalmic HIS1 Conjunctival Biopsy Opthalmic HIS4 Globe / Removal of Eye Opthalmic HIS5 Orbit Contents of Eye Orthopaedic HIS1 Bone Biopsy Orthopaedic HIS2 Bursa	Gynaecology	HIS1	Vulval Biopsy
Haemato-Oncology HIS3 Lymph Node (Lymphoma) Haemato-Oncology HIS5 Spleen Haemato-Oncology HIS5 Spleen Haemato-Oncology HIS5 Thymus Lung Biopsy Lung Biopsy Lung Resections HIS3 Lung Biopsy Lung Resection HIS5 Lung Resection Lung Resections HIS5 Lung Turmour Resection +/- Nodes Neurosurgery HIS3 Brain Biopsy Neurosurgery HIS3 Brain Resection Neurosurgery HIS3 Pituitary Gland - Resection Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS4 Vertebrea Opthalmic HIS1 Cornea Opthalmic HIS1 Cornea Opthalmic HIS2 Lacrimal Gland Biopsy / Excision Opthalmic HIS1 Orbit Contents of Eye Orthopaedic HIS2 Bone Currettings Orthopaedic HIS2 Bursa	Haemato-Oncology	HIS5	Bone Marrow
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 HIS3 Pituitary Gland - Resection Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS3 Spinal Tumour Resection Neurosurgery HIS4 Vertebrea Opthalmic HIS1 Conjunctival Biopsy Opthalmic HIS1 Cornea Opthalmic HIS4 Globe / Removal of Eye Opthalmic HIS2 Lacrimal Gland Biopsy / Excision Opthalmic HIS1 Orbit Contents of Eye Orthopaedic HIS2 Bone Currettings Orthopaedic HIS2 Bone Currettings	Haemato-Oncology	HIS2	Lymph Node
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 HIS3 Brain Resection  Neurosurgery HIS3 Pituitary Gland – Resection  Neurosurgery HIS3 Spinal Tumour Biopsy  Neurosurgery HIS3 Spinal Tumour Resection  Neurosurgery HIS3 Spinal Tumour Resection  Neurosurgery HIS4 Vertebrea  Opthalmic HIS1 Conjunctival Biopsy  Opthalmic HIS1 Cornea  Opthalmic HIS4 Globe / Removal of Eye  Opthalmic HIS4 Globe / Removal of Eye  Opthalmic HIS1 Orbit Contents of Eye  Opthalmic HIS1 Bone Biopsy  Orthopaedic HIS1 Bone Biopsy  Orthopaedic HIS2 Bone Currettings  Orthopaedic HIS2 Bursa	Haemato-Oncology	HIS3	Lymph Node (Lymphoma)
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 HIS3 Pituitary Gland - Resection  Neurosurgery HIS3 Spinal Tumour Biopsy  Neurosurgery HIS3 Spinal Tumour Biopsy  Neurosurgery HIS3 Spinal Tumour Resection  Neurosurgery HIS4 Vertebrea  Opthalmic HIS1 Conjunctival Biopsy  Opthalmic HIS1 Cornea  Opthalmic HIS4 Globe / Removal of Eye  Opthalmic HIS2 Lacrimal Gland Biopsy / Excision  Opthalmic HIS1 Orbit Contents of Eye  Orthopaedic HIS1 Bone Biopsy  Orthopaedic HIS2 Bone Currettings  Orthopaedic HIS2 Bursa	Haemato-Oncology	HIS3	Lymph Node (Metastatic Disease)
Lung – Biopsy HIS3 Lung Biopsy Lung – Resections HIS5 Lung Resection Lung – Resections HIS5+HIS5 Lung Resection Lung – Resections HIS5 Lung Tumour Resection +/- Nodes Neurosurgery HIS3 Brain Biopsy Neurosurgery HIS3 Brain Resection Neurosurgery HIS3 Brain Resection Neurosurgery HIS3 Pituitary Gland – Resection Neurosurgery HIS3 Spinal Tumour Biopsy Neurosurgery HIS3 Spinal Tumour Resection Neurosurgery HIS3 Spinal Tumour Resection Neurosurgery HIS4 Vertebrea Opthalmic HIS1 Conjunctival Biopsy Opthalmic HIS1 Cornea Opthalmic HIS4 Globe / Removal of Eye Opthalmic HIS2 Lacrimal Gland Biopsy / Excision Opthalmic HIS1 Orbit Contents of Eye Orthopaedic HIS2 Bone Biopsy Orthopaedic HIS2 Bone Currettings Orthopaedic HIS2 Bursa	Haemato-Oncology	HIS5	Spleen
Lung - ResectionsHIS3Lung Lesion Small Wedge ResectionLung - ResectionsHIS5 + HIS5Lung Resection + /- NodesNeurosurgeryHIS3Brain BiopsyNeurosurgeryHIS3Brain ResectionNeurosurgeryHIS3 + HIS5Muscle BiopsyNeurosurgeryHIS3 Pituitary Gland - ResectionNeurosurgeryHIS3 Spinal Tumour BiopsyNeurosurgeryHIS3 Spinal Tumour ResectionNeurosurgeryHIS4 VertebreaOpthalmicHIS1 Conjunctival BiopsyOpthalmicHIS1 CorneaOpthalmicHIS4 Globe / Removal of EyeOpthalmicHIS2 Lacrimal Gland Biopsy / ExcisionOpthalmicHIS1 Orbit Contents of EyeOrthopaedicHIS1 Bone BiopsyOrthopaedicHIS2 Bone CurrettingsOrthopaedicHIS2 Bursa	Haemato-Oncology	HIS5	Thymus
Lung – Resections       HIS5+HIS5       Lung Resection         Lung – Resections       HIS5       Lung Turnour Resection +/- Nodes         Neurosurgery       HIS3       Brain Biopsy         Neurosurgery       HIS3       Brain Resection         Neurosurgery       HIS5+HIS5       Muscle Biopsy         Neurosurgery       HIS3       Pituitary Gland – Resection         Neurosurgery       HIS3       Spinal Turnour Biopsy         Neurosurgery       HIS3       Spinal Turnour Resection         Neurosurgery       HIS4       Vertebrea         Opthalmic       HIS1       Conjunctival Biopsy         Opthalmic       HIS1       Cornea         Opthalmic       HIS4       Globe / Removal of Eye         Opthalmic       HIS2       Lacrimal Gland Biopsy / Excision         Opthalmic       HIS1       Orbit Contents of Eye         Orthopaedic       HIS1       Bone Biopsy         Orthopaedic       HIS2       Bone Currettings         Orthopaedic       HIS2       Bursa	Lung – Biopsy	HIS3	Lung Biopsy
Lung - ResectionsHIS5Lung Tumour Resection +/- NodesNeurosurgeryHIS3Brain BiopsyNeurosurgeryHIS5 + HIS5Muscle BiopsyNeurosurgeryHIS3Pituitary Gland - ResectionNeurosurgeryHIS3Spinal Tumour BiopsyNeurosurgeryHIS3Spinal Tumour ResectionNeurosurgeryHIS4VertebreaOpthalmicHIS1Conjunctival BiopsyOpthalmicHIS1CorneaOpthalmicHIS4Globe / Removal of EyeOpthalmicHIS2Lacrimal Gland Biopsy / ExcisionOpthalmicHIS1Orbit Contents of EyeOrthopaedicHIS1Bone BiopsyOrthopaedicHIS2Bone CurrettingsOrthopaedicHIS2Bursa		HIS3	Lung Lesion Small Wedge Resection
NeurosurgeryHIS3Brain BiopsyNeurosurgeryHIS3Brain ResectionNeurosurgeryHIS5+HIS5Muscle BiopsyNeurosurgeryHIS3Pituitary Gland – ResectionNeurosurgeryHIS3Spinal Tumour BiopsyNeurosurgeryHIS3Spinal Tumour ResectionNeurosurgeryHIS4VertebreaOpthalmicHIS1Conjunctival BiopsyOpthalmicHIS1CorneaOpthalmicHIS4Globe / Removal of EyeOpthalmicHIS2Lacrimal Gland Biopsy / ExcisionOpthalmicHIS1Orbit Contents of EyeOrthopaedicHIS1Bone BiopsyOrthopaedicHIS2Bone CurrettingsOrthopaedicHIS2Bursa	Lung – Resections	HIS5+HIS5	Lung Resection
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         Opthalmic       HIS1       Conjunctival Biopsy         Opthalmic       HIS1       Cornea         Opthalmic       HIS4       Globe / Removal of Eye         Opthalmic       HIS2       Lacrimal Gland Biopsy / Excision         Opthalmic       HIS1       Orbit Contents of Eye         Orthopaedic       HIS1       Bone Biopsy         Orthopaedic       HIS2       Bone Currettings         Orthopaedic       HIS2       Bursa	Lung – Resections	HIS5	Lung Tumour Resection +/- Nodes
Neurosurgery HIS3 Pituitary Gland – Resection  Neurosurgery HIS3 Spinal Tumour Biopsy  Neurosurgery HIS3 Spinal Tumour Resection  Neurosurgery HIS4 Vertebrea  Opthalmic HIS1 Conjunctival Biopsy  Opthalmic HIS4 Globe / Removal of Eye  Opthalmic HIS2 Lacrimal Gland Biopsy / Excision  Opthalmic HIS1 Orbit Contents of Eye  Opthalmic HIS1 Bone Biopsy  Orthopaedic HIS2 Bone Currettings  Orthopaedic HIS2 Bursa	Neurosurgery	HIS3	Brain Biopsy
Neurosurgery HIS3 Pituitary Gland – Resection  Neurosurgery HIS3 Spinal Tumour Biopsy  Neurosurgery HIS3 Spinal Tumour Resection  Neurosurgery HIS4 Vertebrea  Opthalmic HIS1 Conjunctival Biopsy  Opthalmic HIS1 Cornea  Opthalmic HIS4 Globe / Removal of Eye  Opthalmic HIS2 Lacrimal Gland Biopsy / Excision  Opthalmic HIS1 Orbit Contents of Eye  Orthopaedic HIS1 Bone Biopsy  Orthopaedic HIS2 Bone Currettings  Orthopaedic HIS2 Bursa	Neurosurgery	HIS3	Brain Resection
Neurosurgery       HIS3       Spinal Tumour Biopsy         Neurosurgery       HIS4       Vertebrea         Opthalmic       HIS1       Conjunctival Biopsy         Opthalmic       HIS1       Cornea         Opthalmic       HIS4       Globe / Removal of Eye         Opthalmic       HIS2       Lacrimal Gland Biopsy / Excision         Opthalmic       HIS1       Orbit Contents of Eye         Orthopaedic       HIS1       Bone Biopsy         Orthopaedic       HIS2       Bone Currettings         Orthopaedic       HIS2       Bursa	Neurosurgery	HIS5+HIS5	Muscle Biopsy
Neurosurgery HIS3 Spinal Tumour Resection  Neurosurgery HIS4 Vertebrea  Opthalmic HIS1 Conjunctival Biopsy  Opthalmic HIS4 Globe / Removal of Eye  Opthalmic HIS2 Lacrimal Gland Biopsy / Excision  Opthalmic HIS1 Orbit Contents of Eye  Orthopaedic HIS1 Bone Biopsy  Orthopaedic HIS2 Bone Currettings  Orthopaedic HIS2 Bursa	Neurosurgery	HIS3	Pituitary Gland – Resection
Neurosurgery HIS4 Vertebrea  Opthalmic HIS1 Conjunctival Biopsy  Opthalmic HIS4 Globe / Removal of Eye  Opthalmic HIS2 Lacrimal Gland Biopsy / Excision  Opthalmic HIS1 Orbit Contents of Eye  Orthopaedic HIS1 Bone Biopsy  Orthopaedic HIS2 Bone Currettings  Orthopaedic HIS2 Bursa	Neurosurgery	HIS3	Spinal Tumour Biopsy
Opthalmic HIS1 Conjunctival Biopsy  Opthalmic HIS1 Cornea  Opthalmic HIS4 Globe / Removal of Eye  Opthalmic HIS2 Lacrimal Gland Biopsy / Excision  Opthalmic HIS1 Orbit Contents of Eye  Orthopaedic HIS1 Bone Biopsy  Orthopaedic HIS2 Bone Currettings  Orthopaedic HIS2 Bursa	Neurosurgery	HIS3	Spinal Tumour Resection
Opthalmic HIS1 Cornea  Opthalmic HIS4 Globe / Removal of Eye  Opthalmic HIS2 Lacrimal Gland Biopsy / Excision  Opthalmic HIS1 Orbit Contents of Eye  Orthopaedic HIS1 Bone Biopsy  Orthopaedic HIS2 Bone Currettings  Orthopaedic HIS2 Bursa	Neurosurgery	HIS4	Vertebrea
Opthalmic HIS4 Globe / Removal of Eye  Opthalmic HIS2 Lacrimal Gland Biopsy / Excision  Opthalmic HIS1 Orbit Contents of Eye  Orthopaedic HIS1 Bone Biopsy  Orthopaedic HIS2 Bone Currettings  Orthopaedic HIS2 Bursa	Opthalmic	HIS1	Conjunctival Biopsy
Opthalmic HIS2 Lacrimal Gland Biopsy / Excision  Opthalmic HIS1 Orbit Contents of Eye  Orthopaedic HIS1 Bone Biopsy  Orthopaedic HIS2 Bone Currettings  Orthopaedic HIS2 Bursa	Opthalmic	HIS1	Cornea
Opthalmic HIS1 Orbit Contents of Eye Orthopaedic HIS1 Bone Biopsy Orthopaedic HIS2 Bone Currettings Orthopaedic HIS2 Bursa	Opthalmic	HIS4	Globe / Removal of Eye
Orthopaedic HIS1 Bone Biopsy Orthopaedic HIS2 Bone Currettings Orthopaedic HIS2 Bursa	Opthalmic	HIS2	Lacrimal Gland Biopsy / Excision
Orthopaedic     HIS2     Bone Currettings       Orthopaedic     HIS2     Bursa	Opthalmic	HIS1	Orbit Contents of Eye
Orthopaedic HIS2 Bursa	Orthopaedic	HIS1	Bone Biopsy
25.00	Orthopaedic	HIS2	Bone Currettings
Orthopaedic HIS2 Duputrenes Contracture	Orthopaedic	HIS2	Bursa
	Orthopaedic	HIS2	Duputrenes Contracture

## Histopathology

CATEGORY	CODE	TISSUE SAMPLE
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
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

## Histopathology

CATEGORY	CODE	TISSUE SAMPLE
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)



## **Non-invasive prenatal testing (NIPT)**

Non-invasive prenatal testing (NIPT) screens for the presence of specific chromosome disorders in the developing fetus. The test analyses fragments of cell-free DNA in maternal plasma that have been released from both maternal and placental cells.

By analysing the proportions of cell-free DNA fragments derived from different chromosomes or chromosome regions, NIPT can screen for the presence or absence of specific chromosome disorders.

NIPT is more accurate than first trimester maternal serum screening and ultrasound in identifying pregnancies with or without these disorders.

TDL Genetics uses the NIPT assay VeriSeq NIPT Solution v2, which is manufactured by Illumina and is processed at our laboratory in London.

# Targeted screening for specific common chromosome disorders

Our NIPT assay is designed to screen for:

 Trisomy 21 (Down syndrome), which is associated with moderate to severe intellectual disability, congenital heart defects and other malformations;

- Trisomy 18 (Edwards syndrome) and trisomy 13 (Patau syndrome), which are associated with severe brain and cardiac malformations. There is a high risk of stillbirth or death during infancy; and
- Sex chromosome aneuploidy (abnormalities in the number of X or Y chromosomes), which can be associated with malformations and infertility, Turner syndrome (45,X) and Klinefelter syndrome (47,XXY). Triple X syndrome and XYY syndrome can also be detected. This screen is optional (no additional cost).

In addition, NIPT can also assess fetal sex. This is optional (no additional cost).

NIPT does not screen for non-chromosome disorders, familial mutations, malformations, fetal growth or fetal viability.



Find out more about NIPT on the TDL website:

www.tdlpathology.com/noninvasive-prenatal-testing/

TEST	CODE	SAMPLE REQS	TAT	PAGE
1,25 Vitamin D	D3	<b>B</b> *	5-8 days	151
1p36 Deletion Syndrome – karyotype + CGH	KARY,	CVS / AF / (1) 9	12-17 days	116
	FISH			
2-Butanone GC	BUTA	RU	7 days	166
5 HIAA	RU5H	PU (collect on acid) <sup>1</sup>	5 days	29
5' Nucleotidase	5NT	В	5 days	29
6-Thioguanine Nucleotides	TGN	AA	2 weeks	29
7 STI Profile by PCR (7 tests from 1 Sample)	DL12	FCRU / PCR Swab / TPV	2 days	72, 77
7 STI Profile by PCR (7 tests from 1 Sample) (Self-collect)	DL12	Aptima urine or multisite swab	2 days	72, 158
7 STI Profile by PCR (7 tests from	PP12	TPV	2 days	170
1 Sample) (Thin Prep)				
11 Deoxycorticosterone	DEOX	8	10 days	57
11 Deoxycortisol	11DC	(Frozen)	10 days	57
16S rRNA Bacterial Gene	16S	J	1 week	47
17 Hydroxyprogesterone	170H	В	5 days	57
18S rRNA Fungal Gene	18S	J	1 week	47
21 Hydroxylase Ab's	21HA	(Frozen)	10 days	29
21-Hydroxylase Deficiency	GENE	A 9,11	5 weeks	116
(Congenital Adrenal Hyperplasia CYP21A2) 22q11 & 10p14 deletion	DGB	CVS / AF / (A) 9	E days	116
(Di George Syndrome) – BOBs only	Dub	GVS/AF/	5 days	110
Acetone – Blood	ACTB	(A) or (1)	2 weeks	166
Acetone – Urine	ACTU	RU	5 days	166
Acetylcholine Receptor Autoantibodies	ACRA	<b>B</b> 4	5 days	29
Achromatopsia NGS Panel	GENE	<b>A A</b> <sup>9</sup>	5 weeks	116
Acid Phosphatase – Total	APT	<b>B</b>	5 days	29
ACTH (Adrenocorticotropic Hormone)	ACTH	(EDTA on ice, Plasma, spun and frozen within 2 hours) <sup>41</sup>	1 day	57
Activated Protein C Resistance	APCR	C (Frozen) <sup>4,18</sup>	3 days	42
Acute Viral Hepatitis Screen	AHSC	В	1 day	83, 89
ADAMTS-13 Activity	CP13	C (Frozen) <sup>4,18</sup>	3 days	42
ADAMTS-13 Antibody	A13A	(Frozen) <sup>9,18</sup>	2 weeks	42
Adenomatous Polyposis NGS Panel	GENE	<b>A</b> 9	4 weeks	116
Adenosine Deaminase	AD	(A) / (B) / Fluid	3 weeks	29
Adenovirus by PCR	ADV	(A) / PCR / VS	7 days	102
Adiponectin	ADIP	<b>B</b>	2 weeks	29
Adrenal Cortex Antibodies	ACTX	B	2 days	83
Aicardi-Goutières Syndrome NGS Panel	GENE	<b>A A</b> <sup>9</sup>	5 weeks	116
Alagille Syndrome NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	116
Albumin	ALB	B	1 day	29
Alcohol (Medical)	ALC0	<b>G</b> <sup>1</sup>	1 day	29
Alcohol (Urine)	UALC	RU	1 day	29
<u> </u>				

TEST	CODE	SAMPLE REQS	TAT	PAGE
Alcohol Profile	AP	<b>ABBG</b>	5-7 days	163, 166
Alcohol Profile 2	ALCP	A B B G RU	5-7 days	163, 166
Aldolase	ALD0	B	5 days	29
Aldosterone	ALDN	(A) or (B)	5 days	57
Aldosterone (Urine)	UALD	PU	5 days	57
ALEX <sup>2</sup> Allergy Test (Self-collect)	ALEX	(TDL Tiny)	3-4 days	141, 158
ALEX <sup>2</sup> Allergy Test (Venous)	ALEX	(Serum)	3-4 days	141, 146
Alkaline Phosphatase	ALP	B	1 day	29
Alkaline Phosphatase Isoenzymes	APIE	B	5 days	29
Allergic Rhinitis/Asthma Profile	ALRN	88	2 days	141, 146
Allergy – Individual Allergens	ALLE	B	2 days	141
Allergy – 5 x Single Individual Allergens	5AL	B	2 days	141
Allergy – 10 x Single Individual Allergens	10AL	B	2 days	141
Allergy Profile 1 (Food & Inhalants)	1A	88	2 days	141, 144
Allergy Profile 2 (UK Aero Allergen)	2A	88	2 days	141, 144
Allergy Profile 3 (Food)	3A	88	2 days	141, 144
Allergy Profile 4 (Nuts & Seeds)	4A	88	2 days	141, 144
Allergy Profile 5 (Children's Panel)	5A	88	2 days	141, 144
Allergy Profile 6 (Shellfish)	6A	88	2 days	141, 145
Allergy Profile 7 (Finfish)	7A	88	2 days	141, 145
Allergy Profile 8 (Cereal – singles)	8A	88	2 days	141, 145
Allergy Profile 9 (Antibiotics)	9A	88	2 days	141, 145
Allergy Profile 10 (Insects)	10A	88	2 days	141, 145
Allergy Profile 11 (Combined Shellfish/Finfish)	11A	88	2 days	141, 145
Allergy Profile 12 (Milk & Milk Proteins)	12A	88	2 days	141, 145
Allergy Profile 13 (Stone fruit/Rosaceae family)	13A	88	2 days	141, 146
Alpha Gal Components (related to red meat)	ZZ37	B	2 days	142
Alpha Thalassaemia – multiplex PCR	GENE	<b>A</b> 9	3 weeks	116
for common large deletions				
Alpha-1-Antitrypsin (Serum)	A1AT	B	1 day	29
Alpha-1-Antitrypsin (Stool)	A1AF	RF	10 days	29
Alpha-1-Antitrypsin Genotype – PI*M, PI*S, PI*Z	GENE	<b>A</b> 9	3 weeks	29, 116
Alpha-1-Glycoprotein	OROS	(Frozen)	5 days	29
Alpha-1-Microglobulin	A1MG	<b>RU</b> 1,22	10 days	29
Alpha-2-Macroglobulins	A2MG	B	5 days	29
Alpha-Fetoprotein	AFP	B	1 day	29, 57,
				106
Alport Syndrome NGS Panel – full sequencing	GENE	<b>A A</b> <sup>9</sup>	5 weeks	116
with deletions and duplications				
ALT (Alanine Aminotransferase) (SGPT)	ALT	В	1 day	29
Alternaria Components	ZZ1	B	2 days	142
Aluminium (Blood)	ALUM	<b>(</b> )	7 days	29, 165
Aluminium (Urine)	ALUU	RU	1-2 weeks	165

TOES, SHBG, FAI) (Self-collect) CHANGE	TEST	CODE	SAMPLE REQS	TAT	PAGE
Amilkacin Level (State dose)         AMIK         3 ⁴         1 day         137           Amino Acid (EDTA Plasma)         AMIN         ② (Frozen EDTA Plasma)         7 days         29           Amino Acid Quantitative (Urine)         UAAQ         RU (Frozen)         7 days         29           Aminolevulinic Acid (Urine)         RUAL         100mis PU         5 days         138           Ammonia         AMTR         4 ⁴         5 days         138           Ammonia         AMTR         4 °         5 days         138           Amoebic (E. histolytica) Antibodies         AFAT         6 °         1 day         29           Ampletamines – Blood         AMPB         3 ©         5 days         163           Amylase (Self-collect)         AMY         2 (TDL Tiny)         1 day         29           Amylase (Venous)         AMY         2 (TDL Tiny)         1 day         29           Amylase (Venous)         AMY         0 (TDL Tiny)         1 day         29           Amylase (Venous)         AMY         0 (TDL Tiny)         1 day         29           Amylase (Venous)         AMY         0 (TDL Tiny)         1 day         29           Amylase (Venous)         AMY         0 (TDL Tiny)	. , , , ,	TAME	(TDL Tiny)	1 day	57, 158
Amino Acid (EDTA Plasma)         AMIN         ⑥ (Frozen EDTA Plasma)         7 days         29           Amino Acid Quantitative (Urine)         UAAQ         RU (Frozen)         7 days         29           Amino Acid Quantitative (Urine)         RUAL         100mls PU         5 days         29           Amitriptyline         AMTR         0⁴         5 days         128           Ammoenia         AMMO         0⅙ (Frozen) 15         1 day         29           Amoebic (E. histolytica) Antibodies         AFAT         3         1 week         92           Amoebic (E. histolytica) PCR         AMAB         RF         2 days         92           Amphase Self-collect)         AMY         3 (TDL Tirry)         1 day         29           Amylase (Urine)         UAMY         CU         1 day         29           Amylase (Venous)         AMY         3 (TDL Tirry)         1 day         29           Amylase (Venous)         AMY         3 (TDL Tirry)         1 day         29           Amylase (Venous)         AMY         3 (TDL Tirry)         1 day         29           Amylase (Venous)         AMY         6 (TDL Tirry)         1 day         29           Amylase (Venous)         AMY         6 (TDL	Amenorrhoea Profile (Venous)	AMEN	B	1 day	57, 62
Amino Acid Quantitative (Urine)         UAAQ         RU (Frozen)         7 days         29           Aminolevulinic Acid (Urine)         RUAL         100mls PU         5 days         23           Amitriptyline         AMTR         4         5 days         138           Ammonia         AMMO         4 (Frozen)¹5¹         1 day         29           Amoebic (E. histolytica) Antibodies         AFAT         0         1 week         92           Amoebic (E. histolytica) PCR         AMAG         RF         2 days         92           Amplases (Gelf-collect)         AMY         0 (TDL Tiny)         1 day         29,158           Amylase (Venous)         AMY         0 (TDL Tiny)         1 day         29,358           Amylase (Venous)         AMY         0 (TDL Tiny)         1 day         29           Amylase (Venous)         AMY         0 (TDL Tiny)         1 day         29           Amylase (Venous)         AMY         0 (TDL Tiny)         1 day         29           Amylase (Venous)         AMY         0 (TDL Tiny)         1 day         29           Amylase (Venous)         AMY         0 (TDL Tiny)         1 day         29           Amylase (Venous)         AMY         0 (TDL Tiny)	Amikacin Level (State dose)	AMIK	<b>B</b> 4	1 day	137
Aminolevulinic Acid (Urine)         RUAL         100mls PU         5 days         29           Amitriptyline         AMTR         0 4         5 days         138           Ammonia         AMMO         0 (Frozen)¹5         1 day         29           Amoebic (E. histolytica) Antibodies         AFAT         3         1 week         92           Ampletamines – Blood         AMPB         3 G         5 days         163           Amylase (Self-collect)         AMY         0 (TDL Tiny)         1 day         29,158           Amylase (Urine)         UAMY         CU         1 day         29           Amylase (Venous)         AMY         0         1 day         29           Amylase (Senenzymes         AMYI         0         1 day         29           Amylase (Venous)         AMY         0         1 day         29           Amylase (Senenzymes         AMYI         0         1 day         29           Amylase (Urine)         UAMY         0         1 day         29           Amylase (Urine)         AMY         0         1 day         29           Amyloidosis (Amyloid A Protein)         SAA         0         5 days         29           Anjecturin (Comip	Amino Acid (EDTA Plasma)	AMIN	(Frozen EDTA Plasma)	7 days	29
Amitriptyline         AMTR         3 days         138           Ammonia         AMMO         3 (Frozen)¹5*         1 day         29           Amoebic (E, histolytica) Antibodies         AFAT         0         1 week         92           Amoebic (E, histolytica) PCR         AMAG         RF         2 days         92           Amplase (Belf-collect)         AMY         0 (TDL Tiny)         1 day         29,158           Amylase (Urine)         UAMY         CU         1 day         29,158           Amylase (Venous)         AMY         0 (TDL Tiny)         1 day         29,158           Amylase (Seenzymes         AMY         0 (TDL Tiny)         1 day         29,158           Amylase (Venous)         AMY         0 (TDL Tiny)         1 day         29,158           Amylase (Seenzymes         AMY         0 (TDL Tiny)         1 day         29,158           Amylase (Seenzymes         AMY         0 (TDL Tiny)         1 day         29           Amylase (Seenzymes         AMY         0 (TDL Tiny)         1 day         29           Amylase (Benezymes         AMY         0 (TDL Tiny)         1 day         29           Amyloides (Brechall Antiboolis (Amyloid A Protein)         AMA         0 (Tolay Seen S	Amino Acid Quantitative (Urine)	UAAQ	RU (Frozen)	7 days	29
Ammonia         AMMO         ⑥ (Frozen)¹5         1 day         29           Amoebic (E. histolytica) Antibodies         AFAT         ③         1 week         92           Amoebic (E. histolytica) PCR         AMAG         RF         2 days         92           Amphate (E. histolytica) PCR         AMAG         RF         2 days         92           Amphate (Electic)         AMY         ⑥ (TDL Tiny)         1 day         29         158           Amylase (Self-collect)         AMY         ⑥ (TDL Tiny)         1 day         29         158           Amylase (Venous)         AMY         0         1 day         29         418         24         29           Amylase (Venous)         AMY         0         1 day         29         414         44         29         414         44         42         29         418         44 </th <th>Aminolevulinic Acid (Urine)</th> <th>RUAL</th> <th>100mls <b>PU</b></th> <th>5 days</th> <th>29</th>	Aminolevulinic Acid (Urine)	RUAL	100mls <b>PU</b>	5 days	29
Amoebic (E. histolytica) Antibodies         AFAT         ①         1 week         92           Amoebic (E. histolytica) PCR         AMAG         RF         2 days         92           Amphetamines – Blood         AMPB         ①         5 days         163           Amylase (Self-collect)         AMY         ① (TDL Tiny)         1 day         29,158           Amylase (Urine)         UAMY         CU         1 day         29           Amylase (Venous)         AMY         ①         1 day         29           Amylase Isoenzymes         AMYI         ①         5 days         29           Amyloidosis (Amyloid A Protein)         SAA         ①         5 days         29           Anaemia Profile         ANAE         ②         ③         2 days         41,44           Anafranii (Clomipramine)         CHLO         ②         7 days         138           ANCA (Anti-Neutrophii (Zytoplasmic Abs)         ANCA         ①         2 days         83           Androstanedioi Glucuronide         ANDR         ①         ①         1 day         5,62           Androstanedioi Glucuronide         ANDR         ①         ①         3 weeks         29           Androstanedioi Enzyme – CSF         A	Amitriptyline	AMTR	<b>A</b> 4	5 days	138
Amoebic (E. histotytica) PCR         AMAG         RF         2 days         92           Amphetamines – Blood         AMPB         3 G         5 days         163           Amylase (Self-collect)         AMY         (In Litry)         1 day         29,158           Amylase (Venous)         AMY         (Unit of the control of the cont	Ammonia	AMM0	(Frozen) <sup>15</sup>	1 day	29
Amphetamines – Blood         AMPB         ③ ⑤         5 days         163           Amylase (Self-collect)         AMY         ④ (TDL Tiny)         1 day         29, 158           Amylase (Urine)         UAMY         CU         1 day         29           Amylase (Venous)         AMY         ⑥         1 day         29           Amylase (Self-collect)         AMY         ⑥         1 day         29           Amylase (Urine)         U. L.         Ø         1 day         29           Amylase (Ivine)         AMY         ⑥         1 day         29           Amyloidesis (Amyloid A Protein)         SAA         ⑥         2 days         41,44           Anaeria Profile         ANAE         ⑥         ⑥         2 days         43           ANCA (Anti-Neutrophil Cytoplasmic Abs)         ANCA         ⑥         2 days         33           Androstanediol Elucuronide         ANDB         ⑥         3 weeks         29           Andr	Amoebic (E. histolytica) Antibodies	AFAT	B	1 week	92
Amylase (Self-collect)         AMY         ③ (TDL Tiny)         1 day         29, 158           Amylase (Urine)         UAMY         CU         1 day         29           Amylase (Venous)         AMY         ①         1 day         29           Amylase Isoenzymes         AMYI         ①         5 days         29           Amyloidosis (Amyloid A Protein)         SAA         ①         5 days         29           Anaeriania Profile         ANAE         ②         ②         2 days         41,44           Anafranial (Clomipramine)         CHLO         ③         7 days         138           Andro Alanti-Neutrophil Cytoplasmic Abs)         ANCA         ②         2 days         83           Androfostanediol Glucuronide         ANDG         ③         3 weeks         29           Androstanediol Glucuronide         ANDG         ③         3 weeks         29           Androstanediole Blucuronide         ANDG         ⑤         3 days         57           Angiotensin Converting Enzyme         ACE         ⑤         1 day         30           Angiotensin Converting Enzyme – CSF         ACEF         CSF (Frozen)         2 weeks         30           Angiotensin Converting Enzyme – CSF         ACEF <th>Amoebic (E. histolytica) PCR</th> <th>AMAG</th> <th>RF</th> <th>2 days</th> <th>92</th>	Amoebic (E. histolytica) PCR	AMAG	RF	2 days	92
Amylase (Urine)         UAMY         CU         1 day         29           Amylase (Venous)         AMY         1 day         29           Amylase Isoenzymes         AMYI         5 days         29           Amyloidosis (Amyloid A Protein)         SAA         3 5 days         29           Anaemia Profile         ANAE         6 2 days         41,44           Anafranii (Clomipramine)         CHLO         7 days         138           ANCA (Anti-Neutrophil Cytoplasmic Abs)         ANCA         2 days         83           Androstanediol Glucuronide         ANDP         6         2 days         83           Androstanediol Glucuronide         ANDR         3 weeks         29           Androstenedione         ANDR         3 weeks         29           Androstenedione         ANDR         3 weeks         29           Angiotensin Converting Enzyme         ACE         1 day         30           Angiotensin Converting Enzyme – CSF         ACEF         CSF (Frozen)         2 weeks         30           Angiotensin I         ANC2         (§ (Frozen plasma)         2 weeks         30           Anti-Actin Antibodies         AAA         3 days         41,44           Anti-Basal Ganglia Antibod	Amphetamines – Blood	AMPB	BB	5 days	163
Amylase (Venous)         AMY         3         1 day         29           Amylase Isoenzymes         AMYI         3         5 days         29           Amyloidosis (Amyloid A Protein)         SAA         10         5 days         29           Anaemia Profile         ANAE         10         5 days         41,44           Anafranii (Clomipramine)         CHLO         7 days         138           ANCA (Anti-Neutrophil Cytoplasmic Abs)         ANCA         2 days         83           Andro Actin Anti-Neutrophil Cytoplasmic Abs)         ANCA         3         2 days         83           Andro Stanediol Glucuronide         ANDR         3         3         44         44           Androstanediol Glucuronide         ANDR         3         3         449         57,62           Androstanediol Glucuronide         ANDR         3         3         48eks         29           Androstanediol Glucuronide         ANDR         3         3         48eks         29           Androstanediolige Liver Profile         ANDR         3         48eks         29           Angiotensin Converting Enzyme - CSF         ACE         CSF (Frozen)         2 weeks         30           Antenatal Profile <th< th=""><th>Amylase (Self-collect)</th><th>AMY</th><th>(TDL Tiny)</th><th>1 day</th><th>29, 158</th></th<>	Amylase (Self-collect)	AMY	(TDL Tiny)	1 day	29, 158
Amylase Isoenzymes         AMYI         3         5 days         29           Amyloidosis (Amyloid A Protein)         SAA         3         5 days         29           Anaemia Profile         ANAE         3         6         2 days         41, 44           Anafranil (Clomipramine)         CHLO         3         7 days         138           ANCA (Anti-Neutrophil Cytoplasmic Abs)         ANCA         3         2 days         83           Andropause Profile         ANDP         3         1 day         57, 62           Androstanediol Glucuronide         ANDG         3         3 weeks         29           Androstanediol Glucuronide         ANDR         3         4 weeks         30           Anti-Rosin Converting Enzyme         CSF         CSF (Forze	Amylase (Urine)	UAMY	CU	1 day	29
Amyloidosis (Amyloid A Protein)         SAA         3         5 days         29           Anaemia Profile         ANAE         3         2 days         41, 44           Anafranil (Clomipramine)         CHLO         7 days         138           ANCA (Anti-Neutrophil Cytoplasmic Abs)         ANCA         2 days         83           Androstanediol Glucuronide         ANDP         1 day         57, 62           Androstanediol Glucuronide         ANDR         3 weeks         29           Androstenedione         ANDR         5 days         57           Angiotensin Converting Enzyme         ACE         1 day         30           Angiotensin Converting Enzyme – CSF         ACEF         CSF (Frozen)         2 weeks         30           Angiotensin II         ANG2         ③ (Frozen plasma)         2 weeks         30           Antenatal Profile         ANTE         ③ ③ 3 ② ③ ③ ⑤         3 days         41, 44           Anti-Basal Ganglia Antibodies         AAA         3         5 days         83           Anti-CP Antibodies         ABGA         3 weeks         83           Anti-Wro Cytosol Antibodies         ALCA         3 days         83           Anti-MUSK Antibodies         MUSK         3 weeks <th>Amylase (Venous)</th> <th>AMY</th> <th>B</th> <th>1 day</th> <th>29</th>	Amylase (Venous)	AMY	B	1 day	29
Anaemia Profile         ANAE         ② ③ ⑤         2 days         41, 44           Anafranil (Clomipramine)         CHLO         3         7 days         138           ANCA (Anti-Neutrophil Cytoplasmic Abs)         ANCA         3         2 days         83           Androstanediol Glucuronide         ANDP         3         1 day         57, 62           Androstanediole Glucuronide         ANDR         3         3 weeks         29           Androstenedione         ANDR         3         5 days         57           Angiotensin Converting Enzyme         ACE         3         1 day         30           Angiotensin Converting Enzyme – CSF         ACEF         CSF (Frozen)         2 weeks         30           Angiotensin II         ANG2         ③ (Frozen plasma)         2 weeks         30           Antenatal Profile         ANTE         ③ ③ ③ ③ ⑤ ⑥ ③ 3 days         41, 44           Anti-Actin Antibodies         AAA         ⑥ ⑤ ③ ③ ⑥ ⑥ ③ 3 days         83           Anti-CCP Antibodies         ABGA         ⑥ ③ 3 weeks         83           Anti-Liver Cytosol Antibodies         ALCA         ⑥ ⑤ 3 weeks         83           Anti-Hous (Intibodies         ALCA         ⑥ ⑤ 6 days         83	Amylase Isoenzymes	AMYI	B	5 days	29
Anafranil (Clomipramine)         CHLO         3         7 days         138           ANCA (Anti-Neutrophil Cytoplasmic Abs)         ANCA         3         2 days         83           Andropause Profile         ANDP         3         1 day         57, 62           Androstanediol Glucuronide         ANDG         3 weeks         29           Androstenedione         ANDR         3         5 days         57           Angiotensin Converting Enzyme         ACE         3         1 day         30           Angiotensin Converting Enzyme – CSF         ACEF         CSF (Frozen)         2 weeks         30           Angiotensin II         ANG2         3 (Frozen plasma)         2 weeks         30           Antenatal Profile         ANTE         3         3 days         41, 44           Anti-Actin Antibodies         AAA         3         3 (Frozen plasma)         2 weeks         30           Anti-Basal Ganglia Antibodies         ABGA         3         3 weeks         83           Anti-Liver Cytosol Antibodies         ALCA         3         3 weeks         83           Anti-Musk Antibodies         ALCA         3         3 weeks         83           Glycoprotein] Antibodies         MUSK         3<	Amyloidosis (Amyloid A Protein)	SAA	B	5 days	29
ANCA (Anti-Neutrophil Cytoplasmic Abs)         ANCA ③         2 days         83           Andropause Profile         ANDP ③         1 day         57, 62           Androstanediol Glucuronide         ANDG ⑤         3 weeks         29           Androstenedione         ANDR ⑥         5 days         57           Angiotensin Converting Enzyme         ACE ⑥         1 day         30           Angiotensin II         ANG2 ⑥         (Frozen)         2 weeks         30           Antenatal Profile         ANTE ⑥         ⑥         3 days         41,44           Anti-Actin Antibodies         AAA         3 seeks         83           Anti-CCP Antibodies         ABGA ⑤         3 weeks         83           Anti-Liver Cytosol Antibodies         ALCA ⑥         3 weeks         83           Anti-MOG [Myelin Oligodendrocyte         AMOG ⑥         3 weeks         83           Anti-Nuclear Antibodies         MUSK ⑥         2 weeks         83           Anti-Phosphatidylserine Antibodies         PHTS ⑥         5 days         83           Anti-Phosphatidylserine Antibodies         PHTS ⑥         5 days         83           Anti-Sta (Soluble Liver Antigen) Abs         LSA ⑥         6 weeks         83           Anti-Stap	Anaemia Profile	ANAE	A A B	2 days	41, 44
Andropause Profile  ANDP 3 3 4 3 4 3 4 3 4 3 4 3 4 3 4 3 4 3 4	Anafranil (Clomipramine)	CHLO	A	7 days	138
Androstanediol Glucuronide  ANDG G 3 weeks 29  Androstenedione  ANDR G 5 days 57  Angiotensin Converting Enzyme  ACE G 1 day 30  Angiotensin Converting Enzyme – CSF ACEF CSF (Frozen) 2 weeks 30  Angiotensin II ANG2 (Frozen plasma) 2 weeks 30  Antenatal Profile ANTE (S 3 3 6 6 3 3 days 41, 44  Anti-Actin Antibodies  AAA G 5 days 83  Anti-CCP Antibodies  ARBGA G 3 weeks 83  Anti-CP Antibodies ALCA G 5 days 83  Anti-Liver Cytosol Antibodies  ANDG G 3 weeks 83  Anti-MOG [Myelin Oligodendrocyte AMOG G 3 weeks 83  Anti-MOG [Myelin Oligodendrocyte AMOG G 3 weeks 83  Anti-MUSK Antibodies  MUSK G 2 weeks 83  Anti-Nuclear Antibodies (titre & pattern) ANAB G 2 days 83  Anti-Nuclear Antibodies PHTS G 5 days 83  Anti-Phosphatidylserine Antibodies PHTS G 6 weeks 83  Anti-SLA (Soluble Liver Antigen) Abs LSA G 5 days 83  Anti-Staphylolysin Titre (SGOT) ASTT G 3 days 83  Anti-Streptolysin Titre (SGOT) ASTT G 3 days 83  Anti-Streptolysin Titre/ASOT ASL G 5 weeks 83  Anti-Sulfatide Antibodies ASA G 5 weeks 83  Anti-Sulfatide Antibodies ASA G 5 weeks 83  Anti-Sulfatide Antibodies ASA G 5 weeks 83	ANCA (Anti-Neutrophil Cytoplasmic Abs)	ANCA	B	2 days	83
Androstenedione  ANDR G  Angiotensin Converting Enzyme  ACE G  ACEF CSF (Frozen)  ANG2 (Apgiotensin II)  ANG3 (Apgiotensin II)  ANG3 (Apgiotensin II)  ANG4 (Apgiotensin II)  ANG5 (Apgiotensin II)  ANG6 (Apgiotensin II)  ANG9 (Apg	Andropause Profile	ANDP	88	1 day	57, 62
Angiotensin Converting Enzyme ACE © 1 day 30 Angiotensin Converting Enzyme – CSF ACEF CSF (Frozen) 2 weeks 30 Angiotensin II ANG2 (Frozen plasma) 2 weeks 30 Antenatal Profile ANTE (A) (A) 30 (Frozen plasma) 2 weeks 30 Antenatal Profile ANTE (A) (A) 30 (B) (Frozen plasma) 3 days 41,44 Anti-Actin Antibodies AAA (B) 5 days 83 Anti-Actin Antibodies ABGA (B) 3 weeks 83 Anti-CCP Antibodies CCP (B) 2 days 83 Anti-Liver Cytosol Antibodies ALCA (B) 5 days 83 Anti-Liver Cytosol Antibodies ALCA (B) 5 days 83 Anti-MOG [Myelin Oligodendrocyte AMOG (B) 3 weeks 83 Anti-MUSK Antibodies MUSK (B) 2 weeks 83 Anti-Nuclear Antibodies (titre & pattern) ANAB (B) 2 days 83 Anti-Phosphatidylserine Antibodies PHTS (B) 5 days 83 Anti-Phospholipase A2 Receptor AA2R (B) 6 weeks 83 Anti-SLA (Soluble Liver Antigen) Abs LSA (B) 5 days 83 Anti-Staphylolysin Titre (SGOT) ASTT (B) 3 days 83 Anti-Streptolysin Titre (ASOT) ASLT (B) 2 days 83 Anti-Streptolysin Titre/ASOT ASLT (B) 2 days 83 Anti-Sulfatide Antibodies ASA (B) 5 weeks 83 Anti-Sulfatide Antibodies ASA (B) 5 weeks 83	Androstanediol Glucuronide	ANDG	B	3 weeks	29
Angiotensin Converting Enzyme – CSF ACEF CSF (Frozen) 2 weeks 30 Angiotensin II ANG2 ((Frozen plasma)) 2 weeks 30 Antenatal Profile ANTE ((Frozen plasma)) 2 weeks 30 Antenatal Profile ANTE ((Frozen plasma)) 2 weeks 31 Anti-Actin Antibodies AAA ((Frozen plasma)) 3 days 41,44 Anti-Actin Antibodies AAA ((Frozen plasma)) 5 days 83 Anti-Basal Ganglia Antibodies ABGA ((Frozen plasma)) 3 weeks 83 Anti-CCP Antibodies ((Frozen plasma)) 2 days 83 Anti-Liver Cytosol Antibodies ALCA ((Frozen plasma)) 3 weeks 83 Anti-MOG [Myelin Oligodendrocyte AMOG ((Frozen plasma)) 3 weeks 83 Anti-MUSK Antibodies MUSK ((Frozen plasma)) 3 weeks 83 Anti-Musclear Antibodies ((Frozen plasma)) 4 weeks 83 Anti-Nuclear Antibodies ((Frozen plasma)) 5 days 83 Anti-Phosphatidylserine Antibodies PHTS ((Frozen plasma)) 5 days 83 Anti-Phospholipase A2 Receptor AA2R ((Frozen plasma)) 2 days 83 Anti-SLA (Soluble Liver Antigen) Abs LSA ((Frozen plasma)) 5 days 83 Anti-Staphylolysin Titre ((Frozen plasma)) 2 weeks 83 Anti-Staphylolysin Titre ((Frozen plasma)) 2 weeks 83 Anti-Streptolysin Titre ((Frozen plasma)) 2 weeks 83 Anti-Streptolysin Titre ((Frozen plasma)) 2 weeks 83 Anti-Sulfatide Antibodies 83	Androstenedione	ANDR	B	5 days	57
Angiotensin II  ANG2 (Frozen plasma) 2 weeks 30  Antenatal Profile ANTE (A) (A) 33 (B) (B) (B) 3 days 41, 44  Anti-Actin Antibodies AAA (B) 5 days Anti-Basal Ganglia Antibodies ABGA (B) 3 weeks Anti-CCP Antibodies CCP (B) 2 days Anti-Liver Cytosol Antibodies ALCA (B) 5 days Anti-Houge Inversional Antibodies Anti-MOG [Myelin Oligodendrocyte AMOG (B) 3 weeks Anti-MUSK Antibodies Anti-Musk Antibodies Anti-Nuclear Antibodies (titre & pattern) ANAB (B) 2 days Anti-Phosphatidylserine Antibodies Anti-Phospholipase A2 Receptor AA2R (B) 6 weeks Anti-SLA (Soluble Liver Antigen) Abs Anti-Staphylolysin Titre (SGOT) ASTT (B) 3 days Anti-Streptolysin Titre (ASOT) ASLT (B) 4 days AsA Anti-Sulfatide Antibodies ASA (B) 5 weeks AsA	Angiotensin Converting Enzyme	ACE	B	1 day	30
Antenatal Profile  ANTE	Angiotensin Converting Enzyme – CSF	ACEF	CSF (Frozen)	2 weeks	30
Anti-Actin Antibodies AAA 3 5 5 days 83 Anti-Basal Ganglia Antibodies ABGA 6 3 3 weeks 83 Anti-CCP Antibodies CCP 6 2 days 83 Anti-Liver Cytosol Antibodies ALCA 6 5 days 83 Anti-MOG [Myelin Oligodendrocyte AMOG 6 3 3 weeks 83 Glycoprotein] Antibodies Anti-MUSK Antibodies MUSK 6 2 weeks 83 Anti-Nuclear Antibodies (titre & pattern) ANAB 6 2 days 83 Anti-Phosphatidylserine Antibodies PHTS 6 5 days 83 Anti-Phospholipase A2 Receptor AA2R 6 6 weeks 83 Anti-SLA (Soluble Liver Antigen) Abs LSA 6 5 days 83 Anti-Staphylolysin Titre (SGOT) ASTT 6 3 days 83 Anti-Streptolysin Titre (SGOT) ASTT 6 2 days 83 Anti-Streptolysin Titre/ASOT ASLT 6 5 weeks 83 Anti-Sulfatide Antibodies ASA 6 5 5 weeks 83	Angiotensin II	ANG2	(Frozen plasma)	2 weeks	30
Anti-Basal Ganglia Antibodies  ABGA © 3 weeks  Anti-CCP Antibodies  CCP © 2 days  Anti-Liver Cytosol Antibodies  ALCA © 5 days  Anti-MOG [Myelin Oligodendrocyte AMOG © 3 weeks  B3  Anti-MUSK Antibodies  Anti-MUSK Antibodies  MUSK © 2 weeks  Anti-Nuclear Antibodies (titre & pattern)  ANAB © 2 days  Anti-Phosphatidylserine Antibodies  PHTS © 5 days  Anti-Phospholipase A2 Receptor  AA2R © 6 weeks  Anti-SLA (Soluble Liver Antigen) Abs  LSA © 5 days  Anti-Staphylolysin Titre (SGOT)  ASTT © 3 days  Anti-Streptolysin Titre/ASOT  ASLT © 2 days  AsSA © 5 weeks  Assa	Antenatal Profile	ANTE	<b>A A</b> <sup>33</sup> <b>B B B G</b>	3 days	41, 44
Anti-CCP Antibodies CCP G 2 days 83 Anti-Liver Cytosol Antibodies ALCA G 5 days 83 Anti-MOG [Myelin Oligodendrocyte AMOG G 3 weeks 83 Glycoprotein] Antibodies  Anti-MUSK Antibodies MUSK G 2 weeks 83 Anti-Nuclear Antibodies (titre & pattern) ANAB G 2 days 83 Anti-Phosphatidylserine Antibodies PHTS G 5 days 83 Anti-Phospholipase A2 Receptor AA2R G 6 weeks 83 Anti-SLA (Soluble Liver Antigen) Abs LSA G 5 days 83 Anti-Staphylolysin Titre (SGOT) ASTT G 3 days 83 Anti-Streptolysin Titre/ASOT ASLT G 2 days 83 Anti-Streptolysin Titre/ASOT ASLT G 3 days 83 Anti-Streptolysin Titre/ASOT ASLT G 5 weeks 83 Anti-Sulfatide Antibodies ASA G 5 5 weeks 83	Anti-Actin Antibodies	AAA	B	5 days	83
Anti-Liver Cytosol Antibodies ALCA 3 5 days 83 Anti-MOG [Myelin Oligodendrocyte AMOG 3 3 weeks 83 Glycoprotein] Antibodies  Anti-MUSK Antibodies MUSK 3 2 weeks 83 Anti-Nuclear Antibodies (titre & pattern) ANAB 3 2 days 83 Anti-Phosphatidylserine Antibodies PHTS 5 days 83 Anti-Phospholipase A2 Receptor AA2R 3 6 weeks 83 Anti-SLA (Soluble Liver Antigen) Abs LSA 5 days 83 Anti-SLA (Soluble Liver Antigen) Abs LSA 5 days 83 Anti-Staphylolysin Titre (SGOT) ASTT 3 days 83 Anti-Streptolysin Titre/ASOT ASLT 3 2 days 83 Anti-Streptolysin Titre/ASOT ASLT 5 5 weeks 83	Anti-Basal Ganglia Antibodies	ABGA	B	3 weeks	83
Anti-MOG [Myelin Oligodendrocyte AMOG © 3 weeks 83 Glycoprotein] Antibodies  Anti-MUSK Antibodies MUSK © 2 weeks 83 Anti-Nuclear Antibodies (titre & pattern) ANAB © 2 days 83 Anti-Phosphatidylserine Antibodies PHTS © 5 days 83 Anti-Phospholipase A2 Receptor AA2R © 6 weeks 83 Anti-SLA (Soluble Liver Antigen) Abs LSA © 5 days 83 Anti-Staphylolysin Titre (SGOT) ASTT © 3 days 83 Anti-Streptolysin Titre/ASOT ASLT © 2 days 83 Anti-Streptolysin Titre/ASOT ASLT © 5 weeks 83 Anti-Sulfatide Antibodies ASA © 5 weeks 83	Anti-CCP Antibodies	CCP	В	2 days	83
Glycoprotein] Antibodies  Anti-MUSK Antibodies  MUSK © 2 weeks 83  Anti-Nuclear Antibodies (titre & pattern) ANAB © 2 days 83  Anti-Phosphatidylserine Antibodies  PHTS © 5 days 83  Anti-Phospholipase A2 Receptor AA2R © 6 weeks 83  Anti-SLA (Soluble Liver Antigen) Abs LSA © 5 days 83  Anti-Staphylolysin Titre (SGOT) ASTT © 3 days 83  Anti-Streptolysin Titre/ASOT ASLT © 2 days 83  Anti-Sulfatide Antibodies ASA © 5 weeks 83	Anti-Liver Cytosol Antibodies	ALCA	B	5 days	83
Anti-Nuclear Antibodies (titre & pattern)  ANAB © 2 days 83  Anti-Phosphatidylserine Antibodies PHTS © 5 days 83  Anti-Phospholipase A2 Receptor AA2R © 6 weeks 83  Anti-Sta (Soluble Liver Antigen) Abs LSA © 5 days 83  Anti-Staphylolysin Titre (SGOT) ASTT © 3 days 83  Anti-Streptolysin Titre/ASOT ASLT © 2 days 83  Anti-Sulfatide Antibodies ASA © 5 weeks 83		AMOG	<b>B</b>	3 weeks	83
Anti-Phosphatidylserine AntibodiesPHTSG5 days83Anti-Phospholipase A2 ReceptorAA2RG6 weeks83Anti-SLA (Soluble Liver Antigen) AbsLSAG5 days83Anti-Staphylolysin Titre (SGOT)ASTTG3 days83Anti-Streptolysin Titre/ASOTASLTG2 days83Anti-Sulfatide AntibodiesASAG5 weeks83	Anti-MUSK Antibodies	MUSK	B	2 weeks	83
Anti-Phospholipase A2 Receptor AA2R © 6 weeks 83 Anti-SLA (Soluble Liver Antigen) Abs LSA © 5 days 83 Anti-Staphylolysin Titre (SGOT) ASTT © 3 days 83 Anti-Streptolysin Titre/ASOT ASLT © 2 days 83 Anti-Sulfatide Antibodies ASA © 5 weeks 83	Anti-Nuclear Antibodies (titre & pattern)	ANAB	B	2 days	83
Anti-SLA (Soluble Liver Antigen) AbsLSA35 days83Anti-Staphylolysin Titre (SGOT)ASTT3 days83Anti-Streptolysin Titre/ASOTASLT32 days83Anti-Sulfatide AntibodiesASA35 weeks83	Anti-Phosphatidylserine Antibodies	PHTS	B	5 days	83
Anti-Staphylolysin Titre (SGOT)ASTT3 days83Anti-Streptolysin Titre/ASOTASLT2 days83Anti-Sulfatide AntibodiesASA35 weeks83	Anti-Phospholipase A2 Receptor	AA2R	B	6 weeks	83
Anti-Streptolysin Titre/ASOT ASLT © 2 days 83 Anti-Sulfatide Antibodies ASA © 5 weeks 83	Anti-SLA (Soluble Liver Antigen) Abs	LSA	В	5 days	83
Anti-Sulfatide Antibodies ASA (3) 5 weeks 83	Anti-Staphylolysin Titre (SGOT)	ASTT	В	3 days	83
	Anti-Streptolysin Titre/ASOT	ASLT	В	2 days	83
Anti-Xa Apixaban Monitoring APIX © (Frozen)*18 3 days 42	Anti-Sulfatide Antibodies	ASA	В	5 weeks	83
	Anti-Xa Apixaban Monitoring	APIX	(Frozen)*18	3 days	42
Anti-Xa Edoxaban Monitoring EDOX	Anti-Xa Edoxaban Monitoring	EDOX	(Frozen)*18	3 days	42

TEST	CODE	SAMPLE REQS	TAT	PAGE
Anti-Xa Fondapariux Monitoring	FOND	€ Frozen)*18	3 days	42
Anti-Xa LMWH Monitoring	LMWX	C (Frozen)*18	3 days	42
Anti-Xa Rivaroxaban Monitoring	RIVA	C (Frozen)*18	3 days	42
Antidiuretic Hormone	ADH	(Plasma Frozen) <sup>4</sup>	10 days	57
Antimony (Urine)	ANTI	RU 30	10 days	30
Antimullerian Hormone (AMH) (Self-collect)	AMH	(TDL Tiny) or (TDL Tiny)	1 day	30, 57, 158
Antimullerian Hormone (AMH) (Venous)	AMH	<b>B</b>	1 day	30, 57
Antithrombin III Activity	A111	(Frozen) <sup>4,9,18</sup>	3 days	42
AP50 Alternative Hemolytic Complement	AP50	(Frozen)	2 weeks	30
Apolipoprotein A1 (Self-collect) NEW	AP0A	(TDL Tiny)	3 days	30, 159
Apolipoprotein A1 (Venous)	AP0A	B	3 days	30
Apolipoprotein B (Self-collect) NEW	APOB	(TDL Tiny)	3 days	30, 159
Apolipoprotein B (Venous)	AP0B	B	3 days	30
Apolipoprotein C	AP0C	B	3 months	30
Apolipoprotein E (12 hours fasting)	AP0E	(fasting)	5 days	30
Apple Components	ZZ36	В	2 days	142
APTT/KCCT	KCCT	<b>C</b> 18	1 day	41
Aquaporin 4 Antibodies (Neuromyelitis Optica)	AQUA	B	2 weeks	83
Arbovirus Antibodies/Abs	ARB0	B 9,14	3 weeks	102
Array-CGH (Comparative Genomic Hybridisation) SNP array	CGH	CVS / AF / (A) (1) 9	10 days	117
Arsenic (Blood)	ARS	A or (1)	5 days	30, 165
Arsenic (Urine)	ARSE	RU 30	5 days	30, 165
Arylsulphatase A	ARYL	<b>1</b> 5,6	8 weeks	30
Ascariasis Serology	ASC	B	5 days	83
Ashkenazi Jewish Carrier Screen	GENE	<b>A</b> 9	4 weeks	117
Aspartate Transaminase (AST) (SGOT)	AST	B	1 day	30
Aspergillus Components	ZZ2	B	2 days	142
Aspergillus Precipitins	ASPP	B	5 days	83
Ataxia NGS Panel	GENE	<b>AA</b> 9	6 weeks	117
Atopic Dermatitis/Eczema Profile (14 allergens)	ALEC	88	2 days	141, 146
Atypical Antibody Screen (handwritten tube label)	AASC	A 22,33	2 days	41
Atypical Pneumonia Screen	APS	B	3 days	102, 104
Autoantibody Profile I	AUT0	B	2 days	83, 89
Autoantibody Profile II	END0	B	3 days	83, 89
Avian Precipitins (11 Species)	AVIA	B	5 days	83
Babesia PCR	PCRB	A	7 days	83
Bancroftia/Oncerciasis/Filarial Antibodies	TFIF	B 14	2 weeks	92
Beckwith-Wiedemann Syndrome – methylation studies on 11p15 imprinting domains KvDMR + H19	GENE	A 9,11	6 weeks	118

TEST	CODE	SAMPLE REQS	TAT	PAGE
Bence-Jones Protein	RBJP	RU or CU	5 days	30
Benzene	BENZ	<b>J</b> 1,6	3 days	166
Beta 2 Glycoprotein 1 Abs	B2GP	В	2 days	83
Beta 2 Microglobulin (Serum)	B2MG	В	2 days	30, 166
Beta 2 Microglobulin (Urine)	UB2M	RU	3 days	30, 166
Beta Carotene	CAR0	В	5 days	151
Beta D Glucan	XBDG	B	3 days	47
Beta Thalassaemia – beta-globin gene	GENE	<b>A</b> 9	4 weeks	118
sequencing + deletions/duplications				
Beta-Glucuronidase (Sly Disease)	BGLU	<b>(1)</b> (1) 9,4	8 weeks	30
Bicarbonate	HC03	В	1 day	30
Bile Acids – Serum	BILE	В	1 day	30
Bilharzia (Schistosome) Antibody Screen	BILH	B 14	10 days	92
Bilharzia (Urine)	USCH	Mid-morning terminal urine following exercise <sup>14</sup>	1-2 days	92
Bilirubin (Direct)	DBIL	В	1 day	30
Bilirubin (Indirect)	IBIL	В	1 day	30
Bilirubin (Total)	BILI	В	1 day	30
Biotin	BIOS	<b>B</b> 7	5 days	151
Biotinidase	BIOT	(Frozen plasma) <sup>4</sup>	3 weeks	30
Birch Components	ZZ3	B	2 days	142
Bismuth	BISM	В	5 days	30
BK Polyoma Virus by PCR	BKPV	A/RU	5 days	102
Blood Culture#	BCUL	2 x <b>BC</b> <sup>4</sup>	6 days +	47
Blood Film Examination	FILM	A	1 day	41
Blood Group †	AB0	A 22,33	2 days	41
BNP (NT-pro BNP)	BNP	B	1 day	30, 57
Bone Alkaline Phosphatase	BALP	(Frozen)	2 weeks	30
Bone Marrow (Aspirate)	BMAS	<b>J</b> <sup>1</sup>	14 days	44
Bone Marrow (Trephine Biopsy)	BMI	<b>J</b> <sup>1</sup>	3 days	44
Bone Screen	BONE	B CU	1 day	30, 38
Bone Screen (Bloods only)	BON2	В	1 day	30, 38
Borrelia Antibodies (Lyme Disease) IgG, IgM	BORR	B 9,14	2 days	83, 92
Borrelia Antibodies (Lyme Disease) IgM	BORM	В	2 days	83, 92
Borrelia Confirmation (Immunoblot)	BORC	B 9,14	10 days	83, 92
Brazil Components	ZZ4	В	2 days	142
Breast Cancer NGS Panel	GENE	<b>A A</b> 9,11	4 weeks	106
Bromide	BROM	В	3 days	166
Brucella Serology	BRUC	<b>B</b> 9	2-3 weeks	83
BUN (Blood Urea Nitrogen) (Calculated)	BUN	В	1 day	30
C Peptide	CPEP	В	3 days	57
C Reactive Protein (Self-collect)	CRP	(TDL Tiny)	1 day	31, 159
C Reactive Protein (Venous)	CRP	В	1 day	31

TEST	CODE	SAMPLE REQS	TAT	PAGE
C Reactive Protein (High	HCRP	(TDL Tiny)	1 day	31, 159
Sensitivity) (Self-collect)				
C Reactive Protein (High Sensitivity) (Venous)	HCRP	В	1 day	31
C1 Esterase Inhibitor	C1EI	В	5 days	83
C1 Esterase: Function & Total	FC1E	(Plasma Frozen) <sup>4,18</sup>	10 days	31
C1q Binding Immune Complex	IMCP	В	5 days	31
C3 Complement	C3	В	1 day	83
C3/C4 Complement	COMP	В	1 day	83
C4 Complement	C4	<b>B</b>	1 day	83
CA 15-3	C153	B	1 day	106
CA 19-9	C199	<b>B</b>	1 day	106
CA 50	CA50	<b>B</b>	5 days	106
CA 72-4	C724	<b>B</b>	5 days	106
CA 125 (Self-collect)	C125	(TDL Tiny)	1 day	106, 159
CA 125 (Venous)	C125	<b>B</b>	1 day	106
CADASIL - NOTCH3 gene sequencing	GENE	<b>A</b> 9	6 weeks	118
Cadmium (Blood)	CADM	A or 🕕	5 days	31, 165
Cadmium (Urine)	URCD	RU 30	5 days	31, 165
Caeruloplasmin	CERU	B	1 day	31, 152
Calcitonin	CAT0	(Frozen) <sup>4</sup>	1 day	57
Calcium (24 hour Urine)	UCA	PU or acid urine	1 day	31
Calcium (Venous)	CA	B	1 day	31
Calcium + Vitamin D (Venous)	CALD	В	1 day	31
Calcium/Creatinine Ratio	CACR	CU B	1 day	31
Calprotectin	CALP	QFIT sample collection device	5 days	31
Calprotectin (Self-collect)	CALP	QFIT sample collection device	5 days	31, 159
Calprotectin/QFIT Profile (Combined) (QFIT)	QCAL	QFIT	5 days	31, 39
Calprotectin/QFIT Profile (Combined) (Self-collect)	QCAL	<b>QFIT</b> sample collection device	5 days	31, 159
Campylobacter Jejuni Antibodies	CJAB	В	5 days	83
Cancer, Comprehensive NGS Panel	GENE	<b>A A</b> 9,11	5 weeks	118
Candida (Culture for ID + Sensitivities)	CAND	STM/CS	2-4 days	47
Candida (Culture for ID Only)	CANC	STM/CS	2-4 days	47
Candida Antibodies	CANA	B	5 days	83
Candida auris Screen	CANS	STM/CS	2-4 days	47
Cannabinoids (Urine) Screen	CANN	RU	1 day	163
Carbamazepine (Tegretol)	CARB	B	1 day	138
Carbapenemase producing organism screen	MDR	STM (rectal)	4-5 days ‡	47
Carbohydrate Deficient Glycoprotein	CDG	B (rectar)	2 weeks	31
	CDT			
Carbohydrate Deficient Transferrin (CDT) (Self-collect)	CDI	(TDL Tiny)	3 days	31, 159
Carbohydrate Deficient Transferrin (CDT) (Venous)	CDT	В	3 days	31

TEST	CODE	SAMPLE REQS	TAT	PAGE
Carboxyhaemoglobin	СВНВ	<b>(A</b>	1 week	41
Carcino Embryonic Antigen	CEA	B	1 day	106
Cardiolipin Antibodies (IgG+IgM)	ACAB	B	2 days	83
Cardiomyopathy, Dilated NGS Panel	GENE	<b>AA</b> <sup>9</sup>	6 weeks	118
Cardiomyopathy, Hypertrophic NGS Panel	GENE	<b>AA</b> <sup>9</sup>	6 weeks	118
Cardiovascular Risk Profile 1	PP10	BB	3 days	31, 38
Cardiovascular Risk Profile 2	PP11	<b>BBB ©</b> 34	3 days	31, 39
Cardiovascular, Comprehensive NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	118
Carotenes	CAR0	B	5 days	151
Carrier Screen (Ashkenazi Jewish)	GENE	<b>A</b> 9	4 weeks	130, 133
Carrier Screen (Ashkenazi Jewish) – Partnered Report	GENE	<b>A</b> 9	4 weeks	133
Carrier Screen (Pan-Ethnic)	GENE	<b>A</b> 9	4 weeks	130, 133
Carrier Screen (Pan-Ethnic) – Partnered Report	GENE	<b>A</b> 9	4 weeks	133
Cashew Components	ZZ35	B	2 days	142
Cat Components	ZZ5	B	2 days	142
Cat Scratch Fever (Bartonella IgG)	CAT	B	5 days	102
Catecholamines (Plasma)	CATE	(Plasma Frozen) <sup>4</sup>	5 days	57
Catecholamines (Urine)	UCAT	PU (collect on acid) <sup>1</sup>	5 days	57
CCP Antibodies (RF)	CCP	B	2 days	84
CD3/CD4/CD8	LYSS	A	1 day	102
CD16	CD16	<b>A</b> 4	1 day	44
CD19 B Cells	CD19	<b>A</b> 4	1 day	44
CD20	CD20	<b>A</b> 10	2 days	44
CD25	CD25	<b>A</b> 10	2 days	44
CD56	CD56	<b>A</b> 4	1 day	44
CD57	CD57	A	1 day	44
Celery Components	ZZ6	<b>B</b>	2 days	142
Cervical Cytology	PAPT	TPV	6 days (combined report)	172
Cervical Cytology + HP20	PAPT + HP20	TPV	6 days (combined report)	172
Cervical Cytology + HPVH	PAPT + HPVH	TPV	6 days (combined report)	172
Cervical Cytology + HPVT	PAPT + HPVT	TPV	6 days (combined report)	172
CH50 (Classical pathway)	CH50	(Frozen) <sup>4</sup>	4 days	84
Chagas Disease Serology	CHGA	B 9,14	10 days	84
(S.American Trypanosomiasis) T. Cruzi			· 	
Charcot-Marie-Tooth Syndrome NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	119
Chest Pain Profile	CPP	<b>B</b>	STAT	31, 39
Chikungunya Virus Abs	CHIK	B 9,14	10 days	102
Chlamydia – PCR swab	SPCR	PCR	2 days	72

TEST	CODE	SAMPLE REQS	TAT	PAGE
Chlamydia – Thin Prep	TPCR	TPV	2 days	72, 170
Chlamydia – Urine	CPCR	FCRU	2 days	72
Chlamydia Species Specific (MIF) Ab Screen	CHAB	B	3 days	84, 89
Chlamydia/Gonorrhoea – PCR Swab	SCG	PCR	2 days	72
Chlamydia/Gonorrhoea – Rectal (PCR)	RSCG	PCR	2 days	72
Chlamydia/Gonorrhoea – Rectal (Self-collect)	RSCG	Aptima multisite swab	2 days	72, 159
Chlamydia/Gonorrhoea – Thin Prep	TCG	TPV	2 days	72, 170
Chlamydia/Gonorrhoea – Throat (PCR)	TSCG	PCR	2 days	72
Chlamydia/Gonorrhoea – Throat (Self-collect)	TSCG	Aptima multisite swab	2 days	72, 159
Chlamydia/Gonorrhoea – Urine (FCRU)	CCG	FCRU	2 days	72
Chlamydia/Gonorrhoea – Urine (Self-collect)	CCG	Aptima urine	2 days	72, 159
Chlamydia/Gonorrhoea – Vaginal (Self-collect)	SCG	Aptima multisite swab	2 days	72, 159
Chlamydia/Gonorrhoea/	SCGT	PCR	2 days	72
Trichomonas – PCR Swab				
Chlamydia/Gonorrhoea/Trichomonas – Thin Prep	TCGT	TPV	2 days	72, 170
Chlamydia/Gonorrhoea/Trichomonas – Urine	CCGT	FCRU	2 days	72
Chloride	CL	В	1 day	31
Cholesterol	CH0	В	1 day	31
Cholesterol (Familial Hypercholesterolaemia)	GENE	<b>A A</b> <sup>9</sup>	7 weeks	31
Cholinesterase (Serum/Pseudo)	CHPS	В	1 day	31, 166
Chromium (Blood)	CHR0	🛕 / Trace metal / 🕕	5 days	31, 165
Chromium (Urine)	URCR	RU 30	4 weeks	31, 165
Chromogranin A	CGA	B	1 week	32
Chromogranin A & B	MTAB	(Frozen plasma)	3 weeks	32
Chromosome Analysis (Blood)	KARY	<b>(1)</b> 9	2-3 weeks	119
Chronic Fatigue Syndrome Profile	VIP1	A + B 10	5 days	84, 89
Citrate (Blood)	CITR	B	5 days	32
Citrate (Urine)	UCIT	CU (Frozen)	5 days	32
CK (MB Fraction)	CKMB	B	1 day	32
CK Isoenzymes	CKIE	B	5 days	32
Clobazam	CLOB	A	5 days	138
Clomipramine (Anafranil)	CHLO	A	7 days	138
Clonazepam	CLON	A	7 days	138
Clostridium Difficile Toxin by PCR	CLOS	RF*	2 days	47
CMV IgM Antibodies	CMVM	(Plasma) or (B) (Serum)	1 day	102
Coagulation Profile 1	CLPF	<b>C</b> 18	1 day	41, 44
Coagulation Profile 2	CLOT	<b>△ ○</b> 18	1 day	41, 44
Cobalt (Blood)	COB	A	5 days	32
Cobalt (Urine)	COBA	RU 30	5 days	32, 165
Cocaine (Urine) Screen	UCOC	RU	1 day	163
Coeliac Disease – HLA DQ2/DQ8 Genotype	Q2Q8	<b>A</b> 9	10 days	84, 119
Coeliac/Gluten Genetic Profile 2	GSA2	<b>AB</b>	10 days	84, 89

TEST	CODE	SAMPLE REQS	TAT	PAGE
Coeliac/Gluten Sensitivity Profile	GSA	В	2 days	84, 89
Coenzyme Q10	CQ10	B	2 weeks	32
Cold Agglutinin	CAGG	J <sup>1</sup>	5 days	32
Colloid Antigen-2 Antibodies	CA2A	B	2 weeks	84
Complement C1q	C1Q	B	5 days	32
Complement C2	C2	B	10 days	32
Complement C3	C3	B	1 day	32
Complement C4	C4	B	1 day	32
Complement C5	C5A	B	2 weeks	32
Complement C6	C6	(Frozen)*	5 weeks	32
Complement C7	C7	B (Frozen)*	5 weeks	32
Complement C8	C8	(Frozen)*	5 weeks	32
Complement C9	C9	B (Frozen)*	5 weeks	32
Complement Factor H	FACH	B	3 weeks	32
Complex PSA (Prostate Specific Ag)	CPSA	B	3 days	106
Congenital Adrenal Hyperplasia NGS Panel	GENE	<b>A</b> 9	6 weeks	120
Congenital Myopathy NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	120
Connective Tissue Disorders NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	120
Coombs (Direct Antiglobulin Test)	COOM	A	2 days	43
Copper (Serum)	COPP	B or (§	5 days	32, 152,
				165
Copper (Urine)	URCU	CU	5 days	32, 165
Cornelia de Lange Syndrome NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	120
Cortisol (Self-collect)	CORT	(TDL Tiny)	1 day	57, 159
Cortisol (Urine)	UCOR	CU	5 days	57
Cortisol (Venous)	CORT	В	1 day	57
Cortisol Binding Globulin	CBG	(Frozen)	1 month	32
Cotinine (Serum)	COT	В	4 days	84
Cotinine (Urine)	COTT	RU	2 days	32
COVID-19 (SARS-CoV-2) (PCR)	NCOV	PCR Swab (nasal/pharyngeal)	1 day	102
COVID-19 (SARS-CoV-2) RNA	NCOV	Aptima multisite swab	1 day	102, 159
by PCR (Self-collect)		of nose/throat		
COVID-19 (SPIKE) Antibodies (Self-collect)	SCOV	(TDL Tiny)	1 day	84, 159
COVID-19 (SPIKE) Antibodies (Venous)	SCOV	SST/Serum (3) (Venous)	1 day	84
Cow's Milk Components	ZZ7	<u>B</u>	2 days	142
Craniosynostosis NGS Panel	GENE	AA <sup>9</sup>	6 weeks	120
Creatine Kinase (CK, CPK)	CKNA	(TDL T: )	1 day	32
Creatinine (including eGFR) (Self-collect)	CREA	(TDL Tiny)	1 day	32, 159
Creatinine (including eGFR) (Venous)	CREA	B	1 day	32
Creatinine (Urine)	UCR	CU	1 day	32
Creatinine Clearance	CRCL	B CU	1 day	32
Crosslaps (Serum DPD)	SDPD	(Freeze within 24 hours)	4 days	32

TEST	CODE	SAMPLE REQS	TAT	PAGE
Cryoglobulins	CRY0	<b>J</b> 6	10 days	32
Cryptococcal Antigen	CRYC	Serum or CSF	1 day	47
Cryptosporidium	0CP	RF	2 days	47
Cryptosporidium Detection by PCR	CRPA	RF	2 days	92
CSF for Microscopy and Culture	CSF	1.5ml CSF	1-3 days	47
CSF Screen by PCR	VPCR	CSF	2 days	102, 104
CT/GC/Trichomonas/Mgen – PCR Swab	SGTM	PCR Swab	2 days	72, 77
CT/GC/Trichomonas/Mgen – Thin Prep	TGTM	TPV	2 days	72, 77
CT/GC/Trichomonas/Mgen – Urine	CGTM	FCRU	2 days	72, 77
Cyclosporin	CYCL	A	1 day	32
Cyfra 21-1	CY21	B	4 days	106
Cystatin C	CYCC	<b>B</b>	5 days	33
Cystine – Quantitative (Beta-CTX)	QCYS	PU	5 days	33
Cytochrome P450 2C19	2C19	<b>A</b> 9	10 days	120
Cytomegalovirus (CMV-DNA) Amnio	CMVD	AF	5 days	102
Cytomegalovirus (IgG/IgM) Antibodies	CMV	B	1 day	102
Cytomegalovirus (PCR) Semen	SCVM	Semen	7 days	102
Cytomegalovirus (PCR) Urine	CMVU	RU	5 days	102
Cytomegalovirus Avidity	CMAV	B	10 days	102
Cytomegalovirus DNA (PCR)	CMVP	A	5 days	102
Cytomegalovirus Resistance	CMVR	(6mls)	21 days	102
D-Dimers (Fibrinogen Degradation Products)	DDIT	<b>C</b> 4	1 day	41
Decidualization Score (DS)	DSRF	J (Contact Referrals)	2-3 weeks	61
Dengue Fever PCR	DPCR	(A) or (B) 9,14	2 weeks	102
Dengue Virus Serology	DENG	B 9,14	5 days	92
Deoxypyridinoline (DPD) – Serum	SDPD	(Freeze within 24 hours)	4 days	33
Deoxypyridinoline (DPD) – Urine	DPD	EMU	4 days	33
DHEA	DHEX	<b>B</b>	7-10 days	57
DHEA – Urine (Dehydroepiandrosterone)	UDHE	CU	3 weeks	57
DHEA Sulphate (Self-collect)	DHEA	(TDL Tiny) or (TDL Tiny)	1 day	57, 159
DHEA Sulphate (Venous)	DHEA	B	1 day	57
Diabetic Profile 1	DIAB	AG	1 day	33, 39
Diabetic Profile 2	DIA2	<b>♠ G</b> RU	2 days	33, 39
Diamine Oxidase Activity	DIAM	B	2 weeks	33
Diazepam (Valium)	DIAZ	A	7 days	138
Digoxin	DIGO	В	1 day	138
Dihydrotestosterone	DHT	88	7 days	57
Dilated Cardiomyopathy NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	120
Diphtheria Antibodies	DIPH	В	5 days	84
DL1-DL12 Screening Profiles				26-27
DL12 7 STI Profile by PCR	DL12	Aptima urine or multisite swab	2 days	159
(7 PCR tests from 1 Sample) (Self-collect)				
DNA (Double Stranded) Antibodies IgG	DNAA	B	2 days	84

TEST	CODE	SAMPLE REQS	TAT	PAGE
DNA (Single Stranded) Antibodies	DNAS	В	5 days	84
Dog Components	ZZ8	B	2 days	142
Down Syndrome Risk Bloods only (Risk to be calculated by clinician)	HCGF/ PAPA	<b>B</b>	1 day	57
Down Syndrome Risk Profile (2nd trimester) Quad	DRP	B DRP form <sup>7,8</sup>	5 days	57
Down Syndrome Risk Profile with risk calculation first trimester	DRP	B DRP form + image of scan <sup>7,8</sup>	5 days	58
Doxepin Level (Sinequan)	DOXE	(A) or (f)	10 days	166
Drugs of Abuse from Blood without Chain of Custody	DOAP	В	5 days	163
Drugs of Abuse Profile – Random Urine Sample/No Chain of Custody	DOA	RU	2 days (5 days with LC- MS/MS confirmation)	163-164
Drugs of Abuse Profile – Random Urine Sample/No Chain of Custody Plus Alcohol	DOA3	RU	2 days (5 days with LC- MS/MS confirmation)	163-164
Drugs of Abuse Profile – With Chain of Custody*	DOAL	RU/CoC Collection Containers 1,2	2 days (5 days with LC- MS/MS confirmation)	163-164
Drugs of Abuse Profile – Without Chain of Custody	DOAN	RU <sup>2</sup>	2 days (5 days with LC- MS/MS confirmation)	163-164
DVT/Pre-travel Screen	DVT1	<b>A B</b> <sup>9</sup>	5 days	41, 45, 92-93, 121, 130
Echinococcus (Hydatid) Antibodies	EFAT	B 9,14	5 days	84, 92
Egg Components	ZZ9	В	2 days	142
Ehlers-Danios Syndrome NGS Panel	GENE	<b>A</b> 9,11	6 weeks	121
Ehrlichiosis Antibodies	EHRL	B 9,14	10 days	84
Elastase (RF)	ELAS	RF	5 days	33
Elastase (Self-collect)	ELAS	Stool/faecal container	5 days	33, 159
Electrolytes	ELEC	B	1 day	33
Electrolytes (Urine)	UELE	CU	1 day	33
ELF/Enhanced Liver Fibrosis	ELF	В	5 days	33
<b>Endometrial Cancer NGS Panel</b>	GENE	<b>A A</b> 9,11	4 weeks	121
Enteric Organism Rapid Detection (RF)	EORD	RF	2 days	92-93
Enteric Organism Rapid Detection (Self-collect)	EORD	Stool/faecal container	2 days	92, 159
Eosin-5 Maleimide Dye binding test for Hereditary spherocytosis (EMA)*	EMA	A	2 days	43
Eosinophil Cationic Protein	ECP	B	7 days	33
Epanutin (Phenytoin)	PHEN	B	1 day	138
Epidermolysis Bullosa NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	121
Epilepsy, Adolescent/Adult Onset Panel	GENE	A	6 weeks	121
Epilepsy, Comprehensive NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	121
Epstein-Barr Virus Antibodies IgG/IgM	EBVA	B	2 days	102
Epstein-Barr Virus PCR	EBVQ	<b>A</b>	5 days	102
Erectile Dysfunction Profile	IMP0	<b>ABBG</b>	3 days	58, 62

TEST	CODE	SAMPLE REQS	TAT	PAGE
Erythropoietin	ERY	B	4 days	33, 138
ESR	ESR	<b>A</b>	1 day	41
Essential Fatty Acid Profile (Red Cell)	EFAR	<b>A</b> 4	10 days	152
Ethosuximide	ETH0	<b>A</b>	7 days	138
Extractable Nuclear Antibodies (nRNP,	ENA	<b>B</b>	2 days	84
Sm, Ro, La, Jo1, Sci70) CENP-B				
Fabry Disease, X-linked – GLA gene sequencing	GENE	<b>A</b> <sup>9</sup>	4 weeks	121
Facioscapulohumeral Muscular Dystropy (FSHD) – D4Z4 repeat deletion	GENE	<b>A A</b> <sup>9</sup>	9 weeks	121
Factor II Assay	FAC2	(Frozen) <sup>9,18</sup>	5 days	42
Factor V Assay	FAC5	(Frozen) <sup>9,18</sup>	5 days	42
Factor VII Assay	FAC7	(Frozen) <sup>9,18</sup>	5 days	42
Factor VIII Assay	FAC8	(Frozen) <sup>9,18</sup>	5 days	42
Factor VIII Inhibiting Antibody	F8IA	<b>0 0</b> 18	2 weeks	42
Factor IX Assay	F1X	(Frozen) <sup>9,18</sup>	5 days	42
Factor IX Inhibiting Antibody	F9IA	<b>C C</b> 18	2 weeks	42
Factor X Assay	FX	C (Frozen) <sup>9,18</sup>	5 days	42
Factor XI Assay	FX1	C (Frozen) <sup>9,18</sup>	5 days	42
Factor XII Assay	FX11	C (Frozen) <sup>9,18</sup>	5 days	42
Factor XIII Assay	FA13	C (Frozen) <sup>9,18</sup>	5 days	42
Faecal Fat (1 day collection)	TFFA	LF <sup>6</sup>	5 days	33
Faecal Fat (3 day)	FFAT	LF <sup>6</sup>	5 days	33
Faecal Lactoferrin	FLAC	RF	5 days	33
Faecal Sugar Chromatography	FCR0	RF (Frozen)	3 weeks	33
Familial Hypercholesterolaemia NGS panel	GENE	<b>AA</b> 9	6 weeks	121
Familial Hypocalciuric	GENE	<b>A A</b> <sup>9</sup>	6-7 weeks	121
Hypercalcaemia (FHH) Panel				
Familial Mediterranean fever	GENE	<b>A</b> 9	5 weeks	121
MEFV gene sequencing				
Farmers Lung Precipitins	FARM	8	5 days	84
Fasciola Hepatica Antibodies (Liver Fluke)	FASC	8	2 weeks	84
Fasting Insulin Resistance Index (FIRI)	FIRI	86	1 day	58
Female Hormone Profile (LH, FSH, PROL, TOES) (Self-collect)	TFIP	(TDL Tiny) and (3) (TDL Tiny)	1 day	58, 159
Female Hormone Profile (Venous)	FIP	B	1 day	58, 62
Ferritin (Self-collect)	FERR	(TDL Tiny)	1 day	33, 159
Ferritin (Venous)	FERR	B	1 day	33
Fibrinogen	FIB	C 4,18	1 day	41
Fibrotest (Liver Fibrosis)	FIBT	B	2 weeks	33
Filaria (Lymphatic and Non-	FIFA	B 9,14	10 days	92
Lymphatic) Antibodies		•		3=
First Trimester Maternal Screen (PAPP-A/Free Beta-hCG)	FTMS	В	1 day	58, 63

TEST	CODE	SAMPLE REQS	TAT	PAGE
Fish Components	ZZ10	B	2 days	142
FK506 (Tacrolimus/Prograf)	FK5	<b>A</b> 4	1-2 days	138
Flecainide (Tambocor)	FLEC	A	5 days	138
Fluid Culture	FLUD	SC	2-7 days	47
Fluid Cytology	CATF	Fluid <sup>4</sup>	3 days	174
Fluoride (Urine)	UFL	RU	5 days	33
Fluoxetine (Prozac)	PROZ	<b>A</b> 4	5 days	138
Folate (Red Cell)	RBCF	A	2 days	33, 152
Folate (Serum)	F0LA	B	1 day	33
Free Fatty Acids	FFA	(Frozen) <sup>1</sup>	10 days	33
Free T3 (Self-collect)	FT3	(TDL Tiny)	1 day	58, 159
Free T3 (Venous)	FT3	В	1 day	58
Free T4 (Self-collect)	FT4	(TDL Tiny)	1 day	58, 159
Free T4 (Venous)	FT4	В	1 day	58
Friedreich Ataxia – frataxin gene repeat analysis	GENE	<b>A</b> 9	5 weeks	121
Fructosamine	FRUC	В	1 day	33
FSH (Self-collect)	FSH	(TDL Tiny) or (TDL Tiny)	1 day	58, 159
FSH (Venous)	FSH	В	1 day	58
Full Blood Count	FBC	A	1 day	41
Full Blood Count* (Haemoglobin,	TFBC	(TDL Tiny)	1 day	41, 159
White Cell Count, Red Cell Count, Platelets)				
(Self-collect) NEW				
Fungal investigations (non-	FUN	All specimens other than	3-21 days	47
superficial extended culture)	DEDM	Skin, Hair and Nails	0.7.1	47
Fungal investigations (superficial/	DERM	Skin, Hair, Nails	3-7 days	47
dermatophyte PCR test) FXIII A Subunit	F13S	(Fragan) 918	14 dovo	42
G6PD	G6PD	(Frozen) <sup>9,18</sup>	14 days 4 days	43
Gabapentin	GABA	B 4	5 days	138
Galactomanan (Aspergillus Antigen)	SGAL	<b>B</b>	2 weeks	47
Galactose-1-Phosphate Uridyltransferase	GAL1	<b>(1)</b> 5,6	2 weeks	33
Galactosidase – Alpha*	GALA		6 weeks	33
Gall Stone Analysis	RSTA	STONE	10 days	33
Gamma GT	GGT	B	1 day	33
Ganglionic Acetylcholine Receptor Antibodies	GACA	<u> </u>	9 weeks	84
Ganglioside GM1, GD1B, GQ1B Abs	GANG	<u> </u>	5 days	84
Gardnerella vaginalis (Thin Prep)	GVPC	TPV	2 days	170
Gardnerella vaginalis by PCR	GVPC	FCRU / PCR / TPV	2 days	72
Gastric Parietal Autoantibodies	GASP	B	2 days	84
Gastrin	GAST	(Plasma frozen)	5 days	33
Gastrointestinal Pathogen Profile	EORD	Stool/faecal container	2 days	92, 159
by PCR (Self-collect)	LUID	סנטטו/ ומסטמו טטוונמוווסו	L udyo	32, 139
Gaucher Disease full gene sequencing	GENE	<b>A</b> 9	6 weeks	121
aaaa 2.00000 iun gono ooquononig	JEITE	•	J	121

	CODE	SAMPLE REQS	TAT	PAGE
Genetic Reproductive Profile (Male)	GRP	<b>A B</b> <sup>9</sup>	10-15 days	121
Genetics: TDL Genetics				109-136
Gentamicin Assay	GENT	<b>B</b> 4	1 day	137
Gilbert Syndrome –	GENE	<b>A</b> 9	2-3 weeks	122
common UGT1A1 repeat variation				
Gliadin Antibodies (IgG) (deamidated) (Self-collect)	AGAB	(TDL Tiny)	2 days	84, 160
Gliadin Antibodies (IgG) (deamidated) (Venous)	AGAB	<b>B</b>	2 days	84
Globulin (Calculated)	GLOB	<b>B</b>	1 day	33
Glomerular Basement Membrane Abs	AGBM	<b>B</b>	2 days	84
Glucagon	GLUG	(Plasma)	10 days	34
Glucose	RBG	G	1 day	34
Glucose Challenge Test/Mini-GTT	RBGM	<b>G</b>	1 day	137
Glucose Tolerance Test (Extended Plus)	GTTX	7 x 📵 , 7 x <b>RU</b>	1 day	137
Glucose Tolerance Test (Extended)	GTTE	5 x 📵 , 5 x RU	1 day	137
Glucose Tolerance Test (Short)	GTTS	2 x 🕒 , 2 x RU	1 day	137
Glucose Tolerance Test/OGTT	GTT	3 x 🕒 , 3 x RU	1 day	137
Glucose Tolerance with Growth Hormone	GTT +	3 x 📵 35,	1 day	137
	GHDF	3 x 🕒 , 3 x <b>RU</b>		
Glucose Tolerance with Insulin	GTTI	3 x 🔒 , 3 x 😉 , 3 x RU	1 day	137
Glucose-6-Phosphate Dehydrogenase (G6PD) Deficiency – full G6PD gene sequencing	GENE	<b>A</b> 9	3-4 weeks	122
Glutamic Acid Decarboxylase Antibodies (GAD 65)	GAD	3	5 days	84
Gluten Sensitivity Evaluation	GSA	B	2 days	84
Gluten Sensitivity Profile	GLUT	ABB	10 days	84, 89,
•			•	141, 146
Gluten/Coeliac Genetic Profile 2	GSA2	AB	10 days	84
Glycan Determinants	ZZ27	В	2 days	142
Glycogen storage disease type 2 (Pompe) variant analysis	POMP	0	4 weeks	122
Gonorrhoea – PCR swab	SGON	PCR	2 days	72
Gonorrhoea – Thin Prep	TGON	TPV	2 days	72, 170
Gonorrhoea – Urine	CGON	FCRU	2 days	72
Gonorrhoea Culture – Cervix	GONC	CS <sup>‡‡‡</sup>	3-5 days	47, 72
Gonorrhoea Culture – Other site	GONO	CS	3-5 days	47, 72
Gonorrhoea Culture – Rectal	GONR	CS	3-5 days	47, 72
Gonorrhoea Culture – Throat	GONT	CS	3-5 days	47, 72
Gonorrhoea Culture – Urethral	GONU	CS	3-5 days	47, 72
Granulocyte Immunology	GRIM	(or 2 x 6ml) (B	2 weeks	84
<del>-</del>	GBSX	Blue gel Amies swab x2	3-5 days	47, 160
Group B Strep - Vaginal and Rectal (Self-collect)	UDOA	Dido goi Aillios sivas Az	o o aayo	71, 100

TEST	CODE	SAMPLE REQS	TAT	PAGE
Growth Hormone (Fasting)	GH	B 7,35	1 day	58
Gut Hormone Profile	GUTP	A (Frozen within	3 weeks	58
H. pylori Antibodies (IgG)	НВРА	15 minutes) <sup>41</sup>	2 days	85
H. pylori Antigen – Stool (RF)	HBAG	RF	3 days	48
H. pylori Antigen – Stool (Self-collect)	HBAG	Stool/faecal container	3 days	48, 160
H. pylori Culture	HPCU	J	1 month	48
Haematology Profile	PP3	A	1 day	41, 45
Haemochromatosis	HMD	<b>A</b> 9	3 days	34, 122
- HFE common variants C282Y + H63D			,	
Haemoglobin	НВ	A	1 day	41
Haemoglobin Electrophoresis	HBEL	A	4 days	43
Haemophilia A (Factor VIII deficiency) – CVS	8CVS	CVS 40	3 days	122
Haemophilia B (Factor IX deficiency) – CVS	9CVS	CVS 40	3 days	122
Haemophilus B Influenzae Antibodies	HINF	В	5 days	85
Haemophilus ducreyi by PCR	DUCR	PCR	7 days	72
Haemosiderin (Urine)	HSID	EMU	2 weeks	34
Hantavirus Serology	HANV	B 9	10 days	102
Haptoglobin	HAPT	В	5 days	34
Hazelnut Components	ZZ11	В	2 days	142
HbA1c (Self-collect)	GHB	(TDL Tiny)	1 day	34, 160
HbA1c (Venous)	GHB	A	1 day	34
HCG (Oncology)	HCGQ	В	1 day	106
HCG (Quantitative) (Self-collect)	QHCG	(TDL Tiny)	1 day	58, 160
HCG (Quantitative) (Venous)	QHCG	В	1 day	58
HDL Cholesterol	HDL	В	1 day	34
HE4 + ROMA	HE4	В	1 day	106, 108
(Earlier Detection of Ovarian Tumour)				
Hearing Loss NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	122
Hepatitis (Acute) Screen	AHSC	В	1 day	95, 104
Hepatitis A (IgM)	HAVM	В	1 day	95
Hepatitis A Immunity (IgG/IgM)	HAIM	В	1 day	95
Hepatitis A Profile	HEPA	В	1 day	72, 95
Hepatitis A RNA by PCR	HAVR	A or B	3 weeks	95
Hepatitis A, B & C Profile	ABC	BB	1 day	95, 104
Hepatitis B 'e' Antigen and Antibody	HEPE	В	1 day	95
Hepatitis B (PCR) Genotype	BGEN	A or B	7 days	95
Hepatitis B Core Antibody – IgM	HBCM	В	1 day	95
Hepatitis B Core Antibody – Total	HBC	В	1 day	95
Hepatitis B DNA (Viral load)	DNAB	(A) or (B)	5 days	95
Hepatitis B Immunity (IgG) (Self-collect)	THBI	(TDL Tiny)	1 day	95, 160
Hepatitis B Immunity (IgG) (Venous)	HBIM	В	1 day	95
Hepatitis B Profile	HEPB	В	1 day	96, 104

TEST	CODE	SAMPLE REQS	TAT	PAGE
Hepatitis B Resistant Mutation	HBRM	(A) or (B)	7 days	96
Hepatitis B Surface Antigen (Self-collect)	THBA	(TDL Tiny)	1 day	72, 96,
				160
Hepatitis B Surface Antigen (Venous)	AUAG	В	1 day	72, 96
Hepatitis C Abs Confirmation (RIBA)	RIBA	<b>B</b>	5 days	96
Hepatitis C Antibodies (Self-collect)	THCV	(TDL Tiny)	1 day	73, 96,
				160
Hepatitis C Antibodies (Venous)	HEPC	<u>B</u>	1 day	73, 96
Hepatitis C Antigen (Early detection)	TCAG	(TDL Tiny)	1 day	73, 96,
(Self-collect)				160
Hepatitis C Antigen (Early detection) (Venous)	HCAG	8	1 day	73, 96
Hepatitis C Genotype	CGEN	A or B	5 days	96
Hepatitis C Quantification (Viral Load)	QPCR	A or B	5 days	96
Hepatitis Delta Antibody	HEPD	8	5 days	96
Hepatitis Delta Antigen	HDAG	<u>B</u>	5 days	96
Hepatitis Delta RNA	DRNA	<b>A</b>	5 days	96
Hepatitis E IgG/IgM	HBE	B	5 days	96
Hepatitis E RNA (PCR)	EHEP	A	2 weeks	96
Hepatitis G (PCR)	HEPG	(Frozen plasma)	2 weeks	96
Hereditary Colorectal Cancer NGS Panel	GENE	<b>A A</b> 9,11	4 weeks	122
Hereditary Comprehensive Cancer NGS Panel	GENE	<b>A A</b> 9,11	5 weeks	122
Hereditary Neuropathy with Liability to	GENE	<b>A</b> 9	6 weeks	122
Pressure Palsy – PMP22 deletion analysis				
Herpes Simplex (HSV) 1 & 2 – Genital lesion	HERS	Aptima multisite swab	5 days	73, 102,
(Self-collect)	HEDC	Antinea modificita conch	E dava	160
Herpes Simplex (HSV) 1 & 2 – Oral lesion (Self-collect)	HERS	Aptima multisite swab	5 days	73, 102, 160
Herpes Simplex (HSV) 1 & 2	HERS	PCR	5 days	73, 102
(PCR) (Oral or Genital)	IILIIO	1 011	Judyo	70, 102
Herpes Simplex I/II (Thin Prep)	HERD	TPV	5 days	170
Herpes Simplex I/II Antibody Profile (IgG)	HERP	ß	2 days	102
Herpes Simplex I/II by PCR (Urine)	HERD	FCRU	5 days	73, 102
Herpes Simplex I/II IgM	HERM	ß	2 days	102
HFE gene (Haemochromatosis) –	HMD	<b>A</b> 9	3 days	43, 122
common variants C282Y + H63D				-,
Hirsutism Profile	HIRP	В	1 day	58, 63
Histamine (Blood)	HITT	(Frozen plasma)	5 days	85
Histamine (Urine)	HITU	RU	5 days	85
Histamine Releasing Urticaria Test	CURT	В	3 weeks	85, 141
Histone Antibodies	HISA	<u>B</u>	5 days	85
Histopathology				175
Histoplasma Antigen	HANT	RU	3 days	48
Histoplasmosis	HISP	B	10 days	85
- ·			•	

HIV 1 & 2 Abs/p24Ag (Self-collect)  HIV 1 & 2/p24Ag (Venous)  HDU0 ③ 1 day  73, 160  HIV-1 Genotypic Resistance (Integrase)  HIV-1 Genotypic Resistance (RT & Protease)  HIVD ② (2 x 6ml)  21 days  101  HIV-1 Genotypic Resistance (RT & Protease)  HIVD ② (2 x 6ml)  21 days  101  HIV-1 Proviral DNA  HIVP ② 7 days  101  HIV-1 RNA Viral Load by PCR  HIV1 ② (2 x 6ml)  3 days  101  HIV-1 Tropism  TRPM ② (2 x 6ml)  28 days  101  HIV-2 RNA by PCR  HIV2 ③ 10 days  101  HIV-2 RNA by PCR  HIVC ③ 10 days  101  HIV Confirmation of Positive  Screens (3 methodologies)  HIV Rapid RNA HIV-1 QUALITATIVE  HIV Q (Vacutainer only)  HIV Rapid RNA HIV-1 QUANTITATIVE  RHIV ② (Vacutainer only)  HIV Screening: HIV1 & 2 Abs/p24 Ag (4th Gen)  HDU0 ⑤ 1 day  101  HIV Screening: HIV1 & 2 Abs/p24 Ag (4th Gen)  HDU0 ⑥ 1 day  101  HIV Therapeutic Drug Monitoring  TDM J¹ 21 days  101  HIV-HIV-HIV-HIV-HIV-HIV-HIV-HIV-HIV-HIV-
HIV-1 Genotypic Resistance (Integrase) HIVD
HIV-1 Genotypic Resistance (RT & Protease) HIVD
HIV-1 Proviral DNA       HIVP       7 days       101         HIV-1 RNA Viral Load by PCR       HIV1       (2 x 6ml)       3 days       101         HIV-1 Tropism       TRPM       (2 x 6ml)       28 days       101         HIV-2 RNA by PCR       HIV2       10 days       101         HIV Confirmation of Positive Screens (3 methodologies)       HIVC       1 day       101         HIV Rapid RNA HIV-1 QUALITATIVE       LHIV       (Vacutainer only)       1 day       73, 78, 101, 104         HIV Rapid RNA HIV-1 QUANTITATIVE       RHIV       (Vacutainer only)       1 day       73, 78, 101, 104         HIV Screening: HIV1 & 2 Abs/p24 Ag (4th Gen)       HDU0       3 days       104       101         HIV Therapeutic Drug Monitoring       TDM       J¹       21 days       101         HIV/HBV/HCV (Early detection by       STXX       3 days       73, 77
HIV-1 RNA Viral Load by PCR HIV1
HIV-1 Tropism         TRPM         ③ ② (2 x 6ml)         28 days         101           HIV-2 RNA by PCR         HIV2         ①         10 days         101           HIV Confirmation of Positive         HIVC         ①         1 day         101           Screens (3 methodologies)         HIV Rapid RNA HIV-1 QUALITATIVE         LHIV         ④ (Vacutainer only)         1 day         73, 78, 101, 104           HIV Rapid RNA HIV-1 QUANTITATIVE         RHIV         ④ (Vacutainer only)         1 day         73, 78, 101, 104           HIV Screening: HIV1 & 2 Abs/p24 Ag (4th Gen)         HDU0         ③         1 day         101           HIV Therapeutic Drug Monitoring         TDM         J¹         21 days         101           HIV/HBV/HCV (Early detection by         STXX         ⑤ ② 2 x 6mls or 2 x 4mls         3 days         73, 77
HIV-2 RNA by PCR         HIV2         10 days         101           HIV Confirmation of Positive Screens (3 methodologies)         HIVC         1 day         101           HIV Rapid RNA HIV-1 QUALITATIVE         LHIV         (Vacutainer only)         1 day         73, 78, 101, 104           HIV Rapid RNA HIV-1 QUANTITATIVE         RHIV         (Vacutainer only)         1 day         73, 78, 101, 104           HIV Screening: HIV1 & 2 Abs/p24 Ag (4th Gen)         HDU0         3         1 day         101           HIV Therapeutic Drug Monitoring         TDM         J¹         21 days         101           HIV/HBV/HCV (Early detection by         STXX         3 2 x 6mls or 2 x 4mls         3 days         73,77
HIV Confirmation of Positive Screens (3 methodologies)       HIVC       ①       1 day       101         HIV Rapid RNA HIV-1 QUALITATIVE       LHIV       ② (Vacutainer only)       1 day       73, 78, 101, 104         HIV Rapid RNA HIV-1 QUANTITATIVE       RHIV       ③ (Vacutainer only)       1 day       73, 78, 101, 104         HIV Screening: HIV1 & 2 Abs/p24 Ag (4th Gen)       HDU0       ③       1 day       101         HIV Therapeutic Drug Monitoring       TDM       J¹       21 days       101         HIV/HBV/HCV (Early detection by       STXX       ③ ② 2 x 6mls or 2 x 4mls       3 days       73, 77
Screens (3 methodologies)
HIV Rapid RNA HIV-1 QUANTITATIVE   RHIV   (A) (Vacutainer only)   1 day   73, 78, 101, 104   101,
101, 104
HIV Therapeutic Drug Monitoring TDM J¹ 21 days 101  HIV/HBV/HCV (Early detection by STXX 3 days 73,77
HIV/HBV/HCV (Early detection by STXX
i ontinunt finan oppinio
HIV/HBV/HCV Screen by PCR/         STDX         2 x 6mls or 2 x 4mls         3 days         73,77,           NAAT (10 days post exposure)         (Vacutainer only)         101-102,           104         104         104
<b>HLA A, B, C</b> 14RF <b>(A) (A)</b> 2 weeks 61
HLA B*57:01 HL57 🐧 9 10 days 101
<b>HLAB 27</b> HLAB <b>Q</b> 9 3 days 85
HLA DQ Alpha Antigens 10RF (2) (A) 2 weeks 61
HLA DQ Beta Antigens 11RF QQ 2 weeks 61
HLA DR Antigens 9RF (2) (2) weeks 61
<b>HLA-C</b> 26RF <b>Q Q</b> 2 weeks 61
Homocysteine (Quantitative) HOMO 1 of 17 or (A) (Plasma) 1 day 34
Homocysteine (Urine) HCYS CU 2 weeks 34
Homovanillic Acid (HVA) HVA PU 5 days 34
Horse Components ZZ38 3 2 days 142
House Dust Mite Components ZZ12 3 2 days 142
HPV (19 high risk DNA subtypes, reported HPVY Qvintip vaginal swab 3 days 73, 160, as types 16, 18 or Others) (Self-collect) 173
HPV (28 individually typed low risk (LR) & HPVT TPV 5 days 73, 172 high risk (HR) DNA subtypes and reflexed mRNA for types 16, 18, 31, 33 and 45)
HPV (28 individually typed HP20 TPV 3 days 73, 172 LR & HR DNA subtypes)
HPV (A group of 14 HR mRNA types) HPVH TPV 3 days 73, 172
HPV (Individually typed high risk DNA subtypes)HPVZQvintip vaginal swab3 days73, 160,(Self-collect)173
HRT Profile 1 HRT (3) 1 day 58, 63

TEST	CODE	SAMPLE REQS	TAT	PAGE
HRT Profile 2	HRT2	BG	1 day	58, 63
HTLV 1 & 2 Abs.	HTLV	B	1 day	101
(Human T Lymphotropic Virus Type I-II)				
HTLV by PCR	HTLP	A	21 days	101
Hughes Syndrome	LUPA	B @ @ 4,18	2 days	42
Human Herpes Virus – 6 by PCR	HHV6	A	5 days	103
Human Herpes Virus – 8 (IgG)	HHV8	В	10 days	103
Human Herpes Virus – 8 by PCR	HV8D	A	5 days	103
Human Parvovirus B19 – DNA	PCRP	A	2 weeks	103
Huntington Disease –	GENE	<b>A A</b> 9,11	5 weeks	123
HD gene repeat analysis PCR				
HVS	HVS	STM/CS	2-4 days	48
Hyaluronic Acid	AHT	В	1 week	34
Hydroxybutyrate Dehydrogenase	HBD	(Frozen)	1 week	34
Hydroxyprolene	UHYD	CU	2 weeks	34
Hyperinsulinism NGS Panel	GENE	<b>A A</b> 9	6 weeks	123
Hyperparathyroidism – CASR sequencing	GENE	<b>A</b> 9	6 weeks	123
IDH1/2 Screening Assay	GENE	<b>A</b>	48 hours	123
IgE (Total)	IGE	В	1 day	85
IGF-1 (Somatomedin)	SOMA	(Frozen) <sup>4,7</sup>	1 day	58
IGF-BP3	IGF3	⊕ (Frozen) <sup>4</sup>	5 days	58
IgG Subclasses	IGSC	В	5 days	34
IgVH variant analysis for CLL	IGMU	<b>A</b>	4 weeks	123
Imipramine	IMIP	<b>A</b> 4	4 days	138
Immune Function Evaluation (Total)	TIE	A + B 5,10	7 days	41
Immune-Complexes	IMCP	В	5 days	85
Immunofluorescence in Skin Biopsies	IHCS	Skin sample in Michels solution	2 weeks	85
Immunoglobulin A	IGA	В	1 day	34
Immunoglobulin D	IGD	8	5 days	34
Immunoglobulin E – Total	IGE	В	1 day	34
Immunoglobulin G	IGG	В	1 day	34
Immunoglobulin M	IGM	В	1 day	34
Immunoglobulins (IgG, IgM, IgA)	IMM	В	1 day	34, 85
Impotence Profile	IMP0	<b>ABBG</b>	3 days	58, 63
Individual Semen Parameters***	SPOD	Semen 1	1 day	68
Inherited bleeding and platelet disorders (R90)	R90U	00	12 weeks	123
Inhibin A	INIA	В	1 month	58
Inhibin B	INIB	(Day 3 of cycle, frozen)	5 days	58
INR	PTIM	<b>C</b> 18	1 day	41
Insect/Worm/Ova/Cysts	FLEA	Send Specimen <sup>9,14</sup>	5 days	92
Insulin	INSU	B 4,7	1 day	58
Insulin Antibodies	INAB	В	5 days	85
Insulin-Like Growth Factor 2	IGF2	<b>B</b> 6	1 month	34

TEST	CODE	SAMPLE REQS	TAT	PAGE
Intellectual Disability NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	123
Interleukin 1 Beta	ILB	(Frozen) <sup>4,7</sup>	1-2 weeks	85
Interleukin 2	IL2	(Frozen) <sup>4,7</sup>	1-2 weeks	85
Interleukin 4	IL4A	(Frozen) <sup>4,7</sup>	1-2 weeks	85
Interleukin 6	IL6	(Frozen) <sup>4,7</sup>	1-2 weeks	85
Interleukin 8	IL8	(Frozen) <sup>4,7</sup>	1-2 weeks	85
Interleukin 10	IL10	(Frozen) <sup>4,7</sup>	1-2 weeks	85
Interleukin 28b Genotype	IL28	A	2 weeks	85
Intrinsic Factor Antibodies	IFAB	B	2 days	85
lodide – Urine	UIOD	RU	1 week	34
lodine – Serum	IODI	B	1 week	34
Ionised Calcium	ICPA	B	5 days	34
Iron (TIBC included) (Venous)	FE	B	1 day	34
Iron Overload Profile	IOP	<b>△ B</b> <sup>9</sup>	3 days	34, 39,
				123, 130
Iron Status Profile (Venous)	ISP	B	1 day	34, 39
ISAC Panel (Self-collect)	ISAC	(TDL Tiny)	3 days	141, 160
ISAC Panel (Venous)	ISAC	В	3 days	141, 146
Islet Cell Antibodies	ICAB	В	3 days	85
IUCD for Culture	IUCD	Send Device	11-12 days	48
JC Polyoma Virus by PCR	JCPV	(A) / CSF	5 days	103
Joubert/Meckel-Gruber Syndrome NGS Panel	GENE	<b>A</b> 9	6 weeks	123
Kallmann Syndrome NGS Panel	GENE	<b>AA</b> <sup>9</sup>	6 weeks	123
Kennedy Disease (Spinal Bulbar	GENE	<b>A</b> 9	5 weeks	123
Muscular Atrophy) – AR repeat expansion				
Ketamine Screen	KETA	RU	7-10 days	163
Kidney/Urinary Tract Comprehensive	GENE	<b>A A</b> 9,11	4 weeks	123
Cancer NGS Panel	17DF	000	0.0 wooko	61
KIR (Killer-like Immunoglobulin-like Receptors) Genotyping	17RF	000	2-3 weeks	01
Kiwi Components	ZZ32	B	2 days	142
Lactate (Plasma)	LACT	<b>G</b> 16	1 day	34
Lactate Dehydrogenase (LDH)	LDH	B	1 day	34
Lactate Pyruvate Ratio	LPR	J <sup>1</sup>	4-6 weeks	34
Lactose Tolerance Test	LTT	By appointment only	1 day	34, 137
Lamotrigine	LAMO	B 4	5 days	138
Latex Components	ZZ13	B	2 days	142
LDL7 Subfractions	LDL7	<u> </u>	10 days	34
Lead (Blood)	LEAD	A	5 days	35, 165
Lead (Urine)	URPB	RU	5 days	35, 165
Leber's Hereditary Optic Neuropathy	GENE	<b>A</b> 9	6 weeks	123
- m.3460G>A + m.11778G>A +		•		.20
m.14484T>C common variants				

TEST	CODE	SAMPLE REQS	TAT	PAGE
Legionella Antibodies	LEG0	В	3 days	85
Legionella Urine Antigen	LEGA	Urine with boric acid	1 day	48, 85
Leishmania Antibodies	LEIS	B	5 days	92
Leptin	LEPT	(height & weight required) 19	5 days	35
Leptospirosis (Weil's Disease) Abs (IgM)	LEP	В	5 days	85
Leucocyte Antibody Detection Panel FEMALE	8RF	В	1 week	61
Leucocyte Antibody Detection Panel MALE	7RF	<b>A A B A</b> 6,34	1 week	61
Leukaemia (Rapid Acute) DNA	ALRP	(3mL minimum) or bone	3 days	124, 131
and RNA NGS Panel		marrow aspirate sample		
Leukaemia Fusion Gene Screening Assay (Q30)	LMPX	A	24 hours	124
Leukaemia Immunophenotyping	LYPT	<b>A</b> 4,5	5 days	44
Leukaemia/Lymphoma RNA Sequencing	PHFP	A	2 weeks	124, 131
(Fusion Gene and SNV/Indel) Panel				
Leukotriene E4	LTE4	CU (Frozen)	3 weeks	85
Levetiracetam (Keppra)	LEVE	<b>B</b> 4	3 days	138
Li-Fraumeni Syndrome (p53-related cancer	GENE	A 9,11	6 weeks	124
predisposition) – TP53 sequencing + MLPA		0.00		
Limb-Girdle Muscular Dystrophy	GENE	<b>A A</b> <sup>9</sup>	6 weeks	124
(LGMD) NGS Panel Lipase (Self-collect)	LIPA	(TDI Tipu)	1 day	35, 160
	LIPA	B (TDL Tiny)		
Lipase (Venous) Lipid Profile (Self-collect)	LIPA	(I) (TDL Tiny)	1 day	35, 40 35, 160
Lipid Profile (Venous)	LIPP	B (TDL TIIIY)	1 day	35, 100
Lipid Transfer Proteins	ZZ23	<u>B</u>	2 days	142
Lipocalins	ZZ28	<u>В</u>		142
Lipoprotein (a) (Self-collect)	LPOA	B (TDL Tiny)	2 days 1 day	35, 160
,,	LPOA	B	1 day	35, 100
Lipoprotein (a) (Venous) Lipoprotein Electrophoresis	LEL	<u>В</u>	5 days	35
Lissencephaly NGS Panel	GENE	<b>AA</b> <sup>9</sup>	6 weeks	124
Lithium (take 12 hours after dose)	LITH	B	1 day	35, 138
Liver Fibrosis (Enhanced Liver Fibrosis ELF)	ELF	<u> </u>		
Liver Fibrosis Fibrotest	FIBT	<u>В</u>	5 days 2 weeks	35 35
Liver Function Tests (Excluding	TLFT	(TDL Tiny) and (B) (TDL Tiny)		35, 160
AST) (Self-collect)	ILITI	(TDE TIIIY) and (TDE TIIIY)	i uay	33, 100
Liver Function Tests (Venous)	LFT	В	1 day	35, 40
Liver Immunoblot	LIVI	В	3 days	85
Liver Kidney Microsomal Antibodies	LKM	В	2 days	85
Long QT Syndrome/Brugada Syndrome	GENE	<b>AA</b> <sup>9</sup>	4-6 weeks	124
NGS Panel	QLIIL.		. 5 .700110	ILT
Lorazepam	LORA	<b>A</b> 4	10 days	138
Lp-PLA2 (PLAC) Test	PLA2	В	2 days	35
LSD	LSD	RU	5 days	163
Lung Disorders NGS Panel	GENE	<b>AA</b> <sup>9</sup>	6 weeks	124
9				

TEST	CODE	SAMPLE REQS	TAT	PAGE
Lupus Anticoagulant and Anticardiolipin Abs	LUPA	B C C 4,18	2 days	42, 85
Lupus Anticoagulant only	LUPC	<b>C C</b> 9,18	2 days	42
Luteinising Hormone (LH) (Self-collect)	LH	(TDL Tiny) or (TDL Tiny)	1 day	58, 160
Luteinising Hormone (LH) (Venous)	LH	B	1 day	58
Lyme Disease (Borrelia Abs) IgG, IgM	BORR	B 9,14	2 days	85
Lyme Disease (Borrelia Abs) IgM	BORM	В	2 days	85
Lymphocyte Subsets (CD3/CD4/CD8)	LYSS	A	1 day	41, 101
Lymphogranuloma Venerium (LGV)	LGVP	Aptima multisite swab	1-2 weeks	73, 160
- Rectal (Self-collect)*				
Lymphogranuloma Venerium (LGV) (PCR)	LGVP	PCR*42	1-2 weeks	73
Lynch Syndrome (HNPCC) NGS Panel	GENE	<b>A</b> 9	4 weeks	124
Lysosomal Enzyme Screen	LE	<b>J</b> <sup>1</sup>	2 months	35
Lysosomal Storage Disorders NGS	LSDS	<b>A A</b> <sup>9</sup>	4-6 weeks	124, 131
Panel – full gene sequencing				
Lysozyme	LYS0	В	5 days	35
Macrolide Resistance Test (Mgen)	MGR	FCRU / PCR	1-2 weeks	73
Macroprolactin	PRLD	B	4 days	58
Magnesium (Serum)	MG	В	1 day	35, 165
Magnesium (Urine)	URMG	PU	1 day	35, 165
Magnesium (Whole blood)	RCMG	A or (1)	4 days	152
Malarial Antibodies (Pl. falciparum)	MALA	B 9,14	5 days	92
Malarial Antibodies (species specific)	MALS	<b>B</b> 9,14	10 days	92
Malarial Parasites	MALP	A 4,9,14	STAT	41
Malarial Parasites (visa, non-urgent)	MP48	<b>A</b>	2 days	41
Male Genetic Reproductive Profile	GRP	<b>A</b> (1) 9	10-15 days	124, 130
Male Hormone Profile	MIPR	В	1 day	59, 63
Manganese (Serum)	MANG	В	5 days	35, 165
Marfan Syndrome – FBN1 sequencing + deletions/duplications	GENE	<b>(A</b> ) 9	6 weeks	124
Marfan Syndrome and Thoracic Aortic	GENE	<b>A A</b> <sup>9</sup>	6 weeks	124
Aneurysm and Dissection NGS Panel				
Maturity-Onset Diabetes of the Young (MODY) Diabetes NGS Panel	GENE	<b>A</b> 9	12 weeks	124
MBOCA in Urine	MBOC	RU	10 days	166
Mean Cell Volume (MCV)	MCV	A	1 day	41
Measles Antibodies (IgG) Immunity	MEAS	В	1 day	95, 103
Measles Antibodies (IgM)	MEAM	<b>B</b> 9	2 days	95, 103
Measles PCR	MEAP	Buccal swab	48 hours	103
Measles, Mumps, Rubella (MMR)	MMR	В	1 day	95
Meckel-Gruber/Joubert Syndrome NGS Panel	GENE	<b>A</b> 9	6 weeks	124
Melanoma Comprehensive Cancer NGS Panel	GENE	<b>A A</b> 9,11	4 weeks	124
Melatonin (Serum)	MEL	B (Frozen)	5 days	59
Melatonin (Urine)	UMEL	CU 13	2 weeks	59
	UNILL		= .100110	

TEST	CODE	SAMPLE REQS	TAT	PAGE
Meningococcal Serology (only serogroup C)	MENI	В	6 weeks	85
Menopausal Profile (FSH, LH, TOES, TSH, FT4) (Self-collect)	TMEN	(TDL Tiny) and (3) (TDL Tiny)	1 day	59, 160
Menopause Profile (Venous)	MENO	В	1 day	59, 63
Mercury (Blood)	MERC	(A) or (B)	5 days	35, 165
Mercury (Urine)	URHG	RU <sup>1</sup>	5 days	35, 165
MERS Coronavirus Test	MERS	J	1 day	103
Metabolic Syndrome Profile	METS	<b>ABBG</b>	9 days	59, 63
Metanephrines (Plasma)	PMET	(Frozen plasma, must be frozen within 2 hours)	7 days	59
Metanephrines (Urine)	UMEX	PU (collect on acid) <sup>1</sup>	5 days	59
Methaqualone	METQ	RU	5 days	35
Methotrexate	METX	В	2 days	138
Methylmalonic Acid – Serum	MMAS	B	5 days	35
Methylmalonic Acid – Urine	MMA	CU	2 weeks	35
Metronidazole Level	METR	<b>B</b> 4	10 days	137
Microfilaria Blood Film	MICF	A	STAT	41
Mineral Screen	MINE	<b>B ((</b>	5 days	152-153
Mineral Screen (Whole blood)	RMIN	00	5 days	152-153
Mineral Screen and Industrial Heavy Metal Screen (Trace Metals)	TRAC	<b>4800</b>	7-10 days	152-153
Miscarriage/Thrombotic Risk Profile	PROP	<b>AABCC</b> 18	5 days	43, 45
Mitochondrial Antibodies	AMIT	В	3 days	85
Mitochondrial Antibodies M2	MTM2	В	2 days	85
Mitochondrial Genome Sequencing	GENE	<b>A</b> 9	6 weeks	125
Molybdenum (Serum)	MOLY	B	5 days	166
Motor Neurone Disease	GENE	<b>A A</b> <sup>9</sup>	5 weeks	125
(Amylotrophic Lateral Sclerosis) NGS Panel				
MPOX (Virus) – Lesion (Self-collect)	MPXV	Aptima multisite swab	2 days	73, 160
MRSA (Rapid PCR) one swab per site	MRSA	Blue liquid Amies swab	1 day	48
MRSA (Rapid PCR) one swab per site x 2	MRS2	Blue liquid Amies swab x 2	1 day	48
MRSA Culture (Self-collect) – Nose/Groin	MRW2	Purple liquid Amies swab x2	2 days	48, 160
MRSA Culture one swab per site	MRSW	Blue liquid Amies swab	2 days	48
MRSA Culture one swab per site x 2	MRW2	Blue liquid Amies swab x 2	2 days	48
MRSA PCR (Self-collect) – Nose/Groin	MRS2	Purple liquid Amies swab x2	1 day	48, 160
MTHFR – common C677T + A1298C variants	MTHF	<b>A</b> <sup>9</sup>	5 days	125
Mucopolysaccharides	MPS	RU (Frozen)	3 weeks	35
Mucopolysaccharidosis NGS Panel	GENE	<b>AA</b> <sup>9</sup>	6 weeks	125
Multiple Endocrine Neoplasia Type 1 – full MEN1 sequencing	GENE	<b>A</b> 9,11	6-7 weeks	125
Multiple Endocrine Neoplasia Type 2  – RET gene hotspot sequencing	GENE	<b>A</b> 9,11	6-7 weeks	125
Mumps Antibodies (IgG)	MUMP	B	1 day	95, 103

TEST	CODE	SAMPLE REQS	TAT	PAGE
Mumps Antibodies (IgM)	MUMM	B	1 day	95, 103
Myasthenia Gravis Evaluation	MGE	B	5 days	85
Mycophenolic Acid (Cellcept)	MYCP	A	5 days	138
Mycoplasma genitalium (Thin Prep)	MGEN	TPV	2 days	170
Mycoplasma genitalium by PCR	MGEN	FCRU / PCR / TPV	2 days	73
Mycoplasma genitalium by PCR – Urine and Vaginal (Self-collect)	MGEN	Aptima urine or multisite swab	2 days	74, 161
Mycoplasma genitalium Resistance  – Urine or Vaginal (Self-collect)	MGR	Aptima urine or multisite swab	1-2 weeks	74, 161
Mycoplasma genitalium/Ureaplasma (Thin Prep)	MUPC	TPV	2 days	170
Mycoplasma genitalium/Ureaplasma by PCR	MUPC	FCRU / PCR / TPV	2 days	74
Mycoplasma pneumoniae IgM and IgG	MYC0	В	2 days	103
Mycoplasma species – DNA	MPCR	A	5 days	103
Myelin Associated Glycoprotein Antibodies	MAG	B	5 days	86
Myelin Basic Protein Antibodies	MBPA	B	2 weeks	86
Myeloid Gene Panel	MVPS	(3mL minimum) or bone marrow aspirate sample	2 weeks	125, 131
Myeloma Screen CHANGE	MYEL	<b>AB</b> 6	5 days	35, 40
Myeloperoxidase Antibodies	MP0	B	2 days	86
Myeloproliferative Neoplasm NGS Screening Panel	MPNS	(3mL minimum) or bone marrow aspirate sample	1 week	125, 131
Myocardial Antibodies	MY0	B	1 week	86
Myoglobin (Serum)	SMY0	B	1 day	35
Myoglobin (Urine)	UMY0	RU	5-10 days	35
Myositis Panel	MYOS	<b>B</b>	3 days	86
Myotonic Dystrophy Type 1 – DMPK repeat PCR	GENE	<b>A</b> 9	5 weeks	125
Myotonic Dystrophy Type 2 (PROMM) – ZNF9 repeat PCR	GENE	<b>A</b> 9	6 weeks	125
Mysoline (Primidone)	PRIM	B 4	3 days	138
Narcolepsy (HLA DQB1*06:02)	GENE	<b>A</b> 9	3 weeks	125
Natural Killer Profile 2	NKP2	<b>A</b> 10	2 days	41, 45
Needle Stick Injury Profile	NSI	88	1 day	103, 105
Nephrotic Syndrome, Steroid- Resistant NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	125
Nervous System/Brain Cancer NGS Panel	GENE	<b>A A</b> 9,11	4 weeks	125
Neurofibromatosis Type 1 – NF1 + SPRED1 sequencing + deletions/duplications	GENE	<b>A A</b> 9,11	8 weeks	125
Neurological Viral Screen	NVIR	88	2 days	103, 105
Neuronal Antibody (Hu, Ri, Yo, Cv2, Ma2)	NEUR	8	10 days	86
Neurone Specific Enolase	NSE	B	5 days	106
Newborn Screening Panel	GUTH	<b>J</b> <sup>1</sup>	2 weeks	35
Nickel (Serum)	NICK	<b>B</b>	5 days	35, 165

TEST	CODE	SAMPLE REQS	TAT	PAGE
Nickel (Urine)	NICU	RU	4 weeks	35, 165
NK (CD69) and NK Cytotoxicity	69C	000*	Send Mon-Thurs only	62
NK (CD69) Cell Assay	CD69	<b>B</b> *	Send Mon-Thurs only	62
NK Assay Follow-Up Panel	5RF	000	1 week	61
NK Assay Panel + Intralipids	16RF	000	1 week	61
NK Assay/Cytotoxicity Panel	4RF	000	1 week	61
NK Cytotoxicity Assay	HSNK	000*	Send Mon-Thurs only	62
NK Cytotoxicity with suppression with steroid,	69CI	000*	Send Mon-Thurs only	62
IVIg and intralipin, and NK (CD69) cell assay				
NK Cytotoxicity with suppression,	NKCY	000*	Send Mon-Thurs only	62
steroid, IVIg & Intralipin				
NMDA Receptor Antibodies	NMDA	В	3 weeks	86
Non-Invasive Prenatal Testing (NIPT)	NIPT	J / Special tube <sup>1</sup>	2-4 days	126, 136
- common aneuploidy screening				
from maternal blood	OFNE	000	0	400
Noonan Syndrome and RASopathies NGS Panel	GENE	<b>AA</b> <sup>9</sup>	6 weeks	126
Nucleic Acid Antigen Antibodies	DNA	B (TDI Time)	2 days	86
Oestradiol-17-Beta (Self-collect)	TOES	(TDL Tiny)	1 day	59, 161
Oestradiol-17-Beta (Venous)	0EST	8	1 day	59
Oestriol (Estriol)	E3	88	4 days	59
Oestrone	E1	88	4 days	59
Olanzapine	OLAN	<b>A</b> <sup>4</sup>	5 days	139
Oligoclonal Bands	CSF0	2ml <b>CSF</b> + 🔞	5 days	86
Oligosaccharides	UOLI	RU	6 weeks	36
Olive Components	ZZ14	B	2 days	142
Omega 3/Omega 6 (Self-collect)	OMG3	(TDL Tiny)	5 days	152, 154,
0	01400	04	F. da	161
Omega 3/Omega 6 (Venous)	OMG3	<b>A</b> 4	5 days	152, 154
Opiate Screen (Urine)	UOPI	RU	2 days	163
Orosomucoid (A1AG – Alpha 1 Glycoprotein)	OROS	(Frozen)	5 days	36
Osmolality (Serum)	OSMO	8	1 day	36
Osmolality (Urine)	ROSM	RU	1 day	36
Osteocalcin	0ST	(Frozen) <sup>4</sup>	4 days	59, 106
Osteogenesis Imperfecta NGS Panel	GENE	<b>AA</b> <sup>9</sup>	6 weeks	126
Osteoporosis Screen	0PS	88	4 days	36, 40
Ovarian Autoantibodies	OVAB	8	3 days	86
Ovarian Cancer NGS Panel	GENE	<b>A A</b> 9,11	4 weeks	126
Oxalate (Plasma)	POXA	(Frozen)	7 days	36
Oxalate (Urine)	UOXA	PU	5 days	36
Oxidative Stress in Semen (ROS + MIOXSYS)	SROS	Semen 1	1 day	68
P2Y12 Receptor Platelet Function	P2Y	J <sup>1</sup>	1 day	43
Analysis (Clopidogrel Resistance)				

TEST	CODE	SAMPLE REQS	TAT	PAGE
p53-related cancer predisposition	GENE	<b>(A)</b> 9,11	6 weeks	126
(Li-Fraumeni Syndrome) –				
TP53 sequencing + MLPA				
PAI-1 4G/5G Polymorphism	PAIP	A	10 days	41, 61
Pancreatic Cancer NGS Panel	GENE	<b>A A</b> 9,11	4 weeks	126
Pancreatic Peptide	PP	J	4 weeks	36
Paracetamol	PARA	В	1 day	139
Paragomius Serology	PRGM	В	2 weeks	86
Parathyroid Antibodies	PTHA	В	3 weeks	86
Parathyroid Hormone (Whole)	PTHI	<b>A</b> 4	1 day	59
Parathyroid Related Peptide	PTRP	2ml \Lambda Plasma frozen	2 weeks	36
		(Freeze immediately) <sup>1</sup>		
Parvalbumins	ZZ29	В	2 days	142
Parvovirus Antibodies (IgG)	PARG	<b>B</b>	2 days	103
Parvovirus Antibodies (IgM)	PARV	В	2 days	103
Parvovirus IgG/IgM Abs	PARP	В	2 days	103
Paul Bunnell (Monospot)	PAUL	(A) or (B)	1 day	41
Peach Components	ZZ15	В	2 days	142
Peanut Components	ZZ16	В	2 days	142
Pemphigus/Pemphigoid Autoantibodies	SKAB	В	2 days	86
Pertussis (Whooping Cough) Antibodies	PERS	В	5 days	86, 95
Pertussis (Whooping Cough) by PCR	PERP	Pernasal or dry swab	2-3 days	86
PEth (Phosphatidylethanol) (Self-collect)	PETH	(TDL Tiny)	5-7 days	36, 161, 163
PEth (Phosphatidylethanol) (Venous)	PETH	A	5-7 days	36, 163
Phencyclidine (PCP)	DUST	RU	5 days	36
Phenobarbitone	PHB	В	1 day	139
Phenytoin (Epanutin)	PHEN	В	1 day	139
Phosphate	PHOS	В	1 day	36
Phosphate (24 hour Urine)	UPH	PU	1 day	36
Pituitary Antibodies	PITU	<b>B</b> 4	1 month	86
Pituitary Function Profile	PITF	<b>BB</b> <sup>7</sup>	1 day	59, 64
PLAC Test (Lp-PLA2) (Self-collect)	PLA2	(TDL Tiny)	2 days	36, 161
PLAC Test (Lp-PLA2) (Venous)	PLA2	В	2 days	36
Plasminogen	PLAS	(Frozen plasma) <sup>4</sup>	5 days	36
Plasminogen Activator Inhibitor – 1	PAI1	(Frozen plasma)	2 weeks	36
Platelet Aggregation Studies	PLAG	<b>J</b> ** <sup>9</sup>	3 days	43
Platelet Function Test Screen - PFA-100/200	PFAT	<b>J</b> ** <sup>1</sup>	1 day	43
Pleural Fluid for Culture	FLUP	SC	7 days	48
Pneumococcal Antibodies – Serotype Specific	PASS	B	5 weeks	86
Pneumococcal Antibody Screen	PNEU	B	5 days	86, 95
Pneumococcal Antigen	PNAG	Urine with boric acid	1 day	48
Pneumocystis jiroveci (PJP) PCR	MPCP	SC BAL#	2-3 days	48

TEST	CODE	SAMPLE REQS	TAT	PAGE
Pneumonia (Atypical) Screen	APS	В	3 days	103
Polcalcins	ZZ25	B	2 days	143
Polycystic Ovary Syndrome Profile	PCOP	<b>ABBB</b> G <sup>7</sup>	5 days	59, 64
Polycystic Ovary Syndrome SHORT	PCOS	ABG	1 day	59, 64
Porphyrin (Blood)	PORP	<b>A</b> <sup>3</sup>	15 days	36
Porphyrin (Stool)	FPOR	RF <sup>3</sup>	3 weeks	36
Porphyrin (Urine)	RPOR	RU <sup>3</sup>	3 weeks	36
Porphyrin Full Screen (Total: Urine, Stool, Blood)	PORS	A RU, RF <sup>3</sup>	3 weeks	36, 40
Post-Travel Screen 1 (Up to 6 weeks post travel)	PTS	<b>AAB</b> G 14	10 days	92-93
Post-Travel Screen 2 (6 weeks after travel)	PTS2	AABBBG14	10 days	92-93
Postnatal SNP Array CGH	CGH	<b>A B</b> <sup>9</sup>	10 days	132
Potassium	K	B	1 day	36
PR-10 Proteins	ZZ22	B	2 days	143
Pre-Travel Screen (DVT)	DVT1	<b>A A B</b> <sup>9</sup>	5 days	42, 45,
,			,	92, 127
Prealbumin	PALB	В	3 days	141
Pregnancy (Serum) [Quantitative]	QHCG	В	1 day	36, 59
Pregnancy Test (Urine)	PREG	RU	1 day	36
Pregnenolone	PREN	В	15 days	59
Prenatal SNP Array CGH	CGH	Amniotic fluid, CVS or POC 9	10 days	132
Primidone (Mysoline)	PRIM	<b>B</b> 4	3 days	139
Procalcitonin	PCAL	(Frozen) <sup>4,7</sup>	1 day	36
Procollagen 1 Peptide N-Terminal (NTX)	P1NP	В	5 days	36
Procollagen 3 Peptide	PRC0	B	5 days	36
Products of Conception (BOBs + Culture)	PBK	Placental Sample 1,9	10-25 days	127, 130
Profilins	ZZ24	B	2 days	143
Progesterone (Self-collect)	PROG	(TDL Tiny) or (TDL Tiny)	1 day	59, 161
Progesterone (Venous)	PROG	В	1 day	59
Proinsulin	PROI	(Frozen plasma) <sup>4</sup>	5 days	59
Prolactin (Macro)	PRLD	В	4 days	59
Prolactin (Self-collect)	PROL	(TDL Tiny) or (TDL Tiny)	1 day	59, 161
Prolactin (Venous)	PROL	В	1 day	59
Propanalol	PR0	<b>B</b> 4	7 days	139
Propoxyphene	DPR0	RU	5 days	36
Prostate Cancer NGS Panel	GENE	<b>AA</b> 9,11	4 weeks	127
Prostate Profile (Total & Free PSA)	PR2	В	1 day	106, 108
Prostate Specific Antigen (Total) (Self-collect)	PSPA	(TDL Tiny)	1 day	106, 161
Prostate Specific Antigen (Total) (Venous)*	PSPA	В	1 day	106
Prostatic Acid Phosphatase	PACP	(Frozen)	3 days	36
Protein (Urine)	UPRT	CU	1 day	36
Protein 14.3.3 (Creutzfeldt–Jakob Disease)	CJD	J	5 weeks	36
Protein C Activity	PRC	(Frozen) <sup>4,9,18</sup>	3 days	43
Protein Electrophoresis incl. immunoglobulin	PRTE	B	5 days	36
		<del>-</del>		

TEST	CODE	SAMPLE REQS	TAT	PAGE
Protein S Activity	PS1	(Frozen) <sup>4,9,18</sup>	5 days	43
Protein S Free Ag	FPRS	(Frozen) <sup>4,9,18</sup>	3 days	43
Protein Total (Blood)	PROT	В	1 day	37
Protein/Creatinine Ratio (Urine)	UCPR	RU	1 day	37
Proteinase 3 Ab	PR3	В	2 days	86
Prothrombin Time	PTIM	<b>C</b> 18	1 day	42
Purkinje Cell Antibody (Hu and Yo)	PURK	В	10 days	86
Pyruvate Kinase (M2-PK)	M2ST	RF <sup>4</sup>	5 days	106
Pyruvate Kinase (M2-PK)	M2PK	(Frozen plasma) <sup>7</sup>	5 days	106
Q Fever (C Burnetti) Antibodies	QFEV	<b>B</b> 9	10 days	86
QFIT/Calprotectin Profile (Combined) (QFIT)	QCAL	QFIT	5 days	36
QFIT/Calprotectin Profile	QCAL	QFIT sample collection device	5 days	37, 161
(Combined) (Self-collect)				
Quantitative Faecal Immunochemical Test (QFIT)	QFIT	QFIT	1 day	37
Quantitative Faecal Immunochemical Test (QFIT) (Self-collect)	QFIT	QFIT sample collection device	1 day	37, 161
Rabies Antibody	RABI	B	20 days	95
Rapid Strep PCR (incl. m/c/s)	RAPS	Blue liquid Amies swab**	1-3 days**	48
Rapid Xpert HIV-1 RNA Qualitative – Early Detection from 10 days	LHIV	(Vacutainer only)	1 day	74
Rapid Xpert HIV-1 RNS Viral Load – Rapid Testing for HIV-Positive Patient Prognosis and Response To Antiretroviral Therapy	RHIV	(Vacutainer only)	1 day	74
Recurrent Miscarriage Profile (female)	RMP	<b>AABCCC</b> 19,18	10-15 days	127, 130
Renal Calculi Screen (Metabolic)	RSPR	<b>J</b> 6	5 days	37
Renal Stone Analysis	RSTA	STONE	10 days	37
Renin	RENI	(Frozen plasma) <sup>36</sup>	5 days	59
Reproductive Immunophenotype Panel	3RF	000	1 week	61
Respiratory PCR Panel (COVID-19, Flu A/B and RSV) (PCR)	FLU4	PCR nasopharyngeal swab	2 days	103-104,
Respiratory PCR Panel (COVID-19, Flu A/B and RSV) (Self-collect)	FLU4	Aptima multisite swab of nose/throat	2 days	103, 161
Reticulocyte Count	RETC	A	1 day	42
Retinol Binding Protein	RBP	В	3 days	37
Retrograde Ejaculation	RTR0	Contact lab	2 days	68
Reverse T3	RT3	B 7,37	15 days	59
Rheumatoid Factor (Latex Test)	RF	В	3 days	86
Rheumatology Profile 1 (Screen)	RH	AB	2 days	86, 90
Rheumatology Profile 2 (Connective tissue)	RH2	AABB	3 days	86, 90
Rheumatology Profile 3 (Rheumatoid/Basic)	RH3	<b>A</b> B	2 days	86, 90
Rheumatology Profile 4 (Systemic Lupus)	RH4	ABB	2 days	86, 90
Rheumatology Profile 5 (Mono Arthritis)	RH5	AABB	3 days	86, 90
Rheumatology Profile 6 (Rheumatoid Plus)	RH6	<b>B</b>	3 days	86, 90

TEST	CODE	SAMPLE REQS	TAT	PAGE
Rheumatology Profile 7 (Sjogren's Syndrome)	RH7	B	15 days	86, 90
Rickettsial Species Antibody Profile	RICK	<b>B</b>	7 days	86, 92
Risperidone	RISP	<b>A</b> 4	7 days	139
RNA Polymerase Antibodies	RNAP	<b>B</b>	3 days	86
Rotavirus in Stool by PCR	ROTA	RF	1 day	103
RPR (Syphilis)	RPR	B	2 days	74, 86
Rubella Antibody (IgG)	RUBE	<b>B</b>	1 day	95, 103
Rubella Antibody (IgM)	RUBM	<b>B</b>	1 day	95, 103
Rubella Avidity	RUAV	<b>B</b>	1 week	103
Rubella PCR	RUBP	Amniotic Fluid	5 days	95
S100 Malignant Melanoma	S100	<b>B</b>	4 days	106
Saccharomyces Cerevisiae Antibodies	ASCA	<b>B</b>	2 weeks	86
Salicylates	SALI	<b>B</b>	1 day	37
Salivary Duct Antibodies	SAB	<b>B</b>	15 days	86
Schistosoma (Urine)	USCH	Mid-morning terminal urine following exercise <sup>14</sup>	1-2 days	48
Schistosome (Bilharzia) Antibodies	BILH	B 14	10 days	92
Scleroderma Immunoblot	SCLI	<b>B</b>	3 days	87
Screening Profile 1 – Biochemistry CHANGE	PP1	<b>3 G</b>	1 day	26
Screening Profile 2 – Haematology/ Biochemistry CHANGE	PP2	<b>43</b> 6	1 day	26
Screening Profile 3 – Haematology	PP3	A	1 day	26
Screening Profile 4 – Haematology/ Biochemistry (Short) CHANGE	PP4	<b>A</b> 36	1 day	26
Screening Profile 5 – Haematology/ Biochemistry (Postal) CHANGE	PP5	<b>A B G</b>	1 day	26
Screening Profile 6 – Well Person CHANGE	PP6	<b>A</b> BG	1 day	26
Screening Profile 7 – Well Man CHANGE	PP7	ABG	1 day	27
Screening Profile 8 – Well Person CHANGE	PP8	ABG	2 days	27
Screening Profile 9F – Senior Female CHANGE	PP9F	ABBGRU4	2 days	27
Screening Profile 9M – Senior Male CHANGE	PP9M	ABBGRU <sup>4</sup>	2 days	27
Screening Profile 10 – Cardiovascular Risk 1	PP10	88	3 days	27
Screening Profile 11 – Cardiovascular Risk 2	PP11	BBB 0 34	3 days	27
Screening Profile 12 – Sexual Health Screen	PP12	FCRU / PCR / TPV	2 days	27
Seed Storage Proteins	ZZ26	B	2 days	143
Selenium (Serum) (Self-collect)	SELE	(TDL Tiny)	4 days	37, 152, 161
Selenium (Serum) (Venous)	SELE	В	4 days	37, 152
Self-collection samples		<del>-</del>		155
Sellotape Test	SELL	Send Sample***	1 day	48
Semen Analysis, Comprehensive*	SPER	Semen 1	2 days*	68
Semen Analysis, Post-Vasectomy**	PVAS	Semen 1	2 days	68
Semen Analysis, Vasectomy Reversal*	SPER	Semen 1	2 days*	68

TEST	CODE	SAMPLE REQS	TAT	PAGE
Semen Culture	SPCU	Semen	2-4 days	48, 68
Semen Fructose (Qualitative assessment)	SPCF	Semen	2 days	68
Semen Leucocytes	PMNS	Semen	2 days	68
Semen Zinc	SPCZ	Semen	up to 10 days	68
Serotonin	SERT	(Frozen whole blood) <sup>1</sup>	10 days	59
Serotonin (Urine)	USER	PU 50mls (Frozen) <sup>1</sup>	5 days	60
Serum Albumins	ZZ30	B	2 days	143
Serum Free Light Chains	SLC	B	5 days	37
Sesame Components	ZZ39	В	2 days	143
Sex Hormone Binding Globulin (Self-collect)	SHBG	(TDL Tiny) or (TDL Tiny)	1 day	60, 161
Sex Hormone Binding Globulin (Venous)	SHBG	B	1 day	60
Short Stature – SHOX variant screening + deletions/duplications	GENE	<b>A</b> 9	8 weeks	127
Short-Chain Acyl-CoA Dehydrogenase Deficiency – ACADS sequencing	GENE	<b>A</b> 9	6 weeks	127
Shrimp Components	ZZ17	В	2 days	143
Silver (Blood)	SILV	В	5 days	37, 165
Silver (Urine)	USIL	RU	5 days	37, 165
Silver-Russell Syndrome – methylation studies on 11p15 imprinting domains KvDMR + H19	GENE	<b>A</b> 9	7 weeks	128
Sinequan (Doxepin)	DOXE	A or (1)	10 days	139
Sirolimus	SIR0	A	3 days	139
Sjogren's Syndrome	RH7	В	15 days	87
Skeletal Dysplasia NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	128
Skin (Pemphigus/Pemphigoid) Autoantibodies	SKAB	В	2 days	87
Skin Antibodies by Immunofluorescence	STSK	В	1 month	87
Skin Scrapings/Mycology by PCR	DERM	Send Sample	3-7 days	48
Sleeping Sickness Serology	TRYP	<b>B</b> 9	10 days	87
(African Trypanosomiasis)				
Smith-Lemli-Opitz Syndrome – DHCR7 sequencing	GENE	<b>A</b> 9	6 weeks	128
Smooth Muscle Antibodies	ASM0	В	2 days	87
Sodium	NA	В	1 day	37
Somatomedin (IGF-1)	SOMA	(Frozen) <sup>4,7</sup>	1 day	60
Soybean Components	ZZ18	В	2 days	143
Spastic Paraplegia NGS Panel	GENE	<b>A A</b> <sup>9</sup>	6 weeks	128
Sperm Aneuploidy	SPPL	Semen 1	4 weeks	68
Sperm Antibodies (Serum)	ASAB	В	5 days	68, 87
Sperm Antibodies/MAR Test (Semen)†	ASPA	Semen	1 day	68
Sperm Comet®	CMET	Semen 1	1-2 weeks	68
Sperm Comet® Exact Focus	CMT2	Semen 1	1-2 weeks	68
Sperm Comet® Extend	CMT3	Semen 1	1-2 weeks	68
Sperm Comet® Extend Focus	CMT4	Semen <sup>1</sup>	1-2 weeks	68

TEST	CODE	SAMPLE REQS	TAT	PAGE
Sperm Count (Post-Vasectomy)	PVAS	Semen 1	2 days	68
Sperm DNA Fragmentation (SCSA type test)	SEXT	Semen 1	1-2 weeks	68
Sperm Morphology (Kruger strict criteria)	MRPH	Semen 1	2 days	68
Spinal Bulbar Muscular Atrophy (Kennedy	GENE	<b>A</b> 9	5 weeks	128
Disease) – AR repeat analysis				
Spinal Muscular Atrophy – SMN1 deletions/duplications	SMA	<b>A</b> 9	10 days	128
Spinocerebellar Ataxia – multiplex SCA1+2+3+6+7+8+10+12 +17	GENE	<b>A</b> 9	5 weeks	128
common repeat expansions				
Spinocerebellar Ataxia NGS Panel	GENE	<b>AA</b> <sup>9</sup>	6 weeks	128
Sports/Performance Profile	SPOR		5 days	152-153
Sputum for Routine Culture	SPU1	SC	2-4 days	48
Sputum for TB Culture (AFB)	SPU2	SC	up to 8 weeks	48
Squamous Cell Carcinoma	SCC	<u>B</u>	4 days	106
STD1 M/F STD Quad (Urine and Serology)	STD1	₿ FCRU	2 days	74, 76
STD2 M/F STI Profile Plus (Urine and Serology)	STD2	B FCRU (If culture	4 days	74, 76
orbz min orritomor nas (ormo ana ostology)	0102	swabs are needed please	i dayo	71,70
		request separately)		
STD3 Female STD Quad (PCR Swab and Serology	) STD3	<b>₿</b> PCR	2 days	74, 76
STD4 Female STI Profile Plus	STD4	BPCR (If culture swabs	4 days	74, 76
(PCR Swab and Serology)		are needed please		
		request separately)		
STD5 Serology only	STD5	В	1 day	74, 76
STD6 Serology only without HIV	STD6	В	1 day	74, 76
STD8 Vaginitis/BV Profile using	STD8	PCR and STM	3 days	74, 76
Culture & PCR Swab				
STD9 Symptomatic lesion sample	STD9	PCR Swab	7 days	74, 77
using PCR Swab from lesion	SCA	<u> </u>	O days	07
Steroid Cell Antibody STI Profile by PCR (7 tests from 1 sample)	DL12	Aptima urine or multisite swab	2 days	74, 161
(Self-collect)	DLIZ	Apunia unile oi mulusite swab	2 uays	74, 101
STI Profile: MSM1 (Blood + Urine/	MSM1	(TDL Tiny) / Aptima Urine	2 days	74, 161
Throat/Rectal Swabs) (Self-collect)		/ Aptima multisite swab x 2		
STI Profile: MSM1 (Venous)	MSM1	3 / FCRU / PCR Swab Throat / PCR Swab Rectal	2 days	74, 78
STI Profile: MSM2 (Blood + Urine/	MSM2	(TDL Tiny) / Aptima urine	3 days	74, 161
Throat/Rectal Swabs) (Self-collect)		/ Aptima multisite swab x 2		
STI Profile: MSM2 (Venous)	MSM2	3 / FCRU / PCR Swab Throat / PCR Swab Rectal	3 days	74, 78
Stockholm3 NEW	STK3	AA	2 weeks	28, 106
Stockholm3 Reflex NEW	STKR	AAB	2 weeks	28, 106
Stool for OCP and Culture by PCR	PENT	RF <sup>††</sup>	2-3 days	48
Stool for OVA Cysts & Parasites by Microscopy	MOCP	RF	2 days	49
Cost. of other open an arabitoo by mioroscopy		•••	- wuju	

TEST	CODE	SAMPLE REQS	TAT	PAGE
Stool Reducing Substances	STRS	RF <sup>7</sup>	2-3 weeks	49
Streptomycin Levels	STRM	•	5 days	139
Striated/Skeletal Muscle Antibody	STRA	B	3 days	87
Strongyloides Antibodies	STGA	B	10 days	87
Sulpiride	SULP	<b>B</b> 4	4 days	139
Superoxide Dismutase	SODI	<b>A</b> / <b>()</b>	5 days	37
Suppression with steroid, IVIg and intralipin, NK (CD69) cell assay, TH1/TH2 cytokines	NCIT	0000*	Send Mon-Thurs only	62
Swab (Cervical)	CERS	STM / CS	2-4 days	49
Swab (Ear)	EARS	STM	2-4 days (Culture) 8-9	49
			days (Fungal) – same swab	
Swab (Eye)	EYES	STM	2-4 days	49
Swab (Nasal)	NASS	STM	2-4 days	49
Swab (Oral)	ORSW	STM/CS	2-4 days	49
Swab (Penile)	PENS	STM/CS	2-4 days	49
Swab (Rectal)	RECG	STM/CS	2-4 days	49
Swab (Skin)	SKIS	STM	2-4 days	49
Swab (Throat)	THRS	STM	2-4 days	49
Swab (Urethral)	URES	STM/CS	2-4 days	49
Swab (Vaginal)	VAGS	STM/CS	2-4 days	49
Swab (Vulval)	VULV	STM/CS	2-4 days	49
Swab (Wound)	WOUS	STM	2-4 days	49
Synacthen Stimulation Test	SYNA	By appointment only	1 day	137
Synovial Fluid	FLU2	SC <sup>†††</sup>	14 days	49
(for microscopy, crystals and culture)				
Syphilis by PCR (chancre)	SYPS	PCR	5 days	74
Syphilis IgG/IgM (Self-collect)	TSYP	3 (TDL Tiny)	1 day	74, 87, 161
Syphilis IgG/IgM (Venous)	SERJ	B	1 day	75, 87
T Regulatory Cells	25RF	0	3 days	61
T3	T3	B	1 day	60
T3 (Reverse)	RT3	B 7,37	15 days	60
Tacrolimus/Prograf (FK506)	FK5	<b>A</b> 4	1-2 days	139
Taipan Snake Venom Time	TTVT	<b>()</b> () 9,18	2-3 weeks	43
Tay Sachs Screen	GENE	<b>A</b> 9	4 weeks	128
TB (Pleural Fluid)	TBCU	SC	up to 8 weeks	49
TB Culture	SPU2	SC	up to 8 weeks	49
TB Culture (Urine)	TBUR	3 x EMU	up to 8 weeks	49
TB PCR (PCR detection of Mycobacterium	TBPC	All samples except blood	1 day	49
tuberculosis complex and mutations for Rifampicin resistance)		cultures and urine, as clinically requested.		
TB Quantiferon®-TB Gold*	TBQ4	Special tubes or (1) 1	3 days	87

TEST	CODE	SAMPLE REQS	TAT	PAGE
TB Slopes – Confirmation and Sensitivity	TBSL	TB slope (LJ medium-green) <sup>6</sup>	up to 8 weeks	49
Tegretol (Carbamazepine)	CARB	В	1 day	139
Teicoplanin Assay	TEIC	B	5 days	137
Temazepam	TEMA	<b>B</b> 4	4 days	139
Testicular Tumour Profile (LDH, AFP, HCQG)	TTP	В	1 day	106, 108
Testosterone (Self-collect)	TEST	(TDL Tiny) or (TDL Tiny)	1 day	60, 161
Testosterone (Total), LC MS Mass Spec NEW	MSTT	B	5-7 days	60
Testosterone (Venous)	TEST	<b>B</b>	1 day	60
Testosterone (Free) (Self-collect)	FTES	(TDL Tiny) or (TDL Tiny)	3 days	60, 161
Testosterone (Free) (Venous)	FTES	B	3 days	60
Tetanus Antibody	TETA	В	5 days	87, 95
TH1/TH2 Cytokine Profile	1TH2	000*	Send Mon-Thurs only	62
TH1/TH2 Cytokine Ratio	6RF	000⁵	1 week	61
TH1/TH2 Intracellular Cytokine Ratios with IVIG	21RF	<b>000</b> <sup>5</sup>	1 week	61
TH1/TH2 Intracellular Cytokine	20RF	<b>000</b> <sup>5</sup>	1 week	61
Ratios with IVIG, Prednisolone				
TH1/TH2 Intracellular Cytokine	22RF	<b>⊕⊕</b> ∮	1 week	61
Ratios with Prednisolone				
Thalassaemia Screen	HBEL	<u>A</u>	4 days	43
Thallium (Blood)	THAL	<b>A</b> / <b>(1)</b>	1 week	166
Thallium (Urine)	URTH	RU	1 week	166
Theophylline	THE0	В	1 day	139
Thiopurine Methyl Transferase	TPMT	<b>A</b> 5	5 days	37
Thrombin Time	THR0	<b>C</b> 18	1 day	42
Thrombotic Risk Profile	PR0P	<b>A B O O O</b> <sup>18</sup>	5 days	43, 45,
	TOAR		4.1	128, 130
Thyroglobulin Abs	TGAB	<u>B</u>	1 day	60
Thyroglobulin Assay	TGA	B (TDL T' )	1 day	60
Thyroid Abs (Thyroglobulin + Thyroid	THAB	(TDL Tiny)	2 days	60, 87,
Peroxidase Abs) (Self-collect) Thyroid Abs (Thyroglobulin + Thyroid	THAB	B	1 day	60, 87
Peroxidase Abs) (Venous)	ITIAD	•	i udy	00, 07
Thyroid Cancer NGS Panel	GENE	<b>A A</b> 9,11	4 weeks	128
Thyroid Peroxidase Antibodies/Anti TPO	TPEX	B	1 day	60, 87
Thyroid Profile 1 (FT4/TSH) (Self-collect)	TF	B (TDL Tiny)	1 day	60, 161
Thyroid Profile 1 (FT4/TSH) (Venous)	TF	B	1 day	60, 64
Thyroid Profile 2 (Venous)	TF2	<u>B</u>	2 days	60, 64
Thyroid Profile 3 (FT3/FT4/TSH) (Self-collect)	TF3	B (TDL Tiny)	1 day	60, 161
Thyroid Profile 3 (FT3/FT4/TSH) (Venous)	TF3	B	1 day	60, 64
Thyroxine (T4)	T4	<u>B</u>	1 day	60
Thyroxine (14) Thyroxine Binding Globulin	TBG	(Frozen)	10 days	60
Timothy Grass Components	ZZ19	B	2 days	143
Tissue for culture	TISS	Tissue sample	up to 14 days	49
1100UG IVI GUILUIG	1100	ı ıoauc adınıpıc	up to 14 days	49

TEST	CODE	SAMPLE REQS	TAT	PAGE
Tissue Polypeptide Antigen	TPA	<b>B</b>	1 week	37
Tissue Transglutaminase IgA	TAA	(TDL Tiny)	2 days	87, 161
(Coeliac) (Self-collect)				
Tissue Transglutaminase IgA	TAA	B	2 days	87
(Coeliac) (Venous)**				
Tissue Transglutaminase IgG	TAAG	B	5 days	87
Tobramycin Assay (Provide Clinical Details)	TOBR	0	3 days	137
Toluene (Blood)	T0L	<b>J</b> (Contact Referrals)	10 days	166
Toluene (Urine)	UTOL	RU	10 days	166
Topiramate (Topamax)	TOPI	B 4	4 days	139
Torch Screen	TORC	В	2 days	103, 105
Total Acid Phosphatase	APT	B	5 days	37
Total Bile Acid/Bile Salts	BILS	<b>B</b>	1 week	37
Total IgE	IGE	<b>B</b>	1 day	37, 141
Total Immune Function Evaluation	TIE	A + B 5,10	7 days	87
Total Immunoglobulin E	IGE	B	1 day	87
Total Testosterone, LC MS Mass Spec NEW	MSTT	B	5-7 days	60
Toxocara Antibodies (IgG)	TFAT	<b>B</b> 9	5 days	87
Toxoplasma Antibodies (IgG, IgM)	TFAM	B 9	1 day	87, 92
Toxoplasma Antibody Full Evaluation	TDYE	B 9	10 days	87
(IgM, Dye Test, IgG Avidity)				
Toxoplasma by PCR	TXAG	A	5 days	87
TPHA	TPPA	B	2 days	75, 87
Trace Metal (Blood) Profile	TRAC	<b>4 8 9 6</b>	7-10 days	165-166
Transferrin	TRAN	B	1 day	37
Transferrin Electrophoresis	TREL	B	2 weeks	37
Treacher Collins Syndrome and	GENE	<b>A A</b> <sup>9</sup>	6 weeks	129
Related Disorders NGS Panel				
Trichinella Serology	TRIC	<b>B</b>	5 days	87
Trichloracetic Acid (Urine)	UTCA	RU	5 days	166
Trichomonas vaginalis (Thin Prep)	TVPC	TPV	2 days	170
Trichomonas vaginalis (PCR)	TVPC	FCRU / PCR / TPV	2 days	75
Trichomonas vaginalis (TV) – Urine or Vaginal (Self-collect)	TVPC	Aptima urine or multisite swab	2 days	75, 162
Triglycerides	TRI	<u> </u>	1 day	37
Trimethylaminuria (Fish Odour Syndrome)	FOS	J	6 weeks	37
Trimipramine	TRIM	<u> </u>	5 days	139
Triple Swab Female STI Profile	3SWA	PCR swab x 3 (label by site)	2 days	75, 78
(Vaginal/Throat/Rectal Swabs) (PCR)	JOWA	i dii swan x 3 (lanci ny Sile)	L uuyo	13, 10
Triple Swab Female STI Profile	3SWA	Aptima multisite swab	2 days	75, 162
(Vaginal/Throat/Rectal Swabs) (Self-collect)	OUIA	x 3 (label by site)	- aayo	70, 102
Tropical Screen (from 6 weeks post-travel)	TROP	B B 9,14	10 days	92-93
Tropomyosins	ZZ31	<u>в</u>	2 days	143
	2201	<u> </u>	- auyo	140

TEST	CODE	SAMPLE REQS	TAT	PAGE
Troponin I (High sensitive)	TROC	B	1 day	37
Troponin T (High sensitive)	TROT	B	1 day	37
Trypanosome (Chagas) Antibodies	CHGA	B 9,14	10 days	88
Tryptase	STRY	B	2 days	37, 141
TSH (Self-collect)	TSH	(TDL Tiny)	1 day	60, 162
TSH (Venous)	TSH	B	1 day	60
TSH-Receptor Antibodies	TSI	B	4 days	60, 88
Tuberous Sclerosis (TSC1 + TSC2)	GENE	<b>A A</b> <sup>9</sup>	7 weeks	129
Tularaemia Antibodies	TULA	B 14	5 days	88
Tumour Necrosis Factor – Alpha	TNF	<sup>□</sup> (Frozen) <sup>4</sup>	2 weeks	37
Urate (Uric acid)	UA	B	1 day	37
Urea (Self-collect)	UREA	(TDL Tiny)	1 day	37, 162
Urea (Urine)	UURE	CU	1 day	38
Urea (Venous)	UREA	B	1 day	37
Urea and Electrolytes	U/E	<b>B</b>	1 day	38, 40
Urea/Creatinine/eGFR (Self-collect)	TCU	(TDL Tiny)	1 day	38, 162
Ureaplasma urealyticum (Thin Prep)	UGEN	TPV	2 days	170
Ureaplasma urealyticum by PCR	UGEN	FCRU / PCR / TPV	2 days	75
Uric Acid (Serum)	UA	B	1 day	38
Uric Acid (Urine)	UURI	CU	1 day	38
Urinary Bladder Cancer Antigen	UBC	RU (Freeze within 48 hours)**	5 days	38, 106
Urinary Methyl Histamine	UHIT	RU (Frozen)	2 weeks	88
Urinary Tract/Renal Cancer NGS Panel	GENE	<b>A A</b> 9,11	4 weeks	129
Urine (Microscopy Only)	UMIC	RU	1 day	49
Urine Chemistry and Microscopy (Self-collect)	UMIC	Urine (Universal). Mid stream.	1-2 days	49, 162
Urine Chemistry, Microscopy	UCEM	Urine (Universal &	1-2 days	50, 162
and Culture (Self-collect)		Boric). Mid stream.		
Urine Cytology (Urine cytology containers	URCY	Urine (30mls) <sup>21</sup>	2 days	174
available from TDL Supplies)				
Urine EtG (Ethyl glucuronide)	ETG	RU	1 week	163
Urine for Extended Culture	UCXD	MSU ††††	up to 7 days	50
Urine for Microscopy and Culture	UCEM	MSU ††††	1-2 days	50
Urine Microalbumin/Creatinine Ratio	UMA	RU	1 day	38
Urine Organic Acids	UORG	RU (Frozen)	3 weeks	38
Urine Steroid Screen (Steroid Hormones)	USTE	CU <sup>9</sup>	2 weeks	38
Urine Sugar Chromatography	UCR0	RU (Frozen)	3 weeks	38
Urticaria Test (Histamine Releasing)	CURT	B	3 weeks	88
Usher Syndrome NGS Panel	GENE	<b>AA</b> <sup>9</sup>	6 weeks	129
Vaginitis/BV Profile (Culture & PCR)	STD8	PCR and STM	3 days	75
Vaginitis/BV Profile using Culture	STD8	Aptima multisite swab and	3-5 days	75, 162
& PCR Swab (Self-collect)		Blue gel Amies swab		
Valium (Diazepam)	DIAZ	<u> </u>	7 days	139
Valproic Acid (Epilim)	VALP	В	1 day	139

TEST	CODE	SAMPLE REQS	TAT	PAGE
Vancomycin Hydrochloride	VANC	B	1 day	137
Varicella zoster – DNA	VZPC	A	5 days	103
Varicella zoster Antibodies (IgG)	VZOS	В	1 day	95, 103
Varicella zoster Antibodies (IgM)	VZOM	B	1 day	95, 103
Vascular Endothelial Growth Factor	VEGF	В	14 days	88
Venom Components	ZZ33	В	2 days	143
Very Long Chain Fatty Acids	VLCF	(A) or (1) (Frozen) 9	4-6 weeks	38
Vigabatrin (Sabril)	VIGA	<b>A</b>	10 days	139
Viral Antibody Screen	VIRA	ВВ	2 days	103, 105
Viral Eye by PCR	VPE	PCR	3 days	103, 105
Viral Respiratory RNA Screen by PCR	VPR	PCR or as specified on the form	2 days	103, 105
Viral Respiratory RNA Screen	VPR	Aptima multisite swab	2 days	103, 162
by PCR (Self-collect)		of nose/throat		
Viral Skin/Mucosa by PCR	VPSK	PCR	5 days	104-105
Viscosity (Plasma)	VISC	<b>A</b> *4	3 days	43
Vitamin A (Retinol)	VITA	В	5 days	151
Vitamin B (Functional)	FUNC	A A or (1)	5 days	151
Vitamin B Profile	VBP	AAB	5 days	151, 153
Vitamin B1 (Thiamine)	VIT1	A	5 days	151
Vitamin B2 (Riboflavin)	VIB2	A	5 days	151
Vitamin B3 (Nicotinamide)	VIB3	В	5 days	151
Vitamin B5 (Pantothenic Acid)	VB5S	В	5 days	151
Vitamin B6 (Pyridoxine)	VITB	A	5 days	151
Vitamin B7 (Biotin)	BIOS	<b>B</b> <sup>7</sup>	5 days	151
Vitamin B9 (Folic acid) – Red cell	RBCF	A	2 days	151
Vitamin B9 (Folic acid) – Serum	FOLA	<b>B</b>	1 day	151
Vitamin B12 (Active) (Self-collect)	B12	(TDL Tiny)	1 day	38, 151,
				162
Vitamin B12 (Active) (Venous)	B12	B	1 day	38, 151
Vitamin B12 (Active)/Red Cell Folate	B12F	<b>Q B</b>	2 days	38, 151
Vitamin B12 (Total)	TB12	8	1 day	38
Vitamin C (Active)	VITC	(spun and frozen within 3 hours)*	5 days	151
Vitamin D (1, 25 Dihydroxy)	D3	B*	5-8 days	151
Vitamin D (25-OH) (Self-collect)	VITD	(TDL Tiny)	1 day	38, 151, 162
Vitamin D (25-OH) (Venous)	VITD	В	1 day	38, 151
Vitamin E (Alpha Tocopherol)	VITE	<b>B</b>	5 days	151
Vitamin K (Nutritional)	VKN	Serum (SST or 🕒) *	5 days	151
Vitamin Profile 1	VITS	<b>A B B</b> <sup>7</sup>	5 days	151, 153
Vitamin Profile 2	VIT2	<b>A A A</b> 38 <b>B B</b> 7,13	5 days	151, 153
VLDL Cholesterol	VLDL	B 13	1 week	38
VMA	UVMA	PU <sup>1</sup>	5 days	38

Voltage Gated Calcium Channel Antibodies  VPCA  VPCA  VPCA  S  Weeks  VPCA  VPCA  S  Weeks  VPCA  VPCA  S  Weeks  VPCA  Weeks  VPCA  S  Weeks  Weeks  Weeks  VPCA  S  Weeks  VPCA  S  Weeks  Weeks  VPCA  S  Weeks  Weeks  VPCA  S  Weeks  Weeks  VPCA  S  Weeks  Weeks  VPCA  S  Weeks  Weeks  Weeks  VPCA  S  Weeks  Weeks  Weeks  VPCA  S  Weeks  W	88 88 129 43, 45
Von Hippel-Lindau Syndrome – GENE (3) 9 6 weeks VHL sequencing + deletions/duplications	129
VHL sequencing + deletions/duplications	
	43, 45
Van Willehvand Drefile FUNE 0 0 4912 5 dags	43, 45
Von Willebrands Multimers VWM	43
Wall Pellitory Components ZZ20 3 2 days	143
Walnut Components ZZ34 3 2 days	143
West Nile Virus Abs WNV 3 2 weeks	104
Wheat Components ZZ21 3 2 days	143
Whole Genome Sequencing (solo/duo/trio) GENE (4) 9,11 5-8 weeks	129
Whooping Cough (Pertussis) Antibodies PERS (3) 5 days	88
Whooping Cough (Pertussis) by PCR PERP Pernasal or dry swab 2-3 days	88
Xanthine – Blood XANB (2) 2 weeks	166
Xylene – Urine UXYL RU 30 2 weeks	166
Yellow Fever Antibodies YELL (3 9.14 10 days	88
Yersinia Antibodies YERS (3) 4 days	88
Zellweger Syndrome NGS Panel GENE (3 (3) 9 6 weeks	129
Zika Abs IgM and IgG – ZKAB (3) Up to 14 days	88, 92,
Antibody detection from 15 days	104
Zika RNA by PCR in Semen ZIKS Semen Up to 14 days	88, 92,
	104
Zika RT PCR – Window of detection from ZIKA (3) Up to 14 days	88, 92
1-7 days from onset of symptoms	
Zika RT PCR – Window of detection from ZIKU RU Up to 14 days	88, 92
1-14 days from onset of symptoms	
Zinc (Serum) ZINC (§ 2 days	152, 165
Zinc (Urine) URZN CU 5 days	152, 165
Zinc (Whole Blood) RBCZ (A) or (1) 5 days	152
Ziwig Endotest® ENDT Endotest saliva collection kit 25 days	60, 129

# **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.

# **TDL Referral laboratories**

A3P Biomedical AB [980]

Addenbrooke's Hospital – BGU and Immunology [899]

Alder Hey Children's NHS Foundation Trust

- Biochemistry Department [880]

Analytical Services International Ltd, St George's University of London – Forensic Toxicology Service [994]

Animal and Plant Health Agency – Veterinary labs [911]

Bio Predictive [Original report]

Bioscientia (Germany) [868]

Birmingham Children's Hospital NHS Foundation Trust – Clinical Chemistry [970]

Birmingham University Hospital NHS Foundation trust [895]

Brucella Reference Unit – Liverpool Clinical Laboratories, Royal Liverpool and Broadgreen Hospital [947]

Cambridge Clinical Laboratory [867]

Cambridge Life Sciences [997]

Cambridge Nutritional Science Ltd [Original report]

Cardiff and Vale University Health Board – Porphyria Service Cardiff [834]

Cardiff and Vale University Health Board — The Analytical Toxicology Department [998]

Douglass Hanly Moir Pathology (Australia) [Original report]

Epsom and St Helier University Hospital NHS Trust

– Biochemistry Department [968]

Epsom and St Helier University Hospital NHS Trust – Immunology Department [968]

Epsom and St Helier University Hospital NHS Trust

– Microbiology Department [951]

Eurofins - Biomnis (France) [950]

Great Ormond Street Hospital –
Department of Chemical Pathology [964]

Great Ormond Street Hospital – Enzyme Unit, Chemical Pathology [964]

Great Ormond Street Hospital – Immunology Department [924]

Great Ormond Street Hospital – Neurometabolic Unit [964]

Guildford RSCH Trace Element Laboratory, SAS Trace Element Centre [955]

HCA Healthcare UK – HCA Laboratories [982]

HFL Sport Science (LGC Group) [861]

Igenomix UK [Original Report]

Imperial College Healthcare NHS Trust – Charing Cross Hospital, Chemical Pathology Department [912]

Imperial College Healthcare NHS Trust – Charing Cross Hospital, Infection and Immunity Department [962]

Imperial College Healthcare NHS Trust — Charing Cross Hospital, Medical Oncology [912]

Imperial College Healthcare NHS Trust – Hammersmith Hospital, Molecular Endocrinology [931]

Imperial College Healthcare NHS Trust, St Mary's Hospital – Virology Department [912]

Institute of Aquaculture – University of Stirling [1000]

Institute of Neurology – Neurogenetics Unit [975]

King's College Hospital – HMDC Laboratory for Molecular Haemato-Oncology [943]

Labor Augsburg MVZ GmbH (Germany) [900]

Latis Scientific [927]

LogixX Pharma Ltd [Original report]

London School of Hygiene & Tropical Medicine

– Diagnostic Parasitology Lab [933]

Matrix Diagnostics [896]

Mayo Clinic Laboratories (Netherlands [894]

# **TDL Referral laboratories**

Meningococcal reference unit (Men RU) Manchester – Manchester Royal Infirmary [949]

Micropathology Ltd [920]

National Blood Service - Colindale,

Red Cell Immuno Haematology Department [910]

NHS Blood and Transplant – Birmingham [856]

NHS Blood and Transplant – H & I Laboratory [855]

NHS Blood and Transplant – Tooting [854]

Norfolk and Norwich University Hospital NHS Foundation Trust – SAS Metabolic Bone Laboratory [993]

Oxford University Hospital NHS Foundation Trust – Churchill Hospital [983]

Queens University Hospital, Belfast – Institute of Clinical Science [853]

Reflab (Denmark) [988]

Reproductive Immunology Centre [839]

Rosalind Franklin University (USA) - [Original report]

Royal Berkshire Hospital NHS Foundation Trust

- Clinical Biochemistry [849]

Royal Devon and Exeter NHS Foundation Trust [838]

Royal Surrey County Hospital – SAS Peptide Hormone Section [959]

Sandwell and West Birmingham NHS Trust

City Hospital Birmingham,

Clinical Biochemistry Department [970]

Sheffield Children's NHS Trust – Clinical Chemistry [847]

Immunology Department [966]

Sheffield Teaching Hospital NSH Foundation Trust – Protein Reference Laboratory Unit and

Southmead Hospital – Antimicrobial Reference Laboratory. Bristol [915]

St George's University Hospital NHS Foundation Trust

- Cell Marker Department [846]

Synnovis – Guy's Hospital,

Biochemistry Genetics Laboratory [930]

Synnovis – King's College Hospital, Clinical Biochemistry [914] Synnovis – St Thomas' Hospital Haemophilia Centre [956]

Synnovis – St Thomas' Hospital Immunohistology [961]

Synnovis – St Thomas' Hospital Purine Research Laboratory [925]

The Epilepsy Society (Chalfont Centre) [837]

The Leeds Teaching Hospital NHS Trust – Endocrinology Laboratory (including SAS Steroid Centre), Department of Specialist Laboratory Medicine, St James University Hospital) [992]

The Leeds Teaching Hospitals NHS Trust

– Mycology Reference Centre [973]

The Newcastle upon Tyne Hospitals – Royal Victoria Infirmary [878]

The Royal Marsden Hospital –
Department of Haematology / Oncology [989]

The Royal Marsden Hospital – Department of Pathology [990]

Toxoplasma Reference Unit, Public Health Wales Microbiology ABM, Singleton Hospital – Swansea [969]

Trace Laboratories Ltd [955]

UCL Great Ormond Street Institute of Child Health [935]

UCL Queen Square Institute of Neurology – Department of Neuroimmunology [975]

UKHSA – Bacteriology Reference Department (BRD), Colindale [910]

UKHSA – Virus Reference Department (VRD) – Colindale [910]

UKHSA Mycology Reference Laboratory – UKHSA South West Laboratory, Southmead Hospital, Bristol [903]

UKHSA National Mycobacterium Reference Service National Infection Service. Colindale [974]

UKHSA Rare and imported pathogens laboratory

– Porton Down [981]

University Hospital Birmingham NHS Foundation Trust – Heartlands Hospital [843]

University Hospital of Wales – Cardiff Medical Immunology Department [842]

# **TDL Referral laboratories**

Wythenshawe Hospital, Manchester University NHS Foundation Trust, Manchester [835]

Ziwig [833]

# **Group laboratories**

Royal Free London NHS Foundation Trust – Haemostasis [984]

University College London Hospitals NHS Foundation Trust (UCLH) — Cytology [Original report]

University College London Hospitals NHS Foundation Trust (UCLH) – Hospital for Tropical disease [933]

University College London Hospitals NHS Foundation Trust (UCLH) – Molecular Virology [999]

University College London Hospitals NHS Foundation Trust (UCLH) – Special Chemistry [953]

# **TDL Genetics Referral laboratories**

All Wales Medical Genetics Service

Anthony Nolan, Histocompatability and Immunogenetics

Asper Biotech

Bioscientia GmBH

Bristol Genetics Laboratory (North Bristol NHS Trust)

CentoGene

DiaGenom GmbH

Douglass Hanly Moir Pathology

East Scotland Regional Genetics Service (NHS Tayside)

Exeter Clinical Laboratory – Department of Molecular Genetics

**Fulgent Diagnostics** 

Institute of Neurology, Queen's Square

International Blood Group Reference Laboratory

London South East Genetics Service

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

Progenika Biopharma Grifols

Protein Reference Unit & Immunology Department
- Sheffield Protein Unit

Purine Research Laboratory – St Thomas' Hospital

Royal Marsden – Haemato-Oncology Unit

Sheffield Diagnostic Genetics Service

SIHMDS – Cytogenetics Laboratory, Great Ormond Street Hospital

South East Scotland Genetics Service (NHS Lothian)

South West Thames Regional Genetics Service

SYNLAB Budapest Diag Center

The Leeds Genetics Laboratory Viapath Analytics LLP

Wessex Region Genetics Service

West Midlands Regional Genetics Laboratory

West of Scotland Genetic Service (NHS Greater Glasgow and Clyde)

The definitions which apply to these Terms and Conditions are set out in clause 19

#### 1 THE SERVICES

- 1.1 These Terms and Conditions and any applicable Service-Specific Terms will apply to any services or consumables that The Doctors Laboratory Limited or TDL Genetics Limited 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 and any applicable Service-Specific Terms 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 Pathology Request, a request for any other services described in the Laboratory Guide or in any other proposal provided by TDL, or an order for any Consumables described in the Laboratory Guide (in each case an 'Order'), the Client offers to purchase those Tests, other services or Consumables on these Terms and Conditions and any applicable Service-Specific Terms from TDL. TDL may accept or reject any Order.
- 1.3 A contract between TDL and the Client for the provision of Services and / or Consumables, incorporating these Terms and Conditions, and any applicable Service-Specific Terms, and the Order (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. In the event of a conflict between the Order and these Terms and Conditions, the Terms and Conditions will take priority.
- 1.4 By Ordering a Service referred to in any Service-Specific Terms, the relevant Service-Specific Terms will apply to that Service in addition to these Terms and Conditions. In the event of a conflict between these Terms and Conditions or the Order and the Service-Specific Terms, the Service-Specific Terms will take priority.
- 1.5 TDL will provide the Services under the Agreement:
- 1.5.1 in accordance with Good Industry Practice;
- 1.5.2 in accordance with the UKAS medical laboratory accreditation standard (ISO 15189); and
- 1.5.3 using suitably skilled and experienced staff.
- 1.6 TDL will use reasonable efforts to achieve the Test turnaround times quoted in the Laboratory Guide, but does not warrant that it will achieve those times in the case of any particular Sample.
- 1.7 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.
- 1.8 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.
- 1.9 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.
- 1.10 In providing the Services, TDL shall comply with all Applicable Law relating to anti-bribery and anti-corruption, including the Bribery Act 2010. TDL shall not, and shall ensure that its staff do not, engage in any activity which would constitute an offence under the Bribery Act 2010.
- 1.11 TDL is committed to trading ethically, with zero tolerance for modern slavery (including forced labour or human trafficking of any kind), human rights violations, and child labour. In performing its obligations under the Agreement, TDL will comply with all Applicable Law and applicable internal policies relating to anti-slavery and human trafficking.
- 1.12 TDL's laboratories are operated by members of the TDL Group. TDL uses those laboratories to undertake the Tests, except where TDL refers the Tests to suitably accredited laboratories operated outside the TDL Group. The UKAS accreditation numbers for the TDL Group laboratories in the UK are as follows: 8059 (HSL Analytics LLP) Genetics and Molecular Sciences, 8169 (HSL Analytics LLP) Blood Sciences, 8860 (HSL Analytics LLP) Infection Sciences, 8812 (The Doctors Laboratory Limited) Haematology, Blood Transfusion, Biochemistry, Microbiology, Molecular Biology, 10199 (The Doctors Laboratory Limited) Andrology, 8511 (HSL Analytics LLP) Cytology, 9706 (The Doctors Laboratory Limited) Urine Cytology.

#### 2 SUPPLY OF CONSUMABLES

- 2.1 TDL shall supply Consumables to the Client in accordance with the terms of the Agreement.
- 2.2 The Consumables shall: (i) be of satisfactory quality (within the meaning of the Sale of Goods Act 1979) and fit for any purpose held out by TDL; and (ii) comply with all Applicable Law.

- 2.3 TDL shall not be liable for Consumables' failure to comply with clause 2.2 if: (i) the Client makes any further use of those Consumables after notifying TDL of such failure; (ii) the defect arises because the Client failed to follow TDL's instructions for the storage, use or maintenance of the Consumables or (if there are none) good practice regarding the same; (iii) the Client alters or repairs those Consumables without TDL's prior written consent; (iv) the defect arises as a result of fair wear and tear, deliberate damage, negligence, or abnormal storage or working conditions; or (v) the Consumables differ from their description as a result of changes made to ensure they comply with Applicable Law.
- 2.4 In the event the Consumables do not comply with clause 2.2, TDL shall provide replacement Consumables without undue delay. This shall be the Client's only remedy for such non-compliance. The terms of this clause 2 shall apply to any such replacement Consumables provided by TDL.
- 2.5 TDL shall ensure that the Consumables are properly packed and secured in a manner to enable them to reach their destination in good condition, and in a manner which complies with Applicable Law.
- 2.6 If the Client or the Client's carrier will collect the Consumables from TDL's premises, delivery shall be completed when TDL places the Consumables at the Client's disposal at TDL's premises. In all other cases, delivery shall be completed on the loading of the Consumables at the premises where they are loaded onto transport for carriage.
- 2.7 TDL may deliver Consumables by instalments, which may be invoiced and paid for separately. Time for delivery of Consumables is not of the essence of the Agreement and delays in the delivery of Consumables shall not entitle the Client to refuse to take delivery. TDL shall have no liability for any failure or delay in delivering Consumables to the extent that any failure or delay is caused by the Client's failure to comply with its obligations under the Agreement.
- 2.8 Title and risk in the Consumables shall pass to the Client on delivery, except that any biofreeze bottles provided by TDL shall remain the property of TDL at all times, regardless of any use by the Client of the biofreeze bottles.
- 2.9 The Client must not resell the Consumables or provide them to any third party without TDL's prior written consent.
- 2.10 The Client shall ensure that: (i) any Consumables provided by TDL are only used by healthcare professionals who are appropriately qualified and trained in the proper use of such Consumables; and (ii) the healthcare professionals use the Consumables in accordance with any instructions relating to the use of the Consumables provided by TDL and in any

event with the degree of skill and care reasonably to be expected of a healthcare professional experienced in the use of such Consumables.

#### 3 PRICE AND PAYMENT TERMS

- 3.1 The price payable by the Client for the Services and / or the Consumables 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 and / or Consumables, the price will be that indicated in the Laboratory Guide.
- 3.2 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 and/or Consumables the Client will pay it at the applicable rate.
- 3.3 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 payments 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.
- 3.4 If the Client disputes any invoice: (i) the Client shall notify TDL in writing as soon as practicably possible and in any event not later than 90 days after the date of the invoice, specifying the reasons for disputing the invoice; (ii) the Client shall pay to TDL all amounts not disputed by the Client as set out in clause 3.3 above; and (iii) the parties shall attempt to resolve the dispute promptly and in accordance with clause 18.1 below.
- 3.5 If the Client does not dispute an invoice in accordance with clause 3.4 above then the amount stated on the invoice shall be deemed payable by the Client and the Client shall not be entitled to dispute the amount invoiced.
- 3.6 Without affecting any of its other rights, TDL may suspend or cease provision of the Services and / or Consumables if the Client fails to pay an invoice due to TDL, or if the total of the sums payable by the Client to TDL under any agreements between the Client and TDL meets or exceeds any credit limit that TDL communicates to the Client from time to time.

### 4 CONFIDENTIALITY

- 4.1 TDL agrees that it will hold and maintain the confidence of:
- 4.1.1 all information of a confidential nature which is received by TDL from the Client or its patients in connection with the Services; and

- 4.1.2 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 the 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.
- 4.2 The restrictions in clause 4.1 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.

# 5 CLIENT RESPONSIBILITIES

- 5.1 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 good clinical practice and, if applicable, TDL's written instructions.
- 5.2 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. Where TDL agrees to arrange transport of the Sample the Client will ensure that the Samples are ready for collection by TDL or its carrier at the agreed times.
- 5.3 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 performance of the Testing, and any other Services, and the use of the Protected Data as contemplated in the Agreement.
- 5.4 The Client will provide TDL with any information reasonably necessary for performing the Services and / or supplying Consumables, 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 and / or Consumables is accurate and complete.

#### 6 LIABILITY

- 6.1 Nothing in the Agreement will limit or exclude any liability that cannot be limited or excluded under Applicable Law, for example liability for death or personal injury caused by negligence.
- 6.2 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, the Consumables or under or in connection with any Agreement.
- 6.3 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.
- 6.4 Neither TDL nor the Client will have any liability for:
- 6.4.1 loss of profit or revenue;
- 6.4.2 loss of anticipated savings:
- 6.4.3 loss of reputation or goodwill; or
- 6.4.4 indirect, special or consequential loss.
- 6.5 TDL will have no liability for any delay or failure in performance of the Services or provision of the Consumables arising from the Client's delay or failure in performing its obligations under the Agreement.
- 6.6 All of the warranties which TDL gives in relation to the Services and / or the Consumables 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.
- 6.7 In this clause 6, references to TDL include the members of TDL's Group, and for the purpose of the limit in clause 6.3 the liabilities of TDL and the TDL Group Members will be counted in aggregate. The members of TDL

#### 7 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.

#### 8 DATA PROCESSOR AND DATA CONTROLLER

- 8.1 When TDL processes Protected Data on behalf of the Client in providing the Services the parties agree that the Client will be the controller and TDL will be the processor. The Annex to these Terms and Conditions sets out when TDL processes Protected Data on behalf of the Client. Clause 17 describes the circumstances where TDL will use Protected Data on its own behalf as controller.
- 8.2 When TDL processes Protected Data as processor, clauses 9 to 16 will apply in relation to the Protected Data. Where TDL processes Protected Data as controller, clause 17 will apply instead.
- 8.3 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.

#### 9 DATA PROCESSING INSTRUCTIONS

- 9.1 When TDL processes Protected Data as processor, TDL will comply with the obligations of processors under the Data Protection Laws.
- 9.2 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 these Terms & Conditions, and in the Annex (the 'Processina Instructions').
- 9.3 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).
- 9.4 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 9.4 do not limit the Client's obligations under clause 8.3.

# 10 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.

#### 11 USING STAFF AND OTHER PROCESSORS

- 11.1 TDL will not engage any 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.
- 11.2 TDL will ensure that each Sub-Processor is appointed under a written contract containing materially the same obligations as clauses 9 to 16 (inclusive).
- 11.3 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).

# 12 ASSISTANCE WITH THE CLIENT'S COMPLIANCE AND DATA SUBJECT RIGHTS

- 12.1 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.
- 12.2 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, (iii) prior consultation with the relevant regulator regarding high risk processing, and (iv) 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 12, 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.

#### 13 INTERNATIONAL DATA TRANSFERS

13.1 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 the 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, subject to clause 13.2

- 13.2 Where the Client requires TDL to transfer Protected Data for the purpose of providing the Services to a country outside the United Kingdom which is not subject to an adequacy regulation under the Data Protection Laws (a Third Country) then:
- 13.2.1 the Client will enter into (or where relevant use reasonable endeavours to procure that the applicable third party recipient of the Protected Data enters into) standard data protection clauses with TDL authorised under the Data Protection Laws for the international transfer of personal data that provide sufficient safeguards for the relevant transfer, on terms acceptable to TDL (acting reasonably); and
- 13.2.2 where the data protection clauses referred to in clause 13.2.1 are not entered into, the Client will procure that prior to the transfer the relevant data subjects provide valid consent to the transfer for the purposes of the Data Protection Laws, and the Client will provide evidence of such consents to TDL on request.

### 14 RECORDS. INFORMATION AND AUDIT

- 14.1 TDL will maintain, in accordance with the Data Protection Laws binding on TDL, written records of all categories of processing activities carried out on behalf of the Client.
- 14.2 TDL will, in accordance with the Data Protection Laws, make available to the Client such information as is reasonably necessary to demonstrate TDL's compliance with its obligations as a 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) to the extent reasonably necessary for that purpose, subject to the Client:
- 14.2.1 giving TDL reasonable prior notice and in any event not less than 30 days' notice of such information request, audit and/or inspection required by the Client;
- 14.2.2 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); and
- 14.2.3 ensuring that such audit or inspection is undertaken during normal business hours, with minimal disruption to TDL's business, any Sub-Processor's business and the business of other customers of TDL.

### 15 BREACH NOTIFICATION

TDL will, without undue delay, notify the Client of a personal data breach involving the Protected Data, and provide the Client with details of the personal data breach.

# 16 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, in which case TDL will inform the Client of any such requirement). Where TDL will process that Protected Data as controller under clause 17, TDL may retain the Protected Data.

# 17 PROTECTED DATA THAT TDL PROCESSES AS A CONTROLLER

- 17.1 TDL may process Protected Data as controller in the circumstances and for the purposes set out in TDL's Privacy Notice. In particular TDL may:
- 17.1.1 retain and submit the 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 a regulator for the purpose of complying with regulatory obligations; and
- 17.1.2 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.
- 17.3 When TDL processes Protected Data to provide Non-Invasive Prenatal Tests, TDL does so as a controller.
- 17.4 When TDL processes personal data on its own behalf as controller, it will do so in accordance with the obligations of data controllers under the Data Protection Laws and with the applicable terms of the Agreement.

#### 18 GENERAL

- 18.1 Disputes
- 18.1.1 If any dispute arises relating to the Agreement or any breach or alleged breach of the 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.
- 18.1.2 Except to the extent clearly prevented by the area of dispute, the parties will continue to perform their respective obligations in respect to any existing Agreements while such dispute is being resolved.
- 18.2 Variation

- 18.2.1 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.
- 18.2.2 Except as set out in clause 18.2.1, any amendments to the Agreement will not be effective unless in writing and signed by an authorised signatory on behalf of each of the parties. The terms of the 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 the Agreement requested by TDL in order to ensure the Services and TDL (and each Sub-Processor) can comply with any change in Applicable Laws.

#### 18.3 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 the 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 the Agreement.

# 18.4 Severability

If any provision of the 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.

#### 18.5 Sub-contracting and Assignment

TDL may assign or sub-contract the performance of the Agreement (in whole or in part) or any one or more of the Tests to be performed hereunder to any member of the TDL Group or any suitably accredited laboratories including those listed in the Laboratory Guide. The Client may not assign the Agreement or any of its rights or obligations hereunder without the prior approval of TDL.

### 18.6 Relationship of the parties

It is acknowledged and agreed that TDL and the Client are independent contractors and nothing in the Agreement will create or be construed as creating a partnership or 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.

#### 18.7 Notices

All notices given under the 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: post 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 business hours, as soon thereafter as such business hours commence. This clause does not apply to the service of any proceedings or any documents in any legal action or, where applicable, any arbitration or other method of dispute resolution.

### 18.8 Entire agreement

The Agreement is the entire agreement between the Client and TDL and supersedes and extinguishes all prior and contemporaneous agreements, promises, assurances, discussions, representations and understandings between them, whether written or oral, relating to its subject matter. Each party acknowledges that it has not entered into the Agreement in reliance on, and will have no remedies in respect of, any statement, representation, assurance or warranty (whether made innocently or negligently) that is not expressly set out in the Agreement except in the case of fraudulent misrepresentation.

#### 18.9 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 6, and where the Client and The Doctors Laboratory Limited agree that these Terms and Conditions will apply to any Orders, that agreement is also for the benefit of and enforceable by TDL Genetics Limited.

#### 18.10 Governing law

The Agreement and any dispute arising out of or in connection with it (including non-contractual disputes and claims) or its subject matter or formation 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.

#### 19 INTERPRETATION

19.1 In these Terms and Conditions and the Annex:

'Agreement' has the meaning given in clause 1.3;

'Annex' means the annex to the Terms and Conditions:

'Applicable Law' means the laws, regulations and 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;

'Business Day' means a day other than a Saturday, Sunday, or public holiday in England;

'Client' means the person or organisation requesting Services and / or Consumables from TDL and for whom TDL has agreed to provide the Services and / or Consumables:

'controller', 'data subject', 'data protection impact assessment', 'personal data', 'personal data breach', 'process' and 'processor' have the meanings given to those terms in the Data Protection Laws;

'Consumables' means any goods to be provided by TDL in order for the Client to benefit from the Services;

'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 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 a request for Testing submitted by the Client in a format TDL accepts from time to time and by any of the methods TDL accepts from time to time, whether in hard copy or via one of TDL's electronic portals;

'Privacy Notice' means TDL's detailed Privacy Notice available at tdlpathology.com;

'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 providing the Services or Consumables;

'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 pathology 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 11.1:

'TDL' means (i) The Doctors Laboratory Limited or, (ii) TDL Genetics Limited in the case of services offered under the TDL Genetics name:

'TDL Group' means TDL Genetics Limited and 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:

'UK GDPR' has the same meaning as it does in section 3(10) of the Data Protection Act 2018, read with section 205(4) of that Act.

19.2 References to the singular include the plural and vice versa.

19.3 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.

19.4 References to paragraphs are to paragraphs of the Annex.

- 19.5 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.
- 19.6 The Annex is incorporated into these Terms and Conditions.

#### ANNEX

#### Subject matter and nature of processing

- 1.1 TDL processes Protected Data as 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 Processing 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 or the provision of Consumables, when TDL is required to process 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:
- Samples and Test results for the purpose of providing clinical pathology Services;
- 1.2.2 information about clinicians who order Tests, for the purposes of reporting the Test results to the Client;
- 1.2.3 information about a patient's health insurance for the purposes of administering payment for the Services; and
- 1.2.4 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 or provide the Consumables.

#### 3 Types of personal data

- 3.1 The Protected Data may 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 Tests conducted
- 3.1.7 results of 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 Tests, and clinicians who request Tests.

#### 5 Reporting 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 9 to 16.

#### SERVICE-SPECIFIC TERMS OF THE DOCTORS LABORATORY FROM 1ST JAN 2025 FOR HISTOPATHOLOGY SERVICES

The definitions which apply to these Service-Specific Terms are set out in clause 6.

#### 1 THE HISTOPATHOLOGY SERVICES

- 1.1 These Service-Specific Terms apply to any histopathology Services that the Client places an Order for, including any support for histopathology cases at multidisciplinary team ('MDT') meetings or preparation and investigation of frozen sections.
- 1.2 These Service-Specific Terms are in addition to the Terms and Conditions.
- 1.3 In the event that TDL's pathologist reasonably determines that it is necessary for a histopathology Sample to be referred for additional analyses (including special staining, immunohistochemistry and molecular diagnostics), the Test Turnaround times referred to in the Terms and Conditions will not apply and instead TDL will carry out the necessary analyses as soon as reasonably possible. TDL will use reasonable endeavours to inform the requesting clinician that additional analyses have been requested on that Sample.
- 1.4 Where a Client requests an Urgent Test, TDL will use reasonable endeavours to contact the referring clinician promptly to provide the initial and supplementary reports as required. The Client is responsible for ensuring that TDL is notified when the Client considers that a Test is Urgent. TDL reserves the right to suspend this aspect of the Service if it considers that the Client is sending an unreasonable number of Urgent Tests and the position cannot be resolved through good faith discussions between the parties within 30 days. 'Urgent' means a Test which has been marked as urgent in the manner and under any conditions that TDL informs the Client that Tests may be marked as urgent.
- 1.5 TDL will use reasonable efforts to make its pathologists available on reasonable notice to discuss histopathology Test results and Reports with the Client.
- 1.6 The Client acknowledges and agrees that TDL's consultants may report on Samples by remotely reviewing digital images, provided that TDL will adopt Good Industry Practice in the production and review of such images.

#### 2 STORAGE

2.1 TDL will store the Samples in accordance with the guidelines of the Royal College of Pathologists for the retention and storage of pathological records and specimens.

#### 3 SUPPORT FOR MULTIDISCIPLINARY TEAM MEETINGS

- 3.1 TDL will, where agreed between the parties, provide the Sample slides and/or images and the relevant Report and associated documentation (together the 'Sample Information') at the relevant MDT meeting, provided that the Client:
- 3.1.1 in relation to Sample Information which has been tested less than six months ago: requests the Sample Information no less than two Business Days in advance of the date of the MDT meeting where the Sample Information is needed;
- 3.1.2 in relation to Sample Information which has been tested six months or more ago: requests the Sample Information no less than three weeks in advance of the date of the MDT meeting where the Sample Information is needed.
- 3.2 TDL will, where between the parties, arrange for a pathologist to attend and participate in MDT meeting(s) at the Client's request, provided that:
- 3.2.1 the request for the pathologist to attend the MDT meeting is made no less than 10 Business Dats in advance of the date of that MDT meeting;
- 3.2.2 where the pathologist is asked to attend and participate in an MDT meeting which relates to a Sample that the pathologist did not personally review or a Report that they did not personally provide, there will be an additional cost to be agreed by the parties in advance of the pathologist's attendance at the MDT meeting.

#### 4 FROZEN SECTIONS

- 4.1 If the Client wishes TDL to provide a frozen section Service, the Client must submit an Order for that Service no less than 10 Business Days in advance of the requested start date for the Service. The Client's Order must include details of the case and the clinical purpose of the frozen section, for example the margin clearance, the location where the Service is to be provided, whether the Client requires TDL to provide a cryostat and consumables, and any other information TDL reasonably considers necessary.
- 4.2 Where TDL provides a cryostat or other equipment or consumables in the course of the Service ('TDL Equipment'), title in the TDL Equipment will remain with TDL or TDL's lessor. Risk in the TDL Equipment will pass to the Client when TDL completes the delivery and (where relevant) installation of the equipment at the Client's Facility (as defined below) and will pass back to TDL when TDL removes the TDL Equipment from the Client's Facility. The Client will not permit any

- person to use, move or modify the TDL Equipment, except a person authorised by TDL. The Client will indemnify TDL against any costs, claims, damage or loss arising from the loss, theft or destruction of, or damage to, the TDL Equipment which occurs whilst risk in the TDL Equipment lies with the Client, except to the extent caused by TDL or TDL's personnel.
- 4.3 Where TDL has not agreed to provide a cryostat or other equipment or consumables necessary for performing the frozen section Service, the Client will be responsible, at its own cost, for providing those items for TDL to use. Title and risk in a cryostat or other equipment or consumables provided by the Client ('Client Equipment') will remain with the Client. The Client will ensure that the Client Equipment is fit for the purpose of providing the Services at all times, provided that the Client will not be liable for any failure in the Client Equipment caused by TDL or TDL's personnel.
- 4.4 Where the Client Orders a frozen section Service the Client will:
- 4.4.1 provide at the location where the Service is to be provided a room with a suitable water supply, sink and power supply, and access to a rest area and bathroom facilities for TDL personnel providing the Services at the location (the 'Facility'). The Client will provide this Facility free of charge for the duration of the Services and a reasonable time after the end of the Services to allow TDL to recover any TDL Equipment from the Facility:
- 4.4.2 TDL has the right to decline to provide this Service with no liability to the Client if the Facility is not to TDL's reasonable satisfaction and/or the Facility does not meet the UKAS accreditation standard ISO15189;
- 4.4.3 ensure that the cryostat and associated Consumables and the Facility it is in are suitable for the purpose of TDL providing the Services required by the Client. to TDL's reasonable satisfaction:
- 4.5 TDL will provide a suitably skilled and experienced biomedical scientist at the Facility to prepare the Samples, including preparing glass slides during the frozen section procedure.

#### 5 PRICING

5.1 The price payable by the Client for the Services will be the price agreed between the parties. If the Client submits an Order before the price has been agreed, TDL will not carry out the Services and will not be responsible for the Sample.

#### 6 INTERPRETATION

- 6.1 In these Service-Specific Terms, any expressions defined in the Terms and Conditions and used in these Service-Specific Terms have the meaning set out in the Terms and Conditions.
- 6.2 In these Service-Specific Terms the 'Terms and Conditions' means the most recent version of the Terms and Conditions of Business of The Doctors Laboratory published on TDL's website.

# **Request forms**

Visit the TDL website to download:

- TDL Request Form
- Maternal Screening Request Form: For Down, Edwards and Patau Syndromes screening
- Leukaemic Studies Request Form (Cytogenetics/Molecular genetics)
- Genetic Request Form
- TDL Supplies Re-order Form



SCAN ME

Download TDL Request Forms from:

www.tdlpathology.com/ tests/request-forms/

Vacutainer	Anticoagulant	Capacity	SAMPI
Lavender	EDTA	4ml/6ml*	A
Gold	SST/Gel	5ml	В
Light blue	Citrate	4.5ml	•
Red	None	6ml	<b>(</b>
Grey	Fluoride oxalate	2ml, 4ml	G
Green	Lithium heparin	6ml	•
Dark blue	Trace metal	7ml	K
* 6ml EDTA tubes are used	for specific PCR assays		
Blood culture bottle: o	ontact laboratory		ВС
Contact laboratory for advice on sample taking			J
Test by appointment			Х
Random faeces			RF
Faecal collection			LF
Random urine			RU
Mid stream urine			MSU
First catch random ur	ine (for DL12/Chlamydia, etc.)		FCRU
30ml aliquot from a 24 hour urine collection – state total volume			CU
•	4 hour urine collection with oric acid added – state total volume	9	PU
Early morning urine (1	st sample of the day)		EMU
60ml container (sterile)			SC
Cytyc thin prep vial			TPV
Orange/Blue swab for	culture – swab in transport mediur	n/Blue microswab	STM
Black charcoal swab			CS
Green viral swab			VS
PCR swab for Chlamy	dia/PCR infection screening		PCR
Tap/bottled water mo	uth wash – 20mls		MW
Ammotic fluid (5mls P	CR – 10mls Karyotype)		AF
Chorionic villus (mediu	ım provided by laboratory)		CVS
Urine cytology contain	er		UCYT

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