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Authorised by: Dr. H. Rizkalla



CAVAN GENERAL HOSPITAL DEPARTMENT OF CLINICAL & LABORATORY SCIENCES

EXTERNAL USER MANUAL

Cavan General Hospital Telephone: +353 (49) 4376293 Fax: +353 (49) 4376993

Director: Dr. Hala Rizkalla E-mail: <u>hala.rizkalla@hse.ie</u>

Laboratory Manager Mr. Brian O'Malley E-mail: <u>brianj.omalley@hse.ie</u>

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CRITICAL VALUES (ADULTS) DIVISION OF BIOCHEMISTRY

DIVISION OF BIOCHEMISTRY			
Analyte	Outside Range		
Sodium	< 120 mmol/L	> 160 mmol/L	
Potassium	< 2.5 mmol/L	> 6.5 mmol/L	
Urea	>30 mn	nol/L (first instance)	
Creatinine	> 354 µr	mol/L (first instance)	
Glucose	< 2.5 mmol/L	> 20 mmol/L	
Adj calcium	< 1.8 mmol/L	> 3.5 mmol/L	
Magnesium	<	< 0.4 mmol/L	
Phosphate	<	< 0.3 mmol/L	
AST		> 510 U/L	
ALT	> 825 U/L		
СК	> 5000 U/L (unless query MI)		
Amylase	> 500 U/L		
Lithium	>	> 1.5 mmol/L	
Triglyceride		> 20 mmol/L	
Total Protein		>100	
Cortisol		<50mmol/l	
CRP	> 300 mg/L		
FT4	>50 pmol/L		
gGFR	<15 mls/min (first instance)		
HBA1c	>100 mmol/mol		
СК		>5000 U/L	

In the case of critical results generated out of hours, the following require communication to NEDOC, Na, K, Urea & Creatinine (First instance), Glucose and CRP. All other critical results can await communication until next working day. Note all critical results generated on Friday evening require phoning to NEDOC

DIVISION OF HAEMATOLOGY		
WBC	$>30.0 \text{ x } 10^9 \text{xl}$	
HGB	<8.5g/l	
PLT	<80 >1,000 x 10 ⁹ x1	
APTT	>120 sec	
INR	>4	
Absolute Neutrophil Count	$<0.5 \text{ x } 10^9 \text{xl}$	
Sickle Screen	Positive in pre-op emergency screen	
Fibrinogen	< 1.0 g/L	
Malaria	Positive	
Infectious Mononucleosis	Positive	

DIVISION OF SURGICAL PATHOLOGY

- 1. Fat in endometrial curettings and endoscopic Biopsies
- 2. 1st Diagnosis of leukaemias/lymphomas, unexpected clinically significant pathology result
- 3. Funisitis
- 4. Necrotizing Fasciitis
- 5. Invasive Organisms in an immunocompromised patient
- 6. Large Vessels in Lung Core Biopsy
- 7. Major disagreement between primary pathologist and external Pathology opinion.
- 8. Acid Fast organisms in any patient
- 9. Pneumocystis in BAL/Washings
- 10. Any organism in CSF

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1. GENERAL INFORMATION

1.1 LABORATORY STAFF & TELEPHONE NUMBERS

All numbers shown are for normal laboratory hours, i.e. Monday to Friday.

Department	Personnel	Telephone No.
Specimen Reception		6840
Consultant Histopathologist	Dr Hala Rizkalla	3108
	Dr Megan Finan	6407
Consultant Microbiologist	Dr. Cathal Collins	Request Consultant
	Dr. Elizabeth Trautt ^(V)	Microbiologist On
		Call at 049-4376000
Consultant Clinical	Dr. Maria Fitzgibbon ^(V)	6298
Biochemist		
Consultant Haematologists	Dr. Anne Fortune (V)	6054
_	Dr. Barry MacDonagh ^(V)	6414
Laboratory Manager	Brian O'Malley	6292
Blood Transfusion	Eamon Hannick*	6271
Haematology &	Anna Dowd*	6294/6296
Coagulation		
Clinical Chemistry	Aileen Reilly*	6297
	Fiona Jennings*	6298
	Elaine Fitzpatrick*	6298
Histopathology/Cytology	Imelda Gibson*	6300
	Brigid Irvine*	
Microbiology	Larry O'Neill*	6295
	Karen Smith *	6295
	Fionnuala Gilmartin*	6295
	Linda Crowe*	6295
	Conor McPhillips*	6295
Histology Reports	Laboratory Secretaries	6293
Microbiology Reports	Laboratory Secretaries	6053
Haematology, Coagulation,	Laboratory Secretaries	6293/6357
Clinical Chemistry Reports		
Surveillance Scientist	Briain McDonald	6916

(V) = Visiting Consultant

* = Chief/Senior Scientists in these departments

All extension numbers are pre-fixed by (049) 437 from outside the hospital.

1.2 LOCATION

The laboratory is located on the first floor of Cavan General Hospital.

1.3 NORMAL LABORATORY HOURS

Monday - Friday	08.00 - 20.00 hours
Lunch	13.00 - 14.00 hours (reduced staff numbers)

1.4 ACCREDITATION

The Laboratory is accredited to ISO15189 through the Irish National Accreditation Board. The current scope of accreditation is available at https://www.inab.ie/fileupload/medical-testing/cavan-general-hospital-231mt.pdf

1.5 GENERAL NOTES

1.5.1 Use of Laboratory

The annual workload of the Laboratory has been increasing. All medical staff are requested to consider carefully the reasons for which they are requesting an investigation before initiating a request. The need for clinical justification applies particularly to emergency investigation requests, especially outside normal working hours. Asher's Catechism (*BMJ*, 2:260; 1954) is still relevant today and its regular application is urged, i.e.

- 1. Why do I request this test?
- 2. What will I look for in the results?
- 3. If I find what I am looking for, will it affect my diagnosis?
- 4. How will this investigation affect my management of the patient?
- 5. Will this investigation ultimately benefit the patient?

1.5.2 Specimens:

The primary responsibility for sample collection lies with the requesting physician. It is imperative that the collector can positively identify the patient from whom a specimen is collected. All specimens must be adequately labelled with the patient's full name and date of birth. It is highly desirable to include the date and time of specimen collection on the specimen also.

If a sample is not taken correctly, the test results may be seriously distorted.

Please ensure that specimen collection containers have not passed their expiry date before use.

1.5.3 Request Forms:

There are 9 Cavan General Hospital request forms for general use:

- General Form for Clinical Chemistry / Endocrinology / Haematology
 / Coagulation / Immunology. Multiple departmental requests may be documented and sent with the appropriate number of specimens to the laboratory.
- Histopathology / Cytology 1 General, 4 anatomical site related
- Microbiology
- Samples for referral to outside laboratories e.g. external test requests

Addressograph labels may be used on all request forms, please affix an addressograph Label to each back copy of the request form.

The request form must contain*:

Patient Details:

- Patient name/identity (*surname & forename*)
- Address
- Date of Birth
- General Practitioner
- Patient Gender

Specimen Details:

- Date and time collected
- Nature of sample requested
- Tests requested

Requesting Details:

- Name of requestor
- Address to send results
- Mode of contact e.g. telephone number

Other Information:

- Full clinical details relevant to investigation
- For microbiology specimens please state any antibiotic used

1.5.4 Transportation of Specimens:

Please seal labelled samples in either the request form with plastic sleeve or in a biohazard plastic bag with the accompanying appropriate request form in the side pocket. Samples may then be sent to the laboratory main laboratory reception area

Where multiple specimens are submitted with a single request form, all specimens must be labelled correct, and where appropriate, the site/source of the specimen.

NOTE: A GP specimen collection service is available to GMS GPs. Information on this service is available from Primary Care on 041-6850700.

1.5.5 Receipt of Specimens

The daily cut off time for receipt of specimens in the laboratory is 5pm on Mondays to Thursdays and 3pm on a Friday.

1.5.6 Storage of Specimens

It is imperative that specimens are transported to the laboratory as soon as possible after collection to ensure that they are stored under optimal conditions. Where a delay in transportation is envisaged, specimens must be stored as follows:

Department Specimens	Store At	Acceptable Time Delay Before Transportation
Haematology	4°C	Same Day
Coagulation	Room Temperature	< 4 Hours /Same Day
Clinical Chemistry	Room Temperature	Same Day
Histopathology	Room Temperature	Same Day
Cytopathology	Room Temperature	Same Day
Microbiology	4°C	Same Day

1.5.7 Rejection of Specimens

Specimens will be rejected for analysis for the following reasons:

- Inadequate/incorrect labelling of specimen or request form
- Inadequate information on the request form or specimen to allow positive patient identification.
- Where specimens have leaked or the container has been damaged during transport.
- Haemolysed blood.
- Where there is obvious inadequacy of specimen.
- Wrong specimen bottle or correct procedure not followed e.g. specimen arrives routinely but should have been sent on ice.
- Specimens requiring prior booking with the laboratory that arrive without arrangement.
- Specimens not meeting the testing criteria e.g. samples for *C. difficile* and Ova and Parasites testing.

1.5.8 Consent

All procedures carried out on a patient need the informed consent of the patient. For most routine procedures, consent can be inferred when the patient presents himself or herself to a medical practitioner and willingly submits to the collecting procedure e.g. venepuncture. Patients should normally be given the opportunity to refuse.

Special procedures, including more invasive procedure, or those with an increased risk of complications to the procedure will need a more detailed explanation and in some cases, written consent.

The requirement for consent for individual tests performed is outlined in the relevant section of this laboratory manual.

1.5.9 Specimen Collection and Patient Preparation Prior to Specimen Collection

Hand hygiene must be performed prior to commencement. Greet the patient and identify yourself and indicate the procedure that will take place. Positive patient identification is **MANDATORY**. Verify that the patient meets and requirements for the testing to be undertaken e.g. fasting status, medication status, predetermined time for specimen collection, etc.

- 1. Standard precautions must be observed when taking blood.
- 2. Disposable non-sterile latex free gloves must be worn by the phlebotomist when taking blood in all circumstances.
- 3. Change gloves between patients
- 4. Wash hands or apply an antimicrobial gel before and after each procedure and on removal of gloves.
- 5. When sampling blood from any patient extreme care must be taken and every patient must be considered as potentially high risk.
- 6. When taking blood ensure the limb is well supported, and the patient is aware to keep it still. The limb may need to be supported by an assistant to achieve this.
- 7. When removing a blunted needle from a limb, ensure that the vacuum bottle has been disconnected from the multi sampler area. Leaving this in situ may cause blood droplets to spray.
- 8. Cover the puncture site with a sterile swab or cotton wool when removing the needle to reduce the risk of blood droplets spraying into the air.

- 9. To remove a blunted needle from the needle holder, press lever on top of vacuette holder, pointing downwards over a sharps bin. Drop into sharps container.
- 10. Avoid spillage of blood. If spillage occurs, clean spillage immediately.
- 11. If a sample bottle breaks, never attempt to pick it up. Avail of the nearest spillage kit and use accordingly to clean the hazardous material.
- 12. The user of 'sharps' is responsible for their safe and appropriate use and disposal. 'Sharps' must never be left for a colleague to tidy up.
- 13. Label the specimen with the appropriate patient details.
- 14. Place the specimen in the bag attached to the request form.
- 15. Take care to prevent needle stick injuries when using and disposing of needles.
- 16. The Pathology Laboratory handles and processes the specimen according to the relevant laboratory method.

Note:

NEVER pour blood from one tube to another since the tubes can have different additives or coatings. Submit the collected specimen, in its entirety to the laboratory. There is no requirement for separation or division of samples.

1.5.9.1 24-Hour Urine Collection

General Information for Patients:

You will receive

- A large plastic container in which to store urine.
- A request form with your details on it.
- A plastic bag in which to return your collection and request form.
- 1. You may need more than one storage container to contain all of your urine for the 24-hour period.
- 2. Make sure each storage container is labelled with your full name and that your hospital number is written on it. If your container is not labelled properly you may be asked to repeat the 24-hour collection.
- 3. Keep your storage container cool throughout the 24-hour collection period until you bring it back
- 4. For certain collections, a blood sample may need to be taken within the 24 hour collection period; you will be informed if this is the case.

Procedure: How to collect your sample.

- 1. Start the 24-hour urine test by urinating directly into the toilet. Do not save this urine.
- 2. After you urinate, write the date and time on your storage container, **<u>this is the start of your test.</u>** Write this time & date on the container.
- 3. For the next 24 hours, collect all your urine into your storage container.
- Exactly 24 hours after you started the test, urinate one last time and place the urine in your storage container. <u>This is the end of your</u> <u>test.</u> Write the date and time the test ended on your storage container.
- 5. If you need to use more than one container during the 24-hour period, use one container at a time. When it is full, collect your urine in the next container.
- 6. Please bring the urine to the hospital as soon as possible. To prevent leaks, make sure the lid is on tightly, and that the container is transported upright inside a plastic bag.
- 7. If you are an inpatient, your nurse will tell you what time to begin and end the collection and will set up more containers, as needed. If you have questions about the procedure, please ask.

1.5.9.2 24-Hour Urine Collection (Acidified)

In the interest of safety, we are no longer issuing pre acidified 24 hour urine containers. Therefore, if you require 24 hr urinary evaluation of any of the following analytes

- Urinary Calcium
- VMA
- Cathecholamines
- Potassium
- Porphyria

Please collect the sample in the plain 24 hour collection bottle. Mark the container clearly 'Acid Required'.

Return to the laboratory as per normal and the sample will then be acidified upon receipt of the sample in Specimen Reception.

1.5.10 Specimen Containers

Refer to section 1.6 below and the inside of the back cover of this manual

1.5.11 Disposal of Sharps:

Please dispose of any sharps in the correct manner, i.e. a SHARPS disposal box. Remember - health and safety is imperative for all.

1.5.12 Results:

Printed reports are returned to all GP practices who opt for this service, on a daily basis. For many GP practices, results are also transmitted electronically via Healthlink. Transmission of results electronically is preferred to printed reports due to the decreased turnaround times and reduction of errors associated with paper systems.

Sample Turnaround Times (TATs)

Those samples whose TAT is usually one working day will be reported within 2 hours of the receipt of sample. Please note that instrument and computer breakdowns may result in delays and TATs may not be attainable for the duration. GP samples that arrive in the afternoon may not be available until the following day. (See page 61 for Microbiology TAT).

Please keep telephone requests for results to a minimum - remember verbal results may lead to transcription errors. Interpretation of results is not always straightforward. Consultations and queries on the interpretation and selection of tests are encouraged. If the validity of a test is doubted it is most important that you contact the Laboratory as soon as possible to repeat the analysis on the stored specimen.

1.5.13 Phoning of Results/Critical Values

Significantly abnormal results i.e. results falling outside defined limits (alert values) will be telephoned, as outlined in the following protocol. These limits are quoted inside the front cover of this manual.



1.5.14 External Tests:

Specimens which need to be sent to external laboratories e.g. Eurofins Biomnis, Mater Misericordiae, NVRL etc must be accompanied by a separate request form. Please ensure all details have been included. Failure to adhere to the above instructions may result in the specimen being rejected for analysis.

Specific forms and containers are used for cervical cytology samples. Specific specimen collection containers are used for Chlamydia testing. Eurofins Biomnis Laboratories perform chromosomal analyses on specimens with the exception of:

Peripheral Blood Samples:

- 1. Newborns & infants of less than 5 years of age for chromosomal analysis
- 2. FISH studies for Microdeletion Syndromes (no age restrictions)
- 3. On-going Family studies
- 4. <u>Solid tissue post natal only</u>

A completed Informed Consent Form for Genetic Testing and a Constitutional Karyotype Information Form is required for all genetic testing. Samples received without these 2 forms cannot be processed. NOTE: The Genetics Laboratory GSTS Pathology 5th Floor Tower Wing Guy's Hospital Great Maze Pond London SE1 9RT analyse specimens for numbers 1 to 4 above.

1.5.15 Information Available

Laboratory management ensure that patients' well-being, safety and rights are the primary considerations. The laboratory provides opportunities for patients and laboratory users to provide helpful information to aid the laboratory in the selection of the examination methods, and the interpretation of the examination results. Any such feedback is welcomed and can be emailed to <u>cavlab1@hse.ie</u>

1.6 SPECIMEN CONTAINERS

This table contains an alphabetical test list of the most common tests requested from the laboratory. Please refer to individual department sections for further test information

Test	Sample required	Department	Container
5' HIAA	Acidified Urine	Outside Laboratory	24 hour Urine (see
			section 1.5.9.2)
Actinomycosis	Pus with "sulphur granules" if present	Outside Laboratory	Sterile Universal
			Container
Acute anterior poliomyelitis	See NVRL User Manual	Outside Laboratory	See NVRL User
			Manual
Acute encephalitis	See NVRL User Manual	Outside Laboratory	See NVRL User
			Manual
AIDS	Clotted blood	Outside Laboratory	
			White
Albumin	Serum	Biochemistry	
			Brown
Albumin Creatinine Ratio	Urine - Random	Biochemistry	
			Plain 🛍
Aldosterone	Serum/Plasma	Outside Laboratory	
			Brown
			(
			Red 💼
Alkaline Phosphatase	Serum	Biochemistry	
			Brown 📖
Alpha 1acid Glycoprotein	Serum	Outside Laboratory	
			Brown

Test	Sample required	Department	Container
Alpha Fetoprotein	Serum	Biochemistry	
			Brown
Alpha1 Antitrypsin	Serum	Outside Laboratory	Brown
ALT	Serum	Biochemistry	Brown
Aluminium	Plasma	Outside Laboratory	Contact phlebotomy
Amikacin	Serum	Outside Laboratory	Brown
Amino Acids	Lithium Heparin	Outside Laboratory	Orange
Amino Acids (Urinary)	Plain- No Boric Acid	Outside Laboratory	Plain
Ammonia	Plasma	Outside Laboratory	EDTA Plasma
Amoebiasis	Warm specimen of faeces (<1hour) for microscopy. Clotted blood for serology	Outside Laboratory	Blue Top
Amylase	Serum	Biochemistry	Brown
Angiotensin Converting Enzyme (ACE)	Serum	Outside Laboratory	Brown
Anthrax	Seek advice from lab	Microbiology	Seek advice from laboratory
Anti Nuclear Antigen	Serum	Outside Laboratory	Brown

Test	Sample required	Department	Container
Anti Streptolysin O titre (ASOT)	Serum	Outside Laboratory	T.
			Brown
Anti Thyroid Antibodies	Serum	Outside Laboratory	
			Brown
Anti Tissue Transglutaminase	Serum	Outside Laboratory	
			Brown
Anti-Hepatitis B titre	Serum	Outside Laboratory	
			Brown
APTT	Sodium Citrate Sample: 3ml (Adults), 1.2ml (Paeds)	Coagulation	(
			Green 💼
Aspergillosis	Clotted blood for precipitin test against A. fumigatus	Outside Laboratory	
			White 📖
AST	Serum	Biochemistry	
			Brown
Atypical pneumonia screen	Serum	Outside Laboratory	
			Brown
Autoantibody Screen(AAS)	Serum	Outside Laboratory	
			Brown
Bacillary dysentery	Faeces	Outside Laboratory	
			Dide Top
Bence Jones Protein.	Urine (Early morning)	Outside Laboratory	
		5	Plain 🗰
Beta 2 Microglobulin	Serum	Outside Laboratory	
			Brown

Test	Sample required	Department	Container
Bilirubin (Direct	Serum	Biochemistry	Brown
Bilirubin (Total)	Serum	Biochemistry	Brown
Blood film	EDTA Sample: 2.7ml (Adults), 1.2ml (Paeds)	Haematology	Red
Blood Grouping	EDTA Sample: 2.7ml or EDTA Sample: 7.5ml	Blood Transfusion	Red
Bone Profile	Serum	Biochemistry	Brown
Brucella Infections	Blood culture. Clotted blood for serology	Outside Laboratory	Blood Culture &
C3/C4	Serum	Outside Laboratory	Brown
CA125	Serum	Biochemistry	Brown
CA15.3	Serum	Biochemistry	Brown
CA19.9	Serum	Biochemistry	Brown
Caeruloplasmin	Serum	Outside Laboratory	Brown
Calcium	Serum	Biochemistry	Brown

Test	Sample required	Department	Container
Calcium	Plain24 Hour Container	Biochemistry	24 hour Urine (see
			section 1.5.9.2)
Candida Infections	Blood culture. Swab form suspected lesion.	Outside Laboratory	Blood Culture &
	Clotted blood for precipitins in suspected systemic		Rlue Ton
	disease.		
Carbornazazina	C armana	Outside Laboratory	white Land
Carbamazapine	Serum	Outside Laboratory	
			Brown
Catecholamines (see Note No.1)	PlainUrine	Outside Laboratory	24 hour Urine (see
			section 1.5.9.2)
Cathecholamines &VMA	Plain Urine	Referral Test	24 hour Urine (see
			section 1.5.9.2)
CEA	Serum	Biochemistry	<u>III</u>
			Brown
Chloride	Serum	Biochemistry	
			Brown
Chloride (Urinary)	Urine	Biochemistry	1 T
			Plain 🏛
Cholera	Faecal sample	Outside Laboratory	
Chalasteral (UDL)	<u>Compute</u>	Dischargister	
Cholesterol (HDL)	Serum	Biochemistry	
	~		Brown
Cholesterol (LDL)	Serum	Biochemistry	
			Brown
Cholesterol (Total)	Serum	Biochemistry	
			Brown

Test	Sample required	Department	Container
Coagulation Screen	Sodium Citrate Sample: 3ml (Adults), 1.2ml (Paeds)	Coagulation	Green
Copper	Plasma	Outside Laboratory	Contact phlebotomy
Cortisol (timed samples)	Serum	Biochemistry	Brown
Creatine Kinase (CK)	Serum	Biochemistry	Brown
Creatinine	Serum	Biochemistry	Brown
Creatinine (Urinary)	Urine	Biochemistry	Plain
Creatinine Clearance	Urine 24 hour collection + Serum (both samples to arrive together)	Biochemistry	24hr Collection + Brown
CRP	Serum	Biochemistry	Brown
Cyclosporin	EDTA	Outside Laboratory	Red
D-Dimers ⁽⁶⁾	Sodium Citrate Sample: 3ml (Adults), 1.2ml (Paeds)	Coagulation	Green
Digoxin	Serum	Outside Laboratory	Brown
Diphtheria	Swab inflamed area/ membranes	Outside Laboratory	Blue Top

Test	Sample required	Department	Container
Drugs Of Abuse	Urine	Outside Laboratory	Plain
Electrolytes (Na, K, Cl)	Serum	Biochemistry	Brown
Electrophoresis	Serum	Outside Laboratory	Brown
Endomysial Antibodies	Serum	Outside Laboratory	Brown
Epanutin (Phenobarbitone)	Serum	Outside Laboratory	Brown
Epilim	Serum	Outside Laboratory	Brown
ESR	Sodium Citrate Sample: 3.5ml (Adults), 2.0ml (Paeds)	Haematology	Mauve 📠
Ethanol	Serum	Biochemistry	Brown
Fasting Lipids	Serum	Biochemistry	Brown
Ferritin	Serum	Biochemistry	Brown
Fetal Genetic RhD Screen	EDTA Sample: 9 ml	Blood Transfusion	Red
Fibrinogen	Sodium Citrate Sample: 3 ml (Adults), 1.2ml (Paeds)	Coagulation	Green

Test	Sample required	Department	Container
Folate	Serum	Biochemistry	
			Brown
Food poisoning	Seek advice from lab	Microbiology	Seek advice from
			lab
FSH	Serum	Biochemistry	
			Brown
fT3	Serum	Biochemistry	
			Brown
fT4	Serum	Biochemistry	
			Brown
Full Blood Count	EDTA Sample: 2.7ml (Adults), 1.2ml (Paeds)	Haematology	
			Red 💼
Gamma GT	Serum	Biochemistry	
			Brown
Giardiasis	Faeces examination for cysts or duodenal aspirate for	Outside Laboratory	
	trophozoites		
	Clotted blood for IFAT		Prown
Glucosa	Eluoride Ovalate	Biochemistry	
Glucose	Fluoride Oxalate	Diochemistry	X7 11
			Yellow
Gonorrhoea & Chlamydia PCR	Specimen Collection Devices (Swab & Urine)	Outside Laboratory	Contact Laboratory
Growth Hormone	Serum	Outside Laboratory	
			Brown
Haemochromatosis Screen	EDTA	Outside Laboratory	
			Red 🚥

Test	Sample required	Department	Container
HbA1C	Potassium EDTA – separate sample required	Biochemistry	Grey
HCG	Serum	Biochemistry	Brown
Hepatitis A	Serum	Outside Laboratory	Brown
Hepatitis A, B & C	Clotted blood	Outside Laboratory	Brown
Hepatitis B	Serum	Outside Laboratory	Brown
Hepatitis B PCR	Serum	Outside Laboratory	Brown
Hepatitis BsAg	Serum	Outside Laboratory	Brown
Hepatitis C	Serum	Outside Laboratory	Brown
Hepatitis C PCR	Serum	Outside Laboratory	Brown
IgE	Serum	Outside Laboratory	Brown
Immunoglobulin A (IgA) g/L	Serum	Biochemistry	Brown
Immunoglobulin G (IgG) g/L	Serum	Biochemistry	Brown

Test	Sample required	Department	Container
Immunoglobulin M (IgM) g/L	Serum	Biochemistry	Brown
Infectious Mononucleosis (Monospot)	EDTA Sample: 2.7ml (Adults), 1.2ml (Paeds)	Haematology	Red
Iron	Serum	Biochemistry	Brown
Lactate Dehydrogenase (LD)	Serum	Biochemistry	Brown
LFT	Serum	Biochemistry	Brown
LH	Serum	Biochemistry	Brown
Lithium	Serum	Biochemistry	Brown
Magnesium	Serum	Biochemistry	Brown
Malaria Testing	EDTA Sample: 2.7ml (Adults), 1.2ml (Paeds)	Haematology	Red
Measles	Seek advice from lab	Microbiology	Seek advice from lab
Meningitis (<i>bacterial</i>)	CSF for culture & PCR for <i>N. meningitidis.</i> Blood culture. Throat or pernasal swab. Tissue fluid aspirate from skin for culture of <i>N.</i> <i>meningitidis / microscopy</i>	Outside Laboratory	Sterile Universal Container Blue Top

Test	Sample required	Department	Container
Meningitis (viral)	CSF and faeces for virology.	Outside Laboratory	Sterile Universal
	Clotted blood		Container, Blue Top
			And the second se
			Brown
Microalbumin	Urine	Biochemistry	
			Plain 🛍
Mycoplasma infections	Paired sera for atypical infections	Outside Laboratory	
			Brown
Oestradiol	Serum	Biochemistry	
			Brown
Oligoclonal Bands	Serum & CSF	Outside Laboratory	
			Brown
Ornithosis	Serum	Outside Laboratory	.
			Brown
Paratyphoid B	Faecal sample	Outside Laboratory	
Phenobarbitone	Serum	Outside Laboratory	
Theneourone	Serum		Brown
Phenytoin	Serum	Outside Laboratory	
			Brown
Phosphate	Serum	Biochemistry	
		j	Brown
Plague	Seek advice from lab	Microbiology	Seek advice from
			lab

Test	Sample required	Department	Container
Pneumocystis jiroveci	Sputum sample	Histopathology	Sterile Universal
(Previously carinii)			Container
Pneumonia (<i>atypical</i>)	Urine	Outside Laboratory	Plain 🏭
	Sputum sample		Sterile Universal Container
	Serology		Brown
Potassium	Serum	Biochemistry	Brown
Potassium	Urine	Biochemistry	Plain
Progesterone	Serum	Biochemistry	Brown
Prolactin	Serum	Referral Test	Brown
Protein (Total)	Serum	Biochemistry	Brown
Protein (Urinary)	Plain Urine or 24 Hour Collection	Biochemistry	Plain or 24 Hr Container
PSA (Free)	Serum	Biochemistry	Brown
PSA (Total)	Serum	Biochemistry	Brown

Test	Sample required	Department	Container
PT/INR	Sodium Citrate Sample: 3ml (Adults), 1.2ml (Paeds)	Coagulation	Green
PTH	Plasma	Outside laboratory	Red
Quantiferon	Specific containers available from Laboratory	Outside Laboratory	Contact Laboratory
Rabies	Seek advice from lab	Microbiology	Seek advice from lab
Rast	Serum	Outside Laboratory	Brown
Renin	Plasma	Outside Laboratory	Red
Reticulocytes	EDTA Sample: 2.7ml (Adults), 1.2ml (Paeds)	Haematology	Red 📕
Rheumatoid Factor	Serum	Biochemistry	Brown
Rubella	Clotted blood	Outside Laboratory	Brown
Salicylate	Serum	Biochemistry	Brown
Sickle Cell Screen	EDTA Sample: 2.7ml (Adults), 1.2ml (Paeds)	Haematology	Red
Sodium (Serum)	Serum	Biochemistry	Brown
Sodium (Urinary)	Plain	Biochemistry	Plain

Test	Sample required	Department	Container
Syphilis	Clotted blood	Outside Laboratory	Brown
Tegretol	Serum	Outside Laboratory	Brown
Testosterone	Serum	Outside Laboratory	Brown
Theophylline	Serum	Outside Laboratory	Brown
Thyroglobulin antibodies	Serum	Outside Laboratory	Brown
Thyroid antibodies	Serum	Outside Laboratory	Brown
TIBC	Serum	Outside Laboratory	Brown
TORCH Screen	Serum	Outside Laboratory	Brown
Toxicology (Blood)	Serum	Outside Laboratory	Brown
Toxicology (Urine)	Urine	Outside Laboratory	Plain
Toxoplasmosis	Clotted blood	Outside Laboratory	Brown
Transferrin	Serum	Biochemistry	Brown

Test	Sample required	Department	Container
Triglyceride	Serum	Biochemistry	
			Brown
Troponin I	Lithium Heparin	Biochemistry	1
			Orange
TSH	Serum	Biochemistry	T .
			Brown
TSH Receptor Antibodies	Serum	Outside Laboratory	
			Brown
Tuberculosis (non-pulmonary)	Lymph node and other biopsy for culture in a sterile	Outside Laboratory	Sterile Universal
	container with no fixative .		Container
Tuberculosis (pulmonary)	3 (minimum) early morning sputum samples. Pleural	Outside Laboratory	Sterile Universal
	fluid for AFB and culture.		Container
Tuberculosis (urinary)	3 (minimum) consecutive early morning urine	Outside Laboratory	50ml Red Top
	samples for AFB and culture. Min. of 50 mls each		
	collection		
U&E	Serum	Biochemistry	
			Brown
Urate	Serum	Biochemistry	
			Brown
Urea	Serum	Biochemistry	
			Brown
Valproate	Serum	Outside Laboratory	
			Brown
Vitamin B12	Serum	Biochemistry	
			Brown

Test	Sample required	Department	Container	
Vitamin D	Serum	Biochemistry		
			Brown	
Whooping cough	NPA or pernasal swab	Outside Laboratory	Specific Wire Swab	
			Available from	
			Microbiology	
Worms	Faeces for ova, cysts & parasites. Whole worm or	Outside Laboratory		
	segment of tapeworm can be sent to laboratory for		Blue Top	
	identification.			
Yellow fever	Contact laboratory	Outside Laboratory		
Yersinia enterocolitica	Faecal sample.	Outside Laboratory		
	Clotted sample for serology		Daoua	
			Brown	
Zinc	Plasma	Outside Laboratory	Contact phlebotomy	

2. BLOOD TRANSFUSION DEPARTMENT

The Blood Transfusion department may be contacted at 049 437 6410. Same day reporting on samples which arrive in the Department after 17.30hrs cannot be guaranteed. Therefore, it is essential to contact the laboratory about samples arriving after this time if the report is required that evening.

2.1 ASSAYS AVAILABLE IN CAVAN GENERAL HOSPITAL

Assay	Specimen	Bottle	Sample	TAT
	-	Colour	Requirements	
Blood Grouping	EDTA (2.7mls) or	RED	Same Day	Same
For Termination	EDTA (7.5mls) – if available		Preferable	Day
of Pregnancy				

2.2 REQUEST FORMS (PINK LF-BT-0015/16)

- 1. Addressograph labels are permitted on Blood Transfusion Request forms.
- 2. It is essential that all sections of the request form are fully and accurately completed with the **essential** information listed below

Essential Information	Additional Important Information
(minimum data required to perform transfusion requests)	•
Patient's: First name	Patient's diagnosis
Surname	Blood group and/or antibodies (if
Date of birth	known)
Address	
Gender	
Signature of the requester &	
bleep number (if applicable)	
Test required	
Date and time of request	
Signature of phlebotomist/	
sampler (Blood Track 'Collect'	
Label)	
Date and time of specimen	
collection	

3. The section on the request form which states, "*Sample taken and patient identification checked by:*" is reserved for the sampler to complete once the blood sample has been obtained. The signature of the sampler and the date and time of sampling must be entered onto the form once the patient's identification has been positively confirmed and the sample has been obtained. This section must be completed manually by the sampler. (Ref: Haemovigilance Procedure CP-HV-0002 Procedure for Labelling Blood Transfusion Samples)

2.3 BLOOD TRANSFUSION SAMPLE TUBES

- 1. All Blood Transfusion samples **MUST be labelled immediately after sampling**. The sample bottles must never be pre-labelled prior to sampling.
- 2. **Positive Patient Identification** must be confirmed by the sampler prior to sampling by:
 - Asking the patient/person to identify themselves by stating their name and date of birth, (if able), by asking, "What is your name and date of birth?"

3 All samples **MUST** be handwritten with the following information:

- Patient's full name (i.e. first name and surname)
- Date of Birth
- Date & time of sample collection
- Location of patient
- Signature/initials of sampler

The **sample tubes must be hand written and signed** by the sampler. Any omissions or errors in sample labelling will result in the sample being rejected by the laboratory.

3. HAEMATOLOGY/COAGULATION DEPARTMENT

The Haematology/Coagulation department may be contacted at 049 437 6296/6294.

Same day reporting on samples which arrive in the Department after 17.30hrs cannot be guaranteed. Therefore, it is essential to contact the laboratory about samples arriving after this time if the report is required that evening.

Assay	Specimen	Bottle	Sample	TAT ⁽⁵	Reference Range		
		Colou	Requirement)		-	
		r	S				
Full Blood	EDTA (2.7mls) – Adult	Red	Same Day	Same	See table below		
Count	EDTA (1.2mls) – Paediatrics	Red	Preferable ⁽¹⁾	Day			
Blood film	EDTA (2.7mls) – Adult	Red	Same Day	Same	Interpretative report		
	EDTA (1.2mls) – Paediatrics	Red	Necessary	Day			
Reticulocytes	EDTA (2.7mls) - Adult	Red	Same Day	Same	0.5-2.5%		
	EDTA (1.2mls) – Paediatrics	Red	Necessary	Day			
ESR	Sodium Citrate (3.5 mls) – Adult	Purple	Same Day	Same	Age mms/hr		
	Sodium Citrate (2.0 mls) – Paediatrics	Purple	Preferable ⁽²⁾	Day		Male	Female
					0-50 yrs	0 - 10	0-12
					50-60 yrs	0 - 12	0-19
					60-70 yrs	0 - 14	0 - 20
					70-120 yrs	0-30	0-35
Infectious	EDTA (2.7mls) – Adult	Red	Same Day	Same	Positive/Negative		
Mononucleosi	EDTA (1.2mls) – Paediatrics		Necessary ⁽³⁾	Day		-	
S			-	-			
Sickle Cell	EDTA (2.7mls) – Adult	Red	Same Day	Same	Positive/Negative		
Screen	EDTA (1.2mls) – Paediatrics		Preferable	Day			
Malaria	EDTA (2.7mls) – Adult	Red	Same Day	Same	Positive/Negative		
Testing ⁽⁶⁾	EDTA (1.2mls) – Paediatrics		Necessary	$Day^{(6)}$			

Assay	Specimen	Bottle	Sample	TAT ⁽⁵	Reference Range
		Colou	Requirement)	
		r	S		
PT/INR	Sodium Citrate (3mls) – Adult ⁽⁴⁾	Green	Same Day	Same	PT 10.2 – 12.1 s
	Sodium Citrate (1.2mls) - Paediatrics ⁽⁴⁾	Green	Necessary	Day	male/female
					Refer to 'CMH Warfarin
					Guidelines'
APTT	Sodium Citrate (3mls) – Adult ⁽⁴⁾	Green	Same Day	Same	APTT 22.9 – 28.6 s,
	Sodium Citrate (1.2mls) - Paediatrics ⁽⁴⁾	Green	Necessary	Day	male/female
Coagulation	Sodium Citrate $(3mls) - Adult^{(4)}$	Green	Same Day	Same	See individual components
Screen	Sodium Citrate (1.2mls) - Paediatrics ⁽⁴⁾	Green	Necessary	Day	
Fibrinogen	Sodium Citrate (3mls) – Adult ⁽⁴⁾	Green	Same Day	Same	1.5 – 4.0 g/L
	Sodium Citrate (1.2mls) - Paediatrics ⁽⁴⁾	Green	Necessary	Day	
D-Dimers ⁽⁶⁾	Sodium Citrate (3mls) – Adult ⁽⁴⁾	Green	Same Day	Same	< 0.5 mg/L FEU considered
	Sodium Citrate (1.2mls) - Paediatrics ⁽⁴⁾	Green	Necessary	Day	Negative
					\geq 0.5 mg/L FEU considered
					Positive

(1) **FBC:** If it is not possible to send the sample to the Laboratory on the day of venepuncture preferably within 4 hours of collection, store it in a refrigerator at 2 to 8°C until it can be sent.

(2) **ESR:** Will be done if sample is received before 17.30 hours. The test takes one hour. Samples stored >4 hours can lead to a false lowering of ESR values.

(3) **Infectious Mononucleosis (Monospot):** Infectious mononucleosis (Monospot) tests are carried out on same day EDTA samples. Infectious mononucleosis (Monospot) requests must be received in the laboratory before 17.30 hours.

- (4) **Coagulation Samples**: It is imperative that coagulation samples are filled to the mark indicated on the container. Coagulation samples must be tested on the day of venepuncture, preferably within 4 hours of collection.
- (5) **D-Dimers**: Due to the introduction of a new D-Dimer assay results will be reported as mg/l FEU (Fibrinogen Equivalent Units) according to the international norm.
- (6) Malaria Testing: Samples requested for Malaria Testing are processed in Cavan General Laboratory with 2 Malaria Rapid Detection Test (RDT) kits. A Malaria RDT kit result will be reported as Positive or Negative.

If the Malaria RDT kit is positive, samples are referred to Mater Hospital (MMUH) for Malaria Microscopy a.s.a.p. and a telephoned result will be phoned to the Clinical team as soon as it becomes available.

If the Malaria RDT is Negative, samples are referred to Eurofins Biomnis Laboratories for testing on the next working day.

All Positive Malaria Microscopy results from the MMUH are referred to PHE Malaria Reference Laboratory, London for confirmatory testing.
Parameter	0 – 3 days	3 day –	1 – 2	2 - 3	3 - 6	6 months	2-6	6 - 12	Adult	Adult
		1 month	months	months	months	- 2 years	years	years	Male	Female
RBC (x $10^{12}/L$)	5.0 - 7.0	4.0 - 6.6	3.0 - 5.4	3.1 – 4.3	4.1 - 5.3	3.9 – 5.1	4.0 - 5.2	4.0 - 5.2	4.5 - 5.5	3.8 - 4.8
Haemoglobin (g/L)	14.0 - 22.0	15.0 - 21.0	11.5 – 16.5	9.4 - 13.0	11.1 - 14.1	11.1 - 14.1	11.0 - 14.0	11.5 – 15.5	13.0 - 17.0	12.0 - 15.0
Hct (1/1)	0.45 - 0.75	0.45 - 0.67	0.33 - 0.53	0.28 - 0.42	0.30 - 0.40	0.30 - 0.38	0.34 - 0.40	0.35 - 0.45	0.40 - 0.50	0.36 - 0.46
MCV (Fl)	100 - 120	92 - 118	92 - 116	87 - 103	68 - 84	72 - 84	75 - 87	77 – 95	83 - 101	83 - 101
MCH (pg)	31 – 37	31 – 37	30 - 36	27 - 33	24 - 30	25 - 29	24 - 30	25 - 33	27 - 32	27 - 32
MCHC (g/L)	30.0 - 36.0	29.0 - 37.0	29 - 37	28.5 - 35.5	30 - 36	32 - 36	31 – 37	31 – 37	31.5 - 34.5	31.5 - 34.5
WBC (X 10 ⁹ /L)	10.0 - 26.0	7.0 - 23.0	5 – 19	5 - 15	6 – 18	6 – 16	5 - 15	5-13	4.0 - 11.0	4.0 - 11.0
Neuts (X 10 ⁹ /L)	4.0 - 14.0	3.0 - 5.0	3 – 9	1-5	1-6	1 – 7	1.5 - 8	2 - 8	2 - 7	2-7
Lymphs (X 10 ⁹ /L)	3.0 - 8.0	2.0 - 8.0	3 – 16	4 -10	4 - 12	3.5 – 11	6-9	1 – 5	1-3	1 – 3
Monocytes	0.5 - 2.0	0.5 – 1.0	0.3 – 1.0	0.4 - 1.2	0.2 - 1.2	0.2 – 1.0	0.2 – 1.0	0.2 – 1.0	0.2 - 1.0	0.2 – 1.0
$(X \ 10^{9}/L)$										
Eosinophils	0.1 – 1.0	0.1 - 2.0	0.2 – 1.0	0.1 – 1.0	0.1 – 1.0	0.1 – 1.0	0.1 – 1.0	0.1 – 1.0	0.02 - 0.5	0.02 - 0.5
(X 10 ⁹ /L)										
Platelets (X 10 ⁹ /L)	150-450	210 - 500	210 - 500	210-650	200 - 550	200 - 550	200 - 450	180 - 400	150 - 400	150 - 400
RDW (%)									11.6 - 14.0	11.6 - 14.0
Basophils (X 10 ⁹ /L)									0.02 - 0.1	0.02 - 0.1

Full Blood Count Specified Ranges

Pregnancy Reference Ranges

Parameter	Units	1st Trimester	2nd Trimester	3rd Trimester
WBC	X 109/L	5.7-13.6	6.2-14.8	5.9-16.9
Haemoglobin	g/L	11.0-14.3	10.0-13.7	9.8-13.7

Reference: Haematological Values during Pregnancy (Blood Cells. A Practical Guide. Barbara J. Bain; 3rd Edition)

4. CLINICAL CHEMISTRY DEPARTMENT

This section attempts to summarise our services and answer the most common questions about testing requirements.

The department may be contacted at 049 437 6298.

Same day reporting on samples which arrive in the Department after 17.30hrs cannot be guaranteed. Therefore, it is essential to contact the laboratory about samples arriving after this time if the report is required that evening.

4.1 ASSAYS AVAILABLE IN CAVAN GENERAL HOSPITAL

	Assays Per	formed in Cavan Genera	l Hospital		
Assay	Refe	rence Range	Sample Type	Additional	TAT
				Information	
Albumin g/L	0 - 4 d	ays 28-44	Serum	Reference Range	72 hrs
	4d - 14	4y 38-54		from Abbott	
	Adult				
	14- 59	y 35 – 50			
	60 - 90	32 - 46			
	90+y	29 - 45			
Albumin Creatinine	Normal	<2.5 mg/mmol male	10ml yellow	Reference Range	72 hrs
Ratio		<3.5 mg/mmol female	topped	from Abbott	
	Microalbuminuria	2.5 - 29 mg/mmol male			
		3.0 - 29 mg/mmol female			
	Macroalbuminuria	>30mg/mmol (both sexes)			
Alkaline Phosphatase	Age	IU/L	Serum	Reference Range from	72 hrs
	0-14 day	70–380		Pathology Harmony	
	14 day – 16yr	60–425			
	Adult	30–130			
ALT		0-55 U/L	Serum	Reference Range	72 hrs
				from Abbott	
Alpha Fetoprotein	0.75	7.4 IU/ml	Serum	Reference Range	72 hrs
				from Abbott	

	Assays Performed in Cavan General Hospital									
Assay	Referen	ce Range	Sample Type	Additional	TAT					
		-		Information						
Amylase	28–10	0 IU/L	Serum	Reference Range	72 hrs					
AST	5-3-	4 U/L	Serum	Reference Range from Abbott	72 hrs					
Bilirubin (Direct	0–8.6 µmol/ L		Serum	Reference Range from Abbott	72 hrs					
Bilirubin (Total)	0 – 7 day Adult	26–205 μmol/L < 21 μmol/L	Serum	Reference Range from Abbott	72 hrs					
CA125	0-35	IU/ml	Serum	Reference Range from Abbott	72 hrs					
CA15.3	0-31	IU/ml	Serum	Reference Range from Abbott	72 hrs					
CA19.9	0-37	IU/ml	Serum	Reference Range from Abbott	72 hrs					
Calcium (Total	0 to 1 year	2.13-2.74	Serum	Reference range up	72 hrs					
Calcium)	1 to 19 year	r 2.29-2.63		to 19 years from						
	>19 year	2.18-2.57		CALIPer. >19 years						
				from CGH study						
Calcium Adjusted	2.2-2.6	mmol/L		Source of evidence	72 hrs					
(Calculation)				Path Harmonisation						
CEA	0-5	ug/L	Serum	Reference range derived from Abbott	72 hrs					

	Assays Performed in Cavan General Hospital						
Assay	Reference Range		Sample Type	Additional	TAT		
				Information			
Cholesterol (HDL)	> 1.0) mmol/L	Serum	ESC Dyslipidaemia	72 hrs		
				Guidelines 2019			
Cholesterol (LDL)	< 3.0) mmol/L	Serum	ESC Dyslipidaemia	72 hrs		
				Guidelines 2019			
Cholesterol (Total)	0-5	.0mmol/L	Serum	ESC Dyslipidaemia	72 hrs		
				Guidelines 2019			
Creatine Kinase (CK)	30-20	0 U/L Male	Serum	Reference range	72 hrs		
	29-168	U/L Female		from Abbott			
Creatinine	0-2 weeks	27 – 81 µmol/ L	Serum	Reference range	72 hrs		
	2w – 5year	14 – 37 μmol/ L		from Abbott			
	5-9 years	25 – 48 µmol/ L					
	9–15 years	30 – 72 μmol/ L					
	Adult	$64 - 104 \mu mol/L Male$					
		49 -90 µmol/L Female					
Creatinine Clearance	80-1	20 ml/min	24 hour urine	Serum sample to be	72 hrs		
			collection	taken within			
				collection period and			
				arrive with the urine			
				sample.			
Cortisol	150-4	455nmol/l	Serum	Reference range	72 hrs		
	For short SynAC	CTHen test, expect an		from Mater In-house			
	increment of >150	nmol/l after 30 minutes.		study.			

	Assays Performed in Cavan General Hospital							
Assay		Reference Range		Sample Type	Additional Information	TAT		
CRP		<5 m	g/L	Serum	Reference range from Abbott	72 hrs		
Electrolytes (Na, K, Cl) (Serum, All in mmol/L)	Sodium Potassium Chloride	0–14 day 15d–1yr 1yr–16yr Adult	133 – 146 3.4–6.0 mmol/L 3.5–5.7 mmol/L 3.5–5.0 mmol/L 3.5–5.3 mmol/L 98– 107	Serum	Reference range from Pathology Harmony (K ⁺ &Na ⁺) Reference range from Abbott	72 hrs		
Ethanol	Up to 100 100-300 >300 mg	Up to 100 mg/dL: euphoric changes, some impairment expected. 100-300 mg/dL: drowsiness, confusion >300 mg/dL : impaired consciousness		Serum	nom robott	72 hrs		
Ferritin	22-275 µş	22-275 μ g/L Male $5 - 205 \mu$ g/L Female		Serum	Reference range from Abbott	72 hrs		
Folate		7 – 46.4 nmol/L		Serum	Reference range from Abbott	72 hrs		
FSH	Follio Mid (cular Cycle	IU/L 3.0-8.1 2.5-16.7	Serum	Reference range from Abbott	72 hrs		

	Assays Performed in Cavan General Hospital								
Assay	I	Referenc	e Range	Sample Type	Additional Information	TAT			
	Lutea	1	1.4 - 5.5						
	Male		0.95–11.95						
fT3	2	2.42-6.00 pmol/L			Reference range from Abbott	72 hrs			
fT4		9.0–19.0 pmol/L			Reference range from Abbott	72 hrs			
Gamma GT	Male 0–55	IU/L	Female 0–38 IU/L	Serum	Reference range from Abbott	72 hrs			
Glucose	Fasting Post prandial	3.9	9 – 5.6 mmol/L 9 – 7.8 mmol/L	Serum	Reference range source from WHO OGTT Diabetic diagnosis range: Fasting >7.0 mmol/L 2hrs glucose ≥11.1 mmol/L	72 hrs			
HbA1C		< 42 mm	nol/mol	Serum	National Reference Range	72 hrs			
HCG	0–5.0 IU	J/L		Serum	Reference range from Abbott Refers to non pregnant females Post-menopausal HCG may be higher (<10u/l)	72 hrs			

	Assays Performed in Cavan General Hospital							
Assay	Referenc	e Range	Sample Type	Additional	TAT			
				Information				
Iron	11 – 31 μm	nol/L male	Serum	Subject to diurnal	72 hrs			
	9 – 30 μmo	l/L female		variation				
Lactate Dehydrogenase	125-220 U/L		Serum	Reference Range	72 hrs			
(LD)				from Abbott				
LH IU/L		IU/L	Serum	Reference range	72 hrs			
	Follicular	1.8 -11.8		derived from Abbott				
-	Mid Cycle	7.6-89.1						
	Luteal	0.6 - 14.0						
	Male	0.6-12.1						
Lithium	0.5 - 1.0	mmol/L	Serum	Tuesdays & Fridays	72 hrs			
				Reference range				
				from Nolen et al.				
				2019				
Magnesium	0.7-1.0	mmol/L	Serum	Reference range	72 hrs			
				derived from				
				Pathology Harmony				
Microalbumin	0–30	mg/L	Urine –	See also ACR	72 hrs			
			random					
Non HDL Cholesterol	<3.4 m	mol/L	NA	Reference range	72 hrs			
(Calculation)				derived from ESC				
				guide 2019				

	Assays Perform	ed in Cavan Genera	al Hospital		
Assay	Reference	e Range	Sample Type	Additional	TAT
				Information	
NT Pro BNP	<75yr <1	125 pg/ml	Serum	Reference range	72 hrs
	>75yr	<450 pg/ml		derived from Abbott	
				NT Pro BNP	
				increases with age	
Osmolality Serum	275-295 m	Osm/Kg	Serum		SD
Osmolality Urine	No range - depends	on fluid intake and	Urine		SD
	balar	nce			
Oestradiol		pmol/L	Serum	Reference range	72 hrs
	Follicular	77 –921		derived from Abbott	
	Mid Cycle	139–2381			
	Luteal	77-1140			
	Male	40–161			
Phosphate	0.8–1.5 1	nmol/L	Serum	Reference range	72 hrs
				derived from	
				Pathology Harmony	
Potassium urine	25 - 125 1	mmol/L	Urine	Interpret with	72 hrs
				respect to both	
				serum and intake	
Progesterone	Luteal 4-	-50 nmol/L	Serum	Reference range	72 hrs
				derived from Abbott	

	Assays Perforn	ned in Cavan Gene	ral Hospital		
Assay	Referenc	e Range	Sample Type	Additional	TAT
		-		Information	
Protein (Total)	60 - 8	0 g/L	Serum	Reference range	72 hrs
				derived from	
				Pathology Harmony	
PSA (Free)* µg/L	Not Inc	licated	Serum	Can only be	72 hrs
				processed on	
				samples less than	
				two hours old.	
PSA (Total) µg/L	Age (years)	Range µg/L	Serum	NPCRG	72 hrs
-	-<49yr	< 2.0		Must be analysed	
	-50 - 59yr	< 3.0		within 24 hours of	
	-60 - 69yr	< 4.0		venepuncture	
	>70yr	< 5.0		National Guideline	
Procalcitonin	0 - 0.2	5 µg/L	Serum	Reference range	
	< 0.1 Bacterial in	nf. Very unlikely		from Abbott	
	0.1 - 0.25 Bacter	rial inf. Unlikely			
	0.1 - 0.5 Bacte	erial inf. Likely			
	>0.5 Bacterial in	nf. Very likely			
Protein/Creatinine Ratio	<22.6 mg/mmol		Urine	Reference range from Abbott See table 1 below for PCR interpretation	24hrs
Rheumatoid Factor	<30 I	U/ml	Serum	Reference range	72 hrs
				derived from Abbott	

	Assay	s Performed in	Cavan Genera	l Hospital		
Assay	Reference Range		Sample Type	Additional	TAT	
		_			Information	
Salicylate	То	xic levels >300 r	ng/L	Serum	Therapeutic range	72 hrs
		Lethal >700 mg/	/1		150 - 300 mg/L	
Sodium urine spot		40 - 220 mmol/l	Ĺ	Urine	Reference range	72 hrs
	Interpret w	ith respect to bot	h serum and		derived from Abbott	
		intake				
Triglyceride		0 - 1.7 mmol/L	,	Serum	ESC Dyslipidaemia	72 hrs
					Guidelines 2019	
Troponin I	< 16 ng/L Female			Serum	Reference Range	72 hrs
		< 34 ng/L Male	;		from Abbott	
Transferrin	0-14 years			Serum		72 hrs
	Male 1.86-3.88 g/L					
	F	Female 1.80-3.91	g/L	_		
		15-60yr				
	N	Iale 1.74-3.64	g/L			
	Fe	emale 1.80-3.82	g/L			
		61-120yr				
]	Male 1.63-3.44 g	:/L			
	F	emale 1.73-3.60	g/L			
Transferrin Saturation		Male	Female	Serum	Reference range	72 hrs
	<1y	4.1-59	9%		source Abbott	
	1-14y	6.5-39	9%			

	Assays Performed in Cavan General Hospital							
Assay		Reference Range			Additional Information	TAT		
	14-19y	9.6-58%	5.2-44%					
	>19y	19-55%	10-50%					
TSH		0.35 – 4.94 mU/L			Reference Range from Abbott	72 hrs		
Urate	Male Fem	210 - 43 nale 140-360	0 μmol/L) μmol/L	Serum	Reference Range from Abbott	72 hrs		
Urea	0 -1 150 2y- Ac	4d 0.8–5.5 l-1yr 1.0–5.5 -16yr 2.5–6.5 lult 2.5–7.5	5 mmol/L .5 mmol/L .5 mmol/L .8 mmol/L	Serum	Reference Range from Pathology Harmony	72 hrs		
Urinary Calcium	2.	5–7.5 mmol/day	(24hr)	Urine	Reference Range from Abbott	72 hrs		
Urinary Creatinine	Ma Fer	le 5100–14200 nale 3900–9400	μmol/L μmol/L	Urine	Reference Range from Abbott	72 hrs		
Urinary Protein 24h		<0.3 g/24h			Reference Range from Abbott	72 hrs		
Urinary Urate	1500-45	500 μmol/L (24h)	r),normal diet		Reference Range from Pathology Harmony	72 hrs		

Assays Performed in Cavan General Hospital				
Assay	Reference Range	Sample Type	Additional	TAT
			Information	
Urate/Creatinine ratio	0 –6wk 0.2 –3.0 mmol/mmol	Urine	Reference Range	72 hrs
(Calculation)	6wk - 2yr = 0.2 - 2 mmol/mmol		Vademecum	
	2–6y 0.2 –1.5 mmol/mmol		Metabolicum	
	6-14yr $0.2-1.0$ mmol/mmol			
	Adult 0.15 –0.6 mmol/mmol			
Vitamin B12	139–651 pmol/L	Serum		72 hrs
Vitamin D	> 50 nmol/L	Serum	Reference Range	72
			from National	hrs
			Osteoporosis Society	7

*: Serum must be separated from cells within 2 hours of venipuncture

**: Reference ranges are derived by manufacturers except where otherwise stated.

PCR	UK CKD	Approximate	Comment
mg/mmol	Interpretation	dipstick equivalent	
<15	Normal	Negative	Normal
16 - 44	Trace Protein	Trace	Trace
			Proteinuria
45 - 100	Clinical	1+	2 or more PCR
	Proteinuria		results >45, in
	or		the absence of
	macroproteinuria		UTI, indicates
			proteinuria
>100	Clinical	2+	Marked
	Proteinuria		proteinuria
	or		
	macroproteinuria		
>450	Nephrotic range	3+	Nephrotic range
	proteinuria		proteinuria

Table-1 shows the PCR Interpretation

Source: UK Kidney Association.

PCR in Pregnancy.

If using protein/creatinine ratio to quantify proteinuria in pregnant women:

• use 30 mg/mmol as a threshold for significant proteinuria

• if the result is 30 mg/mmol or above and there is still uncertainty about the diagnosis of pre-eclampsia, consider re-testing on a new sample, alongside clinical review.

• Source NICE Guideline: Hypertension in Pregnancy, diagnosis & management NG133 . [2019]

Analyte	First Trimester	Second Trimester	Third Trimester	
TSH	0.1 - 3.1	0.2 - 3.3 pmol/L	0.3 - 3.5	
fT ₄	10 - 19 pmol/L	10 - 16 mU/L	9.0 - 15.0	
$fT_3 3.0 - 3.6 \text{ pmol/L} \qquad 3.8 - 5.4 \qquad \qquad 3.5 - 5.5$				
Sample Type is serum in all cases				

Ref: Schneider, H.G. AACB, 2019.

Profile	Tests
U&E	Na, K, Cl, Urea, Creatinine
	Tot Protein, Albumin, ALP, ALT, AST, Total bilirubin
LFT	GGT
Bone	Ca, Po4, ALP, Corrected Ca
Fasting Lipids	Chol., Trig., HDL,LDL

Table -3 Profiles Available in Cavan.

Table-4 Urinary Assays available in Cavan

Urine Assays	Sample	Turnaround time
Albumin/Creatinine Ratio (ACR)	Random Urine	2D
Calcium	Plain	2D
Creatinine	Plain	2D
Sodium	Plain	2D
Potassium	Plain	2D
Chloride	Plain	2D
Microalbumin	Plain	2D
Protein	Plain	2D

* See section 1.5.9.2

4.2 Assays Carried Out at Other Sites

This is an alphabetical list of the commonly requested external tests. For a more comprehensive list please consult the Eurofins Biomnis Laboratories Test Guide, available at

https://www.eurofins.ie/biomnis/test-information/test-guide/.

Test name	Specimen Type	Bottle Colour	Laboratory
Aldosterone (See Note 1)	Serum	Brown	Eurofins Biomnis
Alpha 1acid Glycoprotein	Serum	Brown	Eurofins Biomnis
Alpha1 Antitrypsin	Serum	Brown	Beaumont.
Aluminium	Plasma	Contact phlebotomy	Eurofins Biomnis
Amikacin	Serum	Brown	Eurofins Biomnis
Amino Acids	Lithium Heparin	Brown	Temple St.
Amino Acids	Urine	Plain- No Boric Acid	Temple St.
Ammonia (see note No. 1)	Plasma	EDTA Plasma	Eurofins Biomnis
Angiotensin Converting Enzyme (ACE)	Serum	Brown	Eurofins Biomnis
Anti Nuclear antibody Screen	Serum	Brown	Eurofins Biomnis
Anti Streptolysin O titre (ASOT)	Serum	Brown	Eurofins Biomnis
Anti Thyroid Antibodies	Serum	Brown	Eurofins Biomnis
Anti-Hepatitis B titre	Serum	Brown	NVRL
Atypical pneumonia screen	Serum	Brown	NVRL
Autoantibody Screen(AAS)	Serum	Brown	Eurofins Biomnis
Beta 2 Microglobulin	Serum	Brown	Eurofins Biomnis

Test name	Specimen	Bottle Colour	Laboratory
	Туре		
C3/C4	Serum	Brown	Eurofins
			Biomnis
Caeruloplasmin	Serum	Brown	Eurofins
			Biomnis
Carbamazapine	Serum	Brown	Eurofins
			Biomnis
Catecholamines (see Note	Urine	24 hour	Eurofins
No.1)		acidified (see	Biomnis
		section 1.5.9.2)	
Copper	Plasma	Contact	Eurofins
		phlebotomy	Biomnis
Cyclosporin	EDTA	Red	Eurofins
			Biomnis
Digoxin	Serum	Brown	Eurofins
			Biomnis
Electrophoresis	Serum	Brown	MMUH
Endomysial Antibodies	Serum	Brown	Eurofins
			Biomnis
Epanutin	Serum	Brown	Eurofins
			Biomnis
Epilim	Serum	Brown	Eurofins
			Biomnis
Growth Hormone	Serum	Brown	Eurofins
			Biomnis
Haemochromatosis Screen	EDTA	Red	Eurofins
(Note 2)			Biomnis
Hepatitis A	Serum	Brown	NVRL
Hepatitis B	Serum	Brown	NVRL
Hepatitis B PCR	Serum	Brown	NVRL
Hepatitis BsAg	Serum	Brown	NVRL
Hepatitis C	Serum	Brown	NVRL
Hepatitis C PCR	Serum	Brown	NVRL
IoF.	Serum	Brown	Eurofins
-8-			Biomnis

Test name	Specimen Type	Bottle Colour	Laboratory
Oligoclonal Bands	Serum & CSF	Brown	Eurofins
			Biomnis
Osmolality (Serum)	Serum	Brown	Eurofins
			Biomnis
Osmolality (Urine)	Urine	Plain	Eurofins
			Biomnis
Phenobarbitone	Serum	Brown	Eurofins
			Biomnis
Phenytoin	Serum	Brown	Eurofins
			Biomnis
Prolactin	Serum	Brown	Eurofins
			Biomnis
РТН	EDTA Plasma	Red	MMUH
Rast	Serum	Brown	Eurofins
			Biomnis
Renin (see Note No. 1)	Plasma	Purple	Eurofins
			Biomnis
Tegretol	Serum	Brown	Eurofins
			Biomnis
Testosterone	Serum	Brown	Eurofins
			Biomnis
Theophylline	Serum	Brown	Eurofins
			Biomnis
Thyroglobulin antibodies	Serum	Brown	Eurofins
			Biomnis
Thyroid antibodies	Serum	Brown	Eurofins
			Biomnis
TIBC	Serum	Brown	Eurofins
			Biomnis
Tissue transglutaminase (tTg)	Serum	Brown	Eurofins
			Biomnis
TORCH Screen	Serum	Brown	NVRL
Toxicology (Blood)	Serum	Brown	National
Toxicology (Urine)	Urine	Plain – no boric	Poisons
		acid	Centre

Test name	Specimen Type	Bottle Colour	Laboratory
TSH Receptor Antibodies	Serum	Brown	Eurofins
			Biomnis
Valproate	Serum	Brown	Eurofins
			Biomnis
Zinc	Plasma	Contact	Eurofins
		phlebotomy	Biomnis
Urine assays	Specimen	Bottle	Dept
	type		
Bence Jones Protein.	Urine (Early	Plain- No Boric	Eurofins
	morning)	Acid	Biomnis
Cathecholamines &VMA	Urine	24 hour	Eurofins
		acidified (see	Biomnis
		section 1.5.9.2)	
Drugs Of Abuse	Urine	Plain- No Boric	Beaumont
		Acid	
5' HIAA	Urine	24 hour	Eurofins
		acidified (see	Biomnis
		section 1.5.9.2)	

Notes:

- 1. Frozen specimen, send directly to laboratory for separation.
- For Haemochromatosis Screening, a completed consent form, available from Eurofins Biomnis, is required. This is downloadable from <u>https://www.eurofins.ie/biomnis</u> under Test Information and Test Request/Consent Forms. This form is also available on the Cavan General Hospital Website, under 'Useful Links' (<u>http://www.hse.ie/eng/services/list/3/acutehospitals/Cavan Monaghan/Pathology_Department.html</u>)
- 3. A more comprehensive test guide is available from https://www.eurofins.ie/biomnis/, and follow the links to the online test guide.

4.3 TUMOUR MARKER REQUESTING – USER GUIDELINES

Tumour markers covered by this guideline: CEA, CA-125, CA15-3, CA 19-9, AFP, HCG, PSA

indications for measurement

All requests must be supported by adequate clinical details.

Medical Oncology,	For the monitoring of established
Gastroenterology and related teams	malignancy
	For the monitoring of cirrhosis,
	chronic liver disease, and certain
	premalignant conditions
	For the investigation of Cancers of
	Unknown Primary Origin
	(ESMO/NCCN suggested panel:
	hCG, AFP, PSA, CA 125, CA 15-
	3)
Gynaecology	CA-125 for Ovarian tumours
Surgical Oncology	For the investigation of pancreatic
	tumours and chronic pancreatitis

Clinically justified orders for tumour markers will be accepted from any clinical user where the clinical details meet the above criteria. Other requests need to be discussed on a case by case basis and will not be analysed where a clear indication is lacking.

NB: Any request which does not fulfil the agreed criteria for tumour marker analysis will not be analysed on the day that it is received. These samples will be separated and stored appropriately, for up to 3 months so that they may be analysed at a future date, should the requesting team return with additional information which supports the agreed clinical indications.

Sample Requirements and Stability

Serum samples are required. Samples should be sent to the laboratory as soon as possible after phlebotomy. Samples left unseparated overnight will be unsuitable for analysis. Free PSA is unstable so Free:Total PSA ratio will only be available on samples received within 3 hours of blood taking.

Requesting Tumour Markers:

The following indications are generally recognised in the international literature:-

Tumou	r Ir	ndication	
Marke	<u>r</u>		
*CEA	Colorectal cancer:	Breast cancer:	Ovarian cancer:
	 Staging/Prognosis 	 Prognosis 	• If CA 125 is not
	• Detecting recurrence	• Monitoring	elevated at diagnosis
	• Monitoring therapy	therapy	
	• Screening for Liver Mets		
CA 125	5 Ovarian cancer:		
	 Staging/Prognosis 		
	• Detecting recurrence		
	• Monitoring therapy		
CA 19.	9 Pancreatic Cancer		Ovarian cancer:
	• Prognosis		• If CA 125 is not
	• Monitoring post surgery		elevated at diagnosis
	(may be useful in conjunction		
	with diagnostic imaging)		
CA 15.	3 Breast cancer:		
	• Monitoring therapy		
AFP	Germ Cell Tumour:	Liver tumour:	
	 Diagnosis 	• Monitoring	
	 Staging/prognosis 	high risk patient	
	• Detecting recurrence		
	• Monitoring therapy		
hCG	Germ Cell Tumour:		
	 Diagnosis 		
	 Staging/prognosis 		
	• Detecting recurrence		
	• Monitoring therapy		
PSA	Prostate cancer:		
	• Screening		
	• As an aid to DRE		
	Prognosis		
	• Monitoring for recurrence		

NB: Other Criteria:

- *CEA may also be useful in selected Oncology patients with a diagnosis of prostate, lung and some non-colon malignancy.
- For the investigation of Cancers of Unknown Primary Origin ESMO/NCCN suggested panel: hCG, AFP, PSA, CA 125, CA 15-3
- hCG diagnosis and monitoring of molar disease
- All requests for a specific marker, where there is known or relevant pathology indicated by clinical details, e.g. HCG and AFP with clinical details "Testicular mass detected", should be analysed
- GP requests on patients with known malignancy or previously elevated levels.
- Any other requests, not fitting these criteria, need to be discussed on a case by case basis and will not be analysed where a clear indication is lacking.

4.4 **Reproductive Endocrinology**

Follicle Stimulating Hormone (FSH), Luteinising Hormone (LH), Oestradiol (E2) and Progesterone (Prog)

Woman with a Menstrual Cycle:

Ideally LH, FSH and Oestradiol samples should be taken on day 3 or 4 of the cycle to obtain the best information about ovarian function. Follicular FSH levels tend to rise as a woman gets older and ovarian reserve diminishes and levels may be a little higher than the quoted reference range even in ovulatory cycles. In the early follicular phase of the cycle FSH should be slightly higher than LH. If LH is much higher than FSH at this stage it suggests the possibility of polycystic ovary disease particularly if there is oligoamenorrhoea.

Progesterone should only be measured in the mid-luteal phase to assess ovulation. In a 28 day cycle the mid-luteal time occurs around days 21 to 24 of the cycle. In women with a longer regular cycle, if ovulation occurs, it is later and can be estimated based on the fact that ovulation usually occurs approximately 14 days before the first day of the next cycle. In women with oligoamenorrhoea assessing ovulation is more difficult.

Amenorrhoea (non-pregnant):

LH, FSH and Oestradiol cannot be timed however, when interpreting the results, it is useful to check whether or not the woman has had a period since the blood test was taken when interpreting the results. There is no point in measuring a progesterone level as women with amenorrhoea are not ovulating.

Menopause:

The menopause is a clinical diagnosis and there is no diagnostic blood test. Menopause is defined clinically as 12 months of amenorrhea in a woman over age 45 in the absence of other biological or physiological causes. The average age at menopause is approximately 51 years. Serum FSH concentrations increase across the menopausal transition, but at times may be suppressed into the normal premenopausal range (after a recent ovulation).

4.5 EGFR IN CHRONIC KIDNEY DISEASE

The Chronic Kidney Disease classification is as follows:

Stage	Description	
1	"Normal" GFR	eGFR >90 ml/min/1.73 m ² with other
		evidence of chronic kidney damage*
2	Mild impairment	eGFR 60-89 ml/min/1.73 m ² with other
		evidence of chronic kidney damage*
3A	Moderate impairment	eGFR 45-59 ml/min/1.73 m ²
3B	Moderate impairment	eGFR 30-44 ml/min/1.73 m ²
4	Severe impairment	eGFR 15-29 ml/min/1.73 m ²
5	Established renal	eGFR <15 ml/min/1.73 m ² or on dialysis
	failure	

* "Other evidence of chronic kidney damage" may include:

- Persistent microalbuminuria or persistent proteinuria (in absence of UTI)
- Persistent haematuria (after exclusion of other causes, e.g. Urological disease)
- Structural abnormalities of the kidneys demonstrated on ultrasound scanning or other radiological tests e.g. polycystic kidney disease, reflux nephropathy and/or Biopsy proven chronic glomerular nephritis

NB : Without other evidence, a GFR >90/ml/min **does not indicate CKD**.

Facts about the MDRD eGFR:

- eGFR will be reported in mL/min/1.73m². Since the MDRD formula underestimates GFR in patients with normal or near normal kidney function, eGFRs of ≥90 mL/min/1.73m² will be reported as >90 mL/min/1.73m².
- eGFR is not valid in patients with rapidly changing renal function e.g. acute renal failure. Plasma creatinine should be monitored in these patients.
- The MDRD eGFR calculation was validated in Caucasian and Afro-Caribbean patients with renal disease in the USA. Patients of Afro-Caribbean origin have a higher muscle mass so the eGFR should be multiplied by 1.21 for these patients. Although it has not been validated for all ethnic or population groups, the eGFR has been accepted for use in white and South Asian populations.
- MDRD eGFR has NOT been validated for calculating drug doses.

- Creatinine clearance with timed urine collections is still required for measuring GFR in certain circumstances:
 - Extremes of body size and age e.g. severe malnutrition or obesity, elderly, children < 18 years
 - Pregnancy, Vegan diet, Creatine supplements, Oedematous states
 - Skeletal muscle disease e.g. muscular dystrophy, paraplegia, quadriplegia, amputee
 - Prior to dosing with nephrotoxic/chemotherapy drugs
- Microalbuminuria is still the gold standard for detecting early renal disease in patients with diabetes mellitus.
- <u>eGFR formula varies slightly depending on the method used to</u> <u>analyse creatinine</u>.

4.6 SAMPLE STORAGE

Samples are stored for a of maximum 7 days, (3 days for Hba1c) We try to facilitate add on requests as far as possible within the constraints of sample volume, sample type and the in vitro stability of individual analytes. Consult section 11 page 70 for time limits for requesting additional examinations.

4.7 EXTERNAL QUALITY ASSESSMENT

The department participates in relevant, third party quality assessment schemes. This includes schemes operated by:

- WEQAS (Welsh External Quality Assessment Scheme)
- NEQAS (National External Quality Assessment Scheme, UK)
- IEQAS (Irish External Quality Assessment Scheme)
- RIQAS (Randox External Quality Assessment Scheme)

The Department is committed to ensuring comprehensive external assessment of the test repertoire.

5. MICROBIOLOGY DEPARTMENT

Effective December 2nd 2024, primary care microbiology services are provided by Enfer Medical. For ease of access, the following are the contact details for Enfer Medical:

- Client Services: 045-819 000 and clientqueries@enfermedical.ie.
- GP Clinical Queries: 045-819 000 and gpclinical queries@healthmail.ie

Please note that an Enfer Medical GP User Manual tailored to Cavan users has been developed and can be viewed on the following webpage: <u>https://www.enfermedical.ie/laboratory-testing-cavan/</u>

5.1 INSTRUCTIONS FOR SEMEN ANALYSIS (SPERM TEST) FOR INFERTILITY/POST VASECTOMY.

These tests are now performed in the Rotunda Hospital. Arrangements and appointments can be made by contacting the Rotunda Laboratory at 01-8171739.

5.2 MICROBIOLOGY SPECIFIC SPECIMEN CONTAINERS

Refer to https://www.enfermedical.ie/laboratory-testing-cavan/

6. HISTOPATHOLOGY & CYTOPATHOLOGY DEPARTMENT

6.1 SURGICAL SPECIMENS:

6.1.1 Request Forms:

LF-HIST-0052 Histology/Cytology Request Forms available from laboratory or hospital stores <u>must</u> contain the following details.

- Patients name.
- Patients Hospital Number. If available
- Patient's date of birth
- -Doctors name.
- Examination required
- Date specimen was taken.
- Specimen type(s).
- Clinical details.
- Details of where reports (and copy of report) are to go.
- Date/number of previous histology (if relevant).

Please fill in the request form with as much detail as possible. Use an addressograph label if available. To allow correct matching of previous specimens from a patient, give all identification details, including previous MRN if available. It is mandatory to give sufficient clinical details and to indicate if other specimens have been sent from the present operation (*e.g. wider excision scar from melanoma*) so that all material can be studied together. Otherwise, the specimen will be returned.

6.1.2 Specimen Containers

Place specimens for histopathological examination in an appropriately sized container (available from histopathology laboratory) which allows complete immersion in formalin and which allows the specimen to move freely. 10% formalin is the fixative used for most histopathological specimens.

Ensure that the specimens are correctly labelled with the following information:

Patient's name Patient's Hospital Number if available Date of Birth Specimen type and number (if more than 1 specimen from same patient) Address Date of procedure on container

When several specimens are obtained at one procedure, list them on one single request form and number them to correlate with the relevant specimen container.

Unlabelled containers will <u>NOT</u> be processed and may be returned to their source

6.1.3 Reports/Turnaround Times

A written report is issued on most biopsies within 5 working days of receipt of a specimen in the laboratory. Larger, more complex specimens may take longer. If there are any queries regarding a specimen, either before or after receiving the histopathology report, please ring the secretaries' office (*ext. 6293*) and you will be referred to the appropriate pathologist.

6.1.4 Urgent Biopsies:

If an urgent result is required, please indicate this on the request form, with a contact phone number. It is also helpful to telephone one of the pathologists when sending the specimen. Ensure that the specimen reaches the laboratory without delay.

6.1.5 Referral Rests

Referral tests originate within the Histopathology Laboratory, and the results of these are incorporated into Cavan Pathology Laboratory reports. The list of referral centres is as follows:

Requested Analysis	Referral Laboratory
Colorectal Cancer Mutation Panel	Beaumont Histopathology
Lung Cancer Mutation Panel	Beaumont Histopathology
MSI	Beaumont Histopathology
MLH1 Methylation	Beaumont Histopathology
Melanoma Mutation Panel	Beaumont Histopathology
BRAF Mutation	Beaumont Histopathology
Her2 ISH	Poundbury Cancer Institute
PDL1 (Lung and Melanoma)	Beaumont Histopathology
PDL1 (Breast, Gastric and other)	Poundbury Cancer Institute
PDL1 (Gastric)	Poundbury Cancer Institute
Muscle Biopsy	Beaumont Histopathology
Cobas Pic 3a	Beaumont Histopathology
NTRK Fusion	St. James's Hospital
Flow Cytometry	St. James's Hospital
MTHFR Mutation c677T	Eurofins Biomnis
Prothrombin Mutation G20210A	Eurofins Biomnis
SNP Microarray	Eurofins Biomnis
Chromosome Analysis	Eurofins Biomnis
Chromosome Y Microdeletions	Eurofins Biomnis
CGH Array	Eurofins Biomnis (Viapath)
Cystic Fibrosis Screen	Eurofins Biomnis
JAK2 Mutation	Eurofins Biomnis

6.2 Cytopathology

All Cytology specimens except cervical smears are processed in the Histology Laboratory. A semi-automated liquid-based processor, the Thin Prep processor is used. Request Forms::

LF-HIST-0052 /Cytology /Histology Request Forms available from laboratory or hospital stores <u>must</u> contain the following details. and indicate clearly that the specimen is for cytological examination.

- Patients name.
- Patients Hospital Number.
- Patient's date of birth.
- Consultant's name.
- Examination required
- Date specimen was taken.
- Specimen type(s).
- Ward
- Clinical details.
- Details of where reports (and copy of report) are to go.
- Date/number of previous histology (if relevant).

6.2.1 Specimens Containers::

Almost all specimens except cervical smears should be submitted in a container of Cytolyt (**available from the Histology Laboratory** (Extension 6300). Rinse needle and syringe in cytolyt fluid and inject

into container



Cytolyt Container

Ensure that the specimens are correctly labelled with the following information:

Necessary: Patients name Patients Hospital Number Date of Birth Specimen type and suffix number (if more than 1 specimen from same patient) Address Date of procedure on container

When several specimens are obtained at one procedure, list them on one single request form and number them to correlate with the relevant specimen container.

6.2.2 Turnaround Times

80% of cytology specimens are reported within 5 days as per national Quality Assurance benchmark.

6.2.3 Cervical Smears

If in the screening programme (Cervical Check) the specimen is sent to Quest Diagnostics. Specific request forms are to be used. If the patient is outside the limits of the screening programme the specimen is sent to Eurofins Biomnis Laboratories. See Section 7.

7. EXTERNAL TESTS

These are tests not performed on site in Cavan General Hospital and charges for each test are applied. Route all specimens through the Laboratory with the correct completed forms which will ensure that the sample is sent to the correct location in the correct manner. The following samples are at present dispatched to:

Cervical Cytology	Contact Cervical Check. Samples outside National Screening Programme: Eurofins Biomnis Laboratories Ltd 01 2958545
Toxicology	National Poisons Centre, Beaumont Hospital, Dublin (01) 8379963/8379966
Meningococcal and GroupB strep PCR *	Meningococcal PCR Lab, Children's University Hospital, Temple St., Dublin (01) 8784432 * Sun-Thurs - please contact the Medical Scientist on call who will arrange to have the sample included in the next morning's transport. Fri. & Sat leave samples in lab sample reception for the attention of the Medical Scientist.
Pertussis screening	Children's Health Ireland at Crumlin
Virology	National Virus Reference Laboratory, U.C.D., Belfield Dublin 4. (01) 7161323
Chromosomal Analyses	Eurofins Biomnis Laboratories, see page 13

Eurofins Biomnis Laboratories analyse the majority of the samples referred. Some tests are performed in Dublin, others in France. A full list of tests and specimen requirements for tests analysed by Eurofins Biomnis Laboratories is available at <u>https://www.eurofins.ie/biomnis</u>, under Test Information

For result enquiries phone 1800 252966.

On Call service for technical analysis &/or result enquiry telephone 01 2003825 Main reception number (01) 2958545. For access to the Eurofins results portal, contact client services in Eurofins via email <u>clientservices@eurofins-biomnis.ie</u> or telephone 1800 252 966

8. ADVICE

Scientific and medical advice on issues within the laboratory's range of interest and competence is available. Key contact staff are listed below.

Position	Name	Ext.	Direct Line
Director of Laboratory:	Dr Hala Rizkalla	3108	049
Consultant Histopathologist			4373108
Consultant Microbiologist	Contact Consultant Microbiologist on Call		
	Through Switch (049 4376	000)
Consultant Haematologists*	Dr. Anne Fortune	6054	049
			4376054
Consultant Chemical	Dr. Maria Fitzgibbon	Routine	049
Pathologist		working	4676298
		hours	
Deputy Consultant	Dr. Graham Lee		018032000
Laboratory Manager	Brian O'Malley	6292	049
			4376292

• The Consultant Haematologists provide advice on all clinical and laboratory aspects of Haematology including interpretation of tests and of clinical findings, and recommendations for further clinical, laboratory or therapeutic actions either for specific patients or more generically for development of guidelines and protocols. Advice is given as required to clinicians and patients, management and administrative staff, scientific or nursing staff. Access to advice from the consultant haematologist is available on-site in CGH according to an agreed schedule of attendance at the hospital. Dr. Fortune attends once weekly alternating Mondays & Wednesdays. On call Haematology advice is available by contacting the Haematology Registrar in the Mater Misericordiae Hospital.

9. PROBLEMS / COMPLAINTS/ FEEDBACK

Please do not hesitate to call Cavan General Hospital Pathology Laboratory if assistance is required. Contact details for departments and staff are outlined on page 5 of this manual.

We welcome all submission regarding problems, complaints or feedback on our processes. All such complaints/comments are logged within the quality management system in place, thoroughly investigated and feedback provided to the complainant. Verbal communications are welcomed by the Laboratory Manager (extension 6292) or cavlab1@hse.ie is available for written communications.

The laboratory encourages patients and laboratory users to provide helpful information to aid the laboratory in the selection of the examination methods, and the interpretation of the examination results. Such information can be provided through the HSE Your Service Your Say process, details of which are available at https://www2.hse.ie/complaints-feedback/.

10. DATA PROTECTION POLICY/DATA RELEASE

The Pathology Laboratory complies with the policy of the HSE regarding the legislation pertaining to the rights of the patient and staff and to act in an ethical and responsible manner in maintaining the security and integrity of all personal information

The pathology laboratory retains the following information in relation to each test request received, for a minimum of 30 years, in order to ensure patient history is maintained and that sufficient information is available to staff responsible for the interpretation and reporting of results from the laboratory:

- Patient full name
- Patient medical record number
- Patient date of birth
- For each specimen: date/time of collection, date/time of receipt in the laboratory and date/time of report, specimen type, priority.
- Clinical information provided by clinicians

- The results and where appropriate, interpretation of each test requested.
- Requesting clinician and address

The laboratory is obliged to release patient information to Public Health (Medical Officer of Health/Director of Public Health) in the event that patient samples prove positive for any organism on the Health Protection Surveillance System Notifiable Diseases list. This list and further information is available at

https://www.hpsc.ie/notifiablediseases/listofnotifiablediseases/

11. TIME LIMITS FOR REQUESTING ADDITIONAL

EXAMINATIONS

Department	Time Limit		
Haematology/	Blood film and Infectious Mononucleosis requests must		
Coagulation	be made within 24 hours of sample collection.		
	Additional requests on Coagulation samples must be		
	made within 4 hours of sample collection.		
Biochemistry/	Samples are held in the department for 7 days.		
Endocrinology	Analyte	Time Limit Following Specimen	
		Collection	
	NT Pro BNP	6 days	
	Hba ₁ c	5 days	
	Glucose	7 days	
	CRP	7 days	
	TSH	7 days	
	fT ₄	6 days	
	fT ₃	6 days	
	PSA (Total)	24 hours	
	Iron	7 days	
	Urea/creatinine	7 days	
	Sodium/Potassium	36 hours (once separated)	
	Vitamin B12	7 days	

Department	Time Limit
Histology/	Additional requests on specimens submitted for Cytology
Cytology	must be made within 4 weeks of specimen collection.
	Specimens submitted for Histology are retained in the
	department for 6 weeks. In addition, blocks and slides
	are retained indefinitely on all specimens processed.
	Contact the Histology laboratory for advice regarding
	additional examinations required.

12. REPEAT EXAMINATION DUE TO ANALYTICAL FAILURE OR FURTHER EXAMINATION OF SPECIMENS

Where repeat examinations are required due to analytic failure, every effort will be made to reduce specimen deterioration in the interim, and analyses will be repeated as soon as possible. Where further examinations of specimens are required, where requested by a clinician, these will be carried out provided specimen stability is acceptable. Where deemed necessary, through scientific professional judgement, in the interests of patient care, further tests may be carried out on specimens where results derived initially would warrant further examination.
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BLOOD COLLECTION TUBES (SARSTEDT	')
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Application	Contents	Cap		Order of Draw
Virology, Bacteriology	Serum			1
		White		
Clinical Chemistry	Serum-Gel			2
		Brown		
Coagulation	Trisodium Citrate 1:9			3
		Green		
BNP (Clinical Chemistry)	Potassium EDTA		i	4
		Red		
HbA ₁ c (Clinical Chemistry)	Potassium EDTA			4
· · · · ·		Grey		
Haematology (except ESR)	Potassium EDTA			4
		Red		
Blood Transfusion	Potassium EDTA			5
	(BTS)	Red		
Glucose (Clinical	Fluoride			6
Chemistry)		Yellow		
ESR (Haematology)	Trisodium Citrate 1:4			7
		Mauve		