

Title: SE-HMDS Oncology Cytogenetics User Guide
Subject: Investigations performed in SE-HMDS Cytogenetics

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1. Introduction

Synnovis is a unique partnership of clinical, scientific and operational expertise, with a mission to transform pathology services in the UK. Our organisation is built on scientific expertise, providing a service that helps clinicians create better outcomes for their patients every day.

Our full-service, customer-focused offer is strongly rooted in the patient pathway. We serve our founding NHS Trusts and many other NHS and private hospitals across South-East England.

We are continually focused on innovation, finding new and improved ways to manage the logistics of high-volume pathology testing as well as specialist reference testing. We always strive to improve capabilities to better meet our customers' needs.

The SE-HMDS at King's College Hospital is a regional centre for diagnostic services, providing Immunophenotyping, Cytogenetic, Molecular Diagnostic and Histopathology services covering most of South-East England.

The Cytogenetics laboratory at SE-HMDS offers an extensive testing repertoire, to aid the accurate diagnosis and prognosis of bone marrow disorders, currently utilising on-screen G-banded chromosome analysis, FISH analysis, and the provision of SNP array interpretation.

The Synnovis Cytogenetics laboratory at SE-HMDS is a UKAS accredited medical laboratory, no. 9092.

2. Contact Details

Correspondence Address:

Synnovis Cytogenetics (SE-HMDS)
Ground Floor, Hambleden Wing
King's College Hospital
Denmark Hill
London SE5 9RS

Sample Address:

King's SE-HMDS Laboratory
c/o Central Specimen Reception
Blood Sciences Laboratory
Ground Floor, Bessemer Wing
King's College Hospital
Denmark Hill
London SE5 9RS

General Enquiries:

Email: kch-tr.cytogeneticslaboratory@nhs.net
Phone: 0203 299 7637

3. Main Contacts

Clinical enquiries:

If you have any clinical queries please contact the SE-HMDS Consultants between 9-5pm Monday to Friday.

kch-tr.KHMDC-consultants@nhs.net

Dr Deborah Yallop (SE-HMDS Co-Director)

Deborah.yallop@nhs.net

Dr Shireen Kassam (SE-HMDS Co-Director)

Shireen.kassam@nhs.net

Results and sample requirement enquiries:

If you have any general queries about results not yet received or sample requirements please contact the departmental email that is monitored between 9-5pm Monday to Friday.

Email: kch-tr.cytogeneticslaboratory@nhs.net

Phone: 0203 299 7637

Scientific enquiries:

If you require specific advice or guidance for interpretation of any cytogenetic results received or have other specific queries over assays provided by the department; please contact the Head of laboratory or the departments Principal Clinical Scientists:

Head of Laboratory

Fran Aldridge DipRCPATH

frances.aldridge@synnovis.co.uk

Operations Leads/Principal Clinical Scientists

Remi Oke

Remi.Oke@synnovis.co.uk

Helen Gilbert

Helen.Gilbert@synnovis.co.uk

Phone: 0203 299 7636

4. Hours of Operation

Monday to Friday 9.00am to 5.30pm

Weekends: There is no routine service at weekends. Samples requiring special attention should be arranged in advance.

Bank Holidays: The department is not staffed on Bank Holidays. An email is sent to regular customers in advance detailing arrangements at Christmas and Easter.

5. Sample Types

A Bone Marrow Aspirate (BMA) is the tissue of choice to investigate patients suspected of having leukaemia or related haematological neoplasms. Peripheral Blood (PB) can be sent if disease cells are present in sufficient numbers to allow cell culture. Peripheral blood is suitable for diagnoses of Chronic Lymphocytic Leukaemia (CLL) or T-cell prolymphocytic leukemia (T-PLL).

N.B. BMA should be sent in lithium heparin or sodium heparin vacutainers.

Please DO NOT use other anticoagulants such as EDTA, which may inhibit cell division as required for G-banded chromosome analysis. EDTA is acceptable for FISH only requests. For myeloid referrals please send both a lithium/sodium heparin sample and an EDTA.

6. Dispatch of Samples

To provide an accurate result, samples for the laboratory must be sent in accordance with guidelines to ensure they arrive in a suitable condition to be processed and analysed.

World Health Organisation (WHO) Guidance (2021)¹ states that: "Shippers of infectious substances shall ensure that packages are prepared in such a manner that they arrive at their destination in good condition and present no hazard to persons or animals during transport."

Similarly, under various dangerous goods transport/carriage regulations^{2,3}, it is the responsibility of the consignor (sender/requester) to ensure that all dangerous goods, including diagnostic specimens, are correctly classified and packaged into suitable containers that are correctly marked and labelled.

It is therefore the responsibility of the requestor to ensure that all samples are sent to Synnovis in accordance with the following instructions.

6.1 Packaging requirements

Potentially infectious samples from GPs transported by designated vehicles provided by Synnovis or the local NHS Trust must be carried out in compliance with the UK and European road transport regulations².

Infectious substances include material that is known to contain, or is reasonably expected to contain, pathogens. When in transport, infectious substances must be packaged according to the packing Instruction 650 of ADR as follows:

- All samples in containers (e.g. tube, pot known as the "primary") must be placed in individual sample bags to avoid cross contamination. **Never send samples from different patients in the same sample bag.** Where the primary contains a liquid, then the primary container must be leak proof. Where the primary contains a solid, then the primary container must be sift proof (impermeable to dry contents).
- Individual sample bags should be placed into large, clear, sealable, leak proof, plastic, sample bags (known as the "secondary") that, where the specimen is a liquid, contains

absorbent material sufficient to absorb the entire quantity of the liquid present in the specimen container (e.g. a sufficient amount of paper toweling to absorb any leakage).

- The referral paperwork should be contained in the secondary packaging pocket.
- The large bag should be placed into a suitable rigid sample transport container that meets the testing requirements of the regulations and is correctly marked and labelled.
- Only rigid outer containers supplied by Synnovis or the local NHS Trust may be used to transport samples to the laboratory by road.
- There should be sufficient cushioning lining the outer rigid container to prevent samples becoming unstable.

N.B. Please send samples at the earliest opportunity; samples must be received within 48 hours to ensure sample viability. Samples not sent immediately should be refrigerated at 4°C and sent at the earliest opportunity.

At least 4ml of BMA sample where possible is required for successful cytogenetic studies. It is particularly important that BMA samples for successful Myeloma genomic studies are part of the first draw of the aspirate; as recommended by the European Myeloma Network⁴.

It is advisable to telephone regarding samples that could arrive at the laboratory late in the day or out of hours. A Clinical Scientist may advise sending the sample the following day.

All Friday samples should arrive before 3pm to allow time for culture over the weekend. BMAs from patients with suspected Myeloma need to arrive in the laboratory before 10am on a Friday to allow for the lengthy processing procedure required for CD138+ cell selection before the weekend. As recommended by the European Myeloma Network⁴.

6.2 Request/Referral Forms

Please use the King's SE-HMDS request form which may be retrieved from the South East Genomics website:

<https://southeastgenomics.nhs.uk/glh/cancer-tests/haemato-oncology/>

The reason for referral is important to determine which culture type or other processes need to be set up, which tests to perform, numbers of cells to analyse and sample prioritisation. All relevant clinical and haematological information and likely diagnosis can be included. If the patient is a participant of a research trial, it is important to give details as certain trials can have specific analysis requirements.

The department operates a specimen acceptance policy. The following details are essential requirements for request cards. Samples referred without at least three patient identifiers may not be processed.

Request forms must contain the following information:

- Patient's forename and surname
- Patient's date of birth
- Patient's genetic sex
- Requestor's name and location:

- Internal Request - location (ward code) and clinician details/code
- External Request - address label/surgery and Consultant details.
- NHS and Hospital number
- Type of specimen(s)
- Date & time of specimen collection
- High risk for bacterial or viral infection or confirmed high risk infection; **High risk specimens must be identified to the laboratory using the referral form (Please note: without this information the specimen will not be processed by the laboratory).**
- Test(s) required
- Relevant clinical information, patient history and any transplant donor sex
- Request forms must be dated and signed by those taking the specimen. Please include appropriate contact details. **(Please note: without this information the specimen will not be reported by the laboratory).**

6.3 Rejection of Unacceptable Specimens

Specimens and request forms are checked on receipt to confirm the patient identification (PID) information provided on the form and specimen agree. A minimum of three PID data items (e.g. Full name, DOB, NHS number or hospital number) are required by the laboratory and must match for the specimen to be accepted. Please ensure PIDs and contact details are **clear and legible** on all referral forms sent to SE-HMDS.

Samples without any patient identifiers are discarded and **not processed**.

To avoid a false negative result, **PB** samples that are **more than 4 days old** at receipt will be rejected for investigations such as eosinophilia FISH analysis.

6.4 Policy for High Risk Samples

All samples from patients with a known **danger of infection (DOI)** should be **indicated clearly** on the King's SE-HMDS request form in the appropriate placeholder.

Please note: Specimens indicated with a DOI without further details of the pathogen(s) will not be processed by the laboratory and communication with the referring provider will be attempted. If no response after 48 hours the sample will be disposed of.

Any samples with category 3 pathogen(s) or higher, according to the ACDP (Advisory Committee on Dangerous Pathogens) (such as TB) will not be processed by the laboratory as it does not have the sufficient containment level.

The Health & Safety Executive's approved list of biological agents can be found on their website:

<http://www.hse.gov.uk/pubns/misc208.pdf>

7. Reporting

7.1 Results-online

Kings College Hospital laboratories offer test results online for NHS healthcare professionals. This is a free, secure, electronic, pathology results on-line service and is available to registered users. Please contact Synnovis Customer Support on 0203 299 3576 if you would like to register for access to this service as a new user at an existing referral site.

7.2 Policy for Faxing Reports

SE-HMDS Cytogenetics does not issue reports by fax.

7.3 Accessing Reports

You can access the SE-HMDS results portal here: <https://sehmds.synnovis.co.uk>

If you have not yet requested access, please complete the short form here: <https://forms.office.com/e/vHezuiPLSL>

Using the SE-HMDS portal is the fastest most efficient way to access our results and is available for all NHS users of the SE-HMDS service.

If the portal is not accessible, full copies of authorised reports can be emailed as PDFs.

7.4 Additional Testing

Requests for additional tests on a specimen referral can be made by telephone or email if clinically relevant and agreed with a HCPC registered Clinical Scientist. This is subject to sufficient sample material availability.

7.5 Samples requiring further information

All samples that are not urgent and have an uncertain diagnosis will be held pending further information. Samples referred with urgent referral indications will be processed as appropriate for the disease until additional information is received that indicates tests should be cancelled. Further details are obtained from testing performed by other labs within SE-HMDS, or by telephone or email to the consultant listed on the referral form. This information will be used to decide on the clinical validity of processing the sample; certain samples may have no cytogenetic testing performed, and may be referred for more relevant testing where indicated.

N.B. Consultants are requested to co-operate as fully as possible with this policy; please respond to requests for further clinical information within 7 days otherwise samples will not be analysed. This is to avoid unnecessary work and helps the laboratory to process its large workload.

7.6 Reporting Times

The following table contains turnaround times (TATs) in calendar days. These are within NHSE guidelines, with some TATs that are more stringent due to local agreement*. See section 10 for further details of specific testing for each clinical scenario.

Clinical Scenario	Tests	TAT from test initiation	Priority
Diagnostic ALL or MPAL	<i>BCR::ABL1</i> FISH	3 days*	Very Urgent
	Appropriate reflex testing (see section 10).	14 days	Urgent
Diagnostic AML	APL FISH (<i>PML::RARA/ RARA::</i>)	3 days	Very Urgent
	AML FISH panel (≤80 years)* (inc. sAML FISH) <small>*when eligible for treatment with Gemtuzumab Ozogamicin (Mylotarg®)11 or CPX-351 (Vyxeos®)12</small>	4 days*	Very urgent (local agreement)
	AML FISH panel (>80 years old) (inc. sAML FISH)	14 days	Urgent
	Karyotype	14 days	Urgent
?Burkitt Lymphoma	<i>MYC</i> translocation FISH	3 days	Very Urgent
?other High-grade Lymphoma	High grade lymphoma FISH panel (see section 10)	4 days*	Urgent
?CML (strong indication)	<i>BCR::ABL1</i> FISH (if not previously confirmed by RT-PCR)	3 days	Urgent
	Karyotype	14 days	Urgent
Acute Leukemia with concerns of relapse	Karyotype and/ or FISH as appropriate	14 days	Urgent
CML in transformation	Karyotype	14 days	Urgent

Routine monitoring of AML, ALL, CML	As appropriate	21 days	Routine
MDS	See section 10	21 days	Routine
Myeloma	Myeloma FISH panel (See section 10)	21 days	Routine
CLL	FISH	21 days	Routine
Low grade Lymphoma	FISH as appropriate	21 days	Routine
Other		21 days	Routine
NHSE state that 90% of the samples should be reported within the guideline time (calendar days). All reporting times are subject to change during periods of insufficient staffing levels.			

8. Laboratory Storage of samples

All samples for cytogenetic testing are stored in accordance with the guidelines issued by the Royal College of Pathologists in April 2015⁵.

BMA and PB samples referred to SE-HMDS Cytogenetics are disposed 3 months after receipt of a sample. Cytogenetic preparations (stained slides) are kept for two years after the final report. Digitised images are stored with maintained accessibility for a minimum of 30 years. Fixed cytogenetic cell suspensions are stored for 6 months from receipt of sample. Fluorescence *In-Situ* Hybridisation (FISH) slides are disposed 48 hours after the final written report has been authorised. A representative photographed or digitised image is captured for all patients and stored with maintained accessibility for a minimum of 30 years.

9. Techniques

All cytogenomic investigations and reporting of cytogenomic findings are guided by established best practice guidelines listed in the references ^{6,7,8,9,10,11,12}

9.1 Chromosome analysis

Chromosome analysis is the microscopic examination of chromosomes in dividing cells. Such analysis can detect changes in chromosomal number and structure. Neoplasia may result from acquired cytogenetic abnormalities in otherwise normal individuals. Chromosome analysis allows a whole genome screen at a resolution of 3-10Mb. Tissue needs to be as fresh as possible with viable disease cells present. Cells are processed and stained using banding techniques to produce a karyotype. Abnormalities are defined

and described according to the International System for Human Cytogenomic Nomenclature (ISCN) 2020¹³.

N.B. Analysis may not detect subtle chromosomal abnormalities or clones not represented in dividing cultured cells.

9.2 Fluorescence In-Situ Hybridisation (FISH)

FISH is based on DNA probes annealing to specific target sequence of specimen DNA. Attached to the probes are fluorescent molecules which confirm the presence or absence of a particular genetic aberration when viewed under fluorescence microscopy.

10. Summary of Services Offered for Routine Cytogenetics and FISH

Additional testing required outside of SE-HMDS Cytogenetics testing algorithms may be requested if the patient is a participant of a research trial, however, these requests should be clearly indicated on the referral form.

10.1 Myeloproliferative neoplasms (MPN)

MPN standard investigations:

Referral Indication	Investigations	TAT
?CML	<ul style="list-style-type: none"> Urgent FISH for <i>BCR::ABL1</i> if high suspicion (WCC > 50) and not already done by RT-PCR 	3 days
	<ul style="list-style-type: none"> Full karyotype on BMA 	14 days
Thrombocytosis/ polycythaemia/ ?MPN/ ?ET/ ?PRV	<ul style="list-style-type: none"> Karyotype analysis is NOT routinely performed for MPN referrals.¹⁴ Studies to exclude <i>BCR::ABL1</i> are NOT routinely performed by FISH where CML is not strongly suspected. This can be done by RT-PCR as offered by the Laboratory for Molecular Haemato-Oncology (LMH) in SE-HMDS. 	α
	<ul style="list-style-type: none"> A karyotype study of a BMA will be performed if morphological studies indicate ≥5% of blasts. 	
	<ul style="list-style-type: none"> ○ 5-9% blasts 	21 days
	<ul style="list-style-type: none"> ○ ≥10% blasts 	14 days
	<ul style="list-style-type: none"> Activation of Single nucleotide polymorphism array (SNP-A)* performed by the LMH in SE-HMDS for <ul style="list-style-type: none"> ○ PMF (see below) ○ Post-PV/post-ET MF (see below) 	α

?Myelofibrosis	<ul style="list-style-type: none"> If <5% blasts by morphological studies activation of SNP-A* performed by the LMH in SE-HMDS. 	
	<ul style="list-style-type: none"> A karyotype study of a BMA will be performed if morphological studies indicate ≥5% of blasts. 	
	<ul style="list-style-type: none"> 5-9% blasts 	21 days
	<ul style="list-style-type: none"> ≥10% blasts 	14 days
CML post treatment	<ul style="list-style-type: none"> Where appropriate patients should be monitored using a molecular genetic test to detect gene fusion transcripts instead of cytogenetic methods. Detection of <i>BCR::ABL1</i> [t(9;22)] is offered by the LMH in SE-HMDS[‡]. 	α
	<ul style="list-style-type: none"> Aligned with ELN recommendations¹⁵ the laboratory will perform a karyotype study of a BMA in the following scenarios 	
	<ul style="list-style-type: none"> Treatment failure/resistance to exclude additional chromosome abnormalities (ACA). 	14 days
	<ul style="list-style-type: none"> Progression or suspected progression to CML in accelerated phase or blast phase. 	14 days
	<ul style="list-style-type: none"> FISH (<i>BCR::ABL1</i>) monitoring may be appropriate in rare instances of atypical transcripts that cannot be measured by qPCR 	21 days
Leukaemic Transformation (non-CML MPN → AML)	<ul style="list-style-type: none"> Secondary AML FISH panel 	4 days ^Ω
	<ul style="list-style-type: none"> -5/ del5q -7/ del7q del(17p); TP53 deletion MECOM (3q26) rearrangements 	
	<ul style="list-style-type: none"> Full karyotype on BMA 	14 days

α Refer to LMH Laboratory User's Handbook for TATs

* SNP-A will detect regions of chromosome imbalance at higher resolution than G-banded analysis where present in ≥10% cell population and regions of CN-LOH, but will not detect balanced rearrangements or the presence of independent clones.

‡This test utilises RT-PCR methods (RNA based) so samples must be < 72 hours old when received – refer to the LMH Laboratory User's Handbook Section 8: "Additional Information on Molecular Investigations"

Ω FISH reported in advance of G-banding (4 day TAT) if clinically significant result

10.2 Eosinophilia

Eosinophilia standard investigations:

Referral Indication	Investigations	TAT
Hypereosinophilia (eos count $\geq 1.5 \times 10^9/L$) ?cause	<ul style="list-style-type: none"> • Eosinophilia FISH panel for rearrangements of <ul style="list-style-type: none"> ○ FIP1L1::PDGFRA (4q12) ○ PDGFRB (5q32) ○ FGFR1 (8p11) ○ JAK2 (9p24) ○ ABL1 (9q34) 	21 days
	<ul style="list-style-type: none"> • Full karyotype on BMA only¹⁶. 	21 days
Monitoring of previously detected gene fusion.	<ul style="list-style-type: none"> • Monitoring of FIP1L1::PDGFRA transcripts is available via the LMH in SE-HMDS[‡]. 	α
	<ul style="list-style-type: none"> • Monitoring by FISH can be done for other rare fusions where molecular methods are not available. 	21 days

α Refer to LMH Laboratory User's Handbook for TATs

‡This test utilises RT-PCR methods (RNA based) so samples must be < 72 hours old when received – refer to the LMH Laboratory User's Handbook Section 8: "Additional Information on Molecular Investigations"

10.3 Myelodysplastic/myeloproliferative neoplasms (MDS/MPNs)

MDS/MPN standard investigations:

Referral Indication	Investigations	TAT
?CMML, MDS/MPN or ?atypical CML	<ul style="list-style-type: none"> • FISH to exclude BCR::ABL1 [t(9;22)(q34;q11.2)] for CMML only when not done previously or concurrently by RT-PCR. 	21 days
	<ul style="list-style-type: none"> • Full karyotype on BMA 	
	<ul style="list-style-type: none"> ○ 5-9% blasts 	21 days
	<ul style="list-style-type: none"> ○ $\geq 10\%$ blasts 	14 days
Monitoring	<ul style="list-style-type: none"> • Some monitoring by FISH as indicated to detect diagnostic chromosomal abnormality 	21 days
Leukaemic Transformation (CMML → AML)	<ul style="list-style-type: none"> • Secondary AML FISH panel <ul style="list-style-type: none"> ○ -5/ del5q ○ -7/ del7q ○ del(17p); TP53 deletion ○ MECOM (3q26) rearrangements 	4 days ^Ω
	<ul style="list-style-type: none"> • Full karyotype on BMA 	14 days

α Refer to LMH Laboratory User's Handbook for TATs

Ω FISH reported in advance of G-banding.

* SNP-A will detect regions of chromosome imbalance at higher resolution than G-banded analysis where present in ≥10% cell population and regions of CN-LOH, but will not detect balanced rearrangements or the presence of independent clones.

10.4 Myelodysplastic Neoplasms (MDS)

Please note: referring clinicians must provide the Cytogenetics laboratory with morphology and/or immunophenotyping studies [if not requested to be performed within SE-HMDS]; failure to provide this information will result in the sample being delayed. Fixed cells are stored for 6 months and testing may be requested at a later point upon receipt of this clinical information. Referrers are encouraged to provide an appropriate email address for this communication.

MDS standard investigations:

Referral Indication	Investigations	TAT
?MDS/ pancytopenia/ thrombocytopenia/ neutropenia/ aplastic anaemia/ ITP ?t-MDS	<ul style="list-style-type: none"> If <5% blasts by morphological studies activation of SNP-A* performed by the LMH in SE-HMDS. 	α
	<ul style="list-style-type: none"> Fixed cytogenetic preparations will be stored for 6 months if a G-banded karyotype is necessary to confirm SNP-A findings or to clarify a IPSS-R cytogenetic risk category¹⁷ 	
	<ul style="list-style-type: none"> Full karyotype on BMA if ≥5% blasts by morphological studies 	
	<ul style="list-style-type: none"> ○ 5-9% blasts 	21 days
	<ul style="list-style-type: none"> ○ ≥10% blasts 	14 days
	<ul style="list-style-type: none"> Those with a failed karyotype will have a high risk MDS (translocation) FISH panel performed to detect balanced rearrangements <ul style="list-style-type: none"> ○ MECOM (3q26) rearrangements ○ t(6;9)(p23;q34.1) [DEK::NUP214] ○ t(8;21)(q21.3;q22.1) [RUNX1::RUNX1T1] ○ t(?;12); ETV6 rearrangements ○ KMT2A (11q23) rearrangements including t(9;11)(p21.3;q23.3) [KMT2A::MLLT3] ○ inv(16)(p13.1;q22) or t(16;16)(p13.1;q22) [CBFB::MYH11] <p>...and a SNP-A* will also be activated where possible unless:</p> <ul style="list-style-type: none"> ▪ The patient is known to have previously received a bone marrow transplant. 	As above

	<ul style="list-style-type: none"> In ?t-MDS where the original clonal disorder is persistent within the bone marrow. <p>Where a SNP-A is not feasible FISH for chromosomes 5 and 7 and <i>TP53</i> will also be performed.</p>	
MDS Monitoring	Repeat genomics for MDS in the absence of progression is not routinely performed. But can be requested if required.	21 days
?Leukaemic Transformation/ increased blasts/ ?progression (MDS → ?AML)	<ul style="list-style-type: none"> Secondary AML FISH panel <ul style="list-style-type: none"> -5/ del5q -7/ del7q del(17p); <i>TP53</i> deletion <i>MECOM</i> (3q26) rearrangements 	4 days ^Ω
	<ul style="list-style-type: none"> Full karyotype on BMA 	14 days

α Refer to LMH Laboratory User's Handbook for TATs

Ω FISH reported in advance of G-banding.

* SNP-A will detect regions of chromosome imbalance at higher resolution than G-banded analysis where present in ≥10% cell population and regions of CN-LOH, but will not detect balanced rearrangements or the presence of independent clones.

10.5 Acute myeloid leukaemia (AML) and Acute leukaemias of ambiguous lineage (ALALs)

AML standard investigations:

Referral Indication	Investigations	TAT
APL suspected	<ul style="list-style-type: none"> Rapid Very Urgent FISH for <i>PML::RARA</i> [t(15;17)] FISH for other <i>RARA</i> (17q21.2) rearrangements that can be associated variant APL¹⁸ <ul style="list-style-type: none"> Rearrangement of <i>RARA</i> not with <i>PML</i> will be investigated further by expedited karyotype studies to identify a translocation partner if possible. Normal results will be followed by the appropriate AML FISH panel (see below). 	3 days (target <24 hours)

<p>Diagnostic AML or ALAL (de novo and tAML) patient age ≤80 years (eligible for treatment with Gemtuzumab Ozogamicin (Mylotarg®)¹⁹ or CPX-351 (Vyxeos®)²⁰;</p>	<ul style="list-style-type: none"> • Very Urgent Full AML FISH panel performed to detect: <ul style="list-style-type: none"> ○ MECOM (3q26) rearrangements ○ -5/del5q ○ t(5;11)(q35;p15) [NUP98::NSD1] ○ t(6;9)(p23;q34.1) [DEK::NUP214] ○ -7/del7q ○ t(8;21)(q21.3 q22.1) [RUNX1::RUNX1T1] ○ KMT2A (11q23) rearrangements including t(9;11)(p21.3;q23.3) [KMT2A::MLLT3] ○ t(9;22)(q34;q11.2); [BCR::ABL1] ○ inv(16)(p13.1;q22) or t(16;16)(p13.1;q22) [CBFB::MYH11] ○ del(17p) [TP53 deletion] ○ RARA (17q21.2) rearrangements 	<p>4 days</p>
	<ul style="list-style-type: none"> • For referrals with prominent eosinophils in the absence of other disease defining cytogenetic findings by FISH add the Eosinophilia FISH panel for rearrangements of <ul style="list-style-type: none"> ○ FIP1L1::PDGFRA (4q12) ○ PDGFRB (5q32) ○ FGFR1 (8p11) ○ JAK2 (9p24) ○ ABL1 (9q34) 	<p>14 days</p>
	<ul style="list-style-type: none"> • Full karyotype (with exception of cases with a PML::RARA fusion, for which a karyotype will not be performed unless specifically requested). 	<p>14 days</p>
<p>Diagnostic de novo AML patient age >80 years</p>	<ul style="list-style-type: none"> • Full karyotype (with exception of cases with a PML::RARA fusion, for which a karyotype will not be performed unless specifically requested). 	<p>14 days</p>
	<ul style="list-style-type: none"> • FISH for balanced rearrangements that may be cryptic or subtle by karyotype analysis. <ul style="list-style-type: none"> ○ MECOM (3q26) rearrangements ○ KMT2A (11q23) rearrangements including t(9;11)(p21.3;q23.3) [KMT2A::MLLT3] ○ inv(16)(p13.1;q22) or t(16;16)(p13.1;q22); [CBFB::MYH11] ○ t(?;17)(?;q21); RARA rearrangement 	<p>14 days</p>
	<ul style="list-style-type: none"> • Those with a failed karyotype will have further FISH studies for the following: <ul style="list-style-type: none"> ○ -5/del5q ○ t(6;9)(p23;q34.1) [DEK::NUP214] ○ -7/del7q ○ t(8;21)(q21.3 q22.1) [RUNX1::RUNX1T1] ○ t(9;22)(q34;q11.2); [BCR::ABL1] ○ del(17p) [TP53 deletion] 	

Diagnostic secondary AML	<ul style="list-style-type: none"> • Secondary AML FISH panel 4 days ^α <ul style="list-style-type: none"> ○ -5/ del5q ○ -7/ del7q ○ del(17p); TP53 deletion ○ MECOM (3q26) rearrangements
	<ul style="list-style-type: none"> • Full karyotype on BMA 14 days
Post treatment AML for monitoring	<ul style="list-style-type: none"> • Transcript RNA based quantitative testing[‡] is available via LMH at SE-HMDS for the following. α <ul style="list-style-type: none"> ○ PML::RARA ○ RUNX1::RUNX1T1 ○ CBFB::MYH11 ○ BCR::ABL1 ○ NPM1 mutant (external send away) <p>Cytogenetic studies are not required for these patients.</p>
	<ul style="list-style-type: none"> • In the absence of a molecular MRD marker Karyotype/ FISH is considered if morphology and immunophenotyping results are ambiguous. 21 days
Relapsed AML	<ul style="list-style-type: none"> • Full karyotype 14 days *If no previous cytogenomic testing, case will be treated as per diagnosis
	<ul style="list-style-type: none"> • FISH as indicated to detect diagnostic abnormality/additional abnormalities 14 days

^α Refer to LMH Laboratory User's Handbook for TATs

[‡]This test utilises RT-PCR methods (RNA based) so samples must be < 72 hours old when received – refer to the LMH Laboratory User's Handbook Section 8: "Additional Information on Molecular Investigations"

10.6 Precursor lymphoid neoplasms

10.6.1 ALL

ALL standard investigations:

Referral Indication	Investigations	TAT
Diagnostic B-ALL	<ul style="list-style-type: none"> • B-ALL FISH <ul style="list-style-type: none"> ○ <i>BCR::ABL1</i> [t(9;22)] 	3 days
	<ul style="list-style-type: none"> • If no evidence of a <i>BCR::ABL1</i> fusion, then reflex testing is initiated with the ALL follow-up panel: ○ <i>KMT2A</i> (11q23) <ul style="list-style-type: none"> ▪ If rearranged <i>KMT2A::AFF1</i> [t(4;11)] ▪ If not t(4;11) – karyotype analysis ○ <i>TCF3::PBX1</i> [t(1;19)] ○ <i>TCF3::HLF</i> [t(17;19)] ○ <i>ETV6::RUNX1</i> [t(12;21)] <ul style="list-style-type: none"> ▪ only if patient age ≤40 ○ ABL-class fusions <ul style="list-style-type: none"> ▪ <i>ABL1</i> (9q34) ▪ <i>ABL2</i> (1q25.2) ▪ <i>PDGFRB, CSF1R</i> (5q32) ○ JAK-STAT fusions <ul style="list-style-type: none"> ▪ <i>CRLF2</i> (Xp22.33,Yp11.2) ▪ <i>JAK2</i> (9p24) 	14 days
	<ul style="list-style-type: none"> • Activation of Single nucleotide polymorphism array (SNP-A)* performed by the LMH in SE-HMDS allows detection of microdeletions associated with B-ALL and hyperdiploidy, low hypodiploidy or near-haploidy. 	α
Diagnostic T-ALL	<ul style="list-style-type: none"> • T-ALL FISH panel <ul style="list-style-type: none"> ○ <i>KMT2A</i> (11q23) rearrangements. ○ ABL-class fusions <ul style="list-style-type: none"> ▪ <i>ABL1</i> (9q34) ▪ <i>ABL2</i> (1q25.2) ▪ <i>PDGFRB, CSF1R</i> (5q32) 	14 days

	<ul style="list-style-type: none"> Activation of Single nucleotide polymorphism array (SNP-A)* performed by the LMH in SE-HMDS 	α
ALL monitoring	<ul style="list-style-type: none"> Cytogenetic studies are not usually appropriate for monitoring purposes in ALL. 	
	<ul style="list-style-type: none"> Quantitative detection of BCR::ABL1 [t(9;22)] is offered by the LMH in SE-HMDS[‡]. 	α
ALL relapse	<ul style="list-style-type: none"> If no previous cytogenomic testing, referral will be treated as per diagnosis. 	14 days
	<ul style="list-style-type: none"> FISH and/ or SNP-A as indicated to detect diagnostic abnormality/ additional abnormalities 	14 days/ α

α Refer to LMH Laboratory User's Handbook for TATs

[‡]This test utilises RT-PCR methods (RNA based) so samples must be < 72 hours old when received – refer to the LMH Laboratory User's Handbook Section 8: "Additional Information on Molecular Investigations"

10.7 Mature Lymphoid neoplasms

10.7.1 Chronic Lymphocytic Leukaemia (CLL)

CLL standard investigations:

Referral Indication	Investigations	TAT
CLL Diagnosis & Monitoring (every 12 months)	<ul style="list-style-type: none"> SNP-A* performed by the Laboratory for Molecular Haemato-Oncology (LMH) in SE-HMDS 	α
	<ul style="list-style-type: none"> Lymphoid gene panel including <i>TP53</i> mutation performed by LMH in SE-HMDS 	α
?Richter's transformation	<ul style="list-style-type: none"> HGL FISH panel to detect: <ul style="list-style-type: none"> MYC (8q24.1) rearrangement IGH::MYC [t(8;14)(q24.1;q32)] BCL2 (18q21) rearrangement BCL6 (3q27) rearrangement <ul style="list-style-type: none"> Reflex testing of IGK::MYC / IGL::MYC if MYC rearranged but not with IGH 	4 days

α Refer to LMH Laboratory User's Handbook for TATs

10.7.2 Plasma cell Neoplasms

Please note: Before Myeloma FISH testing can be initiated a diagnosis of Multiple Myeloma (MM) ($\geq 10\%$ clonal bone marrow plasma cells) should be established by either internal SE-HMDS results or external results [if not requested to be performed within SE-HMDS]; failure to provide an external diagnosis will result in the sample being delayed. Fixed CD138+ selected cells are stored for 6 months and testing may be requested at a later point upon receipt of this clinical information. Referrers are encouraged to provide an appropriate email address for this communication.

Genomics is not currently performed for Monoclonal Gammopathy of Uncertain Significance (MGUS), MRD samples or those samples where a diagnosis of MM is not evident from other SE-HMDS studies.

If external trephine results indicate a clear diagnosis of MM, Myeloma FISH studies can be reinstated by referrers provided CD138+ selected cells are available.

MM standard investigations:

Referral Indication	Investigations	TAT
Diagnosis of Plasma cell Myeloma/ Smouldering Myeloma ($\geq 10\%$ plasma cells in the bone marrow)	<p>FISH studies have been selected to enable the detection of cytogenetic aberrations associated with an adverse outcome^{22, 23}</p> <ul style="list-style-type: none"> Myeloma FISH panel on CD138+ selected plasma cells. <ul style="list-style-type: none"> TP53 (17p13) deletion (with 11q22 control) CDKN2C (1p32) deletion CKS1B (1q21) gain IGH (14q32.3) rearrangement MYC (8q24.1) rearrangement <p>Where IGH (14q32.3) is rearranged, sequential reflex FISH for:</p> <ul style="list-style-type: none"> t(4;14)(p16.3;q32); IGH::FGFR3 t(11;14)(q13;q32); IGH::CCND1 t(14;16)(q32;q23); IGH::MAF t(14;20)(q32;q12); IGH::MAFB t(6;14)(p21;q32); IGH::CCND3 	21 days
	<ul style="list-style-type: none"> Where a TP53 deletion is identified by FISH, for transplant eligible patients aged 75 or below (where surplus CD138+ve cells are available); CD138+ve cells will be forwarded to LMH in SE-HMDS for TP53 mutation testing as part of the Lymphoid gene panel. This allows the identification of High risk "double-hit"²³ patients. 	α
Myeloma Relapse or progression	<ul style="list-style-type: none"> FISH on CD138+ selected cells for: <ul style="list-style-type: none"> TP53 (17p13) deletion (with 11q22 control) CDKN2C (1p32) deletion 	21 days

(Smouldering -> MM)	<ul style="list-style-type: none"> ○ CKS1B (1q21) gain ○ Diagnostic IGH t(14;v) rearrangement when present
	<ul style="list-style-type: none"> • Those referrals without known prior successful diagnostic cytogenetic FISH results will be tested as per diagnostic strategy.

10.7.3 B-cell Non-Hodgkin Lymphoma (B-NHL)

FISH analysis for NHL is carried out on uncultured fixed cells from peripheral blood sample or bone marrow aspirate, or from bone marrow smears. Selection of appropriate FISH test(s) will be performed in conjunction with clinical information/request, morphology, flow cytometry, histopathology and immunohistochemistry, and once bone marrow infiltration has been confirmed.

B-NHL Standard investigations:

Referral Indication	Investigations	TAT
?Burkitt Lymphoma	<ul style="list-style-type: none"> • Initial FISH panel to detect <ul style="list-style-type: none"> ○ MYC (8q24.1) rearrangement ○ IGH::MYC [t(8;14)(q24.1;q32)] 	3 days
	<ul style="list-style-type: none"> • Subsequent FISH panel to detect <ul style="list-style-type: none"> ○ BCL2 (18q21) rearrangement ○ BCL6 (3q27) rearrangement ○ +/- Testing of IGK::MYC / IGL::MYC if MYC rearranged but not with IGH 	4 days
High-grade B-NHL / ?DLBCL	<ul style="list-style-type: none"> • HGL FISH panel to detect: <ul style="list-style-type: none"> ○ MYC (8q24.1) rearrangement ○ IGH::MYC [t(8;14)(q24.1;q32)] ○ BCL2 (18q21) rearrangement ○ BCL6 (3q27) rearrangement ○ IGH::CCND1 [t(11;14)(q13;q32)] <ul style="list-style-type: none"> ▪ if specifically requested • Reflex testing of IGK::MYC / IGL::MYC if MYC rearranged but not with IGH 	4 days
?Mantle cell Lymphoma	<ul style="list-style-type: none"> • FISH to detect: <ul style="list-style-type: none"> ○ IGH::CCND1 [t(11;14)(q13;q32)] <ul style="list-style-type: none"> ▪ TP53 (17p13) deletion (if IGH::CCND1 detected) 	21 days
	<ul style="list-style-type: none"> • Lymphoid gene panel including TP53 mutation performed by LMH in SE-HMDS • CCND2 FISH study available at request. 	α

?Follicular Lymphoma	<ul style="list-style-type: none"> FL FISH panel at diagnosis (not required at staging) <ul style="list-style-type: none"> BCL2 (18q21) rearrangement BCL6 (3q27) rearrangement 	21 days
?Splenic Marginal Zone Lymphoma	<ul style="list-style-type: none"> FISH for deletion of 7q available at request.²⁴ 	21 days

10.7.4 Mature T-cell neoplasms

Where indicated FISH analysis can be carried out on uncultured fixed cells from a peripheral blood sample or bone marrow aspirate

PHA-stimulated cultures are used where karyotype analysis is indicated. Selection of appropriate cytogenetic test(s) will be performed in conjunction with clinical information/request, morphology, flow cytometry, histopathology and immunohistochemistry.

Standard investigations:

Referral Indication	Investigations	TAT
?T-PLL	<ul style="list-style-type: none"> Karyotype analysis of PHA-stimulated cultures from peripheral blood – This allows the detection of both major and minor criteria used to establish the diagnosis of T-PLL including the rare t(X;14)²⁵ 	21 days
	<ul style="list-style-type: none"> FISH panel to detect disease defining criteria <ul style="list-style-type: none"> TCL1A rearrangement (14q32) Deletion of 11q (ATM) Detection of gain of 8q (MYC) 	21 days
?Hepatosplenic T cell Lymphoma	<ul style="list-style-type: none"> FISH for isochromosome 7q available at request. 	21 days
Adult T-cell leukaemia/lymphoma (ATL)	<ul style="list-style-type: none"> FISH for TRA/D (14q11) and TP53 (17p13). 	21 days

10.8 Histiocytic cell neoplasms

FISH analysis for Langerhans cell histiocytosis (LCH) is generally carried out on formalin-fixed, paraffin-embedded (FFPE) tissue block sections pre-processed in the histopathology laboratory at King's College Hospital. **Please note that this testing is currently not UKAS accredited.**

Standard investigations:

Referral Indication	Investigations	TAT
Histiocytosis	<ul style="list-style-type: none"> FISH to detect: <ul style="list-style-type: none"> <i>ALK</i> (2p23) rearrangement <i>BRAF</i> (7q34) rearrangement <i>NTRK1</i> (1q23) rearrangement 	21 days

11. Complaints and Compliments

The department has procedures for logging compliments and complaints from service users. Please contact the Head of Service for further details if required.

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