

INTERDISCIPLINARY BLOOD CANCER DIAGNOSTICS



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Please fill out patient data clearly in block letters

Family name

First name

Id. No. Day Month Year

Physician male
 female

Barcode

Client data

SPECIMEN (IMMEDIATE SHIPPING AT ROOM TEMPERATURE)

Peripheral Blood (PB) No. of tubes sent

Bone Marrow (BM) [] []

Slides harvested No. of slides sent
 material (FISH only) [] []

Sampling date Time
 [] [] [] [] [] [] [] [] [] []

ANALYSIS	VOLUME [ml]	ANTICOAGULATION
<input type="checkbox"/> Immunophenotyping and microscopy (Please provide 2-4 unstained slides additionally)	2	EDTA
<input type="checkbox"/> Fluorescence in-situ hybridization (FISH)	2-3	Heparin
<input type="checkbox"/> Chromosome analysis (Cytogenetics)	5-7	Heparin
<input type="checkbox"/> Molecular genetic analysis	2 x 4	EDTA

INDICATION

Non-Hodgkin's Lymphoma B-NHL T-NHL

MGUS/Multiple Myeloma
 (For FISH analysis, "magnetic activated cell sorting" (MACS) will be initially performed to isolate CD138+ plasma cells)

Acute Leukemia: AML B-ALL T-ALL

MDS CML MPN PNH (PB only) Mastocytosis

CLINICAL COURSE

Initial diagnosis:
 Follow up Relapse

Therapy: _____

CLINICAL FINDINGS, REQUEST AND THERAPY (MANDATORY)

Informed patient consent (for research purposes)

I hereby give consent for fully de-identified leftover material of my specimens to be stored or used for the purposes of research in accordance with the EU General Data Protection Regulation. I have been informed that I can revoke this consent at any time in the future without giving any reasons.

 Place, Date Signature of patient or the patient's legal representative

If you have any questions please do not hesitate to contact us: +49 6132 781-240
 A list of our FISH probes can be found online at genetik.bioscientia.de/en/request-forms/

FLUORESCENCE IN-SITU HYBRIDIZATION (FISH) HEMATOLOGICAL DIAGNOSTICS

Please choose and mark the requested probes for the above mentioned patient.

Acute Myeloid Leukemia (AML)

- BCR::ABL1 t(9;22)(q34;q11.2)
- CBF::MYH11 inv(16)(p13q22)/t(16q22)
- D20S108 (20q12), RH7808 (20q13.3), D8Z2 (deletion 20q, trisomy 8)
- DEK::NUP214 t(6;9)(p22;q34)
- EGR1 (5q31), RPS14 (5q33) with control in 5p15 (5q deletion/monosomy 5)
- KMT2A(MLL) t(11q23)
- KMT2A(MLL)::MLLT1(ENL) t(11;19)(q23.3;p13.3)
- KMT2A(MLL)::MLLT3(AF9) t(9;11)(p21.3;q23.3)
- KMT2A(MLL)::AFDN(MLLT4) t(6;11)(q27;q23.3)
- KMT2E (7q22), D7S522/MET (7q31.2) with control D7Z1 (7q deletion/monosomy 7)
- PML::RARA t(15;17)(q24.1;q21.1)
- RPN1::MECOM t(3;3)(q21.3;q26.2)/inv(3)(q26.2)
- RUNX1T1::RUNX1 t(8;21)(q22;q22)
- TP53 (17p13) in combination with ATM (11q22) (deletion 17p/11q)

Chronic Myeloid Leukemia (CML)

- BCR::ABL1 t(9;22)(q34;q11.2)
- TP53 (17p13.1) in combination with ATM (11q22.3) (deletion 17p/i(17q))

Myelodysplastic Neoplasia (MDS)

- Complete MDS panel (deletion 5q/monosomy 5, deletion 7q31/monosomy 7, trisomy 8, deletion 12p, deletion 17p/i(17q), deletion 20q12, trisomy 21, -Y)
- D20S108 (20q12), RH7808 (20q13.3), D8Z2 (deletion 20q, trisomy 8)
- ETV6::RUNX1 (deletion 12p; deletion 21q22/trisomy 21)
- EGR1 (5q31), RPS14 (5q33) with control in 5p15 (deletion 5q/monosomy 5)
- KMT2E (7q22), D7S486 (7q31) with control D7Z1 (deletion 7q/monosomy 7)
- TP53 (17p13) in combination with ATM (11q22) (deletion 17p/i(17q), deletion 11q)
- Loss Y-chromosome (DYZ3)

Chronic Myeloproliferative Neoplasias/ Chronic Myelomonocytic Leukemia/ Chronic Eosinophilic Leukemia/ Hypereosinophilic Syndrome (MPN/CMML/CEL/HES)

- BCR::ABL1 t(9;22)(q34;q11.2)
- ETV6::RUNX1 (deletion 12p/ETV6-rearrangement)
- D20S108 (20q12), RH74808 (20q13.3), D8Z2 (deletion 20q/trisomy 8)
- FGFR1 t(8p11)
- JAK2 t(9p24)
- KMT2E (7q22), D7S522/MET (7q31.2) with control D7Z1 (7q deletion/monosomy 7)
- PCM1::JAK2 t(8;9)(p22;q24)
- PDGFRA/CHIC2/FIP1L1 (deletion 4q12, FIP1L1::PDGFR-rearrangement)
- PDGFRB t(5q32)
- TP53 (17p13) in combination with ATM (11q22) (deletion 17p/i(17q), deletion 11q)

Chronic Lymphocytic Leukemia (B-CLL)/ Small Lymphocytic Lymphoma (SLL)/ Lymphoplasmocytic Lymphoma Waldenstrom Macroglobulinemia

- Complete B-CLL panel (deletion 6q23, deletion 11q22, trisomy 12, deletion 13q14, deletion 13q34, IGH-rearrangement t(14q32), deletion 17p13)
- ATM (11q22) and TP53 (17p13) (deletion 11q, deletion 17p)
- BCL3 t(19q13)
- CCND1::IGH t(11;14)(q13;q32)
- Centromere 12, DLEU (13q14), LAMP1 (13q34) (trisomy 12, deletion 13q14, deletion 13q)
- IGH t(14q32)
- IGH::BCL2 t(14;18)(q32;q21)/trisomy 18
- MYB (6q23) with controls in 6q21(SEC63) and D6Z1

Non-Hodgkin-Lymphoma (NHL)/ B-Cell-Lymphoma (B-NHL)

- Complete NHL panel (deletion 6q23, deletion 11q22, trisomy 12, deletion 13q14, deletion 13q34, IGH-rearrangement t(14q32), deletion 17p13)
- ALK t(2p23) in ALK+ DLBCL
- ATM (11q22) and TP53 (17p13) (deletion 11q, deletion 17p)
- BCL3 t(19q13)
- BCL6 t(3q27)/gain 3q
- CCND1::IGH t(11;14)(q13;q32)
- CCND3::IGH t(6;14)(p21;q32)
- Centromere 12, DLEU (13q14), LAMP1 (13q34) (trisomy 12, deletion 13q14, deletion 13q)
- IGH t(14q32)
- IGH::BCL2 t(14;18)(q32;q21)/gain 18q
- MALT1 t(18q21)
- MYB (6q23) with controls in 6q21(SEC63) and D6Z1
- MYC t(8q24)/gain 8q24

T-Cell-Lymphoma (T-NHL)

- ATM (11q22.3) and TP53 (17p13.1) (deletion 11q, deletion 17p)
- ETV6::RUNX1 (deletion 12p)
- MYC t(8q24)/gain 8q24
- TCR α/δ t(14q11)

Plasma Cell Myeloma/ Multiple Myeloma/MGUS

For FISH analysis, "magnetic activated cell sorting" (MACS) will be initially performed to isolate CD138+ plasma cells

- Complete plasma cell disease panel (deletion 1p32.3, amplification 1q21, deletion/gain 11q22, trisomy 12, deletion 13q14, deletion 13q34, monosomy 13, IGH-rearrangement t(14q32), deletion 17p13)
- Aneuploidies of the chromosomes 5 (5p15), 9 (9q22), 15 (15q22)
- ATM (11q22) and TP53 (17p13) (deletion/gain 11q, deletion 17p)
- CKS1B (1q21), CDKN2C (1p32.2) (deletion 1p/amplification 1q)
- CCND1::IGH t(11;14)(q13;q32)
- CCND3::IGH t(6;14)(p21;q32)
- Centromere 12, DLEU (13q14), LAMP1 (13q34) (deletion 13q/monosomy 13)

- FGFR3::IGH t(4;14)(p16;q32)
- IGH t(14q32)
- IGH::MAFA t(14;16)(q32;q23)
- IGH::MAFB t(14;20)(q32;q12)
- MYC::IGH t(8;14)(q24;q32)

Acute Lymphoblastic Leukemia/B-Cell Acute Lymphoblastic Leukemia (B-ALL)

- Aneuploidies of the chromosomes 4 (D4Z1), 10 (D10Z1), 17 (D17Z1)
- BCR::ABL1 t(9;22)(q34;q11.2)
- CDKN2A (p16, 9p21) with control CEP9 (deletion 9p)
- ETV6::RUNX1 t(12;21)(p13;q22)
- ETV6::RUNX1 (deletion 12p/amplification RUNX1)
- IGH t(14q32)
- KMT2A(MLL) t(11q23)
- KMT2A(MLL)::AFF1(AF4) t(4;11)(q21.3-q22.1;q23.3)
- KMT2A(MLL)::MLLT1(ENL) t(11;19)(q23.3;p13.3)
- KMT2A(MLL)::AFDN(MLLT4) t(6;11)(q27;q23.3)
- MYC t(8q24)
- MYC::IGH t(8;14)(q24;q32)
- TCF3::PBX1 t(1;19)(q23;p13.3) with HLF (17q22)

T-Cell Acute Lymphoblastic Leukemia (T-ALL)

- BCR::ABL1 t(9;22)(q34;q11.2)
- CDKN2A (p16, 9p21) with control CEP9 (deletion 9p)
- KMT2A(MLL) t(11q23)
- TCR α/δ t(14q11)

Severe Aplastic Anemia (AA)

- Centromere 12, DLEU (13q14), LAMP1 (13q34) (trisomy 12, deletion 13q14, deletion 13q)
- D20S108 (20q12), RH7808 (20q13.3), D8Z2 (deletion 20q, trisomy 8)
- EGR1 (5q31), RPS14 (5q33) with control in 5p15 (deletion 5q/monosomy 5)
- KMT2E (7q22), D7S522/MET (7q31.2) with control D7Z1 (7q deletion/monosomy 7)

Chromosome X- and Y-specific FISH after BMT

- Chromosome X- and Y-specific probes (DXZ1, DYZ3)

Other FISH probes (on request)
