

**Diagnostic Laboratories Department of Pediatrics
and Adolescent Medicine**

Medical Director: Prof. Dr. M. Erlacher

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Molecular Diagnostics Laboratory

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Germany

Informed consent of patient

The German law (**Gendiagnostikgesetz – GenDG**) defines under which circumstances genetic testing of a human individual is legal. *Diagnostic* testing can only be conducted with the patient's informed consent which requires documented consultation with a doctor. *Predictive* testing requires genetic counselling by a human genetics specialist prior to and after the investigation, or the patient's written renunciation.

**Please fill out the informed consent carefully and completely and sign it.
Otherwise no molecular genetic testing can be performed!**



Molecular Genetic Testing

3-5 ml EDTA-blood (send by normal mail)

Turnaround time: 2-4 weeks, more than one analysis and rare analyses may take up to 8 weeks

OMIM	Disease	Gene
Immunology		
Immunodeficiency SCID (T-B+)		
<input type="checkbox"/> OMIM 300400	Severe Combined Immunodeficiency (SCID, X-chromosomal, T-, B+, NK-)	IL2RG
<input type="checkbox"/> OMIM 606367	Severe Combined Immunodeficiency, CD25 deficiency	IL2RA
<input type="checkbox"/> OMIM 600802	Severe Combined Immunodeficiency (SCID, T-, B+, NK-), JAK3 deficiency	JAK3
<input type="checkbox"/> OMIM 608971	Severe Combined Immunodeficiency (SCID, T-B+ NK+), IL7R deficiency	IL7R
<input type="checkbox"/> OMIM 608971	Severe Combined Immunodeficiency (SCID, T-B+ NK+), CD3D deficiency	CD3D
<input type="checkbox"/> OMIM 608971	Severe Combined Immunodeficiency (SCID, T-B+ NK+), CD3E deficiency	CD3E
Other defined immune disorders		
<input type="checkbox"/> OMIM 109535	Immunodeficiency - Hyper-IgM-Syndrome (HIGM 3)	CD40
<input type="checkbox"/> OMIM 300386	Hyper-IgM-Syndrome, X-linked (HIGM 1), (SCID, T- B-)	CD40L
<input type="checkbox"/> OMIM 176947	Severe Combined Immunodeficiency (SCID, T-, B-), ZAP70 deficiency, CD8 deficiency	ZAP70
<input type="checkbox"/> OMIM 301000	Wiskott-Aldrich Syndrome	WASP
<input type="checkbox"/> OMIM 603553	familial hemophagocytic lymphohistiocytosis (FHL2; HLH2)	PRF1
<input type="checkbox"/> OMIM 240300	autoimmune polyendocrinopathy-candidiasis-ectodermal dystrophy (APS1), APECED	AIRE
<input type="checkbox"/> OMIM 304790	immune dysregulation, polyendocrinopathy, enteropathy, X-linked (IPEX)	FOXP3
<input type="checkbox"/> OMIM 209950	familial disseminated atypical mycobacterial infection and disseminated BCG infection (FDAMI)	IFNGR1
<input type="checkbox"/> OMIM 601859	Autoimmune lymphoproliferative syndrome (ALPS1a), Canale-Smith-Syndrome	APO1
<input type="checkbox"/> OMIM 601859	Autoimmune lymphoproliferative syndrome (ALPS1b), Canale-Smith-Syndrome	FASL
<input type="checkbox"/> OMIM 603909	Autoimmune lymphoproliferative syndrome (ALPS2a), Canale-Smith-Syndrome	CASP10
<input type="checkbox"/> OMIM 607271	Autoimmune lymphoproliferative syndrome (ALPS2b), Canale-Smith-Syndrome	CASP8
<input type="checkbox"/> OMIM 164790	Autoimmune lymphoproliferative syndrome (ALPS4); N-RAS exon 2	N-Ras
<input type="checkbox"/> OMIM 308240	X-linked lymphoproliferative syndrome (XLP), Purtillo-Syndrome (SH2D1A)	SAP
<input type="checkbox"/> OMIM 613011	EBV-associated lymphoproliferative diseases, autosomal-recessive; IL2-inducible T-cell kinase, ITK deficiency	ITK
<input type="checkbox"/> OMIM 300079	X-linked lymphoproliferative syndrome (XLP2), BIRC4 deficiency	BIRC4
<input type="checkbox"/> OMIM 154545	infections, recurrent, with chronic diarrhea and opsonisation defect (low MBL2), including susceptibility to meningococcal disease, MBL2 deficiency; polymorphisms in promotor and exon 1: -550 C/G, +221 C/G; p.T24A, p.R52C, p.G54D >	MBL2
<input type="checkbox"/> OMIM 123890	Graves disease, susceptibility locus 5 (GDSS), T-cell defect	CTLA4
<input type="checkbox"/> OMIM 305000	Dyskeratosis congenita; X-linked; Codon: p.A353V	DKC1
<input type="checkbox"/> OMIM 305000	Dyskeratosis congenita-1; X-linked	DKC1
<input type="checkbox"/> OMIM 266100	epilepsy, pyridoxine-dependent (EPD); pyridoxine dependency with seizures, ALDH7A1 deficiency	ALDH7A1
<input type="checkbox"/> OMIM 606609	Aicardi-Goutières Syndrom AGS, chilblain lupus CHBL, hereditary endotheliopathy with retinopathy, nephropathy and	TREX1
<input type="checkbox"/> OMIM 300645	chronic granulomatous disease 1 (CGD1); X-linked granulomatous disease; cytochrome b-245, beta polypeptide	CYBB
<input type="checkbox"/> OMIM 608508	CGD; Chronic Granulomatosis; Cytochrome b-245 Alpha Kette - CYBA	CYBA
<input type="checkbox"/> OMIM 612301	CGD; Chronic Granulomatosis; neutrophile cytosolic factor 2 - NCF2	NCF2
<input type="checkbox"/> OMIM 102582	Hyper-IgE-Syndrome, autosomal-dominant (exons 13, 21 und 23; p.K392R; p.N646K; p.K658N; p.T715M)	STAT3
<input checked="" type="checkbox"/> OMIM 300248	*incontinentia pigmenti, hypohidrotic ectodermal dysplasia, (EDA-ID); X-chrom. recessive (NEMO); not: Incontinentia Pigmenti -del. Ex 4-10!	IKBKG
Osteopetrosis		
* not accredited		
<input type="checkbox"/> OMIM 602727	osteopetrosis, Albers-Schonberg disease 4 (OPTB4), recessive	CLCN7
<input type="checkbox"/> OMIM 602727	osteopetrosis, type II (OPTA2), autosomal dominant, ADOII, Albers-Schoenberg disease	CLCN7
<input type="checkbox"/> OMIM 604592	osteopetrosis, lethal B1 (OPTB1)	TCIRG1
<input type="checkbox"/> OMIM 607649	osteopetrosis type 1B 5 (OPTB5), malignant infantile	OSTM1
<input type="checkbox"/> OMIM 602642	osteopetrosis type B2 (POTB2), TNFSF11 deficiency	RANKL
<input type="checkbox"/> OMIM 612301	Osteopetrosis, (TNFRSF11A), Hypogammaglobulinämia	RANK
<input type="checkbox"/> OMIM 612301	Osteopetrosis, (SNX10), Typ B7	SNX10
Periodic Fever Syndromes		
<input type="checkbox"/> OMIM 249100	Familial Mediterranean fever (FMF)	MEFV
<input type="checkbox"/> OMIM 142680	TNF receptor associated periodic syndrome, benign autosomal dominant familial periodic fever (FPF), Hibernian fever	TRAPS
<input type="checkbox"/> OMIM 260920	Hyperimmunoglobulinemia D with recurrent fever (HIDS), mevalonate kinase deficiency	MVK
Hematology		
<input type="checkbox"/> OMIM 300367	dyserythropoietic anemia with thrombocytopenia (X-linked), CDATX	GATA-1
<input type="checkbox"/> OMIM 603474	Diamond-Blackfan anemia (DBA)	RPS19
<input type="checkbox"/> OMIM 603634	Diamond-Blackfan anemia (DBA5)	RPL5
<input type="checkbox"/> OMIM 611184	Dehydrated hereditary Stomatocytosis 1 (DHD1); Hereditary Xerozytosis	PIEZ01
<input type="checkbox"/> OMIM 602754	Dehydrated hereditäre Stomatocytosis 2 (DHD2); Hereditary Xerozytosis	KCNN4
<input type="checkbox"/> OMIM 159530	congenital amegakaryocytic thrombocytopenia (CAMT)	MPL
<input type="checkbox"/> OMIM 600044	thrombocythemia, essential (THPO)	THPO



Request for Molecular Genetic Testing

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OMIM Disease

Hematology

<input type="checkbox"/> OMIM 137295	*MonoMAC Syndrome; acute myeloid Leukemia, AML	GATA2
<input type="checkbox"/> OMIM 615888	*LADII leukocyte adhesion deficiency	RASGRP2
<input type="checkbox"/> OMIM 206200	iron-refractory iron deficiency anemia (IRIDA)	TMPRSS6
<input type="checkbox"/> OMIM 600424	disturbance in MTX clearance, high MTX levels	SLC19A1/GGH
<input type="checkbox"/> OMIM 249270	thiamine responsive megaloblastic anemia (TRMA)	SLC19A2
<input type="checkbox"/> OMIM 600523	pseudo-iron deficiency anemia (PIDA), microcytic anemia with liver iron overload , DMT1 deficiency	SLC11A2/DMT1
<input type="checkbox"/> OMIM147700	isocitrate dehydrogenase 1 (NADP+), soluble, associated with brain tumors and secondary Glioblastoma	IDH1
<input type="checkbox"/> OMIM 613657	isocitrate dehydrogenase 2 (NADP+), mitochondrial; D-2-hydroxyglutaric aciduria (D2HA2)	IDH2
<input type="checkbox"/> OMIM 613673	congenital dyserythropoietic anemia, TYPE IV; CDAN4	KLF1

*not akkreditated

Endocrinology and Diabetology

Disturbance in glucose regulation

<input type="checkbox"/> OMIM 125850	Maturity-Onset Diabetes of the Young, MODY-Diabetes, MODY type 1 (MODY1)	HNF4a
<input type="checkbox"/> OMIM 125851	Maturity-Onset Diabetes of the Young, MODY-Diabetes, MODY type 2 (MODY2)	GCK
<input type="checkbox"/> OMIM 600496	Maturity-Onset Diabetes of the Young, MODY-Diabetes, MODY type 3 (MODY3)	HNF1a/TCF1

Disturbance in body weight regulation

<input type="checkbox"/> OMIM 155541	severe obesity (OBS3)	MC4R
<input type="checkbox"/> OMIM 164160	severe early-onset obesity (OBS1)	LEP
<input type="checkbox"/> OMIM 601007	*early-onset morbid obesity (only after having consulted Prof. Wabitsch)	LEPR
<input type="checkbox"/> OMIM 601047	Berardinelli-Seip congenital generalized lipodystrophy type (BSCL3)	CAV1
<input type="checkbox"/> OMIM 613327	congenital generalized lipodystrophy type 4 (CGL4)	PTRF

Disturbance of growth

<input type="checkbox"/> OMIM 117550	Sotos syndrome (STO)	NSD1
<input type="checkbox"/> OMIM 610978	congenital thyroglobulin defect with respiratory distress (TGDRD)	NKX2-1
<input type="checkbox"/> OMIM 118700	benign hereditary chorea (BHC)	NKX2-1

Polymorphisms (possibly clinical relevant)

<input type="checkbox"/> OMIM 174800	McCune-Albright Syndrome (MAS); polymorphisms in exons 8 & 9: p.R201C,p. R201H, p.R201G, p.Q227H, p.Q227R	GNAS
<input type="checkbox"/> OMIM 167413	*Diabetes; polymorphism in exon 9: rs1712701(c.962A>C; p.P321H)	PAX4
<input type="checkbox"/> OMIM 173360	*Obesity; polymorphism in promotor: rs1799768 (NG_013213.1:g.4328_4329insC; -675_4G/5G)	PAI1
<input type="checkbox"/> OMIM 601487	*Obesity and thromboembolism; polymorphism in exon 1: rs1801282 (c.34C>G; p.Pro12Ala)	PPARG2
<input type="checkbox"/> OMIM 151750	*Obesity; polymorphism in IVS6 : rs57282318 (rs71167395; NM_005357.3:c.2365+142_2365+145delGTGT)	HSL
<input type="checkbox"/> OMIM 170290	*Obesity; polymorphism in IVS6 : rs894160 (NM_002666.3:c.772-799A>G)	PLIN
<input type="checkbox"/> OMIM 190220	*Obesity and Diabetes; polymorphism in exon 1: rs1800470 (c.17T>C; p.Leu10Pro)	TGFB1
<input type="checkbox"/> OMIM 601283	*Obesity; Diabetes Mellitus, Noninsulin-dependent, 1 (NIDDM1); intronic polymorphisms: IVS6; rs3842570 (NM_023083.4:c.997+116Indel), IVS3 rs3792267 (NM_021251.3:c.141+4705G>A) und IVS1 3rs5030952 (NM_023089.1:c.274-13102G>C)	CAPN10
<input type="checkbox"/> OMIM 605441	*Obesity, Adipocyte, C1q, and Collagen Domain containing,(ADIPOQ); polymorphism in exon 9: rs2241766 (NM_004797.2:c.45T>G; p.Gly15Gly)	APM1
<input type="checkbox"/> OMIM 601693	*Obesity; polymorphism in promoter: rs659366 (NG_011478.1:g.4136G>A)	UCP-2
<input type="checkbox"/> OMIM 600716	*Diabetes (type 1); polymorphism in exon 9: rs2476601 (NM_012411.2; c.1858C>TArg620Trp)	PTPN22

Gastroenterology

<input type="checkbox"/> OMIM 608374	hemochromatosis, type 2, Hämocherulin (HVJ); HFE2	HVJ
<input type="checkbox"/> OMIM 604250	hemochromatosis, type 3 ; transferrin receptor 2, TFR2	HFE3/TFR2
<input type="checkbox"/> OMIM 606069	hemochromatosis, type 4 ; solute carrier family 40 (iron-regulated transporter), member 1, FPN1 / SLC40A1	HFE4/FPN1
<input type="checkbox"/> OMIM 602390	hemochromatosis, juvenile, (JH), (HFE2B)	HAMP
<input type="checkbox"/> OMIM 143500	Gilbert-Syndrom (GBTs)	UGT1A1
<input type="checkbox"/> OMIM 146933	chronic inflammatory bowel disease, interleukin 10 receptor, alpha; IL10RA	IL10Ra
<input type="checkbox"/> OMIM 123889	chronic inflammatory bowel disease, interleukin 10 receptor, beta; IL10RB	IL10Rb
<input type="checkbox"/> OMIM 124092	chronic inflammatory bowel disease, interleukin 10; early onset	IL10
<input type="checkbox"/> OMIM 610370	congenital malabsorptive diarrhea 4 (DIAR4); T2DM; hyperproinsulinaemia	NEUROG3

Polymorphisms (possibly clinical relevant)

<input type="checkbox"/> OMIM 266600	chronic inflammatory bowel disease, 1 (IBD1), Morbus Crohn	NOD2
<input type="checkbox"/> OMIM 605956	Hepatic veno-occlusive disease or veno-occlusive disease (VOD)	NOD2
<input type="checkbox"/> OMIM 235200	hemochromatosis (HFE1); polymorphism p.H63D, p.S65C und p.C282Y	HFE
<input type="checkbox"/> OMIM 235200	hemochromatosis (HFE1)	HFE
<input type="checkbox"/> OMIM 613282	*nonalkoholic fatty liver disease : polymorphism in exon3 and 9: rs738409 (c.444C>G;p.I148M) & rs6006460 (c.1358G>T;p.S453I)	PNPLA3
<input type="checkbox"/> OMIM 107720	*Insulin resistance in nonalcoholic fatty liver disease (NG_008949.1:g.4546C>T (c.-482C>T; c.-455T>C)	APOC3
<input type="checkbox"/> OMIM 156845	*Waardenburg syndrome, type 2A", WS2, WS2A; polymorphism in exon 9: rs149617956 (p.E318K; p.Glu318Lys)	MITF

Information about the indication of molecular diagnosis can be achieved via "The portal for rare diseases and orphan drugs" - [orphanet](http://www.orpha.net/consor/cgi-bin/index.php) - : <http://www.orpha.net/consor/cgi-bin/index.php> or http://www.orpha.net/consor/cgi-bin/Disease_Genes.php?lng=EN

External quality assurance (EQA):

our laboratory participates regularly on „Ringversuche“ of the national Reference Institute for Bioanalytics (RfB). These RfB-laboratories determine method-independent target values, so called reference methods values, in control materials for external quality control. In addition we routinely perform laboratory exchanges.

