General Information

Patient		
Surname:		
First name:		
Date of birth:		
Sex:	male female	
Material		
Blood ml (min. 1-2 ml EDTA-blood)	
Dried blood spo	t cards (at least 5 spots)	
□ DNA µg (min. 1-2 µg DNA, concentr. ≥ 50 ng/µl) DNA-No.:		
Source material		
of extracted DNA:	(e.g. EDTA blood, skin biopsy)	
Other specimer	·	
External ID:		
Date of sample col	lection:	
Samples can be sent by mail in a cardboard box or air cushion envelope. Samples should not be exposed to direct sunlight. Dried blood spot cards can be ordered for free (info@cegat.com).		

Declaration of consent

By signing this form, I declare that I have received comprehensive information regarding the genetic background related to the disease in question, as well as the possibilities and limitations of molecular genetic testing. I understand that I have the right to withdraw my consent for genetic analyses.

I have been informed, and agree, that my personal data and the data obtained in the analysis will be recorded, evaluated or stored in an pseudonymized form in scientific databases, and that further, in accordance with data protection and medical confidentiality, the request, or parts thereof, may be transmitted to a specialized cooperating laboratory.

I consent to the re-evaluation of my test results within the data storage period. If significant alterations become apparent, my Physician will be informed by e-mail.

I consent that in addition to the full genetic test as requested, the analysis can be expanded to all pathogenic and likely pathogenic variants (ACMG class 4 and 5) in genes which are related to the indication described for the proband (if applicable, screen for differential diagnosis).

I have been informed, and agree to the electronic storage, processing, use, and transmission of all data collected by CeGaT GmbH.

For more detailed information on data privacy as well as your rights please refer to www.cegat.de/en/privacy-policy

Please Note

Our panels are regularly updated to reflect current scientific research. It should therefore be recognized that there is the possibility that the list of genes on the order form may have changed slightly (genes added or removed) by the time the sample is analyzed in the laboratory. By signing this form, the patient accepts that the list of genes actually analyzed may be slightly different from what is currently listed. When NGS is utilized more than the requested genes are sequenced for each sample.

This declaration of consent can be completely or partially withdrawn at any time. I have had sufficient time to consider giving my consent.

I, the referring physician, confirm that I am authorized to request genetic testing for the above-mentioned patient. For predictive testing, I confirm that I am authorized, and that I have fulfilled the requirements, to request this testing. For minors, I declare that I have the consent of all legal guardians.

If the patient did not sign this order form: I, the referring physician, confirm that the patient received genetic counseling and agrees with the genetic testing. The patient's consent has been obtained in writing.

Sender / Clinic		
Surname:		
First name:		
Institution:		
Street:		
Postcode/City:		
Country:		
Phone:		
Email:		
VAT: If applicable, please includ	e a VAT number or a copy of your business registration certificate.	
Invoice	 to sender / clinic to patient / other (KVA-No.:) 	
Surname:		
First name:		
Street:		
Postcode/City:		
Country:		
Email:		

If you do not check these boxes, your answer will be recorded as "No".

With regard to secondary findings I would like to be informed:	Yes	🛛 No
I consent to the pseudonymous storage and use of surplus genetic material and/or test results for scientific research and in scientific literature.	Yes	🛛 No
I consent to the storage of my test results beyond the timespan of 10 years (as required by German law).	Yes	🛛 No
I consent to the storage of my genetic material for additional tests and/or quality control (for max. 10 years).	Yes	🛛 No

Genetic variation may sometimes be identified, which does not fit within the scope of the requested genetic analysis (so-called secondary findings). The reporting of these variants is limited to pathogenic alterations (ACMG classes 4 and 5) within selected genes, for which a treatment or course of action exists for you or your family (according to the current guidelines of the American College of Medical Genetics and Genomics; details on genes and associated diseases can be found at https://www.cegat.com/acmg-genes/). There is no claim of a comprehensive analysis of this gene set. An absence of secondary findings cannot be used to indicate a reduced disease risk.

Targeted analysis of the ACMG genes according to current recommendations can be requested as "additional analyses".

According to German Genetic Diagnostic Act (GenDG) we will issue the medical report to the counselling physician. Please indicate here the contact email of the counselling physician:

Email:

Physician's stamp / Barcode



CLIA CERTIFIED ID: 9902130225 CeGaT is accredited by DAKS according to DIN EN ISO 15189:2014, the College of American Pathologists (CAP) and CLIA.

Patient / Legal Guardian (Date, Signature)

Patient / Legal Guardian

(Block letters)



Physician

(Date, Signature)

(Surname, First name)

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Indication

Analysis type:	Proband is affected	□ Proband is NOT affected (predictive testing)	
Indication / Suspected diagnosis:			
Major Clinical Symptoms:			
Preliminary genetic diagnostics:			
Transplants (bone marrow, tissue	ə, stem cells) 🛛 No	□ Yes, (please specify)	
Please include a copy of all exist	ing reports of your patien	ıt.	
Pedigree	Consanguinity: 🛛 Yes	No Ethnic origin:	
			○ □ not affected
			affected
			• known carrier
			U deceased
			$\Box_{\overline{1}}$ consanguine parents
			ightarrow unborn child
			abortion, stillborn child
			person of unknown sex
			identical twins (monozygous)
			fraternal twins (dizygous)
Family medical history Are there other family members wh Yes No	o currently have or have ha	ad the same or a similar disease as the patient?	

If yes, please list the affected family members:

Name (not required)	Relationship to the patient (e.g. mother)	Age of onset	Diagnosis / Symptoms

Indication & Clinical Information



Previous genetic diagnostics:			
	Chromosome analysis / Array-CGH:		
	Sequence analysis of the following genes		
Previous operations:	Hematopoietic stem cell transplantation*		
	Splenectomy		
*if no DNA or blood cells have been collected prior to tr	ansplantation, a sample from non-hematopoietic tissue is required.		
Clinical information			
Please provide the following clinical information tifying causative alterations during NGS analysi	for your patient. Detailed clinical information – also on the absence of phenotypic findings - will increase the likelihood of iden- is and significantly improve the interpretation of identified variants. Please specify if applicable:		
General symptoms			
Age at disease onset:			
For newborns / infants: Prenatal medical history:			
	□ preterm-birth:WG		
	Other information on the course of pregnancy (e. g. medication):		
	u with elevated inflammation markers u infection-associated		
	u without further abnormalities		
	accompanied by further symptoms, namely:		
Autoimmunity / Autoinflammation:			
Susceptibility to infection:			
(e.g. skin or lung alterations)			
Miscellaneous: (e.g., skin or lung alterations)			
Previous and current therapy:			

Indication & Inquiry



Blood values & immunological parameters

In case of abnormal test results please specify altered parameters, optionally using symbols such as \uparrow elevated $\uparrow\uparrow$ strongly elevated \downarrow decreased $\downarrow\downarrow$ strongly decreased

Inflammation markers:	normal	D abnormal:	
Immunoglobulins:	normal	abnormal:	
Pathogen detection:	not analyzed	□ none detected	
	detection of the foll	owing pathogens:	
SCID newborn screening (TREC level)	normal	abnormal:	
Hematology:	normal	Iymphopenia	
	thrombocytopenia	anemia Ineutropenia	
Immune phenotyping:	normal	abnormal:	
Autoantibodies:	□ none	□ yes, the following:	
Oxidative burst (DHR assay):	normal	abnormal:	
Complement activity (CH50, AP50):	normal	abnormal:	
Enzyme activities: (e. g. ADA, PNP)	normal	abnormal:	
Other clinical chemistry	normal	abnormal:	
Other specialized (immune) diagnostics: (e. g. lymphocyte function, telomere length, chromosome breakage)			
Physical appearance / other abnormalities			
□ <u>no</u> abnormalities of the physical appearance			
□ Signs of (partial) albinism:			
Facial dysmorphism:			
□ Skeletal anomalies:			
Developmental delay:			
Mental retardation:			
Other abnormalities:			



Indication & Inquiry

Inquiry Immune Disorders

□ Primary antibody deficiencies (40 Genes, PID01)

(incl. Hyper IgM syndrome, CVID, agammaglobulinemia, activated PI3Kdelta syndromes)

ADA2, AICDA, ARHGEF1, ATM, ATP6AP1, BLNK, BTK, CD19, CD40, CD40LG, CD79A, CD79B, CD81, CR2, CXCR4, FNIP1, ICOS, ICOSLG, IGLL1, IKBKB, IKZF1, IRF2BP2, LRBA, MS4A1, NFKB1, NFKB2, NFKBIA, PIK3CD, PIK3R1, PLCG2, PTEN, RAC2, SEC61A1, SH3KBP1, SLC39A7, TCF3, TNFRSF13B, TOP2B, UNG, VAV1

Severe combined immunodeficiencies (SCID) (35 Genes, PID02) (incl. newborn SCID-screening (TREC abnormal))

ADA, AK2, ATM, BCL11B, CD247, CD3D, CD3E, CD3G, CHD7, DCLRE1C, FOXI3, FOXN1, IL2RG, IL7R, JAK3, LAT, LCP2, LIG1, LIG4, MSN, MTHFD1, MTR, NHEJ1, PAX1, PGM3, PNP, PRKDC, PTPRC, RAC2, RAG1, RAG2, RMRP, RPSA, SLC46A1, TBX1

 Combined immunodeficiencies and other T-cell defects (40 Genes, PID03)

B2M, CARD11, CARMIL2, CD27, CD3E, CD3G, CD8A, CIITA, COPG1, CTPS1, DOCK2, FCHO1, ICOS, IKBKB, IL21R, IL2RA, IL2RB, IL2RG, LAT, LCK, LCP2, MAGT1, MALT1, MAP3K14, MSN, ORAI1, PIK3CD, RASGRP1, RELB, RFX5, RFXANK, RFXAP, RIPK1, STIM1, STK4, TAP1, TAP2, TAPBP, TFRC, ZAP70

- □ Hyper-IgE syndrome and differential diagnoses (13 Genes, PID04) ARPC1B, CARD11, DOCK8, DSG1, ERBIN, FOXP3, IL6ST, PGM3, SPINK5, STAT3, STAT5B, TYK2, ZNF341
- Syndromes with deficiencies of the adapative immunity
 (19 Genes, PID05)
 CDC47_CHD7_CHUK_DNMT38_EPG5_EOXI3_EOXN1_HEUS_P4

CDCA7, CHD7, CHUK, DNMT3B, EPG5, FOXI3, FOXN1, HELLS, PAX1, POLD1, POLD2, POLE, POLE2, SEMA3E, SMARCAL1, SP110, TBX1, TBX2, ZBTB24

□ Defects of the complement system (21 Genes, PID06) C1QA, C1QB, C1QC, C1R, C1S, C2, C3, C5, C6, C7, C8A, C8B, CFB, CFD, CFH, CFI, CFP, FCN3, MASP1, MASP2, MBL2

Neutropenia (26 Genes, PID07)

ADA2, CD40, CD40LG, CLPB, CSF3R, CXCR2, CXCR4, DNAJC21, EFL1, ELANE, G6PC3, GATA1, GATA2, GFI1, GINS1, HAX1, JAGN1, SBDS, SMARCD2, SRP54, TAZ, TCIRG1, USB1, VPS45, WAS, WIPF1

- Chronic granulomatous disease (CGD) and differential diagnoses (8 Genes, PID08) CYBA, CYBB, CYBC1, G6PD, MPO, NCF1 (c.75_76delGT), NCF2, NCF4
- □ Other deficiencies of the phagocytes (16 Genes, PID09) CEBPE, CFTR, CLPB, CXCR2, CXCR4, FERMT3, ITGB2, LAMTOR2, MRTFA, RAC2, RMRP, SLC35C1, SMARCD2, TAZ, VPS13B, WDR1
- □ Chronic mucocutaneous candidiasis and susceptibility to other fungal infections (13 Genes, PID10) AIRE, CARD9, CLEC7A, IL12B, IL12RB1, IL17F, IL17RA, IL17RC, MAPK8, RORC, STAT1, STAT3, TRAF3IP2
- □ Susceptibility to mycobacterial infections (18 Genes, PID11) CYBB, GATA2, IFNG, IFNGR1, IFNGR2, IL12B, IL12RB1, IL12RB2, IL23R, IRAK4, IRF8, ISG15, JAK1, RORC, SPPL2A, STAT1, TBX21, TYK2
- Susceptibility to viral infections (29 Genes, PID12)

(incl. Herpes simplex and VZV encephalitis) CXCR4, DBR1, GATA2, GINS1, IFNAR1, IFNAR2, IRF3, IRF7, IRF8, IRF9, MCM10, MCM4, NOS2, PIK3CD, POLR3A, POLR3C, POLR3E, POLR3F, RANBP2, RTEL1, SNORA31, STAT1, STAT2, TBK1, TICAM1, TLR3, TRAF3, TYK2, UNC93B1

- Generalized verrucosis (13 Genes, PID13) CARMIL2, CD4, CIB1, CXCR4, DOCK8, GATA2, IL7, NFKBIA, RHOH, STK4, TAOK2, TMC6, TMC8
- □ Defects of the TLR signaling pathway (4 Genes, PID14) IRAK4, MYD88, TICAM1, TLR4
- □ Defects of the NFkB signaling pathway (15 Genes, PID15) BCL10, CARD11, IKBKB, MALT1, MAP3K14, NFKB2, NFKBIA, RBCK1, REL, RELA, RELB, RIPK1, RNF31, TICAM1, TRAF3
- □ Defects of the type I interferon signaling pathway (13 Genes, PID16)

IFIH1, ÍFNAR1, IFNAR2, IRF7, ISG15, JAK1, STAT1, STAT2, STING1, TICAM1, TLR3, TRAF3, TYK2

Indication & Inquiry



Periodic fever syndromes with/without urticaria (14 Genes, AID01)

F12 (c.859T>A), HTR1A, MEFV, MVK, NLRC4, NLRP12, NLRP3, NTRK1, OTULIN, PLCG2, RIPK1, SLC29A3, TNFRSF1A, WDR1

- □ Inflammation with cardinal symptoms in the connective and supporting tissues (25 Genes, AID02) ADA2, ADAM17, AP1S3, ARPC1B, CARD11, CARD14, CCN6, HAVCR2, IL1RN, IL36RN, LACC1, LPIN2, NFKB1, NLRP1, NOD2, OTULIN, POMP, PSMA3, PSMB4, PSMB8, PSMB9, PSTPIP1, STING1, TNFAIP3, UBA1
- □ Immune dysregulation with colitis, very-early onset (33 Genes, AID03)

ADAM17, BACH2, CARMIL2, CD55, CTLA4, CYBA, CYBB, EGFR, EPCAM, FOXP3, GUCY2C, HSPA1L, IL10, IL10RA, IL10RB, IL21, IL21R, IL2RB, LRBA, NCF1 (c.75_76deIGT), NCF2, NCF4, NFKB1, NLRC4, PLVAP, RIPK1, SKIV2L, STAT3, TGFB1, TTC37, TTC7A, XIAP, ZBTB24

- Hemophagocytic lymphohistiocytosis (HLH) (21 Genes, AID04) AP3B1, CD27, CD48, GATA2, HAVCR2, ITK, LIPA, LYST, MAGT1, NCKAP1L, NLRC4, PIK3CG, PRF1, RAB27A, RC3H1, SH2D1A, SLC7A7, STX11, STXBP2, UNC13D, XIAP
- □ Abnormal Lymphoproliferation (33 Genes, AID05)

incl. autoimmune-lymphoproliferative syndrome (ALPS)

CARD11, CASP10, CASP8, CD27, CD70, CDC42, CTLA4, DEF6, FADD, FAS, FASLG, IL2RA, IL2RB, ITK, KRAS, LRBA, MAGT1, NCKAP1L, NEIL3, NFKB1, NRAS, PIK3CD, PIK3R1, PRKCD, RASGRP1, RELA, SH2D1A, SOCS1, STAT1, STAT3, STK4, TNFRSF9, XIAP

Additional analyses (additional fees may apply)

□ HLA-Typing (HLA01)

I would like to receive an additional report stating the HLA alleles (HLA class I (Gene A, B, C) and HLA class II (Gene DPA1, DPB1, DQA1, DQB1, DRB1, DRB3, DRB4, DRB5)).

□ ACMG genes diagnostics

I would like to be informed of relevant alterations within the list of recommended genes for secondary analysis, according to the current guidelines of the American College of Medical Genetics and Genomics. The analysis is restricted to the sequence data, re-sequencing of regions with poor sequence coverage will not typically be performed. A negative "ACMG genes" report cannot be used to rule out (genetic) disease risk. Additional fees may apply. According to German legislation, predictive tests for minors may not be performed for diseases which have an onset in adulthood. Therefore, some genes will not be analyzed for minors, unless the phenotypic spectrum is within the scope of the primary medical indication of the patient. Details on genes and associated diseases can be found at https://www.cegat.com/acmg-genes/

□ Pharmacogenetics (PGX) (22 genes)

ABCG2, CACNA1S, CYP2B6, CYP2C19, CYP2C9, CYP2D6, CYP3A4, CYP3A5, CYP4F2, DPYD, G6PD, HLA-A, HLA-B, IFNL3, MT-RNR1, NUDT15, POR, RYR1, SLC01B1, TPMT, UGT1A1, VKORC1

I would like to receive an additional report analyzing known variants in 22 genes that are involved in the metabolism of pharmaceutical products.

For further information and advice please do not hesitate to contact our Diagnostic Support team. www.cegat.com/diagnostic-support · diagnostic-support@cegat.com · Phone +49707156544-55

 Defects of the regulatory T-cells and IPEX-like phenotypes (22 Genes, AID06)

BACH2, CARMIL2, CTLA4, DOCK8, FAS, FASLG, FOXP3, IL10, IL10RA, IL10RB, IL2RA, IL2RB, LRBA, MALT1, PIK3CD, PIK3R1, STAT1, STAT3, STAT5B, TGFB1, TTC37, TTC7A

 Type I Interferonopathies and differential diagnoses (27 Genes, AID07)

incl. Interferonopathies with leading neurological symptoms, CANDLE syndrome and chilblain lupus / juvenile systemic lupus erythematosus ADAR, C1QA, C1QB, C1QC, C1R, C1S, C2, C3, DNASE1, DNASE1L3, IFIH1, ISG15, POMP, PRKCD, PSMA3, PSMB10, PSMB4, PSMB8, PSMB9, RNASEH2A, RNASEH2B, RNASEH2C, RNU7-1, SAMHD1, STAT2, STING1, TREX1

□ Syndromes with immune dysregulation (12 Genes, AlD08) ADA2, AIRE, ARPC1B, C2orf69, CDC42, ITCH, NFKB1, RBCK1, RNF31, SLC29A3, STING1, TRNT1