General Information



Patient		Sender / Clinic				
Surname:		Surname:				
First name:		First name:				
Date of birth:		Institution:				
Sex: ☐ male ☐	female	Street:				
Material		Postcode/City:				
□ Blood ml (min. 1-2 ml EDTA-blood)		Country:				
☐ Dried blood spot cards (at least 5 spots)		Phone:				
□ DNA μg (min. 1-2 μg DNA, concentr. ≥ 50 ng/μl) DNA-No.:		Email:				
Source material		VAT:				
of extracted DNA:		If applicable, please include a VAT	number or a copy of your business regi	istration certific	cate.	
☐ Other specimen			sender / clinic patient / other (KVA-No.:		,	
External ID:		Surname:	Jalient / Other (KVA-NO)	
Date of sample collection:						
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).		First name:				
Declaration of consent		Street:				
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		Postcode/City:				
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		Country:				
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.		Email:				
I consent to the re-evaluation of my test results wi	ithin the data storage period. If significant alterations	If you do not check these b	oxes, your answer will be re	ecorded a	ıs "No"	
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		I consent to the storage of my genet quality control (for max. 10 years).		☐ Yes	□ N	
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 consent to the storage of my test res	sults beyond the timespan of 10 years			
I have been informed, and agree to the electronic storage, processing, use, and transmission of all data collected by CeGaT GmbH.		(as required by German law).		☐ Yes	□ N	
For more detailed information on data privacy as well as your rights please refer to www.cegat.de/en/privacy-policy		I consent to the pseudonymous storag and/or test results for scientific research		☐ Yes	□ N	
Please Note		With regard to secondary f	indings I would	□ Yes	□ No	
that there is the possibility that the list of genes of	scientific research. It should therefore be recognized on the order form may have changed slightly (genes	Genetic variation may sometimes be	identified, which does not fit within the		e requeste	
	zed in the laboratory. By signing this form, the patient may be slightly different from what is currently listed.	alterations (ACMG classes 4 and 5	genetic analysis (so-called secondary findings). The reporting of these variants is limited to pathogeni alterations (ACMG classes 4 and 5) within selected genes, for which a treatment or course caction exists for you or your family (according to the current guidelines of the American Colleg			
	completely or partially withdrawn at	of Medical Genetics and Genomics;	details on genes and associated dis There is no claim of a comprehensive	seases can b	e found a	
any time. I have had sufficient time			not be used to indicate a reduced disea			
	ed to request genetic testing for the above-mentioned uthorized, and that I have fulfilled the requirements, to ve the consent of all legal guardians.	dations can be requested a	CMG genes according to cu is "additional analyses".	arrent reco	ommen	
If the patient did not sign this order form: I, the	referring physician, confirm that the patient received testing. The patient's consent has been obtained in		nostic Act (GenDG) we will issue thate here the contact email of the coul			
•		Email:				
		Physician's stemp / Para	ando (
		Physician's stamp / Baro	ode (I	DAKKS Deutsch	he	
				Akkredi	itierungsstel .3206-01-00	
Patient / Legal Guardian	Physician			CAP		
(Block letters)	(Surname, First name)		COLLEG	CREDITED SE of AMERICAN PATH	HOLOGISTS	
Y	Y		CeGa	aT is accredite	ed by	
Patient / Legal Guardian	Physician		DIN E	kS according to EN ISO 15189 College of Ame	9:2014,	
(Date, Signature)	(Date, Signature)			ologists (CAP)		





Analysis type:	☐ Proband is affected	☐ Proband is	NOT affected (predic	ctive testing)	
Indication / Suspected diagnosis	s:				
Major Clinical Symptoms:					
major chilical symptoms.					
Preliminary genetic diagnostics	s:				
Transplants (bone marrow, tiss	ue, stem cells) 🛚 No	☐ Yes, (please	e specify)		
Please include a copy of all exi					
r rougo morado a copy or an oxi					
Pedigree	Consanguinity: Yes	☐ No Ethnic	origin:		
					1 index patient
					not affected
				_	affected known carrier
					deceased
					unrelated parents
					consanguine parents
					unborn child
				1	abortion, stillborn child
				\Diamond	person of unknown sex
				\triangle	identical twins) (monozygous)
				\wedge	fraternal twins) (dizygous)
Family medical history					
Are there other family members w	ho currently have or have h	ad the same or a	a similar disease as th	ne patient?	
☐ Yes ☐ No					
If yes, please list the affected fam	ily members: Relationship to	the patient	Age of onset	Diagnosis / Symp	otoms
(not required)	(e.g. moti		Ago of officer	Diagnosis i Oyin	





Inquiry

All genes listed on this order form, and the entire mitochondrial genome, are sequenced in parallel. This allows you to select multiple gene sets, or choose individual genes in addition to your selected gene set. As there is no additional laboratory expenditure, there is only a moderate price increase due to the subsequent analysis and interpretation. We are happy to answer your questions or send you an individual quote. Please contact us at info@cegat.com.

☐ Usher Syndrome (17 Genes, EYE01) ABHD12, ADGRV1, ARSG, CDH23, CEP250, CEP78, CIB2, CLRN1, ESPN,

HARS1, MYO7A, PCDH15, PDZD7, USH1C, USH1G, USH2A, WHRN

☐ Retinitis pigmentosa, autosomal dominant and X-linked (28 Genes, EYE02)

BEST1, CA4, CACNA1F, CRX, GUCA1B, HK1, IMPDH1, KLHL7, NR2E3, NRL, PRPF3, PRPF31, PRPF4, PRPF6, PRPF8, PRPH2, RDH12, RGR, RHO, ROM1, RP1, RP2, RP9, RPE65, RPGR (incl. ORF15), SEMA4A, SNRNP200, **TOPORS**

Retinitis pigmentosa, autosomal recessive and X-linked (65 Genes, EYE03)

ABCA4, AGBL5, AHI1, ARHGEF18, ARL2BP, ARL6, BBS1, BBS2, BEST1, C8orf37, CDHR1, CEP290, CERKL, CLN3, CNGA1, CNGB1, CRB1, CWC27, CYP4V2, DHDDS, DHX38, EYS, FAM161A, FLVCR1, GNAT1, GUCY2D, HGSNAT, IFT140, IFT172, IMPG2, KIAA1549, KIZ, LRAT, MAK, MERTK, MFRP, NR2E3, NRL, PCARE, PDE6A, PDE6B, PDE6G, POMGNT1, PRCD, PROM1, PRPF31, RBP3, RDH12, REEP6, RGR, RHO, RLBP1, ROM1, RP1, RP1L1, RP2, RPE65, RPGR (incl. ORF15), RPGRIP1, SAG, SLC7A14, SPATA7, TULP1, USH2A, ZNF408

☐ Achromatopsia (6 Genes, EYE04)

ATF6, CNGA3, CNGB3, GNAT2, PDE6C, PDE6H

☐ Bardet-Biedl Syndrome (32 Genes, EYE05)

ALMS1, ARL6, BBIP1, BBS1, BBS10, BBS12, BBS2, BBS4, BBS5, BBS7, BBS9, C8orf37, CCDC28B, CEP164, CEP19, CEP290, CEP41, IFT172, IFT27, IFT74, KIF7, LZTFL1, MKKS, MKS1, NPHP1, SDCCAG8, TMEM67, TRAPPC3, TRIM32, TTC21B, TTC8, WDPCP

□ Congenital Stationary Night Blindness (17 Genes, EYE06)

CABP4, CACNA1F, GNAT1, GNB3, GPR179, GRK1, GRM6, GUCY2D, LRIT3, NYX, PDE6B, RBP4, RDH5, RHO, SAG, SLC24A1, TRPM1

□ Joubert Syndrome (46 Genes, EYE07)

AHI1, ARL13B, ARL3, ARMC9, B9D1, B9D2, C2CD3, CC2D2A, CELSR2, CEP104, CEP120, CEP164, CEP290, CEP41, CPLANE1, CSPP1, EXOC8, FAM149B1, HYLS1, IFT74, IFT172, INPP5E, KIAA0556, KIAA0586, KIAA0753, KIF7, MKS1, NPHP1, OFD1, PDE6D, PIBF1, POC1B, RPGRIP1L, SUFU, TCTN1, TCTN2, TCTN3, TMEM107, TMEM138, TMEM216, TMEM218, TMEM231, TMEM237, TMEM67, TTC21B, ZNF423

☐ Leber Congenital Amaurosis (24 Genes, EYE08)

AIPL1, ALMS1, CEP290, CRB1, CRX, GUCY2D, IDH3A, IFT140, IMPDH1, IQCB1, KCNJ13, LCA5, LRAT, MERTK, NMNAT1, OTX2, PRPH2, RD3, RDH12, RPE65, RPGRIP1, SPATA7, TULP1, USP45

☐ Zellweger Syndrome Spectrum (Refsum/Zellweger/neonatale

adrenoleukodystrophy) (18 Genes, EYE10) ABCD1, ACOX1, HSD17B4, PEX1, PEX10, PEX11B, PEX12, PEX13, PEX14, PEX16, PEX19, PEX2, PEX26, PEX3, PEX5, PEX6, PEX7, PHYH

☐ Senior-Loken Syndrome (13 Genes, EYE11)

CEP164, CEP290, INVS, IQCB1, NPHP1, NPHP3, NPHP4, SCLT1, SDCCAG8, TMEM67, TRAF3IP1, WDR19, ZNF423

□ Stargardt Disease and Macular Dystrophies (23 Genes, EYE12)

ABCA4, BEST1, C1QTNF5, CDH3, CFH, CLN3, CNGB3, CRX, CTNNA1,

DRAM2, ELOVL4, GUCA1A, IMPG1, IMPG2, IRX1, MFSD8, PROM1, PRPH2, RDH12, RP1L1, RPGR (incl. ORF15), TIMP3, TTLL5

□ Cone and Cone Rod Dystrophies (43 Genes, EYE13) ABCA4, ADAM9, AIPL1, ALMS1, ATF6, C8orf37, CABP4, CACNA1F, CACNA2D4, CDHR1, CEP290, CEP78, CERKL, CFAP410, CNGA3, CNGB3, CNNM4, CRB1, CRX, DRAM2, GNAT2, GUCA1A, GUCY2D, KCNV2,

NMNAT1, PCARE, PCYT1A, PDE6C, PDE6H, PITPNM3, POC1B, PROM1, PRPH2, RAB28, RAX2, RDH12, RGS9, RGS9BP, RIMS1, RPGR (incl. ORF15), RPGRIP1, SEMA4A, TTLL5, TULP1

☐ Flecked Retina Disorders (16 Genes, EYE14)

ABCA4, CHM, CYP4V2, EFEMP1, ELOVL4, KCNJ13, OAT, PLA2G5, PROM1, PRPH2, RDH5, RHO, RLBP1, RPE65, RS1, VPS13B

□ Vitreoretinopathies (Familial exudative vitreoretinopathy/ Wagner Syndrome/Norrie syndrome/Knobloch syndrome) (17 Genes, EYE15)

ATOH7, BEST1, CAPN5, COL18A1, COL2A1, CTNNB1, FZD4, KCNJ13, KIF11, LRP5, NDP, NR2E3, P3H2, RCBTB1, TSPAN12, VCAN, ZNF408

☐ Stickler Syndrome (6 Genes, EYE16)

COL11A1, COL11A2, COL2A1, COL9A1, COL9A2, COL9A3

☐ Optic atrophy and Leber hereditary optic neuropathy (23 Genes,

ACO2, AFG3L2, ANTXR1, C12orf65, CISD2, DNM1L, FDXR, MCAT, MFN2, MT-ND1, m.3460G>A; MT-ND4, m.11778G>A; MT-ND6, m.14484T>C; NR2F1, OPA1, OPA3, RTN4IP1, SLC25A46, SPG7, SSBP1, TIMM8A, TMEM126A, WFS1, YME1L1

- □ Ocular and oculocutaneous albinism (8 Genes, EYE18) GPR143, LRMDA, MC1R, OCA2, SLC24A5, SLC45A2, TYR, TYRP1
- ☐ Syndromic albinism (Hermansky-Pudlak/Waardenburg/Vici/ Griscelli/Chediak-Higashi) (23 Genes, EYE19)

AP3B1, AP3D1, BLOC1S3, BLOC1S6, DTNBP1, EDN3, EDNRB, EPG5, HPS1, HPS3, HPS4, HPS5, HPS6, KIT, LYST, MITF, MLPH, MYO5A, PAX3, RAB27A, SNAI2, SOX10, TYR

☐ Ocular malformations (microphthalmia/anophthalmia/ nanophthalmia/coloboma) (31 Genes, EYE20)

ABCB6, ALDH1A3, ATOH7, BCOR, BMP4, CHD7, FOXE3, FREM1, GDF3, GDF6, HCCS, HMX1, MAB21L2, MFRP, OTX2, PAX2, PAX6, PIGL, PRSS56, RARB, RAX, RBP4, SHH, SIX6, SMOC1, SOX2, STRA6, TENM3, TMEM98, VAX1, VSX2

☐ Cataract (61 Genes, EYE21)

ABHD12, ADAMTSL4, AGK, BCOR, BFSP1, BFSP2, CHMP4B, CLPB, COL4A1, CRYAA, CRYAB, CRYBA1, CRYBA4, CRYBB1, CRYBB2, CRYBB3, CRYGC, CRYGD, CRYGS, CTDP1, CYP27A1, CYP51A1, EPG5, EPHA2, EYA1, FAM126A, FOXE3, FTL, FYCO1, GALK1, GALT, GCNT2, GJA3, GJA8, HSF4, LEMD2, LIM2, LISS, MAF, MIP, MIR184, NDP, NF2, NHS, OCRL, OPA3, 22142, BAY6, BEY7, BLYY2, BYDN, BAB2CARA, BFCO14, SCC, ONA, BAYCA, P3H2, PAX6, PEX7, PITX3, PXDN, RAB3GAP1, RECQL4, SC5D, SIL1, SIPA1L3, SLC16A12, TDRD7, VIM, VSX2, WRN

□ Septo-optical dysplasia (6 Genes, EYE22)

FGFR1, HESX1, OTX2, PROKR2, SOX2, SOX3

☐ Glaucoma (12 Genes, EYE23)

CYP1B1, FOXC1, FOXE3, LTBP2, MYOC, NTF4, OPTN, PAX6, PITX2, TBK1, TEK, WDR36

□ Corneal dystrophies (23 Genes, EYE24)

AGBL1, CHST6, COL17A1, COL8A2, CYP4V2, DCN, GSN, KRT12, KRT3, LOXHD1, MIR184, OVOL2, PIKFYVE, PRDM5, SLC4A11, TACSTD2, TCF4, TGFBI, TUBA3D, UBIAD1, VSX1, ZEB1, ZNF469





ADAMTSL4, FBN1	(10 Genes, EYE28) CHN1, COL25A1, DCC, KIF21A, MAFB, PHOX2A, ROBO3, SALL4, TUBB2B, TUBB3
☐ Congenital nystagmus, X-linked (2 Genes, EYE26) FRMD7, GPR143	
□ Progressive external Ophthalmoplegia (12 Genes, EYE27) DGUOK, DNA2, MGME1, OPA1, POLG, POLG2, RNASEH1, RRM2B, SLC25A4, TK2, TWNK, TYMP	
Additional analysis (additional face may such)	
Additional analyses (additional fees may apply)	
□ HLA-Typing (HLA01) I would like to receive an additional report stating the HLA alleles (HLADRB1, DRB3, DRB4, DRB5)).	A class I (Gene A, B, C) and HLA class II (Gene DPA1, DPB1, DQA1, DQB1,
of the American College of Medical Genetics and Genomics. The an sequence coverage will not typically be performed. A negative "ACMO fees may apply. According to German legislation, predictive tests for r	ommended genes for secondary analysis, according to the current guidelines alysis is restricted to the sequence data, re-sequencing of regions with poor G genes" report cannot be used to rule out (genetic) disease risk. Additional minors may not be performed for diseases which have an onset in adulthood. enotypic spectrum is within the scope of the primary medical indication of the os://www.cegat.com/acmg-genes/
□ Pharmacogenetics (PGX) (22 genes) ABCG2, CACNA1S, CYP2B6, CYP2C19, CYP2C9, CYP2D6, CYP3A4, CYP3, SLCO1B1, TPMT, UGT1A1, VKORC1	A5, CYP4F2, DPYD, G6PD, HLA-A, HLA-B, IFNL3, MT-RNR1, NUDT15, POR, RYR1,
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.de/en/diagnostic-support · diagnostic-support@cegat.de · Phone +49 7071 565 44-55