Request Form



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Sample type	Γ	_
🗆 Blood (EDTA)		
□ DNA, Specify Source:		
□ Others, Specify:		

] Prenatal¹⁾

 Native Amniotic Fluid (10-15 ml)
Native Chorionic Villi (20-40 mg)
Fetal DNA
Please add maternal EDTA blood or DNA for maternal cell contamination testing in case of prenatal analysis.

FOR LAB USE ONLY - DO NOT COVER

Requesting Health Care Professional Information

Physician Name:	Institution:	_			
Street:					
Country:					
Patient Information					
First Name:	Las	st Name:			
Date of Birth: Year /Month /Day					
Sex: □ Male □ Female □ Diverse □	∃Unknown Eth	hnicity:			
Patient History					
□ Unaffected □ Affected □ Age of Onse	et:				
Has the patient received hematopoietic stem cell tra	ansplantation?	□ Yes □ N	lo		
Suspected Diagnosis:				ICD-1():
Relevant Clinical Findings (attach copies of clinica	l reports if availa	ible)			
5. (/			
Family History					
Parental Consanguinity: 🛛 Yes 🗆 No)				
Affected Siblings/Family Members: 🗆 Yes 🛛 Relat	ion to Patient: _				_ 🗆 No
Clinical Information of Affected Family Members	(attach pedigree	if available	2)		
Test Information					
We offer a wide spectrum of molecular genetic ar	alvses. In genera	al, we will	perform the most su	iitable ana	lysis and gene composition
testing, based on the patient's medical history yo	ou provide us, or				
contact us at info@medgen-mainz.de in case of a	iny questions.				
Exome (Solo) Exome (Trio) (U	se seperate requ	iest forms f	or each family mem	iber to be	analysed)
Multi-Gene Panel corresponding to the suspect	ted diagnosis				
Customized Panel, Specify:					
Single Gene Testing Gene Name:	🔄 🗆 Sequ	uencing	🗆 Del/Dup Analysi	s (MLPA)	🗆 Repeat Analysis
Family Member Tested by Us: \Box No \Box	Yes Our Patier	nt ID:	Re	lation to P	Patient:
Regarding array CGH or cytogenetics , please find	detailed inform	ation on or	ır website		

Please Remember the Obligatory Declaration of Informed Consent (Back Page).

Billing Information

Invoice to:	🗆 Patient	Institution	
Institution/Last Name:			Department/First Name:
Street:			_ City:
Postal Code:			
Phone:			
Email (mandatory):			

□ The institution or the patient or the legal representative has been informed about the resulting costs and has approved them.

 \Box A quotation is required prior to starting the analysis.

Declaration of Informed Consent

The German Genetic Diagnostics Act (GenDG) requires detailed information and written consent for all genetic analyses as well as additional genetic counselling prior to prenatal and predictive genetic analyses. Please read this form carefully and check any statements that may apply to you.

My attending physician ______ has informed me about the nature, importance and implications of the genetic testing to be performed. All my questions have been answered to my satisfaction and I have had enough time for consideration to make an informed decision. I wish to be informed about the results of the genetic testing and I have been informed of my right not to know. I agree to the requested test being subcontracted to a cooperating specialized medical laboratory, if necessary.

With my signature below I give my consent to genetic analyses according to the enclosed request or which are necessary to clarify the above mentioned disease/disorder/diagnosis in question and I agree with the blood/tissue collection required for this.

I give my additional consent to:

Reporting of secondary findings

I am aware that genetic evaluation focuses on those changes directly related to my clinical indication or suspected diagnosis. However, comprehensive analyses, e.g. whole exome sequencing (WES), examine numerous genes in parallel and can possibly identify genetic variants not related to my existing clinical symptoms, but which could lead to an increased risk of disease and knowledge of which might be of medical value for my personal health and treatment in the future (ACMG guideline: Kalia et al., Genetic Medicine; 2017). I understand the significance of such secondary findings for myself and my biological relatives and I agree to be informed about those changes from which practical consequences may be deduced.

Extended storage and use of my data

I understand that the collected data/test results are stored for 10 years according to legal requirements and must be destroyed afterwards. However, these data may still be of great importance to me or my relatives (e.g. my children)	□ Yes	□ No
even after this period. Therefore, I give my consent to store my data beyond the legal period of 10 years.		
I agree that the collected data/test results may be used in encrypted (pseudonymised) form for scientific and quality assurance purposes, may be published anonymously in scientific journals and transferred in the context of scientific projects.	□ Yes	□ No
Extended storage and use of my sample material		

The Genetic Diagnostics Act requires that unused patient sample material be destroyed after the test is completed. With my consent, however, it may be stored. I authorize any remaining sample material to be transferred to the **Yes No** Medizinische Genetik Mainz, consent to its use for scientific and quality assurance purposes and to the transfer within the framework of scientific projects in encrypted (pseudonymised) form.

I agree that for a more precise assessment/classification of possibly causative changes detected in my child in the course of the genetic analysis performed, testing of my genetic material will be performed to confirm the kinship. The results of this testing will only be used for internal variant assessment, do not correspond to forensic analysis and will not be communicated as part of the results reporting.

I understand that I may withdraw this consent at any time, in whole or in part, without giving reasons and without risk of personal disadvantage. I understand that I have the right not to know the results of the genetic testing. I am aware that I can interrupt the analyses at any time until the results have been reported. I am aware that, once reported, the results are subjected to the legal retention period of 10 years and cannot be destroyed before expiration of this time even if requested.

Date

Date

Signature of Patient or (Legal) Representative

I hereby confirm that the consent as shown above, including all mentioned sub-items, has been declared by the patient or their legal representative.



Signature of Physician