PHYSICIAN INFORMATION	
INSTITUTION/PRACTICE	ADDRESS (STREET NAME, NO., CITY, POSTAL CODE, COUNTRY)
FIRST NAME	TELEPHONE NUMBER (COUNTRY CODE & NUMBER)
LAST NAME	E-MAIL ADDRESS (FOR REPORT ACCESS)
DATIENT INFORMATION	
PATIENT INFORMATION	
FIRST NAME	ADDRESS (STREET NAME, NO., CITY, POSTCODE, COUNTRY)
LAST NAME	TELEPHONE NUMBER (COUNTRY CODE & NUMBER)
DATE OF BIRTH (DD/MM/YYYY)	GENDER (MALE/FEMALE/OTHER - SPECIFY KARYOTYPE)
PERSONAL IDENTIFICATION NO.	SAMPLE COLLECTION DATE (DD/MM/YYYY)
REASON FOR TEST (DIAGNOSIS, PREDICTIVE, CARRIER)	
DECLARATION OF CONSENT (ACCORDING TO GERMAN GEN Applicable only for the determination of genetic (hereditary) characteristics	ETIC DIAGNOSTICS ACT, GenDG)
<ul> <li>By signing the form below I confirm that I:</li> <li>Have been fully informed by my physician about the significance and consequences of the genetic investigation, in compliance with GenDG.</li> <li>Have read/have been read the Information for Patients (page 5) which is attached to this form and which I fully understand.</li> <li>Have been given sufficient opportunity to discuss open questions.</li> <li>Authorize [insert legal entity here] to collect the necessary samples for investigation (blood, tissue, chorionic villus cells or amniotic fluid for prenatal diagnosis) and to send this form to MVZ Martinsried GmbH, Lochhamer Str. 29, 82152 Martinsried, Germany, in order to perform the tests requested through this form.</li> <li>Consent to the genetic test being carried out in order to clarify the disease/dysfunction/suspected diagnosis.</li> <li>YES NO</li> <li>I agree that the investigation or parts of the investigation may be</li> </ul>	<ul> <li>By signing the form below I confirm that:</li> <li>I may stop the investigation at any time and ask for the results available until that time to be destroyed.</li> <li>I may withdraw any of my consents given through this form entirely or in part at any time without giving reasons.</li> <li>I will be charged for the costs incurred until the time of withdrawal of consent.</li> <li>I may choose not to be informed about the test results (right not to know).</li> <li>I know that the genetic investigation and evaluation is limited to the requested indication and no statements will be made about other diseases.</li> <li>All information I have provided is true and correct.</li> <li>Communication of additional findings found during the course of the research  YES, I wish to be informed about additional findings.</li> <li>NO, I do not wish to be informed about additional findings.</li> </ul>
forwarded to collaborating medical laboratories, if necessary.  I agree with the evaluation of additional genes in the same indication group as part of the research.  I agree that the remaining specimens may be stored for further investigations after the examination is completed, yet not claiming storage.  I agree that the specimens, and if applicable DNA sequence information, may be made available anonymously for quality management and scientific purposes.  I agree that the results of the analysis may be stored for a longer period than the statutory period of 10 years, yet not claiming storage of results.  I agree to the storage and use of my test results under the protection of anonymity in a statistical database used for scientific purposes and to help diagnose genetic diseases. I understand that  I will remain under the protection of anonymity and I cannot be	In addition,  • YES NO I agree that a copy of the results of the analysis may be sent to the following physician(s), in accordance with my express requests and according to [insert legal entity here] internal procedures.  DR(S) NAME  STREET  POSTCODE/CITY  COUNTRY  PLACE  DATE  SIGNATURE OF PATIENT OR PARENT/LEGAL GUARDIAN
identified during the analysis of the data and that any personal information will be transformed into information of a non-personal nature.	PHYSICIAN'S SIGNATURE



## CARDIAC AND AORTIC PANELS KNOW&MANAGE

BARCODE

PATIENT INFORMATION		
Indication:		
Is there a pregnancy / partner's pregnancy?  Parental consanguinity:	No Yes No Yes	Gestational week+
CLINICAL SYMPTOMS		
Family history		
Are there other affected family members with similar sy If yes, who?	mptoms? No Yes	
CARDIAC SYMPTOMS  Syncope  Hypertrophic cardiomyopathy  Ventricular septal hypertrophy: mm  Left ventricular hypertrophy: mm  Sudden cardiac death / cardiac arrest  Dilated cardiomyopathy  Left ventricular dilation  ECG FINDINGS  Prolonged QTc interval: ms  Torsades de Pointes  Stress-induced polymorphic ventricular tachycardia	Right ventricular dilation Structural abnormalities of the myocardium:  Left ventricular noncompaction cardiomyopathy Reduced left ventricular ejection fraction: % Reduced right ventricular ejection fraction: % Atrial septal defect  Epsilon wave ST-segment elevation T-wave inversion Delta wave	Ventricular septal defect Tetralogy of Fallot Hypoplastic left heart syndrome Aortic stenosis Pulmonary stenosis Atrioventricular septal defect Valve defect (specify:)  Atrial fibrillation Ventricular fibrillation Atrioventricular block Right bundle branch block
Brugada Type 1 ECG  J wave	Ventricular tachycardia Ventricular extrasystoles	Left bundle branch block
LABORATORY PARAMETERS (please attach corresponded Total Cholesterol: Triglycerides:	ding abnormal lab findings)  HDL-Cholesterol: Abnormal enzyme level:	LDL-Cholesterol:
OTHER ABNORMALITIES  Sensorineural hearing loss Diabetes mellitus	Periodic paralysis Smoker	Systolic blood pressure:  Other abnormalities:
INHERITANCE - INDICATIONS OF:  Autosomal dominant	Autosomal recessive	X-linked
INVESTIGATION ORDER		
Our expert team is available for questions at any time: i Diagnostic Predictive  Targeted diagnostics for a known familial variant?  Gene: Variant:  For ordering Whole Exome Sequencing, please use the	Prenatal  Yes, please include a copy of the findings; alternated a copy of the findings.  Transcript:	atively, specify gene, variant, and transcript exactly:



## CARDIAC AND AORTIC PANELS KNOW&MANAGE

BARCODE

OUR PANELS		
ARRHYTHMOGENIC HEART DISORDERS  Comprehensive arrythmias & cardiomyopathies panel		
ARRHYTHMIA (CHANNELOPATHIES)  Comprehensive arrhythmias panel Atrial fibrillation panel Brugada syndrome panel Catecholaminergic polymorphic ventricular tachycardia panel Early repolarization syndrome panel Long QT syndrome panel Progressive cardiac conduction disorder panel Short QT syndrome panel Sinus node dysfunction panel	CARDIOMYOPATHIES  Comprehensive cardiomyopathies panel Arrhythmogenic cardiomyopathy panel Dilated cardiomyopathy panel Hypertrophic cardiomyopathy panel Noncompaction cardiomyopathy panel Restrictive cardiomyopathy panel	SYNDROMIC FORMS OF HCM WITH EXTRACARDIAC MANIFESTATION  Amyloidosis Friedreich's ataxia Hemochromatosis panel Cardiofaciocutaneous syndrome panel Lysosomal storage disorders panel Mitochondrial syndromes (Leigh syndrome, MELAS, MERRF) Myofibrillar myopathy panel RASopathies panel
AORTIC DISORDERS  Aortic disorders panel Ehlers Danlos syndrome panel Loeys Dietz syndrome panel Marfan syndrome panel Thoracic aortic aneurysm (+/- dissection) panel		
CONGENITAL HEART DEFECTS  Comprehensive congenital heart defects panel Alagille syndrome panel DiGeorge syndrome Heterotaxy panel Isolated & syndromic congenital heart defects panel RASopathies panel Williams Beuren syndrome		
LIPID METABOLISM DISORDERS  Familial hypercholesterolemia panel Hyperlipidemia / Hyperlipoproteinemia panel Hypolipidemia / Hypolipoproteinemia panel		
PHARMACOGENETICS  B-blocker (CYP2D6)  Clopidogrel (CYP2C19)  Mavacamten (CYP2C19)  Statine (SLCO1B1)  Sartane (CYP2C9)		

Please find the up-to-date list of genes for each panel on www.medicover-genetics.com.



## CARDIAC AND AORTIC PANELS KNOW&MANAGE

## **INFORMATION FOR PATIENTS**

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FATENT NAME  LAST NAME  LAST NAME  LAST NAME  LAST NAME  LAST NAME  DATE OF BIRTH (DD/MM/YYYY)  E-MAIL ADDRESS  CLINICAL DIAGNOSIS  Genetic connectling or commelling by the ordering physicion is necessary before ordering a test in order to inform the patient of all of the possible outcome I understand that I will be tested for: (to be filled in by physician)  The role of genetic testing, in many cases, a genetic test can directly detect a genetic alteration. Molecular tests can identify structural changes in the DNA (variants). Cytogenetic tests identify the chromosomal changes (structural or numerical). The sensitivity and specificity of each test varies, the tests offered are complex analyses and are performed using high-end equipment. The methods are externally validated, but there is a minimal possibility of errors.  The significance of the results. If the result is identified as being directly clausative of the clinical manifestations, it is considered to be conclusive. If the tests offered to be incondusive and this does not preclude other genetic changes (or non-genetic factors) responsible for the disease (a genetic disease or succeptibility to a genetic condition is not excluded). Therefore, an inconclusive result is identified as being directly changes (variants) not tested through the current analysis. Interpretation of the genetic custus relies on complete clinical picture of the patient, including clinical manifestations, family medical history and previous diagnoses. An error in diagnosis could occur due to a clinical picture that is different from that declared. In addition, the test can identify a possible nonpaternity. The testus relies on complete clinical picture of the patient, including clinical manifestations, a minimal previous diagnoses. An error in diagnosis could occur due to a clinical picture of the patient, including clinical manifestations, a minimal previous diagnoses. An error in diagnosis could occur due to a clinical picture of the patient in diagnosis could occur due to a clin		
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The significance of the results. If the result is identified as being directly causative of the clinical manifestations, it is considered to be conclusive. If the test does not identify the causative mutations of the clinical manifestations, it is considered to be inconclusive and this does not preclude other genetic changes (or non-genetic factors) responsible for the disease (a genetic disease or susceptibility to a genetic condition is not excluded). Therefore, an inconclusive result (no causative mutation identified) does not exclude the existence of other pathogenic genetic changes (variants) not tested through the current analysis. Interpretation of the genetic results relies on a complete clinical picture of the patient, including clinical manifestations, family medical history and previous diagnoses. An error in diagnosis could occur due to a clinical picture that is different from that declared. In addition, the test can identify a possible nonpaternity. The test results will be forwarded to the patient by the geneticist or ordering physician and are confidential.  By my signature, I hereby certify that:  1. I have been informed of the nature and purpose of the genetic test. by (name of physician).  3. I have been informed that the genetic test can provide information/results which have no connection with the purpose of testing. I understand that only I decide if I want those additional results to be provided.  4. I have received a copy of this form.	e given my written consent.	
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<ul><li>4. I have received clear answers to my questions in relation to the genetic test.</li><li>5. I have received a copy of this form.</li></ul>		
I have explained the risks and benefits of the test as well as alternative test methods to the parent/legal guardian. I have answered all the questions from	the parent/legal guardian.	
Name of the ordering physician		
FIRST NAME LAST NAME		
SIGNATURE OF THE ORDERING PHYSICIAN DATE OF SIGNATURE		

