SOLID TUMOR TESTS HISTOPATHOLOGY & GENETICS DETECT&ACT

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PERSON COMPLETING FORM **CONTACT (PHONE OR E-MAIL)** DATE (DD/MM/YYYY) 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) **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 GENETIC DIAGNOSTICS ACT, GenDG) The GenDG requires provision of detailed information and a written consent for all genetic investigations as well as genetic counselling prior to both predictive (applies to healthy individuals) and prenatal testing (with restrictions: prenatal testing is not performed for late manifesting disorders, including Hereditary Cancer Panels). The German Society of Human Genetics (GfH) and the Association of German Human Geneticists (BVDH) recommend clarifying the issues listed below during the information process. Please read the declaration of consent carefully and tick the boxes, in accordance with your consent. By signing the form below I confirm that I: By signing the form below I confirm that: • Have been fully informed by my physician about the significance and • I may stop the investigation at any time and ask for the results available until consequences of the genetic investigation, in compliance with GenDG. that time to be destroyed. • Have read/have been read the Information for Patients (page 4) which is • I may withdraw any of my consents given through this form entirely or in part attached to this form and which I fully understand. at any time without giving reasons. • I will be charged for the costs incurred until the time of withdrawal of consent. • Have been given sufficient opportunity to discuss open questions. • Authorize [insert legal entity here] to collect the necessary samples for • I may choose not to be informed about the test results (right not to know). investigation (blood, tissue, chorionic villus cells or amniotic fluid for • I know that the genetic investigation and evaluation is limited to the prenatal diagnosis) and to send this form to MVZ Martinsried GmbH, requested indication and no statements will be made about other diseases. Lochhamer Str. 29, 82152 Martinsried, Germany, in order to perform the • All information I have provided is true and correct. tests requested through this form. Communication of additional findings found during the course of the research Consent to the genetic test being carried out in order to clarify the YES, I wish to be informed about additional findings. disease/dysfunction/suspected diagnosis. NO, I do not wish to be informed about additional findings. I agree that the investigation or parts of the investigation may be In addition, forwarded to collaborating medical laboratories, if necessary. • YES NO I agree that a copy of the results of the analysis may be sent I agree with the evaluation of additional genes in the same to the following physician(s), in accordance with my express requests and indication group as part of the research. according to [insert legal entity here] internal procedures. I agree that the remaining specimens may be stored for further DR(S) NAME investigations after the examination is completed, yet not claiming storage. STREET I agree that the specimens, and if applicable DNA sequence POSTCODE/CITY information, may be made available anonymously for quality management and scientific purposes. COUNTRY 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 **PLACE** DATE storage of results. I agree to the storage and use of my test results under the SIGNATURE OF PATIENT OR PARENT/LEGAL GUARDIAN 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 identified during the analysis of the data and that any personal **PHYSICIAN'S SIGNATURE**



non-personal nature.

information will be transformed into information of a

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HISTOPATHOLOGY	
SAMPLE DETAILS	
SPECIMEN:	RETRIEVED DATE:
SPECIMEN ID:	COLLECTION DATE:
BLOCK ID:	COLLECTION TIME:
FIXATIVE/PRESERVATIVE:	BIOPSY DETAILS/BODY SITE:
REASON FOR REFERRAL: New Diagnosis Relapse In Remission Monitoring Other Relevant Clinical Information: Comprehensive information regarding clinical history and diagnosis is essential for interpretation of genomic findings and drug therapy recommendations. Please attach patient's pathology report (if available), clinical history, and other applicable report(s).	
If histopathology was conducted, please fill in: Stage Primary Metastasis - If metastasis, list primary: 0	

Our pathologists will recommend the most appropriate sequencing analysis, if necessary, and provide professional interpretation of the results.



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MOLECULAR ANALYSIS

Please choose the appropriate panel below

TARGETED-THERAPY TESTS

1 BLADDER CANCER

IHC markers: CD44, CK20, CK7, GATA3, Ki67, p53 Genes: FGFR2, FGFR3

2 BREAST CANCER

IHC markers: E-CD, ER, GATA3, HER2, Ki67, PR Genes: BRCA1, BRCA2, PIK3CA Molecular markers: ☐ fusion genes ☐ MSI

3 COLORECTAL CANCER IHC markers: CDX-2, CK20, MLH1, MSH2, MSH6, PMS6, villin Genes: BRAF, KRAS, NRAS

Molecular markers: ☐ fusion genes ☐ MSI

4 GASTRIC CANCER

IHC markers: CDX-2, CG-A, CK20, CK7, HER-2, Ki67, SYN Molecular markers: ☐fusion genes ☐MSI

5 GASTROINTESTINAL STROMAL TUMOR

IHC markers: CD34, c-KIT (CD117), desmin, DOG1, Ki67, S100 Genes: BRAF, KIT, NF1, PDGFRA, SDHA

Molecular markers: ☐ fusion genes

6 MELANOMA IHC markers: HMB45, Ki67, Melan-A, p16, S100, SOX10 Genes: BRAF, KIT, NRAS

Molecular markers: ☐ fusion genes

7 NON-SMALL CELL LUNG CARCINOMA

IHC markers: ALK, CG-A, EGFR, p40, PD-L1, ROS, SYN, TFF1 Genes: BRAF, EGFR, ERBB2, KRAS, MET Molecular markers: ☐ fusion genes

8 OVARIAN CANCER

IHC markers: AFP, calretinin, EMA, ER, hCG, inhibin, napsin A, OCT3/4, p16, p53, PAX8, PR, SALL4, WT1 Genes: BRCA1, BRCA2

Molecular markers: ☐ fusion genes ☐ MSI

9 PANCREATIC CANCER

IHC markers: CK19, CK20, CK7, MUC5AC Genes: BRAF, BRCA1, BRCA2, KRAS, PALB2, SMAD4 Molecular markers: ☐ fusion genes ☐ MSI

10 PROSTATE CANCER

IHC markers: BCC-AMACR, CK34BE12, CK5/6, NKX3.1, p63, PSA, PSAP Genes: ATM, BRCA1, BRCA2, CDK12, CHEK2, FANCA, PALB2, PTEN, RAD51 Molecular markers: MSI

Fusion genes: ALK, NTRK, RET, ROS

IHC, immunohistochemistry MSI, microsatellite instability

SOLID TUMOR PANELS (AND FUSION GENES)

BLADDER CANCER

- 11 Gene Panel
- 11 Fusion Genes

BREAST CANCER

12 Gene Panel

COLORECTAL CANCER

13 Gene Panel

GASTRIC CANCER

14 Gene Panel

GASTROINTESTINAL STROMAL TUMOR

15 Gene Panel

GLIOBLASTOMA

16 Gene Panel

LIVER CANCER

17 Fusion Genes

LUNG CANCER

18 Fusion Genes

MELANOMA

19 Gene Panel

NON-SMALL CELL LUNG CARCINOMA

20 Gene Panel

20 Fusion Genes

OVARIAN CANCER

21 Gene Panel

PANCREATIC CANCER

22 Gene Panel

22 Fusion Genes

PROSTATE CANCER

23 Gene Panel

23 Fusion Genes

SALIVARY GLAND SARCOMA

24 Fusion Genes

SOFT TISSUE SARCOMA

25 Fusion Genes

THYROID CANCER

26 Gene Panel

26 Fusion Genes

27 COMPREHENSIVE SOLID TUMOR PANEL

Please note that other gene fusions in addition to those specified may be detected by this assay and will be reported if they are of potential clinical significance.

For our complete gene list, turnaround times, specimen requirements and more, please visit our website: www.medicover-genetics.com.



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INFORMATION FOR PATIENTS

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PATIENT INFORMATION	
FIRST NAME	GENDER (MALE/FEMALE/OTHER - SPECIFY KARYOTYPE)
LAST NAME	TELEPHONE NUMBER (COUNTRY CODE & NUMBER)
DATE OF BIRTH (DD/MM/YYYY)	E-MAIL ADDRESS
CLINICAL DIAGNOSIS	
Genetic counselling or counselling by the ordering physician is necessary before ordering a test in order to inform the patient of all of the possible outcomes and the limitations of the genetic test. I understand that I will be tested for: (to be filled in by physician)	
I understand that the biological sample will be used to determine if I, or members of my family, are carriers of a genetic variant causing the disease, or are carriers of the disease, or have an increased risk of developing a disease.	
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 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	Incidental findings. Genetic testing can provide information unrelated to the purpose of the test, but that may have medical importance for the patient or family (information correlated with an increased risk for incurable disorders). Use of the sample/result. The sample provided will be used solely for the purpose of the test and for which I have given my written consent. Test results can also be used for research and to improve the diagnosis and treatment of genetic diseases. The genetic material can be used for other purposes only with my prior express written consent. Post-testing genetic counselling. A conclusive result may offer the patient information on the susceptibility, diagnosis, possible prognosis and/or heritability of the disease. An inconclusive result may lead to confusion and anxiety or may suggest the need for further genetic testing. Therefore, post-testing genetic counselling is advised for the clinical interpretation of the results.
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:	Completed by: Parent/Legal Guardian Patient
 I have been informed of the nature and purpose of the genetic test. I have been informed of the benefits and limitations of the genetic test by	FIRST NAME
	LAST NAME
	DATE OF COMPLETION
	SIGNATURE
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

