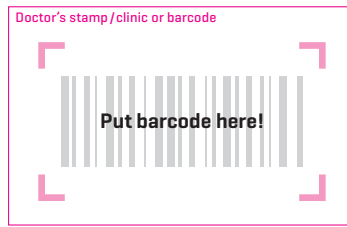


Health Insurance		
Surname, First Name		Date of birth
Health Insurance ID No.	Personal Insurance ID No.	Status
Business No.	Doctor's ID	Date



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Hereditary tumor syndrome consent form according to GenDG

Submitter Physician _____ Phone _____

Cost Unit Statutory health insurance (Please attach referral!) Privately insured Self-payer

Fast Track (Duration approx. 15 working days from sample entry¹), Reason [e.g., surgery appointment]: _____

Patient information

Gender Female Male Diverse Ethnic origin _____

Type of investigation **Affected/Diagnostic** **Predictive/Carrier-screening**

Family Anamnesis _____

Medical history/
Indication _____

Genetic findings available? (Patient/relatives) Yes (Please attach a barcoded document) No

For primarily predictive, please state reasons, e.g., index patient Not tested Deceased Findings of an index not available

Information on the desired analysis/requirement **2 EDTA sample tubes required**

Mamma and Ovarian carcinoma

MAMMA1 prior to Olaparib
Advanced, HER2/neu neg. Mamma-CA or high-grade epithel. Ov-CA (BRCA1, BRCA2)

MAMMA2, hereditary²
(ATM, BARD1, BRCA1, BRCA2, BRIP1, CDH1, CHEK2, PALB2, PTEN, RAD51C, RAD51D, STK11, TP53)

Lynch-syndrome (HNPCC) and Polyposis-syndrome

LYNCH1³
(EPCAM, MLH1, MSH2, MSH6, PMS2)

LYNCH2³
(MLH1, PMS2)

LYNCH3³
(EPCAM, MSH2, MSH6)

Pancreas carcinoma prior to Olaparib
metastasized, platinum-sensitive

PANKC1 (BRCA1, BRCA2)

Pancreas carcinoma, hereditary

PANKC2
(ATM, APC, BRCA1, BRCA2, CDKN2A, EPCAM, MLH1, MSH2, MSH6, PALB2, PMS2, STK11, TP53)

Prostate carcinoma prior to Olaparib
metastasized, castration-resistant

PROSC1 (BRCA1, BRCA2)

Prostate carcinoma, hereditary

PROSC2
(ATM, BRCA1, BRCA2, CHEK2, EPCAM, HOXB13, MLH1, MSH2, MSH6, PALB2, PMS2, RAD51D)

Uterus carcinoma, hereditary

UTERCA
(EPCAM, MLH1, MSH2, MSH6, MUTYH, NTHL1, PMS2, PTEN, POLD1, POLE, STK11)

Stomach carcinoma, hereditary

GASTCA
(BMPR1A, CDH1, EPCAM, MLH1, MSH2, MSH6, PMS2, SMAD4, STK11, TP53)

Familial mutation **FAMMUT** **Confirmation of findings** **MUTVER**

_____ (Please state/attach previous findings)

¹ regular test duration approx. 6 weeks

^{2,3} Indication criteria, see back page

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Einwilligung nach Gendiagnostikgesetz

Herewith in agreement with this consent form and completed consultation, I confirm that I had sufficient time to reconsider my desire to have the requested genetic analyses performed and the sampling material to be taken as required per the German Genetic Diagnostic Act. I was informed in detail about the purpose of this examination, the disease to be examined and its genetic basis, as well as the possibilities and limits of the diagnostics to be carried out in my specific case. I consent to the report being sent to the requesting physician and to the physicians specified by me. I agree to:

- The **forwarding of the request contract**, if necessary, to a specialized cooperating laboratory No
- The **storing of results** for and exceeding the statutory period of 10 years No
- The **storing of material** for possible testing at a later stage No
- The **use of testing material** for the purpose of quality assurance and research No
- The **use of test results** for the purpose of advising and testing of family members No

Information about additional findings: In rare cases, medical findings, which are not related to the initial question can be received, but which have a treatment consequence for me or my family (following the recommendations of the ACMG).

I would like to be informed about such findings (as far as no choice has been made below, "no" is assumed)

Yes No

This declaration of consent in accordance with GenDG is valid for me and for my child on its behalf, and may be revoked in parts or fully at any time.

Surname and First Name of informing physician

Place, Date

Signature of physician

Signature of patient/Legal guardian

Indication criteria for patients of the statutory health insurance

²Hereditary Mamma and Ovarian carcinoma

According to the quality assurance agreement "molecular genetics", the indication for testing the genes BRCA1 and BRCA2 is only given, if **one of the following criteria** is met. Please check the boxes, where applicable:

- ≥ 3 women from the same family lineage had breast cancer, independent of age
- ≥ 2 women from the same family lineage, one of them younger than 51 years, had breast cancer
- ≥ 1 woman had breast cancer, aged younger than 36 years
- ≥ 1 woman had bilateral breast cancer (first diagnosed younger than 51 years)
- ≥ 2 women from the same family lineage had ovarian cancer
- ≥ 1 woman had breast cancer and 1 other woman had ovarian cancer or 1 woman had both breast and ovarian cancer
- ≥ 1 man had breast cancer and 1 woman had breast or ovarian cancer
- ≥ 1 woman had a triple-negative breast cancer (younger than 51 years)

³Lynch-syndrome (HNPCC)

According to the quality assurance agreement "molecular genetics", the indication for direct testing of MMR-genes (MLH1, MSH2, MSH6 and PMS2) without prior analysis of tumor tissue, is bound to the fulfillment of the **Amsterdam-II-criteria**.

All the following criteria must be fulfilled. Please check the boxes, where applicable:

- ≥ 3 family members with HNPCC-associated carcinoma (colon/rectum, endometrium, small intestine, renal pelvis/ureter)
- 1 of the relatives concerned is a first-degree relative of affected persons
- Diseases in at least 2 subsequent generations
- At least 1 person with carcinoma diagnosis before the age of 50
- Exclusion of familial adenomatous polyposis coli (FAP)

Alternatively the testing of the microsatellite instability and, if necessary, testing of MMR-genes – dependent on immunohistological stainings (MLH1 and PMS2 or MSH2 and MSH6) – is bound to the fulfillment of the **revised Bethesda-criteria**.

One criterion must be met. Please check the boxes, where applicable:

- Colorectal carcinoma, initial diagnosis before the age of 50
- Synchronous/metachronous colon-/rectum carcinomas or HNPCC-associated cancer diseases (endometrium, renal pelvis/ureter, small intestine, stomach, pancreas, ovaries, hepatobiliary system, brain [commonly glioblastomas], sebaceous gland adenomas and keratoacanthomas, independent of age
- Colorectal carcinoma with MSI-H-typical morphology, diagnosed before the age of 60
- Patient with colorectal carcinoma and at least 1 direct relative with a HNPCC-associated tumor, who was initially diagnosed before the age of 50
- Patient with colorectal carcinoma and at least 2 first- or second-degree relatives with HNPCC-associated tumors (see above), independent of the age of disease onset