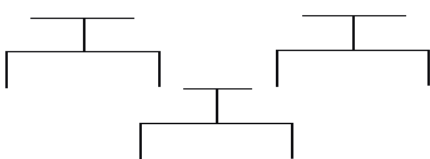


REQUISITION FORM FOR GENETIC DIAGNOSTIC BRCA TESTING

REFERRING DOCTOR (MANDATORY):	
Name (or print label):	ID number:
Hospital:	Service:
E-mail:	Telephone:
Signature:	Date: ____/____/____
Do you authorize the report to be sent by e-mail? Yes <input type="checkbox"/> No <input type="checkbox"/> If yes, please indicate the institutional e-mail address : _____	<i>GenoMed use only:</i> Label _____ Verified by: _____

PATIENT IDENTIFICATION (MANDATORY)	
Name (or print label):	Gender: F <input type="checkbox"/> M <input type="checkbox"/>
Identification number: _____	Date of birth: ____/____/____

CLINICAL DATA AND DIAGNOSIS (we ask a pathology report whenever possible and applicable):		
Affected <input type="checkbox"/>	Not affected (asymptomatic) <input type="checkbox"/>	Ideal expected date: ____/____/____

FAMILIAL INFORMATION	
Index case (affected) <input type="checkbox"/>	Familial variant <input type="checkbox"/>
Known familial variant? NO <input type="checkbox"/> YES <input type="checkbox"/>	
If yes*: Gene/RefSeq: _____ / _____ Variant: _____	
Was the index case studied at GenoMed? NO <input type="checkbox"/> YES <input type="checkbox"/>	
*For carrier studies and predictive testing (familial variant study), please attach a copy of the index case, whenever possible and applicable. According with the Portuguese law, (article 9º, law nº 12/2005), carrier studies and predictive testing (conducted in healthy individuals) it is ESSENTIAL that the clinical geneticist makes the request and that the patient informed consent is obtained.	
Attach whenever possible and can be justified any relevant clinical information, family data including consanguinity, other cases in the family, family tree, etc. Information for the construction of Pedigree:	
<input type="checkbox"/> Man <input type="radio"/> Woman <input checked="" type="checkbox"/> <input checked="" type="radio"/> Affected <input checked="" type="checkbox"/> <input checked="" type="radio"/> Carrier <input checked="" type="checkbox"/> <input checked="" type="checkbox"/> Deceased <input type="checkbox"/> <input type="radio"/> Consanguinity <input checked="" type="checkbox"/> Index case	
	

Sample: Blood (EDTA) <input type="checkbox"/> DNA <input type="checkbox"/> FFPE (10 sections x 10µm) <input type="checkbox"/> Other <input type="checkbox"/>	Collection: Date: ____/____/____ Time: ____:____:____
For somatic studies (in FFPE sample), we strongly recommend sending a blood sample together with the request.	

INFORMED CONSENT (to be filled by the referring doctor):	
I hereby declare that patient informed consent for diagnosis was obtained.	YES <input type="checkbox"/> NO <input type="checkbox"/>
I hereby declare that patient informed consent for investigation was obtained.	YES <input type="checkbox"/> NO <input type="checkbox"/>

Signature of the referring doctor: _____	Date:..... /...../.....
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REQUISITION FORM FOR GENETIC DIAGNOSTIC BRCA TESTING

Available Tests		SNS Code (PT)
<input type="checkbox"/>	Pack 1: <i>BRCA1</i> and <i>BRCA2</i> testing - index case Portuguese founder mutation (blood) + NGS (blood or FFPE) + MLPA (blood) *	34900
<input type="checkbox"/>	Pack 2: <i>BRCA1</i> and <i>BRCA2</i> testing - index case Portuguese founder mutation (blood) + NGS (blood or FFPE) *	34900
<input type="checkbox"/>	<i>BRCA1</i> testing - index case	34543
<input type="checkbox"/>	<i>BRCA1</i> testing - familial case	34544
<input type="checkbox"/>	Portuguese founder mutation <i>BRCA2</i> (insAlu) - index case	36061
<input type="checkbox"/>	Portuguese founder mutation <i>BRCA2</i> (insAlu) - familial case	36062
<input type="checkbox"/>	<i>BRCA2</i> testing - index case	34547
<input type="checkbox"/>	<i>BRCA2</i> testing - familial case	34548
<input type="checkbox"/>	<i>BRCA1/BRCA2</i> testing for CNVs - index case	36059
<input type="checkbox"/>	<i>BRCA1/BRCA2</i> testing for CNVs - familial case	36060

*It includes testing for mutation origin (germline or somatic).

Note: somatic study, on FFPE sample, is only available for pack 1 and 2.

INFORMED CONSENT, according to the Direction of General Health updated standard 015/2013 ([mandatory](#) - to be filled by the patient):

Hereby I _____ [name], ____/____/____ [date of birth], give my consent that my / my child's biological sample will be examined for genetic changes (mutations) in the specified gene(s), related to the diseases / clinical features described above. Herewith I declare that I have been informed about the chances and limitations of the requested testing procedure. I was informed in detail about the consequences resulting from the test results. I agree that the sample may be stored in order to allow repetition of the tests or further related tests in the future. All data about me / my child are subject to medical confidentiality. They can be disclosed to family members or their physicians only with my permission, but not to third parties. I am entitled to revoke this consent at any time.

Agree Do not agree

I agree that my / my child's tests results and/or clinical data may be used in scientific publications in anonymized form in case of approval of the Ethics Committee.

Agree Do not agree

Patient's signature: _____ **Date:** ____/____/____

SAMPLE INFORMATION (FFPE) (REQUIRED FIELD) We ask a pathology report whenever possible, or the tumor cell percentage, for correctly interpret test results.

Sample characterization: Exam nº: _____ Resection/Biopsy Date: ____/____/____

Sections **Macrodissection** Yes No **% Neoplastic Cells** _____
(If tumor cell percentage is lower than 20-30%, please use macrodissection to enrich the sample)

FFPE block

Histology: Biopsy Needle biopsy Surgical piece

Contaminants: Epithelial cells Blood
 Mesenchymal cells Fibrin
 Inflammatory/immune response cells Mucus Other(s) _____

Pathology / Clinical Diagnosis / Relevant information about the sample:

Pathologist: _____ Hospital: _____

Direct contact (☎ or @): _____

Signature: _____

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