Cancer Genetics Pathology Request Form





Patient Information	Requesting Clinician		
Surname:	Name:		
First Name: M F	Address:		
	Postcode:		
DOB: DD MM MY YYYY Phone:	Phone: Fax:		
Address:			
Postcode:	Provider No.		
Medicare No.: No. next to name:	Email (report delivery):		
PATIENT INFORMATION: Your treating practitioner has recommended that you use Genomic Diagnostics. You are free to choose your own pathology provider. However, if your treating practitioner has specified a particular pathologist on clinical grounds, a	Signature:		
Medicare rebate will only be payable if that pathologist performs the service. You should discuss this with your treating practitioner. MEDICARE ASSIGNMENT: (Section 20A of the Health Insurance Act 1973) I offer to	Report Copy Name:		
assign my right to benefits to the approved pathology practitioner ("APP") who will render the requested pathology services and any eligible pathologist determinable service(s)			
established as necessary by the practitioner. In the event that I am issued an account for those services, I also authorise that APP to submit my unpaid account to Medicare so	Address:		
that Medicare can assess my claim and issue me a cheque payable to the APP for the Medicare Benefit.	Postcode:		
Patient Signature: Date:	Phone: Fax:		
Tank Damasakad			
Test Requested Legend: B = Breast, O = Ovarian, P = Prostate, L = Lynch	Family History of Cancer YES NO		
Medicare Criteria Met Private Fee	Conser Type Poletienship		
BRAoVO™ Gene Panel (BRC)	Cancer Type Relationship		
BRAoVO™ Plus Gene Panel (BRC) □ 73296 (B/O) □ 73295 (B/O) □ 73304 (P) □ 73354 (L) □			
BRCA1 & BRCA2 Genes (BRC)			
Lynch Gene Panel (LYN) ☐ 73354			
FAP Gene Panel (AOP)			
Predictive Familial Cancer Test: Gene: Variant:			
□ 73297 (B/O-BRC) □ 73357 (Colorectal – PGT) □			
Please provide a copy of the relative's variant report.			
Germline test for Tumour Gene: Variant:	Clinical Details		
BRCA1/2 variant	Comments:		
Genetic Counselling			
 □ Provided by specialist and written informed consent provided overleaf, OR □ Pre-test counselling required – Note this is only covered for MBS eligible Gene Panel referrals 			
Gene Panel List			
BRAoVO™ (13) Gene Panel - ATM, BARD1, BRCA1, BRCA2, BRIP1, CDH1, CHEK2, PALB2, PTEN, RAD51C, RAD51D, STK11, TP53			
Lynch (5) Gene Panel – MLH1, MSH2, MSH6, PMS2, EPCAM BRAoVO™ Plus (18) Gene Panel – Combined BRAoVO™ and Lynch gene panels FAP (2) Gene Panel – APC, MUTYH			
Payment for non-MBS testing is required prior to testing – see below	Do not send reports to My Health Record		
r aymone for more in the cooling to require a prior to too and to the second			

Payment Information

IF your test is NOT covered by Medicare:

- Full payment is required prior to blood collection.
- · Visit gdpay.com.au to pay or scan QR code.

Receipt Number:

• Call 1800 822 999 (Mon-Fri 0800 - 1800 EST) with any queries



Amount Paid:

Collection Information

Collect 2 x 6mL or 10mL EDTA tube at 10 minute intervals.

PERSON COLLECTING SPECIMEN TO COMPLETE:

I certify I established the identity of the patient named on this request, collected and immediately labelled the accompanying specimen with the patient's details.

Initials:	ACC Code / Location:	
Date of draw:	D D M M Y Y Time:	am/pm

Patient Informed Consent



Informed Consent for Cancer Genetic Testing

This form is to be completed at the conclusion of your Genetic Counseling Consultation				
I (pati	ent),	(print name)		
of		(address)		
hereb	y consent to perform the following gene	tic testing:		
	-	(insert test name)		
I have	e been informed of and understan			
1.	The potential outcomes of the test, includ myself and my relatives.	ling the potential benefits and risks and the implications that this may have for both		
2.	A blood sample will be collected from me	using standard techniques, which carry very little risk.		
3.	The information that I have provided will r de-identified and used for statistical purpo	emain confidential, in accordance with the Privacy Act 1988. My test results may be oses.		
4.	· · · · · · · · · · · · · · · · · · ·	used to assist in improving testing methods.		
5.	Identification of pathogenic variants within treatment protocols and better patient man	this gene(s) may assist clinicians in accurate diagnosis, the selection of appropriate nagement.		
6.	such a variant may exist. This may be du structure, or inability of the technology used	dentify a genetic variant that is associated with increased risk of cancer, even though e to the current lack of knowledge in the scientific community of the complete gene d to identify certain types of changes in genes. In addition, a genetic variant associated be detected because the pathogenic variants may occur in another gene that has not		
7.	If a genetic variant that is associated with i risk of developing cancer in the future.	ncreased risk of any type of cancer is not identified, this does not mean that I am at no		
8.	In some cases, a genetic variant of uncert	ain clinical significance may be detected in one or more genes.		
9.	Test results are based on current knowled	ge, which may change in the future.		
10.		ely voluntary and I may withdraw from the testing at any stage prior to the issue of my in writing. However, if testing is cancelled, a fee may be charged for work completed.		
11.	My test result may have implications for or result may be used to facilitate the counselling	ther members of my family. I have been encouraged to advise them of this result. My ng and testing of other family members.		
12.	The test will not affect my ability to obtain Au insurance and travel insurance.	stralian health insurance but could potentially affect my ability to obtain some types of life		
13.	and, at my future request and with my cons	anded continuously. Genomic Diagnostics will store my sample for a minimum of 5 years sent, may be able to re-test the DNA by a new procedure for additional genes. However, ne availability or integrity of the sample for future use.		
14.	My contact details and test result may be p counselling.	provided to an accredited genetic counsellor for the sole purpose of arranging genetic		
Patie	nt:			
		ed the above to me and I have had the opportunity to ask questions. I am o my questions. I hereby consent to the above statements on this consent form.		
		nformation with health practitioners to help with the genetic testing of blood ation will not be disclosed to the relative wherever possible.		
Patien	nt Signature:	Date:/		
Patien	nt Name (please print):			
Phys	ician/Genetic Counsellor:			
I have		including risks, benefits and alternatives) of the requested genetic test to this		
Practi	tioner Name:	Practitioner Signature:		
Phone	e: Email:	Date:/		

For more information, contact us at info@genomicdiagnostics.com.au

