



Hereditary Cancer Risk Testing

- BILL TO: My Account, Insurance Provided, Lab Card/Select, Patient

PRINT PATIENT NAME (LAST, FIRST, MIDDLE), REGISTRATION # (IF APPLICABLE), DATE OF BIRTH, SEX, LAB REFERENCE #, CELL PHONE

ACCOUNT #: NAME: ADDRESS: CITY, STATE, ZIP: TELEPHONE #:

IMPORTANT! THIS FORM MUST BE FILLED OUT IN ITS ENTIRETY. Reflex tests are performed at an additional charge. Each sample should be labeled with at least two patient identifiers at time of collection. ICD Diagnosis Codes are Mandatory. Fill in the applicable fields below.

PATIENT ID # / MRN, PATIENT PHONE

PATIENT EMAIL ADDRESS

PRINT NAME OF INSURED/RESPONSIBLE PARTY (LAST, FIRST, MIDDLE) - IF OTHER THAN PATIENT

PATIENT STREET ADDRESS (OR INSURED/RESPONSIBLE PARTY) APT. # KEY #

NPI/UPIN ORDERING/SUPERVISING PHYSICIAN AND/OR PAYERS (MUST BE INDICATED)

Form for NPI/UPIN and patient information

PRIMARY INSURANCE: CITY, STATE, ZIP, RELATIONSHIP TO INSURED, PRIMARY INSURANCE CO. NAME, MEMBER / INSURED ID NO. #, GROUP #, INSURANCE ADDRESS

ADDIT'L PHYS.: Dr. NPI/UPIN, NON-PHYSICIAN PROVIDER: NAME, I.D.#, Fax Results to, Client # OR NAME, ADDRESS, CITY, STATE, ZIP

Preauthorization approved, Preauthorization number, Preauthorization not submitted

ABN required for tests with these symbols

- Medicare Limited Coverage Tests: @ = May not be covered for the reported diagnosis. F = Has prescribed frequency rules for coverage. & = A test or service performed with research/experimental kit. B = Has both diagnosis and frequency-related coverage limitations. Provide signed ABN when necessary

Visit QuestDiagnostics.com/MLCP for Medicare coverage guidelines

ICD Codes (enter all that apply)

THIS REQUISITION MUST BE ACCOMPANIED BY THE COMPLETED PATIENT AND FAMILY CLINICAL HISTORY FORM.

For fastest processing, please fax this Requisition and Patient and Family Clinical History Form to 855.422.5181. The Clinical History form can also be completed online at YourHistoryForm.com

Specimen Source: Blood, Saliva, Buccal

Other sample types call 1-866-GENEINFO (1-866-436-3463) prior to collecting and ordering

Breast Cancer Risk

- 91863 BRCA Panel - BRCA1 and BRCA2
92587 BRCA Panel Plus (7 Genes) BRCA1, BRCA2, CDH1, PALB2, PTEN, STK11, TP53
38621 Hereditary Breast Cancer Panel (16 Genes) ATM, BRCA1, BRCA2, CDH1, CHEK2, EPCAM, MLH1, MSH2, MSH6, NBN, NF1, PALB2, PMS2, PTEN, STK11, TP53
91864 BRCA Ashkenazi Jewish Screen (Common founder mutations BRCA1 c.68_69del, BRCA1 c.5266dup, and BRCA2 c.5946del)
92140 BRCA Ashkenazi Jewish Screen with Reflex to BRCA1/2 Sequencing and Deletion/Duplication (BRCA Ashkenazi Jewish Screen, if negative reflex to BRCA Panel-BRCA1 and BRCA2)
91866 BRCA1 and BRCA2 Deletion/Duplication

Colorectal Cancer/Polyposis Risk

- 94053 Juvenile Polyposis Panel - BMPR1A and SMAD4
91461 Lynch Syndrome Panel - MLH1, MSH2 (inc. EPCAM), MSH6, and PMS2
38631 Hereditary Colorectal Cancer Panel (19 Genes) APC, AXIN2, BMPR1A, CDH1, EPCAM, GREM1, MLH1, MSH2, MSH3, MSH6, MUTYH, NTHL1, PMS2, POLD1, POLE, PTEN, SMAD4, STK11, TP53

Expanded Hereditary Cancer Risk Panels

- 38611 Guideline Based Hereditary Cancer Panel (32 Genes) APC, ATM, AXIN2, BMPR1A, BRCA1, BRCA2, BRIP1, CDH1, CDK4, CDKN2A (p16, p14), CHEK2, EPCAM, GREM1, HOXB13, MLH1, MSH2, MSH3, MSH6, MUTYH, NBN, NF1, NTHL1, PALB2, PMS2, POLD1, POLE, PTEN, RAD51C, RAD51D, SMAD4, STK11, TP53
38600 Comprehensive Hereditary Cancer Panel (66 Genes) APC, ATM, AXIN2, BAP1, BARD1, BLM, BMPR1A, BRCA1, BRCA2, BRIP1, CDH1, CDK4, CDKN1B, CDKN2A (p16, p14), CHEK2, DICER1, EGFR, EPCAM, FANCA, FANCC, FANCM, FH, FLCN, GALNT12, GREM1, HOXB13, MAX, MEN1, MET, MIF, MLH1, MRE11 (MRE11A), MSH2, MSH3, MSH6, MUTYH, NBN, NF1, NTHL1, PALB2, PMS2, POLD1, POLE, POT1, PTCH1, PTEN, RAD50, RAD51C, RAD51D, RECQL, RET, SDHA, SDHAF2, SDHB, SDHC, SDHD, SMARCA4, SMAD4, STK11, SUFU, TMMEM127, TP53, TSC1, TSC2, VHL, XRCC2

Other Cancer Risk

- 38661 Tuberosclerosis Complex Panel - TSC1 and TSC2
38651 Nevoid Basal Cell Carcinoma (Gorlin) Syndrome Panel - PTCH1 and SUFU
38641 Hereditary Endocrine Cancer Panel (12 Genes) FH, MAX, MEN1, NF1, RET, SDHA, SDHAF2, SDHB, SDHC, SDHD, TMMEM127, VHL

Single Gene Tests

- (For a known familial mutation please see single site testing)
93797 APC 38805 FH 91458 MSH6 38809 SMARCA4
38802 ATM 38806 FLCN 93944 MUTYH 92565 STK11
38803 BAP1 38807 HOXB13 93941 NF1 92560 TP53
38804 BLM 93942 MEN1 92571 PALB2 93943 VHL
92568 CDH1 38808 MITF 91457 PMS2
93939 CDKN2A 91460 MLH1 92566 PTEN
93940 CHEK2 91471 MSH2 (inc. EPCAM) 93796 RET

Single site testing for any gene in the Comprehensive Hereditary Cancer Panel

93945 Gene Name: Variant Name: Copy of family member's report MUST be submitted with order

REQUIRED SIGNATURES PATIENT ACKNOWLEDGEMENT

I authorize Quest Diagnostics (Quest) to release information received, including, without limitation, medical information, which includes laboratory test results, to my health plan/insurance carrier and its authorized representatives as necessary for reimbursement. I further authorize my health plan/insurance carrier to directly pay Quest for the services rendered. I understand that I may be responsible for portions of this test not covered by my insurance.

SIGNATURE REQUIRED Patient Signature Date

STATEMENT OF MEDICAL NECESSITY AND INFORMED CONSENT

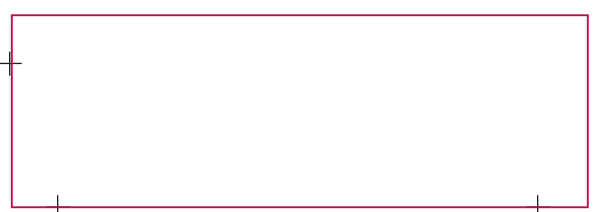
I have supplied information to the patient regarding genetic testing and the patient has given consent for genetic testing to be performed. I further confirm that this test is medically necessary for the diagnosis or detection of disease, illness, impairment, symptom, syndrome, or disorder and the results will be used in the medical management and treatment decisions for the patient. I confirm that the person listed in the Ordering Physician space above is authorized by law to order the test(s) requested herein.

SIGNATURE REQUIRED Medical Professional's Signature X Date

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If you have questions regarding this order, please call 866.GENE.INFO

Many payers (including Medicare and Medicaid) have medical necessity requirements. You should only order those tests which are medically necessary for the diagnosis and treatment of the patient.



SPECIMEN KEY ON BACK

Provide signed ABN when necessary

ICD Diagnosis Codes are Mandatory. Fill in the applicable fields below. Provide signed ABN when necessary

All samples to be shipped ambient, unless otherwise specified.

Specimen Key:

- L = Lavender top tube
- ◆ = See the QuestHereditaryCancer.com Test Selection Guide for the appropriate clinical application of this test.



Patients: Minimize your wait time by scheduling an appointment at a convenient Patient Service Center.

To find a location and make an appointment visit us at QuestDiagnostics.com/appointment or call 1.888.277.8772 or simply download our mobile app. at QuestDiagnostics.com/mobile



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PRIMARY INSURANCE: CITY, STATE, ZIP, RELATIONSHIP TO INSURED: SELF, SPOUSE, DEPENDENT, PRIMARY INSURANCE CO. NAME, MEMBER / INSURED ID NO. #, GROUP #, INSURANCE ADDRESS, CITY, STATE, ZIP

ADDIT'L PHYS.: Dr. NPI/UPIN, NON-PHYSICIAN PROVIDER: NAME, I.D.#, Fax Results to: (), Client # OR NAME: ADDRESS: CITY: STATE: ZIP:

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REQUIRED SIGNATURES PATIENT ACKNOWLEDGEMENT: I authorize Quest Diagnostics (Quest) to release information received, including, without limitation, medical information, which includes laboratory test results, to my health plan/insurance carrier and its authorized representatives as necessary for reimbursement. STATEMENT OF MEDICAL NECESSITY AND INFORMED CONSENT: I have supplied information to the patient regarding genetic testing and the patient has given consent for genetic testing to be performed.

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