

Genetic Risk Evaluation and Testing

Many people have a family member who has had cancer. While most cancer happens by chance, about 5-10% of cancers are hereditary. Hereditary cancers are caused by a harmful variant in a gene that runs in families and may be passed down from parents to children, causing a higher risk for cancer. "Hereditary cancers" may also be called "inherited cancer," or they may be associated with specific genetic syndromes or "familial cancer syndromes." If you have a strong history of cancer in your family, ask your healthcare provider about a genetic risk evaluation. Learning if you have inherited a harmful variant in a cancer gene can help you take steps for early detection and cancer prevention.

Why are Some Cancers Inherited?

All cancers are caused by harmful changes, or variants (also called mutations), in genes. Genes are like blueprints or instructions our body uses to grow and function. Cancer genes work to protect our bodies from developing cancer. When a cancer gene contains a harmful variant, the gene cannot function correctly and does not protect against cancer as well as it should; and this leads to a higher risk for cancer. Usually these harmful variants are acquired over the course of a person's life. However, sometimes a harmful variant can be inherited from one or both parents. These inherited variants increase the chance of developing certain cancers, often at a younger age than expected, and they can be passed down to the next generations of a family.

Factors Associated with Hereditary Cancer

- Multiple cancers in one person
- Cancer diagnosed at a young age
- Multiple family members with the same type of cancer
- Cancer in multiple generations
- Family member with a harmful variant in any cancer gene

Common Hereditary Cancer Syndromes

- Rare cancers or cancers in the less affected gender (e.g., male breast cancer)
- Cancer in paired organs (e.g., both breasts, both ovaries, both kidneys)
- More than one childhood cancer in siblings

Most hereditary cancer syndromes are rare, but some are more common than others. About 1 in 190 people have a harmful variant in a *BRCA1* or *BRCA2* gene which causes Hereditary Breast, Ovarian, and Pancreatic Cancer Syndrome. About 1 in 279 people have Lynch Syndrome, caused by a harmful variant in the *MSH1, MSH2, MSH6, PMS2,* or *EPCAM* genes. Some red flags for these syndromes are listed below. If your or your family members have any of these risk factors, discuss a genetic risk evaluation with your healthcare provider.

Hereditary Breast, Ovarian, and Pancreatic Cancer Syndrome Risk Factors	Lynch Syndrome Risk Factors
 Breast cancer at age 45 or younger Ovarian, male breast, pancreatic, intraductal prostate, metastatic prostate, or metastatic (HER2 negative) breast cancers at any age 2 breast cancers in a person, with 1 before age 65 Triple negative breast cancer at age 60 or younger 2 or more relatives with breast cancer with 1 before age 50 3 or more relatives with breast, ovarian, pancreatic, and/or aggressive prostate cancer at any age A relative with a harmful variant in <i>BRCA1</i> or <i>BRCA2</i> 	 Lynch syndrome related cancers include: colon, rectal, endometrial (uterine), ovarian, stomach, ureter/renal pelvis, biliary tract, small bowel, pancreatic, brain, or sebaceous adenomas Colon or endometrial cancer before age 50 A person with 2 or more of the Lynch cancers listed above 2 or more relatives with a Lynch cancer, with one diagnosed before the age of 50 3 or more relatives with a Lynch cancer A relative with a harmful variant in a Lynch Syndrome gene

What to Expect During a Genetic Risk Evaluation

A genetic risk evaluation is done by a trained medical professional to determine your risk for developing

cancer, to make genetic testing recommendations, and to decide appropriate medical management. A genetic evaluation may include:

- **Medical History:** A detailed review of your personal medical and family history. This information helps determine your risk of developing cancer, appropriate medical management, and if genetic testing would be useful to help clarify your cancer risk.
- **Counseling:** Before having genetic testing, you should be counseled about the risks, benefits, and limitations of genetic testing and cancer risk assessments, to help you make an informed decision about whether or not testing is the right choice for you. After testing, counseling should include a review of your test results and what they mean for you and your family.
- **Testing:** Genetic testing looks at a set of your genes (usually through a blood sample) to see if you have any harmful variants that increase your risk for cancer. Insurance companies usually cover the cost of testing if you have a personal or family history that is concerning for hereditary cancer. Cost and insurance coverage for testing are discussed during your evaluation.
- **Next Steps:** Based on whether or not you have genetic testing and what the results show, you may have different options for cancer screenings and prevention, or even tailored cancer treatments.
- **Family Risk:** If a harmful variant is found in a family, other family members are encouraged to consider genetic testing as well. The information from genetic counseling and testing helps family members make decisions that could prevent cancer and save their lives. This information can also be reassuring to those who do not have the harmful variant found in their family.

Benefits of Genetic Counseling and Testing	Risks of Genetic Testing
 Relief from uncertainty 	 Difficulty coping with cancer risk
 Understand your cancer risk 	 Impact on family and personal relationships
 Make informed medical and lifestyle decisions 	 Concerns about privacy or discrimination
 Provide helpful information for relatives 	 Cost of testing and recommended medical care

Who Should have Hereditary Cancer Genetic Testing?

It is important to identify individuals with hereditary cancers, but not everyone needs genetic testing. Before doing hereditary cancer testing, you should meet with a genetics professional who can review your personal and family history, determine if genetic testing would be helpful for you and/or your family, and decide the best test for you. Individuals with cancer are usually better candidates for genetic testing than their family members who have not had cancer. However, ask your healthcare provider for a referral for genetic evaluation if you or anyone in your family has any of the risk factors listed above or if you have a specific concern about your family cancer history.

Confidentiality of Results

The results of genetic counseling and testing are confidential, with very limited exceptions. Neither your health insurance company nor your employer can require you to have genetic testing. It is also against the law (Genetic Information Nondiscrimination Act) for health insurance companies or employers to discriminate against someone based on genetic information. This law does not apply to life, disability, or long-term care insurance, nor to military members, very small companies, or many federal health plans.

About Texas Oncology

Texas Oncology is an independent private practice with more than 500 physicians and 210 locations across the state. Meeting the oncology needs of Texans for more than 35 years, the practice includes Texas Center for Proton Therapy, Texas Breast Specialists, Texas Oncology Surgical Specialists, Texas Urology Specialists, and Texas Center for Interventional Surgery. As a lead participant in US Oncology Research, Texas Oncology played a role in the development of more than 100 FDA-approved therapies. For more information, visit www.TexasOncology.com.

Sources: American Cancer Society, American Society of Clinical Oncology, National Cancer Institute, and National Comprehensive Cancer Network



www.TexasOncology.com 888-864-4226

