

# Understanding Your VUS Hereditary Cancer Genetic Test Result

## INFORMATION FOR PATIENTS WITH A VARIANT OF UNCERTAIN SIGNIFICANCE

<b>RESULT</b>	The testing found one or more variants of uncertain significance (VUS). There is not currently enough information available to know if the VUS identified is expected to cause an increased risk for cancer or not.
<b>RECLASSIFICATION</b>	Collecting information about a VUS is an ongoing process, so it is possible that your result may be better understood in the future. Ambry regularly reviews the data and published evidence about each VUS, and your healthcare provider will be notified if enough new information becomes available to reclassify your VUS. For this reason, it is recommended that you continue to follow-up with the healthcare provider that ordered your genetic testing.
<b>CANCER RISK</b>	Even though your genetic test result was a VUS, you and your relatives may still have an increased risk of developing cancer based on other factors, including your medical and/or family history. It is important to discuss these risk factors with your healthcare provider.
<b>WHAT YOU CAN DO</b>	Risk management decisions are very personal and depend on many factors. It is important to discuss these options with your healthcare provider and decide on a plan that works for you.
<b>FAMILY</b>	Certain family members may be eligible for genetic testing through our family studies program. In some cases, testing family members may help add to the understanding of your result. However, not all genes are well suited for family studies testing. To determine if your VUS is eligible for family studies testing, your healthcare provider can contact <a href="mailto:FamilyStudies@ambrygen.com">FamilyStudies@ambrygen.com</a> .
<b>RESOURCES</b>	<ul style="list-style-type: none"> <li>American Cancer Society <a href="https://cancer.org">cancer.org</a></li> <li>National Society of Genetic Counselors <a href="https://nsgc.org">nsgc.org</a></li> <li>Canadian Association of Genetic Counsellors <a href="https://cagc-accg.ca">cagc-accg.ca</a></li> </ul>

## WHAT VARIANT CLASSIFICATIONS MEAN

<b>PATHOGENIC VARIANT (POSITIVE TEST RESULT)</b>	Enough evidence showing it can cause a disease
<b>LIKELY PATHOGENIC VARIANT (VLP) (POSITIVE TEST RESULT)</b>	Strong evidence to suggest it causes a disease
<b>VARIANT OF UNCERTAIN SIGNIFICANCE (VUS) (UNCERTAIN TEST RESULT)</b>	Limited and/or conflicting evidence to suggest it may cause a disease
<b>LIKELY BENIGN VARIANT (VLB) (NEGATIVE TEST RESULT)</b>	Strong evidence to suggest it does not cause a disease
<b>BENIGN VARIANT (NEGATIVE TEST RESULT)</b>	Enough evidence to show it does not cause a disease

Please discuss this information with your healthcare provider. The cancer genetics field is continuously evolving, so updates related to your genetic test result, medical recommendations, genetic testing options, and/or potential treatments may be available over time. This information is not meant to replace a discussion with a healthcare provider and should not be considered or interpreted as medical advice.