Hereditary Pancreatitis: Trypsinogen Gene Mutations

Frequently Asked Questions, Including Genetic Testing download this file in PDF format

Topics:
- What is hereditary pancreatitis (HP)?
- Is there a cure for HP?
- What causes HP?
- If I have inherited any of the HP mutations, will I definitely develop pancreatitis?
- What is the risk for my child to inherit HP?
- Genetic testing for HP
- Who might benefit from genetic testing?
- Potential benefits of genetic testing
- Potential risks & limitations of genetic testing
- Disclaimer
- Recommended reading

What is hereditary pancreatitis (HP)?

Hereditary Pancreatitis is a rare genetic condition characterized by recurrent episodes of acute pancreatitis attacks. In about half of these cases the problem progresses to chronic pancreatitis, which is severe scarring of the pancreas. Symptoms of an acute attack include abdominal pain, nausea, and vomiting. Laboratory test during an attack usually detect high blood levels of amylase and lipase, which are enzymes released from the pancreas. The first attack typically occurs within the first two decades of life, but can begin at any age. In the United States, it is estimated that at least 1,000 individuals are affected with hereditary pancreatitis.

Is there a cure for HP?

At this time, there is no cure for HP. Treating the symptoms associated with HP is the choice method of medical management. Patients may be prescribed pancreatic enzyme supplements to treat maldigestion, insulin to treat diabetes, analgesics and narcotics to control pain, and lifestyle changes to reduce the risk of pancreatic cancer (for example, NO SMOKING!).

Dietary recommendations to help control pain with digestion include the consumption of small meals throughout the day that are high in carbohydrates and low in protein and fat. Pancreatic enzymes such as Creon, Pancrease, and Violiase are helpful in providing improved digestion and a reduction in diarrhea and pain for some patients with more advanced disease.

Exposure to smoking and alcohol are known to dramatically increase the risk for pancreatic attacks among individuals with HP. Smoking is strongly discouraged as it doubles the risk for pancreatic cancer. Similarly, alcohol consumption is not recommended for these patients because alcohol is a known risk factor for both acute and chronic pancreatitis. Therefore it is recommended that all HP patients avoid smoking and alcohol consumption.

What causes HP?

HP is a genetic disorder, which means that it is usually passed from one generation to the next.

The symptoms of HP are caused by a change to a specific gene. Genes are the packages of information that control how our bodies look and function. A single gene appears to be involved in 60-75% of hereditary pancreatitis families. This gene produces the "cationic trypsinogen" enzyme, which breaks
down the proteins present in the foods we eat. (In some research papers cationic trypsinogen is called PRSS1).

When a change to a gene occurs, the gene may no longer function properly. These gene changes are called mutations. Currently, there are two common, and more than 6 uncommon cationic trypsinogen gene mutations that are associated with hereditary pancreatitis. The major mutations are known as cationic trypsinogen "R122H", "N29I".

Families with HP might carry one of these mutations, but usually not more than one type of mutation. It is also possible that a family with a strong history of HP may not carry any of the currently known mutations. For this reason, it is believed that additional genes and mutations that cause HP are awaiting discovery. (Every concerned person is encouraged to join on going research studies through the University of Pittsburgh 1-888-PITT DNA, or an affiliated medical center)

If I have inherited any of the HP mutations, will I definitely develop pancreatitis?

HP symptoms can develop at any age, but most patients have their first pancreatic attack before the age of 20. In addition, there is a great deal of variability in the frequency and severity of pancreatic attacks. Some affected relatives may only have a few episodes of pain, while others in the same family experience more severe symptoms. The cause for this variation in symptoms is unknown.

Individuals who have inherited either the R122H or the N29I mutation have an 80% risk of developing clinical symptoms of HP over the course of their lifetime. Although the remaining 20% of these mutation carriers do not show any symptoms, it is important to remember that they still have a risk of having a child who inherits their HP mutation and is affected with HP.

Much less is known about the other mutations, and it is believed that are rare in the population. Without additional knowledge about these mutations, it is not possible to predict how many people will eventually develop symptoms of pancreatitis.

What is the risk for my child to inherit HP?

All of our genes come in pairs. We inherit one set of genes from each of our parents. Hereditary pancreatitis is inherited in a dominant manner, which means that only one copy of the HP gene needs to have a mutation in order for an individual to become affected with HP.

When a parent carries an HP mutation in one of their genes, then each child has a 50% (or 1 in 2) chance of inheriting that mutation. It is important to remember that this risk to inherit an HP mutation is 50% for each pregnancy. The risk to have an affected child is actually less than 50% (it is approximately 40%) since one out of five individuals with the R122H or N29I mutation remain symptom-free over their lifetime.

Genetic testing for HP

Genetic testing for HP is currently available both on a research and commercial basis.

Commercial testing is conducted through a licensed laboratory for a specified fee that may be covered by your insurance plan. A small blood sample is drawn at your doctor's office or hospital laboratory and sent to the commercial laboratory for testing. Results are then provided to your referring physician or counsellor. One commercial laboratory that provides testing for HP is Molecular Diagnostics at the University of Pittsburgh (phone: 412-648-8519). Currently, the fee for commercial
testing through Molecular Diagnostics is $276.50 per person. Check with your insurance carrier to determine whether genetic testing is covered by your health plan. Testing can only be done in a laboratory licensed to perform this test. If testing is being done at other institutions within the United States, please call the 888-PITT DNA number to be sure that the site is approved.

Research testing is available for a reduced fee for those who qualify to be enrolled in the HP research study at the University of Pittsburgh. Research testing is confidential, requires the completion of several forms and questionnaires, but may be associated with a longer turnaround period to obtain results.

Genetic testing for any condition is a complex process. Genetic counselors are available in your local area to help identify the potential risks, benefits, and limitations of genetic testing for HP. Referrals to local genetic counselors can be obtained from your primary care physician. To find a genetic counselor near you, ask your physician or call us for assistance at 888-PITT-DNA.

Who might benefit from genetic testing?

Individuals who have a relative with a documented diagnosis of hereditary pancreatitis.

Individuals with a strong history of unexplained abdominal pain that resembles pancreatitis.

See the Genetic Consensus Statement published in Pancreatology 2001;1:405-415, through the Hereditary Pancreatitis Research Study office 888-PITT DNA, or on http://www.pancreas.org/

What are some of the risks, benefits, and limitations of genetic testing?

Before requesting HP genetic test results, we ask that each participant consider the possible benefits and limitations of genetic testing.

Potential benefits of genetic testing:

a) Obtaining a diagnosis of HP. Some individuals with HP have reported difficulty in obtaining a diagnosis or explanation of their abdominal symptoms. A positive test result (i.e. when a person is found to carry an HP mutation) may help validate or prove that an individual has true medical symptoms that are hereditary in nature. In short, physicians can use genetic testing to diagnose HP.

b) Identifying relatives at risk to develop HP. Once an HP mutation has been found in a family, other relatives can be tested to determine whether they inherited the same HP mutation. This type of testing, "pre-symptomatic testing," is conducted for individuals who have not yet developed pancreatitis but are at risk of having inherited an HP mutation from either parent. Individuals who test positive prior to developing symptoms of HP can then be educated about important lifestyle recommendations to reduce the risk of developing future pancreatic disease such as pancreatic cancer. Although pancreatitis cannot be prevented at this time, these individuals would be recommended to avoid the use of tobacco and alcoholic beverages, since these exposures are risk factors for pancreatic cancer in the general population.

c) Reducing feelings such as anxiety or uncertainty. Test results can often help reduce feelings of anxiety and uncertainty for individuals with a family history of HP. For example, an unaffected relative who receives negative test results (i.e. when a person is found NOT to carry an HP mutation) is likely to feel relieved from the uncertainty of whether symptoms will ever develop in the future. Note: negative test results only eliminate the risk to develop a hereditary form of pancreatitis. It is still possible to develop pancreatitis due to other causes later in life.

4. Prenatal Genetic Testing. Genetic testing can provide information that is useful when making
important reproductive decisions. Prenatal diagnosis is a form of testing that is conducted during pregnancy to determine whether a developing baby has inherited a specific mutation or other health problem. Currently prenatal testing is not being conducted. If both members of the couple have tested negative for a particular HP mutation, then the risk to pass on that HP mutation is essentially zero. For example, if your family is known to carry the R122H* mutation and you test negative for R122H*, then none of your children (or future pregnancies) would be at risk to develop HP.

Potential risks & limitations of genetic testing:

1. Genetic discrimination.
   As with genetic testing for any type of condition, there is a small but potential threat for insurance and/or employment discrimination. We have never heard of a documented case of this type of discrimination among our research population of over 700 individuals. Moreover, it is reassuring to know that there are no published cases of insurance discrimination with regard to hereditary pancreatitis. Individuals who undergo genetic testing as part of a research study are generally well protected because of our strict rules of confidentiality. For instance, no information about an HP participant is ever released to a third party without written permission from a participant. All blood samples are identified using a code number instead of personal information such as names or social security numbers. Because no research organization can completely eliminate the risk for discrimination, this potential threat should be considered before requesting your genetic test results are confirmed and released to you.

2. Adverse psychological emotions.
   Powerful emotions such as anxiety, guilt, and depression can accompany the process of genetic testing. In addition, genetic information has a powerful influence on an individualís reproductive behaviour and a lifelong impact on future descendants.

3. Genetic test results cannot prevent or cure HP.
   At this time, there is no cure for HP, nor is there a way to prevent pancreatic attacks in patients who carry HP mutations. Furthermore, there does not appear to be a difference in medical treatment for patients who have a hereditary form of pancreatitis versus patients who have a non-hereditary form of pancreatitis.4. "Non-informative" test results.

   Non-informative test results include any type of test result that is not conclusive in ruling out a hereditary form of pancreatitis. Negative test results must be interpreted very carefully, especially for individuals undergoing pre-symptomatic testing. Pre-symptomatic testing applied to an individual who is not clinically affected with the disease, but has at least one parent affected with a dominant disorder. When test results for know trypsinogen mutations are negative, then . . .

If a mutation has been identified in another family member.
Interpretation: He/she is not at risk to develop the genetic form of pancreatitis that runs in the family. Since he/she does not carry the mutation, his/her children are not at risk either.

If a mutation has not been identified in another family member.
Interpretation: Since we do not know what mutation runs in the family, it is possible that this family carries a mutation that we cannot yet recognize. Therefore it is possible that this person could carry the unrecognizable mutation as well.

Action: Enter a research study with the extended family to search for a new gene mutation.
This information is based on 2001 data for the cationic trypsinogen gene (PRSS1). Disclaimer: This information is intended for general education of individuals who are interested in hereditary pancreatitis and related disorders. It in no way can substitute for the evaluation and care provided to individuals through their trained health care providers. For further information the following publications will be helpful.

Recommended reading (technical)

Chronic Pancreatitis

Trypsinogen Gene Mutations
The trypsinogen R122H and N29I mutations*

Genetic Testing

Cancer Risk

SPINK1 Mutations (not discussed in this document)

CFTR Mutations (not discussed in this document)


* Note: The trypsinogen R122H and N29I mutations were previously numbered R117H and N21I.

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