Direct-to-Consumer Genetic Testing

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What is direct-to-consumer genetic testing?

Most of the time, genetic testing is done through healthcare providers such as physicians, nurse practitioners, and genetic counselors. Healthcare providers determine which test is needed, order the test from a laboratory, collect and send the DNA sample, interpret the test results, and share the results with the patient. Often, a health insurance company covers part or all of the cost of testing.

Direct-to-consumer genetic testing is different: these genetic tests are marketed directly to customers via television, print advertisements, or the Internet, and the tests can be bought online or in stores. Customers send the company a DNA sample and receive their results directly from a secure website or in a written report. Direct-to-consumer genetic testing provides people access to their genetic information without necessarily involving a healthcare provider or health insurance company in the process.

Dozens of companies currently offer direct-to-consumer genetic tests for a variety of purposes. The most popular tests use genetic variations to make predictions about health, provide information about common traits, and offer clues about a person’s ancestry. The number of companies providing direct-to-consumer genetic testing is growing, along with the range of health conditions and traits covered by these tests. Because there is currently little regulation of direct-to-consumer genetic testing services, it is important to assess the quality of available services before pursuing any testing.

Other names for direct-to-consumer genetic testing include DTC genetic testing, direct-access genetic testing, at-home genetic testing, and home DNA testing. Ancestry testing on page 8 (also called genealogy testing) is also considered a form of direct-to-consumer genetic testing.
For more information about direct-to-consumer genetic testing:


National Human Genome Research Institute: Direct-to-Consumer Genomic Testing (https://www.genome.gov/27570940/april-20-directtoconsumer-genomic-testing/)


What kinds of direct-to-consumer genetic tests are available?

With so many companies offering direct-to-consumer genetic testing, it can be challenging to determine which tests will be most informative and helpful to you. When considering testing, think about what you hope to get out of the test. Some direct-to-consumer genetic tests are very specific (such as paternity tests), while other services provide a broad range of health, ancestry, and lifestyle information.

Major types of direct-to-consumer genetic tests include:

**Disease risk and health**

The results of these tests estimate your genetic risk of developing several common diseases, such as celiac disease, Parkinson disease, and Alzheimer disease. Some companies also include a person's carrier status for less common conditions, including cystic fibrosis and sickle cell disease. A carrier is someone who has one copy of a gene mutation that, when present in two copies, causes a genetic disorder. The tests may also look for genetic variations related to other health-related traits, such as weight and metabolism (how a person's body converts the nutrients from food into energy).

**Ancestry or genealogy**

The results of these tests provide clues about where a person's ancestors might have come from, their ethnicity, and genetic connections between families. For more information, see What is genetic ancestry testing? on page 8

**Kinship**

The results of these tests can indicate whether tested individuals are biologically related to one another. For example, kinship testing can establish whether one person is the biological father of another (paternity testing). The results of direct-to-consumer kinship tests, including paternity tests, are usually not admissible in a court of law.

**Lifestyle**

The results of these tests claim to provide information about lifestyle factors, such as nutrition, fitness, weight loss, skincare, sleep, and even your wine preferences, based on variations in your DNA. Many of the companies that offer this kind of testing also sell services, products, or programs that they customize on the basis of your test results.
Before choosing a direct-to-consumer genetic test, find out what kinds of health, ancestry, or other information will be reported to you. Think about whether there is any information you would rather not know. In some cases, you can decline to find out specific information if you tell the company before it delivers your results.

**Learn more about the available types of direct-to-consumer genetic testing:**

Genetics Generation: Types of DTC Testing (http://knowgenetics.org/kinds-of-dtc-testing/)

Michigan State University: Choosing the right DNA test for your needs (http://msue.anr.msu.edu/news/choosing_the_right_dna_test_for_your_needs)


Applied & Translational Genomics (free full-text via PubMed Central): Only a click away — DTC genetics for ancestry, health, love…and more: A view of the business and regulatory landscape (https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4796702/)

What is genetic ancestry testing?

Genetic ancestry testing, or genetic genealogy, is a way for people interested
in family history (genealogy) to go beyond what they can learn from relatives or
from historical documentation. Examination of DNA variations can provide clues
about where a person's ancestors might have come from and about relationships
between families. Certain patterns of genetic variation are often shared among
people of particular backgrounds. The more closely related two individuals,
families, or populations are, the more patterns of variation they typically share.

Three types of genetic ancestry testing are commonly used for genealogy:

Y chromosome testing

Variations in the Y chromosome, passed exclusively from father to son,
can be used to explore ancestry in the direct male line. Y chromosome
testing can only be done on males, because females do not have a Y
chromosome. However, women interested in this type of genetic testing
sometimes recruit a male relative to have the test done. Because the Y
chromosome is passed on in the same pattern as are family names in many
cultures, Y chromosome testing is often used to investigate questions such
as whether two families with the same surname are related.

Mitochondrial DNA testing

This type of testing identifies genetic variations in mitochondrial DNA.
Although most DNA is packaged in chromosomes within the cell nucleus,
cell structures called mitochondria also have a small amount of their
own DNA (known as mitochondrial DNA). Both males and females have
mitochondrial DNA, which is passed on from their mothers, so this type
of testing can be used by either sex. It provides information about the
direct female ancestral line. Mitochondrial DNA testing can be useful for
genealogy because it preserves information about female ancestors that
may be lost from the historical record because of the way surnames are
often passed down.
Single nucleotide polymorphism testing

These tests evaluate large numbers of variations (single nucleotide polymorphisms or SNPs) across a person's entire genome. The results are compared with those of others who have taken the tests to provide an estimate of a person's ethnic background. For example, the pattern of SNPs might indicate that a person's ancestry is approximately 50 percent African, 25 percent European, 20 percent Asian, and 5 percent unknown. Genealogists use this type of test because Y chromosome and mitochondrial DNA test results, which represent only single ancestral lines, do not capture the overall ethnic background of an individual.

Genetic ancestry testing has a number of limitations. Test providers compare individuals' test results to different databases of previous tests, so ethnicity estimates may not be consistent from one provider to another. Also, because most human populations have migrated many times throughout their history and mixed with nearby groups, ethnicity estimates based on genetic testing may differ from an individual's expectations. In ethnic groups with a smaller range of genetic variation due to the group's size and history, most members share many SNPs, and it may be difficult to distinguish people who have a relatively recent common ancestor, such as fourth cousins, from the group as a whole.

Genetic ancestry testing is offered by several companies and organizations. Most companies provide online forums and other services to allow people who have been tested to share and discuss their results with others, which may allow them to discover previously unknown relationships. On a larger scale, combined genetic ancestry test results from many people can be used by scientists to explore the history of populations as they arose, migrated, and mixed with other groups.

For more information about genetic ancestry testing:

The University of Utah provides video tutorials (http://learn.genetics.utah.edu/content/basics/molgen/) on molecular genealogy.

The International Society of Genetic Genealogy (https://isogg.org/) promotes the use of DNA testing in genealogy.

The American Society of Human Genetics (ASHG) developed a position paper on ancestry testing (http://www.ashg.org/pdf/ASHGAncestryTestingStatement_FINAL.pdf).

What are the benefits and risks of direct-to-consumer genetic testing?

Direct-to-consumer genetic testing has both benefits and limitations, although they are somewhat different than those of genetic testing ordered by a healthcare provider.

Benefits

• Direct-to-consumer genetic testing promotes awareness of genetic diseases.
• It provides personalized information about your health, disease risk, and other traits.
• It may help you be more proactive about your health.
• It does not require approval from a healthcare provider or health insurance company.
• It is often less expensive than genetic testing obtained through a healthcare provider.
• DNA sample collection is usually simple and noninvasive, and results are available quickly.
• Your data is added to a large database that can be used to further medical research. Depending on the company, the database may represent up to several million participants.

Risks and limitations

• Tests may not be available for the health conditions or traits that interest you.
• This type of testing cannot tell definitively whether you will or will not get a particular disease.
• Unexpected information that you receive about your health, family relationships, or ancestry may be stressful or upsetting.
• People may make important decisions about disease treatment or prevention based on inaccurate, incomplete, or misunderstood information from their test results.
• There is currently little oversight or regulation of testing companies.
• Unproven or invalid tests can be misleading. There may not be enough scientific evidence to link a particular genetic variation with a given disease or trait.
• Genetic privacy may be compromised if testing companies use your genetic information in an unauthorized way or if your data is stolen.
• The results of genetic testing may impact your ability to obtain life, disability, or long-term care insurance.

Direct-to-consumer genetic testing provides only partial information about your health. Other genetic and environmental factors, lifestyle choices, and family medical history also affect the likelihood of developing many disorders. These factors would be discussed during a consultation with a doctor or genetic counselor, but in many cases they are not addressed when using at-home genetic tests.

**Learn more about the pros and cons of direct-to-consumer genetic testing:**


American College of Medical Genetics: Position statement on direct-to-consumer genetic testing (https://www.acmg.net/docs/ACMG%20Revised%20DTC%20Statement%20AOP%20Dec%202015.pdf)

How do I choose a direct-to-consumer genetic testing company?

If you are interested in direct-to-consumer genetic testing, do some research into the companies that offer these services. Questions that can help you assess the quality and credibility of a testing company include:

- Does the company’s website appear professional? Does the company provide adequate information about the services it offers, including sample reports, pricing, and methodology?
- Does the company have experienced genetics professionals, such as medical geneticists and genetic counselors, on its staff? Does the company offer consultation with a genetics professional if you have questions about your test results?
- Does the company explain which genetic variations it is testing for? Does it include the scientific evidence linking those variations with a particular disease or trait? Are the limitations of the test and the interpretation of results made clear?
- What kind of laboratory does the genetic testing, and is the laboratory inside or outside the United States? Is the laboratory certified or accredited? For example, does the laboratory meet U.S. federal regulatory standards called the Clinical Laboratory Improvement Amendments (CLIA)? Is the test approved by the U.S. Food and Drug Administration (FDA)?
- Does the company indicate how it will protect your privacy and keep your genetic data safe? Does that information include both current privacy practices and what may happen to your genetic data in the future?
- Does the company indicate who will have access to your data and how it may be shared? Does it share or sell their customers’ genetic data for research or other purposes? For some companies, much of their profit comes from selling large amounts of participant data for research and drug development, not from selling individual test kits.

Be sure to read and understand the “fine print” on the company’s website before purchasing a direct-to-consumer genetic test. This detailed information, which is often called the “terms of use” or “terms of service,” is a legally binding agreement between you and the company providing the testing. It spells out what is included and excluded in the service and details your rights and the company’s
responsibilities. If you still have questions, contact the company to get more information before you make a decision about testing.

More information about factors to consider when choosing a direct-to-consumer genetic testing company:


National Human Genome Research Institute: Regulation of Genetic Tests (https://www.genome.gov/10002335/regulation-of-genetic-tests/)

Genetic Alliance: Genetic Testing (http://www.geneticalliance.org/advocacy/policyissues/genetictesting) (including policy issues and regulations)


How is direct-to-consumer genetic testing done?

For most types of direct-to-consumer genetic testing, the process involves:

1. **Purchasing a test**
   Test kits can be purchased online (and are shipped to your home) or at a store. The price of some test kits includes the analysis and interpretation, while in other cases this information is purchased separately.

2. **Collecting the sample**
   Collection of the DNA sample usually involves spitting saliva into a tube or swabbing the inside of your cheek. You then mail the sample as directed by the company. In some cases, you will need to visit a health clinic to have blood drawn.

3. **Analyzing the sample**
   A laboratory will analyze the sample to look for particular genetic variations. The variations included in the test depend on the purpose of the test.

4. **Receiving results**
   In most cases, you will be able to access your results on a secure website. Other test companies share results in the mail or over the phone. The results usually include interpretation of what specific genetic variations may mean for your health or ancestry. In some cases, a genetic counselor or other healthcare provider is available to explain the results and answer questions.

   The test kit will include step-by-step instructions, so be sure you understand them before you begin. If you have questions, contact the company before collecting your sample.

**Learn more about the direct-to-consumer genetic testing process:**

How much does direct-to-consumer genetic testing cost, and is it covered by health insurance?

The price of direct-to-consumer genetic testing ranges from under a hundred dollars to several thousand dollars. The cost depends on how many genetic variations are analyzed (and it will cost more if whole genome or whole exome sequencing is used), how extensive the interpretation of results is, and whether other products, programs, or services are included. Some companies charge separately for the sample collection kit and the analysis, while others offer the sample collection and analysis as part of a package. In some cases, consultation with a healthcare professional (such as a genetic counselor) is included in the cost of testing; in others, it can be added for an additional fee. Before you proceed with testing, make sure you know the total cost for all of the results, support, and other services you expect to receive.

Direct-to-consumer genetic tests, even tests that provide information about health and disease risk, are not covered by most health insurance plans. Because this testing is done without a referral from a healthcare provider and is not considered “diagnostic” (that is, it cannot be used to diagnose any disease or condition), health insurance companies generally will not pay for it. However, if you share your results with your healthcare provider and he or she recommends additional testing or management, that follow-up care may be covered.

Direct-to-consumer genetic tests that are unrelated (or indirectly related) to health, such as ancestry testing and paternity testing, are typically not covered by health insurance plans.

Learn more about the costs of direct-to-consumer genetic testing:

- Genetics Home Reference: What is the cost of genetic testing, and how long does it take to get the results?
- Genetics Home Reference: Will health insurance cover the costs of genetic testing?
What do the results of direct-to-consumer genetic testing mean?

Direct-to-consumer genetic testing can provide interesting information about your health, traits, and ancestry. However, the results may not be as clear-cut as many people assume. Companies that provide these tests often tell their customers that the results are for information, education, and research purposes only—they are not meant to diagnose, prevent, or treat any disease or health condition. It is useful to keep this distinction in mind when interpreting your own test results.

Health and disease risk

The results of these genetic tests provide information about your chance of developing certain diseases and the likelihood that you have particular traits (such as dimples or lactose intolerance). These results are usually based on an analysis of one or more genetic variations that are known or suspected to be associated with the disease or trait.

The results of tests to predict disease risk do not provide a “yes or no” answer about whether a person will develop a given disease. Other factors, including genetic variations that were not tested, environmental factors, and lifestyle choices (such as diet and exercise) also contribute to disease risk in ways that may not be fully understood. Therefore, a result showing an increased risk does not mean you will definitely develop the disease, and a result showing a reduced risk does not mean you will never develop the disease.

Ancestry

The results of these tests give clues about major geographic areas that are your family’s origins. These results are calculated on the basis of genetic variations that are more common in people from certain areas of the world than in others. You may also choose to receive information about individuals who are likely related to you. (These individuals have also undergone testing, and the predictions are based on similarities among DNA sequences.)

Sometimes the results of ancestry testing are unexpected or inconsistent with what a person understands about his or her family history. These tests can uncover previously unknown information about biological relationships among people (such as paternity). People who are closely related, such as siblings, may receive slightly different information about their ancestry because results are limited by the number and diversity of people who have submitted DNA samples to a given direct-to-consumer genetic testing company. It is important...
to be aware that receiving unexpected or ambiguous information about your background or family is a potential risk with this type of testing.

**Lifestyle**

In most cases, direct-to-consumer lifestyle tests assess genetic variations related to very specific traits, such as how your body converts the nutrients from food into energy (metabolism), day/night (circadian) rhythm, or the senses of taste and smell. The company may recommend specific diet or fitness programs, dietary supplements, skincare products, or other products and services on the basis of your results. However, in most cases the link between a given genetic variation and a complex trait like weight, athletic performance, or sleep is indirect or unknown. Therefore the results of these tests can be challenging to interpret, and it can be difficult to predict whether a recommended product or service will be helpful to you.

If you have questions about the meaning of your test results, professional support (such as guidance from a genetic counselor) may be available from the company that provided the test. You can also share questions about your results with your own healthcare provider. Talk to your doctor before making any major changes in managing your health, diet, or fitness after you receive results of a direct-to-consumer genetic test.

**Read more about the issues raised by direct-to-consumer genetic test results:**


Stanford at the Tech: Going From DNA to Health Risk (http://genetics.thetech.org/ask-a-geneticist/how-gwas-works)

What can raw data from a direct-to-consumer genetic test tell me?

In addition to providing various reports and analyses based on your genetics, some direct-to-consumer genetic testing companies make your raw data available to download. The raw data are your genotype—the particular A’s, C’s, T’s, and G’s of your DNA—extracted from the sample you provided. These data are unique to you. Most companies caution that the raw data are only for research or education and are not suitable for medical purposes, such as diagnosing a disease.

It is challenging to interpret raw genotype data on your own. To help with this, several online “third-party interpretation” services offer analysis and interpretation of the raw data collected by another company. Third-party interpretation services can potentially use your genetic data to provide you with more information about your disease risk, traits, and ancestry. However, these services also have some risks and limitations:

- Relatively often, test results include false positives, which means that the service indicates an increased risk of disease when your risk is not actually higher than that of the general population. False positives and other errors can cause stress and anxiety.
- The results may include unexpected or upsetting information about your disease risk or family relationships without any context or guidance.
- The raw data, once you download it and send it by e-mail or store it on your computer, is no longer protected by the original service’s privacy measures.
- There is little regulation of third-party interpretation services.

As with any kind of genetic testing, it is important to assess the credibility of any company you are considering and find out how it protects your privacy before submitting your genetic information. Your healthcare provider can help you understand your results and determine whether any follow-up testing would be useful.
Learn more about the interpretation of raw genetic data:

Kintalk (University of California San Francisco): Third Party Raw DNA Interpretation Services: Stay in the Know (http://kintalk.org/thirdpartyinterpretationservices/)


Can a direct-to-consumer genetic test tell me whether I will develop cancer?

While a direct-to-consumer genetic test can estimate your risk, it cannot tell you for certain whether you will or will not develop certain forms of cancer. Many other factors, including gender, age, diet and exercise, ethnic background, a history of previous cancer, hormonal and reproductive factors, and family history also contribute to a person’s overall cancer risk.

The U.S. Food and Drug Administration (FDA) has allowed at least one direct-to-consumer genetic testing company, 23andMe, to offer a test for cancer risk. The test looks for three specific variations in two genes: *BRCA1* and *BRCA2*. These variations are associated with an increased risk of breast cancer, ovarian cancer, and potentially other forms of cancer in people of Ashkenazi (eastern European) Jewish ancestry.

Researchers estimate that 5 to 10 percent of all cancers run in families. Some of these cancers are associated with inherited mutations in particular genes, such as *BRCA1* or *BRCA2*. More than 1,000 variations in each of these genes have been associated with an increased risk of cancer. However, the FDA-approved direct-to-consumer genetic test analyzes only three of these variations. The variations included in the test are much more common in people of Ashkenazi Jewish background than in people of other ethnicities, so if you are not of Ashkenazi Jewish heritage, the results may not be useful to you.

Because the variants included in the test are uncommon, most people will have a negative result. A negative result does not mean that you will never get cancer. Similarly, a positive result (one that indicates a cancer-related genetic variation) does not mean that you will definitely develop cancer.

Direct-to-consumer genetic testing for cancer risk can be stressful and anxiety-producing. Health professional organizations and patient advocacy groups strongly recommend that people considering genetic testing for *BRCA1* and *BRCA2* variations, including those included in direct-to-consumer genetic tests, talk with a genetic counselor about the reasons they want to undergo testing and what the results could mean for their health.

**Learn more about direct-to-consumer genetic testing for cancer risk:**

U.S. Food and Drug Administration: FDA Authorizes, with Special Controls, Direct-to-Consumer Test that Reports Three Mutations in the BRCA
Breast Cancer Genes (https://www.fda.gov/NewsEvents/Newsroom/PressAnnouncements/ucm599560.htm)


Can a direct-to-consumer genetic test tell me whether I will develop Alzheimer disease?

While a direct-to-consumer genetic test can estimate your risk, it cannot tell you for certain whether you will or will not develop Alzheimer disease. Variations in multiple genes, together with lifestyle factors such as diet and exercise, all play a role in determining a person’s risk.

The U.S. Food and Drug Administration (FDA) has allowed at least one direct-to-consumer genetic testing company, 23andMe, to offer a test for Alzheimer disease risk. The test analyzes a gene called APOE. Certain variations in this gene are associated with the likelihood of developing late-onset Alzheimer disease (the most common form of the condition, which begins after age 65). Specifically, the test allows you to find out how many copies (zero, one, or two) you have of a version of the gene called the e4 allele. People who have zero copies of the e4 allele have the same risk of late-onset Alzheimer disease as the general population. The risk increases with the number of copies of the e4 allele, so people with one copy have an increased chance of developing the disease, and people with two copies have an even greater risk. However, many people who have one or two copies of the e4 allele never develop Alzheimer disease, and many people with no copies of this allele ultimately get the disease.

Variations in the APOE gene are among many factors that influence a person’s overall risk of developing Alzheimer disease. Variations in many other genes, which are not reported in the FDA-approved direct-to-consumer genetic test, also contribute to disease risk. Additionally, there are risk factors for Alzheimer disease that have yet to be discovered. Therefore the APOE e4 allele represents only one piece of your overall Alzheimer disease risk.

Currently, there are no effective approaches for preventing Alzheimer disease, and while the disease can be treated, it has no cure. For these reasons, the National Institute on Aging recommends against genetic testing for Alzheimer disease except in a research setting (such as a clinical trial). Patient advocacy groups strongly recommend that people considering genetic testing for Alzheimer disease, including direct-to-consumer genetic testing, talk with a genetic counselor about the reasons they want to undergo testing and how they would cope with the results.

Learn more about direct-to-consumer genetic testing for Alzheimer disease:

U.S. Food and Drug Administration: FDA Allows Marketing of First Direct-to-Consumer Tests that Provide Genetic Risk Information for Certain Conditions
Direct-to-Consumer Genetic Testing


What does it mean to have Neanderthal or Denisovan DNA?

Several direct-to-consumer genetic testing on page 4 companies report how much DNA a person has inherited from prehistoric humans, such as Neanderthals and Denisovans. This information is generally reported as a percentage that suggests how much DNA an individual has inherited from these ancestors. The percentage of Neanderthal DNA in modern humans is zero or close to zero in people from African populations, and is about 1 to 2 percent in people of European or Asian background. The percentage of Denisovan DNA is highest in the Melanesian population (4 to 6 percent), lower in other Southeast Asian and Pacific Islander populations, and very low or undetectable elsewhere in the world.

Neanderthals were very early (archaic) humans who lived in Europe and Western Asia from about 400,000 years ago until they became extinct about 40,000 years ago. Denisovans are another population of early humans who lived in Asia and were distantly related to Neanderthals. (Much less is known about the Denisovans because scientists have uncovered fewer fossils of these ancient people.) The precise way that modern humans, Neanderthals, and Denisovans are related is still under study. However, research has shown that modern humans overlapped with Neanderthal and Denisovan populations for a period, and that they had children together (interbred). As a result, many people living today have a small amount of genetic material from these distant ancestors.

Scientists have sequenced Neanderthal and Denisovan genomes from fossils discovered in Europe and Asia. This genetic information is helping researchers learn more about these early humans. Determining which areas of the genome are shared with archaic humans, and which areas are different, will also help researchers find out what differentiates modern humans from our closest extinct relatives.

In addition to the percentage of Neanderthal or Denisovan DNA, direct-to-consumer testing reports may include information about a few genetic variants inherited from these ancestors that influence specific traits. Studies have suggested that certain genetic variations inherited from archaic humans may play roles in hair texture, height, sensitivity of the sense of smell, immune responses, adaptations to high altitude, and other characteristics in modern humans. These variations may also influence the risk of developing certain diseases. However, the significance of Neanderthal or Denisovan genetic variants on disease risk is still an area of active study, and most direct-to-consumer test results currently do not include them.
While knowing how much DNA a person has in common with his or her Neanderthal or Denisovan ancestors may be interesting, these data do not provide practical information about a person’s current health or chances of developing particular diseases. Having more or less DNA in common with archaic humans says nothing about how “evolved” a person is, nor does it give any indication of strength or intelligence. For now, knowing which specific genetic variants a person inherited from Neanderthal or Denisovan ancestors provides only limited information about a few physical traits.

**Scientific journal articles for further reading**


**Learn more about the genetics of Neanderthals and Denisovans:**

The Smithsonian's Human Origins Program provides information about the genetics of archaic humans and its relevance to modern humans:

- Homo neanderthalensis (http://humanorigins.si.edu/evidence/human-fossils/species/homo-neanderthalensis)
- Ancient DNA and Neanderthals (http://humanorigins.si.edu/evidence/genetics/ancient-dna-and-neanderthals)
• Interbreeding (http://humanorigins.si.edu/evidence/genetics/ancient-dna-and-neanderthals/interbreeding)
• DNA: Genotypes and Phenotypes (http://humanorigins.si.edu/evidence/genetics/ancient-dna-and-neanderthals/dna-genotypes-and-phenotypes)

A news release about the complete sequencing of the Neanderthal genome (https://www.genome.gov/27539119/2010-release-complete-neanderthal-genome-sequenced/) is available from the National Human Genome Research Institute.

The Max Planck Institute for Evolutionary Anthropology provides information and data about the Denisovan genome (http://www.eva.mpg.de/denisova/index.html).
How do direct-to-consumer genetic testing companies protect their customers’ privacy?

A person’s genetic data represent personal, private health information. If you are considering direct-to-consumer genetic testing, it is important to know how the testing company will protect your information. In particular, you should know how the company will handle your sample (for example, saliva), how it plans to safeguard your data, and whether and how your data will be used for secondary purposes (such as research or advertising).

Most direct-to-consumer genetic testing companies provide detailed information on their websites about their privacy and security practices. This information may be included in a “privacy policy,” “privacy statement,” or “privacy center.” Be sure to read, understand, and agree with this information before you start the testing process. If you have questions, contact the company to get more information.

Questions that can help you assess a company’s privacy practices include:

- What does the company do with your sample once it has completed the analysis? Will the sample be stored, shared, sold, or destroyed?
- Once you take the test, who owns your genetic data?
- How does the company safeguard your genetic data and other personal information that you provide? Is it stored in a database that is protected from unauthorized access? What happens if the database is hacked or otherwise compromised?
- Can you delete your results from the company’s database if you wish?
- Does the company use your information for internal research, advertising, or other secondary purposes?
- Will the company share your genetic data or sell it to pharmaceutical or biotechnology companies, academic institutions, or nonprofit organizations? If so, will the shared data include other information that could identify you (such as your name or date of birth)? For what purposes will your data be used? Will you be informed when your data are shared or sold?
- If you do not want your genetic data shared, sold, or used for research, can you opt out? What happens if you agree to share your information but want to opt out later?
• Will you be notified in the future if the company changes its privacy policies?
• What would happen to your sample and your genetic information if the company is sold or goes out of business?

It is important to remember that your DNA is unique and specific to you. Current technology makes it possible to link a sequence of DNA to a particular individual (“de-anonymize” a DNA sample). So if you agree to allow a company to use or share your DNA sequence, even if they don’t include your name or other easily identifying information, your genetic information may not remain anonymous and your privacy could be at risk.

Learn more about privacy issues related to direct-to-consumer genetic testing:


The Hastings Center: Read the Fine Print Before Sending Your Spit to 23andMe (https://www.thehastingscenter.org/response-to-call-for-essays-read-the-fine-print-before-sending-your-spit-to-23andme-r/)
Can the results of direct-to-consumer genetic testing affect my ability to get insurance?

Passed in 2008, a federal law called the Genetic Information Nondiscrimination Act (GINA) made it illegal for health insurance providers in the United States to use genetic information in decisions about a person's health insurance eligibility or coverage. This means that health insurance companies cannot use the results of a direct-to-consumer genetic test (or any other genetic test) to deny coverage or require you to pay higher premiums. However, GINA does not apply when an employer has fewer than 15 employees, and it does not cover people in the U.S. military or those receiving health benefits through the Veterans Health Administration or Indian Health Service.

GINA does not apply to other forms of insurance, such as disability insurance, long-term care insurance, or life insurance. Companies that offer these policies have the right to request medical information, including the results of any genetic testing, when making decisions about coverage and rates. Some of these companies request information about genetic testing as part of their application process, but others do not. It is unclear whether genetic information, including the results of direct-to-consumer genetic testing, will become a standard part of the risk assessment that insurance companies undertake when making coverage decisions.

You should weigh the possible benefits and risks of direct-to-consumer genetic testing, including potential impacts on insurance eligibility and coverage, before you start the testing process.

Learn more about the implications of direct-to-consumer genetic testing in insurance coverage:

National Human Genome Research Institute: Genetic Discrimination (https://www.genome.gov/10002077/genetic-discrimination/)

Genetic Alliance: Genetic Discrimination (http://www.geneticalliance.org/advocacy/policyissues/geneticdiscrimination)

Wharton Public Policy Podcast: Why Genetic Testing Is a 'Perfect Storm' for Insurers (http://knowledge.wharton.upenn.edu/article/why-genetic-testing-is-a-perfect-storm-for-insurers/)
Where can I read more about the diseases and traits covered in my direct-to-consumer genetic testing report?

Learn more from Genetics Home Reference about the health conditions and traits included in your report. Each plain-language summary provides information about the condition’s major features, frequency, causes, and inheritance. You will also find links to other reputable sources of online health information.

**Health conditions**

- **BRCA1- or BRCA2-related breast cancer and ovarian cancer**
- Age-related macular degeneration
- Alpha-1 antitrypsin deficiency
- Celiac disease
- Glucose-6-phosphate dehydrogenase deficiency, also called G6PD deficiency
- Hereditary hemochromatosis
- Hereditary thrombophilia: prothrombin thrombophilia and factor V Leiden thrombophilia
- Alzheimer disease
- Parkinson disease

**Wellness**

- Lactose intolerance

**Traits**

- Cheek dimples
- Eye color
- Hair texture
- Hair color (light or dark hair)

**Carrier status**

- Autosomal recessive spastic ataxia of Charlevoix-Saguenay, also called ARSACS
- Andermann syndrome, also called agenesis of the corpus callosum with peripheral neuropathy
Polycystic kidney disease
Beta thalassemia
Bloom syndrome
Canavan disease
PMM2-congenital disorder of glycosylation, also called PMM2-CDG
Cystic fibrosis
D-bifunctional protein deficiency
Dihydrolipoamide dehydrogenase deficiency
Familial dysautonomia
Congenital hyperinsulinism, also called familial hyperinsulinism
Fanconi anemia
GRACILE syndrome
Gaucher disease
Glycogen storage disease type I
Hereditary fructose intolerance
Junctional epidermolysis bullosa
Leigh syndrome
Limb-girdle muscular dystrophy
Medium-chain acyl-CoA dehydrogenase deficiency, also called MCAD deficiency
Maple syrup urine disease
Mucolipidosis type IV
CLN5 disease, also called neuronal ceroid lipofuscinosis (CLN5-related)
CLN1 disease, also called neuronal ceroid lipofuscinosis (PPT1-related)
Niemann-Pick disease
Nijmegen breakage syndrome
Nonsyndromic hearing loss
Pendred syndrome
Phenylketonuria
Primary hyperoxaluria
Rhizomelic chondrodysplasia punctata
Sialic acid storage disease, including Salla disease
Sickle cell disease, also called sickle cell anemia
Sjögren-Larsson syndrome
Tay-Sachs disease
Tyrosinemia
Usher syndrome
Zellweger spectrum disorder, also called Zellweger syndrome spectrum

For more help understanding your test results:
How can I find a genetics professional in my area?
What happens during a genetic consultation?