You’ve probably heard a lot about coronavirus testing recently. If you think you have coronavirus disease 2019 (COVID-19) and need a test, contact your health care provider, local pharmacy, or local health department immediately. The FDA has been working around the clock to increase the availability of critical medical products, including tests for the coronavirus, to fight the COVID-19 pandemic. Learn more about the different types of tests and the steps involved.

There are two different types of tests – diagnostic tests and antibody tests.

<table>
<thead>
<tr>
<th>MOLECULAR TEST</th>
<th>ANTIGEN TEST</th>
<th>ANTIBODY TEST</th>
</tr>
</thead>
<tbody>
<tr>
<td>Also known as...</td>
<td>Diagnostic test, viral test, molecular test, nucleic acid amplification test (NAAT), RT-PCR test, LAMP test</td>
<td>Rapid diagnostic test [Some molecular tests are also rapid tests.]</td>
</tr>
<tr>
<td>How the sample is taken...</td>
<td>Nasal or throat swab (most tests) Saliva (a few tests)</td>
<td>Nasal or throat swab</td>
</tr>
<tr>
<td>How long it takes to get results...</td>
<td>Same day [some locations] or up to a week</td>
<td>One hour or less</td>
</tr>
<tr>
<td>Is another test needed...</td>
<td>This test is typically highly accurate and usually does not need to be repeated.</td>
<td>Positive results are usually highly accurate but negative results may need to be confirmed with a molecular test.</td>
</tr>
<tr>
<td>What it shows...</td>
<td>Diagnoses active coronavirus infection</td>
<td>Diagnoses active coronavirus infection</td>
</tr>
<tr>
<td>What it can’t do...</td>
<td>Show if you ever had COVID-19 or were infected with the coronavirus in the past</td>
<td>Definitively rule out active coronavirus infection. Antigen tests are more likely to miss an active coronavirus infection compared to molecular tests. Your health care provider may order a molecular test if your antigen test shows a negative result but you have symptoms of COVID-19.</td>
</tr>
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Types of Diagnostic Tests

There are some new diagnostic tests available with alternative methods and benefits.

- **Rapid, point-of-care** diagnostic tests use a mucus sample from the nose or throat but can be analyzed at the doctor’s office or clinic where the sample is collected and results may be available in minutes. These may be molecular or antigen tests.

- **At-home collection** tests, available only by prescription from a doctor, allow the patient to collect the sample at home and send it directly to the lab for analysis.

- **Saliva tests** allow a patient to spit into a tube rather than get their nose or throat swabbed. Saliva tests may be more comfortable for some people and may be safer for health care workers who can be farther away during the sample collection.

Molecular Tests

Many companies and labs have developed tests to diagnose COVID-19 based on detection of the virus’s genetic material in a sample from the patient’s nose or throat. These steps may change as new technology becomes available, but currently the typical steps in molecular testing are:

1. A doctor, pharmacist, or other health professional orders a COVID-19 test. All currently authorized COVID-19 tests, including those used with a home collection kit, require a prescription or order from a health professional.

2. You or a health care professional use a specialized, swab to collect mucus from your nose or throat.

3. You or a health care professional put the swab in a sterile container and seal it for transport to a lab.

4. During the shipping process, most molecular test swabs must be kept within a certain temperature range so that the test will be accurate. The sample must arrive at the lab within 72 hours.

5. A lab technician mixes chemicals with the swab to extract the genetic material of any virus that may be on the swab.

6. The lab technician uses special chemicals, called primers and probes, and a high-tech machine to conduct several controlled heating and cooling cycles to convert the virus’s RNA into DNA, and then make millions of copies of the DNA. Some tests use only one warming cycle to make copies of the DNA.

7. When DNA binds to specific probes, a special type of light is produced that can be seen by the machine and the test shows a “positive” result for infection with SARS-CoV-2, the virus that causes COVID-19.

The FDA continues to work with test developers to make more coronavirus tests available to more people in the future.

Molecular diagnostic tests that detect the genetic material of the virus are commonly used for diagnosing COVID-19 or active coronavirus infection. But no test is 100% accurate all of the time. Some things that may affect the test’s accuracy include:

- You may have the virus, but the swab might not collect it from your nose or throat.
- The swab or mucus sample may be accidentally contaminated by the virus during collection or analysis.
- The nasal or throat swab may not be kept at the correct temperature before it can be analyzed.
- The chemicals used to extract the virus genetic material and make copies of the virus DNA may not work correctly.
Antigen tests usually provide results diagnosing an active coronavirus infection faster than molecular tests, but antigen tests have a higher chance of missing an active infection. If an antigen test shows a negative result indicating that you do not have an active coronavirus infection, your health care provider may order a molecular test to confirm the result.

Antibody (Serology) tests may provide quick results, but should not be used to diagnose an active infection. Antibody tests only detect antibodies the immune system develops in response to the virus, not the virus. It can take days to several weeks to develop enough antibodies to be detected by a test.

How the Coronavirus Tests May Be Used

Americans rely on the FDA to provide an independent review of medical products, such as drugs, diagnostic tests and other medical devices. During a public health emergency like the COVID-19 pandemic, there is an urgent need for products to diagnose, treat or prevent a medical threat. There are two ways a coronavirus test might be used for this emergency:

1. Emergency Use Authorization (EUA)
   In certain types of emergencies, the FDA can issue an Emergency Use Authorization, or EUA, to provide more timely access to critical medical products that may help during the emergency when there are no adequate, approved, and available options. The EUA process is different than full approval or clearance because in some emergency situations we cannot wait for all of the evidence needed for full FDA approval or clearance. Instead, the FDA evaluates the options very quickly using the evidence that is available, carefully balancing the risks and benefits of the product as we know them, in addition to evaluating other criteria. FDA has issued many EUAs for diagnostic tests and antibody tests.
   To expand the nation’s COVID-19 testing capacity, the FDA also issued a policy guidance offering regulatory flexibility in certain circumstances. The policies in this guidance apply to commercial manufacturers and to laboratories certified under the Clinical Laboratory Improvement Amendments (CLIA) to perform high-complexity testing and who create laboratory-developed tests (LDTs) for coronavirus. In the guidance, the FDA outlines policies for test developers that offer certain types of COVID-19 tests before they receive an EUA for that test, under the circumstances described in the guidance. Under certain policies in the guidance, the test developers validate their test, give notice to FDA, and submit an EUA request within a specified timeframe.

2. State Authorization of LDTs
   The FDA is providing flexibility to states that want to authorize labs certified to conduct high-complexity testings in that state to develop and perform coronavirus testing. Under this policy, the state or territory takes responsibility for the safety and accuracy of COVID-19 testing by laboratories in its state/territory and the lab does not submit an EUA request to the FDA.

The best way to get a coronavirus test is to contact your health care provider. You may also visit your state or local health department’s website to look for the latest local information on testing.

The FDA encourages health care professionals and patients to report adverse events or side effects related to the use of coronavirus tests to the FDA’s MedWatch Safety Information and Adverse Event Reporting Program:

• Complete and submit the report online through the FDA’s MedWatch website.
• Download the form or call 1-800-332-1088 to request a form, then complete and return to the address on the form or submit by fax to 1-800-FDA-0178.