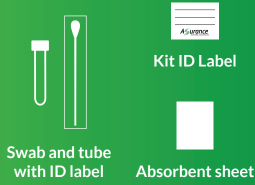


COVID-19 HOME COLLECTION KIT INSTRUCTIONS

INSIDE YOUR KIT:













Poly mailer

Please read all instructions before collecting your sample.

For help with your kit, visit our online Home Collection Video at: AssuranceScientificLabs.com/HomeCollection

1. Register

Register your kit online at AssuranceScientificLabs.com/Register before you collect your sample. The lab is unable to process your sample if you do not register.



2. Arrange for Shipping

Check the return label to identify the shipping carrier either UPS or FedEx. Confirm the nearest drop box location and hours. Ship the sample the same day as collection before the last pickup. Do not ship over the weekend.



3. Fill out Kit ID Label

Remove all contents from the shipping box and bag. With a pen, write your first and last name, date of birth, and date of collection on the sample Kit ID label. Place the sample Kit ID label on the collection tube, making sure the information is easy to read.



4. Prep for Collection

Wash your hands thoroughly with soap and water for at least 20 seconds, then remove the swab from the packaging. Ensure that the tip does not come in contact with any surface.*



5. Sample Collection

- a. Place the swab inside one nostril until it meets resistance (about 1 inch).
- b. Rotate the swab against the nasal wall 3 to 5 times.
- c. Repeat process in second nostril using the same swab.
- d. Place swab in collection tube and break shaft at the breakpoint. (Be careful not to spill the liquid in the tube. If the liquid is spilled, the lab will reject the sample.*)
- e. Leave the swab in the collection tube and screw the lid tightly back on the collection tube. (Ensure the lid is tight. The lab will reject the sample if there is leakage.*)



^{*}If any liquid is spills, leaks, or the swab comes in contact with any surface, please contact ClientServices@ AssuranceScientificLabs.com to order a new kit.



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6. Assemble the Biohazard Bag

Place the absorbent sheet and the collection tube containing your sample in the smaller plastic biohazard bag provided. Ensure the bag is well sealed.



7. Re-Wash Your Hands

Wash your hands thoroughly with soap and water for at least 20 seconds.



8. Assemble the Transport Box

With clean hands, assemble the provided transport box and place the completed biohazard bag into the box. Close the transport box and make sure the content inside is secure.

For help assembling the transport box, please watch our Home Collection Video at: AssuranceScientificLabs.com/HomeCollection.



9. Assemble the Return Mailer

Place the transport box inside the provided poly mailer. Apply the pre-paid shipping label to the poly mailer, ensuring you do not cover up the UN 3373 label. Remove the mailer's cover strip and seal the package closed. Make sure the transport box is secure inside the poly mailer. With the supplied alcohol pad, wipe the outside of the mailer, avoiding the label, before dropping the mailer at the drop box.



10. Ship Your Sample

Immediately after finishing step #9, deliver the package to the appropriate carrier's drop box using the prepaid, pre-addressed return mailer containing your sample. It is important that the sample is shipped on the same day of collection. Ensure you drop your sample off before the last pickup at the drop box location. Do not ship your sample on the weekend.



Rx only

IVD

For use by people 18 years of age or older

This home collection kit has not been FDA cleared or approved.

This home collection kit has been authorized by FDA under an EUA. This home collection kit has been authorized only for the home collection and maintenance of nasal swab specimens as an aid in detection of nucleic acid from SARS-CoV-2, not for any other viruses or pathogens.

This home collection kit in combination with the authorized test is only authorized for the duration of the declaration that circumstances exist justifying the authorization of emergency use of in vitro diagnostic tests for detection and/or diagnosis of COVID-19 under Section 564(b)(1) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. § 360bbb-3(b)(1), unless the authorization is terminated or revoked sooner.