

## SALIVA-BASED COVID-19 TESTING

**CRL RAPID RESPONSE™** is an FDA EUA Approved saliva-based molecular test, ordered at the discretion of a healthcare provider, that is easily self-collected and shipped to our lab. Results of the test confirm if the COVID-19 virus is detected in the saliva of the individual. Since no nasal swab is required, collection is minimally invasive and can be reliably self-administered without the assistance of a healthcare worker.

### 3 EASY COLLECTION STEPS

#### 1. REGISTER KIT

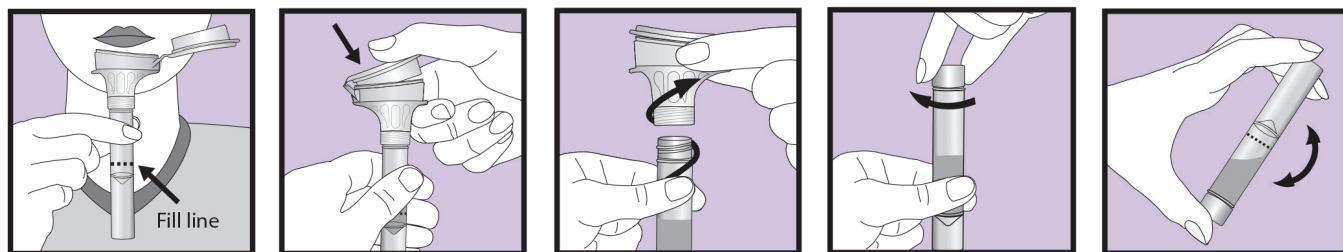
- Register your collection kit at [crlclear.com](http://crlclear.com) and complete a health screening questionnaire.

#### 2. COMPLETE COLLECTION

- Spit into funnel until the amount of liquid (not bubbles) reaches the fill line shown in picture #1.
- Close the funnel lid. The liquid in the lid will be released into the tube to mix with the sample.
- Unscrew the funnel from the tube, and use the small cap to close the tube tightly. Shake for five seconds.

#### 3. RETURN SAMPLE

- Write your first and last name on the barcode label and affix barcode label to the saliva collection device.
- Package the sample and return using the prepaid shipping label.



### FLEXIBLE KIT DISTRIBUTION

- Bulk distribution to specified site(s) for distribution to individuals
- Collection kits sent directly to individuals

### MULTIPLE REPORTING OPTIONS

- Reporting to employer, TPAs, or background screening company
- Web OASIS, WorkForce, HL7, API (XML)
- Reporting to individuals via digital portal



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Emergency Use Authorization (EUA) request for distribution and/or use of the DNA Genotek OMNIgene® ORAL OM-505 saliva collection device for the collection and stabilization of saliva in buffer to transport viral SARS-CoV-2 RNA from patients suspected of COVID-19 by a healthcare provider. The specimen collection device is for use in conjunction with molecular diagnostic testing performed at a clinical laboratory using an in vitro diagnostic (IVD) test for the detection of SARS-CoV-2 that is authorized for use with the home collected kit.