CalREDIE Manual Laboratory Reporting Module (MLRM)

Reporting Antigen Test Results

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Per Title 17 section 2505 of the California Code of Regulations, any entity performing SARS-CoV-2 testing is required to report both positive and non-positive results to public health. Results must be reported to the local health department where the patient resides within eight (8) hours. These laboratory test data are critical for public health disease control and response activities.

Entities performing SARS-CoV-2 point-of-care tests (POC), including PCR and antigen tests are required to report ALL test results. This includes, but is not limited to, the Quidel Sofia SARS Antigen FIA and BD Veritor System.

The CalREDIE Manual Lab Reporting Module (MLRM) can be used to meet these reporting requirements. CalREDIE is the state wide reportable condition reporting system. MLRM allows users to report results for each patient tested directly to the local health department where the patient resides. Entities in San Diego or Los Angeles Counties should not use MLRM and should contact their local health department for instructions on how to report.

Steps to Report via CalREDIE MLRM:

1. Complete the CalREDIE Manual Lab Reporting Account Request Form. Each person who will be reporting results must complete this form.
2. Submit completed account form to calrediehelp@cdph.ca.gov.
3. CalREDIE Helpdesk will process the account and send the user his/her login credentials.
4. Review the CalREDIE Manual Lab Reporting Quick Start Guide. This Guide walks the user through how to log in and submit a result in the MLRM.
5. Login to the CalREDIE MLRM and begin submitting results.
6. Contact calrediehelp@cdph.ca.gov with any questions.