



**Abbott**

Blanchard Valley Health System Site: \_\_\_\_\_ Location: \_\_\_\_\_ Form revised 8/7/2023 LTR

Training / Competency Assessment for Laboratory Point of Care Testing

Date of Hire / Transfer: \_\_\_\_\_

Method: **POCT ID NOW RSV**

Date of Initial Training: \_\_\_\_\_

Test Analyte: **Respiratory Syncytial Virus (RSV) viral RNA**

Associate Name & Title \_\_\_\_\_ Maiden/Former Name: \_\_\_\_\_

(print legibly) Associate ID number: \_\_\_\_\_ Supervisor: \_\_\_\_\_

CLIA Complexity: Waived (2 areas must be assessed) (CAP approved areas 1 – 6) (JC approved areas 1, 3, 5, 6)

\_\_\_ Training assessment after initial training (new hires) and/or new method and/or new test analytes Date: \_\_\_\_\_

\_\_\_ Competency assessment annually Date Due: \_\_\_\_\_

1. \_\_\_ 2. \_\_\_ 3. \_\_\_ **Perform QC testing as required. (Properly running QC under direct observation and recording acceptable test results meets CAP/JC competency requirements for elements 1, 2, 3.)** Record results:

Positive Control: \_\_\_\_\_

Negative Control: \_\_\_\_\_ PASS / FAIL (circle)

4. \_\_\_ **Clean the IDNOW instrument and record on Cleaning Log. (Properly performing instrument maintenance and function checks under direct observation and documenting acceptable results meets CAP/JC competency requirements for element 4.)**

Instrument Serial Number \_\_\_\_\_ Date and Initials to document cleaning \_\_\_\_\_

PASS / FAIL (circle)

5. \_\_\_ **Perform testing on an unknown sample as required. (Analyzing an unknown sample and getting acceptable test results meets CAP/JC competency requirements for element 5.)** Record results.

**UNKNOWN** Test: (Patient or Control or CAP Proficiency sample) \_\_\_\_\_ PASS / FAIL (circle)

Associate Signature: \_\_\_\_\_ Date: \_\_\_\_\_

Signature of Qualified Trainer/Assessor: \_\_\_\_\_ Date: \_\_\_\_\_

Job title of Qualified Trainer/Assessor: \_\_\_\_\_ Satisfactory / Unsatisfactory

6. \_\_\_ **An evaluation of problem solving skills by written exam with 70% passing score will be assigned to meet CAP/JC competency requirements for element 6.**

Assigned: \_\_\_\_\_ Due: \_\_\_\_\_ Completed: \_\_\_\_\_ Score: \_\_\_\_\_ PASS / FAIL (circle)

Reviewed by: \_\_\_\_\_

Date: \_\_\_\_\_

**Note: All annual competency assessment forms are due by September 30.**



## ID NOW™ RSV Quiz

Form revised 8/7/2023 LTR

NAME: \_\_\_\_\_

DATE: \_\_\_\_\_ SCORE: \_\_\_\_\_

**Circle T (True) or F (False) for each Question:**

1. ID NOW RSV can be stored at 2-30°C, but ensure all test components are at room temperature before use. T F
2. To transfer the sample, you should press the white Transfer Cartridge into the blue Sample Receiver until a click is heard. The orange indicator needs to rise to the top of the Transfer Cartridge. T F
3. The white Transfer Cartridge is firmly attached to the orange Test Base by pressing down until the orange indicator descends back down to its starting position. T F
4. Test components can be separated once they are assembled. T F
5. It is acceptable to mix components from different kit lot numbers. T F
6. If the instrument was transported or moved, a performance check using ID NOW™ positive and negative controls is recommended to ensure proper functionality. T F
7. If any assay components are dropped cracked, found to be damaged, or opened when received, they should not be used and should be discarded. T F
8. After a test is completed, discard the components by removing the connected orange Test Base and white Transfer Cartridge and connecting them to the blue Sample Receiver in the ID NOW™ Instrument. Discard the three (3) connected components according to federal, state, and local regulations. T F
9. External positive and negative controls, which are included in the kit, should be tested when an assay is run on the instrument for the first time, once with each new shipment and once for each untrained operator, following a software upgrade, or in order to conform to local, state and/or federal regulations, accrediting groups, or your lab's standard Quality Control procedures. T F
10. Clean the ID NOW™ Instrument daily by spraying with 70% ethanol or 10% bleach. T F
11. Nasopharyngeal swabs may be eluted in saline or approved viral transport media for testing with the ID NOW™ RSV assay. T F
12. Nasopharyngeal swab specimens can be stored at room temperature up to 2 hours prior to testing, or refrigerated at 2-8°C up to 24 hours from time of collection. T F
13. Nasopharyngeal swabs eluted in viral transport media can be stored at room temperature up to 8 hours prior to testing, or refrigerated at 2-8°C up to 24 hours from time of collection. T F
14. Only the nasopharyngeal swabs provided in the kit can be used to collect specimens. T F

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