

## Overview

### Method Name

Quantitative Sandwich Enzyme Immunoassay (EIA)

### NY State Available

No

## Specimen

### Specimen Type

Urine

### Specimen Required

Specimen Type: Urine

Container/Tube: Plastic preservative-free urine container

Specimen Volume: 2 mL

Collection Instructions: Collect 2 mL random urine specimen. Ship specimen refrigerated in a plastic, preservative-free urine container.

**Note:** Sputolysin and Sodium Hydroxide are interfering substances.

### Specimen Minimum Volume

0.5 mL

### Reject Due To

Thawing:	Warm OK; Cold OK
Other:	Specimen that is too viscous to pipette. Tissue, biopsy, sputum, bronchial brush, tracheal aspirate, FNA, bone marrow aspirate, stool or samples in transport media, fixative or Isolator tubes

### Specimen Stability Information

Specimen Type	Temperature	Time	Special Container
Urine	Refrigerated (preferred)	14 days	
	Ambient	48 hours	
	Frozen		

## Clinical & Interpretive

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**Reference Values**

Reference interval: None Detected

Results reported as ng/mL in 0.07 - 8.2 ng/mL range

Results above 8.2 ng/mL are reported as 'Positive, Above the Limit of Quantification'

**Cautions**

Cross-reactions are seen with histoplasmosis, blastomycosis, paracoccidioidomycosis.

Sputolysin, sodium hydroxide, and potassium hydroxide treatment degrade the analyte detected in the assay.

**Performance****PDF Report**

No

**Day(s) Performed**

Monday through Friday

**Report Available**

3 to 5 days

**Performing Laboratory Location**

MiraVista Diagnostics

**Fees & Codes****Fees**

- Authorized users can sign in to [Test Prices](#) for detailed fee information.
- Clients without access to Test Prices can contact [Customer Service](#) 24 hours a day, seven days a week.
- Prospective clients should contact their account representative. For assistance, contact [Customer Service](#).

**Test Classification**

This test was developed and its performance characteristics determined by MiraVista Diagnostics. It has not been cleared or approved by the FDA; however, FDA clearance or approval is not currently required for clinical use. The results are not intended to be used as the sole means for clinical diagnosis or patient management decisions.

**CPT Code Information**

87449

**LOINC® Information**

Test ID	Test Order Name	Order LOINC® Value
FMVCO	MVista Coccidioides Ag, U	Not Provided

Result ID	Test Result Name	Result LOINC® Value
Z2258	MVista Coccidioides Ag, U	93227-7
Z2561	Interpretation	Not Provided