Effective September 10, 2019

NEW FDA-APPROVED TEST FOR DETECTION & QUANTITATION OF CYTOMEGALOVIRUS DNA IN PLASMA

Effective September 10, 2019, Wisconsin Diagnostic Laboratories (WDL) will transition to a new instrument system for Quantitative CMV Nucleic Acid Amplified Testing (CMV NAATQT). This test is traceable to the first WHO International Standard for Human Cytomegalovirus for Nucleic Acid Amplification Techniques (NIBSC 09/162) and reliably monitors CMV viremia.

TESTING SCHEDULE:
- Testing will be performed Monday through Saturday.
- For specimens received prior to 4:00 a.m., results will be available same day, early afternoon.

RANGE OF QUANTITATION:
- The new range of quantitation (Dynamic Range) will be expanded.
  - Current Dynamic Range: 137 – 9,100,000 IU/mL
  - New Dynamic Range: 34.5 – 10,000,000 IU/mL

ORDERING INFORMATION:
- Test ordering will remain the same.

<table>
<thead>
<tr>
<th>Test Name</th>
<th>Mnemonic</th>
<th>Test Code</th>
<th>CPT Code</th>
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</thead>
<tbody>
<tr>
<td>CMV Quantitative Nucleic Acid Amplified Test Quantitative, Plasma</td>
<td>CMV NAATQT</td>
<td>6650018</td>
<td>87497</td>
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FOR QUESTIONS OR ADDITIONAL INFORMATION, PLEASE CONTACT:
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