

Statement of Deficiencies	(X1) Provider/Supplier/CLIA Identification Number 13D1020051	(X3) Date Survey Completed 12/08/2021
Name of Provider or Supplier Rm Lab Llc Db a Express Lab	Street Address, City, State 7988 W Marigold St Ste 100, Boise, ID	
For information on the provider's plan to correct this deficiency, please contact the provider or the state survey agency.		

(X4) ID Prefix Tag	Summary Statement of Deficiencies
D5411	<p>TEST SYSTEMS, EQUIPMENT, INSTRUMENTS, REAGENT CFR(s): 493.1252(a)</p> <p>Test systems must be selected by the laboratory. The testing must be performed following the manufacturer's instructions and in a manner that provides test results within the laboratory's stated performance specifications for each test system as determined under 493.1253.</p> <p>This STANDARD is not met as evidenced by: Based on a review of the Sure-Vue rapid plasma reagin (RPR) package insert, patient test results and quality control (QC) logs and an interview with the general supervisor (GS) on 12/8/2021, the laboratory failed to follow the manufacturer's instructions for Syphilis Serology testing. The findings include: 1. A review of the Sure-Vue RPR package insert, patient test results and QC log identified that the laboratory failed to perform and document an accuracy check on the antigen dispensing needle to ensure that the correct volume of antigen was used as required by the manufacturer. 2. A review of the Sure-Vue RPR package insert, patient test results and QC log identified that the laboratory failed to verify and document that the rotator speed used for testing met the manufacturers requirements. 3. An interview with the GS on 12/8/2021 at 9: 12 am confirmed the above findings. 4. The laboratory reports performing 84 RPR tests annually.</p>
D5451	<p>CONTROL PROCEDURES CFR(s): 493.1256(d)(3)(iii)(g)</p> <p>Unless CMS Approves a procedure, specified in Appendix C of the State Operations Manual (CMS Pub. 7), that provides equivalent quality testing, the laboratory must-- At least once a day patient specimens are assayed or examined perform the following for-- Test procedures producing graded or titered results include a negative control material and a control material with graded or titered reactivity, respectively; 493.1256</p>

(g) The laboratory must document all control procedures performed.

This STANDARD is not met as evidenced by:

Based on a random record review of immunohematology quality control (QC), patient testing logs and an interview with the general supervisor (GS) on 12/8/2021, the laboratory failed to document control material results with graded or titered reactivity and include negative control material for ABO grouping and Rh typing . The findings include: 1. A random record review of immunohematology QC and immunohematology patient testing logs for 2020 and 2021 identified that the laboratory failed to document QC for ABO grouping and Rh typing with a graded or titered reactivity and a negative control on 6/23/2021. 2. ABO grouping and Rh typing was performed and reported on three patients on 6/23/2021. 3. An interview with the GS on 12/8/2021 at 9:27 am confirmed the above findings. 4. The laboratory reports performing 1297 ABO grouping and Rh typing tests annually.