

## **Acknowledgment Letter**

5/4/2020

McKenzie Cato Hyman, Phelps & McNamara, P.C. 700 13th Street NW, Suite 1200 Washington, DC 20005 UNITED STATES

Dear McKenzie Cato:

The Center for Devices and Radiological Health (CDRH) of the Food and Drug Administration (FDA) has received your submission. This submission has been assigned the unique document control number below. All future correspondence regarding this submission should be identified prominently with the number assigned and should be submitted to the Document Control Center at the above letterhead address. Failure to do so may result in processing delays. If you believe the information identified below is incorrect, please notify the Program Operations Staff at (301) 796-5640.

Submission Number: PEUA200955

Received: 5/3/2020

Applicant: Spring Healthcare Services AG

Device: Spring COVID-19 IgM/IgG Rapid Test Cassette

We will notify you when the review of this document has been completed or if any additional information is required. If you are submitting new information about a submission for which we have already made a final decision, please note that your submission will not be re-opened. For information about CDRH review regulations and policies, please refer to <a href="http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/default.htm">http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/default.htm</a>.

Sincerely yours,

Center for Devices and Radiological Health