

July 21, 2021

James A. Hayward, Ph.D. Chairman, President & CEO Applied DNA Sciences, Inc. 50 Health Sciences Drive Stony Brook, NY 11790

Re: EUA200474/S009

Trade/Device Name: Linea COVID-19 Assay Kit

Dated: June 30, 2021 Received: June 30, 2021

Dear Dr. Hayward:

This is to notify you that your request to update the Instructions for se (IFU) a COVID-19 Assay Kit to valuate the include results of the additional post-authorization study to further ytical performance of your product using material representing SARS-CoV-2 sequence val ed to fulfill Condition of Authorization Q. of the May 11, 2021 letter of authorization, and add an associ limitation is granted. Upon review, we concur that the data and information submitted in EUA2004 the requested updates for use with the Linea COVID-19 Assay Kit. In addition, Food and Drug Administration A) have updated the webpage links in the Fact t authorizati Sheet for Healthcare Providers to reflect more rece is. By submitting this EUA revision for review by the FDA, you have complied with the Condition thorization tated in the letter authorizing the emergency use s of X of the Linea COVID-19 Assay Kit reissued on Ma

Si cerely yours,

Uwe Scherf, M.Sc., Ph.D.

Director, Division of Microbiology Devices

OHT7: Office of In Vitro Diagnostics and Radiological Health

Office of Product Evaluation and Quality Center for Devices and Radiological Health