

December 23, 2020

Irma Barr, Principal Regulatory Affairs Specialist Cepheid 904 Caribbean Drive, Sunnyvale, CA 94089

Re: EUA202699/S001

Trade/Device Name: Xpert Omni SARS-CoV-2

Dated: December 11, 2020 Received: December 21, 2020

Dear Irma Barr:

This is to notify you that your request to update the GeneXpert Omni System product labels to include the CE Mark is granted. By submitting these revisions for review by the Food and Drug Administration (FDA), you have complied with the Conditions of Authorization stated in the letter authorizing the emergency use of the Xpert Omni SARS-CoV-2 issued on November 27, 2020.

Sincerely 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