

## **Global Product Development**

09 Aug 2021

Marion Gruber, Ph.D.
Director
Office of Vaccines Research and Review
Food and Drug Administration
Center for Biologics Evaluation and Research
Document Control Center
10903 New Hampshire Avenue
WO71, G112
Silver Spring, MD 20993-0002

THIS DOCUMENT CONTAINS
CONFIDENTIAL AND/OR TRADE SECRET
INFORMATION THAT IS DISCLOSED ONLY
IN CONNECTION WITH THE LICENSING
AND/OR REGISTRATION OF PRODUCTS FOR
PFIZER INC OR ITS AFFILIATED
COMPANIES. THIS DOCUMENT SHOULD
NOT BE DISCLOSED OR USED, IN WHOLE OR
IN PART, FOR ANY OTHER PURPOSE
WITHOUT THE PRIOR WRITTEN CONSENT
OF PFIZER INC.

Re: BLA 125742

**COVID-19 mRNA Vaccine (BNT162/PF-07302048)** 

Response to FDA's 02 August 2021 IR Regarding Validation of Assay Methods and Lot Release

Dear Dr. Gruber,

Reference is made to the Biologics License Application (BLA) submitted 18 May 2021 for the COVID-19 mRNA Vaccine (BNT162/PF-07302048) developed by BioNTech and Pfizer under BB-IND 19736 for the prevention of COVID-19 caused by SARS-CoV-2 in individuals ≥16 years of age.

On 02 Aug 2021, the Agency sent an Information Request regarding validation of assay methods and lot release. The requested information is provided in Response to 02 Aug 2021 FDA Information Request – Assay Methods and Lot Release in Module 1.11.1.

Should you have any questions regarding this submission, or require additional information, please contact me via phone at 215-280-5503; via facsimile at 845-474-3500; or via e-mail at elisa.harkinstull@pfizer.com.

Sincerely,

Elisa Harkins Global Regulatory Lead Global Regulatory Affairs – Vaccines

CC: Ramachandra S. Naik, Ph.D.

CC: Michael Smith, Ph.D. CC: Laura Gottschalk, Ph.D.