From: Gottschalk, Laura

Sent: Wednesday, May 11, 2022 7:52 PM

To: CBER Lot Clearance <cberlotclearance@fda.gov>

Cc: Naik, Ramachandra < Ramachandra. Naik@fda.hhs.gov >; MaguireThon, Meghan

<Meghan.MaguireThon@fda.hhs.gov>; Smith, Michael (CBER) <Michael.Smith2@fda.hhs.gov>

Subject: OVRR: Lot Clearance Request for STN 125742/45

Please confirm that there are no pending lot release issues that would prevent approval of this efficacy supplement.

Applicant Name: BioNTech Manufacturing GmbH (in partnership with Pfizer, Inc.)

Product Names: COVID-19 Vaccine, mRNA (COMIRNATY)

• License Number: 2229

 Address: BioNTech Manufacturing GmbH, An der Goldgrube 12 Mainz, GERMANY

• Application #: 125742/45

• Submission type: Efficacy supplement

- Action Due Date: June 17, 2022, but we may take action by the end of May
- **Summary:** For active immunization to prevent coronavirus disease 2019 (COVID-19) caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) in adolescents 12 through 15 years of age.

Thanks,

Laura

Laura Gottschalk, PhD

Regulatory Project Manager/Primary Reviewer

Center for Biologics Evaluation and Research Office of Vaccines Research and Review U.S. Food and Drug Administration Tel: 301-796-0798 laura.gottschalk@fda hhs.gov