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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

APPLICATION TO MARKET A NEW OR ABBREVIATED NEW DRUG OR BIOLOGIC FOR HUMAN USE

(Title 21, Code of Federal Regulations, Parts 314 & 601)

Form Approved: OMB No. 0910-0338 Expiration Date: March 31, 2020 See PRA Statement on page 3.

1. Date of Submission (mm/dd/yyyy)

05/18/2021

| AF | APPLICANT INFORMATION 2. Name of Applicant BioNTech Manufacturing GmbH | | | | | | | | |
|--|--|---------------------|-----------------------------|----------------|--------------------------|---|--|--|--|
| | Telephone Number (Include country code 9 (0) 6131 9084-7593 | | plicable and area code) | | 4. Facsimile (FAX) N | umber (Include country and area code) +49 (0) 6131 9084-390 | | | |
| 5. | Applicant Address | | | | | | | | |
| | Address 1 (Street address, P.O. box, con An der Goldgrube 12 | npany nam | ne c/o) | | | Email Address | | | |
| | Address 2 (Apartment, suite, unit, building | a. floor. etc | c.) | | | Ruben.Rizzi@biontech.de | | | |
| | () | 3 ,, | - / | | | Applicant DUNS | | | |
| | City Mainz | I | ate/Prov /A | vince/Region | | 117645848 | | | |
| | Country | 14/ | 771 | ZIP or Pos | stal Code | U.S. License Number if previously issued | | | |
| | Germany | | | 55131 | | | | | |
| 6. | 6. Authorized U.S. Agent (Required for non-U.S. applicants) | | | | | | | | |
| Authorized U.S. Agent Name Telephone Number (Include area code) | | | | | | | | | |
| Elisa Harkins, Global Regulatory Lead, Pfizer Globa | | | | ulatory Affa | irs - Vaccines | 215-280-5503 | | | |
| Address 1 (Street address, P.O. box, company name co | | | | | | FAX Number (Include area code) | | | |
| 500 Arcola Road | | | | | | | | | |
| Address 2 (Apartment, suite, unit, building, floor, et | | | | | | 845-474-3500 | | | |
| | | | | | | Email Address | | | |
| City State | | | | | | Elisa.HarkinsTull@pfizer.com | | | |
| | Collegeville | PA | A | | | U.S. Agent DUNS | | | |
| | ZIP Code | | | | | | | | |
| ┕ | 19426 | | | | | | | | |
| PF | ODUCT DESCRIPTION | 7. NDA, A 125742 | ANDA, c | or BLA Appli | cation Number | 8. Supplement Number (If applicable) | | | |
| | Established Name (e.g., proper name, U OVID-19 mRNA Vaccine (nucleoside mo | | name) | | | | | | |
| 10 | Proprietary Name (Trade Name) (If any | | | | | | | | |
| | OMIRNATY | | | | | | | | |
| | Chemical/Biochemical/Blood Product N | | iy) | | | | | | |
| _ | OVID-19 Vaccine (BNT162, PF-07302048 | | Ctrop at | h a | | 14 Davida of Administration | | | |
| | Dosage Form quid | | 13. Strengths 30 mcg | | | 14. Route of Administration Intramuscular | | | |
| | A. Proposed Indication for Use | 301 | | | | | | | |
| ı | • | nd by | IS | s this indicat | ion for a rare disease (| prevalence <200,000 in U.S.)? | | | |
| Active immunization to prevent COVID-19 caused by SARS-CoV-2 in individuals ≥16 years of age | | | Orphan Designation for this | | | If yes, provide the Orphan Designation number for this indication: Continuation Page for #15 | | | |
| | ONOMED OT LABOR. | (1.1 | | | | | | | |
| ı | 3. SNOMED CT Indication Disease Term | | | - | | | | | |
| C | OVID-19; SARS-CoV-2; Disease caused by | by severe a | icute res | spiratory syn | drome coronavirus 2; \$ | SARS-CoV-2 vaccination; COVID-19 vaccination | | | |
| AF | APPLICATION INFORMATION 16. Application Type (Select one) New Drug Application (NDA) Biologics License Application (BLA) Abbreviated New Drug Application (ANDA) | | | | | | | | |
| | | | | | | | | | |
| 19 | If a 351(k), identify the biological referen | nce produc | ct that is | s the basis f | or the submission. | | | | |
| | ame of Biologic: | | | | Holder of Licensed A | pplication: | | | |
| Ь— | If an ANDA, or 505(b)(2), identify the lis | ted drug p | roduct t | that is/are th | | | | | |
| ı | ame of Drug: | arag p | | | | of Relied Upon Product: | | | |
| ı | | Прэ | | D3 | | | | | |
| "' ['] | Indicate Patent Certification: P1 P2 P3 P4 Section viii - MOU Statement of no relevant patents | | | | | | | | |

| | Previous Page Next Page | | | | | | | | |
|-----|---|--|---------------------------|---|---|--|--|--|--|
| 21. | Submission (See instructions) | Labeling Supple upplement Uther (Specif | Postma | CMC Supplement CMC Supplements or CMC Supplement | ☐ Efficacy Supplement ☐ Annual Report Commitments ☐ Periodic Safety Report | | | | |
| 22. | Sub-Type Presubmission Initial Submission | Amendment Resubmission | | 23. If a supplement, identify the appropriate category. CBE Prior Appro | | | | | |
| 24. | For Originals and all Supplements, is the product combination product (21 CFR 3.2(e))? Yes | | | oination Product (See instructions) | Request for Designation (RFD) Number | | | | |
| 25. | Does the submission contain: Only Pediatric data? Yes V No | Status (Select one) luct (Rx) Over-The-Counter Product (OTC) | | | | | | | |
| l | 27. Reasons for Submission Rolling Submission Sequence 0002 for STN/BL 125742 - FINAL ROLL | | | | | | | | |
| 28 | Establishment Information (Full establishment | information shou | ıld be nı | rovided in the body of the | application) | | | | |
| | Establishment Name Pharmacia and Upjohn Company LLC (Pfizer) | | | | | | | | |
| | Address 1 (Street address, P.O. box, company r 7000 Portage Road | | Registration (FEI) Number | | | | | | |
| | Address 2 (Apartment, suite, unit, building, floor | , | | | MF Number | | | | |
| | City Kalamazoo | State/Province/F | | | Establishment DUNS Number 618054084 | | | | |
| | USA USA | ZIP 4900 | 01 | al Code | | | | | |
| | Is the establishment new to the application? | Yes No | | What is the status of the e | | | | | |
| | Establishment Contact Information at the site/fa | acility | | | | | | | |
| | Name of Contact for the Establishment (b) (6) | | | | Telephone Number (Include area code) | | | | |
| (b) | | | | | (b) (6) | | | | |
| | | | | - | FAX Number (Include area code) | | | | |
| | | | | - | (b) (6) Email Address | | | | |
| | | | | - | (b) (6) | | | | |
| | | | | | | | | | |
| | Manufacturing Steps and/or Type of Testing LNP production and bulk drug product formulation, F Drug product testing | ill and finish, Prima | ary packa | aging, Secondary packaging, | Is the site ready very Yes No N/A for inspection? If No, when will site be ready? (mm/dd/yyyy) | | | | |
| | | | | | Continuation Page for #28 | | | | |
| | Cross References (List related BLAs, INDs, NI | | . , | | 1Fs referenced in the current application.) | | | | |
| IN | D 19736, DMF 012683, DMF 9543, DMF 15209, DM | F 011793, DMF 01 | 1820, DN | MF 011321, DMF 10953, | Contin. Page for | | | | |
| 30 | This application contains the following items (\$\frac{5}{2}\$) | Select all that and | o/v) | | #29 | | | | |
| | 1. Index 2. Labeling (Select one) | | | Final Printed Labeling | 3. Summary (21 CFR 314.50 (c)) | | | | |
| | ✓ 4. Chemistry Section ✓ A. Chemistr | y, manufacturing, | and co | ontrols information (e.g., 21 | 1 CFR 314.50(d)(1); 21 CFR 601.2) conly upon FDA's request) | | | | |
| | | | | , 21 CFR 314.50(e)(2)(i); 2 | | | | | |
| | 5. Nonclinical pharmacology and toxicolog (e.g., 21 CFR 314.50(d)(2); 21 CFR 601 | | | | kinetics and bioavailability section .50(d)(3); 21 CFR 601.2) | | | | |
| | 7. Clinical microbiology section (e.g., 21 C. | FR 314.50(d)(4)) | | 8. Clinical data section | on (e.g., 21 CFR 314.50(d)(5); 21 CFR 601.2) | | | | |
| L | Item 30 continued on page 3 | | | | | | | | |

| Previous Page Next Pag | е | | | | | | |
|--|---------------------------|----------------|--------------------|----------------|---|--|---------------|
| 30. This application contains the following | tems (Continued; s | select all tha | at apply) | | | | |
| 9. Safety update report (e.g., 21 CF 21 CFR 601.2) | FR 314.50(d)(5)(vi)(| (b); | 10. Statis | stical section | on (e.g., 21 CFR | 314.50(d)(6); 21 (| CFR 601.2) |
| 11. Case report tabulations (e.g., 2 21 CFR 601.2) | 1 CFR 314.50(f)(1) | ; | 12. Case | report for | ms (e.g., 21 CFR | 314.50 (f)(2); 21 | CFR 601.2) |
| 13. Patent information on any pate biologic (21 U.S.C. 355(b) or (c | | rug/ | | | cation with respect 21 U.S.C. 355 (b)(| t to any patent tha (2) or (j)(2)(A)) | at claims the |
| 15. Establishment description (21 0 | CFR Part 600, if app | plicable) | ☐ 16. Deba | rment cer | tification (FD&C A | Act 306 (k)(1)) | |
| 17. Field copy certification (21 CFF | R 314.50 (I)(3)) | | | | | Form FDA 3397, G 92, or MDUFA Forn | |
| 19. Financial Disclosure Informatio | n <i>(21 CFR Part 54)</i> | | | | | | |
| 20. Other (Specify): | | | | | | | |
| CERTIFICATION I agree to update this application with new safety information about the product that may reasonably affect the statement of contraindications, warnings, precautions, or adverse reactions in the draft labeling. I agree to submit safety update reports as provided for by regulation or as requested by FDA. If this application is approved, I agree to comply with all applicable laws and regulations that apply to approved applications, including, but not limited to, the following: 1. Good manufacturing practice regulations in 21 CFR Parts 210, 211 or applicable regulations, Parts 606, and/or 820. 2. Biological establishment standards in 21 CFR Part 600. 3. Labeling regulations in 21 CFR Parts 201, 606, 610, 660, and/or 809. 4. In the case of a prescription drug or biological product, prescription drug advertising regulations in 21 CFR Part 202. 5. Regulations on making changes in application in FD&C Act section 506A, 21 CFR 314.71, 314.72, 314.97, 314.99, and 601.12. 6. Regulations on Reports in 21 CFR 314.80, 314.81, 600.80, and 600.81. 7. Local, state, and Federal environmental impact laws. If this application applies to a drug product that FDA has proposed for scheduling under the Controlled Substances Act, I agree not to market the product until the Drug Enforcement Administration makes a final scheduling decision. The data and information in this submission have been reviewed and, to the best of my knowledge, are certified to be true and accurate. Warning: A willfully false statement is a criminal offense, U.S. Code, title 18, section 1001. | | | | | | | |
| 31. Typed Name and Title of Applicant's Responsible Official 32. Date (mm/dd/yyyy) | | | | | | | |
| Elisa Harkins, Global Regulatory Lead, Global Regulatory Affairs - Vaccines, Pfizer Inc. 05/18/2021 33. Telephone Number (Include country 34. FAX Number (Include country code if 35. Email Address | | | | | | | |
| 33. Telephone Number (Include country code if applicable and area code) 215-280-5503 | applicable an | , | | | | | |
| 36. Address of Applicant's Responsible Off | | | | 21154.1141 | | ···· | |
| Address 1 (Street address, P.O. box, cor | | | | | | | |
| 500 Arcola Road Address 2 (Apartment, suite, unit, buildin | na. floor. etc.) | | | | | | |
| | | | | | | | |
| City Collegeville | State/Provi | ince/Region | | | | | |
| Country | | ZIP or Pos | ZIP or Postal Code | | | | |
| United States of America | | 19426 | 19426 | | | | |
| 37. Signature of Applicant's Responsible O Other Authorized Official | | Sign | 38. Countersi | gnature of | Authorized U.S. | Agent | Sign |
| Elisa Harkins Digitally signed by Eli DN: o=Pfizer Inc, cn= Reason: I attest to the second | Elisa Harkins Tull | | | | | | |
| Tull integrity of this docum | ent | | | | | | |
| The information | below applies only | y to require | ments of the Pa | perwork F | Reduction Act of 1 | 1995. | |
| The burden time for this collection of information is estimated to average 24 hours per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden to the address to the right: Department of Health and Hu Food and Drug Administration Office of Operations Paperwork Reduction Act (PF PRAStaff@fda.hhs.gov | | | | | | ministration s ion Act (PRA) Staff | |
| "An agency may not conduct or sponsor, an collection of information unless it displays a | | | oond to, a | | | OUR COMPLETED FF EMAIL ADDRE | |

| RST CONTINUATION PAG | GE FOR ITEM 28 – Establisi | nment Information | Provide information for additional establishments below, as needed | |
|------------------------------|--|--------------------|--|--|
| Establishment Name | | | | |
| Pfizer Manufacturing Belgium | | | | |
| Address 1 (Street address, I | Address 1 (Street address, P.O. box, company name c/o) | | | |
| Rijksweg 12 | | | 1000654629 | |
| Address 2 (Apartment, suite | , unit, building, floor, etc.) | | | |
| | | | MF Number | |
| | | | | |
| City | State/Provi | nce/Region | | |
| City Puurs | State/Provi | ince/Region | Establishment DLINS Number | |
| , | | ZIP or Postal Code | Establishment DUNS Number | |
| Puurs | | | Establishment DUNS Number 370156507 | |
| Puurs Country | N/A | ZIP or Postal Code | 370156507 of the establishment? | |

| IRST CONTINUATION PAGE FOR ITEM 20 | - ESTABLIS | innent mi | ormation | establishments below, as needed. | | |
|--|------------------|----------------|---------------------------|--|--|--|
| Establishment Name | | | | | | |
| Pfizer Manufacturing Belgium NV | | | | | | |
| Address 1 (Street address, P.O. box, company | name c/o) | | | Registration (FEI) Number | | |
| Rijksweg 12 | 4- \ | | | 1000654629 | | |
| Address 2 (Apartment, suite, unit, building, floo | r, etc.) | | | MF Number | | |
| City | State/Dray | ince/Regior | | _ | | |
| City Puurs | N/A | ilice/Regioi | ı | | | |
| Country | IV/A | ZIP or Po | stal Code | Establishment DUNS Number | | |
| Belgium | | 2870 | otal code | 370156507 | | |
| Is the establishment new to the application? | | 1 = 0, 0 | What is the status of the | e establishment? | | |
| | / Yes | No | ✓ Pending | ☐ Active ☐ Inactive ☐ Withdrawn | | |
| Establishment Contact Information at the site/ | facility | | | | | |
| Name of Contact for the Establishment | · · · · · | | | Telephone Number (Include area code) | | |
| (b) (6) | | | | (| | |
| (6) | | | | (b) (6) | | |
| | | | | FAX Number (Include area code) | | |
| | | | | AX Number (miciade area code) | | |
| | | | | (b) (6) | | |
| | | | | | | |
| | | | | Email Address | | |
| | | | | (b) (6) | | |
| | | | | (8) (8) | | |
| Manufacturing Steps and/or Type of Testing | | | | Is the site ready Yes No N/A | | |
| LNP production and bulk drug product formulation, | Fill and finish, | , Primary pac | ckaging, Secondary | for inspection? | | |
| packaging, Drug product testing | | | | If No, when will site be ready? (mm/dd/yyyy) | | |
| | | | | ready: (mm/dd/yyyy) | | |
| | | | | | | |
| Establishment Name | | | | | | |
| Wyeth BioPharma Division of Wyeth Pharmaceutica | | | | Desistation (EEI) Number | | |
| Address 1 (Street address, P.O. box, company | name c/o) | | | Registration (FEI) Number | | |
| 1 Burtt Road Address 2 (Apartment, suite, unit, building, floo | r etc) | | | 1222181 | | |
| Address 2 (Apartment, Suite, unit, building, 1100 | i, eic.) | | | MF Number | | |
| City | State/Prov | ince/Regior | า | | | |
| Andover | MA | iiioc/i (cgioi | 1 | E L L II L L E L II L L | | |
| Country | ., | ZIP or Po | stal Code | Establishment DUNS Number | | |
| United States | | 01810 | | 174350868 | | |
| Is the establishment new to the application? | | 1 | What is the status of the | e establishment? | | |
| | / Yes | No | ✓ Pending | Active Inactive Withdrawn | | |
| Establishment Contact Information at the site/ | facility | | | | | |
| Name of Contact for the Establishment | | | | Telephone Number (Include area code) | | |
| (b) (6) | | | | | | |
| (6) | | | | (b) (6) | | |
| | | | | FAX Number (Include area code) | | |
| | | | | | | |
| | | | | (b) (6) | | |
| | | | | Email Address | | |
| | | | | Email Address | | |
| | | | | (b) (6) | | |
| | | | | | | |
| Manufacturing Steps and/or Type of Testing | | | | Is the site ready Yes No N/A | | |
| Manufacture of drug substance, Drug substance testi | ng, Drug prod | uct testing | | for inspection? If No, when will site be | | |
| | | | | ready? (mm/dd/yyyy) | | |
| | | | | | | |
| | | | | Add Second Continuation Page for #28 | | |
| | | _ | | | | |

Remove Continuation Page Return to Form Provide information for additional SECOND CONTINUATION PAGE FOR ITEM 28 – Establishment Information establishments below, as needed.

| | Pfizer Inc | | | | | | | | |
|-------|---|--------------------------------------|--------------|---------------------------|--|--|--|--|--|
| | Address 1 (Street address, P.O. box, company | name c/o) | | | Registration (FEI) Number | | | | |
| | 875 Chesterfield Parkway West | | | | 1940118 | | | | |
| | Address 2 (Apartment, suite, unit, building, floo | r, etc.) | | | | | | | |
| | | | | | MF Number | | | | |
| | City | | ince/Region | ı | | | | | |
| | Chesterfield | MO | T= | | Establishment DUNS Number | | | | |
| | Country | | ZIP or Pos | stal Code | 004954111 | | | | |
| | United States Is the establishment new to the application? | | 63017 | What is the status of the | ootablishmont? | | | | |
| | is the establishment new to the application: | / Yes | No | Pending | Active Inactive Withdrawn | | | | |
| | Establishment Contact Information at the site/ | | | | | | | | |
| | Name of Contact for the Establishment | Telephone Number (Include area code) | | | | | | | |
| | (b) (6) | | | | releptione Number (include area code) | | | | |
| (b) | | (b) (6) | | | | | | | |
| (~) | | | | | FAX Number (Include area code) (b) (6) Email Address | | | | |
| | | | | | | | | | |
| | | | | | | | | | |
| | | | | | | | | | |
| | | | | _ | | | | | |
| | | | | | (b) (6) | | | | |
| | Manufacturing Steps and/or Type of Testing | | | | Is the site ready Yes No N/A | | | | |
| | Drug substance testing, Drug product testing | | | | for inspection? | | | | |
| | | | | | If No, when will site be ready? (mm/dd/yyyy) | | | | |
| ı | | | | | | | | | |
| ı | | | | | | | | | |
| | Establishment Name | | | | | | | | |
| ı | Pfizer Ireland Pharmaceuticals | | | | | | | | |
| ı | Address 1 (Street address, P.O. box, company | name c/o) | | | Registration (FEI) Number | | | | |
| ı | Grange Castle Business Park Clondalkin | | | | 3004145594 MF Number | | | | |
| ı | Address 2 (Apartment, suite, unit, building, floo | r, etc.) | | | | | | | |
| ı | City | State/Broy | rince/Region | , | | | | | |
| ı | Dublin 22 | N/A | ince/Region | ı | | | | | |
| ı | Country | 14/71 | ZIP or Pos | stal Code | Establishment DUNS Number | | | | |
| ı | Ireland | | N/A | | 985586408 | | | | |
| ı | Is the establishment new to the application? | | _ | What is the status of the | establishment? | | | | |
| ı | <u> </u> | / Yes | No | ✓ Pending | Active Inactive Withdrawn | | | | |
| ı | Establishment Contact Information at the site/ | facility | | | | | | | |
| ı | Name of Contact for the Establishment | | | | Telephone Number (Include area code) | | | | |
| ı | (b) (6) | | | | | | | | |
| (b) (| (6) | | | | (b) (6) | | | | |
| | | | | | FAX Number (Include area code) | | | | |
| | | | | | 4 | | | | |
| | | | | | (b) (6) | | | | |
| | | | | | Email Address | | | | |
| | | | | | (I.) (Z) | | | | |
| | | | | | (b) (6) | | | | |
| | Manufacturing Steps and/or Type of Testing | | | | Is the site ready Veg No No NA | | | | |
| | Drug product testing | | | | for inspection? | | | | |
| 1 | | | | | If No, when will site be | | | | |
| 1 | | | | | ready? (mm/dd/yyyy) | | | | |
| 4 | | Add Third Continuation Page for #28 | | | | | | | |

FORM FDA 356h (08/18 - PREVIOUS EDITIONS OBSOLETE)

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| Remove Continuation Page | Return to Form |
|---------------------------|-----------------|
| | |
| HIRD CONTINUATION PAGE FO | R ITEM 28 - Est |

| TH | IIRD CONTINUATION PAGE FOR ITEM 28 | | Provide information for additional establishments below, as needed. | | | | | | |
|-----|---|-------------|---|--|--------|--|----------------|--------------------|------|
| | Establishment Name Hospira Zagrab Ltd. | | | | | | | | |
| | Address 1 (Street address, P.O. box, company | name c/o) | | | Re | egistration | (FEI) Numbe | r | |
| | Prudnicka cesta 60 | | | | _ 30 | 010630287 | | | |
| | Address 2 (Apartment, suite, unit, building, floo | r, etc.) | | | MI | MF Number | | | |
| | City | State/Provi | ince/Region | 1 | | | | | |
| | Prigorje | Brdovecko | | | Fs | stablishmer | nt DUNS Nur | nber | |
| | Country | | ZIP or Pos | stal Code | | 00625201 | | | |
| | Croatia | | 10291 | _ | | | | | |
| | s the establishment new to the application? What is the status of the Yes No Pending | | | | | establishment? Active Inactive Withdrawn | | | |
| | Establishment Contact Information at the site/facility Name of Contact for the Establishment (b) (6) | | | | | | | | |
| | | | | | | lephone N | umber (Inclu | de area code) | |
| | | | | | | | | | |
| (b) | | | | | (t | (b) (6) | | | |
| | | | | | FA | X Number | (Include are | a code) | |
| | | | | | _ (k | o) (6) | | | |
| | | | | | Er | mail Addres | SS | | |
| | | | | | (k | o) (6) | | | |
| | Manufacturing Steps and/or Type of Testing | | | | | Is the site i | ready V | es 🗌 No 🔲 | N/A |
| | Drug Product Release Testing (Sterility) | | | | | for inspect | ion? | es NO | N/A |
| | Brag Froduct Release Testing (Stermity) | | | | | If No, when | n will site be | | |
| | | | | | - | ready? (mi | m/aa/yyyy) | | _ |
| | | | | | | | | | |
| | Establishment Name SGS Lab Simon SA | | | | | | | | |
| | | | | | | | | | |
| | Address 1 (Street address, P.O. box, company name c/o) | | | | Re | egistration | (FEI) Numbe | r | |
| | Vieux Chemin du Poete 10 | | | |]] 30 | 3004186644 | | | |
| | Address 2 (Apartment, suite, unit, building, floor | r, etc.) | | | | | | | |
| | | | | | _ MI | MF Number | | | |
| | City | State/Provi | ince/Region | 1 | | | | | |
| | Wavre | N/A | | | Es | stablishmer | nt DUNS Nur | nber | |
| | Country | | ZIP or Pos | stal Code | 1 28 | 83063907 | | | |
| | Belgium | | 1301 | \\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\ | | | | | |
| | Is the establishment new to the application? | / Yes | No | What is the status of the Pending | e esta | Active [| Inactive | Withdrawn | |
| | Establishment Contact Information at the site/ | facility | | | | | | | |
| | Name of Contact for the Establishment (b) (6) | | | | Те | lephone N | umber (Inclu | de area code) | |
| (b) | | | | | (k | o) (6) | | | |
| | | | | | - FA | X Number | (Include are | a code) | |
| | | | | | _ (k | o) (6) | | | |
| | | | | | Er | mail Addres | SS | | |
| | | | | | (k | o) (6) | | | |
| | Manufacturing Steps and/or Type of Testing | | | | | Is the site i | ready V | es No No | N/A |
| | Drug Product Release Testing (Sterility) | | | | | for inspect | ion? | ES INO I | IN/A |
| | Solution (States) | | | | | | n will site be | | |
| | | | | | - | ready? (mi | n/uu/yyyy) | | _ |
| | | | | | | Add Fo | urth Continu | ation Page for #28 | |