DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

PREMARKET TOBACCO PRODUCT APPLICATION (PMTA) SUBMISSION

Form Approved: OMB No. 0910-0879

Expiration Date: xx/xx/20xx

Paperwork Reduction Act Statement: The Paperwork Reduction Act of 1995 provides that an agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a valid OMB control number. The valid OMB control number for this information collection is 0910-0879. The time required to complete this information collection is estimated to average 35 minutes per response. Send comments regarding this burden estimate or any other aspects of this collection of information, including suggestions for reducing burden to PRAStaff@fda.hhs.gov.

Marketing without a Marketing Granted Order (MGO) is illegal and may be subject to enforcement. Please carefully read the instructions located in the Appendix before completing this form.

SECTION I – APPLICANT IDENTIFICATION

Part A: Applicant Information¹

Complete for either an organization or an individual, NOT both. Organization applicants should complete fields 1–20 only. Individual applicants should complete fields 21–36 only.

| If applicant is an orga | Part A fi | elds 1- | elds 1–20 and proceed to Part B. | | | | | | | | |
|--|---|-----------|----------------------------------|-------------------|---|--------------------|--------|------------------------------------|--------------------------------|------------------------|--|
| 1. Organization Name | | | | 2. Oth | 2. Other Organization Name (if applicable) | | | | | | |
| Organization FDA-Assigned Facility Establishment Identifier (FEI Number) | | | | 4. Org | 4. Organization D&B DUNS® Number | | | | | | |
| 5. Submit Date (mm/dd/yyyy) 6. Street Address | | | | Line 1 | ine 1 7. Street Address Line 2 (Apt., Suite | | | | Line 2 (Apt., Suite, Bldg., #) | | |
| 8. City | 8. City 9. State, Provi | | | | ce, or T | erritory | / | 10. 0 | Country | 11. ZIP or Postal Code | |
| Point of Contact for O | rganizati | ion | | | | | | | | | |
| 12. First Name 13. Middle | | | | Middle | Initial | | 14. L | ast Na | ame | | |
| 15. Generational Suffix | 15. Generational Suffix 16. Professional Suffix | | | | 17. Position Title | | | | | | |
| 18. Phone Number 19. Fax Number | | | umber | 20. Email Address | | | | | | | |
| If applicant is an indiv | idual, co | mplete | Par | t A field | ls 21–3 | 6 and | proce | ed to | Part B. | | |
| 21. First Name | 22. N | liddle Ir | nitial | 23. La | st Name | | | | 24. Submit Date (mm/dd/yyyy) | | |
| 25. Generational Suffix | 26. Professional Suffix | | | | 27. Po | 27. Position Title | | | | | |
| 28. Street Address Line 1 | | | | | 29. S | treet A | Addres | ss Line 2 (Apt., | Suite, Bldg., #) | | |
| 30. City | 30. City 31. State, Provi | | | | nce, or Territory | | | 32. Country 33. ZIP or Postal Code | | | |
| 34. Phone Number | | 35. Fa | ax Nı | umber | | | 36. E | mail <i>F</i> | Address | | |

| Part B: Authorized Representative ¹ or U.S. Agent Information ¹ | | | | | | | | | | | |
|---|-------------------------------|-----------------------------------|----------|------------------|----------------------|-----------|---|------------------------------------|---|--|--|
| ☐ 1. Select if authorized representative or U.S. agent is the same as the applicant identified in Part A. If the same, skip Part B fields 2–18 and proceed to Part C. | | | | | | | | | | | |
| 2. Identify the authorized representative OR a U.S. agent (select one): | | | | | | | | | | | |
| ☐ Authorized representative (responsible official authorized to represent the domestic applicant) | | | | | | | | | | | |
| OR | | | | | | | | | | | |
| ☐ U.S. agent (responsible official who either resides or maintains a place of business in the U.S. who is authorized to represent the foreign applicant) | | | | | | | | | | | |
| Contact Information for t | he Auth | orized Re | present | tative o | r U.S. | Agent | | | | | |
| 3. First Name | | | | | | | | | | | |
| 6. Generational Suffix | 7. Profe | essional S | uffix | 8. Pos | ition T | itle | | | | | |
| 9. Organization Name | 9. Organization Name 10. Stre | | | | eet Address Line 1 1 | | | | 11. Street Address Line 2 (Apt., Suite, Bldg., #) | | |
| 12. City | | 13. State, Province, or Territory | | | | · | 14. Country 15. ZIP or Postal Code | | | | |
| 16. Phone Number | | 17. Fax | Number 1 | | | | 18 | 18. Email Address | | | |
| Part C: Alternate Point of Use the Continuation Page by | | | | | | of contac | ct. | | | | |
| 1. Select alternate: | | | | | | | - | | | | |
| ☐ Applicant ☐ Auth | norized re | epresentat | tive [| J U.S. | agent | | Oth | ner | | | |
| 2. First Name | | | | | | st Name | | | | | |
| 5. Generational Suffix | sional Suffix 7. Position Ti | | | | Title | | | | | | |
| 8. Organization Name 9. Street Ad | | | | Address Line 1 1 | | | 10. Street Address Line 2 (Apt., Suite, Bldg., #) | | | | |
| 11. City 12. State, Province, o | | | | | e, or Territory | | | 13. Country 14. ZIP or Postal Code | | | |
| 15. Phone Number | | 16. Fax | Numbei | r | | | 17 | . Email Address | | | |
| | | | | | | | | | | | |

¹ Required content and format as per §1114.7 (Standard PMTA), 1114.15 (Supplemental PMTA), and 1114.17 (Resubmission).

Continuation Page for Part C

| Part D: Manufacturer Information ¹ | | | | | | | | | | | |
|--|-----------------------------|-----------------------------|----------|---------|-----------|--------------------|------------------------|---|-----------------------------|--------------------------|----------------------------------|
| ☐ Select here if manuf proceed to Part E. | fact | urer is | the sar | ne as | applic | ant ider | ntified | in Part | A. If | the same, skip I | Part D fields 2–20 and |
| Organization Name 3. Organization FDA-Assigned Flapplicable) | | | | | • | | | . Organization D&B DUNS® Number (if applicable) | | | |
| 6. Street Address Line | 1 | | | | | | 7. St | reet Ad | ddres | s Line 2 <i>(Apt., S</i> | uite, Bldg., #) |
| 5. Select here if manufacturer address is the same as applicant address provided in Part A; if the same, skip fields 6–11 and proceed to field 12. | | | | | | | | | | | |
| 8. City | | | 9. Stat | te, Pro | ovince | , or Terr | itory | 10. C | ountr | у | 11. ZIP or Postal Code |
| Point of Contact for M | lanı | ufactu | rer | | | | | | - | | |
| 12. First Name | | 13. Mi | iddle In | itial 1 | 14. Las | st Name |) | | 15. 0 Suffi | Generational x | 16. Professional Suffix |
| 17. Position Title | | 18. Phone Number 19. Fa | | | | 19. Fa | x Number 20. Email Add | | | 20. Email Add | ress |
| Part E: Manufacturer/Packaging/Storage/Control Facility Information Use the Continuation Page button below for each additional site. | | | | | | | | | | | |
| 1. Select type of site: | | | | | | | | | | 1 | |
| ☐ Manufacturer | | Contr | act mai | nufact | urer | | Repa | cker/re | elabel | er 🗌 Other | |
| 2. Organization Name | | 3. Org applic | | on FD | A-Ass | igned F | El Nu | El Number (if 4. Organization D&B DUNS® Nur applicable) | | | 9&B DUNS [®] Number (if |
| , , | | | | | irer/pa | ıckagir | ng/stc | orage/control fac | ility ready for inspection? | | |
| 7. Street Address Line 2 (Apt., Suite, Bldg., #) | | | | | | Suite, Bldg., #) | | | | | |
| 9. City | 10. State, Province, or Tel | | | | e, or Ter | ritory 11. Country | | | у | 12. ZIP or Postal Code | |
| Point of Contact for M | lanı | ufactu | rer/Pac | kagin | g/Sto | rage/Co | ontrol | Facili | ty | | |
| 13. First Name | | . Middle 15. Last Name tial | | | | ime | 16. Su | | | Generational x | 17. Professional Suffix |
| 18. Position Title | | 19. Ph | none Nu | umber | | 20. Fa | x Num | ber | | 21. Email Add | ress |
| | | | | | | | | | | | |

Continuation Page for Part E

SECTION II - NEW TOBACCO PRODUCT INFORMATION¹

Use required Form FDA 4057b – Premarket Tobacco Product Application Grouping Product Submission Spreadsheet to provide new product information. The form is available on the FDA website.

| SECTION III – SUBMISSION INFORMATION | | | | | | | | | | |
|--|--|--|--|--|--|--|--|--|--|--|
| Part A: General Submis | sion Information | | | | | | | | | |
| 1. Identify submission type (select one):1 | | | | | | | | | | |
| ☐ Standard PMTA ☐ Resubmission ☐ Supplemental PMTA | | | | | | | | | | |
| 2. For products that were previously commercially marketed in the United States, provide the product names and corresponding marketing date(s):1 | | | | | | | | | | |
| | | | | | | | | | | |
| | Part B: Cross-Reference Information (Optional) Complete one row per cross-reference submission. Use the Continuation Page button below to list additional cross-references. | | | | | | | | | |
| 1. Cross-Referenced STN | 2. Is the content relevant to all products within this submission? 3. Information and sections to be referenced (e.g., all sections, sections I–IV) | | | | | | | | | |
| | ☐ Yes | | | | | | | | | |
| | ☐ No (list applicable product name[s]): | | | | | | | | | |
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| | | | | | | | | | | |
| | | | | | | | | | | |
| | ☐ Yes | | | | | | | | | |
| | \square No (list applicable product name[s]): | | | | | | | | | |
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| | | | | | | | | | | |
| | ☐ Yes | | | | | | | | | |
| | | | | | | | | | | |
| | ☐ No (list applicable product name[s]): | | | | | | | | | |
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| | | | | | | | | | | |

Continuation Page for Part A

Complete fields 1-5 for each TPMF cross-reference. Use the Continuation Page button below to list additional TPMFs. 1. TPMF Owner 2. TPMF STN (assigned by FDA) 3. Is the content applicable to all ☐ Yes ☐ No (list applicable product name[s]): products within this submission? 4. Information and sections to be referenced (e.g., all sections, sections I-III) 5. Right of reference included1 ☐ Yes ☐ No 1. TPMF Owner 2. TPMF STN (assigned by FDA) 3. Is the content applicable to all ☐ Yes ☐ No (list applicable product name[s]): products within this submission? 4. Information and sections to be referenced (e.g., all sections, sections I-III) 5. Right of reference included¹ ☐ Yes ☐ No 1. TPMF Owner 2. TPMF STN (assigned by FDA) 3. Is the content applicable to all ☐ Yes ☐ No (list applicable product name[s]): products within this submission? 4. Information and sections to be referenced (e.g., all sections, sections I-III) 5. Right of reference included1 ☐ Yes ☐ No

Part C: Referenced Tobacco Product Master File(s) (TPMF) (Optional)

Continuation Page for Part C

Part D: Formal Meetings Held with FDA Pertaining to the New Product(s) (Optional)

Indicate one meeting per row. As needed, enter the STN and meeting held date. Use the Continuation Page button below to list additional meetings.

| 1. Submission STN | 2. Meeting Held Date¹ (mm/dd/yyyy) | 2. Is the meeting relevant to all products within this submission? | | | | | | | |
|-------------------|------------------------------------|--|---|--|--|--|--|--|--|
| | | ☐ Yes | □ No (list applicable product name[s]): | | | | | | |
| | | ☐ Yes | □ No (list applicable product name[s]): | | | | | | |
| | | □ Yes | □ No (list applicable product name[s]): | | | | | | |
| | | ☐ Yes | ☐ No (list applicable product name[s]): | | | | | | |

Continuation Page for Part D

SECTION IV – APPLICATION CONTENTS CHECKLIST

This application contains the following items (select all that apply and indicate file name and location of application content).

| Part A | A: Administrative Content | Part I | D: Scientific Content |
|--------|--|--------|--|
| 1. | ☐ Cover Letter Location: | 1. | ☐ General Information¹ Location: |
| 2. | ☐ Comprehensive Index¹ and Table of Contents¹ Location: | 2. | ☐ Descriptive Information¹ |
| 3. | ☐ English¹ Translations for Non-English Information Location: | 3. | Location: ☐ Product Samples² |
| 4. | ☐ Request for FDA to refer PMTA to TPSAC¹ Location: | | Location: |
| | | 4. | ☐ Statement of Compliance with 21 CFR part 25¹ Location: |
| Part I | B: Labeling and Marketing Plans | - | Currence 1 |
| 1. | ☐ Specimens of all Proposed Labelling¹ Location: | 5. | ☐ Summary¹ Location: |
| 2. | ☐ Description of Marketing Plans¹ | 6. | ☐ Product Formulation¹ Location: |
| 3. | Location: ☐ Statement of Compliance with Applicable | 7. | ☐ Manufacturing¹ |
| | Tobacco Product Standards Location: | 8. | Location: |
| Part (| C: Inspections | 0. | Location: |
| 1. | □ Location and Contact Information for Each Location Subject to Potential Inspection | 9. | ☐ Organized References Location: |
| | Location: | 10. | ☐ Health Risk Investigations¹ Location: |
| | | 11. | ☐ Study Reports¹ Location: |

² FDA generally expects that product samples will be a required part of a PMTA and that an applicant should be prepared to submit them in accordance with FDA instructions.

| SECTION V – STATEMENTS OF COMPLIANCE WITH THE FEDERAL FOOD, DRUG, AND COSMETIC (FD&C) ACT |
|---|
| Provide a brief description of how the PMTA satisfies content requirements of section 910(b)(1) of the FD&C Act in the space below: |
| |
| |
| |
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| |
| 2. Provide a brief description of how marketing the new tobacco product would be appropriate for the protection of publi health as determined with respect to the population as a whole including users and non-users of the tobacco product and taking into account: |
| The increased or decreased likelihood that existing users of tobacco products will stop using such products; and The increased or decreased likelihood that those who do not use tobacco products will |
| start using such products. |
| |
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SECTION VI - CERTIFICATION STATEMENTS

Applications must contain the following certifications, as appropriate for the specific type of PMTA, with the appropriate information inserted, as described in each parenthetical, signed by an authorized representative of the applicant:

- i Certification Statement for Standard PMTAs
- ii Modified Tobacco Product Certification for Supplemental PMTAs
- iii Same Product Certification for Resubmissions
- iv Different Product Certification for Resubmissions
- v Financial Interest and Arrangements of Clinical Investigators Certification Statement

For the following section, insert the name of the authorized representative as identified in Section I Part B or Part C, the name of the organization being represented as identified in Section I Part A, the individual new tobacco product(s), and the name of the previously submitted PMTA product(s). Complete the information for all applications. If you choose to print and wet sign the certification statements, upload them as a separate document from 4057 to maintain the dynamic fields in Adobe and ensure all content is available for FDA to process, read, review, and archive.

| i. Certification Statement for Standard PMTAs: |
|---|
| I, (insert name of responsible official), on behalf of the applicant, (applicant name), hereby certify that the applicant will maintain all records to substantiate |
| the accuracy of this application for the period of time required in 21 CFR 1114.45 and ensure that records remain |
| readily available to FDA upon request. I certify that this information and the accompanying submission are true |
| and correct, that no material fact has been omitted, and that I am authorized to submit this on the applicant's |
| behalf. I understand that under section 1001 of title 18 of the United States Code anyone who knowingly and |
| willfully makes a materially false, fictitious, or fraudulent statement or representation in any matter within the |
| jurisdiction of the executive, legislative, or judicial branch of the Government of the United States is subject to |
| criminal penalties. |
| 4. Circumstance and Data (compatibility) |
| 1. Signature and Date (mm/dd/yyyy) |

| I, (name of responsible official), | on behalf of the applicant, (applicant name), |
|--|--|
| certify that (new | tobacco product name) has a |
| different (describe each modification to the pro | oduct) |
| | acco product(s)) described in (STN |
| | but is otherwise identical to (name of previously |
| submitted PMTA(s)) | I certify that (name of applicant) |
| understands this means there is no other mod | lification to the materials, ingredients, design, composition, |
| heating source, or any other feature of the orig | ginal tobacco product. I also certify that (name of applicant) |
| will maintain all r | records that substantiate the accuracy of this application, and |
| ensure that such records remain readily availa | able to FDA upon request for the period of time required in 21 CFR |
| 1114.45. I certify that this information and the | accompanying submission are true and correct, and that I am |
| authorized to submit this on the applicant's be | half. I understand that under section 1001 of title 18 of the United |
| States Code, anyone who knowingly and willfu | ully makes a materially false, fictitious, or fraudulent statement |
| or representation in any matter within the juris | diction of the executive, legislative, or judicial branch of the |
| Government of the United States is subject to | |
| ignature and Date (mm/dd/yyyy) Same Tobacco Product Certification for R | · |
| ignature and Date (mm/dd/yyyy) | · |
| ignature and Date (mm/dd/yyyy) Same Tobacco Product Certification for R | · |
| ignature and Date (mm/dd/yyyy) Same Tobacco Product Certification for R I, (insert name of responsible official), | Resubmission |
| ignature and Date (mm/dd/yyyy) Same Tobacco Product Certification for R I, (insert name of responsible official),, certify that this s | Resubmission on behalf of (name of applicant) |
| ignature and Date (mm/dd/yyyy) Same Tobacco Product Certification for R I, (insert name of responsible official),, certify that this seresponds to all deficiencies outlined in the main | Resubmission on behalf of (name of applicant) submission for (new tobacco product name) |
| ignature and Date (mm/dd/yyyy) Same Tobacco Product Certification for R I, (insert name of responsible official),, certify that this seresponds to all deficiencies outlined in the managereviously submitted PMTA) | on behalf of (name of applicant) submission for (new tobacco product name) rketing denial order issued in response to (STN of the |
| ignature and Date (mm/dd/yyyy) Same Tobacco Product Certification for R I, (insert name of responsible official),, certify that this seresponds to all deficiencies outlined in the many previously submitted PMTA) herein is identical to the product described in the many previously submitted PMTA) | on behalf of (name of applicant) submission for (new tobacco product name) rketing denial order issued in response to (STN of the and the new tobacco product described |
| ignature and Date (mm/dd/yyyy) Same Tobacco Product Certification for R I, (insert name of responsible official),, certify that this seresponds to all deficiencies outlined in the main previously submitted PMTA) herein is identical to the product described in the applicant) understanding the product described in the applicant) understanding the product described in the applicant) understanding the product described in the applicant of the product descr | on behalf of (name of applicant) submission for (new tobacco product name) rketing denial order issued in response to (STN of the and the new tobacco product described the previously submitted PMTA. I certify that (name of |
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| ignature and Date (mm/dd/yyyy) Same Tobacco Product Certification for R I, (insert name of responsible official),, certify that this seresponds to all deficiencies outlined in the many previously submitted PMTA) herein is identical to the product described in the applicant) understing regions, design, composition, heating sour will maintain all responsition. | on behalf of (name of applicant) submission for (new tobacco product name) rketing denial order issued in response to (STN of the and the new tobacco product described the previously submitted PMTA. I certify that (name of stands this means there is no modification to the materials, rece, or any other feature. I also certify that (name of applicant) records that substantiate the accuracy of this statement, and |
| ignature and Date (mm/dd/yyyy) Same Tobacco Product Certification for R I, (insert name of responsible official),, certify that this series and the main previously submitted PMTA), herein is identical to the product described in the applicant) understing ingredients, design, composition, heating sour will maintain all rensure that such records remain readily available. | on behalf of (name of applicant) submission for (new tobacco product name) rketing denial order issued in response to (STN of the and the new tobacco product described the previously submitted PMTA. I certify that (name of stands this means there is no modification to the materials, rce, or any other feature. I also certify that (name of applicant) |
| ignature and Date (mm/dd/yyyy) Same Tobacco Product Certification for R I, (insert name of responsible official),, certify that this seresponds to all deficiencies outlined in the many previously submitted PMTA) herein is identical to the product described in the applicant) understing redients, design, composition, heating sour will maintain all resoure that such records remain readily available 1114.45. I certify that this information and the applicant in the such records remain readily available 1114.45. I certify that this information and the applicant in the such records remain readily available 1114.45. I certify that this information and the applicant in the such records remain readily available 1114.45. I certify that this information and the applicant in the such records remain readily available 1114.45. I certify that this information and the applicant in the such records remain readily available 1114.45. I certify that this information and the applicant in the such records remain readily available 1114.45. I certify that this information and the applicant in the such records remain readily available 1114.45. I certify that this information and the applicant in the such records remain readily available 1114.45. | on behalf of (name of applicant) submission for (new tobacco product name) rketing denial order issued in response to (STN of the and the new tobacco product described the previously submitted PMTA. I certify that (name of estands this means there is no modification to the materials, rece, or any other feature. I also certify that (name of applicant) records that substantiate the accuracy of this statement, and able to FDA upon request for the period of time required in 21 CFR |
| ignature and Date (mm/dd/yyyy) Same Tobacco Product Certification for R I, (insert name of responsible official),, certify that this seresponds to all deficiencies outlined in the many previously submitted PMTA) herein is identical to the product described in the applicant) undersing redients, design, composition, heating sour will maintain all rensure that such records remain readily availated authorized to submit this on the company's be | on behalf of (name of applicant) submission for (new tobacco product name) rketing denial order issued in response to (STN of the and the new tobacco product described the previously submitted PMTA. I certify that (name of estands this means there is no modification to the materials, rece, or any other feature. I also certify that (name of applicant) records that substantiate the accuracy of this statement, and able to FDA upon request for the period of time required in 21 CFR accompanying submission are true and correct, and that I am |
| ignature and Date (mm/dd/yyyy) Same Tobacco Product Certification for R I, (insert name of responsible official),, certify that this seresponds to all deficiencies outlined in the many previously submitted PMTA) herein is identical to the product described in the applicant) ingredients, design, composition, heating sour will maintain all resource that such records remain readily availated authorized to submit this on the company's be States Code, anyone who knowingly and willful. | con behalf of (name of applicant) submission for (new tobacco product name) rketing denial order issued in response to (STN of the and the new tobacco product described the previously submitted PMTA. I certify that (name of stands this means there is no modification to the materials, rece, or any other feature. I also certify that (name of applicant) records that substantiate the accuracy of this statement, and able to FDA upon request for the period of time required in 21 CFR accompanying submission are true and correct, and that I am shalf. I understand that under section 1001 of title 18 of the United |

| I, (name of responsible official), on behalf of (name of applicant) | iv. Different Tobacco Product Certificat | tion for Resubmission |
|--|---|--|
| responds to all deficiencies outlined in the marketing denial order issued in response to (STN of the previously submitted PMTA) and the new tobacco product described herein has a different (describe each modification to the product) than (name of original tobacco product) described in (STN of the previously submitted PMTA) but is otherwise identical to (name of original tobacco product) described in (STN of the previously submitted PMTA) but is otherwise identical to (name of original tobacco product) described in (STN of the previously submitted PMTA) locatify that (name of applicant) understands this means there is no modification to the materials, ingredients, design, composition, heating source, or any other feature of the original tobacco product except for the (describe each modification to the tobacco product) I also certify that (name of applicant) will maintain all records that substantiate the accuracy of this statement, and ensure that such records remain readily available to FDA upon request for the period of time required in 21 CFR 1114.45. I certify that this information and the accompanying submission are true and correct, and that I am authorized to submit this on the company's behalf. I understand that under section 1001 of title 18 of the United States Code, anyone who knowingly and willfully makes a materially false, fictitious, or fraudulent statement or representation in any matter within the jurisdiction of the executive, legislative, or judicial branch of the Government of the United States is subject to criminal penalties. 1. Signature and Date (mm/dd/yyyy) v. Financial Interest and Arrangements of Clinical Investigators Certification Statement: I, (name of responsible official), certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest and documentation is provided (please specify in the | I, (name of responsible official) | , on behalf of <i>(name of applicant)</i> |
| previously submitted PMTA) and the new tobacco product described herein has a different (describe each modification to the product) than (name of original tobacco product) described in (STN of the previously submitted PMTA) but is otherwise identical to (name of original tobacco product) described in (STN of the previously submitted PMTA) learning for the previously submitted PMTA understands this means there is no modification to the materials, ingredients, design, composition, heating source, or any other feature of the original tobacco product except for the (describe each modification to the tobacco product) I also certify that (name of applicant) will maintain all records that substantiate the accuracy of this statement, and ensure that such records remain readily available to FDA upon request for the period of time required in 21 CFR 1114.45. I certify that this information and the accompanying submission are true and correct, and that I am authorized to submit this on the company's behalf. I understand that under section 1001 of title 18 of the United States Code, anyone who knowingly and willfully makes a materially false, fictitious, or fraudulent statement or representation in any matter within the jurisdiction of the executive, legislative, or judicial branch of the Government of the United States is subject to criminal penalties. 1. Signature and Date (mm/dd/yyyy) 1. V. Financial Interest and Arrangements of Clinical Investigators Certification Statement: 1. (name of responsible official), on behalf of (name of company), certify that there are no financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). 1. No, there are no financial conflicts of interest and documentation is provided (please specify in the | | |
| has a different (describe each modification to the product) | responds to all deficiencies outlined in t | the marketing denial order issued in response to (STN of the |
| has a different (describe each modification to the product) | previously submitted PMTA) | and the new tobacco product described herein |
| | | |
| | original tobacco product) | described in (STN of the previously |
| I certify that (name of applicant) understands this means there is no modification to the materials, ingredients, design, composition, heating source, or any other feature of the original tobacco produce except for the (describe each modification to the tobacco product) I also certify that (name of applicant) will maintain all records that substantiate the accuracy of this statement, and ensure that such records remain readily available to FDA upon request for the period of time required in 21 CFR 1114.45. I certify that this information and the accompanying submission are true and correct, and that I am authorized to submit this on the company's behalf. I understand that under section 1001 of title 18 of the United States Code, anyone who knowingly and willfully makes a materially false, fictitious, or fraudulent statement or representation in any matter within the jurisdiction of the executive, legislative, or judicial branch of the Government of the United States is subject to criminal penalties. 1. Signature and Date (mm/dd/yyyyy) v. Financial Interest and Arrangements of Clinical Investigators Certification Statement: I, (name of responsible official), on behalf of (name of company) , certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). Do, there are no financial conflicts of interest. Yes, there are financial conflicts of interest and documentation is provided (please specify in the | | |
| the materials, ingredients, design, composition, heating source, or any other feature of the original tobacco product except for the (describe each modification to the tobacco product) | described | in (STN of the previously submitted PMTA) |
| except for the (describe each modification to the tobacco product) | I certify that (name of applicant) | understands this means there is no modification to |
| that (name of applicant) will maintain all records that substantiate the accuracy of this statement, and ensure that such records remain readily available to FDA upon request for the period of time required in 21 CFR 1114.45. I certify that this information and the accompanying submission are true and correct, and that I am authorized to submit this on the company's behalf. I understand that under section 1001 of title 18 of the United States Code, anyone who knowingly and willfully makes a materially false, fictitious, or fraudulent statement or representation in any matter within the jurisdiction of the executive, legislative, or judicial branch of the Government of the United States is subject to criminal penalties. 1. Signature and Date (mm/dd/yyyy) v. Financial Interest and Arrangements of Clinical Investigators Certification Statement: I, (name of responsible official), on behalf of (name of company) , certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). Does not be accuracy of the period of time and the accompany of the executive, legislative, or judicial branch of the executive, legislative, or judicial financial to a function of the executive, legislative, or judicial financial to a function of the executive, legislative, or judicial financial to a function of the executive, legislative, or judicial financial to a function of the executive, legislative, or judicial financial to a function of the executive, legislative, or judicial financial to a function of the executive, legislative, or judicial financial to a function of the executive, legisla | the materials, ingredients, design, comp | position, heating source, or any other feature of the original tobacco product, |
| this statement, and ensure that such records remain readily available to FDA upon request for the period of time required in 21 CFR 1114.45. I certify that this information and the accompanying submission are true and correct, and that I am authorized to submit this on the company's behalf. I understand that under section 1001 of title 18 of the United States Code, anyone who knowingly and willfully makes a materially false, fictitious, or fraudulent statement or representation in any matter within the jurisdiction of the executive, legislative, or judicial branch of the Government of the United States is subject to criminal penalties. 1. Signature and Date (mm/dd/yyyy) V. Financial Interest and Arrangements of Clinical Investigators Certification Statement: I, (name of responsible official), on behalf of (name of company) , certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). \[\begin{arrange} No, there are no financial conflicts of interest. \] \[\begin{arrange} Yes, there are financial conflicts of interest and documentation is provided (please specify in the | except for the (describe each modification | ion to the tobacco product) I also certify |
| required in 21 CFR 1114.45. I certify that this information and the accompanying submission are true and correct, and that I am authorized to submit this on the company's behalf. I understand that under section 1001 of title 18 of the United States Code, anyone who knowingly and willfully makes a materially false, fictitious, or fraudulent statement or representation in any matter within the jurisdiction of the executive, legislative, or judicial branch of the Government of the United States is subject to criminal penalties. 1. Signature and Date (mm/dd/yyyyy) v. Financial Interest and Arrangements of Clinical Investigators Certification Statement: I, (name of responsible official), on behalf of (name of company) , certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). \[\begin{arrange} No, there are no financial conflicts of interest. \] \[\begin{arrange} Yes, there are financial conflicts of interest and documentation is provided (please specify in the | that (name of applicant) | will maintain all records that substantiate the accuracy of |
| and that I am authorized to submit this on the company's behalf. I understand that under section 1001 of title 18 of the United States Code, anyone who knowingly and willfully makes a materially false, fictitious, or fraudulent statement or representation in any matter within the jurisdiction of the executive, legislative, or judicial branch of the Government of the United States is subject to criminal penalties. 1. Signature and Date (mm/dd/yyyy) v. Financial Interest and Arrangements of Clinical Investigators Certification Statement: I, (name of responsible official), on behalf of (name of company), certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). No, there are no financial conflicts of interest. Yes, there are financial conflicts of interest and documentation is provided (please specify in the | this statement, and ensure that such rec | cords remain readily available to FDA upon request for the period of time |
| of the United States Code, anyone who knowingly and willfully makes a materially false, fictitious, or fraudulent statement or representation in any matter within the jurisdiction of the executive, legislative, or judicial branch of the Government of the United States is subject to criminal penalties. 1. Signature and Date (mm/dd/yyyy) v. Financial Interest and Arrangements of Clinical Investigators Certification Statement: I, (name of responsible official), on behalf of (name of company), certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). \[\begin{arrange} No, there are no financial conflicts of interest. \] \[\begin{arrange} Yes, there are financial conflicts of interest and documentation is provided (please specify in the | required in 21 CFR 1114.45. I certify that | at this information and the accompanying submission are true and correct, |
| statement or representation in any matter within the jurisdiction of the executive, legislative, or judicial branch of the Government of the United States is subject to criminal penalties. 1. Signature and Date (mm/dd/yyyy) v. Financial Interest and Arrangements of Clinical Investigators Certification Statement: I, (name of responsible official), on behalf of (name of company), certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). \[\begin{arrange} \text{No, there are no financial conflicts of interest.} \] \[Yes, there are financial conflicts of interest and documentation is provided (please specify in the section). | and that I am authorized to submit this o | on the company's behalf. I understand that under section 1001 of title 18 |
| Government of the United States is subject to criminal penalties. 1. Signature and Date (mm/dd/yyyy) v. Financial Interest and Arrangements of Clinical Investigators Certification Statement: I, (name of responsible official), on behalf of (name of company), certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). \[\Boxedom{No, there are no financial conflicts of interest.} Yes, there are financial conflicts of interest and documentation is provided (please specify in the | of the United States Code, anyone who | knowingly and willfully makes a materially false, fictitious, or fraudulent |
| 1. Signature and Date (mm/dd/yyyy) v. Financial Interest and Arrangements of Clinical Investigators Certification Statement: I, (name of responsible official), on behalf of (name of company), certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). \[\begin{arrange} \text{No, there are no financial conflicts of interest.} \] \[\begin{arrange} Yes, there are financial conflicts of interest and documentation is provided (please specify in the | statement or representation in any matt | ter within the jurisdiction of the executive, legislative, or judicial branch of the |
| v. Financial Interest and Arrangements of Clinical Investigators Certification Statement: I, (name of responsible official), on behalf of (name of company), certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). No, there are no financial conflicts of interest. Yes, there are financial conflicts of interest and documentation is provided (please specify in the | Government of the United States is sub | eject to criminal penalties. |
| I, (name of responsible official), on behalf of (name of company), certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). \[\textsquare \text{No, there are no financial conflicts of interest.} \] \[\text{Yes, there are financial conflicts of interest and documentation is provided (please specify in the} \] | 1. Signature and Date (mm/dd/yyyy) | |
| I, (name of responsible official), on behalf of (name of company), certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). \[\begin{align*} \text{No, there are no financial conflicts of interest.} \] \[\begin{align*} Yes, there are financial conflicts of interest and documentation is provided (please specify in the | | |
| , certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). \[\textstyle \text{No, there are no financial conflicts of interest.} \] Yes, there are financial conflicts of interest and documentation is provided (please specify in the | v. Financial Interest and Arrangements | of Clinical Investigators Certification Statement: |
| , certify that there are no financial conflicts of interest or have included documentation fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). □ No, there are no financial conflicts of interest. □ Yes, there are financial conflicts of interest and documentation is provided (please specify in the | | |
| fully disclosing any potential financial conflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). No, there are no financial conflicts of interest. Yes, there are financial conflicts of interest and documentation is provided (please specify in the | , | |
| □ No, there are no financial conflicts of interest. □ Yes, there are financial conflicts of interest and documentation is provided (please specify in the | | |
| ☐ Yes, there are financial conflicts of interest and documentation is provided (please specify in the | fully disclosing any potential financial co | nflicts of interest required by 21 CFR § 1114.7(k)(3)(ii). |
| | ☐ No, there are no financial conflic | cts of interest. |
| table of contents where the documentation is located). | | |
| 1. Signature and Date (mm/dd/yyyy) | 1. Signature and Date (mm/dd/yyyy) | |

SECTION VII – APPENDICES

CONTINUATION PAGES

Appendix A: Alternate Point of Contact Information

| SECTION I, Part C: Alter | nate P | oint c | of Contact Informa | tion (| Optional |) | | | |
|--------------------------|---|--------|------------------------------|--------|---|-------------------|----------------|--|--|
| 1. Select alternate: | | | | | | | | | |
| ☐ Applicant ☐ Author | zed rep | resen | ntative 🔲 U.S. age | ent | ☐ Other | | | | |
| 2. First Name | | | 3. Middle Initial | 4. La | 4. Last Name | | | | |
| | | | | | | | | | |
| 5. Generational Suffix | 5. Generational Suffix 6. Profession | | | | | 7. Position Title | | | |
| 8. Organization Name | | | 9. Street Address L | ine 1 | | | 10. S Bldg. | treet Address Line 2 (Apt., Suite, , #) | |
| 11. City | City 12. Terr | | | 13. C | Country | | | 14. ZIP or Postal Code | |
| 15. Phone Number 16. Fa | | | ax Number | | 17. Email Address | | | s | |
| SECTION I, Part C: Alter | nate P | oint c | of Contact Informa | tion (| Optional |) | | | |
| 1. Select alternate: | | | | | | | | | |
| ☐ Applicant ☐ Author | zed rep | resen | tative 🔲 U.S. age | ent | t 🛘 Other | | | | |
| 2. First Name | | | 3. Middle Initial 4. Last Na | | | st Name | | | |
| 5. Generational Suffix | Suffix 6. Professional Suffix | | | | | 7. Position Title | | | |
| 8. Organization Name | 9. Street Address L | ine 1 | | | 10. Street Address Line 2 (Apt., Suite, Bldg., #) | | | | |
| 11. City | 1. City 12. State, Province, or Territory | | | | Country | | | 14. ZIP or Postal Code | |
| 15. Phone Number | 15. Phone Number 16. Fax Number | | | | | 17. Email Address | | | |

Appendix B: Manufacturer Information

| Section I Part E: Man | ufa | cturer/Pack | aging/Stora | ge/Cor | ntrol F | acilit | y Info | rmation | | |
|---|------------------------------------|---|-----------------|---------------------------------|-------------------|---------------------------------------|-------------------|--|-----------------------------|--|
| 1. Select type of site: | | | | | | | | | | |
| ☐ Manufacturer ☐ Contract manufacturer ☐ Repacker/relabeler ☐ Other | | | | | | · · · · · · · · · · · · · · · · · · · | | | | |
| Organization Name 3. Organization F applicable) | | | | , | | | | 4. Organization D&B DUNS® Number (if applicable) | | |
| 5. Division Name (if app | 6. Is the ma | 6. Is the manufacturer/packaging/storage/control facility ready for inspection? ☐ Yes ☐ No | | | | | | | | |
| 7. Street Address Line | 1 | | | | 78. S | Street A | Addres | ss Line 2 (Apt., | Suite, Bldg., #) | |
| 9. City | | 10. 5 | State, Province | e, or Ter | ritory | 11. C | ountry | / | 12. ZIP or Postal Code | |
| Point of Contact for M | lan | ufacturer/Pa | ckaging/Sto | rage/Co | ontrol | Facili | ity | | | |
| 13. First Name | 14 Ini | . Middle tial | ame | | | 16. G Suffix | Generational K | 17. Professional Suffix | | |
| 18. Position Title | on Title 19. Phone i | | | lumber 20. Fax Number | | | | 21. Email Address | | |
| Section I Part E: Man | ufa | cturer/Pack | aging/Stora | ge/Cor | ntrol F | acilit | y Info | rmation | | |
| 1. Select type of site: | | | | ' | | | | | | |
| ☐ Manufacturer | | Contract m | anufacturer | | Repa | cker/re | elabele | er 🗌 Other | - | |
| 2. Organization Name | | 3. Organiza applicable) | tion FDA-Ass | ion FDA-Assigned FEI Number (if | | | | 4. Organization D&B DUNS® Number (if applicable) | | |
| 5. Division Name (if app | olica | able) | 6. Is the ma | anufactu | ırer/pa | ckagii | ng/sto | rage/control fac | ility ready for inspection? | |
| | ☐ Yes ☐ No | | | | | | | | | |
| 7. Street Address Line 1 | | | | | 78. S | Street A | Addres | dress Line 2 (Apt., Suite, Bldg., #) | | |
| 9. City | 9. City 10. State, Province, or Te | | | | Territory 11. Cou | | | / | 12. ZIP or Postal Code | |
| Point of Contact for M | lan | ufacturer/Pa | ckaging/Sto | rage/Co | ontrol | Facili | ity | | | |
| 13. First Name | 14. Middle 15. Last Name Initial | | | | 16 Si | | | Generational K | 17. Professional Suffix | |
| 18. Position Title | | 19. Phone Number 20. Fax | | | | nber | • | 21. Email Address | | |

Appendix C: Cross-Referenced Information

| 1. Cross-Referenced STN | 2. Is the content relevant to all products within this submission? | 3. Information and sections to be referenced (e.g., all sections, sections I–IV) |
|-------------------------|--|--|
| | ☐ Yes | |
| | ☐ No (list applicable product name[s]): | |
| | | |
| | ☐ Yes | |
| | ☐ No (list applicable product name[s]): | |
| | | |
| | ☐ Yes | |
| | ☐ No (list applicable product name[s]): | |
| | | |
| | ☐ Yes | |
| | ☐ No (list applicable product name[s]): | |
| | | |
| | ☐ Yes | |
| | ☐ No (list applicable product name[s]): | |
| | | |
| | ☐ Yes | |
| | ☐ No (list applicable product name[s]): | |
| | | |
| | ☐ Yes | |
| | ☐ No (list applicable product name[s]): | |
| | | |
| | | |

Appendix D: Referenced Tobacco Product Master File(s) (TPMF)

| 1. TPMF Owner | | | 2. TPMF STN (assigned by FDA) |
|---|-------|------------|-------------------------------|
| 3. Is the content applicable to all products within this submission? | ☐ Yes | □ No (list | applicable product name[s]): |
| 4. Information and sections to be referenced (e.g., all sections, sections I–III) | | | |
| 5. Right of reference included | ☐ Yes | □ No | |
| 1. TPMF Owner | | | 2. TPMF STN (assigned by FDA) |
| 3. Is the content applicable to all products within this submission? | ☐ Yes | ☐ No (list | applicable product name[s]): |
| 4. Information and sections to be referenced (e.g., all sections, sections I–III) | | | |
| 5. Right of reference included | ☐ Yes | □ No | |
| 1. TPMF Owner | | | 2. TPMF STN (assigned by FDA) |
| 3. Is the content applicable to all products within this submission? | ☐ Yes | □ No (list | applicable product name[s]): |
| Information and sections to be referenced (e.g., all sections, sections I–III)) | | | |
| 5. Right of reference included | ☐ Yes | □ No | |

| 1. TPMF Owner | | | 2. TPMF STN (assigned by FDA) |
|--|-------|------------|-------------------------------|
| 3. Is the content applicable to all products within this submission? | ☐ Yes | □ No (list | applicable product name[s]): |
| 4. Information and sections to be referenced (e.g., all sections, sections I–III) | | | |
| 5. Right of reference included | ☐ Yes | □ No | |
| 1. TPMF Owner | | | 2. TPMF STN (assigned by FDA) |
| 3. Is the content applicable to all products within this submission? | ☐ Yes | ☐ No (list | applicable product name[s]): |
| 4. Information and sections to be referenced (e.g., all sections, sections I–III)) | | | |
| 5. Right of reference included ¹ | ☐ Yes | □ No | |
| 1. TPMF Owner | | | 2. TPMF STN (assigned by FDA) |
| 3. Is the content applicable to all products within this submission? | ☐ Yes | ☐ No (list | applicable product name[s]): |
| 4. Information and sections to be referenced (e.g., all sections, sections I–III) | | | |
| 5. Right of reference included | ☐ Yes | □ No | |
| 1. TPMF Owner | | | 2. TPMF STN (assigned by FDA) |
| 3. Is the content applicable to all products within this submission? | ☐ Yes | ☐ No (list | applicable product name[s]): |
| 4. Information and sections to be referenced (e.g., all sections, sections I–III)) | | | |
| 5. Right of reference included | ☐ Yes | □ No | |

Appendix E: Formal Meetings Held with FDA Pertaining to the New Product(s)²

| 1. Submission STN | 2. Meeting Held Date (mm/dd/yyyy) | 3. Is the content relevant to all products within this submission? | |
|-------------------|-----------------------------------|--|---|
| | | ☐ Yes | ☐ No (list applicable product name[s]): |
| | | ☐ Yes | □ No (list applicable product name[s]): |
| | | ☐ Yes | ☐ No (list applicable product name[s]): |
| | | ☐ Yes | ☐ No (list applicable product name[s]): |
| | | ☐ Yes | ☐ No (list applicable product name[s]): |

Appendix F: Instructions for Completion of PMTA Form

This form and the instructions for use are solely intended to provide the applicant an organized format to supply information required for submission of a Premarket Tobacco Product Application (PMTA).

Form FDA 4057 - Premarket Tobacco Product Application (PMTA) Submission is a required form for applicants to use when submitting a PMTA to FDA. The numbered items in the below instructions correspond to those provided on the form. Prior to submitting to FDA, ensure all information entered in each field is readable after saving. For more information on what to include in a PMTA submission, see 21 CFR § 1114.7

SECTION I — APPLICANT IDENTIFICATION

Section I should include information regarding the identity of the applicant, including the following parts:

- Part A: Applicant Information
- Part B: Authorized Representative or U.S. Agent Information
- Part C: Alternate Point of Contact Information
- Part D: Manufacturer Information
- Part E: Manufacturer/Packaging/Storage/Control Facility Information

Part A: Applicant Information

Part A should include information regarding the applicant for the submission. An applicant may be a manufacturer or importer that submits an SE Report to receive marketing authorization for a new tobacco product. Part A should be completed for either an organization applicant or an individual applicant, NOT both.

Note: Organization applicants should complete fields 1–20 only. Individual applicants should complete fields 21–36 only.

If applicant is an organization, complete Part A fields 1-20 and proceed to Part B.

For these fields, provide the following information for the organization:

- **I.A.1.** The organization name is the party who takes responsibility for and initiates the submission of an SE application to FDA. The legal name of the organization may be an individual or company name (private or otherwise) and should match the applicant's Dun & Bradstreet Data Universal Numbering System D-U-N-S® (D&B DUNS®) number.
- **I.A.2.** All other names the applicant operates under (e.g., any "Doing Business As" [D.B.A.]), if applicable.
- **I.A.3.** The organization FDA-assigned Facility Establishment Identifier (FEI) number, if applicable. *Note:* To obtain or retrieve an FEI number, applicants can use the FEI Search Portal at https://www.accessdata.fda.gov/scripts/feiportal/index.cfm?action=portal.login, and/or contact FEI Search Portal support at feiportal@fda.hhs.gov for any FEI number-related questions.
- **I.A.4.** The organization D&B DUNS number, if applicable. *Note:* To obtain or retrieve a DUNS number, applicants can contact Dun and Bradstreet directly by phone at 1.866.705.5711 or through their website at https://www.dnb.com/duns/get-a-duns.html.
- **I.A.5.** The submit date, or date you are formally submitting the application to FDA (e.g., submitting via the CTP Portal, the FDA Electronic Submissions Gateway [ESG], or handed to courier).
- **I.A.6.** The street address for the organization (including street number, street name and type, suffix direction, etc.). The street address cannot be a P.O. Box.
- **I.A.7.** Additional street address information for the individual applicant (including apartment, suite, building number, #) that you were not able to include in I.A.6.
- **I.A.8.** The city of the individual applicant.
- **I.A.9.** The state, province, or territory of the individual applicant.
- **I.A.10.** The country of the organization location.
- **I.A.11.** The ZIP or postal code of the organization location.

Point of Contact for Organization (only complete if applicant is an organization)

- **I.A.12.** The first name of the organization point of contact.
- **I.A.13.** The middle initial of the organization point of contact, if applicable.
- **I.A.14.** The last name of the organization point of contact.

- I.A.15. The generational suffix (e.g., Jr., III) of the organization point of contact, if applicable.
- **I.A.16.** The professional suffix (e.g., M.D., Ph.D.) of the organization point of contact, if applicable.
- **I.A.17.** The professional position title of the organization point of contact.
- I.A.18. The phone number of the organization point of contact (include country code, if applicable, and area code).
- **I.A.19.** The fax number of the organization point of contact, if applicable (include country code, if applicable, and area code).
- **I.A.20.** The email address of the organization point of contact.

If applicant is an individual, complete Part A fields 21–36 and proceed to Part B.

For these fields, provide the following information for the individual applicant:

- **I.A.21.** The first name of the individual applicant.
- **I.A.22.** The middle initial of the individual applicant, if applicable.
- **I.A.23.** The last name of the individual applicant.
- **I.A.24.** The submit date, or date you are formally submitting the application to FDA (e.g., submitting via the CTP Portal, the FDA Electronic Submissions Gateway [ESG], or handed to courier).
- **I.A.25.** The generational suffix (e.g., Jr., III) for the individual applicant, if applicable.
- I.A.26. The professional suffix (e.g., M.D., Ph.D.) for the individual applicant, if applicable.
- **I.A.27.** The professional position title of the individual applicant.
- **I.A.28.** The street address for the individual applicant (including street number, street name and type, suffix direction, etc.). The street address cannot be a P.O. Box.
- **I.A.29.** Additional street address information for the individual applicant (including apartment, suite, building number, #) that you were not able to include in I.A.28.
- I.A.30. The city of the individual applicant.
- **I.A.31.** The state, province, or territory of the individual applicant.
- **I.A.32.** The country of the individual applicant.
- **I.A.33.** The ZIP or postal code of the individual applicant.
- **I.A.34.** The phone number of the individual applicant (include country code, if applicable, and area code).
- I.A.35. The fax number of the individual applicant, if applicable (include country code, if applicable, and area code).
- **I.A.36.** The email address of the individual applicant.

Part B: Contact Information for the Authorized Representative or U.S. Agent

Part B should include information for either an authorized representative OR U.S. agent (for a foreign applicant).

For these fields, provide the following information for the authorized representative or U.S. agent:

- **I.B.1.** Select the checkbox if the authorized representative/U.S. agent information is the same as the applicant information identified in Part A. If the same, skip Part B fields 2–18 and proceed to Part C.
- **I.B.2.** Select only one checkbox to indicate whether you are completing Part B for either an authorized representative or an U.S. agent.
- **I.B.3.** The first name of the authorized representative or the U.S. agent.
- **I.B.4.** The middle initial of the authorized representative or the U.S. agent, if applicable.
- **I.B.5.** The last name of the authorized representative or the U.S. agent.
- **I.B.6.** The generational suffix (e.g., Jr., III) of the authorized representative or the U.S. agent, if applicable.
- **I.B.7.** The professional suffix (e.g., M.D., Ph.D.) of the authorized representative or the U.S. agent, ifapplicable.
- **I.B.8.** The professional position title of the authorized representative or the U.S. agent.
- **I.B.9.** The legal name of the organization that the authorized representative or U.S. agent is associated with, if applicable.
- **I.B.10.** The street address for the authorized representative or the U.S. agent (including street number, street name and type, suffix direction, etc.). The street address cannot be a P.O. Box.
- **I.B.11.** Additional street address information for the authorized representative or the U.S. agent (including apartment, suite, building number, #) that you were not able to include in I.B.10.
- **I.B.12.** The city of the authorized representative or the U.S. agent.
- I.B.13. The state, province, or territory of the authorized representative or the U.S. agent.

- I.B.14. The country of the authorized representative or the U.S. agent.
- **I.B.15.** The ZIP or postal code of the authorized representative or the U.S. agent.
- **I.B.16.** The phone number of the authorized representative or the U.S. agent (include country code, if applicable, and area code).
- **I.B.17.** The fax number of the authorized representative or the U.S. agent, if applicable (include country code, if applicable, and area code).
- I.B.18. The email address of the authorized representative or the U.S. agent.

Part C: Alternate Point of Contact Information (Optional)

Part C is an optional space for information for individuals not previously listed in Section I Parts A and/or B. Use the Continuation Page button within the form for additional alternate points of contact, as needed.

For these fields, provide the following information for the alternate point of contact:

- **I.C.1.** Indicate whether the alternate point of contact is one of the following:
 - Applicant
 - Authorized representative*
 - U.S. agent*
 - Other

*Note: Only contacts listed as the authorized representative and/or U.S. agent are authorized to act on behalf of the applicant for the submission.

- **I.C.2.** The first name of the individual.
- **I.C.3.** The middle initial of the individual, if applicable.
- I.C.4. The last name of the individual.
- **I.C.5.** The generational suffix (e.g., Jr., III) for the individual, if applicable.
- **I.C.6.** The professional suffix (e.g., M.D., Ph.D.) for the individual, if applicable.
- **I.C.7.** The professional position title of the individual.
- **I.C.8.** The legal name of the organization of the individual, if applicable.
- **I.C.9.** The street address for the individual (including street number, street name and type, suffix direction, etc.). The street address cannot be a P.O. Box.
- **I.C.10.** Additional street address information for the individual (including apartment, suite, building number, #) that you were not able to include in I.C.9.
- **I.C.11.** The city of the individual.
- **I.C.12.** The state, province, or territory of the individual.
- I.C.13. The country of the individual.
- **I.C.14.** The ZIP or postal code of the individual.
- I.C.15. The phone number of the individual (include country code, if applicable, and area code).
- I.C.16. The fax number of the individual (include country code, if applicable, and area code).
- I.C.17. The email address of the individual

Part D: Manufacturing Information

Only complete this section if the manufacturer information is different from the applicant. If this information is the same as the applicant, check the checkbox in I.D.1, skip Part D fields 2–20 and proceed to Part E.

- **I.D.1.** Select the checkbox if the manufacturer and applicant information is the same. If the same, skip Part D fields 2–20.
- **I.D.2.** The legal name of the manufacturer as it appears in the manufacturer's Dun & Bradstreet Data Universal Numbering System D-U-N-S® (D&B DUNS®) number.
- I.D.3. The manufacturer FDA-assigned Facility Establishment Identifier (FEI) number, if applicable.

 Note: To obtain or retrieve an FEI number, applicants can use the FEI Search Portal at https://www.accessdata.fda.gov/scripts/feiportal/index.cfm?action=portal.login or click here, and/or contact FEI Search Portal support at feiportal@fda.hhs.gov for any FEI number-related questions.
- **I.D.4.** The manufacturer D&B DUNS number, if applicable.

 Note: To obtain or retrieve a DUNS number, applicants can contact Dun and Bradstreet directly by phone at 1.866.705.5711 or through their website at https://www.dnb.com/duns/get-a-duns.html.

- **I.D.5.** Select the checkbox if the manufacturer address is the same as the applicant address, skip Part D fields 6–11 and proceed to Part D field 12.
- **I.D.6.** The street address for the manufacturer (including street number, street name and type, suffix direction, etc.). The street address cannot be a P.O. Box.
- **I.D.7.** Additional street address information for the manufacturer location (including apartment, suite, building number, #) that you were not able to include in I.D.6.
- **I.D.8.** The city where the manufacturer is located.
- **I.D.9.** The state, province, or territory where the manufacturer is located.
- **I.D.10.** The country where the manufacturer is located.
- **I.D.11.** The ZIP or postal code where the manufacturer is located.

Point of Contact for Manufacturer

- **I.D.12.** The first name of the manufacturer point of contact.
- **I.D.13.** The middle initial of the manufacturer point of contact, if applicable.
- **I.D.14.** The last name of the manufacturer point of contact.
- **I.D.15.** The generational suffix (e.g., Jr., III) of the manufacturer point of contact, if applicable.
- I.D.16. The professional suffix (e.g., M.D., Ph.D.) of the manufacturer point of contact, if applicable.
- I.D.17. The professional position title of the manufacturer point of contact.
- **I.D.18.** The phone number of the manufacturer point of contact (include country code if applicable and area code).
- **I.D.19.** The fax number of the manufacturer point of contact, if applicable (include country code if applicable and area code).
- **I.D.20.** The email address of the manufacturer point of contact.

Part E: Manufacturer/Packaging/Storage/Control Facility Information (Optional)

Part E is an optional space for information for additional manufacturing sites. Use the Continuation Page button within the form for additional sites, as needed.

For these fields, provide the following information for the Manufacturer/Packaging/Storage/Control Facility

- **I.E.1.** Select the appropriate checkbox to indicate the type of site.
- **I.E.2.** The organization name for the Manufacturer/Packaging/Storage/Control Facility (required by 21 CFR 1114.7(j)).
- **I.E.3.** The manufacturer FDA-assigned Facility Establishment Identifier (FEI) number, if applicable (required by 21 CFR 1114.7(j)).
- **I.E.4.** The manufacturer D&B DUNS number, if applicable.

 Note: To obtain or retrieve a DUNS number, applicants can contact Dun and Bradstreet directly by phone at 1.866.705.5711, at their website https://www.dnb.com/duns/get-a-duns.html, or click <a href="https://www.dnb.com/duns/get-a-duns.h
- **I.E.5.** The Manufacturer/Packaging/Storage/Control Facility, if applicable (required by 21 CFR 1114.7(j)).
- I.E.6. The Manufacturer/Packaging/Storage/Control Facility is ready for inspection by checking "Yes" or "No."
- **I.E.7.** The street address for the Manufacturer/Packaging/Storage/Control Facility (include street number, street name, and street type, and suffix direction, etc.) (required by 21 CFR 1114.7(j)). The street address cannot be a P.O. Box.
- **I.E.8.** Additional street address information (including apartment, suite, building number, #) that you were not able to include in I.E.7. (required by 21 CFR 1114.7(j)).
- **I.E.9.** The city where the Manufacturer/Packaging/Storage/Control Facility is located (required by 21 CFR 1114.7(j)).
- **I.E.10.** The state, province, or territory where the Manufacturer/Packaging/Storage/Control Facility is located (required by 21 CFR 1114.7(j)).
- **I.E.11.** The country where the Manufacturer/Packaging/Storage/Control Facility is located (required by 21 CFR 1114.7(j)).
- **I.E.12.** The ZIP or postal code where the Manufacturer/Packaging/Storage/Control Facility is located (required by 21 CFR 1114.7(j)).

Point of Contact Information for Manufacturer/Packaging/Storage/Control Facility

For these fields, provide the following information for the Manufacturer/Packaging/Storage/Control Facility point of contact.

- **I.E.13.** The first name of the Manufacturer/Packaging/Storage/Control Facility point of contact (required by 21 CFR 1114.7(j)).
- **I.E.14.** The middle initial of the Manufacturer/Packaging/Storage/Control Facility point of contact, if applicable (required by 21 CFR 1114.7(j)).
- **I.E.15.** The last name of the Manufacturer/Packaging/Storage/Control Facility point of contact (required by 21 CFR 1114.7(j)).
- **I.E.16.** The generational suffix (e.g., Jr., III) of the Manufacturer/Packaging/Storage/Control Facility point of contact, if applicable.
- **I.E.17.** The professional suffix (e.g., M.D., Ph.D.) of the Manufacturer/Packaging/Storage/Control Facility point of contact, if applicable.
- **I.E.18.** The position title of the Manufacturer/Packaging/Storage/Control Facility point of contact, if applicable (required by 21 CFR 1114.7(j)).
- **I.E.19.** The phone number of the Manufacturer/Packaging/Storage/Control Facility point of contact (include country code if applicable and area code) (required by 21 CFR 1114.7(j)).
- **I.E.20.** The fax number of the Manufacturer/Packaging/Storage/Control Facility point of contact, if applicable (include country code if applicable and area code) (required by 21 CFR 1114.7(j)).
- **I.E.21.** The email address of the Manufacturer/Packaging/Storage/Control Facility point of contact ((required by 21 CFR 1114.7(j)).

SECTION II — NEW TOBACCO PRODUCT INFORMATION

Utilize form FDA-4057b Premarket Tobacco Product Application Grouping Product Submission Spreadsheet available on the FDA website to provide new tobacco product(s) information..

SECTION III – SUBMISSION INFORMATION

Section IV should include submission information, including the following parts:

- Part A: Submission Type
- Part B: Cross-Referenced Information
- Part C: Referenced Tobacco Product Master File(s) (TPMF)
- Part D: Formal Meetings Held with FDA Pertaining to this Tobacco Product

Part A: General Submission Information

- **III.A.1.** Select the appropriate PMTA submission type: Standard, Resubmission, or Supplemental (as defined below).
 - A standard PMTA is a submission from an applicant seeking a marketing granted order to introduce a new tobacco product into interstate commerce. 21 CFR 1114.7
 - A resubmission PMTA is submitted to seek a marketing granted order for a new tobacco product by
 providing new information to address the deficiencies outlined in a marketing denial order and crossreferencing applicable content from the denied PMTA. 21 CFR 1114.17
 - A supplemental PMTA may be submitted by an applicant that is seeking authorization for modifications made to a new tobacco product for which they have already received a previous marketing granted order. 21 CFR 1114.15.15
- **III.A.2.** If the new products were previously commercially marketed in the United States, list the product name and provide the date(s) during which the product(s) were previously marketed.

Part B: Cross-Referenced Information (Optional)

Complete Part B if the application includes one or more cross-reference(s) to another PMTA or MRTPA 21 CFR 1114.7(b), 1114.15(b), or 1114.17(b). Supplemental PMTAs and resubmissions may cross-reference content in standard PMTAs. Standard PMTAs should not cross-reference another Standard PMTA or other pending applications with the exception of a pending MRTPA for the same tobacco product. Within the table, utilize a single row for each cross-reference. Use the Continuation Page button within the form to provide additional cross-references, as needed.

- III.B.1. In column 1, provide the FDA submission tracking number (STN) for the cross-referenced submission.
- **III.B.2.** In column 2, identify if the cross-reference provided in column 1 is for all products in the submission. If the cross-reference is only for some of the new products in the submission, select "no" and list the name of the product(s) that reference the cross-reference.
- **III.B.3.** In column 3, identify what information in the cross-reference submission you are seeking to reference for your new submission. For example, if you have the specific file name, document name, and page number, please list them.

Part C: Referenced Tobacco Product Master File(s) (TPMF) (Optional)

Complete Part C if the application includes a Tobacco Product Master File (TPMF).

Boxes 1–5 should be provided for each TPMF. Use the Continuation Page button within the form for additional TPMFs, as needed.

- **III.C.1.** In field 1, identify the TPMF owner.
- **III.C.2.** In field 2, provide FDA the submission tracking number (STN) of the TPMF. When a TPMF is established by FDA, the TPMF STN is provided to the owner and can be referenced by the TPMF owner and/or an authorized party. If the TPMF is not established at time of application submission, insert "N/A."
- **III.C.3.** In field 3, identify if the TPMF is applicable to all products in the submission. If the TPMF is only for some of the new products in the submission, select "no" and list the name of the product(s) that reference the TPMF.
- **III.C.4.** In field 4, identify what information in the TPMF you are seeking to reference for the new submission(s).
- **III.C.5.** In field 5, indicate if the right of reference is included in the submission. The TPMF owner may authorize another party to reference information contained within a TPMF through a right of reference such as a letter of authorization (LOA).

Part D: Formal Meetings Held with FDA Pertaining to the New Product(s) (Optional)

Complete Part D if FDA and the applicant held one or more meetings related to the new product(s). This can include meetings for study design, earlier versions of the product, etc. Within the table, utilize a single row for each meeting. Use the Continuation Page button within the form to list additional meetings, as needed.

- III.D.1. In column 1, provide the FDA submission tracking number (STN) for the industry meeting.
- **III.D.2.** In column 2, identify the date of the meeting between FDA and the applicant.
- **III.D.3.** In column 3, identify if the meeting topic was for all products in the submission. If the meeting relates only to some of the new products in the submission, select "no" and list the name(s) of the product(s) to which the meeting pertained.

SECTION IV – APPLICATION CONTENTS

Section IV is intended to help applicants organize their submission per 21 CFR 1114.7. For each item included in your submission, select the corresponding checkbox in the list and provide the location of the document. (For example, the file name, document name and page number). Select all that apply.

Part A: Administrative Content

- IV.A.1. Cover Letter.
- **IV.A.2.** Comprehensive Index (i.e., a listing of files and data associated with those files) and Table of Contents (21 CFR 1114.(b)(1)).
- **IV.A.3.** Written in English or accompanied by an English translation for non-English information (21 CFR 1114.(b)(1)). For any document that contains content that is not in English, translation is required. If all contents of the application are in English, leave the box blank. If you are providing translations for non-English information, select the checkbox.
- **IV.A.4.** Request for FDA to Refer PMTA to Tobacco Product Scientific Advisory Committee is optional. Select the checkbox if you are requesting a referral to TPSAC (21 CFR 1114.7(c)(5)).

Part B: Labeling and Marketing Plans

- **IV.B.1.** Specimens of all proposed labeling (21 CFR 1114.7(f)(1)).
- IV.B.2. Marketing plans (21 CFR 1114.7(f)(2)).

Part C: Inspections

IV.C.1. Location and contact information for each location subject to potential inspection (21 CFR 1114.7(k)(3) (vii)). Inspections may be conducted for manufacturing, clinical, or nonclinical sites.

Part D: Scientific Content

- **IV.D.1.** General information (e.g., product name, product category, subcategory and product properties) (21 CFR 1114.7(c)).
- IV.D.2. Descriptive information (21 CFR 1114.7(d)).
- IV.D.3. Product samples (21 CFR 1114.7(e)). FDA generally expects that product samples will be a required part of a PMTA and that an applicant should be prepared to submit them in accordance with FDA instructions within 30 days after submitting a PMTA. There may be situations in which sample submission may not be necessary, including, in some circumstances, PMTAs that are resubmitted for the same product after a marketing denial order (such as resubmissions as described in 21 CFR 1114.17) or PMTAs submitted for modifications to an authorized product where the modifications do not require review of new samples as part of the PMTA evaluation process. Pre-submission meetings with FDA may help provide additional information about whether product samples will need to be included in a PMTA; however, in most situations, FDA will only be able to determine the need for product samples after a PMTA is accepted for review.
- IV.D.4. Statement of compliance with 21 CFR part 25 (e.g., Environmental Assessment) (21 CFR 1114.7(g)).
- IV.D.5. Summary (21 CFR 1114.7(h)).
- **IV.D.6.** Product formulation (e.g., components, ingredients, additives, properties, and principles of operations) (21 CFR 1114.7(i)).
- IV.D.7. Manufacturing (e.g., methods, facilities, controls) (21 CFR 1114.7(j)).
- **IV.D.8.** Literature search (21 CFR 1114.7(k)(2)). A literature search is a search of available documents that includes: (1) clear search objectives, (2) a description of methodologies used in the search in detail, (3) an identification of relevant documents, (4) a formal or informal evaluation of study quality, and (5) a bibliography of referenced publications.
- IV.D.9. Organized references used to compile information in the submission.
- **IV.D.10.** Health risk investigations (21 CFR 1114.7(k)). Examples of health risk investigations include but are not limited to: Toxicological Risk Evaluation, Health Impact (e.g., use behavior, health risk), and Tobacco Product Perception and Intention Studies.

IV.D.11. Study Report(s) – Examples of documents include:

- Study protocol
- · Statistical analysis plan
 - Study report
 - Statistical software programming code
 - □ Study instruments (e.g., surveys/questionnaires)
 - □ Informed consent form
- Case Report Forms (CRFs):
 - In general, CRFs from clinical studies are not needed for filing a PMTA. However, FDA will require them for filing the CRFs from clinical studies that have been made to show the health risks of the PMTA product and whether such product presents less risk than other tobacco products where the CRF: (1) relates to participant deaths, other serious and unexpected adverse experiences, or participant discontinuation (including withdrawals) AND (2) where the study participant was exposed to the tobacco product(s) which is/are the subject of the PMTA(s) or to a similar/related product that the applicant is using to show that the PMTA product meets the standard for marketing authorization under section 910.
 - Additional information may be requested on a case-by-case basis during FDA review. FDA expects all CRFs would be available for review during agency inspections of clinical and/or nonclinical study sites.
- · Analyzable data sets:
 - In general, raw data such as raw chromatograms/spectra/mass spectra arising from analytical chemistry testing and raw (meaning no integration of the data) output from high-throughput (e.g., genomic) studies are not needed for filing a PMTA. Line data/ analyzable datasets that are representative chromatograms/spectra/mass spectra that demonstrate the adequacy of separations/specificity, standard solution, and sample solutions should be included. The line data/ analyzable data sets may be used to replicate findings or conduct alternative analyses of the underlying data. Additional information may be requested on a case-by-case basis during FDA review. FDA expects all raw data would be available for review during Agency inspection of clinical and/or nonclinical study sites.

SECTION V – Statements of Compliance with the Federal Food, Drug, and Cosmetic (FD&C) Act

- **V.1.** Provide information for how the application meets the requirements and addresses the question(s) in each of the statements according to the requirements section 910(b)(1) of the FD&C Act as required by 21 CFR 1114.7(c)(10) and (11). Your descriptions should address:
 - Full reports of all information, published or known to, or which should reasonably be known to, the applicant, concerning investigations which have been made to show the health risks of such tobacco product and whether such tobacco product presents less risk than other tobacco products.
 - Full statement of the components, ingredients, additives, and properties, and of the principle or principles of operation, of such tobacco product.
 - Full description of the methods used in, and the facilities and controls used for, the manufacture, processing, and, when relevant, packaging and installation of, such tobacco product.
 - An identifying reference to any tobacco product standard under section 907 which would be applicable
 to any aspect of such tobacco product, and either adequate information to show that such aspect of such
 tobacco product fully meets such tobacco product standard or adequate information to justify any deviation
 from such standard.
 - Specimens of the labeling proposed to be used for such tobacco product.
 - Such other information relevant to the subject matter of the application as the Secretary may require.
- **V.2.** Provide a brief description of how marketing the new tobacco product would be appropriate for the protection of public health as determined with respect to the population as a whole, including users and non-users of the tobacco product, and taking into account:
 - The increased or decreased likelihood that existing users of tobacco products will stop using such products.
 - The increased or decreased likelihood that those who do not use tobacco products will start using such products.

SECTION VI - CERTIFICATION STATEMENTS

The application must contain the following certifications, as appropriate for the specific type of PMTA, with the appropriate information inserted, as described in each parenthetical, signed by an authorized representative of the applicant.

Provide the name of authorized representative who is signing the certification. Insert name of authorized representative as identified in Section I Part B or Part C. Provide the name of the applicant being represented in the certification. Insert name of applicant as identified in Section I Part A.

The required Certification Statements (i.-iv.) are based on the specific type of PMTA you are submitting, as follows:

- i. Certification statement for standard PMTAs is appropriate when submitting a standard PMTA
 - Insert the name of the authorized representative as identified in Section I Part B or Part C, the name of the organization being represented as identified in Section I Part A.
- ii. **Modified tobacco product certification for supplemental PMTAs** is appropriate when submitting a supplemental PMTA.
 - Insert the name of the authorized representative as identified in Section I Part B or Part C, the name
 of the organization being represented as identified in Section I Part A, the individual new product(s)
 names(s), a description of each modification, and the name and STN of the previously submitted PMTA
 where appropriate in the statement. If submitting multiple products, it is recommended a separate
 certification is submitted for each product.
- iii. **Same product certification for resubmissions** is appropriate when submitting a resubmission PMTA where the product is unchanged, and the applicant is addressing deficiencies outlined in the marketing denial order (MDO).
 - Insert the name of the authorized representative as identified in Section I Part B or Part C, the name of the organization being represented as identified in Section I Part A, the individual new product(s) names(s), and the STN of the previously submitted PMTA where appropriate in the statement.
- iv. **Different product certification for resubmissions** is appropriate when submitting a resubmission PMTA where the product is a modification of the previously submitted PMTA that results from changes necessary to address the deficiencies outlined in the marketing denial order (MDO).
 - Insert the name of the authorized representative as identified in Section I Part B or Part C, the name of the organization being represented as identified in Section I Part A, the individual new product(s) names(s), the name and the STN of the previously submitted PMTA, and a description of each modification where appropriate in the statement. If submitting multiple products, it is recommended a separate certification is submitted for each product.
- v. **Financial Interest and Arrangements of Clinical Investigators Certification Statement** is appropriate when submitting any type of PMTA and must be included if your application includes any type of study in support of this application. This certification covers all actions taken to ensure the reliability of the study.
 - Insert the name of the authorized representative as identified in Section I Part B or Part C, and the name of the organization being represented as identified in Section I Part A.

For each certification statement applicable, insert the signature of the authorized representative and the date the certification is signed.

OTHER INFORMATION

Identify and provide information for any additional information not captured in the PMTA submittal form that is pertinent to your application.

We remind you that all regulatory correspondence can be submitted via the FDA Electronic Submission Gateway (www.fda.gov/esg) using eSubmitter.

We are unable to accept regulatory submissions by electronic mail.