TFDA Risk No. 1101107078 Announcement dated September 28, 2021 Notices for the Plant Master File (PMF) Application Form C-1

Plant Master File (PMF) Checklist for Foreign Pharmaceutical Manufacturer

Form C-1: Documents in Common review (For the expansion of manufacturing site, the items which are marked with asterisk are required, and the relevant documents shall be enclosed.)

Applicant:	Receipt No.	Case Number
Item	Please complete the Checklist item by item and indicate the attachment numbers or the page numbers of <u>submitted</u> <u>documents</u>	Reviewer' s comment
*1.1 Name of manufacturer (which shall be consistent with that shown in the official supporting documents)	Р.	
*1.2 Address of manufacturer (which shall be exact detailed and consistent with that shown in the official supporting documents; and give both contact address and site address, if different)	Р.	
 1.3 Legalization requirements: Based on Article 5, Paragraph 2 of the "Regulations of Medicament Manufacturer Inspection, the PMF application dossier for submission shall meet one of the following requirements for submission: PMF or SMF, which is to be certified by the highest competent health authority or the chamber of commerce in the country of origin or legalized by the embassy, representative office or agencies authorized by the Ministry of Foreign Affairs of R.O.C. Original copy of documents from the hosting competent authority or certified photocopy of the said documents certified by the hosting competent authority or certified photocopy of the said documents certified by the manufacturer is in compliance with local pharmaceutical GMP standards. (if the two original documents mentioned above were already submitted to the TFDA through other cases, a photocopy of the whole document may be enclosed and indication of the case number that the original copies were submitted.) Original copy of Certificate of Pharmaceutical Product (CPP) from the hosting competent authority or chamber of the said documents certified photocopy of the said documents certified photocopy of the said documents certified photocopy of the said documents certified by the hosting competent authority or certificate of Pharmaceutical Product (CPP) from the hosting competent authority or chamber 	P.	

of commerce in the country of origin clearly stating		
that said manufacturer is in compliance with local		
pharmaceutical GMP standards. (if the two original		
documents mentioned above were already submitted		
to the TFDA through other cases, a photocopy of the		
whole document may be enclosed and indication of		
the case number that the original copies were		
submitted.)		
If the paper based GMP certification is no longer		
available in the country of origin, or the drug product		
is contracted manufacturing, the statement of license		
holder of the imported drug shall be submitted		
explaining why the GMP certification and CPP is not		
available. Therefore, the applicants could submit the		
original copy or certified photocopy of the CPP issued		
by any one of the A-10 countries, EMA, or by the		
highest competent health authority in the country		
where the contractor is located, in accordance with the		
letter TFDA Risk No. 1051105400 dated October 17,		
2016.		
*1.4 Dosage form/product/manufacturing process applied in		
the application		
*1.4.1 Completed by the Taiwan pharmaceutical company,		
with indication of the manufacturing process stage		
being applied for. For biological medicinal products,		
and blood products derives from human blood or		
plasma, the product/dosage form and manufacturing		
process stage shall be specified, and the following		
items shall be checked:		
Animal sourced products		
Allergen products	Р.	
Animal immunosera products		
Recombinant products		
Monoclonal antibody products		
Transgenic animal products		
Transgenic plant products		
Gene therapy products		
Somatic and xenogeneic cell therapy products and		
tissue engineered products		
Blood products		
*1.4.2 Explain if special products (biological medicinal		
products, highly sensitizing, highly pharmacological		
active, toxic, or hazardous substances) are included		
in the Dosage form/product/manufacturing process	Р.	
applied in the application, such as β -lactam		
antibiotics (e.g., penicillin, cephalosporins, Penems,		
Carbacephem, Monobactams), Hormone (include sex		
hormones and non-sex hormones),		

cytotoxics/cytostatic, or radioactive medicinal	
products.	
*1.4.3Explain whether the manufacturing and testing of the	
Dosage form/product/manufacturing process applied	
in the application is full-manufacturing process or	
phased. If the production or testing is phased, the	Р.
implementation stage in the manufacturing site	
shall be specified separately.	
For applicants applying for simplified review for	
non-sterile dosage form except for secondary	
packaging, the flowchart of major manufacturing	Р.
steps for the applied dosage form/manufacturing	
process are required.	
*1.4.4 Enclose the layout of production area (from	
weighing to secondary packaging and shall include	
personnel/material flow, air flow/pressure difference	
and room cleanness) and address the area for the	Р.
Dosage form/product/manufacturing process applied	
in the application.	
1.5 The approval letter holds by applicant or other	
pharmaceutical companies.	
The TFDA issued approval letter holds by applicant	
already, photocopies enclosed.	
	Р.
When applying for quote review: The TFDA-issued	
approval letter holds by applicant or other companies,	
photocopies enclosed.1.6 Overview of the manufacturing site	
1.6.1Briefly describe the premises (area, location,	P.
surroundings)	
*1.6.2Site layout, with indication of purposes of respective	
buildings and each floor in the site. Address the	
buildings and floors for the Dosage	P.
form/product/manufacturing process applied in the	
application.	
1.6.3 Describe whether the outsourced activities and the	
contract meet the regulations in PIC/S GMP, Chapter	Р.
7.	
1.7 Manufacturing activity at the site approved by the	
competent authority in the original country (photocopy	Р.
of the official document)	
*1.8 Description of the all production activity in the site	
*1.8.1 List the products currently manufactured in the	
manufacturing site by their dosage forms and their	
active ingredient (including contract manufacturing	
given and accepted), and enclose a list. The type of	Р.
the products shall also be included in the list, for	· ·
example, human medicinal product, human	
investigation medicinal product, veterinary	
myosugation motional product, vetermary	

aquinment in the shared and heating areas an	
equipment in the shared production areas or	
shared facilities and equipment with human	
medicinal products, etc.), and indicate	
production areas for said products in the	
layout.	
*1.8.3.2.1 For <u>dedicated equipment in the shared</u>	
production areas with human medicinal products,	
the following shall be described:	
A. Is the active ingredient of certain products	
archived in the pharmacopoeia? If yes, enclose	
relevant bases.	D
B. Is the manufacturing in compliance with the	Р.
PIC/S GMP standards? (The statement shall be	
written and signed by the manufacturer)	
C. Describe the measures to prevent cross-	
contamination and how their effectiveness being	
periodically reviewed.	
*1.8.3.2.2 For shared facilities and equipment with	
human medicinal products, the following shall be described in detail:	
A. Is the active ingredient of certain products	
archived in the pharmacopoeia? If yes, enclose	
relevant bases.	
B. <u>Is the manufacturing in compliance with the</u>	
PIC/S GMP standards? (The statement shall be	
written and signed by the manufacturer)	
C. Describe the measures to prevent cross-	
contamination and how their effectiveness being	
periodically reviewed.	
D. If the veterinary medicinal products and human	
medicinal products are manufactured at the same	
facilities and share same equipment, and the said	
veterinary medicinal products are not used in	Р.
Human, the following dossiers shall be	Г.
submitted: the risk assessment report including	
toxicological data, Health Based Exposure Limit	
(HBEL)PDE /ADE, and correspondence	
measurements to prevent from cross	
contamination .(If the veterinary medicinal	
products are not manufactured from weighing to	
primary packaging at the same facility which	
also produce human medicinal products, this	
item is not required.)	
E. If the specific products shared facilities and	
equipment (from weighing to primary	
packaging) with human medicinal products are	
veterinary medicinal products, herbal medicine,	
homeopathic drugs or common products(anti	
mosquito products, tooth paste, mouthwash,	

shampoo, etc.), the following requirements shall be enclosed: the list of shared equipment, the list of dosage form and components for all products, describe the implementation of cleaning validation (e.g. single product approach or grouping approach). Where grouping approach is used, describe the		
categories in detail, and also list the APIs of each		
product in each group, and the target ingredient		
for cleaning validation. (If there are no specific		
products manufactured at the facility of human		
medicinal products (from weighing to primary		
packaging), this item is not required.)		
	Signature	
	(including date of	
	signing)	