Product Registration (PR)

List of Documents to be enclosed with MAI Application

- 1. **Product Registration Reimbursement Application form. Annexure-I** duly completed & authenticated by CEO/CHAIRMAN/MD / PARTNER/PROPRIETOR /Authorized Person.(click here-refer page no: 2)
- 2. **Declaration** On the letter head of the Company, duly completed & authenticated by CEO/CHAIRMAN/MD/PARTNER/PROPRIETOR (click here-refer page no: 3)
- 3. **Affidavit** Completed & authenticated by CEO/CHAIRMAN/MD/ PARTNER/PROPRIETOR <u>click here-refer page</u> <u>no: 4)</u> Note: Declaration and Affidavit should be signed by the same person.
- 3a. If Declaration and Affidavit both were signed by other than CEO/CHAIRMAN/MD/PARTNER/PROPRIETOR, Power of Attorney (or) Board Resolution in the name of signed person.
- 4. CA Certificate mentioning the Product Expenses (With Exchange rate) (click here-refer page no: 5)
- 5. FOB Value of Exports for the preceding Financial Year. (click here-refer page no: 6)
- 6. Product Registration Certificates "Original+Copy" (Registration Certificate SHOULD BE on the Name of the Applicant as MARKETAUTHORIZATION HOLDER / MANUFACTURER)
 - a. Original Certificates need to be Verified/Attested by PHARMEXCIL Office (Hyderabad/Mumbai/Delhi).
 - I. As in some cases where the Original Product Registration certificates were not issued by the Particular Regulatory Authority then, we will verify those certificates through on-line providing us the Online Links to verify in Official website of drug regulatory authority, and Login credentials, if required (Login id and Password of the company for Verification).
 - II. Now-a-days, some countries were issuing online Renewal/Retention Certificates, then all the Invoice and Receipt copies which were issued in original need to be Verified/Attested by PHARMEXCIL Office.
 - b. If the product is re-registered, then the details of previous registration along with date of registration and its validity period and a copy of the registration certificate.
 - c. After Verification, the Originals will be returned back immediately.
- 7. Self-Attested Translation Copies in English wherever necessary (when documents are in foreign language):
 - a. Registration Certificates.
 - b. Previous registration certificate.
 - c. Invoice.
 - d. Receipt.
 - e. Registration guidelines reflecting and highlighting the registration fee structure.
- 8. Copies of Invoices & Receipts raised by Drug Regulatory Authorities as Proof of Payment.
- 9. Debit Note raised by Agent (if registered through agent), payment is made by the agent.
- 10. Bank Transfer Remittance (**SWIFT** copy) showing the Customer Name, Beneficiary Name & Purpose of Payment details a. Attestation/Round Seal by the Banker is MANDATORY on the Bank Transfer Remittance (Swift Copy) with Employee code.
- 11. Foreign Outward Remittance to known the exchange rate as on payment date.
- 12. In case of **Cash Payment**, the Payment Invoice Number of each product along with product name against which the payment was made should be reflected on Cash Receipt.
- 13. **Valid Manufacturing Drug Licenses** (Form 25/28) and Renewal License (Form 26) reflecting and highlighting the name of claiming products for export purpose.
- 14. **Manufacturing Agreement** (if manufacturing is done through Third Party- enclosing Annexure with name of the claimed products).
- 15 .Copy of the Product Registration **Guidelines of FDA**, showing/reflecting the details of Registration Fee to be paid for the Subject product, procedure of product registrations etc.
- ${\bf 16.\ Applicant\ name\ should\ be\ clear\ from\ DGFT\ DEL\ STATUS\ (Denied\ Entity\ List)}.$

Note:

- Financial year of the claim, is considered basing on the <u>DATE OF ISSUE OF PRODUCT REGISTRATION CERTIFICATE</u> (Financial year period: 1st April to 31st March).
- > Products Registered in different Countries need to be submitted in different Applications.
- Products pertaining to different Financial Years to be submitted in different Applications.
- For all the MAI claims filed by Exporters to Pharmexcil from 26th June 2023, the following two provisions would apply:
 - 90 days' timeline for submission of MAI claims by Exporter to Pharmexcil
 - Limitation of reimbursement to the exporters having f.o.b value of exports up to Rs. 100 crore during the preceding financial year.
- In case of unaudited F.O.B Values of Exports (April-September) applicants have to submit the Chartered Accountant Certificate w.r.t FOB value of Exports during the preceding FY along with GST returns filed during the preceding year.
- Application will be accepted only if the company submit all the mandatory documents and confirm the application online.
- Processing fee of 5% will be charged on sanction of the amount.

FORM FOR CLAIMING REIMBURSEMENT OF REGISTRATION CHARGES PAID FOR REGISTRATION ABROAD OF PHARMACEUTICAL/ BIOTECHNOLOGICAL/ AGRO CHEMICAL PRODUCTS

Date:

1.	Name of the firm with full address	
		IEC No.
2.	EH/TH Certificate	No. & Date:
	Whether SSI/Non-SSI/Trader	Valid Up to:
3.	FOB Value of exports for the preceding financial year	Rs. In Crores:
4.	Particulars of products registered by the Company	Name of the Product: Category: Pharma/Biotech/Agro-chem
		Place:
		Country:
5.	Particulars of certificate procured from Ministry of Health/ Agricultural etc. of Foreign Country along with receipt for payment of registration fee	Date of letter/certificate
		Amount paid in US(\$)/Euro(€)/ Foreign currency and Indian (Rs.)
6.	Whether certificate from Indian mission concerned certifying the amount paid towards registration charges is enclosed	
7.	Details of registration fees claimed in past:	
	1.Name of the product (Pharma/Biotechnological/agro- chemical)	
	2. The country of registration	
	3. Date of receipt of reimbursement	
	4. Amount received	
8.	Details of claims of reimbursement for the year	
9.	Enclose copy of registration certificate issued by foreign agency (if the certificate is in a language other than English, enclose attested translated copy)	
10.	Enclose copy of manufacturing license issued by State Drugs Controller/Licensing Authority (in India) for the subject product	

On Company Letter head

DECLARATION

I solemnly declare that the particulars given in the above statement are correct. I bound myself and the company accountable and responsible for any incorrect information given in the above statement and shall immediately refund amount received on the basis of wrong information provided in the above statement.

Signature:			
Name:		-	
Designation:			
Office Seal:			
Counter signed by CEO/CH	AIRMAN/MD/PARTNER/PROP	RIETOR of the Company	r:
		Signature:	
		Name :	
		Designation:	
		Office seal:	
Place:			
Date			

Affidavit to be submitted on Rs.50/- or Rs.100/- Non-Judicial Stamp Paper

AFFIDAVIT

I, S/o	Years Resident of
do here by affirm on oath as under.	
claiming reimbursement of Registration charges paid Products to the Department of Commerce, Ministr Under MAI Scheme. I, solemnly declare that the particulars given in the a accountable and responsible for any information	on whose behalf an application is made for d for Registration Abroad of Pharmaceutical /Biotechnological ry of Commerce and Industry, Government of India, New Delhi bove application are correct. I bound myself and the company on given in the application. If any of the information and insible for any action initiated by the Government under any law is, immediately.
	Signature: Name:
	Designation:
	Office seal:
Place:	
Date:	

On CA Letter Head

S.NO	Particulars of expenses		Amounts in	
	(each products names in individual cells)	USD/EURO	Exchange rate	Rupees
1.				
2.				
3.				
4.				
5. 6.				
Signatur	e & Stamp/Seal of the Signatory:			
0	,			
Signatur	e:			
Name:				
varrie.				
∕lember	ship No.:			
-ull Addı	·occ·			
uli Auui	<u> </u>			
Name ar	d address of the Institution where registered.			
Place:				
D-4-				
Date:				

On CA Letter Head

TO WHOM SO EVER IT MAY CONCERN

		Financial Year	Amount in Crores (Only)
	1		
ure & Stamp/ ure:	Seal of the S	ignatory:	
<u> </u>		_	
rship No.:		_	
lress:		_	
nd address o	of the Institu	tion where registered.	

Date:

SELF DECLARATION

TO WHOM SOEVER IT MAY CONCERN

 Th	e following produc) was submitted to	by confirm that the following MAI Control Pharmexcil towards reimbursement ed/initiated to the Drug Regulatory And the trailing table.	of Product Registration Charges
	Product Name	Product Registration number	Category: Registration/Re- Registration/Renewal/Retentio n	Date of submission/initiatio n of application to Drug Regulatory Authority towards particular category.
Su		• •	particular product registration/re-re of the country of on or after 0	-
	uthorize person sig	nature)		
De	esignation:			

ON Company letter head

To Whom So Ever It May Concern

We M/s	5		(Compa	ny name)	hereby pr	oviding t	the follov	ving details t	owards payr	nent for
the claimed products.										
S.NO	Product	Invoice.	Invoice	Receipt	Receipt	Debit	Debit	Bank	TT	Amount
	Name	No.	Date	No	Date	Note	Note	statement	reference	in
						No.	date	Date	Number	Foreign
										currency
1										
2										
3										
And als	so mentior	n the Purp	ose of pay	/ment:						
Yours s	incerely,									
(Author	ize person	signature)							
Designa	ition:									