

Application Form for the Registration of Drugs

Form DA-1/88

Application Form for the Registration of Drugs (which are included as monographs in BP/BPC/USP-NF/Int. Ph. or are already Introduced in Bangladesh)

1. Name and address of the Manufacturer of the Drug :
 2. Manufacturing Licence Number (for locally manufactured drugs):
 - (a) Biological:
 - (b) Non-Biological:
 3. Name of the Drug:
 - (a) Generic name (use INN name if included in INN List) :
 - (b) Name under which the Drug is proposed to be sold :
 4. Product Data Sheet
(Including Presentations, Uses, Dosage & Administration, Contra-indication, Use in pregnancy and lactation, Side-effects, Precaution, Warning, Drug Interaction, Absorption, Fate, Distribution, Excretion, Elimination, Package Quantities, etc.)
 5. Technical Data:
 - (a) Composition/Formula
 - (b) Manufacturing Instruction
 - (c) Control Data for the Active Constituents
 - (d) Pharmacopoeial References or Control Data for other constituents
 - (e) Control data for Finished Product
 - (f) Stability data (if not done, then to be submitted at the time of inclusion)
 - (g) Proposed shelf life (must be expressed on Finished Product in the form of Manufacturing Date and Expiry Date).
- Note :
1. Excipients should always be mentioned in generic/chemical names; but may be followed by brand names, if desired.
 2. Overage to be shown separately in Composition/Formula, eg., 2.5% for antibiotics Capsule/Tablet, 5% for antibiotics Dry syrup/Injection, 10% for Vitamins, etc.
 3. Capsule size to be mentioned by number; name/Monogram should be printed on capsule or engraved in tablet.
 4. Coating material should be shown separately.
6. (a) Number of manufacturer/importer already manufacturing/importing the product in Bangladesh; and
(b) Estimated market of this product/product group in Bangladesh.
 7. (a) Proposed Maximum Retail Price (MRP);
(b) Estimated Price-per dose; per day treatment; cost for the recommended course of treatment.
 8. For locally manufactured drugs :
Particulars of quality Control manager and Factory/Production manager.
Full name, Qualifications, Date of Joining in the applicant's company; Total experience in the pharmaceutical industries, Registration Number and Signatures.
 9. In case of imported drugs, the following additional information are to be provided :
 - (a) Name and address of the Indentor/or Manufacturer's authorized agent.
 - (b) Registration status in the country of origin (including Free Sale Certificate)
 - (c) Registration status in other countries (include Sale Certificates from atleast 2 other development countries)
 - (d) Signature of the Indentor/or Manufacturer's authorized agent:

10. Date of submission :

11. Additional information (if any):

Please Note : Information supplied if found wrong will lead to immediate cancellation of registration of the product.