

**FORM CT-21***(See rule 80)***APPLICATION FOR GRANT OF PERMISSION TO MANUFACTURE NEW DRUG FORMULATION FOR SALE OR FOR DISTRIBUTION**

I/We,.....(*name and full postal address of the applicant*) of M/s..... hereby apply for grant of permission to manufacture new drug for sale or distribution.

The details of the application are as under:

1. Name of applicant:	
2. Nature and constitution of applicant:  (i.e. proprietorship, partnership including limited liability partnership, company, society, trust, other to be specified)	
3.(i) Corporate or registered office address including telephone number, mobile number, fax number and e-mail id:  (ii) Manufacturer's address including telephone number, mobile number, fax number and e-mail id:  (iii) Address for correspondence:	
4. Details of new drug to be manufactured (Active pharmaceutical Ingredient or Finished Formulation or both) [As per Annexure].	
5. Details of the manufacturer and manufacturing site [As per Annexure].	
6. Fee paid on _____Rs__receipt or challan or transaction ID.	
7. I hereby state and undertake that:  (i) I shall comply with all the provisions of the Drugs and Cosmetics Act, 1940 and Chapter X of the New Drugs and Clinical Trials Rules, 2019.	

Place:.....

Date: .....

Digital Signature

(Name and designation)

**Annexure:**

Details of new drug:

Name of the new drug:	
Dosage form:	
Composition of the formulation:	

Therapeutic class of the new drug:	
Indications for which proposed to be used:	

Details of manufacturer and manufacturing site of new drug:

Name and address of manufacturer (full address with telephone, fax and e-mail address of the manufacturer).	Name and address of manufacturing site (full address with telephone, fax and e-mail address of the manufacturing site).