Duly filled specimen pro forma for submission of application in Form MD-12 for obtaining Manufacturing Test license



FORM MD-12 1. The name of the applicant (firm) shall be 2. The applicant may select the option as provided in the portal e.g., proprietorship, mentioned as registered under applicable law [See sub-rule (1) of rule 31] LLP. Pvt. Ltd. etc. like company act, proprietorship etc. Application for License to Manufacture Medical Device for Purpose of Clinical 3.(ii) 3.(i) The address shall be mentioned as Investigation, Test, Evaluation, Examination, Demonstration of Training In case of testing/ evaluation/ examination. registered under applicable law like company the applicant shall mention the name and act, proprietorship etc. Name of Applicant: address of the testing centre(s) where the proposed test is to be carried out. Nature and constitution of manufacturer: · In case of demonstration/ training, the 3. (iii)The applicant shall mentioned the name and address of the site where the actual test applicant shall mention the name and (i) Corporate/registered office address: address of the site(s) where batches will be developed/manufactured demonstration/ training to be performed. (ii) Testing or evaluation site address: In case of Clinical investigation, the 4.1.1 Only the common name or generic name applicant shall mention the name and of device shall be mentioned. (iii) Address for correspondence: address of each site(s) where the clinical study is to be carried out. 4. Details of medical device(s) to be manufactured S. No. Details of Device(s) 4.1.2 The specific brand name of the device 4.1.3 Should mention the model number/name shall be mentioned (If any). or catalogue number as mentioned in Generic Name: DMF & IFU/ UM/ PI and labels of the device(s). Brand Name (if any): if available. Model No. (if any): 4.1.5 The proposed intended use of the device Proposed Class of Medical Device: as mentioned in manufacturer's draft IFU/ UM/ PI (If applicable). Proposed Intended Uses **_6**. Whether Sterile or Non-sterile: 4.1.4 Proposed risk class of the devices shall Proposed Quantity: be as per classification list published on the 4.1.7 The proposed quantity (in metric units) CDSCO website or as per the claim of the shall be properly justified in the line of test manufacturer (Click here) protocol submitted 5. Please refer details in Payment receipt attached. 6.1 hereby state and undertake that, I shall comply with all applicable provisions of the Drugs and Cosmetics Act, 4.1.6 Only applicable for sterile product 5. Payment shall be made as per the Second otherwise the applicant shall mention schedule of Medical Devices Rules, 2017 1940 (23 of 1940) and the Medical Devices Rules, 2017. 'Not Applicable'. Place: Signature 🚤 It should be digitally signed using digital (Name & Designation) Date: signature certificate as per IT Act. Place where the address of the applicant Shall be in DD/MM/YYYY (firm) is located IFU/ UM/ PI: Instructions for use/ User Manual/ Package Insert Abbreviations: DMF: Device Master File CLA: Central Licensing Authority

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