



Government of Pakistan
Ministry of National Health Services
Regulations & Coordination
Drug Regulatory Authority of Pakistan

Sr. No. 045



FORM-4

[see rule 5(2)]

LICENCE TO IMPORT MEDICAL DEVICES

Licence No. ELI-00072

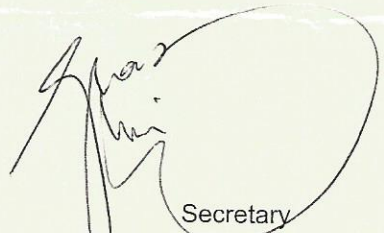
Date of issue: 03/08/2018

F.No: 12-36/2017-MD

M/s Medco Healthcare, is hereby licensed to import registered medical devices at the following premises: 155, Block -3 C.P Barar Society, Alamgir Road Karachi.

2. Name(s) of proprietor(s) along with the residential address and CNIC Number(s)
Atif Ahmed Khan,
B-41/11, F. B. Area, Karachi.
CNIC No: 42101-4193983-5
3. Name(s) of the person(s) incharge who will personally supervise the import and sale of medical devices by way of wholesale along with registration No, residential address and CNIC No.
 - (i) Muhammad Kashif Shamim
House No.667-A, Mohallah North Nazimabad, Block L, Karachi.
CNIC # 42101-9028255-9
4. Addresses of godowns , if any, where medical devices shall be stored
155, Block -3 C.P Barar Society, Alamgir Road Karachi.
5. The licence will be in force for a period of five years from the date of issue unless earlier suspended or cancelled.
6. This licence shall, in addition to the conditions specified in the Medical Devices Rules, 2017 made under the DRAP Act, 2012, be subject to the following conditions namely:-
 - (i) The persons mentioned above shall personally supervise the sale of medical devices.
 - (ii) The licence and registration certificate from the Pharmacy council of the person(s) incharge, personally supervising the sale of medical devices shall be displayed in a prominent place in the premises open to public.
 - (iii) No medical device requiring special storage conditions of temperature and humidity shall be stored or sold unless the precaution necessary for preventing the properties of the components have been observed throughout the period during which it remained in possession of the licensee.
 - (iv) Importer shall be responsible for labeling requirements as per Medical Devices Rules, 2017 including Importer Licence details, Products Registration Numbers and MRP.

Renewal Date: 03/08/2023


Secretary
Medical Devices Board
(DR. GHAZAFAR ALI)
Additional Director (MD&C)
Drug Regulatory Authority of Pakistan
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