## FORM 25D

Renewal Licence

## (See Rule 154) Licence to manufacturing for sale of Ayurvedic, siddha or Unani drugs

Number of licence GA/1696 Date of Issue : 08-05-2022

- DIRECTOR OF VIRGO UAP PHARMA PVT. LTD. is / are hereby licenced to manufacture the following Ayurvedic, Siddha or Unani drugs on the premises situated at SURVEY NO 423/98-B,MAHAGUJARAT IND. ESTATE,SARKHEJ-BAVLA HIGHWAY, VILL-MORAIYA,AHMEDABAD,TALUKA : SANAND,DISTRICT : AHMEDABAD,382213 under the direction and supervision of the following technical staff
  - a. Competent Technical staff (Names).-(AS ATTACHED)
  - b. Names of drugs categorized as per Schedule T (each item to be separatly specified) with specific Product Code/QR Code for each approved drug.(AS ATTACHED)
- 2. The licence shall be in force from 08-05-2022
- 3. The licence is subject to the conditions stated below and to such other conditions as may be specified in the Rules for the time being in force under the Drugs and Cosmetics Act, 1940.

Joint Commissioner (Ayurved) Food & Drugs Control Administration Gujarat State.

## Date:- 08-05-2022

## **Conditions of Licence**

- 1. Any change in the Technical staff named in the licence shall be forthwith reported to the Licensing Authority.
- 2. This licence shall be deemed to extend to such additional items as the licencee may intimate to the Licensing Authority from time to time, and as may be endorsed by the Licensing Authority.
- 3. The licencee shall inform the Licensing Authority in writing in the event of any change in the constitution of the firm operating under the licence. Where any change in the constitution of the firm takes place, the current licence shall be deemed to be valid for a maximum period of three months from the date on which the change takes place unless, in the meantime, a fresh licence has been taken from the Licensing Authority in the name of the firm with the changed constitution.
- 4. The licence unless sooner suspended or cancelled shall remain valid perpetually. However, the compliance with the conditions of licence and the provisions of the Drugs and Cosmetics Act 1940 (23 of 1940) and the Drugs Rules, 1945 shall be assessed not less than once in five years or as needed as per risk based approach.
- 5. The licence is issued only after fulfillment of the requirements of Good Manufacturing Practices (GMP) of Ayurveda, Siddha or Unani drugs as laid down in Schedule T of the Drugs Rules, 1945.".

