Certificate of Pharmaceutical Product(s)¹ This Certificate conforms to the format recommended by the World Health Organization (General instruction and explanatory notes attached) : 26/1/औषधि/44/2019/ उ०५९ Date : 26/03/2022 No. of Certificate Exporting (Certifying) Country : INDIA Importing (requesting) Country : All Country 1. Name and dosage form of product : Midaver [Midazolam Injection BP 1mg/ml-5ml] 1.1 Active ingredient(s)² and amount(s) per unit dose³: Each ml contains: Midazolam BP 1mg Water for Injection BP q.s 1.2 Is this product licensed to be placed on the market for use inexporting country?⁵ 1.3 Is this product actually on the market in the exporting country?⁵ The answer to 1.2 is Yes continue with section 2A and omit section 2B The answer to 1.2 is No Omit Section 2A and Continue with section 2B⁶ 2A.1 Number of product license 34/UA/SC/P-2010 (Form-28) 2B.1 Applicant for certificate (name and address): And date of issue: 05-11-2020 2B.2 Status of Applicant 2A.2 Product License holder (Name & Address) 2B.2.1 For categories b and c the name and Verve Human care Laboratories. address of the manufacturer producing Plot No. 15-A, Pharmacity, Selagui, dosage form are9 Dehradun, Uttarakhand (India). 2A.3 Status of product License holder8: 2B.3 Why is marketing authorization lacking?: 2A.3.1 For categories b and c the name and address of the manufacturer producing the dosage form are 9: N.A 2A.4 Is Summary Basis of Approval appended?¹⁰ 2B.4 Remarks¹³: 2A.5 Is the attached, officially approved product Information complete and consonant with the License?: Not Provided 2A.6 Applicant for certificate different from license holder (name and address)12: N.A 3. Does the certifying authority arrange for periodic inspection of the manufacturing plant in which the dosage form is Produced? Yes 3.1 Periodicity of routine inspections (years): Annual 3.2 Has the manufacturer of this type of dosage form been inspected? Yes 3.3 Do the facilities and operations confirm to GMP as recommended by World Health Organization?15 Yes 4. Does the information submitted by the applicant satisfy the certifying authority on all aspects of the manufacture of the product?16 N.A If no, explain: This certificate valid upto: 27-02-2025 Address of certifying authority: Drug Controller, Directorate General of Health Services. Sahastradhara Road, Dehradun, Uttarakhand, India, Hemant Singh Negi) Drug Controlling & Licensing Authority Name of the authorized Person: Mr. Hemant Singh Negi

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