

# CERTIFICATE OF A PHARMACEUTICALS PRODUCT

Certificate No : MFG/STATE GMP/COPP/2023/2242239352

Valid Up To :02/08/2025

Exporting (Certifying) Country : **India**

Importing (Requesting) Country : **Moldova**

1 Name and dosage form of products : (Brand Name if any) : **Carbamazepine Tablets BP 400mg**

1.1 Active ingredient(s) and amount(s) per unit dose :

**Each uncoated Tablet Contains:**

Composition	Ingredients	Standards	Strength	UOM	Equivalent to
API	Carbamazepine	BP	400	Milligram	
Excipients	Excipients:	--	0	QS	

1.2 Is this product licensed to be placed on the market for use in the exporting country?

**Yes**

1.3 Is the product actually on the market in the exporting country?

**Yes**

2A.1	No. of Product license and Date of issue <b>Product License in Form 25 bearing no. G/25/163 Date of Issue : 03/11/2023</b>	2B.1	Applicant for certificate (name and address): <b>Not Applicable</b>
2A.2	Product License holder :(Name and address) <b>WEST-COAST PHARMACEUTICAL WORKS LTD., F.P.NO.- 17&amp;16/5,MELDI ESTATE,B/S. MELDI MATA TEMPLE, NR.GOTA RAILWAY CROSSING, AT &amp; POST.- GOTA , TAL : CITY &amp; DIST : AHMEDABAD - 382 481</b>	2B.2	Status of Applicant: <b>Not Applicable</b>
2A.3	Status of Product-license holder: <b>manufactures the dosage form</b>	2B.2.1	For categories b & c have the name and address of the Manufacturer producing the dosage form are: <b>Not Applicable</b>
2A.3.1	For category b and c the name and address of the manufacturer producing the dosage form are <b>Not Applicable</b>	2B.3	Why is marketing authorization lacking? <b>Not Applicable</b>
2A.4	Is Summary Basis of Approval appended? <b>Not Applicable</b>	2B.4	Remarks: <b>Not Applicable</b>
2A.5	Is the attached, officially approved product information complete and consonant with the license? <b>Not Applicable</b>	2B.5	Applicant for certificate, if different from license holder: <b>Not Applicable</b>
2A.6	Applicant for certificate, if different from license holder: <b>No</b>		

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 inspection (years) :

**Once in a Year**

3.2 Has the manufacturer of this type of dosage form been inspected?

**Yes**

3.3 Do the facilities and operations conform to State GMP as per Schedule M of Drugs and Cosmetic Act 1940 and Rules there under?

**Yes**

4 Does the information submitted by the applicant satisfy the certifying authority on all aspects of the manufacture of the products?

**Not Applicable**

**Address of Certifying Authority :**

The Commissioner  
Food & Drugs Control Administration,  
Gujarat State, Jivraj Mehta Bhavan,  
Block No. 8, 1st Floor, Gandhinagar (INDIA)  
Tel:+91-79232-53 417,  
Fax: +91-79232-253400

**Name & Signature :**

(This Document is Digitally Signed)

**Bhavika N. Vyas**

**Asstt. Commissioner**

Food & Drugs Control Administration  
Gujarat State - Gandhinagar

Date of Issue : **11-Dec-2023**



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