	o. of Certificate: 26/1/Drug/20/20	19/8456 31678			Exporting (Certifying) C	ountry: INDIA
In	Date: Importing (Requesting) Country: Afghanistan, Africa, Algeria, Azerbaijan, Bahrin, Baelarus, Bhutan, Bolovia, Brazil, Burkina Faso, Cambodia, Cameroon, Chad, Costa Rica, D.R. Congo Egypt, Equatorial Guines, Guyana, Gabon, Ghana, Gorgia, Indonesia, Iran, Iraq, Israel, Ivory Coast, Jamaica, Jordan Kazakhastan, Kenya, Korea, Kuwait, Kyrgystan, Madagascar, Malawi, Malaysia, Mangolia, Mozambique. Myanmar, Nepal, Nigeria, Oman, Panama, Peru, Philipines, Qatar, Republic of Congo, Republic of Guinea, Romania, Russia, Rwanda, Saudi Arabia, Senegal, Serbia, Singapore, South Africa, Srilanka, Sudan, Syria, Tajikistan, Tanzania, Thailand, Togo, Turkmenistan, UAE, Uganda. Ukraine, Uzbekistan, Venezuela, Vietnam, Yemen.					
		CEI	RTIFICATE OF A PHAR	MACEUTICAL PRODUCT ¹		
1.	1. Name & dosage form of product: COLOVERINE - 135 TABLETS (Mebeverine Tablets BP) 1.1 Active ingredient (s) ² and amount (s) per unit dose ³ (complete qualitative composition including excipients, see attached) Composition: Each film coated tablet contains: Mebeverine Hydrochloride BP Eq. to Mebeverine					
1	3 Is this product actually on the	market in the exporting country	/!	YES	В	The same of the same of
2 F 2 F 2 F	A.2 Product licence holder: M/A.3 Status of product licence holder. A.3.1 For categories b and c the nother dosage form are?: Not A.4 Is summary basis of Approximately.	ame and address of the manufa Applicable	AL INDUSTRIES. REA, DEHRADUN (c)	2B.2 Status of applicant: (2B.2.1 For categories b and dosage form are ⁹ : 2B.3 Why is marketing au Not □	c the names and address of the thorization lacking?	Refused
1	consonant with the licence	¹? Yes \[\] No \[\]	Not provided			
	 A.6 Applicant for certificate if of 3. Does the certifying authority Periodically of routine inspe 3.2 Has the manufacture of this to 3.3 Do the facilities and operation 4. Does the information submitted 	ction (years): Once in ype of dosage form been inspect as conform to GMP as recomm	n of the manufacturing plan a year cted ended by the World Health	organization? ¹⁵ spects of the manufacture of the	Ye	_No
W Ad Di Sa Di	WHO-GMP-CERT.NO. 26/1/Drug/2 ddress of certifying authority: drug Licensing & Controlling Authority: directorate General of Medical Heal ahastradhara Road, Dehradun-248 directorate General of Medical Heal ahastradhara Road, Dehradun-248	0/2019/8456, Valid up to: 14/0 ority th & Family Welfare, 001 (Uttarakhand) (INDIA) th & Family Welfare,	N. M. and St. Company	Name of the author Signature: Stamp & Date Stamp & Date	The Control of the Co	76/20 Asher S

