	\$-15 bis and \$10 b	
No. of Certificate: 26/1/Drug/20/2019/8456 111677		Exporting (Certifying) Country: INDIA
Date: Importing (Requesting) Country:	Rica, D.R. Congo Egypt, Equatorial Guines, Guyana, Gabo Kazakhastan, Kenya, Korea, Kuwait, Kyrgystan, Madagas	Bhutan, Bolovia, Brazil, Burkina Faso, Cambodia, Cameroon, Chad, Costa on, Ghana, Gorgia, Indonesia, Iran, Iraq, Israel, Ivory Coast, Jamaica, Jordan car, Malawi, Malaysia, Mangolia, Mozambique. Myanmar, Nepal, Nigeria, o, Republic of Guinea, Romania, Russia, Rwanda, Saudi Arabia, Senegal, Serbia, Tanzania, Thailand, Togo, Turkmenistan, UAE, Uganda.
	CERTIFICATE OF A PHARM	IACEUTICAL PRODUCT <sup>1</sup>
1.1 Active ingredient (s) <sup>2</sup> and amo Composition:  Each hard Pantopraz Eq. to Par (As enteri	uct: CYCLODOM CAPSULES (Pantoprazole & Dompount (s) per unit dose <sup>3</sup> (complete qualitative composition included gelatin capsule contains: cole Sodium Sesquihydrate attoprazole	Attested  Attested
Excipient Colour: A	sq.s. pproved colours used in empty capsule shells laced on the market for use in the exporting country? complete box B:	YES SUSHMITA SECRETARY YES JUN 2001
THE SHOWING THE RESIDENCE OF THE PERSON OF T	A	B
dt. 24/08/2016  2A.2 Product licence holder: M/ 122,  2A.3 Status of product licence holder: M/ 2A.3.1 For categories b and c the management the dosage form are 9: Not a summary basis of Appro  2A.4 Is summary basis of Appro  2A.5 Is the attached, officially appropriate the dosage form are 9: Not a summary basis of Appro  2A.6 Applicant for certificate if of the dosage form are 9: Not a summary basis of Appro  2A.6 Applicant for certificate if of the dosage form are 9: Not a summary basis of Appropriate and a summary basis of Appropria	name and address of the manufacturer producing Applicable val appended <sup>10</sup> ? Yes	2B.1 Applicant for certificate (name & address)  2B.2 Status of applicant: (a)
Periodically of routine inspe 3.2 Has the manufacture of this to 3.3 Do the facilities and operation	r arrange for periodic inspection of the manufacturing plant in ction (years):  Once in a year  The property of dosage form been inspected as conform to GMP as recommended by the World Health Countries of the applicant satisfy the certifying authority on all aspected.	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
WHO-GMP-CERT.NO. 26/1/Drug/20/20 Address of certifying authority: Drug Licensing & Controlling Authority Directorate General of Medical Health & Sahastradhara Road, Dehradun-248001 ( Directorate General of Medical Health & Sahastradhara Road, Dehradun-248001 (	Family Welfare, (Uttarakhand) (INDIA) Family Welfare,	Name of the authorized person:  Signature:  Stamp & Date

