No. of Certificate: 26/1/Drug/20/2019/8456 11649	Exporting (Certifying) Country: INDIA
Rica, D.R. Congo Egypt, Equatorial Guines, Guyana, Gab Kazakhastan, Kenya, Korea, Kuwait, Kyrgystan, Madaga Oman, Panama, Peru, Philipines, Qatar, Republic of Cong	, Bhutan, Bolovia, Brazil, Burkina Faso, Cambodia, Cameroon, Chad, Costa on, Ghana, Gorgia, Indonesia, Iran, Iraq, Israel, Ivory Coast, Jamaica, Jordan scar, Malawi, Malaysia, Mangolia, Mozambique. Myanmar, Nepal, Nigeria, to, Republic of Guinea, Romania, Russia, Rwanda, Saudi Arabia, Senegal, Serbia, n, Tanzania, Thailand, Togo, Turkmenistan, UAE, Uganda.
CERTIFICATE OF A PHARM	MACEUTICAL PRODUCT ¹
 Name & dosage form of product: T – ZOL 500 TABLETS (Tinidazole Tablets 500 r Active ingredient (s)² and amount (s) per unit dose³ (complete qualitative composition incl Composition: 	uding excipients, see attached)
Each film coated tablet contains:	Attested ATTERICAN CHAMBER OF
Tinidazole USP500 mg	alm Sol
Excipientsq.s.	Sushinity.
Colour: Indigo Carmine Lake & Indigo Carmine Supra	SUSHMITA (2 6 JUN 2000) 3
1.2 Is this product licensed to be placed on the market for use in the exporting country? If yes, complete box A, if no, complete box B:	YES SUSHMITA Secretary 0 0 2 8 1 8
1.3 Is this product actually on the market in the exporting country? ⁵	YES WEWDELM
2A.1 Number of product licence & date of issued: 65/UA/2016 & 65/UA/SC/P-2016	2B.1 Applicant for certificate (name & address)
dt. 24/08/2016 2A.2 Product licence holder: M/s SIGNATURE PHYTOCHEMICAL INDUSTRIES. 122, MI, SELAQUI INDUSTRIAL AREA, DEHRADUN	2B.2 Status of applicant: (a)
2A.3 Status of product licence holder ⁸ : (a,	2B.3 Why is marketing authorization lacking? Not
2A.4 Is summary basis of Approval appended 10? Yes No 2A.5 Is the attached, officially approved product information complete and consonant with the licence 11? Yes No Not provided 2A.6 Applicant for certificate if different from licenced holder: Not Applicable 12	2B.4 Remark: ¹³
 Does the certifying authority arrange for periodic inspection of the manufacturing plant Periodically of routine inspection (years): Once in a year Has the manufacture of this type of dosage form been inspected Do the facilities and operations conform to GMP as recommended by the World Health Q Does the information submitted by the applicant satisfy the certifying authority on all aspects. 	Yes ✓ No ☐ Yes ✓ No ☐ Yes ✓ No ☐
WHO-GMP-CERT.NO. 26/1/Drug/20/2019/8456, Valid up to: 14/05/2023 Address of certifying authority: Drug Licensing & Controlling Authority Directorate General of Medical Health & Family Welfare, Sahastradhara Road, Dehradun-248001 (Uttarakhand) (INDIA)	Name of the authorized person: Signature: Stamp & Date Stamp & Date Stamp & Date

