No. of Certificate: 26/1/Drug/20/2019/8456 31676		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 scar, Malawi, Malaysia, Mangolia, Mozambique. Myanmar, Nepal, Nigeria, o, Republic of Guinea, Romania, Russia, Rwanda, Saudi Arabia, Senegal, Serbia, t, Tanzania, Thailand, Togo, Turkmenistan, UAE, Uganda.
A STAR STAR	CERTIFICATE OF A PHARM	MACEUTICAL PRODUCT <sup>1</sup>
1.1 Active ingredient (s) <sup>2</sup> and amo Composition:	uct: DUSTALOSINE CAPSULES (Dutasteride & Tamsount (s) per unit dose <sup>3</sup> (complete qualitative composition included)	sulosin Hydrochloride Capsules) uding excipients, see attached)
Each har	rd gelatin capsule contains:	
Tamsulos	sine hydrochloride BP	IAMBER OF
(as Sustai	ned Release Pellets)	Attested
Dutasterio	de BP	Milestea
	diate Release Pellets)	
Excipient	tsq.s.	Sughuits. (2 6 JUN 2020) (5)
Colour: A	Approved colours used in empty capsule shells	SUSHMITA
to the special and the second	Control of the control of the second of the	Socretory
	placed on the market for use in the exporting country?	
If yes, complete box A, if no		NO 0 0 0 0 1 5 WEWDELH
1.3 Is this product actually on the	market in the exporting country?	YES
The Strate has some board and the	The state of the Arms and the state of the s	B
2A.1 Number of product licence <sup>7</sup> & date of issued: 65/UA/2016 & 65/UA/SC/P-2016		2B.1 Applicant for certificate (name & address)
dt. 24/08/2016		2E.2 Status of applicant: (a) (b) (c) (d)
2A.2 Product licence holder: M/s SIGNATURE PHYTOCHEMICAL INDUSTRIES. 122, MI, SELAQUI INDUSTRIAL AREA, DEHRADUN		2B.2.1 For categories b and c the names and address of the manufacturer producing the dosage form are 9:
2A.3 Status of product licence h	older <sup>8</sup> : (a)	
2A.3.1 For categories b and c the name and address of the manufacturer producing		2B.3 Why is marketing authorization lacking?
the dosage form are?: Not Applicable		Not
2A.4 Is summary basis of Appro		Required Requested Consideration
	pproved product information complete and	2B.4 Remark: <sup>13</sup>
consonant with the licence	Proved product information complete and 1.11? Yes $\square$ No $\square$ Not provided $\square$	2D.4 Remark.
24.6 Applicant for cortificate if	different from licenced holder: Not Applicable 12	
2A.6 Applicant for certificate if	y arrange for periodic inspection of the manufacturing plant	in which the dosage form is produced? Yes √ No NA
		in which the dosage form is produced? Yes V No NA
Periodically of routine inspe		Yes ☑ No □
3.2 Has the manufacture of this	type of dosage form been inspected	
3.3 Do the facilities and operation	ons conform to GMP as recommended by the World Health (	
4. Does the information submit	tted by the applicant satisfy the certifying authority on all asp	sects of the manufacture of the product?
WHO-GMP-CERT.NO. 26/1/Drug/20/20	019/8456. Valid up to: 14/05/2023	Name of the authorized person:
Address of certifying authority:		
Drug Licensing & Controlling Authority		Signature:
Directorate General of Medical Health & Family Welfare, Sahastradhara Road, Dehradun-248001 (Uttarakhand) (INDIA)		Stamp & Date
Directorate General of Medical Health & Family Welfare,		19/6/2000 Silve Authority I'm
Sahastradhara Road, Dehradun-248001 (Uttarakhand) (INDIA)		Stamp & Date  Stamp & Date  Stamp & Date  Stamp & Licensing Authority (Miles)  (Uttarakhand)
		Controlling Ittarakhario
		Third -

