	No. of Certificate: 26/1/Drug/20/2019/8456 11685	Exporting (Certifying) Country: INDIA
	Date:	
	Kazakhastan, Kenya, Korea, Kuwait, Kyrgystan, Mada Oman, Panama, Peru, Philipines, Qatar, Republic of Co	nus, Bhutan, Bolovia, Brazil, Burkina Faso, Cambodia, Cameroon, Chad, Costa abon, Ghana, Gorgia, Indonesia, Iran, Iraq, Israel, Ivory Coast, Jamaica, Jordan gascar, Malawi, Malaysia, Mangolia, Mozambique. Myanmar, Nepal, Nigeria, ngo, Republic of Guinea, Romania, Russia, Rwanda, Saudi Arabia, Senegal, Serbia, tan, Tanzania, Thailand, Togo, Turkmenistan, UAE, Uganda.
		RMACEUTICAL PRODUCT <sup>1</sup>
	<ol> <li>Name &amp; dosage form of product: ASIGREL – A FORTE CAPSULES (Aspirin, Atorvastatin Calcium &amp; Clopidogrel Bisulphate Capsules)</li> <li>Active ingredient (s)<sup>2</sup> and amount (s) per unit dose<sup>3</sup> (complete qualitative composition including excipients, see attached)</li> <li>Composition:</li> </ol>	
	Each hard gelatin capsule contains: Aspirin USP (enteric coated pellets)	5 mg
	Atorvastatin Calcium USP  Equivalent to Atorvastatin (as pellets)	YES  YES  Sushin Ta  Sushin Ta  Secretary  YES  O 0 2 8 2 4  NEW DELM:
	A	В
	<ul> <li>2A.1 Number of product licence<sup>7</sup> &amp; date of issued: 65/UA/2016 &amp; 65/UA/SC/P-2016 dt. 24/08/2016</li> <li>2A.2 Product licence holder: M/s SIGNATURE PHYTOCHEMICAL INDUSTRIES.         <ul> <li>122, MI, SELAQUI INDUSTRIAL AREA, DEHRADUN</li> </ul> </li> </ul>	2B.1 Applicant for certificate (name & address)  2B.2 Status of applicant: (a)
	2A.3 Status of product licence holder <sup>8</sup> : (a)	2B.3 Why is marketing authorization lacking?  Not Not Under Required Requested Consideration  2B.4 Remark: 13
	3. Does the certifying authority arrange for periodic inspection of the manufacturing plant in which the dosage form is produced?  Periodically of routine inspection (years):  Once in a year  3.2 Has the manufacture of this type of dosage form been inspected  3.3 Do the facilities and operations conform to GMP as recommended by the World Health Organization?¹⁵  4. Does the information submitted by the applicant satisfy the certifying authority on all aspects of the manufacture of the product?¹⁶  Yes ✓ No ☐  Yes ✓ No ☐	
4	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) Directorate General of Medical Health & Family Welfare, Sahastradhara Road, Dehradun-248001 (Uttarakhand) (INDIA)	Name of the authorized person:  Signature:  Stamp & Date

