GOVERNMENT OF TELANGANA DRUGS CONTROL ADMINISTRATION Vengalarao Nagar, Hyderabad 500 038

CERTIFICATE OF PHARMACEUTICAL PRODUCT

This certificate conforms to the format recommended by the World Health Organization (General Instructions and explanatory notes attached.)

No. of Co	ertificate	: 1032/DI/ML	K/TST/	COPP/09	1219		Valid up to: 08/12/2	2021			
Exporting	g (certify	ing) country:	INDLA	A							
Importing	g (request	ting) country:	ТАЛІ	KISTAN							
1. Name	e and dos	age form of the p	roduct:	DOXORU	BICIN HY	DROCHLORI	IDE INJECTION USP	10 mg			
1.1 Activ	e Ingredient (S) ² and amounts (S) per unit dose ³ : Each Lyophilized vial contains										
	Doxoru	bicin Hydrochlor	ride	USP	10 mg						
	Lactose	e Monohydrate		USP	50 mg						
1.2	Is this product licensed to be placed on the market for use in the exporting country? 5 (Key in as appropriate)										
	Yes	\boxtimes			No 🗌						
1.3	Is this product actually on the market in the exporting country?										
	Yes	\boxtimes			No 🗌		Unknown				
	If the answer to 1.2 is yes, continue with section 2A and omit section 2B.										
	If the answer to 1.2 is no, omit section 2A and continue with section 2B6										
S	ECTION	i 2A									
2.A.1	Number of product Licence ⁷ and date of issue: 22/RR/TS/2015/F/G, Dated: 13.01.2015										
2.A.2	Product license holder (Name and address) :			Plot.No Jeedime	GLS PHARMA LIMITED Plot.No. 10,IDA, Phase-I Jeedimetla, R.R.Dist, Hyderabad, Telangana, INDIA						
2.A.3	Status of product – license holder ⁸ (Key is appropriate category as defined in note (8)										
	a) 🛛			b)			c) 🔲				
2A.3.1	For categories b and c the name and address of the Manufacturer producing the dosage form is ⁹ ?										
	Yes	l		No			Not applicable				
2.A.4	Is summary basis for approval appended 10 ? (enclosed at the time of product approval)										
	Yes 🗵	I		No			Not applicable				
2.A.5	Is the attached, officially approved product information complete and consonant with the license? ¹¹ (key as appropriate)										
2. A.6	Yes 🛭 Applica	nt for certificate, i	if differe	No nt from lice	-	Name & Address	social applicable				
	Yes 🗌			No		WEDE (SOL)	Not applicable FEREN PROPERTY OF THE PROPERTY				

SECTION 2B IS TO BE OMITTED

2. B.1	Applicant for certificate (Name & address)										
2. B.2	Status of applicant: (Key in the appropriate category as defined in note 8)										
2. B.2.1	For categories b and c the name and address of the manufacturer producing the dosage from is 9:										
2. B.3	Why is marketing authorization lacking? Not required / Not requested / under consideration / Refused (Key in as appropriate)										
2. B.4	Remarks: ¹³										
3,	Dose the certifying authority arrange for periodic inspection of the manufacturing plant in which the dosage form is produced?										
	Yes	\boxtimes		No 🗌	Not applicable ¹⁴						
	If not or not applicable, proceed to question 4.										
	Periodicity of routine inspections (years): NOT LESS THAN ONCE A YEAR										
	Has the manufacturer of this type of dosage form been inspected Yes/No (Key in as appropriate)										
	Yes			No 🗌	Not applicable						
	Do the facilities and operations conform to GMP as recommended by the World Health Organisation 15?										
	Yes			No 🗌	Not applicable						
l .	Does the information submitted by the applicant satisfy the certifying authority on all aspects of the manufacturer of the product ? ¹⁶										
	Yes	\boxtimes		No 🗌	Not applicable						
	Address	of certifying authority	: Office of the Deputy Director Drugs Control Administration, Vengalarao Nagar, Hyderabad 500 038, Telangana, INDIA.								
	Telephone and Fax numbers		:	TEL: +91 40 23814119	FAX: +91 40 23814360						
	Name of	Authorized Person	Dr. B. VENKATESHWARLU DEPUTY DIRECTOR & CERTIFYING AUTHORITY								
	Signatur	e									
	Stamp a	nd Date	DEP	Dr. B. VENKATESHWARLU DEPUTY DIRECTOR & CERTIFYING AUTHORIT							









DRUGS CONTROL ADMINISTRATION Government of Telangana



Dated 15-02-2020

To

M/s. GLS Pharma Limited, Plot.No.10, Phase – I, IDA., Jeedimetla, Medchal-Malkajgiri District – 500 055, Telangana, INDIA.

Sirs.

Sub: Drugs and Cosmetics Act, 1940 and Rules made thereunder – Issue of World

Health Organization Good Manufacturing Practice Certificate - Regarding.

Ref: 1. Your application dated 05.11.2018.

2. Joint Inspection Report dated 20.02.2019 & 21.02.2019.

3. Compliance Verification Report dated 05.12.2019.

4. Lr.No. 5-6(490 A1)/2018/7492, dated 13.02.2020 of Deputy Drugs Controller(India), CDSCO, Hyderabad

(a, a)

I forward herewith WORLD HEALTH ORGANIZATION GOOD MANUFACTURING PRACTICE CERTIFICATE for the products recommended by the Joint Inspection Team consisting of officers of Central Drugs Standard Control Organization and officer from Drugs Control Administration, Telangana for Export Purpose.

This Certificate is valid for a period of Three years from the date of issue.

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Yours faithfully,

Dr. B. VENKATESHWARLU

JOINT DIRECTOR(FAC)

DRUGS CONTROL ADMINISTRATION

