



## **CERTIFICATE OF ANALYSIS**

Product:

Ivermectin

Batch no .:

05KB21.HM00173

Retest Date:

January 2023

Manufacturing Date: January 2021

TEST	REF.	SPECIFICATION	RESULT
Description:	AATG018_001	White to yellowish white powder	Conforms
Crystallinity:	USP, Monograph	The product is crystalline	Conforms
Identification (by IR):	EP, Monograph	Conforms to the spectrum of the European Pharmacopeia	Conforms
Identification (by HPLC):	CRLC177_001	Chemical Reference Substance In the chromatogram obtained with the sample solution, the principal peaks should have the same RTs as those of the principal	Conforms
Appearance of solution:	EP, Monograph	peaks in the chromatogram obtained with the reference standard A 2% w/v solution in toluene (1g/50mL) is clear and not more intensely coloured than the reference solution BY7.	Conforms
Water:	AA001835_002	Not more than 1.0 % w/w (2g)	0.1 % w/w
Residue on ignition:	USP, Monograph	Not more than 0.1 % w/w (1 g; 600°C; constant weight)	0.0 % w/w
Specific optical rotation:	AA001833_003	Not less than -26° and not more than -17°, calculated with reference to the anhydrous and solvent-free substance (c=2.5; methanol; t=20° C; 589.3nm)	-19 °
Ratio (by HPLC): Iv. B1a / (Iv. B1 a + B1b):	CRLC177_003	Not less than 90.0 %	98.6 %
Related substances (by HPLC):			
EP Impurity A	CRLC177_025	Not more than 1.0 % w/w	0.4 % w/w
EP Impurity H	CRLC177_026	Not more than 1.0 % w/w	0.2 % w/w
EP Impurity D	CRLC177_027	Not more than 1.0 % w/w	Less than 0.05 % w/w
EP Impurity E	CRLC177_036	Not more than 1.0 % w/w	0.2 % w/w
Impurity with RRT 0.87 (Impurity 1)	CRLC177_028	Not more than 0.2 % w/w	0.1 % w/w
Impurity with RRT 0.55	CRLC177_011	Not more than 0.4 % w/w	0.2 % w/w
Impurity with RRT 0.77	CRLC177_012	Not more than 0.4 % w/w	0.1 % w/w
Impurity with RRT 0.59	CRLC177_038	Not more than 0.5 % w/w	Less than 0.05 % w/w
Any unspecified impurity	CRLC177_034	Not more than 0.1 % w/w	0.1 % w/w
Sum of Impurities between RRT 1.3 -1.5 (Impurity K)	CRLC177_035	Not more than 2.5 % w/w	1.4 % w/w
Total impurities	CRLC177_018	Not more than 5 % w/w	3 % w/w
Residual solvents (by GC):		AND DESCRIPTION OF THE PROPERTY OF THE PROPERT	
Formamide	CRGC098_008	Not more than 3.0 % w/w	2.3 % w/w
Ethanol	CRGC098_009	Not more than 5.0 % w/w	4.2 % w/w
Methanol	CRGC175_001	Not more than 0.3 % w/w	0.0 % w/w
Assay (by HPLC):	CRLC177_002	Not less than 95.0 % w/w and not more than 102.0 % w/w , calculated with reference to the anhydrous and solvent-free substance	98.6 % w/w

The batch number 05KB21.HM00173 of Ivermectin has been tested as above and conforms to the latest EP and Hovione specifications.	Approved by:  Kim Lao 29.Jan.2021 12:50:57  Quality Control			
Storage conditions: Protected from the incident of sunlight and/ or artificial light; store below 25 °C and 65% RH				
The batch was manufactured according to Good Manufacturing Practices.	Released by:  Bruna Nunes 04.Feb.2021 12:09:56  Quality Assurance			
Reference: 198428, 198390	GQSP1314.8 E_NORMAL			

This document has been signed electronically in compliance with 21CFR Part 11.

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By: Jonadel Hufana