Sun Pharmaceutical Industries Ltd.

A-7/A-8, M.I.D.C Industrial Area,

Ahmednagar 414 111, Maharashtra, INDIA. Tel.: (91-241) 2777329, 2777330, 2777359 Fax: (91-241) 2777231

www.sunpharma.com

CIN: L24230GJ1993PLC019050



CERTIFICATE OF ANALYSIS

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Order No.: 073	30025468		1460 01 01 02
Product	MESALAZINE Ph. Eur.	A.R.No.	QFP/17/1592
Batch No.	ASANF17028	Mfg date	October 2017
Release date	24.10.17	Exp. Date	September 2022

Sr.	Test	Dorolle	C .c
1	Characteristics	Results	Specification
387		A1	
1.1	Appearance	Almost white powder.	Almost white or light Grey or light pink powder or crystals.
1.2	Solubility	Very slightly soluble in water. Practically insoluble in Ethanol (96%). It dissolves in dilute solutions of alkali hydroxides and in dilute hydrochloric acid.	Practically insoluble in Ethanol (96%).
2(\$)	Identification		
2.2	B) Infrared absorption:	Infrared absorption spectrum in KBr dispersion of sample is concordant with the spectrum of Mesalamine working standard.	Infrared absorption spectrum in KBr dispersion of sample should be concordant with the spectrum of Mesalamine working standard.
3	Appearance of solution		
3.1	Appearance of solution	Solution is clear	Solution should be clear
3.2	Absorbance at 440 nm	0.047 Au	Not more than 0.15 Au.
3.3	Absorbance at 650 nm	0.01 Au	Not more than 0.10 Au.
4	Reducing Substances	The colour of solution is blue	The colour of solution should be blue or violet brown.
5	Related Substances (By HPLC)		
	Known Impurities		
	Impurity E + Impurity D	0.008 %	Not more than 0.05%
	Impurity F	BQL	Not more than 0.1%
	Impurity G	0.002 %	Not more than 0.05%
	Impurity H	BQL	Not more than 0.1%
	Impurity J	BQL	Not more than 0.1%
	Impurity L	BDL	Not more than 0.05%
	Impurity M	BDL	Not more than 0.05%
	Impurity O	BQL	Not more than 0.1%
	Impurity P	BQL	Not more than 0.1%
	Impurity R	BDL	Not more than 0.05%
	UnKnown Impurities		
	Unspecified impurities	BDL	Not more than 0.05 %
	Total impurities (known + Unknown	0.01 %	Not more than 0.5%
6	Content - by HPLC		
	Impurity A	Not Detected	Not more than 200ppm
	Impurity C	Not Detected	Not more than 200ppm

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Batch No.	ASANF17028	Mfg date	October 2017
Release date	24.10.17	Exp. Date	September 2022

Specification No.: BD0270P0DA Rev. No.: 7.0					
Sr.	Test	Results	Specification		
7.	Aniline (Impurity K) (By HPLC) *	No potential	Not more than 10 ppm		
8	Chloride	0.0 % w/w	Not more than 0.1 % w/w.		
9	Sulphates .	Less than 200 ppm.	Not more than 200 ppm.		
10	Heavy metals	Less than 10 ppm.	Not more than 10 ppm.		
11	Loss on drying	0.25 % w/w	Not more than 0.5 % w/w.		
12	Sulphated Ash	0.06 % w/w	Not more than 0.2 % w/w.		
13	Assay by titrimetric	100.4 % w/w	Between 98.5 w/w % and 101.5 % w/w. (On dried basis)		
14	Residual Solvents (By GC)				
	Methanol	BQL	Not more than 100 ppm		
15	Bulk Density				
15.1	Bulk Density	0.28 gm/ml	Between 0.2 gm/ml and 0.35 gm/ml		
15.2	Tapped Density	0.55 gm/ml	Between 0.35 gm/ml and 0.60 gm/ml		
16	Sulphanilic acid * (By HPLC)	No potential	Not more than 0.10 %		
17	Hydrogen sulfide and sulfur dioxide	Complies	Lead acetate test paper should not become discoloured.		
18.	pН	4.05	Between 3.50 and 4.50		
19.	Particle size (By Malvern)	10 % = 3 micron 50 % = 11 micron 90 % = 44 micron	10 % between 2 to 6 micron 50 % between 8 to 20 micron 90 % between 25 to 50 micron		
20	Total aerobic microbial count	Less than 10 CFU/gm	Not more than 1000 CFU/gm		
21	Total combined yeast & Mould count	Less than 10 CFU/gm	Not more than 100 CFU/gm		
22	Escherichia coli	Not detected	Should be absent		
23	Salmonella species	Not detected	Should be absent		
24	Pseudomonas aeruginosa	Not detected	Should be absent		
25	Staphylococcus aureus	Not detected	Should be absent		

* = No potential for this impurity as it is not being used in the manufacturing process.

Conclusion: Product complies with the quality standards as per Ph. Eur., In-house & Customer's specification.

Date of issue: 30.10.17

Prepared by Vunsule

Checked by 30.10-17

Approved h

V. V. Sangale - Executive QA

S.N. Dhandar - Manager QA

K.D. Gaikwad - Manager QA

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