

ISO 9001: 2008 Certified

NATCO PHARMA LIMITED

Regd. Off.: 'NATCO HOUSE', Road No. 2, Banjara Hills, Hyderabad - 500 034. Telangana, INDIA. Tel: +91 40 23547532, Fax: +91 40 23548243 CIN: L24230TG1981PLC003201, www.natcopharma.co.in



CERTIFICATE OF ANALYSIS

Product name: Velpanat (Sofosbu	Batch No.: 1900983	
Batch size: 100,000 Tablets	Sampling Date: 22/06/2018	Mfg. Date: 07/2018
Qty. Sampled: 20 Tablets	Analysis Date: 22/06/2018	Exp. Date: 06/2020
Sampled by: A.Anvesh Kumar	Reporting Date: 28/06/2018	A.R.No.: U4/FP/1014/18

S. No.	TEST	R	ESULT	SPECIFICATION		
1	Description	Blue coloured, oval shaped, film-coated tablets debossed with 'SL' on one side and plain on other side.		Blue coloured, oval shaped, film-coated tablets debossed with 'SL' on one side and plain on other side.		
2	Identification					
	a) By HPLC	•	tention time with the standard as obtained in the	The retention time of the major peak in the chromatogram of the sample preparation corresponds to that in the chromatogram of the standard preparation, as obtained in the Assay.		
	b) By UV	•	ima of the standard ectra exhibit at ghts.	The UV absorption spectrum of the sample solution and standard solution shall exhibit maxima at the same wavelenght.		
3	Uniformity of dosage units (By content uniformity)	Velpatasvir = 1.3 Sofosbuvir = 1.4		The acceptance value of the first 10 dosage units is less than or equal to L1 (L1 is 15.0 and L2 is 25.0)		
4	Average weight	1026.8 mg		1030.0mg±5.0%		
5	Water content	1.75 % w/w		Not more than 5.0% w/w		
	Dissolution (By HPLC)					
6	Velpatasvir	100.2%	99.7%	Not less than 80% (Q) of the labeled amount of Ledipasvir and Sofosbuvir are dissolved in 45 minutes.		
		98.7%	97.3%			
		98.8%	99.9%			
	Sofosbuvir	102.3%	100.8%			
		100.0%	99.0%			
		100.3%	102.2%			

Prepared by:

Reviewed by:

Approved:

1080

Date: 28/06/2018

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S. No.	TEST	RESULT	SPECIFICATION			
7	Assay (By HPLC): Each film coated tablet contains.					
	Velaptasvir: 100 mg	98.9%	Not less than 95.0% and Not more than 105.0% of the labeled amount of Velpatasvir.			
	Sofosbuvir: 400 mg	99.6%	Not less than 95.0% and Not more than 105.0% of the labeled amount of Sofosbuvir.			
8	Related impurities (% w/w, By HPLC)					
	a) Sofosbuvir					
	Any individual impurity	Less than 0.05%	Not more than 0.30%			
	Total impurities	Less than 0.05%	Not more than 1.0%			
	b) Velpatasvir					
	Keto impurity	Less than LOQ (0.048%)	Not more than 0.8%			
	Any Individual unspecifield impurity	Less than 0.05%	Not more than 0.20%			
	Total impurities	Less than 0.05%	Not more than 1.20%			
9	Microbial Enumeration tests and Test for specified microorganisms					
	Total aerobic microbial count	Less than 10 gfu/g	Not more than 1000 cfu/g			
	Total combined molds and yeasts	Less than 10 gfu/g	Not more than 100 cfu/g			
	Escherichia coli	Adsent	Should be absent/g			
	Salmonella species	Adsent	Should be absent/10g			
	Pseudomonas aeruginosa	Adsent	Should be absent/g			
	Staphylococcus aureus	Adsent	Should be absent/g			

Remarks: The product Conforms / Does not conforms to Specification No.: K/FPS/436

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