Gandla. Kumara Swamy. et al. /Asian Journal of Research in Chemistry and Pharmaceutical Sciences. 5(3), 2017, 114-122.

Research Article



Asian Journal of Research in Chemistry and

Pharmaceutical Sciences Journal home page: www.ajrcps.com



NOVEL STABILTY INDICATING RP-HPLC METHOD SIMULTANEOUS DETERMINATION OF SOFOSBUVIR AND VELPATASVIR IN BULK AND COMBINED TABLET DOSAGE FORMS

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ABSTRACT

A accurate and precise RP-HPLC method has been developed for the validated of Sofosbuvir and Velpatasvir in bulk and combined Tablet dosage form. Separation was carried out on a Primesil C₁₈ (4.6 x 250mm, 5 μ m) column using a mixture of Acetonitrile: 0.1% perchloricacid (50:50 v/v) as the mobile phase at a flow rate of 1.2mL/min, The detection was carried out at 262 nm. The retention time of the Sofosbuvir and Velpatasvir 4.25, 5.91 min respectively. The method produce linear responses in the concentration range of 25-150 µg/mL for Velpatasvir, and 100-600µg/ml of Sofosbuvir. The method precision for the determination of assay was below 2.0 %RSD. The method is useful in the quality control of bulk and pharmaceutical formulations.

KEYWORDS

Sofosbuvir and Velpatasvir, RP-HPLC, UV-VIS Detection and Validation.

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INTRODUCTON

Sofosbuvir

Sofosbuvir is a medication used for the treatment of hepatitis C. It is only recommended with some combination of ribavirin, peginterferon-alfa, simeprevir, ledipasvir, or daclatasvir. Cure rates are 30 to 97% depending on the type of hepatitis C virus involved. Safety during pregnancy is unclear; while, some of the medications used in combination may result in harm to the baby. It is taken by mouth and chemically it is Isopropyl (2S)-2-[[[(2R, 3R, 4R, 5R)-5-(2, 4-dioxopyrimidin-1-yl)-4-fluoro-3hydroxy-4-methyl-tetrahydrofuran-2-yl] methoxyphenoxy-phosphoryl] amino] propanoate Molecular formula C₂₂H₂₉FN₃O₉P Molecular Weight 529.453

g/mol and Soluble in Methanol, Acetonitrile and water.

Velpatasvir is an NS5A inhibitor which is used together with sofosbuvir in the treatment of hepatitis C infection of all six major genotypes.

MATERIAL AND METHODS¹³⁻¹⁴

Material and Reagents

The reference standards of Sofosbuvir and Velpatasvir were procured from Sura Pharma Labs, Dilshuknagar, Hyderabad, India. The branded tablet formulation Epclusa (sofosbuvir 400 mg and velpatasvir 100 mg) was purchased from the local market. All the HPLC solvents and analytical reagent grade chemicals were purchased from S.D. Fine Chemicals, Hyderabad, India.

Instrumentation

A Shimazu HPLC system equipped with a SPD-20AD binary pump, an auto sampler and a UV-VIS detector was employed for the study. The output signal was monitored and processed with LC-Solution software.

Chromatographic conditions

The separation of the drugs was achieved on a Discovery® C18 HPLC Column (250 x 4.6 mm; 5µ particle size) by running a mobile phase containing a 50:50 v/v mixture of 0.1% perchloric acid in water and acetonitrile at a flow rate of 1.2 mL/min. The injection volume was 10μ L. The column temperature was maintained at 30°C and the analytes in the eluates were monitored at 262 nm. The run time was 9.0 min. A 50:50 v/v mixture of 0.1% perchloric acid and acetonitrile was used as the diluent to prepare drug solutions.

Preparation of standard solution:

Accurately weighed and transferred 25 mg of Velpatasvir and Sofosbuvir working standard into 25 mL of clean dry volumetric flask add about 15mL of Diluent and sonicate to dissolve and removal of air completely and make volume up to the mark with the same Methanol.

Tablet sample solution

Twenty tablets of "Epclusa" (velpatasvir 100 mg and sofosbuvir 400 mg) were accurately weighed and the average weight of the tablet was calculated. The tablets were finely powdered and a quantity of

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the powder equivalent to one tablet was transferred into a 100mL volumetric flask. 70mL of the diluent was added to it and sonicated for 5 minutes. Then the volume was made up with the diluent and mixed well to prepare the sample stock solution. This solution was filtered through a 0.45µm nylon filter. 2.0mL of the filtrate was transferred to a 20mL volumetric flask and the volume made up to give final theoretical concentrations of 100µg/mL and 400µg/mL of Sofosbuvir and Velpatasvir respectively

Method development and Validation

Different mobile phases were considered for simultaneous separation of the two drugs on a Primesil C_{18} HPLC Column. Selection of the mobile phase was done on the basis of ideal resolution among Sofosbuvir and Velpatasvir and also their impurities formed during forced degradation studies. The required chromatographic conditions were optimized. The developed method was validated for precision, specificity, accuracy (recovery), linearity and robustness as per the ICH guidelines⁷.

System Suitability

System suitability was established for initial evaluation of the method before running the sample for the validation parameters. The test was performed according to the USP¹⁰. The standard solutions prepared as per the proposed method were analyzed. The results of the system suitability study are presented in Figure 3. The acceptance criterion is % RSD \leq 2.0. A percent % RSD Sofosbuvir and Velpatasvir of 0.62 and 0.92 indicates good system precision of the method. The tailing factor obtained from the standard injection is 1.69 and 2.18 and Theoretical plates obtained from the standard injection are 4710 and 5840 respectively.

Linearity

The linearity were observed for in the concentration rages from $25-150\mu$ g/mL for Velpatasvir and 100- 600μ g/mL. The Linearity of the method was demonstrated by preparing different concentrations of drug substance and analyzing as per the proposed method. A plot of the area of the peak as a function of analyte concentration was prepared and its regression equation computed. The linearity data of

the two drugs are given in Table No.6 and Figure No.4.

From the above the LOD values of Sofosbuvir and Velpatasvir were found to be 7.71 and $02.74\mu g/m l$ respectively. The LOQ values of Sofosbuvir and Velpatasvir were found to be 23.37 and $8.31\mu g/m l$ respectively. Thus the method developed was found to be sensitive.

Precision

In the precision study,% RSD was found to be less than 2 % for Velpatasvir 0.6% and Sofosbuvir 0.2 which indicates the system has a good reproducibility for precision studies 5 replicate studies of Sofosbuvir and Velpatasvir formulation (method precision) was performed.% RSD was determined for peak areas of Sofosbuvir and Velpatasvir and the acceptance limits should be NMT 2% and the results were found to be within the acceptance limits The chromatograms of precision were showed in Figures No.7.22-7.26. The results were reported in Table No.7.25

Accuracy

The accuracy studies were shown as % recovery for Sofosbuvir and Velpatasvir at 50%, 100%, 150% ,the limits of recovery should be in range of 98-102% the limits obtained for Sofosbuvir and Velpatasvir were found to be within the limits. Hence the method was found to be accurate. The accuracy studies shows % recovery of the Velpatasvir 100% and Sofosbuvir and the limits of % recovery of drugs were 98-102% and from the above results its indicates that the method was accurate and also revealed that the commonly used exciepients present in the pharmaceutical information do not interfere in the proposed method. The chromatograms of shown in results were shown Tables No.7 and 8.

FORCED DEGRADATION STUDIES Acid degradation

Degradation was observed by the additon of 0.5 N HCl.

Alkaline degradation

Degradation was observed by the additon of 0.5N NaoH.

Thermal degradation

Degradation was observed when the sample solution was kept under heat at $60-80^{\circ}$ C for 3 hours.

Peroxide degradation

Degradation was observed by the additon of 3% H₂O₂.

Hydrolysis degradation

Degradation was observed by sunlight exposre.

Table No.1: Optimized Chromatographic condition					
S.No	Parameters	Chromatographic conditions			
1	Mobile phase ratio	Acetonitrile: Water(50:50% v/v)			
2	Column	Primesil C ₁₈ (4.6×250r	nm) 5µ		
3	Detector	UV-VIS Detector)r		
4	Column temperature	Ambient	Ambient		
5	Wavelength	262 nm			
6	Flow rate	1.2 ml/min			
7	Injection volume	10 µl			
8	Run time	Run time 9 minutes			
Table No.2: Result of system suitability parameters					
S.No	Parameter	Sofosbuvir	Velapatasvir		
1	Retention time	4.23	5.91		
-					

 Table No.1: Optimized Chromatographic condition

Tuble 1002. Result of system sufability parameters					
S.No	Parameter	Sofosbuvir	Velapatasvir		
1	Retention time	4.23	5.91		
2	Theoretical plates	4710	5840		
3	Tailing factor	1.68	2.18		
4	Area	2275297	1815105		

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		Table N	o.3: Lin	earity Resu	lts of Velpatasvir			
S.No	Linearity	y Level		Concentration (µg/mL)		Pe	ak Area	
1	1				25			80022
2	2				50		1443902	
3	3				75		2	139373
4	4				100		2	856272
5	5				125		3.	589707
6	6				150		42	220592
	Correlation	coefficient			0.99	9%		
		Table N	0.4: Lin	earity Resu	lts of Sofosbuvir			
S.No	Linearity	y Level		Co	ncentration (µg/mL)		Pea	ık Area
1	1				100		13	16161
2	2				200		27	30754
3	3				300		42	57943
4	4				400	5904229		
5	5				500	7185983		
6	6				600		87	90138
7	Correlation of	coefficient	0.999%					
Lim	it of detection and Li	mit of Quanti	fication	(LOD and	LOQ)			
		Tab	le No.5:	Data of LO	OD and LOQ		-	
S.No	Drug			LOD		Ι	JOQ	
1	<u>^</u>	Velpatasvir 2.74				8.31		
2	Sofosbuvir 7.71			2	3.37			
]	Fable No	o.6: Data of	precision	-		
S.No	No. Inj	ections		Velpatasvir Peak Area		Sof	osbuvir Po	eak Area
1	Injec	Injection1			2475114	6397432		32
2	Injec	tion2			2475284		6396243	
3	Injec	tion3		2528371			6468822	
4	Injec			2515584			6462955	
5	8			2515624			6463145	
	8	Injection5						
6	Injection6		2515624		6462841			
7		Average 2504268		6438573				
8	S.	D		23052		34985		5
9	% R	RSD		0.92		0.62		
I		Table N	0.7: Acc	uracy Resu	lts of Velpatasvir			
S.No	% Concentration (at specification Level)	Peak Area	Amou	int added opm)	Amount found (ppm)	% Re	ecovery	Mean Recover

50

100

150

2075760

2676350

3064030

50%

100%

150%

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1 2

3

99.8

99.8

100.1

99.8%

117

49.9

99.8

151.0

S.No	% Concentration (at specification Level)	Peak Area	Amount Added (ppm)	Amount Found (ppm)	% Recovery	Mean Recovery
1	50%	8971513	50	49.9	99.8%	00.70/
2	100%	12607635	100	99.8	99.4%	99.7%
3	150%	14838255	150	150.1	99.2%	

Table No.8: Accuracy Results of Sofosbuvir

Robustness

Table No.9: System suitability data of Results for Sofosbuvir and Velpatasvir

	Flow Rate (ml/min)		System Suitability Results			
S.No		Name of drugs	USP Plate count	USP Tailing	Retention time(min)	
1	1.2mL/min	Sofosbuvir	8971513	2.18	4.22	
		Velpatasvir	2075760	1.68	5.91	

Table No.10: Data of degradation studies

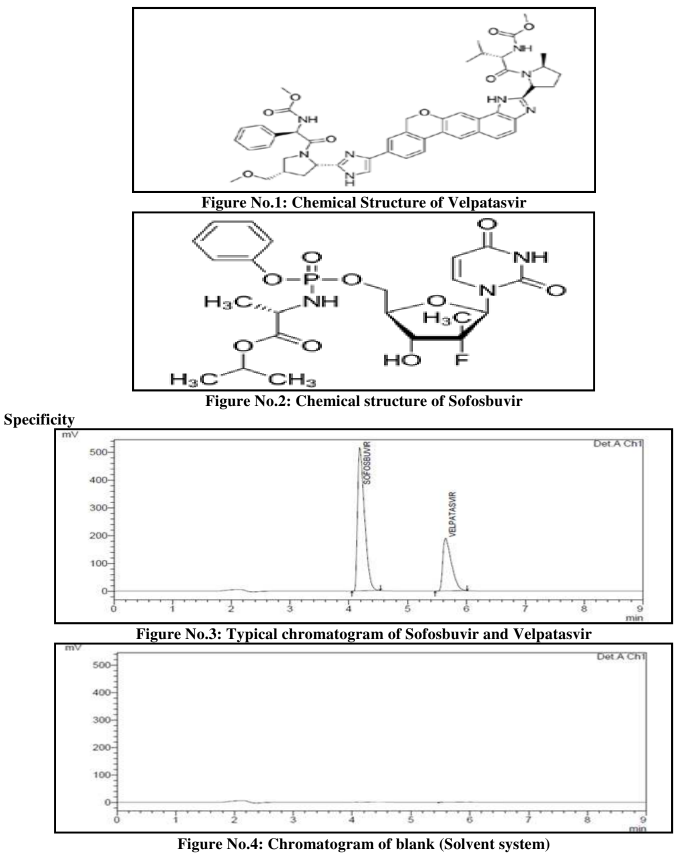
S.No	Type of degradation	Area of	sample	Assay content (% w/w)		
		Velpatasvir	Sofosbuvir	Velpatasvir	Sofosbuvir	
1	Acid (0.5N HCl)	859527	2831919	89.02	94.6	
2	Base (0.5N NaOH)	968847	2904346	91.4	96.3	
3	Peroxide $(3\% H_2 O_2)$	890779	28036451	91.2	94.9	
4	Thermal (at $60-80^{\circ}$ c)	975570	2884978	93.2	95.3	
5	Hydrolysis	1038182	3036541	92.9	95.7	

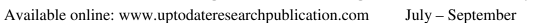
Table Summary for RP-HPLC Method

S.No	PARAMETERS	ACCEPTANCE CRITERIA	RESULTS OBTAINED
		Theoretical Plates- NLT 2000	Velpa- 5840
			Sofos-4710
1	System suitability	Tailing factor - NMT 2	Velpa -2.18
1	System suitability		Sofos -1.68
		Retention time	Velpa -5.91
		Retention time	Sofos -4.22
2	Dragision	% RSD of Velpa -NLT 2	Velpa -0.92
2	Precision	% RSD of Sofos -NLT 2	Sofos -0.62
	ID Precision		DAY-1 Velpa -0.83
3		% RSD of Velpa -NLT 2	Sofos -0.73
5		% RSD of Sofos -NLT 2	DAY-2 Velpa -1.02
			Sofos -0.87
4	Lincority	Correlation coefficient NLT 0.999	Velpa -0.999
4	Linearity	Contention coefficient NL1 0.999	Sofos -0.999
5	Accuracy	$\mathbf{P}_{\text{ansanto}, \alpha \alpha} = \mathbf{P}_{\alpha \alpha \alpha \gamma \alpha \gamma \gamma} = \frac{1020}{2}$	Velpa -99.6%
3		Percentage Recovery 98-102%	Sofos -99.4%
(Limit of Detection	1.2	Velpa -2.71µg/ml
6		1:3	Sofos – 7.71μ g/ml
7	Limit of quantitation	1.10	Velpa -8.31µg/ml
7		1:10	Sofos – 23.37μ g/ml

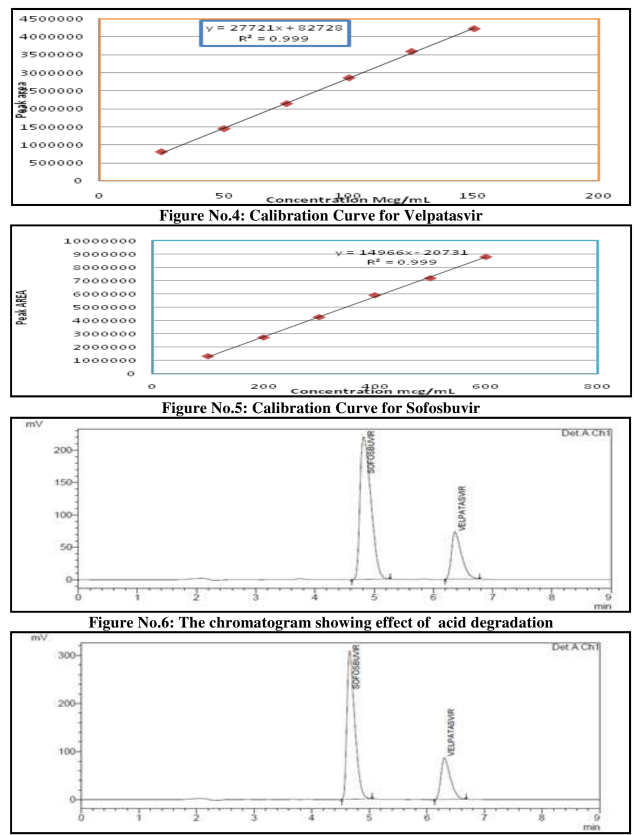
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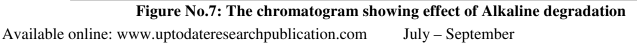
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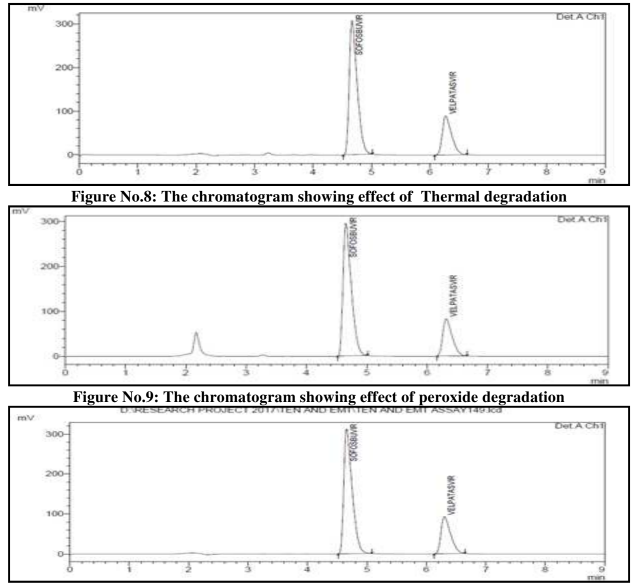


Figure No.10: The chromatogram showing effect of Photolytic degradation

SUMMARY AND CONCLUSION SUMMARY

RP-HPLC method was developed for simultaneous estimation of Sofosbuvir and Velpatasvir in pharmaceutical dosage form. Chromatographic separation was performed on Premisil C18 (4.6×250 mm) 5µ column, with mobile phase comprising of mixture of Acetonitrile: 0.1% Perchlioric acid in the ratio of 50:50% (v/v), at the flow rate 1.2ml/min. The detection was carried out at 262nm.

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CONCLUSION

The proposed HPLC method was found to be precise, specific, accurate, rapid and economical for simultaneous estimation of Sofosbuvir and Velpatasvir in tablet dosage form. It was also proved to be convenient and effective for the determination of Sofosbuvir and Velpatasvir in the bulk and combined dosage form. It inferred the method found to be simple, accurate, precise and linear. The method was found to be have a suitable application in routine laboratory analysis with high degree of accuracy and precision.

ACKNOWLEDGEMENT

The authors are very thankful to Sura Pharma Lab. Pvt., Ltd., Dilshuknagar, Hyderabad, for offered required reference samples for our research project and also very thankful to Chairman and Principal of Care College of Pharmacy-Warangal, Telangana, for providing necessary facilities and all required standard samples and continuous support for entire research work.

CONFLICT OF INTEREST

We declare that we have no conflict of interest.

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Please cite this article in press as: Gandla. Kumara Swamy *et al.* Novel stability indicating RP-HPLC method simultaneous determination of Sofosbuvir and Velpatasvir in bulk and combined tablet dosage forms, *Asian Journal of Research in Chemistry and Pharmaceutical Sciences*, 5(3), 2017, 114-122.

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