Development and Characterization of Fast Dissolving Buccal Strip of Frovatriptan Succinate Monoydrate for Buccal Delivery

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ABSTRACT

Objectives: Aim of the present research work is to prepared fast mouth dissolving film of frovatriptan succinate monoydrate. Methods: Fast dissolving buccal film is prepared by Solvent casting method, hydroxypropyl methylcellulose E3 and E15 are helpful in the film-forming polymer, plasticizer in polyethylene glycols disintegrant is croscarmellose sodium, the artificial sweetener in Sodium saccharin as, citric acid as saliva stimulant, fructose as diluents and natural sweetener, wild cherry as a flavour and Brilliant blue dye for elegance was selected for fast dissolving film preparation. By prism, software result is obtained and then evaluated using analysis of variance (ANOVA). Results: The result suggested that the formulation containing 20% w/w Sodium saccharin, 10% w/w fructose and 5% citric acid was found to effectively obscure the bitter taste of drug with best overall acceptability. The same composition of Sodium saccharin, citric acid and fructose were used for further optimization using design of an experiment to continue obscuring the bitter taste of frovatriptan succinate monoydrate. Simple lattice mixture design which is helpful in drug

formulation by using polymer plasticizer and disintegrant concentration for disintegration time-independent variable tensile strength and percentage elongation for the response. **Conclusion:** The effect of each variable, two and three-factor interactions were studied. The batches were numerically optimized to give a design space. fast mouth dissolving films are also found to behave better patient compliance in all the age groups. **Key words:** Migraine, Fast Dissolving Film, Solvent casting method,

Stability studies, Frovatriptan succinate monoydrate.

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INTRODUCTION

Migraine: Migraine is more commonly found in women than in men. In percentage, it is 20.7% in female and 9% in male. It is an episodic neurovascular disorder that occurs by severe attacks; the symptoms of unilateral pulsatile headache are nausea, photophobia and phonophobia.1 As it is related to prolonged disability and seen in Healthcare place. They are very painful and make a person weak.² It is the main issue responsible for absenteeism in the workplace or school.³ Headache that occurs due to tension is maximum among people, followed by a migraine headache, but it is excruciating out of all these migraines are found most common.⁴ The oral route is the most preferred route of administration for systemic effect. About 60% of all formulations are of the solid dosage form. Tablet is the most preferred dosage form due to ease of transportation, manufacturing and more patient compliance. After swallowing the conventional tablet, some patient faces some difficulties like a paediatric, geriatric and forbidden patient. To rectify this issue, a formulation was being developed, i.e. oral fast dissolving film.⁵ Oral dissolving films can be used anytime in the absence of water. Fast dissolving film shows faster dissolution due to least integration time. Sweeteners and flavours are added in preparation fast dissolving films which help in curing the patients.^{6,7}

Frovatriptan succinate monohydrate is a 5-HT receptor agonist which binds with high affinity to 5-HT_{1B} and 5-HT_{1D} receptors.⁸ *In vitro*, frovatriptan succinate monohydrate shows a moderate affinity for the receptor 5-HT7, which is believed to contribute to its distinctive pharmacologic properties.⁹ Frovatriptan succinate monoydrate demonstrated higher binding affinity than sumatriptan at the human 5-HT_{1B} receptor (~4-fold) and a comparable affinity at the human

 $5-HT_{1D}$ receptor.¹⁰ The absolute oral bioavailability of Frovatriptan succinate monohydrate is 22%-30%. The mean time to reach peak plasma concentration (T_{max}) is approximately 2-3 hr. Food does not influence the pharmacokinetics of frovatriptan succinate monoydrate.¹¹

MATERIALS AND METHODS

Materials

Frovatriptan succinate monohydrate, Hydroxypropyl Methylcellulose, Hydroxypropyl cellulose, Polyethylene glycol 400, polyethylene glycols, Glycerol, Fructose, Maltodextrin, Mannitol, Sodium Starch Glycolate, Crospovidone, Croscarmellose Sodium, Citric Acid, Sodium saccharin were used of analytical grade.

Methods

The drug and excipients were individually weighed and dispensed. The polymer was slowly dissolved in water under constant stirring at 600 rpm, followed by the addition of plasticizer, sweetener, citric acid, disintegrant, drug, colour and flavour. The solution was stirred for 2 hr to ensure a perfectly homogeneous mixture is obtained. The foam formed after stirring was removed by sonicating and degassing the solution, to obtain the final solution for fast dissolving buccal casting. The calculated amount of the solution was poured on Petri plates and specially fabricated rectangular glass plate. The Petri plates and rectangular glass plate was kept in an oven at 60°C for 4 hr for drying the solution. Fast dissolving buccal film formed after drying were carefully removed and cut into the desired size. The fast dissolving buccal film were packed in Aluminium

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wraps and sealed with the help of tagger sealer and stored until further evaluation was carried out.

Polymer selection, diluent and disintegrant

On drug-polymer compatibility, diluent and disintegrant studies Hydroxypropyl methylcellulose (HPMC), Hydroxypropyl cellulose (HPC), Polyvinylpyrrolidone (PVP) of different grades and ratios were used for the primary selection between the polymers diluent and disintegrant.

Characterization of the fast dissolving buccal film

Physical appearance/surface texture: After visual inspection of the fast dissolving buccal Film evaluation was done according to the surface texture or by feel or touch.

Thickness uniformity: By using digital Vernier calliper thickness was measured.

Uniformity of dosage units: The films were tested for uniformity of dosage form by the UV-Spectrophotometric method. Three different fast dissolving film of the same formulation was dissolved in 100 ml of Phosphate buffer pH 6.8 medium; 2 ml of this solution was diluted up to 10 ml with Phosphate buffer pH 6.8 medium to give a 7.8 ppm solution. The absorbance of the solution was measured at 244nm using a UV visible spectrophotometer. The standard solution of the raw drug of the same concentration was used as a bracketing standard after every six samples. By finding a standard graph percentage of drug content was being calculated and the same steps are repeated to other fast dissolving buccal film and the average drug content and standard deviation of each formulation was determined. This procedure was repeated for all 14 DOE formulations.

Assay

Formulation of 20 films were taken to dissolve it in 100 ml volumetric flask with purified water as medium (Stock I). From the Stock I solution equivalent to a single dose was calculated and was diluted with 100 ml of purified water. 2 ml of the solution is diluted till 10 ml with distilled water and its absorbance was measured at 244 nm using a UV visible spectrophotometer. The standard solution of the raw drug of the same concentration was used as a bracketing standard after every ten samples. The % drug content in each formulation was calculated using a standard graph. This procedure was repeated for every formulation and the assay was calculated.

In vitro disintegration

Each film that contains 10 ml of Phosphate buffer with pH 6.8 is placed in a petri dish. The time taken by each film when it starts to disintegrate was recorded. Average and standard deviations from 3 fast dissolving buccal films were measured and recorded.

In vivo disintegration

Fast dissolving buccal film is carried out in healthy volunteers (ageing 25-40 years, n=6) *in vivo* disintegration time. Before performing the test, the volunteers were aware of the steps and aim of the test. They were told that before placing the piece of fast dissolving buccal film on their tongue, they need to rinse their mouth. They were asked to monitor for the time required by the film to disintegrate and wash off thoroughly. *In vivo* disintegration times of the selected optimized batches were only carried out due to safety reasons.

In vitro dissolution

A suitable house method was used for *in vivo* drug release studies of fast dissolving buccal film, for dissolution to be carried out in USP type-I

(basket apparatus) of pH 6.8 medium with 500 ml of Phosphate buffer. The medium was maintained with a temperature of 37±0.5°C stirred at 100 rpm. The samples were withdrawn at an interval of 2, 5, 10, 15, 20 min and 30 min time. At each time interval, 10 ml of the sample was collected and is replaced with the same amount of medium. The samples were analysed in UV/Visible spectrophotometer, bracketing standard after every six samples were measured which was prepared by dissolving the same amount of drug in 500 ml Phosphate buffer pH 6.8 medium.

Solid-state form of drug by X-Ray Diffractometer (XRD): The form of input drug and the drug in the formulation was studied with the help of an X-ray diffractometer. The drug in the formulation must not recrystallize; if the drug in the formulation recrystallizes it gives a non-elegant appeal to the film.

Surface morphology of the formulation by Scanning Electron Microscopy (SEM): The surface morphology of the optimized formulation was studied with the help of Scanning Electron Microscopy (LEO 400).

Stability studies: Stability studies of the formulation were carried out under accelerated stability conditions (40 ± 2 , $75\pm5\%$ R.H.) and normal condition ($25\pm2^{\circ}$ C, $75\pm5\%$ R.H.) for four weeks. The stability samples were withdrawn at two weeks and four weeks duration and were tested for physical appearance, assay and *in vitro* disintegration time are shown in Table 5. The samples were loaded in LDPE (open condition) and sealed aluminium packs (closed condition).

RESULTS

Selection of Polymer, Diluent and Disintegrant

Polymer, diluent and disintegrant Selection are shown in Table 1. PEG 400, Citric acid, Sodium saccharin shows 60% that's for the best polymer. HPMC E3:E15 (85:15) shows 50% that's for the best diluent and best disintegrant.

Result of Polymer, Plasticizer, Diluent Selection Trials

Polymer has tested for different properties like Film-forming property, Appearance, Peelability, *in vitro* disintegration time and results are in Table 2. Plasticizers has tested for different properties like Folding endurance, Tensile strength (N), Percentage elongation, *in vitro* Disintegration time (sec) and results are in Table 2. Diluent has tested for different properties like recrystallization, texture, *in vitro* disintegration time (sec) and results are in Table 2.

Disintegration time, tensile strength, Percent elongation response as a response

The Analysis of Variance (ANOVA) result for Disintegration time, tensile strength, Percent elongation response presented in Table 3.

In vitro disintegration

The *in vitro* disintegration time of film was increases as the amount of Citric acid and Sodium saccharin increases and increase in concentration of plasticizer decreased disintegration time but after excessive amount of polymer increase, the film became brittle so there was slight decrease in disintegration time. Results of different batches are shown in Table 4.

In vivo disintegration

The *in vitro* disintegration time of different batches was found to be between 31-124 sec. Results of different batches are shown in Table 4.

Table 1: Initial trials for polymer, diluents and disintegrant selection.

Excipients	Ingredients			FDF -	I		
		1/01	1/02	1/03	1	/04	1/05
	Drug	6.72%	6.72%	6.72%	6	.72%	6.72%
	HPC EF	6.72%	-	0.727	, 0	.7 2 /0	-
	HPC LF	0.7270	- 50%	-	-		
	HPMC E3LV	-	50%	- 50%	-		-
	HPMC E15LV	-	-	-		0.0/	
	PVP	-	-			0%	-
	PEG 400	-	-	-	-	00/	50%
Polymer	Citric acid	10%	10%	10%		0%	10%
	Sodium saccharin	10%	10%	10%		0%	10%
	Brilliant Blue	10%	10%	10%		0%	10%
	Wild Cherry	0.02%	0.02%			.02%	0.02%
	white cherry	2.19%	2.19%	2.19%	2	.19%	2.19%
		3/01	3/02	3/03			
	Drug	6.72%	6.72%	6.72%			
	HPMC E3 : E15 (85:15)	50%	50%	50%	,		
	polyethylene glycols	10%	-	5070			
	Maltodextrin	10%	-	-			
	Mannitol	-	- 10%	-			
	Fructose	-	-				
Diluent	Citric acid			10%			
Jucit	Sodium saccharin	10%	10%	10%			
	Colour	10%	10%	10%			
		0.02%	0.02%				
	Flavor	2.19%	2.19%	2.19%)		
		3/01	3/01	3/01	3/01	3/01	
	Drug	6.72%	6.72%	6.72%	6.72%	6.72	%
	HPMC E3, E15 (85:15)	40%	40%	40%	40%	40%	
	polyethylene glycols	10%	-	-	-	-	
	Fructose	10%	-	-	-	-	
	Sodium starch glycolate	4%	-	-	-	-	
	Croscarmellose sodium	-	4%	-	-	-	
	Crospovidone XL	-	-	4%	-	-	
	Crospovidone XL 10	-	-	-	4%	-	
Disintegrant	Kollidone CLSF	-	-	-	-	4%	
	Citric acid	10%	10%	10%	10%	10%	
	Sodium saccharin	10%	10%	10%	10%	10%	
	Brilliant Blue	0.02%	0.02%	0.02%	0.02%	0.02	
	Wild Cherry	2.19%	2.19%	2.19%	2.19%	2.19	

The polymers selected for the initial trial are HPC EF, HPC LF, HPMC E3LV, HPMC E15LV, PVP, PEG 400, Citric acid, Sodium saccharin, Brilliant Blue, Wild Cherry.

The diluents selected for the initial trial are HPMC E3: E15 (85:15), polyethylene glycols, Maltodextrin, Mannitol, Fructose, Citric acid, Sodium saccharin, Colour, Flavour.

The suitable disintegrates for the initial trial are HPMC E3, E15 (85:15), polyethylene glycols, Fructose, Sodium starch glycolate, Croscarmellose sodium Crospovidone XL, Crospovidone XL 10, Kollidone CLSF, Citric acid, Sodium saccharin, Brilliant Blue, Wild Cherry.

In vitro dissolution

In vitro drug release study of batches was performed in USP Dissolution Apparatus-I at $37 \pm 0.5^{\circ}$ C and 100 rpm using 500 ml of PBS pH 6.8 buffer for 30 min. Cumulative % drug release of different batches at the end of 5 min was found to be between 85.9 to 97.6%. Results of different batches are shown in Table 5.

Solid-state form of drug by X-Ray Diffractometer (XRD)

The X-Ray Diffractometer (XRD) results also confirm the interactions and formation of a single homogenous phase by the appearance of the amorphous halo characteristic of single co-amorphous dispersions.

Scanning Electron Microscopy (SEM)

SEM images of frovatriptan succinate monoydrate fast dissolving buccal film are presented in Figure 1. SEM of frovatriptan succinate monoydrate fast dissolving buccal film exposes discrete, elongated flake-like structures with rough edges covered on their surfaces by fine particles. Some structures are large with parallelogram shape. It also reveals the hard and thick nature of the drug particles. In contrast, frovatriptan succinate monoydrate complex observed by SEM is soft and thin.

Stability studies

Accelerated stability studies optimized after 4 weeks and it showed good stable condition. Results of stability studies of different batches are shown in Table 6.

DISCUSSION

The physicochemical characteristics of the drug were investigated and had a similarity with the internal specifications. The working λ_{max} was determined by using a UV spectrophotometer and was found to be 244nm, which complied with the internal specifications (λ_{max} 245nm).¹²

The taste masking of the films was done by optimizing the concentration of Sodium saccharin (sweetener), citric acid (saliva stimulant and flavour) and fructose (natural sweetener and diluents). The tastes of the

Table 2: Result of Polymer, Plasticizer, Diluent Selection Trials.

optimized films were panel tested by a group of 5 healthy volunteers. The selected formulations after the panel testing were evaluated statistically to see if there was any significant difference in the three formulations.¹³ The evaluation was done applying one way ANOVA using the Kruskal-Wallis test. Post-test, the formulations were compared to each other using Dennett's multiple comparison tests. Prism 5.0 software was used for calculation and analysing the results. The concentration of Sodium saccharin, citric acid and fructose from the most desirable batch was selected and kept constant in further formulation optimization, to mask the taste of the drug effectively.

For the optimization procedure, a three-level simplex lattice mixture design was used. Three factors, concentration of Polymer (HPMC) (A), Plasticizer (polyethylene glycols) (B) and Disintegrant (C), were used in the design and the responses were *in vitro* disintegration time (Y1), Tensile strength (Y2) and Percentage elongation (Y3).¹⁴

Based on a Simplex lattice mixture design, a total of 14 batches were prepared by solvent casting method and evaluated based on *in vitro* disintegration time, Tensile strength and Percentage elongation. The *in vitro* disintegration time of different batches was found to be between 32-122 sec. The tensile strength of various batches was found to be in the

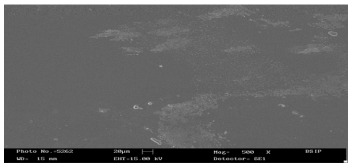


Figure 1: Scanning Electron Microscope (SEM) Image of fast dissolving buccal film at 400x magnification.

Excipients	Property tested		FDF -I	
		1/06	1/07	1/08
	Film-forming property			
	Appearance	2	2	2
Polymer	Peelability	1	1	1
	In vitro disintegration time	2	1	2
		172	169	166
		1/08	1/09	1/10
	Folding endurance	127	231	153
	Tensile strength (N)	5.95	11.57	8.81
Plasticizer	Percentage elongation	2.3%	8.7%	3.7%
	<i>In vitro</i> Disintegration time (sec)	165	168	172
		3/02	3/02	3/03
	Recrystallization Texture	No	Yes	No
Diluent	<i>In vitro</i> disintegration time (sec)	Rough	-	Smootl
	in the domestication time (see)	158	-	138

The in vitro disintegration time and texture of FDF -I-3/03 was found to be acceptable.

The different formulations like FDF -II-1/6, FDF -II-1/7, FDF -II-1/8 were selected for panel testing and found to be overall accepted

Time Response	Source		ANOVA	Result		Comment
		Sum of	Mean	F	p-value	
		Squares	Square	Value	Prob > F	
Disintegration Time Response (ANOVA for Mixture Special	Model Linear Mixture AB AC BC ABC	9747.933 6442.64 772.5344 570.6028 682.3541 137.832	1541.324 3172.22 870.5254 470.5038 682.3541 137.9362	68.38752 128.2156 34.71706 21.64333 28.85092	<0.0001 <0.0001 0.0005 0.002 0.0012	Significant
Cubic Model	Residual Lack of Fit Pure Error Core Total	155.4472 79.05724 78.5 10112.4	22.45102 24.32231 22.525	5.255006 1.118608	0.0406	not significant
Tensile Strength Response	Model Linear Mixture AB AC BC ABC Residual Lack of Fit Pure Error Cor Total	34.1323 31.24952 1.158323 0.107102 0.687857 0.675196 1.101973 0.813673 0.2883 35.21436	5.752722 14.52988 1.158323 0.107102 0.687857 0.675196 0.157425 0.271224 0.072075	35.1762 107.5355 6.727325 0.650373 5.06888 4.289005 3.763086	< 0.0001 < 0.0001 0.0250 0.3141 0.0591 0.0662 0.1165	Significant not significant
Percentage Elongation Response	Model Linear Mixture AB AC BC Residual Lack of Fit Pure Error Cor Total	14.81713 11.98774 2.118837 0.028382 0.533003 0.602808 0.573358 0.10944 15.51885	2.163428 6.493868 2.218838 0.028382 0.543001 0.087851 0.143339 0.022363	36.00903 73.91914 25.25684 0.323075 5.180928 3.429183	< 0.0001 < 0.0001 0.0010 0.5854 0.0377 0.0893	Significant not significant

Table 3: ANOVA for Mixture Special Cubic Model for Disintegration Time Response, Tensile Strength Response and Percentage Elongation Response.

Different model values were presented as mean \pm S.D. (*n*=3).

range of 4.35 to 9.20 MPa. The Percentage elongations of various batches were found to be in the range of 2.48 to 5.88%. *In vitro* drug release study of batches was performed in USP Dissolution Apparatus-I at 37 ± 0.5 °C and 100 rpm using 500 ml of PBS pH 6.8 buffer for 30 min. Cumulative % drug release of different batches at the end of 5 min was found to be between 84.9 to 98.4%.¹⁵

The Optimized batch (OB) fast dissolving film (FDF)-III-1/7 was determined by using the software Design Expert. The optimized batch (OB) was prepared with concentrations of Polymer, Plasticizer and disintegrate at 43.33, 8.33 and 3.33 respectively by solvent casting method. Optimized batch (OB) showed *in vitro* disintegration time of 42 sec, Tensile strength of 7.36 MPa and Percentage elongation of 4.43%. The results of Scanning Electron Microscope (SEM) (Figure 1) showed that uniformity in size and shape of frovatriptan succinate monohydrate fast dissolving buccal film. The Scanning Electron Microscope (SEM) of fast dissolving buccal film at 500X demonstrated the characteristic amorphous aggregates and presence of the defined shape of crystals. The fast dissolving buccal film prepared from solvent casting method showed an almost clear, transparent glassy and homogenous film with wholly dissolved into the polymer matrix.¹⁶

Table 4: Result of in vitro/ in vivo Disintegration DOE Batches.

Evaluation parameter	Method/time	Formulation	D.T. (sec)
		FDF -III-1/1	30.5±0.5
		FDF -III-1/2	46±1
		FDF -III-1/3	52±1
		FDF -III-1/4	66.5±2
		FDF -III-1/5	34.5±1.4
		FDF -III-1/6	32.5±0.5
Disintermetica	T ''	FDF -III-1/7	41±1
Disintegration	In vitro	FDF -III-1/8	65.5±1.4
		FDF -III-1/9	51.5±0.5
		FDF -III-1/10	66.5±0.5
		FDF -III-1/11	123.5±1.4
		FDF -III-1/12	46.5±1.5
		FDF -III-1/13	110.5±0.5
		FDF -III-1/14	56.5±0.5
		FDF -II-1/6	16.16
	In vivo	FDF -II-1/7	17.66
		FDF -II-1/8	15.33

In vitro/ in vivo disintegration time of different batches FDF III and FDF II

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Table 5: F

In vitro Dissolution	Time	FDF -III- 1/1	FDF -III- 1/2	FDF -III- 1/3	FDF -III- 1/4	FDF -III- 1/5	FDF -III- 1/6	FDF -III- 1/7	FDF -III- 1/8	FDF -III- 1/8	FDF -III- 1/9	FDF -III- 1/10	FDF -III- 1/11	FDF -III- 1/12	FDF -III- 1/13	FDF -III- 1/14
	2 min	73.8%	82.6%	72.4%	84.2%	84.4%	78.2%	76.7%	83.2%	95.2%	83.2%	84.3%	85.5%	91.1%	77.1%	84.6%
	5 min	86.3%	92.4%	96.1%	94.7%	92.2%	86.7%	84.8%	95.2%	98.8%	96.1%	91.6%	92.8%	96.6%	93.5%	93.6%
	10 min	96.1%	98.7%	101.6%	96.2%	97.6%	93.3%	94.1%	98.8%	100.3%	99.7%	98.2%	98.4%	97.6%	99.6%	99.3%
	15 min	96.5%	100.2%	102.1%	98.5%	98.2%	97.6%	98.5%	100.3%	100.3%	100.5%	99.3%	100.6%	99.5%	100.1%	101.6%
	20 min	98.2%	101.1%	102.2%	98.6%	101.5%	101.2%	98.5%	100.3%	100.2%	101.3%	99.5%	101.3%	%2.66	100.3%	102.5%
	30 min	98.2%	101.4%	102.2%	98.2%	101.5%	101.5%	98.8%	100.2%		101.5%	<i>66</i> %	101.6%	99.8%	100.3%	102.7%

In vitro Dissolution profile of different DOE Batches FDF 1 to FDF 14

Table 6: Result of Accelerated Stability	Studies.
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Storage condition	Duration	Туре	Colour change	Assay	<i>In vitro</i> disintegration time (sec)
40±2°C, 75±5% R.H.	Two weeks	Open	Yes	94.42±0.66	34±0.34
40±2°C, 75±5% R.H.	Two weeks	Closed	No	97.86±2.65	32±0.5
40±2°C, 75±5% R.H.	Four weeks	Open	Yes	93.78±0.93	36±0.72
40±2°C, 75±5% R.H.	Four weeks	Closed	No	96.61±1.92	34±1.4

Stability data of different formulations were presented as mean \pm S.D. (*n*=3).

The X-Ray Diffractometer (XRD) spectrum of Optimized Batch Placebo Formulation of fravotriptan Succinate monohydrate results also confirm the interactions and formation of a single homogenous phase by the appearance of the amorphous halo characteristic of single coamorphous dispersions.¹⁶ The incorporation of fast dissolving buccal film ascorbic acid co-amorphous dispersions in fast dissolving buccal film was intended to provide a simple, easily administered dosage form for the application inside the mouth. The formulated film was found to have an acceptable transparent appearance and mechanical properties, disintegration and dissolution.¹⁷ Almost complete dissolution from the film was attained within 10 min. Also, the drug was found to be stable in the film for up to 12 weeks at 40°C and 75% RH which could be attributed to compatibility between polymers used in the film. This increased phase stability possibly occurred through extending the super saturation state. The optimized fast dissolving buccal film formula obtained by the model was chosen for testing the physicochemical stability. Stability study of the optimized batch was carried out at accelerated condition and was found to be stable with no colour change when packed and sealed in aluminium foil, for four weeks.17

CONCLUSION

From the present study, it may be concluded that the fast dissolving films of frovatriptan succinate monohydrate can be prepared by the solvent casting method using HPMC E3 and E15 in 85:15 ratio respectively by using polyethylene glycols as plasticizer, croscarmellose sodium as a disintegrant and fructose as a natural sweetener and diluents. The final formula was optimized by simplex lattice mixture design using Stat-Ease design expert software to give the optimized batch. The optimized batch (OB) prepared with concentrations of Polymer, Plasticizer and disintegrant at 42.32, 7.32 and 4.32 respectively by solvent casting method. Optimized batch (OB) showed *in vitro* disintegration time of 44 sec, Tensile strength of 6.35 MPa and Percentage elongation of 3.62%.

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CONFLICT OF INTEREST

The authors declare that there is no conflict of interest.

ABBREVIATIONS

FTIR: Fourier Transform Infrared Spectroscopy; FDF: fast dissolving films; DOE: Design of experiment; SEM: Scanning Electron Microscope;

HPMC: Hydroxypropyl methylcellulose; **HPC:** Hydroxypropyl cellulose; **PVP:** Polyvinylpyrrolidone.

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