



FORMULATION AND EVALUATION OF FAST DISSOLVING FILMS OF RIZATRIPTAN



PRASANNA KUMAR DESU, BRAHMAIAH BONTHAGARALA,
SREEKANTH NAMA, A.NAGALAKSHMI



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Department of Pharmaceutics, Priyadarshini Institute of Pharmaceutical Education &
Research (PIPER).

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Corresponding Author

Mr. Prasanna Kumar Desu

The present study was aimed to formulate and evaluate fast dissolving films of Rizatriptan using hydroxyl propyl methyl cellulose. 5-HT_{1B} and 5-HT_{1D} antagonist which is an antimigraine. Hydroxyl propyl methyl cellulose is used as film forming agent. Fast dissolving films are meant to be dissolved in saliva and remain in oral cavity until swallowed. Hence taste masking becomes critically important. The films are prepared by solvent evaporation method and characterized by UV studies. The suitable plasticizer and its concentration were selected on the basis of flexibility, folding endurance and stickiness of the film. In the present study propylene glycol was used as plasticizer. Films were evaluated for drug content and the drug loading capacity was found to be 99.96% per 2cm². The dissolution profile, disintegrating time and folding endurance were found to be satisfactory. Further, the optimized films were evaluated and it was found that the films disintegrate within 1 min and absence of bitterness in the film. Hence it is concluded that Rizatriptan H.P.M.C fast dissolving films are successfully developed and evaluated.

INTRODUCTION:

For the last two decades, there has been an enhanced demand for more patient-compliant dosage forms. As a result, there are now approximately 350 drug delivery corporations and 1000 medical device companies. The demand for their technologies was approximately \$14.20 billion in 1995 and, according to industry reports; this is expected to grow to \$60 billion annually. Oral administration is the most popular route due to ease of ingestion, pain avoidance, versatility (to accommodate various types of drug candidates), and most importantly, patient compliance. Also, solid oral delivery systems do not require sterile conditions and are, therefore, less expensive to manufacture. Several novel technologies for oral delivery have recently become available to address the physicochemical and pharmacokinetic characteristics of drugs, while improving patient compliance. Electrostatic drug deposition and coating, and computer-assisted three-dimensional printing (3DP) tablet manufacture have also recently become available.

Fast-dissolving drug-delivery systems were first developed in the late 1970s as an alternative to tablets, capsules, and syrups for pediatric and geriatric patients who experience difficulties swallowing ⁽¹⁾ traditional oral solid-dosage forms. The novel technology of oral fast-dispersing dosage forms is known as fast dissolve, rapid dissolve, rapid melt and quick disintegrating tablets. However, the function and concept of all these dosage forms are similar

By definition, a solid dosage form that dissolves or disintegrates quickly in the oral cavity, resulting in solution or suspension without the need for the administration of water, is known as an oral fast-dispersing dosage form. Difficulty in swallowing (dysphasia) is common among all age groups, especially in elderly, and is also seen in swallowing conventional tablets and capsules. Dysphasia is associated with many medical conditions, including stroke, Parkinson's, AIDS, thyroidectomy, head and neck radiation therapy, and other neurological disorders, including cerebral palsy. The most common complaint was tablet size, followed by surface, form and taste. The problem of swallowing tablets

was more evident in geriatric and pediatric patients, as well as travelling patients who may not have ready access to water⁽¹⁾.

Research and development in the oral drug delivery segment has been led to transition of dosage forms from simple conventional tablets/capsules to oral disintegration tablet (ODT) to wafer to the recent development of oral films (ODF) can be considered as an ultra thin strip of postage stamp size with an active agent or active pharmaceutical ingredient and other pharmaceutical excipients. The advantage of convenience of dosing probability of ODF has led to wider acceptability of this dosage form by pediatric as well as geriatric population equally.

Salient features of fast dissolving drug delivery systems

1. Ease of administration for patients who are mentally ill disabled and uncooperative.
2. Require no water.
3. Over comes unacceptable taste of the drugs.
4. Can be designed to leave minimal or no residue in the mouth after

administration and also provide a pleasant mouth feel.

5. Ability to provide advantages of liquid medication in the form of solid preparation.
6. Adaptable and amenable to existing processing and packaging
7. Cost effective.

Advantages⁽²⁾:

These rapid dissolving films offer several advantages like,

- Convenient dosing.
- Fast disintegration or dissolution followed by quick effect which is desirable in some cases such as pain.
- No water needed.

Disadvantages⁽³⁾:

- The disadvantage of OTF is that high dose cannot be incorporated into the strip.
- Expensive packaging of oral film.

MATERIALS AND METHODS:

Rizatriptan, Hydroxy propyl methyl cellulose, xylitol, Acesulfame potassium, propylene glycol and peppermint flavor were arranged by pharmatrai, Hyderabad.

Formulation of drug films^(4,5):

The formulation of films by using solvent casting method. The following steps are used in the manufacturing of films by this method. The polymers were dissolved in hot water. The drug and other excipients were dissolved in ethanol. This solution was added to the polymeric solution. Then the solution was mixed by using mixing device for 45 minutes with rotating speed 60-80 rpm. The entrapped air is removed by vacuum. The resulting solution was casted slowly and with continuous flow on a glass plates. The plates were kept in a hot air oven at 60°C for 24 hours. The dried film was gently separated from glass plate and cut into desired sizes

RESULTS AND DISCUSSIONS:

1. Thickness:

The thicknesses of the films were calculated by using screw gauge. The thickness was varied from 0.025 to 0.06 mm. The values

obtained for all the formulations were given in the table 2.

2. Folding Endurance

The folding endurance was found to be in the range of 21 ± 0.57 to 25 ± 0.57 . The values for all eight formulations are given in the table 1. This data revealed that the films had good mechanical strength along with flexibility. The values obtained for all the formulations were given in the table 2.

3. Weight Variation

The three films of 2*2 cm² was cut and weighed on electronic balance for weight variation test. The results obtained for all formulations were tabulated in the table 2.

4. Invitro disintegration:

2ml of pH 6.8 phosphate buffer was placed in a petri plates with a film on the surface of buffer the time taken for the disintegration of the film was measured. The results obtained all the formulations were tabulated in the table no.2.

5. Assay: This test was performed by dissolving a 4cm² area of film in 50ml of pH 6.8 phosphate buffer with stirring. This solution was filtered using a Whatmann

filter paper, and the filtrate was diluted to 100ml with the same buffer in a volumetric flask. This solution was analyzed by U.V. Spectrophotometer and all vales were tabulated in the table no. 2.

6. Invitro dissolution Time:

900ml of phosphate buffer(pH6.8)was used as a media, and was maintained at $37\pm 0.5^{\circ}\text{C}$ while the basket was set at 100 rpm a film sample of $4\text{cm}^2(2*2\text{cm})$ was cut and taken in to the basket. 5ml of the sample were taken every 2 minutes and the same amount was replaced with fresh buffer. The withdrawn samples were filtered and analyzed by using a U.V Spectrophotometer at a wavelength of 226nm. The results obtained for all the formulations are tabulated in the table 2 Plot a graph between % cumulative drug release on y- axis and time on x- axis, the obtained graph was represented in figure.No.1.

SUMMARY AND CONCLUSION:

The Rizatriptan is a serotonin (5-HT₁)

agonist used for the treatment of migraine with or without aura. The half-life of Rizatriptan is 2.5 to 3 hrs and it undergoes hepatic metabolism, the absolute oral bioavailability is about 40 to 50%. So, in order to improve the bioavailability and efficacy, we have prepared rapidly dissolving films of Rizatriptan. In the present research work, films were prepared using H.P.M.C 5cps polymer by Solvent Casting Technique.. Rizatriptan is water soluble drug belongs to class III of BCS classification of drugs. Estimation of the drug by U.V method at 226nm.. Formulation study involves F-4 shows good mechanical properties and less disintegration time. The drug release from this formulation was good.

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Table 1

Composition of all Formulations

INGREDIENTS	F1	F2	F3
Drugs	125	125	125
H.P.M.C 5cps	625	750	875
Xylitol	1375	1250	1125
Propylene glycol	250	250	250
Acesulfame potassium	100	100	100
Peppermint flavour	25	25	25
Total	2500	2500	2500

Table 2

Physicochemical evaluation data of Rizatriptan films

Formulation	Thickness (mm)	Folding endurance	Weight variation(mg)	Disintegration Time	Assay (%)
F-1	0.60	16	0.975	35	97.23
F-2	0.56	12	0.985	30	98.46
F-3	0.58	14	0.986	32	96.12
F-4	0.52	10	0.986	28	99.23
F-5	0.54	12	0.983	25	97.58

Table3

In vitro dissolution studies of all formulations

Time(min)	F-1	F-2	F-3	F-4	F-5
2	47.62	46.92	42.26	47.88	45.56
4	63.23	65.46	62.12	64.52	54.56
6	68.23	72.40	69.12	72.74	71.46
8	84.46	83.23	80.52	89.57	88.23
10	92.23	90.45	91.46	99.96	94.56

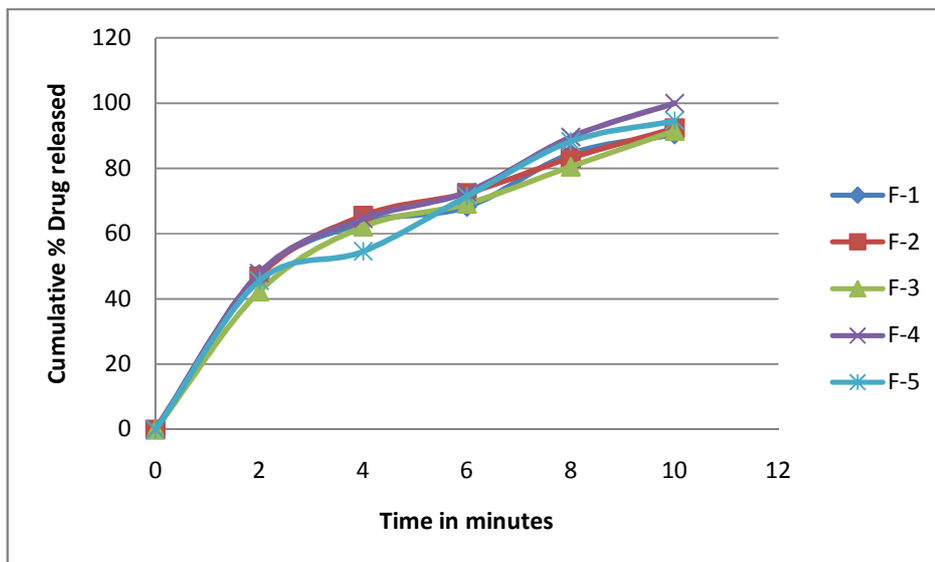


Fig1 Comparative Invitro Drug release from all formulations

REFERENCES:

1. Committee for medicinal and products for human use, European medicines agency EMEA, Reflection paper; formulation of choice of the pediatric population, sep, 2006.
2. Technology catalysts International Corporation, accessed on jun 15th 2011 Available from <http://www.technologycatalyst.com>
3. "oral thin films," in orally Disintegrating Tablet and Film Technologies, 4th ed. pp:18-31.
4. Technical brief 2010. Vol 3 Particle sciences Drug Development Services.
5. Coppens, K.A., M>j. Hall, S.A. Mitchell and M.D. Read, 2005. Hypermallose, Ethyl cellulose and polyethylene oxide used in the hot melt extrusion. Pharmaceutical Technol., pp:1-6
6. Felton L, Donnell P.O, McGinity J, Mechanical properties of polymeric films prepared from aqueous dispersions, in: Aqueous polymeric coatings or pharmaceutical dosage form, 3rd edition, McGinity j, Felton L Drugs and the pharmaceutical sciences, p.108.
7. Fulzels S.V, sattuar P.M and Dorie A.K, Polymerized rosin: novel film forming polymer for the drug delivery, Int J Pharm. 2002; 249:175-184.