INTRODUCTION

Oral drug delivery system represents one of the frontier areas of controlled drug delivery system; such dosage forms are having a major advantage is patient compliance. Floating drug delivery system belongs to oral controlled drug delivery system group that are capable of floating in the stomach by bypassing the gastric transit. These dosage forms are also defined as gas powered system (GPS), which can float in the contents of the stomach and release the drug in a controlled manner for prolonged periods of time. The release rate will be controlled depending upon the type and concentration of the polymer that swells, leads to diffusion and erosion of the drug.1

The investigation was concerned with design and characterization of Imatinib Mesylate floating matrix tablets for controlled release in order to improve efficacy and better patient compliance. Imatinib Mesylate is an anti-cancer agent which is used to treat chronic myelogenous leukemia (CML), gastrointestinal stromal tumors (GISTs) and a number of other malignancies. In the present research work an attempt has been made for the formulation of floating tablet containing Imatinib Mesylate as a drug candidate which would remain in stomach or upper part of GIT for prolonged period of time thereby maximizing the drug release at the desired site within the stipulated time. In the present study Imatinib Mesylate floating tablets were prepared by wet granulation method. The floating tablets were subjected to preformulation studies, in-vitro drug release, kinetic studies and stability studies. FTIR studies showed there was no interaction between drug and polymers. The percentage of Imatinib Mesylate content from the tablets was determined by UV-Spectroscopy and ranged from 98.25±1.8 to 98.91±1.5. The in-vitro percentage release of Imatinib Mesylate from the optimized tablets at the end of 12 hours was 99.46 %. The kinetic studies revealed that the drug was released by zero-order kinetics. The optimized formulation was subjected to stability studies and shown there were no significant changes in drug content, physicochemical parameters and release pattern. From this study, it is concluded that, the formulations retained for longer period of time in the stomach and provides controlled release of the drug. Hence, it will be increasing the bioavailability of the drug and patient compliance.

MATERIALS AND METHODS

Materials

Imatinib Mesylate is obtained as gift sample from Celon labs pvt.Ltd., Hyd. HPMC K4M, HPMC K15M was obtained as gift sample from Lupin Laboratories, Pune. NaHCO₃ was obtained from Fisher Scientifics, Pvt ltd. Lactose was obtained as gift sample from Dr. Reddy's labs India ltd, Hyd. All other materials used were of analytical grade.

Experimental Methods

Pre formulation study

Almost all the drugs which are active orally are marketed as tablets, capsules or both. Prior to development of dosage forms with a new drug candidate, it is essential that certain fundamental physical and chemical properties of the drug molecule and other derived properties of the drug powder are determined. This information will dictate many of the subsequent events and possible approaches in formulation development.

Solubility study

The solubility study was used to identify the suitable solvent that possesses good solubilizing capacity for Imatinib Mesylate. Solubility of Imatinib Mesylate in various solvents was determined by adding excess of Imatinib Mesylate in each selected solvents in each conical flask containing 20ml and is shaken for 24h using Rotary shaking apparatus. Then the solubility of drug in each solvent is observed visually and by UV spectrophotometrically at maximum wavelength 236 nm after relevant dilutions.

Excipients compatibility study

The successful formulation of a suitable and effective solid dosage form depends upon the careful selection of the excipients. Excipients are added to facilitate administration, promote the consistent release and bio availability of drug. It's necessary to study the compatibility of excipients with drug. Here IR spectroscopy was used to investigate and predict any physicochemical interaction between components in a formulation and to the selection of suitable compatible excipients.

Infrared spectrophotometer (IR)

Infrared (IR) spectroscopy was conducted and the spectrum was recorded in the wavelength region of 4000 to 400cm⁻¹. The
procedure consisted of, dispersing a sample (drug alone, polymers alone and mixture of drug and polymers in KBr and compressing into discs by applying a pressure of 7 tons for 5 min in a KBr press. The pellet was placed in the light path and the spectrum was obtained².

**Preparation of floating matrix tablets of Imatinib Mesylate**

Formulation development study was carried out for the preparation of floating tablet by using wet granulation method. Total weight of each floating matrix tablet is 400 mg. Imatinib Mesylate and all the ingredients are accurately weighed and passed through sieve #60. Imatinib Mesylate is well mixed (as shown in Table 1) with half quantity of lactose, polymer and then mixed with remaining ingredients in geometric proportions. Mixed homogeneously in a polybag for about 5-10 min. The obtained homogenous powder was taken in a glass mortar and granules were prepared by using isopropyl alcohol as granulating agent. The wet mass was passed through sieve # 14 and dried in hot air oven at a temperature of 50°C, dried granules were sieved through sieve # 16. Then the obtained dry granules are lubricated with the previously weighed and sieved magnesium state, talc and aerosil to obtain the blend for compression. Then the lubricated blend is subjected to compression by 12 mm circular standard flat faced punch (Rimek mini press, model RSB-4, M/S: Karnavathi engineering, Ahmadabad).

**Evaluation of dry granule characteristics**

Imatinib Mesylate dry granules of different formulas from F1 to F13 were evaluated for angle of repose, bulk density, tapped density, Hausner ratio, Carr's index.

### Table 1: Composition of floating matrix tablet of Imatinib Mesylate

<table>
<thead>
<tr>
<th>Formulation Code</th>
<th>Imatinib Mesylate (mg)</th>
<th>SBC (mg)</th>
<th>HPMC K4M (mg)</th>
<th>HPMC K15M (mg)</th>
<th>Lactose (mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>120</td>
<td>70</td>
<td>105</td>
<td>25</td>
<td>68</td>
</tr>
<tr>
<td>F2</td>
<td>120</td>
<td>70</td>
<td>90</td>
<td>40</td>
<td>68</td>
</tr>
<tr>
<td>F3</td>
<td>120</td>
<td>70</td>
<td>75</td>
<td>55</td>
<td>68</td>
</tr>
<tr>
<td>F4</td>
<td>120</td>
<td>70</td>
<td>60</td>
<td>70</td>
<td>68</td>
</tr>
</tbody>
</table>

SBC=Sodium bicarbonate, HPMC=Hydroxy propyl methyl cellulose, All formulations contain 1% of Talc, 1% of Magnesium stearate and 1% of Aerosil.

**Angle of repose**

The fixed funnel and free standing cone methods employ a funnel that is secured with its tip at a given height, h, which was kept 2cm above graph paper that is placed on a flat horizontal surface. With r being the radius, of base of conical pile, angle of repose can be determined by following equation: θ = tan⁻¹ (h/r)

Where, θ is the angle of repose, h is height of pile, r is radius of base of the pile

**Bulk density and tapped density**

Both loose bulk density and tapped bulk density were determined. A quantity of 2gm of granules from each formula, previously light shaken for the break of any agglomerates formed, was introduced into the 10ml of measuring cylinder. After the initial volume was observed, the cylinder was allowed to fall down its own weight from the hard surface from a height of 2.5cm at 2 sec Intervals. The tapping was continued until no further change in the volume was noted LBD and TBD were calculated using the following formulas:

- **LBD:** Weight of the powder/volume of the packing.
- **TBD:** Weight of the powder/Tapped volume of the packing.

**Compressibility index**

The compressibility index of the granules was determined by Carr's Compressibility index.

- **Carr’s index (%) = ([TBD-LBD] / TBD) × 100)**
- **TBD:** Weight of the powder/Tapped volume of the packing.

**Hauser’s ratio**

Hauser’s ratio can be determined by the following equation:

- Hauuer's ratio = TBD / LBD

Where, TBD -Tapped bulk densities & LBD- Loose bulk densities

**Evaluation of tablet characteristics**

Evaluation was performed to assess the physicochemical properties and release characteristics of the developed formulations. Following parameters were evaluated.

**Weight variation**

Twenty (20) tablets from each batch were individually weighed in grams (gm) on an analytical balance. The average weight and standard deviation were calculated and the results were expressed as compliance or non-compliance of set limits.

**Weight variation tolerance**

<table>
<thead>
<tr>
<th>Average weight (mg)</th>
<th>% Deviation allowed</th>
</tr>
</thead>
<tbody>
<tr>
<td>130 or less</td>
<td>10</td>
</tr>
<tr>
<td>130-323</td>
<td>7.5</td>
</tr>
<tr>
<td>More than 324</td>
<td>5</td>
</tr>
</tbody>
</table>

**Tablet hardness**

Tablet hardness was measured using a Monsanto hardness tester. The crushing strength of the 10 tablets with known weight and thickness of each was recorded in kg/cm² and the average hardness and standard deviation was reported.

**Friability**

Twenty (20) tablets were selected from each batch and weighed. Each group of tablets was rotated at 25 rpm for 4 minutes (100 rotations) in the Roche friabilator. The tablets were then dusted and re-weighed to determine the loss in weight. Friability was then calculated as percent weight loss from the original tablets and % friability was calculated using formula.

\[ F = \left[ \frac{W_t - W_r}{W_t} \right] \times 100 \]

Where, \( W_t \) - Weight of tablet before test, \( W_r \) - Weight of tablet after test

**Content uniformity**

From each batch of prepared tablets, ten tablets were collected randomly and powdered. A quantity of powder equivalent to weight of one tablet was transferred in to a 100 ml volumetric flask, to this 100 ml of 0.1N HCL was added and then the solution was subjected to sonication for about 2 hours. The solution was made up to the mark with 0.1N HCL. The solution was filtered and suitable dilutions were prepared with 0.1N HCL. Same concentration of the standard solution was also prepared. The drug content was estimated by recording the absorbance at 236 nm by using UV-Visible spectrophotometer.

**Buoyancy / Floating Test**

The in vitro buoyancy was determined by floating lag time, as per the method described by a Rosa et al., 1994. Here, the tablets...
were placed in a 100-ml beaker containing 0.1N HCL. The time required for the tablet to rise to the surface and float was determined as floating lag time and total duration of time by which dosage form remain buoyant is called Total Floating Time (TFT).

**Water uptake studies**

The swelling behavior of dosage unit can be measured either by studying its dimensional changes, weight gain or water uptake. The water uptake study of the dosage form was conducted by using USP dissolution apparatus II in a 900ml of distilled water which was maintained at 37°C + 0.5°C, rotated at 50 rpm. At selected regular intervals the tablet was withdrawn and weighed. Percentage swelling of the tablet was expressed as percentage weight uptake (%WU).

\[ \%\text{WU} = \left( \frac{Wt - Wo}{Wo} \right) \times 100 \]

Where, Wt is the weight of the swollen tablet, Wo is the initial weight of the tablet.

**Dissolution Study of tablets**

The tablet was placed inside the dissolution vessel. 5ml of sample were withdrawn at time intervals of 30, 60, 120 and 180, 240, 300, 360, 420, 480, 540, 600, 660, and 720 minutes. The volume of dissolution fluid adjusted to 900 ml by replacing 5ml of dissolution medium after each sampling. The release studies were conducted with 3 tablets, & the mean values were plotted versus time. Each sample was analyzed at 236 nm using double beam UV and Visible Spectrophotometer against reagent blank.

The tablet was placed inside the dissolution vessel. 5ml of water uptake study of the dosage form was conducted by using USP dissolution apparatus II in a 900ml of distilled water which was maintained at 37°C + 0.5°C, rotated at 50 rpm. At selected regular intervals the tablet was withdrawn and weighed. Percentage swelling of the tablet was expressed as percentage weight uptake (%WU).

**Kinetics of in-vitro drug release**

To study the release kinetics in-vitro release data was applied to kinetic models such as zero-order, first order, Higuchi and Korsemeyer-Peppas.

**Zero-order,**

\[ C = K \times t \]

Expressed in units of concentration/ time, \( K \) is zero order release constant and t is the time in hrs.

**First-order,**

\[ \log C = \log C_o - \frac{K_o}{2.303} t \]

Where \( C \) is the concentration, \( C_o \) is the initial concentration of drug, \( K \) is the first-order rate constant, and t is the time.

**Higuchi,**

\[ Q_s = K_{H} \times t^{1/2} \]

Where \( Q_s \) is the amount of release drug in time t, K is the kinetic constant and t is the time in hrs.

**Korsemeyer peppas,**

\[ M_t / M_{\infty} = K \times t^{n} \]

Where Mt represents amount of the released drug at time t, M is the overall amount of the drug (Whole dose). The value of n indicates the drug release mechanism related to the geometrical shape of the delivery system, if the exponent n = 0.5, then the drug release mechanism is Fickian diffusion. If n < 0.5 the mechanism is quasi-Fickian diffusion, and 0.5 < n < 0.5, then it is non-Fickian or anomalous diffusion and when n = 1.0 mechanism is non-Fickian case II diffusion, n > 1.0 mechanism is non-Fickian super case II.

**Stability study**

In present study, stability studies were carried out at 40°C and 75% RH for a specific time period up to 90 days for optimized formulation. For stability study, the tablets were sealed in aluminium packing coated inside with polyethylene. These sample containers were placed in desiccators maintained at 75%RH.

**RESULTS AND DISCUSSION**

**Preformulation Studies**

Imatinib mesylate is very soluble in DMSO, 0.1 NHCL and it is also soluble in water due to its salt form. So, we select 0.1 N HCL is solvent for further studies as it is acidic medium and its drug solubility (as represented in Figure 1).

![Fig. 1: Solubility of Imatinib Mesylate in different solvents](image)

**Drug-excipient interaction by using IR graphs**

From below spectras, it was found that there is no significant change in plain drug spectra and drug with excipient spectra wavelengths (as shown in Figure 2). So there is no Drug-Excipient interaction.

**Micromeritic properties of Imatinib Mesylate and polymers**

Micromeritic properties of pure drug and polymers indicating that the flow properties of drug and polymers were low (as shown in Table 2). To increase the flow properties of drug and polymers granules were prepared by wet granulation method.

**Granule properties of all batches**

The granulation characteristics (as shown in Table 3) are the most important interest to formulation scientist and therefore most universally measured. These basic measurements of the granulation have been used to develop and monitor the manufacture of many successful pharmaceutical dosage forms.

The bulk densities of granules were found to be between 0.446±0.02 to 0.482±0.04 g/cm³. This indicates good packing capacity of granules. Carr’s index evaluated interparticulate cohesive properties with angle of repose measurements and studied the effects of packing geometry of solids with bulk and tapped density. Bulk density and tapped density measurements found that density of a powder depends on particle packing and that density changes as the powder consolidates. Values of Carr’s index below 15% usually show good flow characteristics, but readings above 25% indicate poor flowability. Carr’s index was found to be between 11.3±0.10 to 146.6±0.07.

Hausner’s ratio is simple method to evaluate stability of powder column and to estimate flow properties. Low range was observed of Hausner’s ratio that indicates good flow ability. Many different types of angular properties have been employed to assess flow ability. Angle of repose is suited for particle > 150 μm. Values of angle of repose ≤ 30º generally indicate the free flowing material and angle of ≥ 40º suggest a poor flowing material. The angle of repose is indicative of the flowability of the material. The angle of repose of all formulations fell within the range of 27.09±0.17 to 29.9±0.15 i.e. granules were of good flow properties.
Fig. 2: FTIR Spectra of a) Imatinib Mesylate b) HPMC K4M c) HPMC K15M d) Formulation F4

Table 2: Micromeritic properties of Imatinib Mesylate and polymers

<table>
<thead>
<tr>
<th>Material</th>
<th>Bulk density (gm/cc ± SD, n=3)</th>
<th>Tapped density (gm/cc ± SD, n=3)</th>
<th>Carr’s index (%) ± SD, n=3</th>
<th>Hausner’s ratio ± SD, n=3</th>
<th>Angle of repose (degree) ± SD, n=3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Imatinib Mesylate</td>
<td>0.233</td>
<td>0.352</td>
<td>33.8</td>
<td>1.51</td>
<td>36.72</td>
</tr>
<tr>
<td>HPMC K4M</td>
<td>0.341</td>
<td>0.557</td>
<td>39</td>
<td>1.63</td>
<td>41.28</td>
</tr>
<tr>
<td>HPMC K15M</td>
<td>0.322</td>
<td>0.543</td>
<td>38</td>
<td>1.62</td>
<td>38.32</td>
</tr>
</tbody>
</table>

Table 3: Granule properties of all batches

<table>
<thead>
<tr>
<th>Batch Code</th>
<th>Bulk density (gm/cc ± SD, n=3)</th>
<th>Tapped density (gm/cc ± SD, n=3)</th>
<th>Carr’s index (%) ± SD, n=3</th>
<th>Hausner’s ratio ± SD, n=3</th>
<th>Angle of repose (degree) ± SD, n=3</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>0.45±0.07</td>
<td>0.525±0.09</td>
<td>14.5±0.06</td>
<td>1.2±0.03</td>
<td>28.60±0.05</td>
</tr>
<tr>
<td>F2</td>
<td>0.446±0.02</td>
<td>0.514±0.04</td>
<td>13.2±0.09</td>
<td>1.15±0.05</td>
<td>28.67±0.08</td>
</tr>
<tr>
<td>F3</td>
<td>0.482±0.04</td>
<td>0.544±0.02</td>
<td>11.3±0.10</td>
<td>1.12±0.06</td>
<td>28.08±0.13</td>
</tr>
<tr>
<td>F4</td>
<td>0.476±0.07</td>
<td>0.551±0.06</td>
<td>14.6±0.07</td>
<td>1.15±0.05</td>
<td>29.9±0.15</td>
</tr>
</tbody>
</table>

All values are expressed average ± SD; (n=3)
Evaluation of Tablet characteristics

The floating tablets of Imatinib Mesylate were prepared by effervescent technique. The tablets were evaluated (as shown in Table 4) for weight variation, thickness, hardness, friability and drug content.

The total weight of each formulation was maintained constant; the weight variations of the tablets were within the permissible limits of 5%, as specified for tablet weighing more than 324mg. Weight of the tablet was fixed at 400mg and the weight variation for every batch was tested and found within the acceptance limits.

Hardness of the tablets of all batches was found to be between 6.2±0.4 to 6.5±0.5 kg/cm² and was maintained for all the batches in order to minimize the effect of hardness on the drug release because, the effect of polymer concentration is the only area of interest.

Tablet thickness was also used to assess the quality of tablets. Under uniform conditions of manufacture, the total weight of tablet and thickness were linearly related. The thickness of floating tablets ranged from 3.52±0.03 to 3.57±0.05 mm and linearly correlated with the weight of the tablets.

Friability test of all the formulations was found satisfactory showing enough resistance to the mechanical shock and abrasion. The values of friability are within the limit.

Drug content uniformity in all formulations was calculated and the percent of active ingredient ranged from 98.25±1.8 to 98.91±1.5 indicating good content uniformity in the prepared formulations as per within the limits as per IP. (Not less than 98% and not more than 102%).

<table>
<thead>
<tr>
<th>Formula code</th>
<th>Weight variation(mg)</th>
<th>Hardness (kg/cm²)</th>
<th>Thickness (mm)</th>
<th>Friability (%)</th>
<th>Drug content (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>405±1.6</td>
<td>6.5±0.5</td>
<td>3.52±0.03</td>
<td>0.64±0.06</td>
<td>98.46±2.3</td>
</tr>
<tr>
<td>F2</td>
<td>404±1.5</td>
<td>6.5±0.2</td>
<td>3.57±0.03</td>
<td>0.95±0.06</td>
<td>98.91±1.5</td>
</tr>
<tr>
<td>F3</td>
<td>402±1.6</td>
<td>6.3±0.6</td>
<td>3.52±0.06</td>
<td>0.71±0.06</td>
<td>98.75±2.3</td>
</tr>
<tr>
<td>F4</td>
<td>403±1.5</td>
<td>6.2±0.4</td>
<td>3.57±0.05</td>
<td>0.66±0.02</td>
<td>98.25±1.8</td>
</tr>
</tbody>
</table>

All values are expressed average ± SD; (n=3)

Effect of Sodium bicarbonate concentration on floating lag time

Placebo tablets were prepared with changing the concentration of sodium bicarbonate concentration along with same concentration of drug and polymer. Finally, it can be concluded that the concentration of sodium bicarbonate increases the floating lag time decreased.

Buoyancy determination

The formulated tablets on immersion in 0.1N Hydrochloric acid media they remain buoyant for 12 h with lag time of 96 to 257 seconds. Sodium bicarbonate was added as a gas-generating agent. The optimized concentration of effervescent mixture utilized aided in the buoyancy of all tablets. This may be due to the fact that effervescent mixture in tablets produced CO₂ that was trapped in swollen matrix, thus decreasing the density of the tablet below 1 making the tablets buoyant. All the batches showed good in vitro buoyancy.

Water uptake studies

The swelling index was calculated with respect to time. As time increases, the swelling index was increased, because weight gain by tablet was increased proportionally with rate of hydration. Later on, it decreased gradually due to dissolution of outermost gelled layer of tablet into dissolution medium (as represented in Figure 3). The direct relation was observed between swelling index and polymer concentration.

In vitro dissolution testing

The in vitro dissolution testing was performed (as represented in Figure 4,5) and the results of the formulations were expressed as mean ± S.D (n=3).

First the dissolution studies of pure Imatinib Mesylate (IM), marketed formulation CELONIB (100mg) (FM) were conducted, 0.1 N HCL used as dissolution medium. Imatinib Mesylate highly soluble in 0.1N HCL, so the complete drug release was observed within 10 minutes. In case of marketed formulation, it's a conventional dosage form that's why drastic drug release was takes place within 45 minutes. Because of this drastic release, dose related side effects will occurred in long term therapy, the dosing frequency also high due to this, fluctuation in plasma concentration also takes place. To reduce these consequences controlled release Imatinib Mesylate floating matrix tablets were fabricated.

In vitro dissolution study of formulations F1, F2, and F3 prepared with combination of HPMC K4M and HPMC K15M done in 0.1N HCL and the percent of drug release from formulations F1, F2, and F3 was 98.26 in6hr, 99.69 in 8hr and 98.50 in 10hr respectively. These
formulations unable to sustain the drug release desired period of time. This is because, being water soluble polymers and being available in less concentrations, they dissolve and form pores filled liquid, in which drug can there after diffuse in dissolution medium and the drug release was more in these formulations. Drug release was retarded for 12hrs in F4 formulation due to the presence of increased concentration of HPMC K15M polymer and also shown 99.46% of drug release. Hence, it was considered as the best formulation. All these four formulations floated for 12hr.

In all formulations as the concentration of HPMC K15M polymer increased the rate of drug release from the formulations was decreased.

\[
\begin{array}{|c|c|c|}
\hline
\text{Formulation} & \text{F1} & \text{F2} & \text{F3} & \text{F4} \\
\hline
\text{R}^2 \text{ value of Zero order} & 0.9631 & 0.975 & 0.973 & 0.974 \\
\hline
\text{R}^2 \text{ value of Higuchi} & 0.9631 & 0.975 & 0.973 & 0.974 \\
\hline
\end{array}
\]

Dissolution Profile Modeling

The mechanism of release for the all formulations was determined by finding the \( R^2 \) value for each kinetic model viz. Zero-order, First-order, Higuchi, and Korsmeyer-Peppas corresponding to the release data of formulations. For Imatinib Mesylate (IM) \( R^2 \) value of Higuchi model near to 1 and marketed formulation (FM) \( R^2 \) value of Higuchi and zero-order models are very near to 1 than the \( R^2 \) values of other kinetic models (as shown in Table 5). Thus it can be said that the drug release follows Higuchi and zero-order model mechanism.

For all formulations F1 to F4, \( R^2 \) value of Zero order and Higuchi models are very near to 1 when compare with other models. And optimized formulation F4, \( R^2 \) values of Zero order and Higuchi are 0.9631, 0.975 respectively.
It can be concluded that all formulations F1 to F4 drug release patterns were following the Zero order and Higuchi release kinetic models. The n values of Korsmeyer-Peppas model of the all formulations are in between 0.38 to 0.46. Therefore, the most probable mechanism that the release patterns of the formulations followed was Fickian diffusion.

<table>
<thead>
<tr>
<th>Formulation</th>
<th>Zero order</th>
<th>First order</th>
<th>Higuchi</th>
<th>Korsmeyer &amp; Peppas</th>
<th>Peppas(n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>0.9498</td>
<td>0.9047</td>
<td>0.9966</td>
<td>0.7796</td>
<td>0.38</td>
</tr>
<tr>
<td>F2</td>
<td>0.967</td>
<td>0.8185</td>
<td>0.9936</td>
<td>0.7704</td>
<td>0.44</td>
</tr>
<tr>
<td>F3</td>
<td>0.9602</td>
<td>0.8839</td>
<td>0.9803</td>
<td>0.7897</td>
<td>0.45</td>
</tr>
<tr>
<td>F4</td>
<td>0.9631</td>
<td>0.7449</td>
<td>0.975</td>
<td>0.8051</td>
<td>0.46</td>
</tr>
</tbody>
</table>

Stability studies

Stability studies revealed that there was no significant change in the drug content, the release rate of the drug and buoyancy characters of the optimized formulation F4 kept on stability studies(400°C/75% RH).

CONCLUSION

Imatinib Mesylate tablets can be formulated to increase the gastric residence time and thereby achieve the slow release of the drug in a constant manner. Formulation F4 gave better control led drug residence time and thereby achieve the slow release of the drug in a constant manner. Formulation F4 gave better controlled drug release and floating properties in comparison to the other formulations. Formulation F4 contains combination of hydrophilic polymers; it is better polymer combination to produce promising results for their evaluation like Hardness, weight variation, floating lag time, floating time, and in vitro drug release. Finally, it can be concluded that Imatinib Mesylate was good candidate for the preparation of Floating drug delivery system due to its gastric stability, gastric absorption.

ACKNOWLEDGEMENT

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