DEVELOPMENT OF NATURAL AND MODIFIED GUM BASED SUSTAINED-RELEASE FILM-COATED TABLETS CONTAINING POORLY WATER-SOLUBLE DRUG

AJAY KUMAR SHUKLA*, RAM SINGH BISHNOI, MANISH KUMAR, C.P. JAIN

Department of Pharmaceutical Sciences, MohanLal Sukhadia University, Udaipur, Rajasthan, India. Email: ashukla1007@gmail.com

Received: 16 October 2018, Revised and Accepted: 19 December 2018

ABSTRACT

Objectives: The objective of this work was to develop the characteristics of natural and modified gum based sustained-release film-coated tablets containing poorly water-soluble drug.

Methods: Tamarind seed gum (TSG), fenugreek seed gum (FSG), sodium trimetaphosphate, hydroxypropyl methylcellulose (HPMC), sodium alginate (SA), and nifedipine (NFD) were used. The core tablets of nifedipine were prepared and evaluated for weight, diameters, thickness, hardness, and disintegration time. The coated tablets were prepared using 2% w/v solution of TSG, MTSG, FSG, and MFSG. The in vitro release rates of nifedipine from these coated tablets were compared with the release rate of drug from the tablets coated with 2% w/v of HPMC.

Results: The tablets coated with natural and modified TSG sustained the release of the drug up to 11 h and 14 h, respectively, and natural and modified FSG sustained release the drug up to 9 h and 11 h, respectively, The tablets coated with HPMC sustained released the drug up to 15 h. The drug release profile of tablets coated with modified TSG was comparable to the release profile of tablets coated with HPMC.

Conclusion: On the basis of the release profile, it is concluded that unmodified and modified TSG can be used as release rate controlling membrane.

Keywords: Tamarind seed gum, Fenugreek seed gum, Pharmaceutical Excipient, Natural polymer.

INTRODUCTION

Film coating is a most significant unit operation in the pharmaceutical industry [1]. Film coatings are used for various purposes such as improvement of visual qualities of dosage forms, masking disagreeable taste or odor, improving stability, and modifying the release characteristics of the drug [2]. Film coating is applied to a variety of pharmaceutical products such as tablets, beads, pellets, granules, capsules, and drug crystal [3]. Polymer is the main ingredient in the majority of film-coated formulations and it may be from different origin (natural, synthetic, or semisynthetic), including cellulose, acrylics, vinyl, and combination polymers [4,5].

Fenugreek gum is a natural polymer and is extracted from the seeds of Trigonella foenum-graecum (Family Leguminosae). It is cultivated in Northern Africa, the Mediterranean, Western Asia, Northern India, and in Canada [6,7]. Application of fenugreek seed gum (FSG) has been reported as oral drug release retardant [7-9], binder [10-12], mucoadhesive [13,14], emulsifiers [15-17], suspending agent [18], gelling agent [19], and formulation of nanoparticles [20,21].

Tamarind seed gum (TSG) has wide application in the drug delivery. It has been reported that gum can be successfully extracted from tamarind seed using water solvent extraction method. The extracted gum is reported to be used as a gelling agent. It is used in the formulation of sustained-release dosage form of water-soluble and water-insoluble drugs [21,22]. For water-soluble drugs, the release of the drug can also control by partially cross-linking the matrix [23-25], and the extent of release can be varied by controlling the degree of cross-linking.

METHODS

Tamarind seed, fenugreek seed, sodium trimetaphosphate (STMP), nifedipine, hydroxypropyl methylcellulose (HPMC), and all other chemicals used were of analytical grade.

Isolation of FSG
The seeds of fenugreek were collected and washed with water. Then, after dried in hot air oven and seeds were crushed, soaked in water for 6 h, boiled for 30 min, and left for 1 h to allow complete release of the gum into the water. The gum was separated using a 4-fold muslin cloth bag to remove the marc from the solution. Then, ethanol (in the volumes of 3 times to the volume of filtrate) was added for precipitation of gum, the collected dried gum then grinded and passed through a # 120 sieve and stored in a desiccator at room temperature.

Isolation of TSG
The seeds of tamarind were collected and washed with water. Then, after dried in hot air oven, the dried tamarind seeds were soaked in distilled water for 1 week, and then, the external cover was removed and obtained white part of seeds was crushed. The crushed seeds of tamarind were soaked in water for 12 h and boiled for 30 min and left for 1 h to allow complete release of the gum into the water. The gum was separated using a 4-fold muslin cloth bag to remove the marc from the solution. Then, ethanol (in the volumes of 3 times to the volume of filtrate) was added for precipitation of gum, the collected dried gum then grinded and passed through a # 120 sieve and stored in a desiccator at room temperature.

Modification of isolated gum with STMP
About 1 g of STMP and 1 g of natural seed gum were taken and dissolved separately in 50 ml of distilled water. Then, after prepared STMP and 5 ml of 0.1 NaOH solutions were slowly added with stirring to 1 g of natural seed gum solution. The prepared solution (100 ml) was stirred for 2 h and poured into Petri dish and dried at 60°C for 24 h. The dried complex (modified gum) was grinded and passed through a # 120 aperture sieve and stored in an air-tight container at room temperature [26].

Characterization of gum
The physical nature of gum was characterized using Fourier transform infrared spectroscopic (FTIR) and X-ray diffraction (XRD). The scanning
range of FTIR was 4000–500 cm$^{-1}$ used. FTIR spectra are shown in Figs. 1-4.

Surface morphology of natural and modified seed gum was studied by XRD. The XRD diffractions spectra were recorded. XRD diffractions spectra of natural and modified seed gum are shown in Figs. 5-9.

**Drug-excipient compatibility study by FTIR**
The compatibility of the drug with excipients was studied by FTIR spectroscopy (FTIR, Bruker). All characteristic peaks of the drug were observed in the FTIR spectra of drug and excipients. Drug-excipient compatibility study by FTIR spectra is shown in Fig. 10.

**Evaluation of blends for core tablets [27,28]**
The powder mixture before compression was evaluated for the angle of repose, bulk density g/cm$^3$, tapped density g/cm$^3$, Carr’s index, and Hausner ratio. Composition of core tablet and the results of evaluation are shown in Tables 1-2.

---

**Preparation of core tablets**
Core tablets of NFD were prepared using the composition given in Table 5 by direct compression method. Accurately weighed quantities of the drug, polymer mixture were passed through sieve no #80 and mixed well for 10 min. Sufficient quantity of the diluent microcrystalline cellulose was used to raise the total bulk of the tablets to a weight of 400 mg each. The resulting powder blend was compressed on single tablet punching machine. The tablets were evaluated for parameters such as weight uniformity, friability, and hardness.

**Table 1: Composition of core tablet**

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drug</td>
<td>20 mg</td>
</tr>
<tr>
<td>PVP</td>
<td>10 mg</td>
</tr>
<tr>
<td>Lactose</td>
<td>265 mg</td>
</tr>
<tr>
<td>Mg. stearate</td>
<td>5 mg</td>
</tr>
</tbody>
</table>

---

**Fig. 1:** Fourier transform infrared spectra of fenugreek seed gum

**Fig. 2:** Fourier transform infrared spectra of tamarind seed gum

**Fig. 3:** Fourier transform infrared spectra of MFSG

**Fig. 4:** Fourier transform infrared spectra of MTSG
Evaluation of core tablets
The core tablets were evaluated for the following parameters such as weight uniformity, diameters (cm), thickness (cm), hardness kg/cm², and disintegration time (min.) [23]. The results of evaluated coated tablets are shown in Table 3.

Dimensions
The diameter and the thickness of tablets were determined using Vernier caliper.

Weight uniformity
A total of 20 tablets were sample out from each batch. The individually weight of tablets was taken and the average weight of tablets was calculated using the formula. The results are shown in Table 3.

Average weight of tablets = (total weight of tablets)/number of tablets

Hardness test
The resistance of tablets to shipping or breakage under the conditions of storage, transportations, and handling before usage depends on its hardness. The hardness of tablet was measured by Pfizer hardness tester. The hardness was measured in terms of kg/cm². The results are shown in Table 3.

Preparation of coating solution
The coating solution was prepared by dissolving coating materials 2% w/v natural seed gum and 1% sodium alginate, 2% w/v modified seed gum and 1% sodium alginate, and 2% w/v HPMC and 1% sodium alginate, in warm water using a magnetic stirrer and 0.5% w/v titanium oxide (as opacifier) were added to each of above coating solution. The solutions were stirred for 60 min at room temperature. Composition of coating solutions is given in Table 3.

Viscosity of coating solution
The viscosity of prepared coating suspensions was determined with Brookfield LVDV-IV+ digital rheometer using spindle 4 at 100 rpm. The results of evaluation are shown in Table 4.

Preparation of film-coated tablets
The coating solution was prepared by dissolving the gum in warm water using a magnetic stirrer. On complete solubilization of the polymer and gum (2% w/v of selected gum and 1% w/v of sodium alginate polymer), talc (0.1% w/v) as antiadherent and titanium dioxide (0.5% w/v) as opacifier were added. The solution was stirred for 60 min at room temperature. Then, prepared core tablets were dipped into prepared coating solution for 5 min and after coated tablets were dipped into 5% w/v solution of CaCl₂ for 5 min and after were dried in hot air oven.

Evaluation of film-coated tablets [27]
The film-coated tablets were evaluated for the quality parameters, namely tablet thickness, uniformity of weight, crushing strength, and friability. A Vernier caliper was used to measure the thickness and diameters of film-coated tablets.

Table 2: Evaluation of blends for core tablet

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>TSG</th>
<th>FSG</th>
<th>MTSG</th>
<th>MFSG</th>
<th>HPMC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Angle of repose⁶</td>
<td>24.95±0.01</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bulk density g/cm³</td>
<td>0.428±0.01</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tapped density g/cm³</td>
<td>0.500±0.04</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carr’s index%</td>
<td>14.40±0.03</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hausner ratio</td>
<td>1.4±0.07</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hardness kg/cm²</td>
<td>6±0.01</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean±SD, n=3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3: Composition of film-coating solution

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>TSG</th>
<th>FSG</th>
<th>MTSG</th>
<th>MFSG</th>
<th>HPMC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Batch code</td>
<td>A</td>
<td>B</td>
<td>C</td>
<td>D</td>
<td>D</td>
</tr>
<tr>
<td>Gum %</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Sodium alginate%</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Titanium dioxide</td>
<td>10 mg</td>
<td>10 mg</td>
<td>10 mg</td>
<td>10 mg</td>
<td>10 mg</td>
</tr>
<tr>
<td>Mean±SD, n=3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 4: Viscosity of coating solution

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>TSG-SA</th>
<th>FSG-SA</th>
<th>MTSG-SA</th>
<th>MFSG-SA</th>
<th>HPMC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Viscosity cp</td>
<td>220±0.01</td>
<td>236±0.03</td>
<td>107±0.07</td>
<td>103±0.02</td>
<td>202±0.06</td>
</tr>
<tr>
<td>Mean±SD, n=3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>4</td>
</tr>
</tbody>
</table>

Table 5: Evaluation of film-coated tablet

<table>
<thead>
<tr>
<th>Evaluation of coated tab.</th>
<th>FSG</th>
<th>TSG</th>
<th>MFSG</th>
<th>MTSG</th>
<th>HPMC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hardness Kg/m²²</td>
<td>1.0±0.01</td>
<td>1.2±0.04</td>
<td>9.1±0.02</td>
<td>11.1±0.06</td>
<td>11.7±0.03</td>
</tr>
<tr>
<td>Friability %</td>
<td>0.7±0.02</td>
<td>0.29±0.08</td>
<td>0.49±0.01</td>
<td>0.39±0.02</td>
<td>0.35±0.01</td>
</tr>
<tr>
<td>Diameter cm</td>
<td>0.9±0.02</td>
<td>0.9±0.07</td>
<td>0.9±0.04</td>
<td>0.9±0.09</td>
<td>0.9±0.08</td>
</tr>
<tr>
<td>Thickness cm</td>
<td>0.9±0.08</td>
<td>0.9±0.09</td>
<td>0.9±0.08</td>
<td>0.9±0.07</td>
<td>0.9±0.06</td>
</tr>
<tr>
<td>Disintegration time (min)</td>
<td>12.55±0.01</td>
<td>17.04±0.07</td>
<td>12.01±0.08</td>
<td>14.10±0.04</td>
<td>15.06±0.09</td>
</tr>
<tr>
<td>Mean±SD, n=3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Disintegration time
Disintegration testing of tablets was carried out in the six tablet basket rack USP disintegration apparatus. One tablet was introduced into each tube of the basket rack assembly of the disintegration apparatus without disc. The assembly was positioned in the beaker containing 900 ml distilled water. The temperature was maintained at 37±2°C and operated the apparatus for 2 h.

Dissolution studies
*In vitro* drug release from natural and modified TSG, natural and modified FSG, and HPMC film-coated tablets was studied, using eight-station (USP) Type II dissolution apparatus at 37±0.5°C and at 50 rpm speed in 0.1 N HCl as dissolution media for 2 h. From the dissolution medium, 5 ml of the samples was withdrawn at the specific time intervals and replaced with an equal volume of fresh medium (5 ml) to maintain sink conditions. After filtration and dilution, each sample was analyzed using double-beam UV visible spectrophotometer at selected wavelength 238 nm λ max. This study was performed thrice for each batch. After 2 h, dissolution media was replaced by phosphate buffer pH 7.4 [23]. *In vitro* drug release data and graph of NFD from film-coated tablets are shown in Table 5.

**RESULTS AND DISCUSSION**
The FTIR spectra of the natural gum (FSG and TSG) are given in Figs. 1 and 2, respectively, which indicated that the FSG and TSG were carbohydrates in nature. Modified natural gum successfully cross-linked with STMP, characterization of modified natural gums, MFG and MTSG were investigated by FTIR spectra of modified natural gums, MFG and MTSG are shown in Figs. 3 and 4.

The absence of sharp peak at 1700–1800 cm⁻¹ in the FTIR spectrum indicates that there is no carboxyl group in the extracted sample. On the other hand, the presence of peak at 1000–1200 cm⁻¹ indicates the presence of secondary alcohols. The range of wave numbers 800–1200 cm⁻¹ represents carbohydrates nature.

The surface characterization of natural (FSG and TSG) and modified (MFG and MTSG), using XRD spectra of the gum was taken. The XRD spectra of natural FSG and TSG exhibited high rough surface compare to modified natural gum with pores and crevices on it, XRD spectra of natural and modified gums are shown in Figs. 5-8.

From the XRD, it was also evident that the particle size of the powders was not uniform and the size distribution was not within a narrow range. The powder contains larger to ultrafine particles. This might be the reason for the “heavy” nature of the powders. The powders exhibit a “closer” packing arrangement, in which, the smaller particles fill the voids between larger particles and reduce the bulkiness. The low porosity values also indicate this packing arrangement. The close packing can also be responsible for poor flow properties of FSG. However, after modified natural gum MTSG and MFG is exhibited good flowing proper.

The compatibility study of the drug with excipients was studied by FTIR spectroscopy. All characteristic peaks of the drug were observed in the FTIR spectra of drug and excipients. The results showed no chemical interaction between drug and various excipients alone or in combination. The FTIR spectra of drug nifedipine (NFD) showed characteristic peaks at 3858.75 cm⁻¹ (hydroxyl group secondary of NFD), 3670.03 cm⁻¹ (N-H overlap of amide with O-H of MTSG), 3555.66 cm⁻¹ (N-H stretching, OH alcohol of MFSG), 3359.77 cm⁻¹ (aromatic CH NFD), 3120.28 cm⁻¹ (C≡N, C≡C of NFD), 2173.78 cm⁻¹ (C≡C stretching of TSG), 1916.93 cm⁻¹ (C≡C stretching of MFG), 1790.93 cm⁻¹ (tertiary amide of NFD), 1681.94–1655 cm⁻¹ (CH OH represents carbohydrates nature.

The FTIR spectra of the natural gum (FSG and TSG) and modified (MFG and MTSG), using XRD spectra of the gum was taken. The XRD spectra of natural FSG and TSG exhibited high rough surface compare to modified natural gum with pores and crevices on it, XRD spectra of natural and modified gums are shown in Figs. 5-8.

From the XRD, it was also evident that the particle size of the powders was not uniform and the size distribution was not within a narrow range. The powder contains larger to ultrafine particles. This might be the reason for the “heavy” nature of the powders. The powders exhibit a “closer” packing arrangement, in which, the smaller particles fill the voids between larger particles and reduce the bulkiness. The low porosity values also indicate this packing arrangement. The close packing can also be responsible for poor flow properties of FSG. However, after modified natural gum MTSG and MFG is exhibited good flowing proper.
Shukla et al.


stretching vibration of FSG), 1576.73 cm⁻¹ (CH₂ stretching of TSG), 1521.64 cm⁻¹ (N-O nitro compound NFD), 1423.99 cm⁻¹ (OH bending carboxylic acid MTSG), 1277.67 cm⁻¹ (C-O alkyl aryl ether of NFD), 1097.50 cm⁻¹ primary alcohol of NFD), and 1053 cm⁻¹ (CH stretching vibration of FSG). It is clear that functionalities of drug have remained unchanged including intensities of the peak. Hence, in physical mixture, no interaction was found, which supports the formulation of the formulation of film-coating tablets. Compatibility study of nifedipine and excipients FTIR spectra are shown in Figs. 9 and 10.

Blends of nifedipine core tablets were prepared. All the ingredients mixed well and evaluated as parameters such as angle of repose, bulk density, tapped density, Car’s index, and Hausner ratio, composition of core tablets is shown in Table 1. The results of these parameters were found in the range of angle of repose 24.95±0.01, bulk density 0.428±0.11, tapped density 0.500±0.04, Car’s index 14.40±0.03, Hausner ratio 1.4±0.07, and hardness kg/cm² ±0.01. The results of evaluated parameters represented that the property of blends was excellent. Evaluation results of nifedipine core tablets are shown in Table 2.

Composition of coating blends is shown in Table 3. The viscosity of coating solution was determined using Brookfield viscometer spindle-4 at 100 rpm. Results are shown in Table 4.

The evaluation results of film-coated nifedipine tablets are shown in Table 5. The coated nifedipine tablets with HPMC had better hardness compared to those coated with fenugreek and TSG, and the disintegration time of core tablets increased from 3 min (core) to 12.33, 10.61, 8.93, and 9.40 min after coating with FSG. The order of disintegration time of formulation was found: HPMC > TSG > FSG > core. Developed film-coating tablets using FSG, TSG, MFSG, MTSG, HPMC, diameter (cm) 0.9±0.02, 0.9±0.07, 0.9±0.04, 0.9±0.09, and 0.9±0.08, and thickness (cm) 0.9±0.08, 0.95±0.09, 0.9±0.08, 0.9±0.07, and 0.9±0.06 were found.

The drug dissolution profile of the coated tablets is shown in Table 6. Coated tablets, coated with natural seed gum combination of 2% w/v FSG-1% w/v-SA and 2% w/v TSG-1% w/v-SA, modified seed gum 2% w/v MFSG-1% w/v-SA, 2% w/v MTSG-1% w/v-SA, and 2% w/v HPMC-1% w/v-SA, respectively, shown. The drug release rate of coated tablets was found up to (FSG) 8 h, (TSG) 11 h, (MFSG) 12 h, (MTSG) 14 h, and (HPMC-SA) 15 h, respectively. Hence, drug release rate of TSG film-coated tablets was sustained as compare to core tablets. Hence, TSG can be used for the formulation of sustained release of drugs. The drug dissolution profile of the coated tablets is shown in Fig. 11.
CONCLUSION
The potential of natural and modified FSG and TSG was investigated using nifedipine as model drug and found that natural and modified form of TSG better film-coating potential than fenugreek gum and similar to near standard film-forming agent HPMC. The coated tablets were evaluated using parameters as uniformity of weight, friability, disintegration time, and dissolution profiles. TSG can be used for the improvement of visual qualities of dosage forms, masking disagreeable taste or odor, improving stability, and modifying the sustained-release characteristics of the drug.

ACKNOWLEDGMENTS
The authors would like to thank the Department of Pharmaceutical Sciences, Department of Physics, and Department of Chemistry, Faculty of Sciences, Mohanlal Sukhadia University, Udaipur, Rajasthan, India, for providing all necessary facilities.

AUTHORS’ CONTRIBUTIONS
All the authors have contributed equally.

CONFLICTS OF INTEREST
No conflicts of interest related with this work.

REFERENCES