Development of the Modified *Ocimum gratissimum* Seeds for Orally Disintegrating Tablets

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Abstract: Background: Natural materials have been encouraged in controlled drug release and improved drug bioavailability.

Objective: This study aimed to develop a modification process for the use of a natural material, *Ocimum gratissimum* seeds (OGS), in Orally Disintegrating Tablets (ODTs).

Methods: The OGS was investigated with four different modification processes including only milling, swelling, swelling/milling, and swelling/milling/incubation. The ODTs containing the modified OGS as a disintegrant were prepared by the wet granulation method. Furthermore, an evaluation to assess parameters of tablets, such as weight variation, hardness, friability, wetting time, disintegration time, drug content, and dissolution studies, was performed.

Results: The modification of OGS using the swelling/milling process resulted in a completion of OGS modification, leading to an ideal wetting time, disintegrating time, and dissolution rate. The OGS concentrations also affected the wetting and disintegrating time with the optimal range of ODTs from 15% to 20%. On the other hand, the modification with the incubation processes varied by temperature and time increased the wetting time and disintegrating time.

Conclusions: The modified OGS demonstrated that it is a potential material with the advantages of cost-effectiveness, non-toxicity and easy manufacture in the preparation of ODTs.

Keyword: Orally disintegrating tablets, *Ocimum gratissimum* seeds, disintegrant, modification, oral delivery, natural material.

1. INTRODUCTION

The development of natural products has been encouraged in recent pharmaceutical research of controlled drug release and improved drug bioavailability [1-8]. *Ocimum gratissimum* seeds (OGSs) were investigated in several studies as a suspending agent or a disintegrant [9, 10]. On the contact with water, a mucilaginous layer is formed surrounding the seeds due to its capability of excellent water uptake, demonstrating the swelling power of OGSs. A large amount of polysaccharide with a high capacity for hydration was attributed to the formation of the mucilaginous layer of the swollen seeds, showing the potential application in pharmaceutical formulations [11, 12].

Although tablets and capsules are extensively used as solid oral dosage forms in manufacturing operations, for some patients with persistent trouble in swallowing these kinds of dosage forms pose a real problem. Orally Disintegrating Tablets (ODTs) would be an appropriate choice for these patients [13]. ODTs are normally known as patient-friendly dosage forms, which are particularly suitable for pediatric and geriatric patients. ODTs rapidly disintegrate in the oral cavity by absorbing a small amount of water [14]. The absorption site of ODTs may improve the pharmacokinetics of therapeutic agents by reducing first-pass metabolism [15].

To develop an ODT formulation, a disintegrant should be noticed as a critical excipient in the formulation as it plays an important role in ensuring the appropriate disintegration time and dissolution rate of the ODTs [16, 17]. ODTs hence provide a comfortable and safe drug delivery to therapeutic and patient compliance requirements [15]. Furthermore, ODTs also face the challenge of high production costs [15, 18]. Therefore, the present research aims to develop a modified OGS and investigate its role in formulating and manufacturing ODTs. This study may pave a way to use natural materials for the development of ODTs entailing great properties of safety and cost-effectiveness.
Development of the Modified Ocimum gratissimum Seeds

2. MATERIALS AND METHODS

2.1. Materials

Acetaminophen was purchased from Hebei Jiheng Pharmaceutical Co., Ltd. (China). Polyvinylpyrrolidone K30 (PVP K30) was purchased from BASF Group (Germany). Ocimum gratissimum seeds (OGS) were purchased at the local market and stored at room temperature (25°C) for the study. Aspartame was obtained from Anhui Wanhe Pharmaceutical Co., Ltd (China). Magnesium stearate (MgS) was purchased from Nitika Pharmaceutical Specialties Pvt. Ltd. (India). Mannitol (Pearlitol®) was obtained from Roquette Pharma Company (France).

2.2. Methods

2.2.1. Preparation of Modified OGSs for Tableting

**Method 1 (M1):** The pure seeds were simply milled and sieved with a 500 μm sieve.

**Method 2 (M2):** The powder of M1 was swelled in the distilled water for 1 h. The swelled OGS was dried at 60°C.

**Method 3 (M3):** The powder of M1 was also swelled in the distilled water for 1 h and immediately subjected to milling and sieving with a 500 μm sieve. The samples were then dried at 60°C.

**Method 4 (M4):** The powder of M1 was also swelled in the distilled water for 1 h and immediately subjected to milling and sieving with a 500 μm sieve. However, the samples were incubated at 50°C (and 90°C) for 3 h or 6 h before subjected to dry at 60°C.

In all methods mentioned above, the batch size equivalent to 10 g OGS seeds was milled by the blender machine (Sunari KT-BL 28, China) in 15 min to ensure all particles passed through a 500 μm sieve. Moreover, the samples after drying at 60°C were passed again through a 500 μm sieve to get the final powder for tablet preparations. All formulations are described in below Table 1. The ODT tablets were prepared by wet granulation with a total weight of 200 mg (acetaminophen 40%, PVP K30 4%, aspartame 3%, magnesium stearate 1%, modified OGS from 5 to 20%, and the amount of mannitol was adjusted according to modified OGS). Briefly, after wet granulation, the dried granules containing acetaminophen, PVP K30, and aspartame were mixed with modified OGS, mannitol and magnesium stearate for tableting (batch size of 100 tablets). The hardness of the tablet was controlled in the range of 20-40 N.

2.2.2. Uniformity of Tablet

Thirty tablets were collected from each formulation and weighed individually using a digital balance, and then compared with the theoretical tablet weight (200 mg). According to the United States Pharmacopeia 29 – NF 24, the formulation passes:

Table 1. Modified OGS for tableting formulations of ODTs.

| No. | OGS (%) | Method | Hardness (N) | OGS:water (w/v) | Incubation Temp (°C) | Modified Time (h) | Dried Temp (°C) |
|-----|---------|--------|--------------|-----------------|----------------------|-------------------|-----------------|
| F1  | 5%      | M1     | 40           | 1:50            | -                    | -                 | 60              |
| F2  | 5%      | M1     | 30           | 1:50            | -                    | -                 | 60              |
| F3  | 5%      | M1     | 20           | 1:50            | -                    | -                 | 60              |
| F4  | 10%     | M1     | 20           | 1:50            | -                    | -                 | 60              |
| F5  | 5%      | M2     | 40           | 1:50            | -                    | -                 | 60              |
| F6  | 5%      | M2     | 30           | 1:50            | -                    | -                 | 60              |
| F7  | 5%      | M2     | 20           | 1:50            | -                    | -                 | 60              |
| F8  | 10%     | M2     | 20           | 1:50            | -                    | -                 | 60              |
| F9  | 15%     | M2     | 20           | 1:50            | -                    | -                 | 60              |
| F10 | 5%      | M3     | 40           | 1:50            | -                    | -                 | 60              |
| F11 | 5%      | M3     | 30           | 1:50            | -                    | -                 | 60              |
| F12 | 5%      | M3     | 20           | 1:50            | -                    | -                 | 60              |
| F13 | 10%     | M3     | 20           | 1:50            | -                    | -                 | 60              |
| F14 | 15%     | M3     | 20           | 1:50            | -                    | -                 | 60              |
| F15 | 20%     | M3     | 20           | 1:50            | -                    | -                 | 60              |
| F16 | 15%     | M3     | 20           | 1:25            | -                    | -                 | 60              |
| F17 | 15%     | M3     | 20           | 1:100           | -                    | -                 | 60              |
| F18 | 15%     | M4     | 20           | 1:50            | 50                   | 3                 | 60              |
| F19 | 15%     | M4     | 20           | 1:50            | 50                   | 6                 | 60              |
| F20 | 15%     | M4     | 20           | 1:50            | 90                   | 3                 | 60              |
| F21 | 15%     | M4     | 20           | 1:50            | 90                   | 6                 | 60              |
• If no more than 2 tablets are outside the 7.5% of tablet weight
• And if no more than 2 times the 7.5% of tablet weight

2.2.3. Friability Test

Thirty tablets were weighed and then, they were placed in a friability tester (25 rpm for 4 min). The tablets were reweighed, and the friability was calculated using the following formula.

\[
\text{Friability} = \frac{\text{Initial weight} - \text{final weight}}{\text{Initial weight}} \times 100\%
\]

2.2.4. In vitro Wetting Study

A filter paper was put in a Petri-dish (diameter of 10 cm). Then, 10 mL of water and methylene blue was added to the Petri dish. A tablet was carefully dropped on the surface of the filter paper. The time when water reaches the upper surface of the tablets was noted as the wetting time [19].

2.2.5. In vitro Disintegration Test

The tablet was carefully put in the center of the Petri dish (10 cm in diameter) containing 10 mL of water. The disintegration was noted at the time for the tablet to disintegrate completely into fine particles [19].

2.2.6. Drug Content

Six tablets were weighed and crushed to a fine powder which was then dissolved in a 50 mL volumetric flask containing distilled water. The sample solutions were diluted with methanol for the HPLC test as described below.

2.2.7. High-Performance Liquid Chromatography (HPLC)

The quantification of acetaminophen in tablets was performed using Ultimate 3000 HPLC (USA) on a C18 (150 x 4.6 mm, 5 µm, Phenomenex, USA) column. The mobile phase containing buffer solution (pH 3.5, adjusted by phosphoric acid) and acetonitrile at the ratio 3:1 was controlled at a flow rate of 1 mL/min. The UV/ VIS detector was set at a wavelength of 207 nm. The sample injection volume was 20 µL.

2.2.8. Dissolution Test

Dissolution tester (DT70 Pharmatest, Germany) was used for dissolution studies. The tests were conducted at 37 ± 0.5°C in a USP specification dissolution test type II apparatus (Paddle apparatus) according to the United States Pharmacopoeia 29 - NF 24. 900 mL of buffer pH 5.8 was added into each dissolution vessel. The apparatus was set up at 50 rpm of rotation speed. 1 mL of sample was collected from the media after 30 min and replaced by an equal volume of the fresh media. 100 µL sample solutions were diluted with 900 µL distilled water for the HPLC test.

3. RESULTS AND DISCUSSION

3.1. Effect of the Hardness and Modification Method on the Disintegrating Time and Wetting Time

Firstly, the ODTs were investigated upon the effect of the hardness on the disintegrating time and wetting time. The concentration of OGS in formulations was fixed at 5% and screened with the three main hardness ranges 40 ± 2 N, 30 ± 2 N, and 20 ± 2 N. Fig. (1) shows the effect of hardness on

![Graph showing the effect of hardness on disintegrating time and wetting time](image)
the disintegrating time and wetting time of ODTs in three modified methods. Generally, the data indicated that the decrease in hardness improves the disintegrating time and wetting time of the tablets. Specifically, in Fig. (1A) which shows the 5 % OGS in formulations of method 1, the disintegrating times at 40 N, 30 N, and 20 N were 290 ± 10 s, 257 ± 3 s, 230 ± 5 s, respectively, and the corresponding wetting times were 336 ± 5 sec, 294 ± 5 sec, and 217 ± 4 s. Although the disintegrating and wetting times were proportional to the hardness of the tablets, they were 3 minutes longer than the acceptable range of the disintegrating and wetting times for common ODTs [20-22]. In contrast, Fig. (1B and 1C) indicated that there was a significant change in the time of wetting and disintegration from more than 5 min at the hardness of 40 N to below 2 min when the hardness decreased to 20 N. Therefore, method 2 and method 3 demonstrated that the hardness had a direct effect on ODTs preparations using OGS as a disintegrant.

Moreover, Table 2 shows the average weight of all formulations. According to the United States Pharmacopeia, the percentage limit is ± 7.5 % due to the average weight of tablets less than 250 mg. There were no more than 2 tablets that were out of ±7.5 % and no more than ± 15 % percentage limit (date not shown). Hence, all formulations were acceptable in the uniformity range of tablets. In addition, Table 2 indicated that all formulations (except formulation F3 which was slightly out of the limitation) showed good mechanical strength because the friability values were less than 1.0 %.

In addition, formulation F3 in method 1 (containing 5 % OGS) had a friability of 1.03 %, which was out of 1% of the limit acceptance for the friability test. Although the increase of OGS up to 10% could increase the mechanical strength in F4 formulation, the GGS without modification in method 1 was not suitable for the preparation of ODT tablet based on the friability, disintegration, and wetting test. In contrast, in method 2 and method 3, with the same concentration of OGS (5 %) the friability of these tablets was observed between 0.1 % and 0.76 %, which was less than 1 %, indicating that tablets had good mechanical resistance [23]. Therefore, the modification of OGS and the hardness were selected for further studies of ODT.

### 3.2. Investigation of OGS Concentration on the Disintegrating Time and Wetting Time

At the hardness of 20 N, method 3 (F12 – F15) had an ability to form tablets with OGS concentrations from 5 % to 20 % while method 2 only formed tablets at 15 %. The

| No. | OGS (%) | Method | Hardness (N ± SD, n=30) | Uniformity (mg ± SD, n=30) | Friability (%) |
|-----|---------|--------|-------------------------|-----------------------------|---------------|
| F1  | 5%      | M1     | 40                      | 205 ± 4                     | 0.34          |
| F2  | 5%      | M1     | 30                      | 196 ± 3.8                   | 0.57          |
| F3  | 5%      | M1     | 20                      | 198.7 ± 4.9                 | 1.03          |
| F4  | 10%     | M1     | 20                      | 191 ± 3.2                   | 0.47          |
| F5  | 5%      | M2     | 40                      | 198.2 ± 3.5                 | 0.1           |
| F6  | 5%      | M2     | 30                      | 193.8 ± 3.1                 | 0.36          |
| F7  | 5%      | M2     | 20                      | 198.2 ± 3.4                 | 0.76          |
| F8  | 10%     | M2     | 20                      | 202.9 ± 4.5                 | 0.23          |
| F9  | 15%     | M2     | 20                      | 190.6 ± 4.2                 | 0.72          |
| F10 | 5%      | M3     | 40                      | 205 ± 5.4                   | 0.48          |
| F11 | 5%      | M3     | 30                      | 190.5 ± 4.6                 | 0.28          |
| F12 | 5%      | M3     | 20                      | 197.3 ± 3.1                 | 0.1           |
| F13 | 10%     | M3     | 20                      | 197 ± 3.2                   | 0.15          |
| F14 | 15%     | M3     | 20                      | 208.6 ± 3.8                 | 0.28          |
| F15 | 20%     | M3     | 20                      | 191.5 ± 3.7                 | 0.4           |
| F16 | 15%     | M3     | 20                      | 193.4 ± 3.1                 | 0.38          |
| F17 | 15%     | M3     | 20                      | 197.2 ± 3.6                 | 0.18          |
| F18 | 15%     | M4     | 20                      | 198 ± 3.4                   | 0.1           |
| F19 | 15%     | M4     | 20                      | 199 ± 4                     | 0.32          |
| F20 | 15%     | M4     | 20                      | 203 ± 6.5                   | 0.31          |
| F21 | 15%     | M4     | 20                      | 195 ± 4.3                   | 0.02          |
maximal concentration of OGS. As shown in Fig. (2) (top), only slightly changed disintegrating time was observed: 98s (5 % OGS), 95s (10 % OGS), and 92s (15 % OGS) with method 2. However, both wetting time and disintegrating time significantly changed with an increase of OGS concentration in method 3. Specifically, the disintegrating times were 100s (5 % OGS), 85s (10 % OGS), 73s (15 % OGS), and 50s (20 % OGS) in method 3 (Fig. 2, bottom). This data demonstrated that the modification of method 3 was more effective than method 2. In method 2, only the shell of OGS was swelled in water, and the unswelled core still remained inside (Fig. 3). In method 3, the powder OGS was swelled and then milled, leading to a complete swelling of all parts of seeds. Therefore, method 3 showed a higher swelling property by increasing water absorption with an increase of OGS concentration. The swelling property of OGS was due to the presence of polysaccharide, the main composition of the mucilage layer of OGS seeds [12]. The fast increased disintegration with increased polysaccharide content was attributed to the swelling of polysaccharide powder, which leads to penetration of water in the pores of tablets and hence, generate a hydrodynamic pressure for the quick and completed disintegration of tablets. In general, the modification of OGS with concentration from 5 % to 20 % using method 3 was ideal for ODTs. Moreover, 15 % and 20 % were the best two concentrations that facilitate the disintegrating and wetting time around 60s.

3.3. Effect of Amount of Water in the Modification Process on Wetting Time and Disintegrating Time

In this investigation, the amount of water for the swelling of OGS during the modification process was evaluated based on the effect on disintegrating time and wetting time. In fact, the formulations were fixed at 15 % OGS for the comparisons because of the ability of tablet formation for all formulations and the shortest disintegrating time and wetting time at this OGS level. Three ratios of OGS/water (w/v) 1:25 (F16), 1:50 (F14), and 1:100 (F17) were examined. In Fig. (4), the investigation clearly showed that the ratio suitable for OGS swelling completely was 1:50 w/v with the time for disintegrating and wetting were 73.4 ± 4 s and 49.7 ± 2 s, respectively. Meanwhile, the disintegrating time and wetting time were 125 ± 3 s and 54 ± 2 s at ratio 1:25 w/v; and 145 ± 2 s and 102 ± 2 s at ratio 1:100, respectively. Hence, less or more than 50 mL of water in the modified process also

Fig. (2). Effect of OGS concentration on disintegrating time and wetting time in method 2 and method 3. (A higher resolution / colour version of this figure is available in the electronic copy of the article).

Fig. (3). Ocimum grassium seeds (a) dry, (b) swollen, (c) magnified. (A higher resolution / colour version of this figure is available in the electronic copy of the article).
affected directly the disintegrating time and wetting time of the tablets, indicating that F14 was still the best formulation for ODTs.

3.4. Effect of the Incubation Processes on the Disintegrating Time and Wetting Time

In a further study on the modification process, temperature and time of the incubation were investigated. Fig. (5) demonstrates that the temperature and time of incubation process also directly affected the time of disintegrating and wetting. Firstly, the incubation temperatures of the swelling process at 50°C and 90°C were investigated and compared to the formulation prepared without incubation (F18, F20 vs. F14). All formulations in this test were fixed at the same concentration of 15 % OGS and the same range of 20 ± 2 N of the hardness. Fig. (5) indicated that the incubation process could lead to an increase in the wetting time and the disintegrating time. Specifically, the disintegrating time and wetting time of the incubation at 50°C were 132 ± 2.5 s and 73 ± 3.5 s, respectively. Similarly, the increase of the temperature to 90°C could also increase the disintegrating time to 148 ± 4.2 s and the wetting time to 92 ± 2.5 s. These values were out of the standard time for ODTs.

Secondly, at each temperature investigation above 50°C and 90°C, the incubation time was increased from 3h to 6h (F18 vs. F19 and F20 vs. F21). As a result, the increase of incubation time up to 6h resulted in 171 ± 3.2 s disintegrating time, 102 ± 3.6 s wetting time at 50 °C (F19) and 183 ± 3.5 s disintegrating time, 109 ± 2.5 s wetting time at 90 °C (F21). The data suggested that the modification of the incubation process might result in changing the ability swelling of OGS in ODT. Therefore, method 3 in the modification of OGS was optimized as an ideal fabrication for ODT.

3.5. Drug Content and Dissolution Studies

Four formulations of method 3 were tested with regards to the drug content and dissolution to ensure that the modification process is proper for the ODT quality. Table 3 shows the percentage drug content of the formulations in method 3 (F12 – F15) was 101.3 %, 103.5 %, 103.4 %, and 98.8 %, respectively. These values were found to be within an acceptable range and confirmed the use of OGS as a disintegrant. Furthermore, the percent of drug release in 30 min of formulations in method 3 (F12 – F15) was more than 80 %, which is acceptable according to the standard for ODTs by USP29/NF 24. Especially, the F15 formulation contained 20 % OGS in formulation had the drug release more than 90 %. Overall, the modified OGS concentration in ODT affected not only the disintegrating time but also the dissolution rate.

Table 3. The percentage of drug content and drug release from ODT using modified OGS (F12–F15).

| No. | Method | Concentration OBL (%) | Drug Content (%) | Drug Release in 30 min (%) |
|-----|--------|------------------------|-----------------|---------------------------|
| F12 | M3     | 5                      | 101.3           | 83 ± 2.5                  |
| F13 | M3     | 10                     | 103.5           | 86 ± 2.3                  |
| F14 | M3     | 15                     | 103.4           | 88 ± 3                    |
| F15 | M3     | 20                     | 98.8            | 94 ± 1.4                  |
CONCLUSION

This study successfully modified the OGS for ODTs. Method 3, in which OGS was swelled and milled during the modification process, demonstrated that OGS was an effective disintegrant in an ODT formulation. Of note, formulations containing 15% OGS and 20% OGS with the hardness 20 ± 2 N showed a rapid wetting time and disintegration time. The present work revealed that the modified OGS is a potential candidate for use as a disintegrant in the formulation of ODTs. Furthermore, taking the advantages of cost-effectiveness, non-toxicity, compatibility, and easy to manufacture, the modified OGS would be a promising material in ODT formulations as compared to synthetic disintegrants. Further in vivo studies of modified OGS will evaluate its application as a disintegrant in ODT tablets.

ETHICS APPROVAL AND CONSENT TO PARTICIPATE

Not applicable.

HUMAN AND ANIMAL RIGHTS

No Animals/Humans were used for studies that are base of this research.

CONSENT FOR PUBLICATION

Not applicable.

AVAILABILITY OF DATA AND MATERIALS

Not applicable.

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CONFLICT OF INTEREST

The authors declare no conflict of interest, financial or otherwise.

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