Stability analysis of primary emulsion using a new emulsifying agent gum odina

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ABSTRACT

Gum odina and various parts of the plant Odina wodier are traditionally used in Indian folk medicine. Here an effort was made to investigate the efficacy of gum odina as new pharmaceutical excipients, in particular, as an emulsifying agent. Primary emulsion was prepared using wet gum method taking oil: water: gum (4:2:1) with gum acacia powder as an emulsifying agent. This was used as a standard control formulation. In case of experimental emulsions the primary emulsion was prepared by same wet gum technique taking oil: water: gum (4:2:0.5) (gum content was just a half of gum acacia) by using gum odina powder as an emulsifier. The gum odina as emulsifying agent provided a stable emulsion at a very low concentration as compared to the amount required for other conventional natural emulsifying agents. Stability studies of the emulsion were made as per the ICH guideline to study thermal stability, photosensitivity, pH related stability and stability in presence of oxygen. The emulsion type was identified by staining techniques (dye test by using Sudan III) as o/w type preparation without creaming or cracking even after long storage for 24 months at 25°C. It was found that the emulsion containing gum odina produced more stable emulsion at a much lower amount as compared to the emulsion stabilized by gum acacia.

Keywords: Emulsifying Agent, Gum Odina, Odina wodier

1. INTRODUCTION

Use of various gums as pharmaceutical excipients is nothing new. As a stabilizer and thickening agent, use of natural gum has been found in the literature about five thousand years back [1]. Some natural or induced-exudation of normally neutral or slightly acidic complex of polysaccharides or partially acetylated polysaccharide or heterogeneous polysaccharide are obtained as a mixture with calcium, potassium and magnesium salts [2-3]. As a natural defense mechanism to prevent infection or dehydration many trees and shrubs are known to produce an aqueous thick exudation when the plants bark is injured [4]. Eventually the solution dries up in contact with sunlight and air and a hard transparent brown-tint glass like mass is formed. This solid exudation is commonly known as natural gum [4-5]. Some of the gums used frequently now-a-days as pharmaceutical excipients and/or in food industry are gum acacia, gum tragacanth, gum Karaya etc. Gum acacia is mainly used in the confectionary industry. Traditionally it is used in candies to provide the appropriate texture so that they do not adhere to the teeth. Gum acacia is used in chewing gum as a coating agent [6-7] and is also used as emulsifier in soft drink industries [8]. Pharmaceutically gum acacia is still used as a suspending agent, emulsifier, adhesive and tablet binding agent [9-11]. In cosmetic industry it is used as a stabilizer in lotions and protective creams, where it increases viscosity, imparts spreading properties and maintains a protective coating [4].

Gum tragacanth is used in ice creams to provide texture to the product [12] and acts as a thickener and provides texture for chewy sweets such as lozenges [13]. Gum tragacanth is widely used in pharmaceutical industry as an effective suspending agent. Gum tragacanth is used as a stabilizer in dermatological creams and lotions and it also provides a protective coating [14-15]. Suspending properties are used in jellies and tooth paste giving spreadability and a shiny creamy appearance [16-17].

Gum Karaya is well-studied for stabilizing low pH emulsion such as sauces [18]. Due to the water binding capacity of Gum karaya it extends the shelf-life of baked goods. It is widely used as stabilizer, thickener, texturiser and emulsifier in foods. Powdered Gum karaya is widely applied on dental plates as an adhesive [19]. It is used as a bulk laxative, and also used as an adhesive in leak-
proof sealing rings for post surgical drainage pouches or osotomy bags and in skin lotions [20-22].

In recent past we described the use of gum odina (Figure 1(a)) as an excellent substitute of starch paste as a tablet binder [23]. Odina wodier, Roxb. family Anacardiaceae is a large tall tree (Figure 1(b)) found in deciduous forest in India, Myanmar, Sri Lanka, China, Malaysia, Cambodia and Philippine Islands [24]. It is popularly known as Kashmala, Odimaram, Jiol in local language and in English it is called Rhus olina [25]. Various parts of this plant have been found to be used as medicines in Ayurveda. The leaves have been reported to use in Elephantiasis of the legs [25]. Juice of green branches is used as an emetic in case of coma or insensibility produced by narcotic. The dried and powdered bark is found to use as tooth powder by poor villagers [24]. The bark extract has been reported to be useful in vaginal trouble, curing ulcer, heart diseases etc. [26].

In the present study we investigated and compared the emulsifying property of the gum odina (obtained from Odina wodier, Roxb. Family Anacardiaceae) with respect to that of a well-known natural gum emulsifier (gum acacia) and the stability aspects of emulsion prepared with the gum.

1.1. Materials and Methods

Chemicals procured for preparing emulsion were cod liver oil (E. Merck Ltd, Mumbai, India) and acacia powder (E. Merck Ltd, Mumbai, India). All other chemicals were of analytical grade and used as received if not otherwise mentioned.

1.2. Collection of Gum Odina

Gum was collected from the tree Odina wodier, Roxb., family Anacardiaceae during Autumn in the month of August from the Mandal Ghat of Jalpaiguri, West Bengal, India. The gum was the natural exudates on the bark of the tree. It was collected in a dry condition. After collection of the gum, the entire work was carried out in the Department of Pharmaceutical Technology, Jadavpur University.

Figure 1. (a) Transparent reddish brown needle shape gum liberating from the bark of the plant; (b) Tree of Odina wodier, Roxb., family Anacardiaceae.
2. FORMULATION DEVELOPMENT

2.1. Formula for Preparation of Primary Emulsion

Formulation was developed by conventional “wet gum” technique [27]. Formula for primary emulsion was prepared using “wet gum” method taking oil: water: gum (4:2:1) with gum acacia powder as an emulsifying agent. This was used as standard control formulation. In case of experimental emulsions (test sample) the primary emulsion was prepared by the same “wet gum” technique taking oil: water: gum (4:2:0.5) (gum content is just a half of gum acacia) by using gum odina powder as an emulsifier (Table 1).

2.2. Procedure for the Preparation of Emulsion

3.75 gm of experimental gum (gum odina) was taken in a mortar and thick mucilage was prepared by taking 15 ml of water using a pestle. Then to it, required volume (30 ml) of cod liver oil was added drop-wise with constant and uniform clockwise trituration to make a primary emulsion [27]. Final volume was adjusted to 90 ml with water (Table 1).

2.3. Stability Study of Emulsion

2.3.1. FTIR Study

IR grade KBr with a drop of respective emulsion was compressed into pellets by applying 5.5 metric tons of pressure in a hydraulic press and scanned over a wave number of 4000 cm⁻¹ - 400 cm⁻¹ in a FTIR spectrophotometer 8400S Shimadzu.

2.3.2. Thermal Stability

Prepared emulsions were kept (test and control) at different temperatures namely 20°C, 40°C and 60°C for one month by following ICH guideline [28]. Samples were taken out and FTIR spectroscopy was done.

2.3.3. Stability at variable pH

Initial pH of the prepared emulsion was 4.75. To determine the stability at different pH values, the emulsion were adjusted at different pH conditions namely 2, 7.4 and 10 by using 0.1(N) HCl and 0.1(N) NaOH as applicable and kept for one month both for test and control. Then the FTIR spectroscopic studies of the sample were done.

<table>
<thead>
<tr>
<th>Control</th>
<th>Experimental</th>
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<tbody>
<tr>
<td>Cod-liver oil-30 ml</td>
<td>Cod-liver oil-30 ml</td>
</tr>
<tr>
<td>Water q.s - 90 ml</td>
<td>Water q.s - 90 ml</td>
</tr>
<tr>
<td>Acacia powder-7.5 g</td>
<td>Gum odina powder - 3.75 g</td>
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</table>

2.3.4. Photo-Stability

To determine the photo-stability, the formulations (the test and control samples) were exposed to 40 watt (2216.16 CP), 60 watt (3656.66 CP) and 100 watt (5540.4 CP) using electric bulbs. That was adjusted 6 inches above the formulations kept in transparent glass bottle capped tightly in a closed chamber for one month. Following this study FTIR spectra were determined and compared with the samples not exposed to light (i.e. kept in a dark place at 4°C) [29].

2.3.5. Oxygenation of Emulsion and FTIR Study

To analyze the susceptibility of the prepared emulsion containing gum to oxidation, 30 ml of emulsion in a glass bottle of 50 ml capacity was continuously exposed to a stream of O₂ (5 L/min) for 1h and the samples were capped tightly and kept for 15 days before being analyzed by studying their FTIR spectra.

30 ml of emulsion in a glass bottle of 50 ml capacity was continuously exposed to inert environment by using a stream of N₂ (5 L/min) (considered as control against oxidation) for 1 h and the samples were kept for 15 days before being analyzed by studying their FTIR spectra.

3. CHARACTERIZATION OF EMULSION

3.1. Viscosity of Emulsion

Dynamic viscosity of the prepared emulsion was measured using Brook-field rotational viscometer TV-10 (Toki Sangyo Co. Pvt.Ltd, Tokyo, Japan) rotated at 60 rpm for one minute. The length and diameter of the cylinders were 10.5 cm and 3 cm respectively. Length and diameter of the spindles were 6.4 cm and 1.8 cm respectively.

3.2. Test for Identifying Emulsion Type (Dye Test)

Several tests are available for distinguishing between o/w and w/o type emulsions. They include tests of miscibility, dye test, electrical conductivity measurements etc. We adopted for dye test here.

3.3. Dye Test

Prepared emulsion (10 ml) was triturated with Sudan III (0.05 g) and a drop of it was placed on a microscope slide, covered with a cover-slip and examined under a microscope.

3.4. Cracking of Emulsion

This involves coalescence of the dispersed globules and separation of the disperse phase as a separate layer. Redispersion cannot be achieved by shaking and the ad-
advantages of emulsification are lost and accurate dosage is impossible. Simple visual observation (of the stored samples about 24 months) was the means to detect cracking.

3.5. Creaming of Emulsion
Creaming may be defined as the formation of a layer of relatively concentrated emulsion and this conditions favours breakdown of the interface and consequent coalescence of the oil globules and therefore, the emulsion may eventually crack. By shaking, creaming may disappear in many cases. Simple visual observation technique (of the stored sample about 24 months) was the method adopted here to determine creaming.

4. RESULTS
The various parts of *Odina wodier* have been used in Ayurveda and traditional Indian folk medicine (24). We have recently reported the gum of this plant as a tablet binder, effective at a much lower concentration as compared to the other available natural binders and further, the gum is devoid of toxicity [23]. In the present study we have mainly focused on the utility of the gum as an emulsifying agent of natural origin and the stability aspects of emulsions prepared using this emulsifying agent.

The dynamic viscosity of prepared emulsion (4:2:0.5) was measured using Bookfield type rotational viscometer TV-10, rotated at 60 rpm for one minute and the viscosity was 14 centipoises.

Several tests are available for the differentiation of types of primary emulsions *i.e.* o/w and w/o type emulsions. These tests are miscibility with water, Dye test and Electrical conductivity measurements etc. [27]. Dye test is a very common test to determine the types of emulsion. The dispersed globules were appeared ‘red’ due to oil soluble dye Sudan III and the continuous phase was ‘colourless’ (*Figure 2*) in the present study.

Stability of emulsion was analyzed by comparing the FTIR spectra of the freshly prepared experimental emulsion (*Figure 3*) and the stored (24 months) experimental emulsion (*Figure 4*). Physical interactions were detected between wave number 700 cm\(^{-1}\) and 600 cm\(^{-1}\) upon prolonged storage as compared to the freshly prepared samples.

*Figure 2.* Determination of o/w type of emulsion *i.e.* dispersed oil globules appeared ‘red’ and continuous phase ‘colourless’.
Figure 3. FTIR spectra of freshly prepared experimental emulsion.

Figure 4. FTIR spectra of stored (24 months) experimental emulsion kept at room temperature.
Thermal stability of emulsion was analyzed by studying the FTIR spectra of the emulsions stored at 20°C, 40°C and 60°C for 30 days (Figures 5-7). Interactions were detected in the wave numbers between 2700 cm⁻¹ and 720 cm⁻¹ for the samples stored at 40°C and 20°C; and wave number at 3461 cm⁻¹, 2099 cm⁻¹, 1646 cm⁻¹, 718 cm⁻¹ in case of the sample stored at 60°C (Figures 7-9). The types of interaction have been discussed in details in discussion section.

The impacts of variable pH on emulsion stability were detected by changing the pH of emulsion at 2, 7.4 and 10; these were stored for 30 days at room temperature. This was followed by the FTIR spectroscopy and the data indicate that there were variations in wave numbers in the range between 3500 cm⁻¹ and 2600 cm⁻¹ and also at 1744 cm⁻¹ at pH 7.4 (Figures 8-10).

For studying the photostability, the emulsion was exposed to 60 Watt (3656.66 CP) for 30 days and the FTIR spectra were compared with the experimental emulsion stored in the dark for the same period. There were physical interactions detected in the range of wave numbers between 3600 cm⁻¹ and 2800 cm⁻¹ (Figure 3 and Figure 11) and also in the range between 2400 cm⁻¹ and 1700 cm⁻¹. Otherwise no predominant variations in the FTIR spectra were detected when photo-exposed samples were compared.

To know the stability of emulsion exposed to oxidation, samples were exposed either to oxygen or nitrogen (which was used as control) as specified earlier. No interactions were detected upon oxygenation except some minor peak variation at the wave numbers 2665 cm⁻¹, 3457 cm⁻¹, 1437 cm⁻¹, 1148 cm⁻¹ (Figures 12 and 13). However, all the characteristic peaks of the gum were present.

Creaming and cracking of the stored emulsions (24 months) were also visually observed but no such phenomena were detected.
Figure 6. FTIR spectra of thermal stability of emulsion at 40°C.

Figure 7. FTIR spectra of thermal stability of emulsion at 60°C.
Figure 8. FTIR spectra of emulsion with pH 2.

Figure 9. FTIR spectra of emulsion with pH 7.4.
Figure 10. FTIR spectra of emulsion with pH 10.

Figure 11. FTIR spectra of photo stability of emulsion at 60 Watt. (3656.66 CP).
Figure 12. FTIR spectra of oxygenated emulsion.

Figure 13. FTIR spectra of nitrogenated emulsion.
5. DISCUSSION

In this study capability of gum odina as an emulsifying agent was investigated. Viscosity of the experimental emulsion was found to be 14 cP which suggests that it was a thicker emulsion and the emulsion would remain stable for a longer period.

Dye test with Sudan III (oil soluble dye) showed that dye was distributed in the form of droplets throughout the colourless continuous phase. This proves that oil formed the dispersed phase and water was the continuous phase and it was an o/w type of emulsion. Thus o/w emulsion may be prepared using gum odina as emulsifying agent.

When the spectra were compared, in some cases peak height varied. It may be due to the presence of variable amounts of ingredients present in the pellet. There were interactions detected in the wave range numbers between 700 cm\(^{-1}\) and 600 cm\(^{-1}\). This zone is the known stretching vibration zone of CH-alkane and OH (H bonded and normally out of plane). Hydrogen of the fatty acid might have formed weak hydrogen bond or bond due to Van der Waal force or dipole moment with OH- group of water predominantly upon the long storage.

In the cases of thermal stability analysis, samples were kept at 20°C, 40°C and 60°C for 1 month (Figures 5-7) and compared with the freshly prepared sample (Figure 3). Changes in peaks in wave numbers between 2700 cm\(^{-1}\) and 720 cm\(^{-1}\) may be possibly due to interaction between ketonic and aldehyde groups in the fatty acids by formation of H-bonding or weak bondings such as Van der Waal forces or dipole moments, since the zone between 2690 cm\(^{-1}\) and 2840 cm\(^{-1}\) are the medium intensity and strong intensity carbomile stretching vibration zone. The zone predominant functional groups were OH and C=O. Peak variations at 2665 cm\(^{-1}\) and 3457 cm\(^{-1}\) of samples exposed to oxygen were probably due to the possible weak bond formation between OH and COOH of water and fatty acid respectively, since these are the known stretching vibration zones of OH and C=O [30]. Peak variations at 1437 cm\(^{-1}\) and 1148 cm\(^{-1}\) could be due to α-CH\(_2\) bending or C-C-C bending of fatty acid carbons, since, 1400 cm\(^{-1}\) - 1450 cm\(^{-1}\) are the bending vibration zone of α-CH\(_2\) and 1148 cm\(^{-1}\) is the medium intensity C-C bending vibration zone [30]. Thus, upon FTIR spectrum analysis, it may be stated that the emulsion is not susceptible to oxidation, since the reactions were due to physical bond formations. However, oxygenation might have a role to induce such bond formations as they were not noticed in case of the samples exposed to N\(_2\).

Cracking may be caused by any chemical, physical or biological effect that changes the nature of the interfacial film that exists between oil and water [31]. These tend to make it less stable. But here, after long storage of prepared emulsion for 24 months, no coalescence of dispersed globules of oil was noticed. Hence, no cracking was observed in the said period.

Creams may be formed as a layer of relatively concentrated emulsion and this condition favors breakdown of the interface and consequent coalescence of the oil globules and therefore, the emulsion may eventually crack [31]. After a long storage of the emulsion for 24 months there were no cream formations on the upper surface of emulsion.

When experimental emulsions were compared with the prepared acacia emulsion (considered here as control), it was found that requirement of gum odina was 50% of the amount of acacia required for preparation of primary emulsion. Further, gum odina produces a stable emulsion which can be stored at least for 2 years.

Thus, gum odina may be used as an emulsifying agent to prepare o/w primary emulsion.

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REFERENCES


