INTRODUCTION

Medicated chewing is a new drug delivery system for the prevention and treatment of chemotherapy-induced nausea and vomiting (CINV), radiation therapy-induced nausea and vomiting (RINV), and also in post-operative nausea and vomiting (PONV) conditions. Drugs are usually formulated in variety of dosage forms like tablets, capsules, injections, inhalers and ointments etc. The oral drug delivery system is most acceptable route of drug administration due to ease of administration than other dosage forms. In addition to confectionary role, nowadays, chewing gum is also showing best and convenient drug delivery system due to rapid absorption of agents which can be absorbed by oral cavity. Medicated chewing gums are more accepted by the parents for children as comparison to tablets, capsules, or liquids. Medicated chewing gum offers a wide range of advantages that make it an excellent alternative [11-14]. Ondansetron hydrochloride (HCl) is the drug of choice to prevent nausea and vomiting giving rise to a systemic effect. Medicated chewing gum offers a wide range of advantages that make it an excellent alternative [11-14]. Ondansetron hydrochloride (HCl) is the drug of choice to prevent nausea and vomiting giving rise to a systemic effect. Medicated chewing gum offers a wide range of advantages that make it an excellent alternative [11-14]. Ondansetron hydrochloride (HCl) is the drug of choice to prevent nausea and vomiting giving rise to a systemic effect. Medicated chewing gum offers a wide range of advantages that make it an excellent alternative [11-14]. Ondansetron hydrochloride (HCl) is the drug of choice to prevent nausea and vomiting giving rise to a systemic effect. Medicated chewing gum offers a wide range of advantages that make it an excellent alternative [11-14]. Ondansetron hydrochloride (HCl) is the drug of choice to prevent nausea and vomiting giving rise to a systemic effect.

Oil-holding capacity (OHC) is a competitive serotonin type 3 receptor that may cause by cancer surgery, chemotherapy, and radiation treatment.

Medicated chewing gum can be prepared (1) fusion method, (2) cooling, grinding, and tableting method, and (3) direct compression [8-10].

Formulation and Evaluation of New Medicated Chewing Gum for the Treatment of Nausea and Vomiting Induced by Chemotherapy, Radiation Therapy, and Post-Operative Conditions in Cancer

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ABSTRACT

Objective: The objective of the present study was formulation and evaluation of new medicated chewing gum for the treatment of nausea and vomiting induced by chemotherapy, radiation therapy, and post-operative conditions in cancer using ondansetron hydrochloride as drug candidate.

Methods: The medicated chewing gum of ondansetron hydrochloride (OHC) was prepared by direct compression mold method. Four formulations were selected for study which was showing good physicochemical properties and drug release. Formulations were characterized for physical evaluation, weight variation, stickiness, hardness/plasticity, in vitro drug release, estimation of chewing gum consistency and drug release study in saliva.

Results: All the formulations gave satisfactory results in physical evaluation, weight variation, stickiness, and hardness. The formulation medicated chewing gum II (MCG II) showed best in vitro drug release which is 97% in 30 min and it is also more accepted by the people. Drug release in saliva also indicated that more than 50% drug release occurs within 15 min.

Conclusion: Medicated chewing gum is a cost-effective product and also showed better compliance and increase in bioavailability.

Keywords: Ondansetron hydrochloride (OHC), MCGs, Chemotherapy-induced nausea and vomiting, Radiation therapy-induced nausea and vomiting, Post-operative nausea and vomiting, Cancer, Chewing gum.

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Medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained. According to the European Pharmacopoeia, 4th edition, 2002 medicated chewing gum is intended to be chewed but not swallowed, providing a slow steady release of the medicine contained.

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| S. No. | Ingredients/quantity | MCG I | MCG II | MCG III | MCG IV |
|-------|---------------------|-------|--------|---------|--------|
| 1.    | Ondansetron HCl     | 4.0 mg| 4.0 mg | 4.0 mg  | 4.0 mg |
| 2.    | Ascorbic acid       | 0.2 g | 0.2 g  | 0.2 g   | 0.2 g  |
| 3.    | Beeswax             | 1.0 g | 1.0 g  | 1.0 g   | 1.0 g  |
| 4.    | PVP                 | 4.5 g | 5.0 g  | 5.5 g   | 6.0 g  |
| 5.    | Dextrose            | 0.7 g | 0.8 g  | 0.9 g   | 1.0 g  |
| 6.    | Peppermint oil      | 0.5 ml| 0.5 ml | 0.5 ml  | 0.5 ml |
| 7.    | PEG-400             | 0.8 g | 1.0 g  | 1.2 g   | 1.2 g  |
| 8.    | Calcium carbonate   | 0.5 g | 0.5 g  | 0.5 g   | 0.5 g  |

PEG: Polyethylene glycol, HCl: Hydrochloride, PVP: Polyvinylpyrrolidone
antagonists and effective in the treatment of nausea and vomiting caused by cytotoxic agents. OHC blocks the actions of chemicals in the body that can also trigger nausea and vomiting. Having been developed in the 1980s by GlaxoSmithKline and approved by the USFDA since January 1991, OHC has demonstrated a long history of use and efficacy. Commonly formulated as oral tablets, orally disintegrating tablets, and injections and available as generic products as well, OHC continues to see contemporary innovations in its formulation and uses [15-19].

**MATERIALS AND METHODS**

**Materials**

Ondansetron HCl was a gift sample. Excipients polyvinylpyrrolidone (PVP), beeswax, dextrose, calcium carbonate, peppermint, ascorbic acid, and polyethylene glycol 400 (PEG-400) were of analytical or pharmaceutical grade were purchased.

**Method of preparation**

MCGs were prepared by direct compression mold method. In this method, each ingredient was weighed accurately and separately. OHC, PVP, beeswax, dextrose, calcium carbonate, peppermint oil, and ascorbic acid, all ingredients were thoroughly mixed in ascending order of their weights in a mortar. After proper mixing, ingredients smoothly grounded in a mortar pestle and then previously weighed quantity of PEG-400 were added. Then, the whole mixture was again mixed thoroughly in pestle mortar. After mixing and grinding, the mixture was subjected for compression into the desired molds and presses to form medicated chewing gum, as shown in Figs. 1 and 2a-c. After removing from mold, formulated chewing gums were weighed and wrapped properly. Optimization of various selected batches of MCGs by changing concentration of different excipients is depicted in Table 1.

**Physical evaluation**

Physical properties such as size, shape, thickness, color, and odor must be evaluated. It is very important to investigate the MCGs physically, which plays a key role and these should not be disregarded, it is necessary for acceptance by individuals and even also in marketing [1].

**Weight variation test**

Weight variation plays an important role in evaluation parameters, it ensures that each of the medicated chewing gum contains the proper amount of drug. The test was carried out by weighing the 20 mediated chewing gum individually using analytical balance, then calculating the average weight, and comparing the individual medicated chewing gum to the average [1].

**Stickiness**

On plain surface, medicated chewing gum was placed, it is subjected to collide with Teflon hammer with mass of 250 g for a period of 10 min. Hammering frequency was 30/min. After specified time, amount of mass stick to hammer was observed and reported [1].

**Test for hardness/plasticity**

There is no one reported method for the determination of hardness; hence, it was decided to use Pfizer type hardness tester for the determination of hardness/plasticity of all MCG formulations [1].

**In vitro drug release studies**

A medicated chewing gum was immersed in pH solution maintained at pH 6.8 (buccal cavity) then placed on magnetic stirrer and subjected to stirring and after every 5 min interval 2 ml of solution taken out and replaced with fresh buffer solution. Sample was withdrawn at regular intervals of 5, 10, 15, 20, 25, and 30 min. On completion of process, all the collected samples were UV spectrophotometrically analyzed on maximum wavelength of 310 nm.
Table 5: Cumulative percentage drug release of MCG II formulation in saliva

| S. No. | Time (min) | % drug release |
|--------|------------|----------------|
|        | Volunteer A | Volunteer B | Volunteer C | Volunteer D | Volunteer E | Volunteer F |
| 1.     | 0          | 0            | 0           | 0           | 0           | 0           |
| 2.     | 0.5        | 31.2±1.45    | 31.5±2.01   | 32.5±1.46   | 32.7±2.08   | 32.1±1.93   | 32.1±1.74   |
| 3.     | 1          | 34.9±1.94    | 35.6±1.23   | 36.6±1.84   | 35.2±1.67   | 35.8±1.07   | 34.1±1.28   |
| 4.     | 2          | 41.9±1.67    | 39.5±2.23   | 41.6±1.69   | 41.5±2.05   | 40.2±2.12   | 42.0±1.90   |
| 5.     | 5          | 43.4±2.31    | 44.2±1.40   | 42.7±1.98   | 43.3±1.87   | 43.4±1.06   | 44.1±1.07   |
| 6.     | 10         | 47.3±1.68    | 48.3±1.58   | 47.2±1.56   | 47.9±1.52   | 40.1±1.61   | 49.0±1.83   |
| 7.     | 15         | 52.4±1.23    | 52.6±1.62   | 53.2±1.73   | 52.1±1.64   | 53.6±1.37   | 52.5±1.46   |

Fig. 3: (a) In vitro drug release study. (b) Ultraviolet spectrophotometric analysis of samples

Fig. 4: Cumulative percentage drug release of MCG II formulation in saliva

Estimation of chewing gum consistency

This study was carried out using chew out method. Dummy chewing (without drug) was prepared according to optimized formula to check consistency. Then, they were given for certain human volunteer to chew it [11].

Drug release study in saliva

Medicated chewing gums are having quite different drug release process compared to conventional oral drug delivery system. In Medicated chewing gums not only dosage form but also chewing activity of patient may also affects the drug delivery. Mechanical treatment is required to deliver the drug by the teeth but not involve in dissolution. Release of drug from MCGs in saliva was studied by recruiting a panel of six members of volunteers and designed chew out studies. One sample was given to each volunteer for chewing for a particular time interval period, i.e., 0.5, 1, 2, 5, 10, and 15 min. After chewing, chewed out chewing gum samples collected from volunteer, stretched maximum, and cut into small pieces after that dispersed in a 100 ml volumetric flask having phosphate buffer pH 6.8, which was then heated and sonicated for 10 min. These samples were then analyzed by UV spectrophotometer at absorption maxima 310 nm for residual drug content in MCGs [1,2].

Amount of drug release during mastication = The total drug content – Residual drug after chewing

RESULTS AND DISCUSSION

In the present work, an attempt was made to develop medicated chewing gum containing OHC drug. Formulation MCG II was selected as optimized formulation for making medicated chewing gums, where ascorbic acid was used as antioxidant, dextrose was used as sweetening agent as well as bulking agent, beeswax/PVP as elastomer and gum, peppermint oil was used as flavoring agent. Optimized formulation was physically evaluated having weight 5.5 g, shape was cubic having thickness 1.4 cm with whitish-cream color and peppermint odor. Weight variations study revealed that all formulations are having weights in normal range. All formulations showed negligible stickiness and hardness also found within range, as shown in Table 2.

In vitro drug release, it was found that almost all formulations release more than 97% drug after 30 min. However, the optimized formulation MCG II has shown the best release as comparison to other formulations. Hence, it was selected for further study. Fig. 3a and b. The prepared formulations were analyzed for drug content and it was found that MCG II was having highest drug content, i.e., 97.01%, as shown in Table 3. MCGs were studied for consistency on human volunteer and MCG II was more accepted by human volunteers data which are tabulated in Table 4.

Drug release study in saliva is shown in Fig. 4, cumulative percentage drug release in each individual is also given in Table 5, it was observed that within 15 min, more than 50% of drug was released from the optimized formulation MCG II. This study revealed that the drug release was depended on the chewing frequency of the volunteer.

CONCLUSION

Medicated chewing gum OHC was successfully prepared. OHC is a potent and highly selective 5-HT3 receptor antagonist having important antiemetic activity and good tolerability. It is a cost-effective formulation and having better patient compliance and bioavailability. OHC is completely absorbed by GIT which makes this as a choice of drug in preparing medicated chewing gum of OHC. Due to hepatic first-pass metabolism, only 60% OHC is bioavailable compared with IV route so this can also be avoided by preparing MCGs.

In vitro drug release study indicated that drug release from formulation MCG II is best among others so it is the best formulation than others. All the parameters found to be satisfactory; hence, the therapeutic dose of OHC can be given in medicated chewing gum with optimized formula, i.e., MCGII. This study concluded that it is possible to make medicated chewing of OHC for prevention and treatment of vomiting induced by CINV, RINV, and also in PONV conditions.

AUTHORS’ CONTRIBUTIONS

Contribution of all authors is equal in this work.

CONFLICTS OF INTEREST

The authors declare that there are no conflicts of interest regarding the publication of paper.
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Nil.

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