Review Article

Chewing Gums as a Drug Delivery Approach for Oral Health

Morteza Banakar,1,2,3 Sedigheh Moayedi,4 Erfan Shamsoddin,5 Zahra Vahedi,6 Mohammad Hasan Banakar,7 Seyyed Mojtaba Mousavi,8 Dinesh Rokaya,9 and Kamran Bagheri Lankarani1

1 Dental Research Center, Dentistry Research Institute, Tehran University of Medical Sciences, Tehran, Iran
2 Health Policy Research Center, Institute of Health, Shiraz University of Medical Sciences, Shiraz, Iran
3 Department of Pediatric Dentistry, Faculty of Dentistry, Shahed University, Tehran, Iran
4 Department of Orthodontics, Mashhad University of Medical Sciences, School of Dentistry, Mashhad, Iran
5 Cochrane Iran Associate Centre, National Institute for Medical Research Development (NIMAD), Tehran, Iran
6 School of Dentistry, Islamic Azad University Tehran Medical Sciences, Tehran, Iran
7 School of Dentistry, Yasuj University of Medical Sciences, Yasuj, Iran
8 Department of Chemical Engineering, National Taiwan University of Science and Technology, Taipei, Taiwan
9 Department of Clinical Dentistry, Walailak University International College of Dentistry, Walailak University, Bangkok 10400, Thailand

Correspondence should be addressed to Dinesh Rokaya; dinesh.ro@wu.ac.th

Received 28 February 2022; Revised 19 May 2022; Accepted 26 May 2022; Published 20 June 2022

Academic Editor: Andrea Scribante

Copyright © 2022 Morteza Banakar et al. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Background. Drug delivery approaches with the shortest therapeutic period and the lowest side effects have always been considered a sublime target in the medical sciences. Among many delivery methods, chewing gum could be perceived as a promising drug carrier that can carry several types of drugs for oral health. These drug carriers could represent optimal therapeutic time and lower side effects due to their sustained release capability and lower required thresholds for the drug compared with other delivery approaches. The convenient use in the oral cavity’s local environment and the ability to locally carry multiple drugs are considered the main advantages of this delivery approach. Aim. This review aimed to explore chewing gum as a promising drug carrier that can carry several types of drugs for oral health. Materials and Methods. Articles were searched for on PubMed, ISI, SCOPUS, Google Patents, the Royal Society of Chemistry website, and electronic databases using MESH terms and the following keywords: (“Gum” OR “Chewing gum”) and (“Drug delivery OR Drug delivery systems”) in the English language. No time limit was applied, and all documents as of August 30th, 2020 were retrieved. Results. Gum-drug interactions, mechanisms of release, and formulations of the drugs might all play a role in this versatile delivery method. Accordingly, chewing gum-based carriers may be presented as a plausible candidate for drug delivery in oral diseases. Conclusion. Gum-driven drugs could be introduced as promising candidates for treating oral diseases due to their ability to deliver the proper local dosages of active ingredients, short contact time, biocompatibility, and biodegradable chemical structures.

1. Introduction

The field of drug delivery is one of the most fascinating ones in medicine. Numerous medication delivery methods have been developed, sharing several advantages. The delivery process could make many advantages for patients, from the low period of therapy to less side effects due to their low usage dose [1]. Typically, the delivery approaches the target drug-loaded carriers and the routes for drug transmission. In recent decades, many families of carriers and their routes in the human body have been appropriately delineated and their advantages and disadvantages have been illuminated [2–4]. The polymeric and nanomaterial-based carriers with appropriate biocompatibility [5] and drug loading efficiency have attracted much attention for in vivo delivery in various cancer types. Noticeably, their toxicity is an issue that has
remained in debate for many years. This challenge using delivery methods with a safe route is certainly promising [5–7]. Among the routes of administration, the oral types are attractive, mostly because of their comfortable appliances. As a result, approaches based on chewable carriers, such as drug-loaded gums and tablets, show some promise for many oral, esophageal, and GI-related diseases [8, 9]. Considering its unique characteristics, oral-based chewy delivery is highlighted as a promising candidate. Rapid onset of action, facile administration, low side effects, and appropriate local impact on oral diseases are all major factors contributing to this salience [9–11]. In this study, we discussed the characteristics of oral conditions for the effect of medications and the application of chewing gums in drug delivery for oral health. Additionally, medically applicable chewing gum types are further scrutinized in more detail.

2. Materials and Methods

In this narrative review, we searched PubMed, SCOPUS, Google Patents, the Royal Society of Chemistry website, and electronic databases using MESH terms and the following keywords: (“Gum” or “Chewing gum”) and (“Drug delivery” OR “Drug delivery systems”) with a language filter (English). No time limit was applied and all documents by August 30th, 2020 were retrieved. We did not use other filters. All articles and patents that satisfied our selection criteria were retrieved. After omitting the duplicates, we identified 30390 papers. Three independent reviewers assessed the article titles and abstracts, applying eligibility criteria. Articles were omitted if they were deemed irrelevant based on our keyword research. We defined the following criteria for inclusion using the PICO model:

Population: There is no identifiable reference population.
Intervention: Chewing gums are a vehicle for drug delivery in clinical trials, animal studies, and in vitro investigations.
Comparison: Placebo-controlled or intra-individual pre-post comparison.

Outcome: Cavity fighters, antibiotics, antibacterial, antifungal, antiviral, antiplaque, and remineralization are some clinical effects.

References to these articles were also reviewed. Figure 1 illustrates the procedure for conducting a literature search. After removing the duplicates, 26671 papers were obtained, of which only 60 studies made it through the eligibility assessments. One study was excluded after full-text reading and was deemed irrelevant to our inclusion criteria. This study followed the recommendations of SANRA (a scale for the quality assessment of narrative review articles) to ensure internal consistency and proper presentation of the manuscript.

3. Results

3.1. Influencing Parameters in Chewing Gum-Based Drug Delivery

3.1.1. Saliva Flow Rate. The accessibility of drug delivery is an essential factor in designing carriers. The oral cavity potentially provides systemic and local delivery accessibility. Saliva, a complicated multifunctional mixture that can solve the drug and delivery process’s ability from the gum to the oral mucosa cells, acts as an intermediate platform. The saliva flow rate, which has been stimulated by chewing gum, has a positive impact on delivery; for instance, a study showed the beneficial effect of saliva flow on xerostomia [12]. Some reports have considered a plateau phase for saliva secretion rate while is being stimulated by chewing gum [11–13].

3.1.2. Local Effect. Drug molecules released into the mucosal membranes during an equilibrium that occurred in minutes could be absorbed from the oral cavity microenvironment [14]. The buccal epithelium cells with a 20-40-cell thickness and a turnover of two weeks could play a vital role in the delivery process [15]. In the oral cavity, highly vascularized mucus membranes can provide an active drug circulation in jugular veins and act as a suitable drug reservoir [14]. The results showed that clearance is better in sublingual parts than in the labial vestibule. The main reason for these observations is the difference in the anatomy of the oral cavity [16–18]. The extended delivery time in the mouth causes the appropriate drug release rate and the maintenance of drug concentration for better therapy. Additionally, the local effects of therapeutic agents could be altered by the quality of drug distribution in the oral mucosa. The residence time of sublingual tablets showed the best time activity while chewing gum was better than the lozenge [19].

3.1.3. Contact Time. Contact time is a significant determinant of the therapeutic period and side effects of the treatment regime. During chewing, the dissolution of ingredients occurs in the first few minutes of the process. Nevertheless, there is no standard time for chewing in general, but some case studies have suggested a 30-minute-period as a reasonable means of chewing time [20].

Figure 1: Study selection flow diagram.
3.1.4. Ingredients and Formulation. When noticing drug formulation and active ingredients, hydrophobicity, stiffness, chemical structure, and interaction types are essential. The hydrophobicity/hydrophilicity of gum influences the drug-gum interactions and reflects the quality of interactions between the drug and gum structures [21]. The hydrophilicity of carriers in delivery could be mentioned as one of the most important physical properties that potentially control the drug release rate and mechanism. Furthermore, for chewing gums, this parameter could be
3.3. Types of Medicaments That Can Be Used in Chewing Gums. To date, various drugs and substances with various therapeutic properties have been introduced for use in chewing gum, some of which and their therapeutic properties are listed in Table 2. These materials based on the source are divided into synthetic and natural, which are mentioned below.

3.3.1. Synthetic Material. Various types of synthetic gum-driven agents have been suggested so far. The ingredients that fight cavities include Ca phosphate [36–39] and bicarbonate [40,41]. They also include chlorhexidine [42] and copper chlorophyllin [43]. Hydrogen peroxide [44] and zinc [45] are antibacterial and antiviral agents, as are pycnogenol [46], stannous EDTA [47], sulfathiazole [11], urea [48,49], zirconium silicate [50–52], and also fluoride [53–56]. Among these drugs, chlorhexidine, hydrogen peroxide, sulfathiazole, and zinc have more potential to be loaded in chewing gum and as an oral disease remedy [11]. Chlorhexidine was used in nanocapsules to treat dentin substrates that had been decalcified. Gum could be used as a carrier for this drug [57].

3.3.2. Natural Material. Some natural ingredients can be used in chewing gum with medical properties. Garlic, for instance, shows antiviral, antibacterial, and antifungal properties [58]. Ginger can also counteract respiratory viruses [59]. Oregano has powerful antiviral properties. In high concentrations, it could inactivate viral agents within one hour of exposure [60]. Lemon balm and green tea have antiviral properties and effectively against various viruses, including influenza, herpes, adenovirus, and HIV [61, 62]. Elderberry exhibits antiviral and antibacterial effects and is considered a remedy for the common cold in traditional medicine [63]. Coconut oil has also shown strong antiviral properties. It can either eradicate or inactivate harmful bacteria in the body [64]. Black walnut has antiviral, antifungal, antimalarial, and antiparasite properties [65]. Turmeric could inhibit viral replication and interfere with the virus-cell binding process [66].

3.4. Chewing Gums as Drug Carriers for Oral Health. Chewing gum can be used to provide a controlled dose of an active component to the mouth. Chewing gum active compounds are released in a variety of ways, depending on parameters such as chewing speed, gum base concentration, and active ingredient solubility in water, allowing them to remain in the mouth for a longer period of time. Chewing gums could transport chlorhexidine, calcium, and carbamide-based medications such as captopril, nitroglycerin, methadone, antihistamines, and antifungal-based compounds as drug carriers [67–69]. Figure 3 illustrates some of
### Table 1: Carriers that can be used in chewing gum and their properties.

| Carriers             | Origin/components                                           | Properties                                                                 |
|----------------------|------------------------------------------------------------|-----------------------------------------------------------------------------|
| Acacia               | Stems of tree *acacia arabica*                            | Antimicrobial activity                                                      |
| Alginic acid         | Natural polysaccharides isolated from the brown seaweed   | Antianaphylaxis effect, immunomodulatory activity, and antioxidant activity |
| Dextrin              | Produced by the hydrolysis of starch and glycogen         | Applications as a targetable carrier and bioadhesive                        |
| Gelatin              | Linear anionic high molecular weight exopolysaccharide,   | Antibacterial drug delivery systems                                          |
| Guar                 | Biopolymer extracted from the seeds of *Cyanopsis        | Sustained-release systems                                                   |
|                      |     tetragonolobus beans (Leguminosae family)             |                                                                             |
| Lecithin             | The mixture of fats that are essential to cells, derived from | Oral and aerosol delivery systems                                            |
| Sodium alginate      | Brown seaweeds (*Phaeophyceae*)                           | pH-sensitive carrier                                                        |
| Xanthan              | Hydrophilic, anionic-bacterial heteropolysaccharide, derived from | Antioxidant, anti-inflammatory, antibacterial, and biofilm inhibitor         |
|                      | the fermentation of gram-negative bacteria *Xanthomonas   |                                                                             |
|                      |     campestris                                           |                                                                             |
| Gellan               | Exocellular polysaccharide secreted by *Pseudomonas       | Anti-inflammatory                                                           |
|                      |     elodea                                              |                                                                             |
| Rosin                | Clear, pale yellow to dark amber thermoplastic resin present in | Film-forming, coating properties, and sustained and controlled drug release systems |
|                      |     oleoresins of the tree *Pinus roxburghii* and *Pinus  |                                                                             |
|                      |     taeda                                               |                                                                             |
| Chitosan             | Invertebrates, insects, and yeast                        | Antifungal, wound healing acceleration, and immune system stimulation       |
| Tamarind seed        | Galactoxyloglucan, tamarind seed polysaccharide           | Noncancerigenicity, mucoadhesive nature                                     |
| polysaccharide       |                                                           |                                                                             |
| Carrageenan          | Extract from a red seaweed commonly known as Irish moss  | Immunomodulatory and antioxidant activity                                   |
| Terminalia catappa   | *Terminalia catappa* leaves                              | Antimicrobial, antioxidant, anticancer, antiviral, anti-inflammatory properties |
| Pectin               | Methoxyester of pectic acid derived from plant cell walls | Anticancer, immunostimulation, anti-inflammatory, antibacterial, antiadhesive effects |

### Table 2: Therapeutic effects and materials used in chewing gum on oral health.

| Therapeutic effect                              | Material used                                                                 |
|-------------------------------------------------|-------------------------------------------------------------------------------|
| Analgesic                                       | Aspirin, benzocaine, and eugenol                                             |
| Acid neutralization                             | Antacid, calcium carbonate, carbamide, bicarbonate, xylitol                  |
| Antiplaque (biofilm control)                    | Chlorhexidine gluconate, eucalyptus, mastic, xylitol, sorbitol sulfonamides, neomycin, gramicidin, hydrogen peroxide, zinc, sulfa thiazole, magnolia bark extract, fluo ride, and propolis |
| Anticalculus formation                          | Vitamin C and polyphosphates                                                 |
| Antioxidant, antiseptic, and healing            | Green tea and aloe vera                                                      |
| Dental caries prevention                        | Fluoride, calcium phosphate, bicarbonate, copper, chlorophyllin, and xylitol |
| Antibacterial agent                             | Chlorhexidine gluconate, sulfonamides, neomycin, gramicidin, hydrogen peroxide, zinc, sulfathiazole, fluoride, and propolis |
| Antiallergy                                     | Cetirizine, diphenhydramine hydrochloride                                   |
| Gingival inflammation                           | Green tea, amyloglucosidase combined with glucosidase, egg-white lysozyme, and rhozyme P-11 |
| Deficiency of vitamin C                         | Vitamin C                                                                    |
| Plaque removal                                  | Zirconium silicate, decapetide-based antiseptic, and sodium bicarbonate     |
| Dental enamel strengthening agent               | Potassium aluminum sulfate, calcium, CPP-ACP, and fluoride                  |
| Oral candidiasis                                | Miconazole, nystatin                                                        |
| Periodontal disease                             | Sodium bicarbonate/sorbitol                                                  |
| Vincent disease                                 | Metronidazole                                                                |
| Reduction of planktonic bacteria in saliva      | Chlorhexidine, xylitol, chitosan, mastic, magnolia bark extract, and propolis |
| Mitigation in a volatile sulfur compound        | Eucalyptus, zinc, and magnolia bark extract                                  |
| Tooth stain prevention                          | Polyphosphates and hydrogen peroxide                                        |
the uses of chewing gum as a drug delivery approach for oral health.

Mouth dryness, also known as xerostomia, is a condition that occurs when salivary glands do not produce enough saliva to keep the mouth moist. Chewing gum has long been recognized to increase saliva production. The first five minutes of chewing generate a 10-fold increase in salivary flow over unstimulated salivation [70]. Since gum chewing has been shown to alleviate the symptoms of xerostomia in certain conditions, such as Sjogren's syndrome, various clinical experiments have been conducted to support this claim. With pilocarpine added to the chewing gum, salivary secretion can be boosted to an even greater extent. Saliva has a buffering capacity and may be able to lower the acidity of the stomach juice. Gum without active ingredients has been shown to prevent postprandial reflux in clinical trials. In order to get the most out of this, an antacid should be added to the gum.

Gum chewing stimulates the flow of saliva. Mechanical mastication is considered a key element in this boosting effect. Findings show that chewing gum could potentially decrease gingivitis and carious lesions. Additionally, calcium-containing gums could remineralize incipient caries [71]. Tooth plaques are the most critical risk factor affecting dental and periodontal health. Periodontal disease could be partly treated by sodium bicarbonate/sorbitol mixture-based gums. These types of gum could increase the pH of saliva and reduce dental plaque accumulation [72]. Patients with weakened immune systems are more likely to develop bacterial or fungal infections of the mouth. Dental and oral infections can be alleviated by chewing chlorhexidine gum. Dental plaques can be treated with chlorhexidine and decapeptide-based antiseptic gums. Chlorhexidine/xylitol-based gums have the potential to significantly lower the load of S. mutans and lactobacilli in the mouth. [73–76]. Since the harsh taste of chlorhexidine in mouthwash is easily disguised by the sweet flavor of chewing gum, it is a better choice for daily oral hygiene than a chlorhexidine mouthwash [77].

Chewing gums with antibacterial actions in the oral cavity, such as gramicidin and neomycin, are similar to sulfathiazole chewing gums. For Vincent’s illnesses, metronidazole gums could substitute penicillin-loaded gums in terms of bacterial resistance. The miconazole chewing gum has been demonstrated to be at least as effective as a miconazole oral gel in treating oral candidiasis in clinical trials involving patients. In addition, patients favored chewing gum over oral gel since it was more convenient and had fewer adverse effects [78–80]. Namibian chewing gum, which contains Diospyros lycioides, indicates an antibacterial effect on Streptococcus mutans and Streptococcus sanguis [81, 82]. Another natural ingredient used in gum is the magnolia bark extract, which shows an antibacterial effect against Porphyromonas gingivalis, Fusobacterium nucleatum, and S. mutans [83].

Bacterial colonization has long been claimed to be prevented by fluoride-containing chewing gums. For example, fluoride ions block the metabolism of plaque bacteria when used in a dental therapy, as might other chemicals with therapeutic applications that act locally or are absorbed through the oral and buccal capillaries. Patients with xerostomia and children in fluoride-deficient areas, as well as adults with a high incidence of dental caries, may benefit from chewing fluoride-containing gum. Adults can also benefit from its use in preventing dental cavities [53–56].
Other commercially available options include vitamin C chewing gums and tablets. Chewing vitamin C-fortified gum at least five times a day for three months reduced the production of supragingival calculus in comparison to a control gum and no chewing gum at all. Calcium phosphate deposits are thought to be reduced because of the acidic characteristics of vitamin C [84]. However, dental enamel damage due to high local vitamin C concentrations is the main drawback of these gums [85]. Chewing gum with pyro/triphosphate supplementation showed similar benefits after six weeks of use, which may be due to the calcium-sequestering activities of polyphosphates on the enamel. However, the reduction in calculus formation was only observed on supragingival surfaces [86].

Oral health is becoming more and more concerned with appearance, notably the appearance of white teeth. Stains caused by chromogens from food, drink, or smoking can be extrinsic or intrinsic, depending on the source (or calculus). Polypseudophosphates have been added to sugar-free chewing gums to help prevent and remove extrinsic stains. In short-term (2 days) trials, a sugar-free gum containing sodium hexametaphosphate outperformed a control gum at reducing stain formation [87, 88].

Oral malodor is caused by anaerobic gram-negative bacteria adhering to the tongue or associated with periodontitis, which produces VSCs such as hydrogen sulfide and methyl mercaptan [89]. Gums that contain active compounds that target bacteria that cause bad breath have been shown to reduce the amount of VSCs in the mouth and the amount of VSCs in the mouth. To reduce VSC levels after chewing, the zinc-allyl isothiocyanate combination works particularly well because of zinc’s affinity for sulfur compounds [90, 91]. To reduce oral odor, chewing gum with the magnolia bark extract or eucalyptus essential oil has been successful when paired with zinc, which inhibits the viability of the bacteria that produce VSC [91, 92].

Probiotics (L. reuteri, L. salivarius, and L. plantarum) have been studied to minimize dysbiosis and maintain a balanced microbiota in the form of chewing gum. Because of the prevention of antibiotic adverse effects, probiotics are also indicated as a supportive treatment alongside scaling and root planning [93].

Minor pain treatment can benefit from the use of chewing gums as a drug delivery mechanism because of its rapid onset of action and reduced risk of digestive side effects. Up to 63% of the normal dose of acetylsalicylic acid can be delivered by chewing an acetylsalicylic gum for 15 minutes. Drug absorption rates are faster in the liquid form compared to a tablet form with the same dosage of the same medications. Toothaches could possibly be relieved faster if drugs had a faster absorption rate [94–96].

There are various limitations to using chewing gum as a medicine delivery device, including the fact that it prevents you from eating, drinking, and conversing while you have a delivery system in your mouth. Due to saliva dilution, the mouth cavity always shrinks, and the medicine secreted in saliva soon disappears as a result of the swallowing process. When it comes to the release of drugs from chewable formulations, chewing habits have been shown to have a substantial impact.

3.5. Future Trends. The science of using different carriers as a drug delivery system is advancing daily. Chewing gum, mouse [97], exosomes [98], and micro- and nanorobot [99] have been considered. More attempts will be made in the future to develop drugs that use chewing gum as a drug delivery mechanism. The treatment of fungal infections, prevention of cavities and other dental health problems, remineralization of teeth, cold relief, increased energy, antinausea, and a slew of other benefits of this novel drug delivery technique are all likely to play a key role in future research. Chewing gum does, in fact, take some time to gain acceptance as a method of drug delivery. Alternative delivery mechanisms for administering pharmaceuticals locally to the oral cavity may be replaced by medications incorporated into chewing gums. The reason is simple: the chewing gum administration system is convenient, easy to deliver anywhere, at any time, and its pleasant taste encourages patient compliance, particularly among children.

4. Conclusions

Gum-driven drugs could be introduced as promising candidates for treating oral diseases. This is due to their ability to deliver the proper local dosages of active ingredients, short contact time, biocompatibility, biodegradable chemical structures, and ability to maintain a state of eubiosis. These benefits have spurred many people to research to make a lot of different kinds of medicated chewing gum commercials.

Data Availability

The data used to support the findings of this study are available from the corresponding author upon request.

Conflicts of Interest

The authors declare that there are no conflicts of interest regarding the publication of this article.

Authors’ Contributions

Conceptualization or design of the work was done by M.B. and K.BL. Acquisition of data was carried out by E.S, K.BL., and S.M. Analysis and interpretation of data were done by M.B., SM.M., Z.V., and E.S. The draft was written by M.B., D.R., S.M, and Z.V. Revision of the manuscript for important intellectual content was done by MH.B., D.R., Z.V, M.B, K.BL, SM.M., and S.M. All authors gave final approval and agreed to be accountable for all aspects of the work.

References

[1] F. Oroojalian, F. Charbgoo, M. Hashemi et al. “Recent advances in nanotechnology-based drug delivery systems for the kidney,” Journal of Controlled Release, vol. 321, pp. 442–462, 2020.
[2] O. S. Fenton, K. N. Olafson, P. S. Pillai, M. J. Mitchell, and R. Langer, “Advances in biomaterials for drug delivery,” Advanced Materials, vol. 30, 2018.
[3] M. R. Prausnitz, S. Mitragotri, and R. Langer, “Current status and future potential of transdermal drug delivery,” Nature Reviews Drug Discovery, vol. 3, pp. 115–124, 2004.

[4] A. Z. Wang, R. Langer, and O. C. Farokhzad, “Nanoparticle delivery of cancer drugs,” Annual Review of Medicine, vol. 63, pp. 185–198, 2012.

[5] J. Hrkach and R. Langer, “From micro to nano: evolution and impact of drug delivery in treating disease,” Drug Delivery and Translational Research, vol. 10, pp. 567–570, 2020.

[6] J. Shi, A. R. Votruba, O. C. Farokhzad, and R. Langer, “Nanotechnology in drug delivery and tissue engineering: from discovery to applications,” Nano Letters, vol. 10, pp. 3223–3230, 2010.

[7] O. C. Farokhzad, S. Jon, A. Khademhosseini, T.-N. T. Tran, D. A. LaVan, and R. Langer, “Nanoparticle-aptamer bioconjugates,” Cancer Research, vol. 64, pp. 7668–7672, 2004.

[8] A. S. Surana, “Chewing gum: a friendly oral mucosal drug delivery system,” International Journal of Pharmaceutical Sciences Review and Research, vol. 4, pp. 68–71, 2010.

[9] D. Khatiwara, P. Ranabhat, M. Paul, and A. Bagchi, “An overview of calcium phosphates used in biomimetic oral care,” International Journal of Pharmacy, vol. 320, pp. 26–32, 2004.

[10] F. Meyer, B. T. Amaechi, H.-O. Fabritius, and J. Enax, “Novel and emerging techniques of medicated chewing gum in drug delivery system,” The Journal of the American Dental Association, vol. 139, pp. 190–196, 2008.

[11] Y.-E. Lee, J.-H. Kim, M.-J. Cho, K.-B. Song, and Y.-H. Choi, “Effect of xylitol on dental caries prevention: a literature review,” Journal of Korean Society of Dental Hygiene, vol. 19, pp. 449–465, 2019.

[12] E. Aluckal and A. V. Ankola, “Effectiveness of xylitol and xylitol-sweetened chewing gum—a potential drug delivery system,” Expert Opinion on Drug Delivery, vol. 7, pp. 871–885, 2010.

[13] B. A. Burt, “The use of sorbitol-and xylitol-sweetened chewing gum in caries control,” The Journal of the American Dental Association, vol. 139, pp. 190–196, 2008.

[14] A. Z. Wang, R. Langer, and O. C. Farokhzad, “Nanoparticle delivery systems: recent developments and future prospects,” Journal of Research in Medical Sciences, vol. 6, pp. 437–445, 2021.

[15] M. J. Rathbone, J. Hadgraft, and M. S. Roberts, “Modified-release drug delivery technology,” Drugs and the Pharmaceutical Sciences, vol. 126, pp. 101–114, 2003.

[16] A. A. Aslani and F. Rostami, “Medicated chewing gum, a novel drug delivery system,” Journal of Research in Medical Sciences, vol. 6, pp. 437–445, 2021.

[17] M. T. Newton, O. Awojobi, M. Nasseripour et al., “A systematic review and meta-analysis of the role of sugar-free chewing gum in dental caries,” JDR Clinical & Translational Research, vol. 5, pp. 214–223, 2020.

[18] Y. Morjaria, W. J. Irwin, P. X. Barnett, R. S. Chan, and B. Conway, “In vitro release of nicotine from chewing gum formulations,” Dissolution Technologies, vol. 11, pp. 12–15, 2004.

[19] K. K. Chenab, B. Sohrabi, and A. Rahmanzadeh, “Superhydrophobicity: advanced biological and biomedical applications,” Biomaterials Science, vol. 7, pp. 3110–3137, 2019.

[20] E. Aluckal and A. V. Ankola, “Effectiveness of xylitol and xylitol-sweetened chewing gum in dental caries,” JDR Clinical & Translational Research, vol. 5, pp. 214–223, 2020.

[21] Y. Morjaria, W. J. Irwin, P. X. Barnett, R. S. Chan, and B. Conway, “In vitro release of nicotine from chewing gum formulations,” Dissolution Technologies, vol. 11, pp. 12–15, 2004.

[22] L. T. Newton, O. Awojobi, M. Nasseripour et al., “A systematic review and meta-analysis of the role of sugar-free chewing gum in dental caries,” JDR Clinical & Translational Research, vol. 5, pp. 214–223, 2020.

[23] Y.-E. Lee, J.-H. Kim, M.-J. Cho, K.-B. Song, and Y.-H. Choi, “Effect of xylitol on dental caries prevention: a literature review,” Journal of Korean Society of Dental Hygiene, vol. 19, pp. 449–465, 2019.

[24] E. Aluckal and A. V. Ankola, “Effectiveness of xylitol and xylitol-polyol chewing gum on salivary streptococcus mutans in children: a randomized controlled trial,” Indian Journal of Dental Research, vol. 29, p. 445, 2018.

[25] M. Upadhyay, S. K. R. Adena, H. Vardhan, S. Pandey, and B. Mishra, “Development and optimization of locust bean gum and sodium alginate interpenetrating polymeric network of capecitabine,” Drug Development and Industrial Pharmacy, vol. 44, pp. 511–521, 2018.

[26] U. M. Deogade, V. N. Deshmukh, and D. M. Sakarkar, “Natural gums and mucilage’s in NDDS: applications and recent approaches,” International Journal of PharmTech Research, vol. 4, 2012.

[27] L. Chow, S. Takagi, R. Shern, T. Chow, K. Takagi, and B. Sieck, “Effects on whole saliva of chewing gums containing calcium phosphates,” Journal of Dental Research, vol. 73, pp. 26–32, 1994.

[28] F. Meyer, B. T. Amaechi, H.-O. Fabritius, and J. Enax, “Overview of calcium phosphates used in biomimetic oral care,” The Open Dentistry Journal, vol. 12, p. 406, 2018.
[71] V. Carrard, M. Roxo Gonçalves, J. Rodriguez Strey et al., “Telediagnosis of oral lesions in primary care: the EstomatoNet program,” *Oral Diseases*, vol. 24, pp. 1012–1019, 2018.

[72] D. H. Na, J. Faraj, Y. Capan, K. P. Leung, and P. F. DeLuca, “Chewing gum of antimicrobial decapate (KSL) as a sustained antiplaque agent: preformulation study,” *Journal of Controlled Release*, vol. 107, pp. 122–130, 2005.

[73] E. Caglar, S. Kavaloğlu, O. Kuscü, N. Sandalli, P. Holgerson, and S. Twetman, “Effect of chewing gums containing xylitol or probiotic bacteria on salivary mutans streptococci and lactobacilli,” *Clinical Oral Investigations*, vol. 11, pp. 425–429, 2007.

[74] K. K. Mäkinen, K. P. Isotupa, P. L. Mäkinen et al., “Six-month polyol chewing-gum programme in kindergarten-age children: a feasibility study focusing on mutans streptococci and dental plaque,” *International Dental Journal*, vol. 55, pp. 81–88, 2005.

[75] G. H. Hildebrandt and B. S. Sparks, “Maintaining mutans streptococci suppression: with xylitol chewing gum,” *The Journal of the American Dental Association*, vol. 131, pp. 909–916, 2000.

[76] P. Milgrom, K. Ly, M. Roberts, M. Rothen, G. Mueller, and D. Yamaguchi, “Mutans streptococci dose response to xylitol chewing gum,” *Journal of Dental Research*, vol. 85, pp. 177–181, 2006.

[77] Y. Patel, A. Shukla, V. Saini, N. Shrimal, and P. Sharma, “Chewing gum as a drug delivery system,” *International Journal of Pharmaceutical Sciences and Research*, vol. 2, p. 748, 2011.

[78] R. Emslie, “Treatment of acute ulcerative gingivitis. A clinical trial using chewing gums containing metronidazole or penicillin,” *British Dental Journal*, vol. 122, p. 307, 1967.

[79] J. L. Rindum, P. Holmstrump, M. Pedersen, M. R. Rassing, and K. Stoltze, “Miconazole chewing gum for treatment of chronic oral candidosis,” *European Journal of Oral Sciences*, vol. 101, pp. 386–390, 1993.

[80] R. Pavelic, “Use of an analgesic-antibiotic chewing troche (orabitec) in post-tonsillectomy and adenoidectomy complications,” *Eye, Ear, Nose & Throat Monthly*, vol. 39, p. 644, 1960.

[81] L. Cai, G.-X. Wei, P. van der Bijl, and C. D. Wu, “Namibian chewing stick. Diospyros lycoidea, contains antibacterial compounds against oral pathogens,” *Journal of Agricultural and Food Chemistry*, vol. 48, pp. 909–914, 2000.

[82] K. C. Chinsembu, “Plants and other natural products used in the management of oral infections and improvement of oral health,” *Acta Tropica*, vol. 154, pp. 6–18, 2016.

[83] M. Greenberg, P. Urnezis, and M. Tian, “Compressed mints and chewing gum containing magnolia bark extract are effective against bacteria responsible for oral malodor,” *Journal of Agricultural and Food Chemistry*, vol. 55, pp. 9465–9469, 2007.

[84] P. Lingström, S. Fure, B. Dinitzen, C. Fritznez, C. Klebom, and D. Birkhed, “The release of vitamin C from chewing gum and its effects on supragingival calculus formation,” *European Journal of Oral Sciences*, vol. 113, pp. 20–27, 2005.

[85] T. Garg and A. Goyal, “Medicated chewing gum: patient compliance oral drug delivery system,” *Drug Delivery Letters*, vol. 4, pp. 72–78, 2014.

[86] P. F. Porciani, S. Grandini, and S. Sapiò, “Anticalculus efficacy of a chewing gum with polyphosphates in a twelve-week single-blind trial,” *Journal of Clinical Dentistry*, vol. 14, pp. 45–47, 2003.

[87] P. A. Walters, A. R. Biesbrock, and R. D. Bartizel, “Benefits of sodium hexametaphosphate-containing chewing gum for extrinsic stain inhibition,” *American Dental Hygienists’ Association*, vol. 78, p. 8, 2004.

[88] P. F. Porciani, C. Perra, and S. Grandini, “Effect on dental stain occurrence by chewing gum containing sodium tripolyphosphate—a double-blind six-week trial,” *Journal of Clinical Dentistry*, vol. 21, pp. 4–7, 2010.

[89] K. Hampel ska, M. M. Jaworska, Z. Ł Babalska, and T. M. Karpiński, “The role of oral microbiota in intra-oral halitosis,” *Journal of Clinical Medicine*, vol. 9, p. 2484, 2020.

[90] T. Blom, D. Slot, M. Quirynen, and G. Van der Weijden, “The effect of mouthrinses on oral malodor: a systematic review,” *International Journal of Dental Hygiene*, vol. 10, pp. 209–222, 2012.

[91] P. F. Porciani and S. Grandini, “The effect of zinc lactate added tablets on volatile sulfur-containing compounds in the oral cavity,” *The Open Dentistry Journal*, vol. 13, 2019.

[92] P. F. Porciani and S. Grandini, “The effect of zinc acetate and magnolia bark extract added to chewing gum on volatile sulfur-containing compounds in the oral cavity,” *Journal of Clinical Dentistry*, vol. 23, pp. 76–79, 2012.

[93] A. Butera, S. Gallo, C. Maiorana et al., “Probiotic alternative to chlorhexidine in periodontal therapy: evaluation of clinical and microbiological parameters,” *Microorganisms*, vol. 9, p. 69, 2020.

[94] L. L. Christrup, J. Bonde, H. Eriksen, S. N. Rasmussen, M. R. Rassing, and K. Simonsen, “Chewing gum as a drug delivery system III: bioavailability of salicylamlid administered in tablets and chewing gum,” *Farmaco Science Education*, vol. 16, 1988.

[95] M. J. Desborough and D. M. Keeling, “The aspirin story—from willow to wonder drug.” *British Journal of Haematology*, vol. 177, pp. 674–683, 2017.

[96] S. Indhumathi and K. S. Kumar, “A review on medicated chewing gum and its role in mouth ulcers,” *Research Journal of Pharmacy and Technology*, vol. 13, pp. 481–484, 2020.

[97] A. Butera, S. Gallo, M. Pascadopoli et al., “Paraprobiotics in non-surgical periodontal therapy: clinical and microbiological aspects in a 6-month follow-up domiciliary protocol for oral hygiene,” *Microorganisms*, vol. 10, p. 337, 2022.

[98] C. Gutierrez-Millan, C. Calvo Diaz, J. M. Lanao, and C. I. Colino, “Advances in exosomes-based drug delivery systems,” *Macromolecular Bioscience*, vol. 21, Article ID 2000269, 2021.

[99] M. Hu, X. Ge, X. Chen, W. Mao, X. Qian, and W.-E. Yuan, “Micro/nanorobot: a promising targeted drug delivery system,” *Pharmaceutics*, vol. 12, p. 665, 2020.