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Highlighting the impact of chitosan on the development of gastroretentive drug delivery systems

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1. Introduction

Chitosan (CS) is a biopolymer obtained from the deacetylation of chitin that presents several applications in photography [1], cosmetic/ skin regeneration [2], food and nutrition [3], environmental protection [4] and drug delivery systems [5]. Specifically, CS is very useful for pharmaceutical applications due its non-toxic nature, biodegradability, bioadhesivity, biocompatibility besides the capability to form films, gels, nano and microparticles [6,7]. Thus, floating, mucoadhesive, porous and gastric pH resistant pharmaceutical systems may be originated using pristine CS or associated with other materials [8,9].

Investigations have been carried out with the purpose of suggesting, developing and testing microspheres, micelles, hydrogels and nanoparticles using this biopolymer [10–12]. An innovative application of CS is to develop systems with the capacity to remain and release drugs in the stomach. These systems are labelled gastroretentive drug delivery systems (GRDDSs), which demarcate an important frontier in gastric and drug transport therapies that require gastric absorption and/or local action. Consequently, these systems circumvent the short retention time caused by gastric emptying and some physicochemical limitation, such as low solubility of some drugs.

Despite the numerous advantages of controlled drug delivery systems, many drugs with high solubility at stomach pH, instability in intestinal pH, as well as those with narrow absorption window and that are intended for the local treatment of stomach disorders, require additional mechanisms for spatial control of the release, for which GRDDSs classification has been attributed. This topic has been extremely relevant according to the number of preclinical and clinical studies in which their effectiveness have been demonstrated by increasing the drugs absorption despite the low absorption window [15].

Many characteristics attributed to CS are directly related to the preservation of protonable amino groups in the polymeric chain, which are nucleophilic and reactive regions of the molecule. Therefore, being able to interact with a large number of materials, and therefore be widely applied in the most diverse fields, from water treatment to the development of controlled drug release systems, its versatility can be explored individually or in combination with other materials [16].

This review highlights recent works addressing to the CS-based materials to fabricate GRDDSs. In the Scopus database, when requested to search using the keywords “chitosan”, “gastroretentive” and “drug delivery”, 1544 documents were found from 2009 to 2019. (Fig. 1). It is worth noting that since 2013 the number of publications on this topic has increased, showing great interest on the part of scientific community.
2. Chitin and chitosan

Discovered in 1811 by Henri Braconot, chitin is a polysaccharide formed by $\beta(1\rightarrow4)$-2-acetamide-2-deoxy-$\alpha$-glucopyranose units [17]. The orientation of these polymeric chains gives rise to different allomorphs of chitin, forming anti-parallel and parallel arrangements known as alpha, beta and gamma-chitin. Chitin is a hydrophobic polymer, with fewer applications than CS, because it is not dispersible in aqueous, acids and alkaline media, being possible its dispersion only in hexafluoroisopropanol and chloroalcohols. Chitin is usually found in exoskeletons of invertebrate animals as crabs, lobsters, shrimps, cockroaches and scorpions. In addition to these sources, chitin and CS also occur naturally in some fungi like *Mucor rouxii* (30% of total mass) and *Choanephora cucurbitarum* (28% of total mass) [18,19].

CS was first observed in 1859 by Rouget [20]. Later, authors showed that CS is obtained by alkaline deacetylation of chitin (Fig. 2) under heating [20]. CS consists of monomeric units of $\beta(1\rightarrow4)$-2-acetamide-2-deoxy-$\alpha$-glucopyranose and $\beta(1\rightarrow4)$-2-amino-2-deoxy-$\alpha$-glucopyranose, predominantly the last one [17,21,22].

CS presents interesting characteristics like biocompatibility, biodegradability and low toxicity being a versatile biomaterial used in many research areas like textiles, food industry, environmental, agriculture,

![Figure 1](image1.jpg)

**Fig. 1.** Publications from the last 10 years containing the terms “chitosan”, “gastroretentive” and “drug delivery”.

![Figure 2](image2.png)

**Fig. 2.** Deacetylation of chitin producing CS.
medical products, cosmetics and mainly, in pharmaceutical applications [23–26].

In relation to the different CS applications, its dispersion in acidified aqueous media promotes the increase of liquid formulations viscosity, making CS also applicable as a thickener. Additionally, the high viscosity of these systems associated with their polycationic nature gives to the CS impressive mucoadhesive properties. This can be attributed to the CS interaction with sialic acid, an anionic component present in the mucin [27]. The properties above mentioned make CS a special biopolymer used to develop specific (mucoadhesive) drug delivery systems [28].

CS amino groups in acidified aqueous media can become protonated, giving rise to a positively charged material, which can interact with negative charges from anionic polymers, cell membranes, mucus and complex reactive oxygen species (ROS). When not protonated, amine groups can still complex metallic ions like Cu$^{2+}$ and are mainly used in water treatment and clarification of beverages [29–31].

For biological applications, CS also requires dispersion in acidic aqueous solutions with pH values lower than 6.5. In this condition, the CS amino groups (NH$_2$) present in the N-acetyl-d-glucosamine units act as nucleophiles becoming progressively protonated due to the ionization of acids present in the medium, forming a positively charged polyelectrolyte, as shown in Eq. (1) [32].

\[ ChNH_2 + H_3O^+ = ChNH_3^+ + H_2O \] (1)

Changes in the CS chain can directly affect its physicochemical properties, leading to different technological applications changing also its biological activity [33,34]. Some types of CS have different degrees of deacetylation (DD) and molecular weight (MW), which are determinant for their polycationic nature at pH < 6.5. Variations, such as high DD and low MW can increase inter- and intra-chain electrostatic repulsion, favoring biopolymer-water interactions and reducing biopolymer-biopolymer interactions. This effect is also responsible for increasing interactions with negative surfaces like lipids, cells, drugs and anionic molecules, as well as mucus conferring mucoadhesive properties [35].

Interestingly, the CS interaction with anionic polymers can lead to gels formation by complex coacervation [36]. Interaction with smaller polyanions such as sodium tripolyphosphate leads to the formation of bridges (junction zones) between the polymer chains forming gel by ionicotropic gellation [37]. These kinds of interactions make protonated CS interesting biopolymer for the development of pharmaceutical systems such as micro- and nanoparticles, gels and films [38].

The ability to form particles applied as GRDDSs is associated with characteristics of the polymeric chain, such as DD and MW, as well as pH value of the dispersion medium. CSs with low DD are more efficiently dispersible at high pH due to their high charge density that favors more intensely interactions with polymers, drugs and cross-linking agents, such as sodium tripolyphosphate. On the other hand, CS with high MW shows greater capacity than low MW to increase the media viscosity even at low concentrations, making the diluted/semi-diluted dispersion concentration regime suitable for particles formation [37].

The knowledge about structural and biological properties may provide a basis for exploring the potential and versatility of CS-based materials applications as smart drug delivery systems. For instance, changes in CS rheological behavior in response to pH changes can favor the fabrication of suitable GRDDSs [39]. At acidic pH, CS can increase the viscosity in the medium, promoting good adhesion. Beyond that, the modification of CS structure can increase the possibilities of its biological applications [6] as shown in Table 1.

Friedman et al. [56], demonstrated that CS anti-inflammatory activity is related to the blocking of NK-κ-B in human type 1 mast cells inhibiting the production of pro-inflammatory cytokines. This effect can occur due to a downregulation of Ca$^{2+}$-dependent pathways [57]. In addition, CS antioxidant activity is directly associated with the chemical reactivity of the hydroxyl and amino groups from the C-6 and C-2 carbons, respectively [58]. Substitutions performed on CS by reducing the DD promote the exposure of more amino groups in the polymer chain increasing its antioxidant activity. The bio-polymer chain reduction (MW) also exerts a positive effect on the CS antioxidant activity. Smaller biopolymer chains are very rigid, so the intra-chain hydrogen bonds that involve hydroxyl groups and amino acids are less intense than the same interactions in larger chains with more conformational freedom [45]. Thus, smaller polymer chains can interact more efficiently with oxidant molecules inhibiting their action.

CS hypolipidemic activity is associated with its polycationic nature, interacting with dietary lipids inhibiting their absorption. Additionally, CS can reduce the liver damage associated with the oxidation of fatty acids, leading to a considerable reduction in liver enzymes aspartate aminotransferase (AST) and alanine aminotransferase (ALT) and elevation in activities of antioxidant enzymes such as superoxide dismutase (SOD), catalase (CAT), and glutathione peroxidase (GSH-Px). Beyond that, diets rich in CS and chitosan oligosaccharides have shown the reduction of serum triglyceride and LDL-cholesterol levels as well as regulate the expression of liver peroxisome proliferator-activated receptor α (PPARα) [48,59].

Most of studies have demonstrated that CS and their derivatives display anticoagulant properties. These biopolymers show low toxicity reducing side effects and tumor size by inhibiting tumor cells proliferation and, consequently, provoking apoptosis [52,60].

Considering CS applicable as new biomaterials, this biopolymer could be used to fabricate scaffolds for regenerative medicine [61]. CS-based scaffolds exhibit advantages as biocompatibility, biodegradability and malleability. In addition, these biomaterials can easily interact with growth factors, glycosaminoglycans, and DNA making CS-based scaffolds excellent for bone, cartilage and intervertebral discs regeneration [62]. CS-based scaffolds allow an efficient angiogenic response, favoring the oxygen supply to cells, nutrients and chemical mediators for cells proliferation and differentiation, promoting osteogenesis in vitro and in vivo. CS hydrophilicity promotes cells adhesiveness by reducing immune reactions when applicable in implants [63].

NH$_3$ groups available in CS interact with erythrocytes promoting the aggregation and the formation of a physical barrier that prevents the blood loss without rejection by the organism [64]. CS with low DD and dispersed in acidified water can increase the complement pathway by recruiting more erythrocytes, promoting rapid coagulation. Additionally, the CS MW showed less impact on the coagulant effect. In this scenario, CS with MW between 10$^5$ and 10$^6$ Da displayed less coagulant capacity whereas CS with higher MW in physiological saline solution were more efficient in inducing erythrocyte aggregation. The effect of MW in CS suggests that long chains can interact with more platelets, promoting greater efficiency in hemostasis. Therefore, when DD is high the influence of MW on platelet aggregation is reduced. This effect is more pronounced by reducing DD, indicating that DD is the most important factor for platelet aggregation [54,65]. In light of this, the importance of CS to build GRDDSs orally administered is evident. However, considering the harsh barriers imposed by the gastrointestinal tract (GIT).

An important aspect about the use of CS to develop drug delivery systems for GIT consists on understand CS gastrointestinal digestion process in humans. This process still lacks in-depth studies that describe it systematically. However, up to now, some studies have shown that CS cannot be digested in the upper GIT by human digestive enzymes whereas CS digestion can occur mainly in the intestine, through depolymerization catalyzed by enzymes [66–71].

In 1990, Hirano et al. [67], from in vitro assays, showed that CS can be degraded by chitosanase enzymes produced and released by bacteria from the intestinal microbiota. Later, Varum et al. [68] demonstrated that CS can also be depolymerized in humans by the lysozyme present in the serum.
Table 1
Biological applications of distinct and modified CS-based materials

| Applications          | Materials/systems                                                                 | Remarks                                                                 | Refs. |
|-----------------------|----------------------------------------------------------------------------------|------------------------------------------------------------------------|--------|
| Antimicrobial         | HTCC (N-(2-hydroxypropyl)-3-trimethylammonium chitosan chloride), degree of substitution of 57, 62, 63 and 77%. | The inhibitory effect of HTCC against SARS-CoV-2 and MERS-CoV viruses was tested in vitro with Vero and Vero 6 cells, as well as ex vivo on human airway epithelium (HAE). | [40]   |
| CS nano- and microparticles | CS dispersed-Different DD (51.04%–100%), and ultra high MW CS (Mw > 10⁶ g mol⁻¹) | - strongest inhibition by HTCC-62 or HTCC-77  
- SARS-CoV-2: strong inhibition 24 h after virus inoculation.  
- HTCC blocks interaction between MERS-CoV with its cellular entry receptor DPP4  
- Inhibition of the virus may depend on the repeating units of the chitosan polymer  
Low MW and DD presents to higher antimicrobial activity. At basic pH, CS loses antimicrobial activity. | [41]   |
| Anti-inflammatory      | CS with different MWs and chitooligosaccharides (COS)                            | Larger MW (~29.2 kDa) CS usually exhibit anti-inflammatory activity. | [42]   |
| In-vivo (rats) test of anti-inflammatory activity in ulcer model of CS high and low MW. | Lambda-MW CS had the best activity on ulcer healing. | HMW CS helped retain the stomachal folds rendering a gastro protective effect, also can stimulate collagen synthesis. | [44]   |
| Antioxidant           | CS of MW (0.5–1000 kDa) and DD (50%–low then 10%)                              | Reduction of both DD and MW make CS more efficient in removing in vitro reactive oxygen species (ROS). | [45]   |
| Gamma irradiated (5–50 Kgy) and not irradiated CS DD (72–75%), dispersed in lactic acid | Gamma irradiation increased CS antioxidant activity by 10%. | Reduction of CS molecular weight by gamma irradiation at 50Kgy, increased antioxidant capacity of CS dispersion | [46]   |
| Lipid-lowering effects | Dietary supplementation with CS DD (70.03%) and MW (estimated 500–1000 kDa)      | Comparison of the lipid-lowering and intestinal morphological effects of cholestyramine, CS and olat gnm in rats. CS has hypolipidemic effects similar to cholestyramine without deleterious changes in the intestinal mucosa. Oat gnm was less effective | [47]   |
| Scaffold for tissue regeneration | Simvastatin-loaded CS nanoparticles | CS induced new osteoid tissue formation, showing biodegradability and controlled simvastatin release. After 14 days, CS promoted increase of the enzyme ALP (indicator of osteoblast mineralization). CS nanostructures presented porosity suitable for angiogenesis and bone nutrition | [48]   |
| CS-based hyaluronan hybrid polymer fiber | Fibroblasts from patellar tendon of Japanese white rabbit. Fibroblasts had adhesion onto hybrid fibers and produced collagen fibers after 14 days of culture | CS and chitooligosaccharides have the ability to regulate the body weight, liver and cardiac indices, fat/body ratio, as well as serum, liver, and fecal lipids. Simultaneously, maintain the appropriate activity of liver and serum superoxide dismutase (SOD), alanine aminotransferase (ALT), aspartate aminotransferase (AST), as well as liver and fecal total bile acids (TBA) | [49]   |
| Anticancer            | CS and chitooligosaccharides                                                    | Reduction of DD and WW of CS and its derivatives exhibiting good in vivo results, been efficient against prostate cancer, carcinogenic human alveolar basal epithelial cells, and hepatocellular carcinoma, been nontoxic for healthy cells | [50]   |
| Crab (Chionoecetes opilio) CS and Shiitake mushroom (Lentinula edodes) CS | Shiitake CS had the best inhibitory effect on the growth of the human neuroblastoma cell line (IMR 32, BCRC 60014) and the human liver hepatocellular carcinoma cell line (Hep G2, BCRC 60025), occurring because Shiitake CS is more deacetylated. | | [51]   |
| Hemostatic            | CS hydrogel containing nano bioglass                                            | Coagulation was twice faster than pure blood in in vitro hemocompatibility assay. | [52]   |
| CS-based hemostatic dressing ChitoGauze® | A commercial hemostatic dressing for temporary external control of wounds with severe bleeding. It is a CS dressing composed of a non-woven medical gauze of polyester/rayon mixture coated with CS (HemCon Medical Technologies, Portland, OR, USA). Used by US military forces, emergency medical services | | [53]   |

The digestion by enzymes action from intestine is related to the degree of deacetylation (DD) and molecular weight (MW) of CS. Authors reported that as higher the CS DD, lower is its degradability [68]. However, new studies have already shown that there are enzymes, but only present in vegetables and some bacteria, capable of cleaving the glycosidic bond, both in CS with high and low DD including chitin [72].

Another important factor in the CS degradability and digestibility is MW. A study performed by Chae et al. [73], in which it was administered orally 20 mg kg⁻¹ marked (fluorescent) CS varying MW and DD in rats, exhibited that CS with 230 kDa presented absorption 20 times lower than CS of 3.8 kDa. These results indicate that greater CS digestibility and consequent absorption could be achieved decreasing CS MW.

Considering above mentioned properties, CS is widely used to design colon-specific systems. Unlike when CS is associated with GRDDSs that release the drug in gastric media, in colon-specific systems the dissolution of CS in acidic pH is overcome by association or coating with other polymers. These systems pass through the stomach quickly without significant drug release, protecting the drugs until they reach the intestine. In this organ, CS undergoes the action of enzymes from the intestinal microbiota and from its degradation can release the drug [74].

Thus, a deep anatomophysiology knowledge of the different portions that compose GIT is essential for the correct performance of GRDDSs.
### 3. Anatomical and physiological aspects of the stomach

CS-based systems that present mucoadhesive, swelling, and acid erosion properties emerge as an alternative to explore different delivery strategies for the gastric environment. The oral route is the most used and convenient because it is a noninvasive way to access the stomach, being a safe administration route that allows self-administration and consequently, greater acceptance from the patient to the treatment [75]. However, the bioavailability of orally administered drugs may be influenced by several parameters such as gastric emptying and gastrointestinal transit, gastric mucosa status, stomach pH [76].

The stomach is located in the upper left part of the abdominal cavity below the diaphragm. Stomach size can change according to the amount of distention. After meal, approximately 1500 mL whereas the empty volume range from 25 to 50 mL [77]. Gastric pH in healthy conditions during fasting and fed state are 1.1 ± 0.15 and 3.6 ± 0.4, respectively. The pH returns to basal level between 2 and 4 h [78]. Anatomically, the stomach is divided in two parts: the proximal stomach, consisting of fundus and body, and the distal stomach, consisting of pylorus. The proximal stomach acts as a food reservoir while the distal stomach has a role in food processing, forming the chyme, which will be released later into the small intestine. The distal stomach also acts as a propelling pump that assists in gastric emptying (GE). The rate of GE is influenced by both food volume and gastric content composition [79,80]. Table 2 describes the main characteristics of upper GIT.

GE occurs during fasting and in the fed state. However, the motility pattern is different in both states. During fasting, a series of interdigestive electrical events occur every 2 or 3 h [81]. As described by Wilson and Washington [82], this process is called migratory myoelectric cycle (MMC), which is divided into the following 4 phases, illustrated in Fig. 3:

1. Phase I (basal phase) lasts from 30 to 60 min with rare contractions;
2. Phase II (preburst phase) lasts from 20 to 40 min with progressive high frequency and intensity contractions;
3. Phase III (burst phase) lasts from 10 to 20 min. Intense contractions occur and all undigested material is carried from the stomach to the small intestine;
4. Phase IV lasts from 0 to 5 min and occurs between phases III and I.

After meal, the contraction pattern changes from the fasting state to the fed state. It is known as digestive motility and comprises continuous

| Table 2 | Main characteristics of upper GIT. |
|---------|-----------------------------------|
|         | Length (m) | Transit time (h) | pH |
| Stomach | 0.2        | 0.5–2            | <3 |
| Small intestine | 6–10  | 3–4            | 5–8 |

Fig. 3. Schematic representation of the gastric emptying stages.
contractions as in phase II. These contractions result in size reduction of the food particles, which are propelled into the suspended pylorus. During the onset of the fed state, MMC is delayed resulting in decreased emptying rate [83].

The short gastric retention time of dosage forms and the unpredictable rate of GE are limiting factors that can be overcome by GRDDSs. These systems can increase gastric retention time contributing to an enhancement of drugs solubility and, consequently, bioavailability [79].

4. Mechanisms of gastroretention

During the planning of GRDDSs using CS, some particularities should be taken into account like length of retention on the drug/dosage form in the stomach, penetration in the gastric mucosal layer, stomach pH and stability of the drug/pharmaceutical system and CS protonation, drug’s molecule physicochemical parameters, as well as its solubility. Another important factor is the presence of lipids in the stomach. CS may interact with these molecules and cause changes in the drug release profile or even not release it [84].

The use of CS to obtain these types of systems has already been shown to be effective in terms of improving the gastric distribution of some drugs. Modi et al. [85] developed mucoadhesive CS nanoparticles (NPs) with ketoconazole and observed a 5-hour increase in residence and release time and, consequently, absorption of this drug by the stomach. In other study, Arora et al. [86] designed CS NPs containing alginate and pluronic F-127 for gastric amoxicillin release. The authors observed that the drug was protected from acid degradation, CS NPs showed adherence and mucopenetration releasing amoxicillin to deep regions of gastric mucus, increasing its therapeutic effectiveness. In addition, several research have already proven that CS is a polysaccharide that promotes the improvement of gastric delivery of the most diverse drugs, such as beberine [87], ranitidine hydrochloride [88], carvedilol [89] and moxifloxacin hydrochloride [90].

One of the main challenges that motivate the development of new GRDDSs is associated to the drugs physicochemical characteristics such as narrow absorption windows, local activity in the stomach, instability in the colon or distal small intestine and with low solubility at high pH [91]. The use of GRDDSs can circumvent these limitations mainly when associated with biopolymers as CS. This kind of association leads to the design of promising controlled drug delivery systems [92].

GRDDSs in contact with gastric medium can release the drug into stomach. GRDDSs can furnish controlled release profile, more efficient and adapted to different needs, promoting important advances in pharmacotechnical aspects [92].

Regarding the design of pharmaceutical systems for gastric diseases treatment, controlled drug release, specific absorption in stomach and pharmaceutical dosage form with tunable retention time are required. One of the strategies comprises the fabrication of GRDDSs with mucoadhesive properties. These systems can be obtained by using CS or CS combined with other polymers (polymeric blends). CS displays important and desirable characteristics for these smart systems such as biocompatibility, biodegradability, mucoadhesiveness, buoyancy, resistance to gastric pH and, in some cases, pharmacological activity [18,77,92–94].

4.1. GRDDSs mucus/bioadhesiveness

Bioadhesive systems may be comprised of natural or synthetic polymers that can interact attractively with a biological substrate. When the substrate is the mucus or mucosa, such systems are called mucoadhesive [95]. CS has become a popular component of this type of system because it is able to establish various types of mucus interactions with numerous hydrogen bonds and electrostatic interactions, promoting the dosage form adhesion [96]. The application of this system comes with several advantages: gastric drug release, including

Fig. 4. Schematic representation of mucoadhesion stages.
increased bioavailability, pharmaceutical dosage form (PDF) residence time in the stomach and reduction of first pass metabolism [89].

For the adhesion process, the GRDDSs and mucus molecules must interact at the interface. These interactions can occur through ionic and covalent bonds, Van der Waals interactions, hydrogen bonds and hydrophobic interactions [97]. The adhesion is marked by two main steps, contact and consolidation stages, as shown in Fig. 4.

Stage I - The initial contact between the PDF and the mucus begins with interpenetration of the chains in the mucus and adhesion to the surface [97]. At this stage, the polymer chains begin to interact with the medium and mucus. In cases where GRDDSs is developed with CS, it is essential that the acidic medium solution acts by protonating and hydrating the CS chains expecting more effective interactions with mucus and PDF.

Stage II - PDF adhesion consolidation occurs because the polymers constituting the system are mostly hydrated, in the case of CS, protonated and, therefore, presenting greater conformational freedom, being able to establish cooperative interactions with the mucus. The main interactions that occur in this stage are the hydrogen bonds, Van der Waals and electrostatic interactions [97]. Due to the high conformational freedom of the polymer chains and the close contact of the mucus, these chains diffuse through the mucosal layer, deepening and establishing more lasting interactions. Other important contribution to this stage is the suction-type effect promoted using polymers with high affinity for water that causes dehydration of the mucus, promoting the adhesion of the system; this mechanism is explained by the dehydration theory [98].

Nowadays, the phenomenon of mucoadhesion can be explained in part by six theories that individually or together are not able to exhaust discussions on this topic, since it is a complex phenomenon [99]. For CS-based mucoadhesive systems, the principal theories that explain adhesion are wetting, diffusion, electronic interlocking, adsorption and fracture, which are briefly described below.

4.1.1. Wetting theory
This approach is applicable to low viscosity liquid and semi-solid mucoadhesive systems. This theory considers the interfacial tension to estimate the spreadability degree of the system in the mucus. After the spreading phenomenon, the system penetrates into deformations and mucus recesses modifying the surface characteristic, interactions and interfacial tension leading to the adhesion phenomenon [99].

A method to estimate whether a liquid will have good spreadability and adhesion on a surface is the scattering coefficient ($S_b$) measure, using the Eq. (2) where: energies of interfaces are represented by $\gamma_B$ and $\gamma_T$ - interfacial tension between the system and the mucus, and the interfacial tensions of each of the phases by $\gamma_B$ - surface tension of the system and $\gamma_T$ - surface mucosal tension [97].

As the contact angle is easily calculated, a relation is established between it and the system-mucus interfacial tension. In this technique, a system with equal or close contact angles of 0° is sought [100].

$$S_b = \gamma_T - \gamma_B - \gamma_{BT}$$  \hspace{1cm} (2)

4.1.2. Diffusion theory
This theory proposes that adhesion is a two-way process (Fig. 5) due to the diffusion of the polymer chains into the mucus glycoprotein network, as well as mucus diffusion into mucoadhesive system [100]. It is a process in which the penetration rate depends on the diffusion coefficient, MW, cross-linking density, chain mobility/flexibility and expansion capacity of both [101].

From results based on rheological analysis and spectroscopic techniques, the diffusion process is interpreted as a kinetic process, i.e., time-dependent, so that it is possible to estimate the time required for maximum adhesion between two substrates during interpenetration [102], as depicted in Eq. (3).

$$t = \frac{L^2}{Db}$$  \hspace{1cm} (3)

where (L) is the depth of interpenetration, and (Db) the diffusion coefficient.

The adhesion strength of a polymer is maximal when the penetration is close to the size of its chain. It is assumed that the depth of
interpenetration required to produce an efficient mucoadhesion bond lies in the range 0.2–0.5 μm [99].

4.1.3. Electronic theory

In this theory, when chains with different electrical charges, solvated or not, approach to form ionic pairs, the mucoadhesive process is established. However, the most accepted approach is the occurrence of bidirectional [103,104] electrons transference from the mucus to the polymers. This bidirectional electronic transfer at Fermi level is responsible for promoting the formation of an attractive electric double layer at the muco-polymer interface. The strength of adhesion is proportional to the attraction intensity exerted on this interface [99,104,105].

4.1.4. Adsorption theory

According to this theory, after contact between the interfaces, adhesion occurs as the product of two groups of forces, primary and secondary. The primary ones are ionic and covalent, while the secondary forces are Van der Waal's forces, hydrogen bonding, and hydrophobic forces [106].

Interface characteristics determine which types of forces will predominante in mucoadhesion. In most systems, adhesion occurs through secondary type forces, which are individually weak, but due to their high number they become globally significant [107].

4.1.5. Fracture theory

This theory is substantially different from the others, because it measures the necessary force to separate one material from the other one, correlating it with the intensity of the adhesive forces. Thus, it is assumed that adhesion failure occurs between interfaces, ignoring the fact that failure usually occurs due to the low cohesion within one of the components involved in the adhesion process [97,108] (Fig. 6). Larger chains as well as systems with lower cross-linking densities have higher fracture forces, indicating higher adhesion strength [109].

The adhesion force is calculated indirectly by Eq. (4), where: \( S_m \) is the adhesion force, \( F_m \) is the force required for the detachment of the materials and \( A_0 \) is the total area of contact between the interfaces.

\[
S_m = \frac{F_m}{A_0}
\]

It is a very suitable approach for solid and semi-solid systems in which there is little penetration of the mucus chains [110].

4.2. CS-related factors that interfere with mucoadhesion

The theoretical approach to be considered when working with mucoadhesive systems will depend on factors that may influence PF adhesion, and it is necessary to take into account some parameters for the development of the system. In this session, we will discuss the main factors to consider when the polymer used is CS.

4.2.1. Concentration

During the development phase of the mucoadhesive systems it is important to establish the optimum concentration of the polymer used, especially in semi-solid systems [111]. In solid systems, it is observed that, to some extent, the adhesion force increases proportionally with the polymers concentration. Systems obtained with reduced polymer concentrations can lead to poor adhesion and short duration due to the reduced number of interactions between the systems and polymer. Very high concentrations of polymer impart an excessive stability to PF, in which the chains are folded and less accessible to the solvent, reducing the freedom of the chains, in order to reduce the adhesion [112,113].

4.2.2. Crosslinking

The crosslinking degree is an important factor to consider when using CS to fabricate mucoadhesive systems, because the main method of obtaining solid systems such as nano and microparticles is ionotropic gelation using sodium tripolyphosphate (TPP) or glutaraldehyde [114–117]. A high crosslinking density immobilizes the polymer chain, reducing its conformational freedom, affecting its flexibility and in turn on penetration and entanglement with the mucus. When CS is crosslinked, it can hydrated and swell. Swelling is positive for the adhesion of these systems by increasing the area of contact with the mucus, it can additionally provide control characteristics in the drug release [98] pH/DA and hydration.

Acidic media (pH < pKa 6.5) exert a strong effect on the adhesive- ness of CS-containing mucoadhesive systems. This is due to the polycatalytic CS character, which contributes to the establishment of electrostatic interactions and hydrogen bonding with solvent and mucus. The CS DD is the main factor responsible for the cationic nature of this polymer, so that the reduction of the DD is directly associated with the increase in the positive charge's density of the chain, which contributes to the inter- and intra-chain CS repulsion. Thus, in acidic media, the chains with the lowest DD will be more protonated and separated from each other, because of the electrostatic repulsion. The polymer protonation exposes groups previously inaccessible to the solvent, increasing the capacity to establish more hydrogen bonds, having an
### Table 3
Mucoadhesive CS-based GRDDSs.

| Polymer(s)/drug | Objective | Methods | Results | Refs. |
|----------------|-----------|---------|---------|-------|
| CS; Scorpion Venom *Mesobuthus eupeus* | Obtain CS nanoparticles containing scorpion venom in order to promote the animals hyperimmunization. | Nanoparticles (NP) were fabricated by ionotropic crosslinking with TPP in different CS concentrations. NPs were characterized. In vitro release assays were performed and the efficiency and encapsulation capacity determined. | Particles obtained using CS 2 mg mL\(^{-1}\) and CS/TPP mass ratio of 2 containing 500 μg mL\(^{-1}\) of *M. eupeus* venom presented encapsulation efficiency of 91.3%, loading capacity of 76.3% and size mean of the 300–400 nm range (polydispersity index: 0.429). The release occurred in two stages: an initial stage with release of 60% of the venom in the first 10 h, followed by the controlled release for 60 h. | [119] |
| CS, Tween 80, pluronic F127/Emodin | Obtain, characterize and test the inhibition effect of nanomicelles loaded with mucoadhesive beads (NFM-Beads) on tumor cells. | CS-coated emodin-loaded pluronic F127/Tween 80 mixed nanomicelles were prepared by thin film hydration method, tested against human gastric carcinoma. NFM-Beads were prepared by ionotropic gelation using sodium carboxymethylcellulose (CMC) and aluminum chloride. Release, floating and mucoadhesion of the samples were evaluated in vitro and gastric retention evaluated in vivo. | The encapsulation efficiency was approximately 97.74% using 140 mg Pluronic F127 and 10 mg Tween 80. The size of the nanomicelles were in a range of 206.8–288.6 nm, PDI of 0.162–0.295, and their zeta potential was around +30.76 mV, suggesting physical stability. Emodin-loaded nanomicelles demonstrated better antimosis efficacy compared to corresponding emodin suspensions. NFM-Beads showed suitable buoyancy. Emodin release occurred by an anomalous process guided by swelling and erosion. | [120] |
| CS-HPMC/flavonoid taxofolin-Syloid AL-1 | Obtain, characterize and test CS mucoadhesive GRDDS microparticles in the inclusion complex of Taxofolin (TAX) and Syloid® AL-1, with proton pump inhibitory action. | Inclusion complex taxofolin-Syloid AL-1 was prepared using the solvent evaporation method, in Tax/Syloid® AL-1 ratio of 30:70, w/w. Microparticles were prepared using spray-dried method, in 6:1 ratio, CS:HPMC. Mucoadhesion and the drug release profile from the systems were tested. | In vitro 90% of the TAX permeated pig mucus in 2 h, during the 45 to 120 min interval, the average permeation rate was 3.3 mg cm\(^{-2}\) min\(^{-1}\). Ex vivo The average flux calculated across the mucus in the interval of 45–120 min was 0.8 10\(^{-1}\) mg cm\(^{-2}\) min\(^{-1}\). CS microparticles released taxofolin for 5 h in simulated gastric fluid. Microparticles adhered to gastric mucus for 5 h avoiding TAX intestinal degradation. Microparticles favored Tax absorption in the stomach, achieving rapid therapeutic onset in the treatment of gastric ulcer, and made it possible to avoid TAX degradation in the small intestine. | [121] |
| CS/Ranitidine | Preparation and characterization of CS mucoadhesive microparticles containing ranitidine. | The microparticles were obtained by means of CS ionotropic gelation using TPP, both in different concentrations. Size, shape, encapsulation efficiency, in vitro biodistribution using agar plate method, in vitro release kinetics were successfully performed. | The particles could be obtained with concentrations between 4–5% of TPP and 2% of CS showing an average size of 620–720 μm. These systems encapsulated between 41.67 and 87.58% of ranitidine. The bioadhesion test showed that between 62 and 83% of the particles remained adhered for 8 h. At least, 75% of the formulations showed buoyancy for 12 h. The release of ranitidine obeyed zero-order kinetics, with 86% of ranitidine being released in 120 h. | [122] |
| Micromotors coated with CS/Clarithromycin | Preparation and characterization of Clarithromycin (CLR)-loaded Mg-micromotors coated with CS in *H. pylori* treatment. | Mg-micromotor was prepared by an asymmetrical coating of the Mg microspheres with a thin TiO\(_2\) layer using atomic layer deposition, after Mg-TiO\(_2\) Janus microparticles were then coated with a PEGDA film containing the CLR antibiotic payload. After the drug-loading step, the microparticles were coated with an outer thin CS layer (thickness ~ 100 nm). Micromotors were evaluated for drug carrying capacity. In vivo: Capacity to retain micromotors in the mouse stomach, anti-*H. pylori* therapeutic efficacy and toxicity assessment. | The systems demonstrated security in animal model (mouse). Although the therapeutic efficacy of standard treatment and micromotors have been similar. The systems further reduced the bacterial load of *H. pylori* and showed an efficient distribution and retention in the mouse stomach. The micromotors were capable to load antibiotic cargo with high-loading efficiency-1032 ± 37 μg per 2 mg micromotor. | [123] |
| Interpolimeric blend (IPB) | Develop and test IPB gastroretentive | Poly-x-lipo nanoparticles: Both IPB and PXLNET matrices were | | [124] |
impact on PF hydration. The hydrated chains swell, increasing the adhesion area, interpenetration and interaction with mucus and water, through electrostatic interactions and hydrogen bonds. These interactions are the result of a pronounced increase in the conformational freedom of these hydrated chains [112,115].

4.2.3. Molecular weight (MW), shape and flexibility of the polymer chains

The increase in polymers MW can enhance the adhesiveness of mucoadhesive pharmaceutical systems as well as chains flexibility [118]. Long and flexible polymer chains such as high MW CS are able to enter deeper into mucus and establish more interactions because it presents greater mobility than polymers of very short and excessively enveloped chains. However, the use of polymers with excessively long chains is deleterious to mucoadhesive systems because they lose the ability to diffuse and entangle the mucus. In the development of these systems, it is essential to establish an optimum size for each polymer, since the shape and flexibility of the polymer chain is also a factor to be considered along with the size [97,107].

Table 3 lists some mucoadhesive GRDDSs obtained from CS and their particularities.

Considering the Table 3, CS applicability is vast, and it can integrate different types of gastroretentive systems. Despite the use of this biopolymer as carrier since 1980s [125], recent advances in nanotechnology favored the development of efficient drug delivery systems such as CS-based micromotors and nanosystems. CS remains contemporaneous and promising biopolymer to develop gastroretentive/mucoadhesive systems with controlled release profile. Although CS gastroretention properties are usually ascribed to its mucoadhesivity, other action mechanisms will be discussed in the next sessions.

4.3. Expandable GRDDSs

Expandable gastroretentive systems were originally designed to treat veterinary pathologies with high dimensions of about 15 × 3 cm (length × diameter) by Laby [126] in order to avoid regurgitation after administration and arrival of the pharmaceutical dosage form in the rumen. Posteriorly, they were also adapted for treatment of humans by Johnson and Rowe [127]. Early, veterinary devices exploited the unfolding mechanism and had a ring-shape or multilayer insoluble polymeric sheet from Griffin and Brewer [128]. On the other hand, the first device for human use was a tablet based on thiolated gelatin which after hydration in the stomach and swelling to a larger size, making it impossible to pass through the pyloric sphincter (mean diameter, 12.8 ± 7 mm) [129]. However, some authors relate that the cut-off size of pylorus cannot be determined exactly [130]. For example, Timmermans [131] stated that the size of GRDDSs can vary from 7 to 10 mm, while Kholia and Davis [132] noted that sizes up to 15 mm in diameter were not sufficient to provide gastric retention.

According to Klausner et al. [133], the expandable systems should combine a number of characteristics, of which retention in the stomach is cited after facilitated oral administration, high degradability of the dosage form preferentially in the stomach, no interference in the gastric motility, prolongation of the shelf-life, among others. These systems, also called plug type systems [134] receive this classification precisely because after administration they undergo a great expansion of their structure, which may be related to both a high swelling or unfolding, blocking their passage through the pylorus. In this way, the permanence of the system in the stomach is increased until drug release begins with subsequent reduction in size as the system is evacuated through the GIT.

It is important to point out the three basic configurations associated with expandable GRDDSs that make clear the understanding of its gastroretention mechanism. First one, an initial state represented by expandable GRDDSs reduced dimensions that facilitate oral administration, followed by a second state represented by its maximum state of expansion which is responsible for gastroretention, and then, the third and last state of minimal dimensions related to the release and depuration of the drug from the system [101,133,135].

Several mechanisms of gastroretention for expandable GRDDSs have been explored, but it should be mentioned that they all show many advantages and also disadvantages over each other [134]. Two basic mechanisms are responsible for the expansion of structures, which are swelling and unfolding, the former being related to an osmotic effect, and the latter to shape memory.

The main mechanism for swelling and drug release is diffusion through hydrophilic polymers, which can absorb water from the gastric fluid through the appearance of pores in the device surface, creating channels along its entire length, leading to a capillary effect [134]. In unfolding systems, the previously large system (uncompressed) with adequate mechanical properties is folded (compressed) to just open after reaching the stomach when capsules are dissolved in the gastric juice [133]. Different geometrical shapes have been attributed to unfolding systems, such as tetrahedron, ring or planner membrane [15].

A recurrent problem that can occur with unfolding systems is the fact that their shape memory is often not sufficient, in addition to being influenced by storage stress, and may lead to performance problems in the stomach [136]. Some strategies have emerged to overcome

| Polymer(s)/drug | Objective | Methods | Results | Refs. |
|-----------------|-----------|---------|---------|-------|
| and Poly-x-lypo CS nanoparticles and Eudragit-enabled tablets (PXLNET)/levodopa | system as well as nano-enabled gastroretentive levodopa delivery system. | Eudragit and CS were dispersed in HCl. After levodopa (L-dopa) and benzerazide were added into dispersion, lecithin was dissolved in chloroform and added to the L-dopa-loaded polymeric solution. TPP dissolved in acetic acid was added under stirring and the nanoparticles produced were thereafter frozen and lyophilized. IPB - Dissolved methacrylate copolymer was added to the NaCMC solution under vigorous stirring with locust bean addition and nanoparticles incorporation. PXLNET was achieved by incorporating L-dopa-loaded nanoparticles into IPB with other components and compressed directly with carver hydraulic press. | mucoadhesive, the adhesion force and adhesion work for IPB matrices were found to be significantly more than the values observed for a PXLNET matrix. IPB presented triple mechanism of gastroretention (high density, swelling and gastro-adhesive). Levodopa release from IPB exhibited a more linear profile than Madopar® HBS and Sinemet® CR. In pH = 1.5 medium, IPB matrices released 90% of levodopa in 24 h vs Madopar® HBS 100% of the levodopa was released by the 16th h. In comparisons to the conventional dosage forms, both matrices have exhibited constant delivery over a prolonged time period and L-dopa-loaded and IPB matrices were the best fit for zero-order release. | a ADMadopar® HBS - hydrodynamically balanced system Madopar (levodopa+benzerazide). b Sinemet® CR - controlled release Sinemet (carbidopa = levodopa). |
this problem, mainly by exploiting polymers that do not present plastic deformation, but only elastic deformation.

Other forms of expandable GRDDSs have been reported, such as the use of hydrogels that swell to the significant extent primarily as a function of polymer content, such as those made of polyvinyl pyrrolidone cross-linked with albumin [137]. Superporous hydrogels (those with pores of a size of hundreds of micrometers) have been also used as expandable GRDDSs. In this case, the high porosity of these systems has been related to a very rapid rate of swelling (generally 1 min and an increase of up to 1000– initial weight), as well as the ability to absorb a greater volume of liquids, both factors contributing effectively to a fast size-increasing [138].

Regardless of the mechanism responsible for the expansion, after suffering a large increase in size, these systems must have adequate mechanical properties so that immediate erosion does not occur as a result of loosening the polymer chains together with the intense peristalsis of the stomach. Thus, the extensive expansion for swelling GRDDSs, for example, must be attributed to the presence of physical/chemical crosslinking in the hydrophilic polymer network. These crosslinks prevent the dissolution of the polymer and hence maintain the physical integrity of the dosage form [139].

Despite the effectiveness of these systems in increasing the residence time in the stomach, they must have some properties to ensure patient safety. Therefore, such devices should not interfere with gastrointestinal motility, should not irritate the stomach mucosa due to its local retention, the device must have rounded edges, and administrations should only be performed after the biodegradation of the previous system present in the stomach to avoid accumulation with high doses.

As it is known, alendronate – a drug belonging to the class of bisphosphonates – is the drug of choice for treating osteoporosis, acting mainly through specific inhibition of bone resorption. However, its bioavailability after oral administration is significantly low (<1%) due to reduced intestinal absorption and the fact that when administered with food it leads to the formation of complexes that cannot be absorbed [140]. Although alendronate is mainly absorbed in the upper portions of the GIT, the short transit time further contributes to low absorption and bioavailability. Another marked limitation is the local irritation caused in the upper portions of the GIT. Thus, treatment requires daily intake of the GIT, the short transit time further contributes to low absorption and bioavailability. Moreover, alendronate is also known to decrease the bioavailability due to the presence of positive charges in its structure. Several studies have combined different strategies for development of gastroretentive systems, in order to obtain a larger time of gastric residence [142].

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Although there are successful works, few systems with CS using the magnetic approach have been reported in literature. Magnetic hollow hydrogels of rosiglitazone loaded with CS were synthesized using glyoxal as a crosslinking agent for Gupta et al. [145]. The authors explored CS precisely because it is a polyelectrolyte that gives rise to pH-dependent hydrogels which, when in contact with acidic solutions, show a high swelling due to the presence of positive charges in the network, expanding its volume. Swelling rates were dependent on the extent of crosslinking, so that the higher the amount of crosslinking agent, the lower the swelling ratio. Ionic strength was also another factor that affected swelling, which ranged from 11% for HCl solution (pH 1.2) with 1 M NaCl to 156% for lowest ionic strength (0.0001 M).

4.5. Magnetic systems

Among the strategies used to increase the gastric residence time, magnetic systems are the only gastroretentive delivery systems based on the attraction between two magnets. This approach contain a small internal magnet present in dosage form, and an external magnet attached to the abdomen over the position of the stomach, which retains the system in the gastric region [101,139,146]. The extracorporeal magnet allows the control of gastrointestinal transit of the dosage form for a prolonged period of time [139].

Ito et al. [147] described the first magnetic system in 1990, performed for application to targeting therapy for esophageal cancer. Magnetic granules containing ultrafine ferrite (γ-Fe₂O₃) were guided to the esophagus with an external magnet for the initial 2 min and almost the entire amount of granules were retained in the region after 2 h of administration [147]. Although there are successful works, few systems with CS using the magnetic approach have been reported in literature. Magnetic hollow...
spheres coated with multilayer of CS/PAA developed by Zhang et al. [118] showed a sustained drug release and a strong magnetic response to magnetic field, which should provide the drug targeting to a desired tissue and/or organ through the external magnetic field [148].

The main advantages of magnetic systems are absolute drug targeting for target organs and tissues, the increase absorption and bioavailability of encapsulated drug, as well as the reduction of the concentration of the drug at non target sites [149]. However, the applicability of these systems depends on an external magnet, which must be positioned with high precision degree to allow drug release in the appropriate place, without compromising patient compliance [101,139]. Moreover, magnetic technical is expensive and requires specialized manufacturing, correct position degree on the stomach is very difficult to achieve and the magnets must have relatively constant gradients in order to avoid overdosing of toxic drug [149].

4.6. Density drug delivery-based systems

4.6.1. Floating drug delivery systems

Specifically, the design of controlled drug delivery systems (CDDSs) with efficient action on GIT is still a challenge due to the inability to restrain DDSs in specific regions of the GIT as well as the dependence of the unpredictable gastric emptying time [150]. These drawbacks can be overcome by GRDDSs, showing great potential for improving the bioavailability and site-specific absorption of drugs [150,151]. Among several GRDDSs approaches, floating drug delivery systems (FDDSs) are of particular interest due to their local active and narrow drugs absorption window for specific area.

FDDS was first reported by Coupe [152] and comprises a system with density lower (usually lower than 1 g cm\(^{-3}\)) than gastric fluids that remain buoyant in the stomach for prolonged time and, consequently, leading to slow drug release at the specific rate. Additionally, gastric retention time is increased as well as the adsorbed drug concentration [150,151,153]. FDDS can be designed and classified according to the dosage forms and mechanism of buoyance: (1) single unit floating dosage forms comprising a) effervescent and b) non-effervescent systems; (2) multiple unit floating dosage forms comprising a) effervescent, b) non-effervescent and c) hollow microspheres systems and, finally, (3) raft forming systems [79,151,154].

4.6.1.1. Single unit dosage systems. Single unit dosage systems are easier fabricated than multiple unit dosage forms. However, the application of single unit dosage forms are less effective due to their all or nothing emptying process from stomach generating distinct bioavailability and high drug concentration delivered in others sites causing irritation [79,154].

Effervescent systems (or gas-generating systems) can be prepared by using swellable polymers and compounds that can generate gas bubbles [79,135,154]. The most common useful swellable polymers as matrix in these systems are methylcellulose, hydroxypropylmethylcellulose (HPMC), hydroxyethylcellulose (HEC), hydroxypropylcellulose (HPC), sodium carboxymethylcellulose (NaCMC), polyacrylated polymers, polyvinyl acetates, polycarbonates, agar, CS and alginate [79,135,154]. As effervescent compounds sodium bicarbonate, tartaric acid and citric acid are the most commonly chosen [79,135,154]. The mechanism of action occurs when the fabricated system (swellable polymer + gas generator compounds + drug) is placed in contact with the fluid gastric leading to CO\(_2\) elimination while the polymer swelling with water. The combination of these actions generate effective density less than gastric fluid increasing the time buoyancy besides to favor the controlled and site-specific drug delivery as shown in Fig. 7 [79,155].

Jiménez-Martínez et al. [156] designed effervescent single unit dosage systems (SUFS) based on the introduction of captopril into Metolose SH 4000 SR (matrix) containing sodium bicarbonate (NaHCO\(_3\)) as gas generator. Properties as compounds proportion (Metolose and sodium bicarbonate) and distinct compaction pressures can influence floating and drug release from proposed matrix. \textit{In vitro} assays showed that at lower pressure (55 MPa) floating time higher than 8 h were achieved. The system density decreases by reducing the compaction pressure.
and increasing the sodium bicarbonate content. On the other hand, authors displayed that increasing sodium bicarbonate and polymer contents favor the reduction of drug dissolution rate from this system. Tadros [157] prepared effervescent SUSs using HPMC K15M and/or sodium alginate as release-retarding polymers and NaHCO3 or calcium carbonate (CaCO3) as a gas former agents containing ciprofloxacin hydrochloride (Cipro HC). The tablet formulations (F7: HPMC K15M (21.42%, w/w), Na alginate (7.14%, w/w), NaHCO3 (20%, w/w) and F10: HPMC K15M (21.42%, w/w), Na alginate (7.14%, w/w), CaCO3 (20%, w/w)) exhibited good floating total and lag time, swelling ability, adhesion and sustained drug release rate acting as a promising GRDDS.

On the other hand, non-effervescent systems (such as hydrodynamically balanced systems, HBS) present one or more gel-forming or high swellable polymers. This system is prepared by mixing drug and polymer generally encapsulated by gelatin. The mechanism comprises fast dissolution of the capsule in gastric fluid followed by swelling of the polymers producing a buoyance gelatin mass with density lower than 1 g cm−3. The formed gelatinous barrier retains the capsule shape, avoiding the system disintegration promoting controlled drug release. The gelatinous surface erosion favors water permeation into inner layers preserving the surface hydration and the air trapped by the swollen polymers promotes buoyancy to the dosage forms, as depicted in Fig. 8 [79,135,151]. In addition, drug release is controlled by diffusion and dependent on the gel barrier erosion/dissolution. If the formulation is composed by fatty or viscous excipients, low-density systems could be achieved, decreasing water penetration and erosion process, which can reduce drug release rates [107,132].

Soni et al. [158] developed HBS systems by using natural and biodegradable polymer crushed puff rice (CPR, as buoyance agent) combining with auxiliary polymers as high molecular weight CS (HMWCS, as swellable polymer) and HPMC (as gel-forming agent) as a single-unit floating capsule for Metoprolol Succinate (MS) sustained delivery. Authors reported that all evaluated samples displayed zero-order kinetics followed by Fickian diffusion model. This means that drug release from these HBS systems are controlled by drug diffusion through the gel barrier. Although the variation of pH and polymers could be evaluated, authors concluded that CPR associated with HPMC and HMWCS comprising a potential single-unit HBS systems for specific sustained release of hydrophilic drugs showing absorption window upper GIT.

The main drawback of single layer HBS consists on the dependence of the air arrested in the dry mass and the characteristics of the swellable polymers [135]. Considering the above mentioned, some approaches [135,159–161] have been developed to improve the efficiency of these dosage forms such as the design of bilayer formulations, as displayed in Fig. 9. In these dosage forms, one layer promotes the fluctuation in gastric liquid whereas the other one control the drug release [135]. Oth et al. [161] fabricated a bilayer floating dosage unit containing misoprostol aiming to treat gastric and duodenal ulcers. As shown in Fig. 9A, both layers are composed by the same swellable polymer (in this case HPMC) but only one containing the misoprostol allowing this a bifunctional system performed buoyancy and drug delivery independently in gastric media. Authors demonstrated that the production of large capsules lead to an increasing in gastric retention time (GRT). In vivo studies exhibited GRT dependent to the meal regimen. After a single meal, GRT was higher than 3 h whereas after several meals, the GRT was higher than 10 h. He et al. [160] produced bilayer floating tablet formulation (Fig. 9B) containing the combination of two distinct drugs such as metformin hydrochloride (MH) and pioglitazone hydrochloride (PG) to treat type 2 diabetes mellitus (T2DM). This formulation comprises the incorporation of each drug into two separate layers in order to achieve sustained MH release followed by immediate PG release. The bilayer tablet was fabricated by using wet granulation method with HPMC E5 for both layers as matrix and then compressed into tablet. In vitro assays showed buoyancy time up to 24 h and floating lag time of 5 min besides sustained MH release controlled by diffusion manner (for 12 h) and fast PG release. In vivo tests performed in dogs suggested good absorption of PG and the enhancement of MH bioavailability with steady plasma concentration, decreased maximum plasma concentration and reduced time of maximum concentration appears as effective and promising GRDDS to treat T2DM. Krögel and Bodmeier [159] developed multifunctional floating drug delivery systems using HPMC as hydrophilic polymer containing drugs (chlorpheniramine maleate or ibuprofen). One of the designed GRDDSs consists of two tablets (HPC + drug) fixed within two orifices of the cylinder (Fig. 9C) with air entrapped in the middle of the two tablets resulting in low density and, consequently, buoyance to the proposed system. Authors reported interesting GRDDSs that provides an enhancement on drug release as a function of HPMC viscosity and content, aqueous drug solubility and surface area of the matrix.

Harrigan [162] described the fabrication of intragastric FDDSs with drugs present inside microporous compartment with pores along its top and bottom walls. In order to prevent a direct contact with the gastric surface, the peripheral walls of these systems are sealed aiming to avoid the direct contact of undissolved drugs with gastric mucosal surface. The containing entrapped air in the microporous compartment promotes buoyancy over gastric media. The gastric fluid permeates...
the apertures favoring the drug dissolution and absorption in the stomach [139,163].

4.6.1.2. Multiple units dosage systems. Multiple units floating systems (MUFS) appear as an interesting way to overcome high variability of gastrointestinal transit time favoring a regular absorption. Beyond that, fast and high drug concentration release could be avoided and, consequently, gastrointestinal tract irritation [79,154]. Hooda et al. [122] reported the preparation of MUFS based on CS and sodium tripolyphosphate (TPP) via ionotropic gelation method containing Ranitidine hydrochloride (RHCl) as microspheres GRDDSs. RHCl has a narrow absorption window and is mainly absorbed in the proximal areas of GIT. Thus, conventional sustained-release dosage form reaches the colon, where it gets metabolized, resulting in low absorption and poor bioavailability (52%). In this way, Hooda et al. [122] proposed to design microspheres GRDDS based on chitosan (CS) and sodium tripolyphosphate (TPP) via ionotropic gelation containing RHCl. In vitro buoyance and mucoadhesive tests suggested that these microspheres present interesting floating (at least 12 h of buoyance) and bioadhesive properties. The above mentioned properties allowed the fabricated microspheres adhere to the gastric mucosal surface remaining prolonged period in stomach ensuring RHCl stability in gastric environment which eventually leads to better bioavailability at much lower dose. Additionally, in vitro drug release assays displayed a zero-order model suggesting that the drug (RHCl) transport across polymeric matrix occurred by a Fickian diffusion process [164].

Talukder [165] fabricated MUFS based on crosslinking reactions among sodium alginate, calcium chloride and low methoxylated pectin (anionic polysaccharide), in aqueous solution. The alginate beads show porous structure that promotes buoyancy for over 12 h besides increasing the residence time in gastric tract substantially (>5.5 h) [139,163]. Dey et al. [166] designed floating and mucoadhesive beads based on sodium alginate and hydroxypropyl methylcellulose (HPMC) as matrix polymers and CS as coating polymer. Sunflower oil was entrapped in this system to improve floating time. All these polymers were chosen to encapsulate amoxicillin trihydrate aiming to treat Helicobacter pylori infection. The floating and mucoadhesive system were prepared by ionotropic gelation method and CS used as polymer barrier promoting bioadhesion. Authors reported that these beads as interesting GRDDSs exhibiting good mucoadhesion properties, excellent efficiency of drug encapsulation increasing buoyance time and drug release for >24 h and 7 h, respectively [166].

As multiple units floating effervescent systems, Jyang et al. [167] fabricated a blend of Eudragit L100 and Eudragit RLPO containing dipyridamole (antithrombotic agent) via solid dispersion technique and afterwards, the dispersion was incorporated into alginate beads prepared by ionotropic gelation method using CaCO3 as gas-forming agent. Buoyancy tests were performed showing that 92% of alginate beads remained floating after 9 h suggesting that drug encapsulated can be retained in the stomach indicating specific drug delivery. In vitro release studies displayed the controlled dipyridamole release using the proposed formulation (ratio of dipyridamole:Eudragit L100: Eudragit RLPO = 1:2:3) with 70% of release up to 9 h. In vivo results exhibited higher (about 2.52-fold) bioavailability for the FDDSs than commercial tablets corroborating this platform for stomach-specific drug delivery by the oral route.

4.6.1.3. Hollow microspheres systems. Hollow microspheres (or microballoons) comprise low-density polymers with low-density core relating to the shell. Most of them are MUFS fabricated by emulsion drying.
solvent diffusion method that entraps oil or air in hollow core [135,163]. Polymers such as polycarbonate, Eudragit S, cellulose acetate, calcium alginate, agar, CS and low methoxylated pectin are commonly used to fabricate hollow microspheres and foam-particles [168,169]. Great buoyancy and drug release from these systems depend on polymers amount, the plasticizer polymer ratio and the solvent used for formulations [135]. Fig. 10 displays hollow (low-density) microalloons (Fig. 10a) and foam-particles (Fig. 10b) obtained by solvent evaporation.

Pawar et al. [153] formulated microalloons as GRDDS based on mucoadhesive and floating mechanisms to deliver norfloxacin (NFX). These systems were fabricated by using HPMC and ethylcellulose (EC) as core matrix by solvent evaporation method. CS coating was prepared via ionotropic gelation method to improve mucoadhesion properties and increasing gastroretention time of the microalloons. Good in vitro buoyance was achieved for these systems. Pharmacokinetics results displayed an enhancement of NFX bioavailability for these systems. In vivo assays showed that the microcarriers could prolong half-life of NFX and increasing plasma drug concentration.

4.6.1.4. Raft-forming systems. Another system that has attracted great attention in the administration of drugs to treat gastrointestinal infections is the raft forming systems. The mechanism of raft preparation consists on the formation of viscous cohesive gel containing entrapped CO2 gas in contact with gastric fluid [79,135]. Usually, the formulation containing gel forming agent and antacids such as alkaline bicarbonates or carbonates to reduce gastric acidity. The buoyance of raft systems occurs by CO2 formation and elimination, acting as a blocker to preclude reflux of gastric media [79,135]. Fig. 11 depicted a schematic illustration of raft forming systems as barrier in specific stomach region.

Raft forming systems usually form a low-density layer on gastric media with high surface area regarding tablets, favoring drug release and the bioavailability enhancement. Beyond that, these GRDDSs show fast buoyance than other floating dosage forms, improving therapeutic efficacy been easily administrated (once a day) to the patient [139]. On the other hand, the stability of these GRDDSs are susceptible to chemical modifications, like: oxidation and hydrolysis. Exposure to the temperature and pH variation as well as types of radiation (ultraviolet-visible, X-ray, among others) could also promotes modification in these systems [139]. El Nabawari [170] demonstrated the preparation of a floating raft system with Mebeverine HCl (MbH) using sodium alginate, HPMC K100M and precirol. The optimized formulation (FRS-11: sodium alginate 3%/HPMC K100M 1%/precirol 2%) displayed floating lag time and total buoyance time near to 15 s and 12 h (in simulated gastric media), respectively. In contrast, FRS-11 formulation exhibited sustained drug release of 82% up to 8 h showing good MbH bioavailability compared to the commercial product. These results suggest the designed raft-forming system as a potential GRDDS to prolong drug action aiming clinical trials [170,171].

4.6.2. High-density drug delivery systems

Different of low-density systems, dosage forms with density next to 3 g mL\(^{-1}\) can sink to the bottom of the stomach to withstand in vivo peristaltic movement remaining intact independently of the GIT disturbance as depicted in Fig. 12 [110]. By using high-density systems (from 2.5 to 3.0 g mL\(^{-1}\) the gastric retention time can increase up to 25 h [155]. These systems are formulated with inorganic compounds such as barium sulphate, zinc oxide,
iron powder and titanium dioxide increasing the density of the dosage form. However, up to now no effective action of these systems were detected in human just in pre-clinical assays [101,151,155]. In addition, the main drawbacks regarding high-density systems consist on the fabrication of pellets with high drug concentration and achieve systems with a density next to 3 g mL$^{-1}$ [101].

5. Conclusions

CS is a biopolymer with a wide applicability, highlighted as a gastroretentive drug delivery system-forming material, because it brings together special and desirable biochemical properties in this type of system. These characteristics enable the fabrication of CS-based drug delivery systems for the stomach with mucoadhesive, floating, expandable, biocompatible, nontoxic and biodegradable properties. Advances in gastroretentive drug delivery system development technology associated with the versatility of CS will be able to deliver increasingly efficient and selective dosage forms with new mechanisms of gastroretention, as well as capable of increasing the capacity of drug internalization using lower drug doses to reach the therapeutic range, ensuring greater safety.

Declaration of competing interest

The authors declare no conflict of interest.

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