REVIEW ARTICLE

Biotechnological production and application of fructooligosaccharides

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Abstract

Currently, prebiotics are all carbohydrates of relatively short chain length. One important group is the fructooligosaccharides (FOS), a special kind of prebiotic associated to the selective stimulation of the activity of certain groups of colonic bacteria. They have a positive and beneficial effect on intestinal microbiota, reducing the incidence of gastrointestinal infections and also possessing a recognized bifidogenic effect. Traditionally, these prebiotic compounds have been obtained through extraction processes from some plants, as well as through enzymatic hydrolysis of sucrose. However, different fermentative methods have also been proposed for the production of FOS, such as solid-state fermentations utilizing various agro-industrial by-products. By optimizing the culture parameters, FOS yields and productivity can be improved. The use of immobilized enzymes and cells has also been proposed as being an effective and economic method for large-scale production of FOS. This article is an overview of the results considering recent studies on FOS biosynthesis, physicochemical properties, sources, biotechnological production and applications.

Keywords

Fructooligosaccharides, fructosyltransferase, functional foods, inulin, prebiotics

History

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Introduction

Modern nutrition focus on a relationship between food quality and health promotion. For this reason, this discipline is now oriented to provide information on foods with an emphasis in their nutrients and bioactive constituents (García-Casal, 2007). These compounds give additional benefits because of their consumption, conceptualizing them as functional foods, forcing several changes from their manufacture, including nutritional, microbiological, technological and sensorial qualities (García-Casal, 2007).

Bioactive compounds present in functional foods, are mainly phytochemicals with complex and diverse chemical structures, such as carotenoids, isoflavones, cumestans, polyphenols, phytostanols, conjugated linoleic acid and epigallocatechin gallate (EGCG), among thousands of chemical compounds with beneficial biological activity (Alvídez-Morales et al., 2002). There is significant evidence for a reduction of health risks due to regular consumption of these bioactive compounds, including cardiovascular diseases, cancer, osteoporosis, hyperlipidemia and neurodegeneration (Alvídez-Morales et al., 2002).

Although, there is not universal consensus about the term of functional food, it is applied to those foods with one or more bioactive components which satisfactorily demonstrate a benefit in one or more determined functions of the organism (Mussatto & Mancilha, 2007). The fundamental effects of functional foods are excellent alternatives to improve health condition and well-being and/or to reduce the risk of some diseases (Mussatto & Mancilha, 2007). A functional food must be a food primarily and must demonstrate its effects in amounts that normally are consumed in the diet. Within the ample context of functional foods, prebiotics have been intensely studied, due to their diversity and magnitude of beneficial effects on health that their consumption generates (Sabater-Molina et al., 2009).

Prebiotics

The concept of prebiotics was introduced by Gibson & Roberfroid in 1995 with a slightly alternative approach which consists of regulation of the gut microbiota (Gibson & Roberfroid, 1995). At present according to the FAO and several researchers, ‘‘a prebiotic is a selectively fermented ingredient that allows specific changes, both in the composition and/or activity in the gastrointestinal microbiota that confers benefits upon host wellbeing and health’’ (Al-Sheraji et al., 2013; Charalampopoulos & Rastall, 2012; Dominguez et al., 2013; Gibson et al., 2004, 2010; Pineiro et al., 2008; Sarbini & Rastall, 2011; Slavin, 2013; Walton et al., 2013).

These compounds are the trophic substrate of probiotics, generally as a strategy to improve balance, growth and activity of the various kinds of intestinal bacteria including bacteria of the colon (Dominguez et al., 2013;
Gibson et al., 2004; Macfarlane et al., 2008; Sarmiento Rubiano, 2006; Silveira Rodríguez et al., 2003).

Thus, to be a nutritional ingredient classified as prebiotic, it must fulfill the following requirements: a) low sensitivity to hydrolysis by saliva, pancreatic and intestinal enzymes or absorption along the gastrointestinal tract; b) constitute a fermentable substrate for the intestinal microflora as established by scientific studies as are: inulin, fructooligosaccharides (FOS), galactooligosaccharides (GOS), isomaltooligosaccharides (IMO), xylooligosaccharides (XOS) and soybean oligosaccharides (SOS) and to selectively stimulate growth and metabolism of one or more beneficial bacteria to the colon c) to modify the composition of the colon flora, facilitating development of beneficial species, and d) to induce beneficial effects into the lumen or are systemically relevant for the health of the individuals (Charalampopoulos & Rastall, 2012; Dominguez et al., 2013; Gibson, 1999; Scheid et al., 2013; Slavin, 2013).

Non-digestible carbohydrates (oligosaccharides and polysaccharides), some peptides and proteins, and certain lipids (esters and ethers) are considered to be prebiotics. Due to their chemical structure, these compounds are not absorbed into the gastrointestinal tract and are not hydrolyzed by human digestive enzymes due to their configuration β in C₂ (Gibson et al., 2004). These compounds can be named colon foods, since they enter into the colon where they are released, allowing their absorption. Moreover, the short-chain carbohydrates increase colonic absorption of zinc, calcium and magnesium ions when causing water attraction by osmosis, in which such minerals are dissolved, providing energy, metabolic substrates and essential micronutrients to the organism (Pérez Conesa et al., 2004; Roberfroid et al., 1998, Silveira Rodríguez et al., 2003).

Carbohydrates can be classified according to their degree of polymerization in oligosaccharides (between 2 and 10 units of monosaccharides) and polysaccharides (more than 10 monosaccharides) as indicated by IUB-IUPAC terminology (Nomenclature, 1982). Englyst & Hudson (1996) proposed the name of short chain carbohydrate for a new nutritional carbohydrate group that included oligosaccharides and the smallest polysaccharides. The main available oligosaccharides are carbohydrates in which the monosaccharidic unit is fructose, galactose, glucose, and/or xylose (Crittenden & Playne, 1996; Delzenne & Roberfroid, 1994), including in this important group the FOS.

**Fructooligosaccharides (FOS)**

FOS are non-digestible carbohydrates that represent one of the major classes of bifidogenic oligosaccharides, FOS is the common name for fructose oligomers, chemically composed mainly of chains of fructose units with a terminal glucose molecule unit linked by glycosidic bridges β-(2-1) (Chacón-Villalobos, 2006; Monsan & Ouarné, 2009; Sabater-Molina et al., 2009). Also, they are known as fructans, oligofructans, glupofructans, inulins or oligosaccharides, where their structure is formed by repetitive unions of disaccharides such as sucrose, inulobiose and levanoibiose (Chacón-Villalobos, 2006). FOS are reserve phytochemicals present in many plants and vegetables as reserve carbohydrates such as Jerusalem artichoke, onion, asparagus, chicory, leek, garlic, wheat, yacon, tomatoes, banana and honey (Monsan & Ouarné, 2009; Mussatto et al., 2009a). According to their structural differences, there are four important groups of FOS: inulin, levans, mixed levans and neo-FOS (Monsan & Ouarné, 2009).

Inulin is a fructooligosaccharide with a polymerization degree of 2 to 60 monomers of sucrose (Murphy, 2001; Roberfroid, 2007b; Watzl et al., 2005). It has been defined as polydisperse fructans, constituted mainly, but not exclusively, of β-(1-2)-fructofuranosyl linkages. Inulin can be obtained from several plant families (mono and dicotyledonous). Nevertheless, only chicory (*Chicorium intybus*) is used to produce inulin at the industrial level, this process is similar to that used for sugar production from sugar beet. Native inulin is processed and transformed into FOS or short chain fructans (scFOS) with a degree of polymerization between 2 and 10 (normally 5) as a result of partial enzymatic hydrolysis with inulinas (Gibson & Rastall, 2006).

Oligofructose is chemically defined as linear non-digestible oligosaccharide of β-(2-1)-linked fructose fraction with a terminal glucose residue unit of oligosaccharides with a degree of polymerization between 2 and 20. Due to the structural conformation of their osidic bridge (3 2–1), this resists the hydrolysis by human alimentary enzymes (Roberfroid, 1993). Oligofructose is the strict definition to oligosaccharides obtained naturally from the enzymatic hydrolysis of inulin that can otherwise be obtained by enzymatic synthesis (transfructosylation) using sucrose as a substrate and consists of a mixture of fructosyl chains (maximum of 5 units), with terminal glucose and fructose FOS of short chain (Roberfroid, 2002, 2007a, Venter, 2007; Walton et al., 2013; Watzl et al., 2005).

Short chain FOS (scFOS) are a mixture of oligosaccharides containing of glucose linked to fructose units: bonds between fructose units are β (2-1) forming the fructooligosaccharides: 1-kestose, nystose and 1-fructofuranosyl-nystose (Figure 1; Roberfroid & Delzenne, 1998; Sánchez et al., 2008; Vega & Zuniga-Hansen, 2014). Kestose is formed by addition of a fructose molecule to one of sucrose. Nystose is formed by the later addition of a fructose molecule, while the addition of another molecule of fructose will give rise to the formation of fructofuranosyl-nystose (Dorta et al., 2006; Rivero-Urgell & Santamaria-Orleans, 2001).

Depending on the linkage type between the monosaccharide residues, different types of FOS series can be distinguished. Neo-FOS consists mainly of neo-kestose (neo-GF₃) and neo-nystose (neo-GF₅), in which fructosyl units are β-(2-6)-linked to the fructofuranosyl residue of sucrose (Chen et al., 2011; Kilian et al., 2002; Lim et al., 2007; Linde et al., 2012; Park et al., 2005; Plou Gasca et al., 2009). However, neo-FOS have not been widely explored, probably because they are not produced by fructofuranosidases in microorganisms (Alvaro-Benito et al., 2007; Chen et al., 2011; Ghazi et al., 2007) or they represent only a minor biosynthetic product (Chen et al., 2011; Farine et al., 2001).

**Physico-chemical properties of FOS**

FOS are compounds soluble in water and their sweetness oscillates between 0, and 6 times to that of sucrose, having the
Clostridium observed that consumption of FOS decreases populations of (*Cummings et al.*, 2001; *Rastall & Maitin*, 2002). It was also growth increase when FOS were used as a carbon source (*Macfarlane et al.*, 2006; *Reyed*, 2007).

The normal microbial accounts (*Cummings et al.*, 2001; *Rastall & Maitin*, 2002). After comparing healthy humans with daily ingestion of low doses of FOS (5–20 g/d) against placebos with high amounts of sucrose, it was found that bifidobacteria increased in order of magnitude above of the normal microbial accounts (*Cummings et al.*, 2001; *Rastall & Maitin*, 2002). During fermentation, prebiotics can promote some specific physiological functions through liberation of metabolites, especially short chain fatty acids (acetate, propionate, butyrate, lactate, etc.) to the intestinal lumen (*Olvera et al.*, 2007b). Short chain fatty acids may act directly or indirectly on intestinal cells and can participate in control of several processes like mucosal proliferation, inflammation, colorectal carcinogenesis, mineral absorption and nitrogen compounds elimination. This FOS property is recognized in several European countries and named as the prebiotic effect. Among the FOS clearly identified with this beneficial effect are kestose and neokestose (*Gibson & Rastall*, 2006).

**Prebiotic effect of FOS**

In the last decades, the effect of bacterial microflora on human health has been one of the main research topics. The large intestine contains more than 500 different types of bacteria, which contribute to an important number of biological functions. Recent studies in *vivo* and *in vitro* with *Bifidobacterium longum*, *B. infantis* and *B. angulatum* revealed an important growth increase when FOS were used as a carbon source (*Cummings et al.*, 2001; *Rastall & Maitin*, 2002). It was also observed that consumption of FOS decreases populations of *Clostridium* and reduces production of flatulences (*Cummings et al.*, 2001; *Rastall & Maitin*, 2002). After comparing healthy humans with daily ingestion of low doses of FOS (5–20 g/d) against placebos with high amounts of sucrose, it was found that bifidobacteria increased in order of magnitude above of the normal microbial accounts (*Cummings et al.*, 2001; *Macfarlane et al.*, 2006; *Reyed*, 2007).

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**Biotechnological production of FOS**

Currently, the main route of chemical synthesis of glycosylated compounds is unattractive due to high chemical sensitive of sugars, which consequently brings the necessity of protection and lack of protection of substrates and products (*Tomotani & Vitolo*, 2007). In addition, most of the used chemical catalysts are toxic with low specificity, which limits their application in the pharmaceutical and food industries (*Tomotani & Vitolo*, 2007). Alternatively, the increasing presence of biocatalysis in different processes of chemical synthesis has opened the possibility to develop selective, efficient and less aggressive alternative processes (*Nemukula et al.*, 2009).

FOS are produced by transfructosylation of sucrose which is carried out via the breaking of the β-(2-1)-glycosidic bond and the transfer of the fructosyl moiety onto any acceptor other than water, such as sucrose or a fructooligosaccharide. In the enzymatic synthesis of FOS, microbial enzymes with transfructosilase activity should be utilized. This synthesis is a complex process in which several reactions occur simultaneously, both in parallel and in series, because FOS are also potential substrates of fructosyltransferases (FTases) (*Vega & Zuniga-Hansen*, 2014). These enzymes are usually classified as β-D-fructofuranosidases (FFase, EC 3.2.1.26), with high transfructosylating activity or fructosyltransferase (FTase, EC 2.4.1.9) (*Maiorano et al.*, 2008b). These enzymes may be produced intra and extracellular by several microorganisms, including bacteria and fungi, as shown in Table 1. Fructofuranosidases production may occur by two kinds of processes: submerged fermentation (SmF) and solid-state
fermentation (SSF). Most research studies on the experimental conditions for FTase production have been conducted under submerged fermentation conditions, being mainly based on shake-flask experiments with some fungal strains. Fermentation parameters such as culture medium, aeration, agitation, pH and temperature must be established for each microorganism, but general conditions for the enzyme production are well known.

Most enzymes for industrial use are produced by SmF but the growing trend of using SSF represents a great alternative to small-scale occurring extracellular. Some advantages of SSF processes are: high volumetric productivity and enzyme concentration, low production costs and energy consumption, the risk of contamination is minimal but the most important are more stable products (Balasubramaniem et al., 2001; Longo et al., 2008; Sangeetha et al., 2004).

Various agro industrial by-products are considered good substrates in SSF processes, especially for enzymes production (Graminha et al., 2008; Rodríguez Couto & Sanromán, 2005). Some of these substrates include cereal brans, sugarcane bagasse, wheat straw, rice husk, soybean shell, corn cobs, cassava waste, apple pomace and waste from tea and coffee industries (Sangeetha et al., 2004). There are many reports on FTase production using SmF and very few using SSF (Sangeetha et al., 2004). Table 2 summarizes some studies using SSF with agro-industrial wastes for the production of FTase.

SSF is defined as the cultivation of microorganisms on wet or semi-moistened solid media (Botella et al., 2007; Lateef et al., 2008b), supports can be inert and insoluble or substrates that can be used as a carbon and energy source (Rodríguez Couto & Sanromán, 2005). SSF is carried out in the absence or with minimum amount of free water trying to adapt the culture conditions of each microorganism (Rodríguez Couto & Sanromán, 2005). SSF is used to produce various chemicals and enzymes. This technique has several advantages such as more stable products and in high concentration, low catabolic repression, and aeration easy, requiring little energy production fermenters which can be used on small and large scale thereby decreasing pollution effluents (Hölker et al., 2004; Longo et al., 2008). Another advantage is that SSF is developed under low moisture content, which is a limitation for several microorganisms. Therefore, this kind of fermentation can be performed mainly by some fungi and yeast, and seldom by bacteria (Hölker et al., 2004; Longo et al., 2008). In addition, using SSF has the possibility of using mixed cultures and thereby exploit the synergism of metabolism between the microorganisms (Hölker et al., 2004).

Generally, the FTase is obtained by liquid or solid fermentation liquid and is used for production of FOS. However, new processes such as fermentation with biofilms have been developed to produce biotechnologically important molecules, combining the advantages of the solid fermentation and high productivity (Aziani et al., 2012). The growth of filamentous fungi in their natural environment is given by the direct association with the substrate, which is extremely important as this allows adhesion and spore germination to form mycelium (Aziani et al., 2012).

There are reports of FOS production by colonization of Aspergillus japonicus in synthetic media (Mussatto et al., 2009a). Colonization occurred in the holder during fermentation, and thus the production of FOS is influenced by the metabolic action of free and immobilized cells. Aziani et al. (2012) immobilized cells in an aqueous solution and not in the culture medium for the production of FOS (Aziani et al., 2012). This procedure has several advantages among which stands out the separation of the cells, recovery of products (FOS) of the fermentation broth and reuse of the catalysts (biofilm) employed for the production of FOS can reduce process costs (Aziani et al., 2012).

FTases are enzymes with potential to be used during the glycosylation process of molecules. Bacterial FTases generally have molecular weights between 45 and 64 kDa, although those produced by lactic acid bacteria usually have superior molecular weights (from 80 to 170 kDa) (Olvera et al., 2007a). Most of these enzymes are extracellular, i.e. they are secreted into the culture medium during the growth of bacteria. The fungal FTases have molecular weights between 60 and 75 kDa, although other enzymes with a higher molecular weight have also been reported (Maiorano et al., 2008a). In both fungal and plants FTases, six conserved regions are clear, in three of these are located the possible amino acids implied on catalysis. One of these regions allows

### Table 1. Microorganisms that produce FTase.

<table>
<thead>
<tr>
<th>Microorganism</th>
<th>Reference</th>
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<tbody>
<tr>
<td>Aspergillus oryzae CRF 202</td>
<td>(Sangeetha et al., 2004)</td>
</tr>
<tr>
<td>Rhizopus stolonifer LAU07</td>
<td>(Lateef et al., 2008a)</td>
</tr>
<tr>
<td>Rhodotorula sp</td>
<td>(Hernalsteens &amp; Maugeri, 2008)</td>
</tr>
<tr>
<td>Aspergillus japonicus ATCC 20236</td>
<td>(Mussatto et al., 2009b)</td>
</tr>
<tr>
<td>Rhodotorula dairenensis</td>
<td>(Gutiérrez-Alonso et al., 2009)</td>
</tr>
<tr>
<td>Penicillium expansum</td>
<td>(Prata et al., 2010)</td>
</tr>
<tr>
<td>Penicillium purpureogenum</td>
<td>(Dhake &amp; Patil, 2007)</td>
</tr>
<tr>
<td>Aspergillus phoenicis</td>
<td>(Rustiguel et al., 2011)</td>
</tr>
</tbody>
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### Table 2. Agro-industrial residues used as substrates in SSF for the production of FTase.

<table>
<thead>
<tr>
<th>Microorganism</th>
<th>Substrate</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. foetidus NRRL 337</td>
<td>Commercial apple pomace</td>
<td>(Hang et al., 1995)</td>
</tr>
<tr>
<td>A. niger NRRL 330</td>
<td>Sugarcane bagasse</td>
<td>(Balasubramaniem et al., 2001)</td>
</tr>
<tr>
<td>A. oryzae CFR 202</td>
<td>Cereal brans like wheat bran, rice bran and oat bran; Corn products like corn cob, corn bran, corn germ, corn meal, corn grits and whole corn powder (coarse); Coffee- and tea-processing by-products like coffee husk, coffee pulp, spent coffee and spent tea; Sugarcane bagasse, Cassava bagasse (tppi).</td>
<td>(Sangeetha et al., 2004)</td>
</tr>
<tr>
<td>A. japonicus ATCC 20236</td>
<td>Corn cobs, coffee silverskin and cork oak</td>
<td>(Mussatto et al., 2009b)</td>
</tr>
<tr>
<td>Rhizopus stolonifer LAU07</td>
<td>Cassava wastes</td>
<td>(Lateef &amp; Gueguim Kana, 2012)</td>
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</table>
the enzyme to be bound to sucrose (Kurakake et al., 2007). Because of the structural similarity between fungal and plant FTases, and that these enzymes hydrolyze sucrose (invertase) and other fructosides, they are classified within the 32 family of glycoside hydrolases (Wallis et al., 1997).

The properties of the microbial FTases may vary according to the microorganism and culture medium composition; specifically the carbon source may act as an inducer (Maiorano et al., 2008a). FOS are produced by the action of microbial and plant transfructosyltransferases (Sánchez et al., 2010), by two processes, generating products slightly different end products. In the first method, FOSs are obtained from the disaccharide sucrose using the activity of transfructosylation of the fungal fructosyltransferase enzyme (Park & Almeida, 1991). The FOS thus formed contains from 2 to 4 units of the fructose bond with connections β (2→1), a terminal remainder to-D-glucose, and among them it is possible to emphasize: 1-kestose (Glu-Fru2), 1-nystose (Glu-Fru3) and 1-fructosylnystose (Glu-Fru4). The second method is controlling the enzymatic or chemical hydrolysis of inulin. In this case, all fructosyl chains β (2→1) do not finish in a terminal glucose and the produced oligosaccharides mixture contains chains of fructo-oligomers longer than those produced by the process of sucrose transfructosylation (Crittenden & Playne, 1996). This product is known as oligo-fructose (Crittenden & Playne, 1996).

In presence of sucrose, reaction conditions and substrate concentrations, FTases are able to carry out several reactions. They can synthesize a polymer, transferring fructose to the growing chains or hydrolyze it to sucrose (Antošová et al., 2002; Maiorano et al., 2008a; Olvera et al., 2007b). When an outside molecule is added to the reaction medium, this molecule is called an acceptor, the enzyme may also transfer fructose, producing a fructosylated molecule also called a fructoside. An interesting mechanism regarding the reaction of these enzymes was proposed by Chambert et al. (1974). They suggested, from kinetic studies on initial velocity, that the behavior is of the Ping-Pong BiBi type (Figure 2). In this behavior, it is proposed that enzyme and sucrose form an enzyme-fructose intermediary (Ping), and the glucose is released (Pong). This complex interacts, for example, with a water molecule (Ping), and fructose is then transferred releasing a second product – fructose (Pong). This second product can be the growing chain of fructoses or only a fructoside (Olvera et al., 2007b; Vega & Zuniga-Hansen, 2014).

The biochemical mechanism used by these fructosylases occurs in two steps: the sucrose (glucose–fructose) enters to the active site where it interacts and a covalent bond between fructose and the enzyme is formed. It is known that this connection is with aspartic 86, being in this way that the enzyme unites covalently to fructose. In the second step, a new sucrose molecule is recognized at the active site, the site that was occupied before by glucose: fructoses together with D86 then are transferred. Aspartic 247 is used to stabilize the state of transition between these two steps (Kim et al., 1996).

Recent research on industrial enzymology has succeeded in producing large-scale FOS by enzymatic processes. Industrial processes for FOS production can be divided into two classes: a batch system using free enzyme and the second a continuous system with immobilized enzyme or cells.

Immobilization of enzymes for use in reactors allows for a high-enzyme load with high activity within the bioreactor, hence leading to high-volumetric productivities. This enables control of the extension of the reaction. Downstream processing is simplified, since the biocatalyst is easily recovered and reused. The product stream is with a biocatalyst where continuous operation and process automation are employed and substrate inhibition can be minimized. Along with this, immobilization prevents enzyme denaturation by autolysis or organic solvents, and can bring enhance thermal, operational and storage stabilization, provided the immobilization is adequately designed (Fernandes, 2010; Mateo et al., 2007; Sheldon, 2007).

Immobilization can be performed by several methods, namely, entrapment or microencapsulation, binding to a solid carrier, and cross-linking of enzyme aggregates, resulting in carrier-free macromolecules (Fernandes, 2010; Sheldon, 2007). For the large-scale production of FOS, β-fructofuranosidase has been immobilized on porous glass, porous silica (Hayashi et al., 1991, 1992, 1993) and ion-exchange resins (Yun & Song, 1996), gluten (Chien et al., 2001), polymethacrylate (Ghazi et al., 2005), macroporous beads (Tanriseven & Aslan, 2005), calcium alginate (Jung et al., 2011; Lin & Lee, 2008; Sheu et al., 2013; Yun et al., 1990), ambersite (Csanadi & Sisak, 2008), niobium and graphite (Alvarado-Huallancos & Maugeri-Filho, 2010). The immobilization of the FTase offers a lot of practical advantages, e.g. the easy separation of enzyme and product, the opportunity to realize a continuous process, the enhancement of volumetric productivity of the reactor, more stability to changes in pH and temperature than free FTase and increased operational stability.

Recent processes for FOS production are shown in Table 3 where it is compared to the batch and processes used to yield FOS production. The Company Meiji Seika began the industrial production of FOS from Aspergillus niger cells immobilized in calcium alginate and recently the company Cheil Food and Chemicals (Seoul Korea) developed a continuous process with immobilized cells of Aspergillus pullulan in a calcium alginate gel.

**FOS applications in the food industry**

In food industries, because chemical additives are becoming less and less welcome by consumers, there is an increasing
interest in the use of saccharidic natural substances known as prebiotic and bio-preservative FOS (Barreteau et al., 2006). The term biopreservative includes a wide range of natural products from both plants and microorganisms, which are able to extend the shelf life of foods, reduce or eliminate pathogenic microorganisms and increase the overall quality of food products. These natural occurring antimicrobials can be, for example, peptides such as bacteriocins or lipophilic substances such as essential oils (Cleveland et al., 2001). Compared to these two kinds of antimicrobial molecules, sugar molecules seem to be less investigated as potential food preservatives (Barreteau et al., 2006). Different functional properties of FOS are due to the difference in their chain lengths. FOS contribute to give body to dairy products and humectancy to soft baked products, decreases the freezing point in frozen desserts, provides crispness to low fat cookies and acts as a binder in nutritional or granola bars in much of the same way as sugar, but with the added benefit of fewer calories, fiber enrichment and other nutritional properties (Kaur & Gupta, 2002). Industries producing FOS commercially from the transfructosylation of sucrose or inulin hydrolysis are listed in Table 4.

Concluding remarks

It is important to emphasize that many research groups have indicated that deficiencies in the diet can lead to disorders and diseases, which can be avoided through an adequate intake of relevant dietary nutrients and functional molecules. FOS are generally used as Generally Recognized As Safe (GRAS) components of functional foods and play a key role in the improvement of the gut microbiota balance and in individual health. For this reason, recently, a great interest on FOS and their influence on dietary modulation (including moderate sweetness, low carcinogenicity, low calorimetric value, and low glycemic index, etc.) of the human gut has been registered.

During the last years, developing products with prebiotic effects has been in the spotlight, and these new products include FOS obtained from sucrose or inulin. FOS have an important economic role in view of the high demand for obtaining and producing quickly on a large scale and at low costs. Conventionally, they are produced through a two-stage process that require an enzyme production and purification step in order to proceed with the chemical reaction itself. Several studies have been conducted on the production of FOS, aimed at optimizing the development of more efficient production processes and their potential as food ingredients. The improvement of FOS yield and productivity can be achieved by the use of different fermentative methods and different microbial sources of FOS-producing enzymes and the optimization of nutritional and culture parameters. Therefore, this review focuses on the latest progress in FOS research such as its production, functional properties, and market data. This is why biotechnology is developing new processes capable of increasing industrial production of enzymes associated in the production of FOS, as there are few investigations related to the production of these enzymes in large scale bioreactors. Therefore, studies have been focused on establishing biochemistry parameters and biochemical engineering in order to improve the production process as well as its economic viability.
Declaración de interés

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