

Ice Cream as a Vehicle for Incorporating Health-Promoting Ingredients: Conceptualization and Overview of Quality and Storage Stability

Christos Soukoulis, Ian D. Fisk, and Torsten Bohn

Abstract: Ice cream is a product with peculiar textural and organoleptic features and is highly appreciated by a very broad spectrum of consumers. Ice cream's structure and colloidal design, together with its low-temperature storage, renders it a very promising carrier for the stabilization and *in vivo* delivery of bioactive compounds and beneficial microorganisms. To date, many applications related to the design and development of functional ice cream have been documented, including products containing probiotics, prebiotics, synbiotics, dietary fibers, natural antioxidants such as polyphenols, essential and polyunsaturated fatty acids, and low glycemic index blends and blends fortified with mineral or trace elements. In this review, promising strategies for the incorporation of innovative functional additives to ice cream through the use of techniques such as microencapsulation, nanoemulsions, and oleogels are discussed, and current insights into the implications of matrix, processing, and digestion on bioactive compounds in frozen dairy desserts are comprehensively reviewed, thereby providing a holistic overview of the current and emerging trends in this functional food sector.

Keywords: antioxidants, fiber, frozen dairy desserts, nutritional value, prebiotics, probiotics

Introduction

Functional foods: definitions, concepts, and trends

Over recent years, there has been significant interest in the development of innovative food products conferring customized benefits to the consumer, that is, improving physical and mental well-being, prevention of diet-associated health complications in addition to fulfilling basic dietary function (hunger satisfaction and fulfillment of the daily nutritional need of consumers). The increased awareness of the consumer regarding health and nutrition related issues as well as the role of several food regulatory bodies to promote the production and consumption of minimally processed, healthier and more nutritious food products, appear to be steering a transformation within the food industry. Moreover, cultural, educational, and economic effects together with food quality and safety criteria have also been highlighted as drivers of consumer demand for healthy and safe food products (Fogliano and Vitaglione 2005).

Regardless of the expanding trends of the health-promoting food market, the term “functional food” still remains arbitrary, and thus several definitions have been introduced by both the research community and governmental regulatory authorities. For example, according to the American Council on Science and Health,

“Functional foods can be considered to be those whole, fortified, enriched, or enhanced foods that provide health benefits beyond the provision of essential nutrients (for example, vitamins and minerals), when they are consumed at efficacious levels as part of a varied diet on a regular basis” (Hasler 2002). Similarly, Arihara and others (2004) and Robefroid (2000) claimed that functional foods are those that contain 1 or more compounds that can support important or limited functions in the organism, promote welfare, health, and protection from degenerative chronic disease, without, however, curing diseases. From a food product standpoint, functional foods can be generally classified into the following categories (Bigliardi and Galati 2013):

- food fortified by nutrients such as vitamins, minerals and trace elements, antioxidants, and so on,
- food enriched with health-promoting ingredients that commonly do not exist in the enriched food matrix such as probiotics or prebiotics,
- reformulated food, that is, food where 1 or several ingredients associated with the adverse health impact is reduced, substituted, or removed, for example, trans-fatty acids, saturated fatty acids, antinutrients, using low glycemic index (GI) ingredients and fat replacers,
- food that is naturally enriched by 1 or more components, for example, butter containing omega-3 fatty acids or conjugated linoleic acid (CLA).

The rapidly expanding functional food market has required a significant expansion in research and innovation within the food industry, a sector traditionally considered as having a rather low innovation level compared to pharmaceuticals. However,

MS 20140231 Submitted 10/2/2014, Accepted 7/4/2014. Authors Soukoulis and Bohn are with Environment and Agro-biotechnologies Dept., Public Research Centre – Gabriel Lippmann, 41, rue du Brill, L-4422, Belvaux, Luxembourg. Author Fisk is with Div. of Food Sciences, School of Biosciences, Univ. of Nottingham, Sutton Bonington Campus, Leicestershire, Loughborough, LE12 5RD, UK. Direct inquiries to author Soukoulis (E-mail: soukouli@lippmann.lu; chrsoukoulis@yahoo.gr).

functional new product development (NPD) required changes to many traditional food production stages such as product strategy development, product design and development, product commercialization, product launch, and postlaunch, though it is generally accepted that many practices employed in every particular stage of NPD critically differ (Khan and others 2013). For instance, the orientation toward innovation, state-of-the-art and knowledge generation, the commercialization strategy, and establishment of collaborative networks and arrangements are only some of the differing points in the case of functional NPD (Khan and others 2013). Thus, it can reasonably deduced that the functional food industry is generally recognized by a higher level of innovation than the traditional one where innovation refers to the incremental improvement of the product rather than to a radical innovation approach.

To date, there are numerous examples of functional foods within the dairy sector, whilst noticeable innovation is also being demonstrated in the confectionary, soft-drink, bakery, and the infant food sector (Bigliardi and Galati 2013). With respect to the type of product functionalization, probiotics are the dominating functional/health-promoting ingredients for food applications followed by vitamins, minerals and trace elements, and prebiotics. Although most government regulatory bodies aim mainly to ensure the success of major dietary enhancing/balancing strategies (for example, fat, sugar, or salt reduction, increase of dietary fiber (DF) intake, and so on), the acceptance of functional food by the consumer is mainly governed by the awareness and confidence of the incorporated active substances and the overall organoleptically perceived quality and appearance. This implies that functional foods must diverge only minimally from regular food formulations in terms of sensory quality. Moreover, adding active compounds of low consumer popularity, for example, flavonoids, carotenoids, stanols, and so on, may result in poor market penetration (Granato and others 2010b). On the other hand, functional food promotion via health benefit claims may drastically increase interest, enhancing the buying intent of consumers; however, this is often considered not feasible in the EU, due to the existing legislation restrictions as, for example, for probiotics.

Incorporating bioactive compounds can be carried out by their direct addition to the food in free form, by using natural food matrices rich in the active compound (for example, spices, fruit, or vegetable pulp) or by adding specifically developed carriers from food grade materials (for example, microcapsules, edible films and coatings) where the active compound is physically or chemically entrapped. Finally, whatever the strategy, the end product should also meet a series of additional criteria such as enhanced stability and prolonged shelf life, cost efficiency, absence of safety concerns, and proven (ideally by means of clinical trials) bioavailability and functionality (Granato and others 2010b).

Ice cream as a vehicle for incorporating health-promoting ingredients

Ice cream is a complex colloidal food system that in its frozen state consists of ice crystals, air cells, and partially coalesced fat droplets dispersed in a continuous freeze-concentrated aqueous (serum) phase containing polysaccharides such as galactomannans, carrageenans, cellulose, sugars (sucrose and lactose), proteins, and minerals (especially calcium, but also sodium and potassium) (Goff 2008). Ice cream structure development and stabilization is a dynamic process where the main components, namely, biopolymers (proteins and polysaccharides), fat droplets, and water undergo significant colloidal and physical changes such as biopolymer hydra-

tion, fat droplet crystallization, ice nucleation, and crystallization, fat droplet partial coalescence, freeze-concentration, formation of cryogels, protein-polysaccharide phase separation, formation of biopolymer entanglement (Goff 1997; Bolliger and others 2000; Chang and Hartel 2002; Patmore and others 2003; Regand and Goff 2003; Soukoulis and others 2009; Cook and Hartel 2011). In addition, it is well established that phenomena such as ice recrystallization, air cell Ostwald ripening, and lactose crystallization govern the organoleptic quality, including creaminess, grittiness, coarseness, mouth-coating, and tongue lubrication as well as shrinkage (Muse and Hartel 2004; Soukoulis and others 2008, 2010b; Varela and others 2014; Soukoulis and Fisk in press).

Over the last 2 decades, ice cream science and technology has undergone a remarkable progress exploring and understanding structure—texture—storage stability interactions on a mechanistic basis. This has enabled food technologists to fairly realize the incorporation of novel or functional ingredients in model or real ice cream systems in order, to not only provide customized technofunctionality such as viscosity enhancement, cryoprotection, emulsification, water-binding, but also to improve health-related and nutritional aspects. For example, health-promoting compounds such as probiotics (Cruz and others 2009), prebiotics (AkalIn and others 2008), DF (Soukoulis and others 2009), natural antioxidants (Sun-Waterhouse and others 2013), fat sources rich in polyunsaturated fatty acids (PUFAs) (omega 3/6 fatty acids) (Song and others 2011), minerals (Erkaya and others 2012), and low GI sweeteners (LGSs) (Whelan and others 2008b) have been incorporated into ice cream systems. This trend for employing ice cream as a vehicle for health-related compounds appears to be supported by the consumers' demand for healthier and more nutritious food products that lack food additives and are able to confer health benefits beyond fulfilling basic dietary demands.

Due to its almost world-wide availability, high consumer acceptability and appealing sensory attributes, resulting in a high sales rate (5.8 billion L for the U.S. market in 2011, USDA 2012) ice cream can be regarded as a favorable vehicle for the delivery of bioactive compounds. Moreover, the implementation of generally low-temperature manufacturing steps (except for the homogenization-heat treatment), including frozen storage typically under dark conditions, renders ice cream as a good substrate for the long-term preservation of the functional traits of added health-promoting compounds, including beneficial living cells such as probiotics, reducing, for example, oxidative damage induced by light or heat.

Recently, major advances in ice cream technology, in terms of structure—texture development and stabilization, have been comprehensively reviewed (Soukoulis and Fisk in press). The aim of this review is to present recent advances in functional ice cream, a product enriched with bioactive compounds, probiotics, and their respective combinations. Strategies for the efficient incorporation of bioactive compounds and their potential impact on structure, organoleptic features, and storage stability are highlighted. In addition, recent studies on the stability/viability, digestion, and bioavailability of the particular bioactive compounds are also concisely discussed in the present work.

Probiotics

According to the FAO/WHO (2002), the term probiotic refers to live microorganisms that when orally administered in adequate amounts (10^6 to 10^7 CFU/g) confer health benefits to the host. Administration of probiotics to the human host has been reported to be associated with immune system modulation, reduction of

symptoms related with irritable bowel syndrome (IBS), diarrhea treatment, serum cholesterol reduction, anti-inflammatory action, and the prevention of cancer and mutagenesis (Saad and others 2013). Although health claims associated with the presence of viable bacteria in processed food are not yet having been ratified by the European Food Safety Association (EFSA), the body of evidence that certain bacteria strains from *Lactobacilli* and *Bifidobacteria* species could be health beneficial is rapidly growing.

In an attempt to satisfy the increasing market demand of probiotic-enriched food products, a vast number of applications related to dairy products such as yogurt (Cruz and others 2012; Cruz and others 2013; Pimentel and others 2013), ice cream (Cruz and others 2009), cheese and dairy spreads (Gomes and others 2011; Alves and others 2013; Esmerino and others 2013), breakfast cereals (Saarela and others 2006b), bakery (Soukoulis and others 2014) and confectionary products (Malmo and others 2013), emulsion-based preparations (Mantzouridou and others 2012), meat products (Khan and others 2011), as well as dairy and fruit beverages (Saarela and others 2006a; Granato and others 2010a, 2010b; Antunes and others 2013; Castro and others 2013; Maganha and others 2014) have been demonstrated. With respect to market share, yogurt and fermented milk beverages and fruit juices remain the largest global market share holders, while ice cream appears to be a promising product for probiotic food development (Euromonitor 2012).

The viability and therefore the functionality of probiotics is impacted by several intrinsic (species/strain, morphological characteristics of the bacteria, and their ability to produce intracellular cryoprotectants) and extrinsic (food substrate composition, exposure to heat, mechanical treatment, oxygen toxicity, and pH) parameters (Fu and Chen 2011). Depending on the species, probiotic bacteria are regarded as strictly anaerobic (*Bifidobacteria*) or microaerophilic (most *Lactobacilli*), and therefore susceptible to high partial oxygen pressure. To date, microencapsulation of probiotics in biopolymer matrices by mechanical or physicochemical means (dehydration, emulsification, extrusion, ionic or heat-induced gelation, cross-linking, and coacervation) are the most common strategies for the delivery of probiotic efficacy to food products able to withstand severe processing steps such as thermal (heat or freezing) and mechanical (mixing, size reduction, and pumping) treatments, thermomechanical processing, storage conditions, and so on (Champagne and Fustier 2007; Burgain and others 2011). Moreover, changing the composition of the fermentation medium (such as polyols, sugars, Tween 80, and inorganic salts), controlled pH and temperature, growth state of the culture (logarithmic or stationary), cryopreservation and subjecting bacterial cells to heat, cold, acid, or osmotic stress and starvation, have also been proposed as alternative strategies for the cryopreservation of lactic acid bacteria (LAB) and probiotics (Pehkonen and others 2008; Siaterlis and others 2009).

Methods for probiotic incorporation into ice cream

The first attempts to incorporate viable probiotic bacteria into frozen dairy desserts were made decades ago using ice cream, fermented frozen desserts as well as their ethnic analogs (Holcomb and others 1991; Laroia and Martin 1991; Hekmat and McMahon 1992; Westerbeek 1995; Kebary 1996; Inoue and others 1998). Probiotics can be incorporated into ice cream either in free or microencapsulated form. In the first case, probiotics can be supplied by either blending an acidified/fermented milk base (such as probiotic yogurt, acidified milk, or cream) with the ice cream mix base at the onset of production, or by direct inoculation of the ice

cream mix with a single or a symbiotic culture starter prior to the whipping-freezing step (Tamime and Robinson 2007; Soukoulis and Tzia 2008). Direct inoculation into the final ice cream mix might allow none, partial, or full fermentation depending on the flavor–texture quality characteristics required (Soukoulis and Tzia 2008). On the other hand, using microencapsulated probiotic bacteria, that is, in biopolymer cross-linked or spray/freeze-dried matrices facilitates the manufacturing process (no need for cultured milk base preparation). However, it can also reduce effectively the mechanical or osmotic stress that would induce injury to living cells (Hekmat and McMahon 1992; Kailasapathy and Sultana 2003; Homayouni and others 2008a). Employing microencapsulated probiotics for the production of functional ice cream has gained much attention during the last few several years due to the versatility of the method, the prolonged shelf life attained for both microcapsules and ice cream, and the minimized impact of the carrier material on the sensory, texture, and structural aspects of the finished product (Mohammadi and others 2011).

Factors affecting probiotic viability in ice cream

Several factors have been reported to significantly impact the viability of probiotic bacteria throughout processing, such as freezing processes and postfreezing (hardening and frozen storage) conditions, and additional parameters such as ice cream composition (sugars, polysaccharides, protein type, and content), pH, and the presence of oxygen (Cruz and others 2009; Mohammadi and others 2011). Although the sublethal impact of each factor might range from negligible to fairly considerable, their combination can result in severe lethality of the probiotic living cells. It is therefore deduced that the strict control of the implemented processing practices and the compositional and structural product are essential to ensure an end-to-end cryopreservation of the probiotic microbiota (Table 1).

Freezing process conditions. Depending on their freeze tolerance, the sublethal injuries of probiotic bacterial cells due to the implemented freezing process may range from minor to severe. Generally, the lethality of probiotics throughout freezing is primarily associated with the occurrence of osmotic and mechanical stresses. From a mechanistic point of view, extremely high freezing rates (cold shock) may lead to increased permeability of the membranes, resulting in leakage of intracellular material due to intracellular and extracellular ice formation, inducing mechanical damage of the cell membrane. The freeze concentration of the intracellular solutes can also modify pH, ionic strength, and concentration of several compounds, leading to toxic effects (Bremer and Ridley 2004). On the other hand, low freezing rates associated with slow supercooling and low ice nucleation have been also reported as being associated with plasmolysis, for example, intracellular water loss driven by high osmotic pressure gradients. In addition, intrinsic factors inherently associated with the freezing process, such as type of freezer (batch or continuous), freezing rate, dasher type, and whipping conditions, are well known to have a strong impact on probiotic viability (Sheu and others 1993; Ferraz and others 2012). It is generally accepted that adequately high freezing rates are required to minimize sublethal injuries of the bacterial cells. Although data relating to the effect of freezing processes are rather scarce, Sheu and others (1993) showed that continuous freezing enhances only slightly the survival of *Lactobacillus delbrueckii* spp. *bulgaricus* compared to batch freezing (55% and 50% survival after freezing, respectively). The former may be attributed mainly to higher ice nucleation in the case of continuous freezing, allowing the formation of a large number of

Table 1—Compositional, technological, and sensory characteristics of probiotic ice cream and stability of probiotic microbiota under frozen storage and *in-vitro* gastrointestinal conditions.

Composition and processing tested	Probiotic bacterial strain	Incorporation method	Overrun (%) and pH	Freezing conditions and survival throughout freezing process	Frozen storage conditions and losses during storage	Survival under <i>in-vitro</i> gastrointestinal conditions/ <i>in vitro</i> adhesion	Reference
Percentage of probiotic cultured milk added to ice cream mix base (25% to 75%)	<i>L. acidophilus</i> (La-5)	Free cells	74% to 88%	Batch (−5 °C)	−20 °C for 16 wk	Nd	Christiansen and others (1996)
Ice cream with/ without 2% glycerol	<i>B. bifidum</i> (Bb-12) <i>L. reuteri</i> <i>L. acidophilus</i> (La-5) <i>L. rhamnosus</i> GG <i>B. bifidum</i> (Bb-12)	Free cells	5.4 to 6.6 Nd 6.1 to 6.3	0.6 to 1.0 logCFU/g Batch (nd) 0.7 to 0.8 logCFU/g	0.1 to 0.7 logCFU/g −20 °C for 52 wk nonsignificant loss	Nd	Hagen and Narvhus (1999)
Milk fat (3.8% and 10%) Resistant starch	<i>L. acidophilus</i> Lafti™ L10 <i>B. lactis</i> BLC-1 <i>L. paracasei</i> LCS-1	Free cells	48% to 58% 6.3 to 6.4	Continuous (−5 °C) 0.3 to 0.7 logCFU for <i>L. acidophilus</i> 0.2 to 0.5 logCFU/g for <i>B. lactis</i> 0.1 to 0.4 logCFU/g for <i>L. paracasei</i> Batch (nd) 0.2 to 0.3 logCFU/g	−25 °C for 52 wk 0.8 to 1.25 logCFU/g for <i>L. acidophilus</i> 0.4 to 0.65 log CFU/g for <i>B. lactis</i> 0.75 to 1.15 log CFU/g for <i>L. paracasei</i> −16 °C for 30 d nonsignificant loss −28 °C for 240 d nonsignificant loss	Nd	Haynes and Playne (2002)
Fat (5% and 10%) and sugar (15% and 22%)	<i>L. jonsonii</i> La 1	Free cells	23% to 27% 6.5 to 6.6	Batch (nd) 0.2 to 0.3 logCFU/g	Incubation in gastric juice for 3 h 3 logCFU/g Growth in MRS containing 0% to 2% w/v bile salts <0.3 logCFU/g for 0% to 0.4% <0.6 logCFU/g for 0.8% to 2.0% Nd	Alamprese and others (2002)	
Plain ice cream	<i>L. acidophilus</i> 2401 <i>B. infantis</i> 1912 and their symbiotic blend	Free cells Calcium alginate—resistant starch microbeads (fresh or freeze-dried)	40% to 50% Nd	Batch (nd) Nd	−20 °C for 24 wk nonsignificant loss	Godward and Kalliasapathy (2003)	
0%, 1%, or 2% inulin 15%, 18%, or 21% sugar	<i>B. lactis</i> BL-01 <i>L. acidophilus</i> La-14 <i>L. delbrueckii</i> ssp <i>bulgaricus</i> <i>S. thermophilus</i> <i>L. rhamnosus</i> GG	Free cells	28% to 32% 5.1 to 5.3	Batch (nd) 0.5 to 1.0 logCFU/g	−18 °C for 90 d 1.5 to 3.0 logCFU/g	Nd	Akin (2005)
Fat (5% and 10%) and sugar (15% and 22%)		Free cells	24% to 29% 6.4 to 6.6	Batch (nd) 0.2 to 0.3 logCFU/g	−16 °C for 30 d nonsignificant loss −28 °C for 240 d nonsignificant loss	Incubation in gastric juice for 3 h 4 logCFU/g Growth in MRS containing 0% to 2% w/v bile salts <0.8 log CFU/g only at 2% w/v	Alamprese and others (2005)

Continued

(Continued)

Table 1—Continued.

Composition and processing tested	Probiotic bacterial strain	Incorporation method	Overrun (%) and pH	Freezing conditions and survival throughout freezing process	Frozen storage conditions and losses during storage	Survival under <i>in-vitro</i> gastrointestinal conditions/ <i>in vitro</i> adhesion	Reference
Acerola ice cream with different pH (4.5 and 5)	<i>B. longum</i> <i>B. lactis</i>	Free cells	Nd 4.5 and 5.0	Continuous (−6 °C) 0.32 and 0.13 logCFU/g for <i>B. longum</i> 0.77 and 0.32 logCFU/g for <i>B. lactis</i>	−18 °C for 15 wk no loss and 0.12 log CFU/g for <i>B. longum</i> no losses observed for <i>B. lactis</i>	Nd	Favaro-Trindade and others (2006)
FOS free, 4% oligofructose, 4% inulin	Mixed culture: <i>L. acidophilus</i> (La-5) and <i>B. animalis</i> ssp. Bb-12	Free cells	27.6% to 50.6% approximately 5.5	Batch (nd) Nd	−18 °C for 90 d <0.6 logCFU/g for <i>L. acidophilus</i> <0.4 logCFU/g for <i>B. animalis</i>	Nd	Akalin and Erişir (2008)
Resistant starch addition (1%)	<i>L. casei</i> (LC-01) <i>B. lactis</i> (Bb-12)	Free cells Cells encapsulated in resistant starch-calcium alginate microbeads	95% 6.6	Batch (nd) Nd	−20 °C for 180 d 2.3 and 1.3 logCFU/g for free and encapsulated <i>L. casei</i> 3 and 0.7 logCFU/g for free and encapsulated <i>B. lactis</i>	Nd	Homayouni and others (2008)
5% and 10% milk fat	<i>L. acidophilus</i> (DMSZ 20079) <i>B. bifidum</i> (DMSZ 200456) and their symbiotic blend	Free cells	35% to 44% 5.8 to 6.3	Batch (nd) 0.2 log CFU/g for <i>L. acidophilus</i> 0.26 for CFU/g <i>B. bifidum</i>	−20 °C for 90 d 0.38 log CFU/g for <i>L. acidophilus</i> 0.23 log CFU/g for <i>B. bifidum</i> Similar results for double strain systems	Nd	Turgut and Cakmakci (2009)
Fruit (2.5% fat, 30.2% carbohydrates, 12.4% MSNF) or vanilla flavor (8.8% fat, 22.2% carbohydrates, and 10.8% MSNF)	<i>L. rhamnosus</i> <i>L. casei</i>	Free cells	41% to 43% Nd	Batch (nd) Nd	−20 °C for 16 wk <0.6 logCFU/g for fruit flavor <0.2 logCFU/g for vanilla flavor <0.5 logCFU/g for synbiotic formulations	Growth in MRS containing 1% bile salt 2.1 logCFU/g for fruit flavor 1.5 logCFU/g for vanilla flavor 2.0 to 2.1 log CFU/g for synbiotic formulations	Di Ciccio and others (2010)
Nonactivated culture (lyophilized culture suspended in sterile milk)	<i>L. acidophilus</i> LMGP 21381	Free cells	53% to 55% 6.7	Batch (−6 °C) 0.86 logCFU/g for nonactivated 0.03 logCFU/g for activated	−15 °C for 45 wk 0.12 logCFU/g for nonactivated no loss for activated −25 °C for 45 wk 0.13 logCFU/g for nonactivated no loss for activated	Nd	Nousia and others (2011)

(Continued)

Table 1—Continued.

Composition and processing tested	Probiotic bacterial strain	Incorporation method	Overrun (%) and pH	Freezing conditions and survival throughout freezing process	Frozen storage conditions and losses during storage	Survival under <i>in-vitro</i> gastrointestinal conditions/ <i>in vitro</i> adhesion	Reference
Not-altered	<i>L. acidophilus</i> <i>L. rhamnosus</i>	Free cells	90% 6.1 to 6.2	Batch (−4 °C) 0.28 logCFU/g for <i>L. acidophilus</i> 0.33 logCFU/g for <i>L. rhamnosus</i>	−19 °C for 12 wk 0.82 logCFU/g for <i>L. acidophilus</i> nonsignificant losses for <i>L. rhamnosus</i>	Incubation in gastric juice for 3 h at pH = 1.5, 2.5, and 3.5 pH = 1.5: No viable cells for both strains pH = 2.5: No viable cells for <i>L. acidophilus</i> , <7 logCFU/g for <i>L. rhamnosus</i> GG pH = 3.5 No significant losses for both strains Viability of both strains reduced at 0.2% of bile salts	Abghari and others (2011)
Ice cream containing phenolic compounds (ellagic acid, gallic acid, grape seed extract, pomegranate peel extract, and peppermint essential oil)	<i>L. casei</i> Shirota	Free cells	Nd 5.88 (PPE), 6.21 to 6.29 for rest phenolics	Batch (−5 °C) Nd	−18 °C for 60 d 0.86, 0.37, 0.27, and 0.34 logCFU/g for free phenolics or for ice cream with EA, PEO, and GSE insignificant losses in ice creams with PPE or GA	Nd	Song and others (2011) Sagdic and others (2012)
Plain ice cream	<i>L. acidophilus</i> (La-5) <i>B. animalis</i> (Bb-12) <i>P. jensenii</i> 702	Free cells	Nd	Batch (nd) Nd	Nd	Incubation in gastric juice for 3 h pH = 2, 3, 5, 2.8, and 5.3 log CFU/g for <i>L. acidophilus</i> , <i>B. animalis</i> , and <i>P. jensenii</i> , respectively No significant losses at pH = 3 and 4 Incubation in bile salts (0.3% w/v) for 4 h 1.6, 0.84, and 2.2 logCFU/g respectively 0.54, 0.73, and 0.61% adhesion in Caco-2 cells	Ranadheera and others (2012)

(Continued)

Table 1—Continued.

Composition and processing tested	Probiotic bacterial strain	Incorporation method	Overrun (%) and pH	Freezing conditions and survival throughout freezing process	Frozen storage conditions and losses during storage	Survival under <i>in-vitro</i> gastrointestinal conditions/ <i>in vitro</i> adhesion	Reference
0% and 3% inulin 0% and 0.46% whey protein concentrate	<i>L. acidophilus</i> NCDC 14	Free cells	Nd 5.5	Batch (nd) 0.61 to 0.77 logCFU/g	−18 to −23 °C for 15 d 0.5 to 0.79 logCFU/g	Nd	Pandiyan and others (2012a)
0%, 3%, 4%, and 5% FOS 0% and 0.46% whey protein concentrate	<i>L. acidophilus</i> NCDC 14 <i>S. boulardii</i>	Free cells	Nd 5.5	Batch (nd) 0.49 to 0.68 logCFU/g	−18 to −23 °C for 15 d 0.56 to 0.7 logCFU/g	<i>In vivo</i> testing of probiotic bacteria survival in human faecal samples 5.96 to 6.82 log CFU/g after 7 d 6.61 to 7.10 logCFU/g after 15 d	Pandiyan and others (2012b)
45%, 60%, and 90% overrun	<i>L. acidophilus</i> DOWARU®	Free cells	45% to 90%	Continuous (nd) 0.02 to 0.05 logCFU/g	−18 °C for 60 d 0.1, 0.2, and 1.98 log CFU/g for 45%, 60%, and 90%, respectively	Nd	Ferraz and others (2012)
1.5% to 5% fat, 0% to 8% oligofructose	<i>L. acidophilus</i> (La-5)	Calcium alginate microbeads	31% to 43%, 5.1 to 5.5	Batch (nd) 0.5 to 1.0 log CFU/g	−18 °C for 60 d 1.5 to 2 log CFU/g	Nd	Ahmadi and others (2012)
Packaging material (glass, polyethylene, and polystyrene)	<i>L. acidophilus</i> (La-5) <i>B. animalis</i> (Bb-12) <i>P. jensenii</i> 702	Free cells	6.5 to 6.7 26% to 34%	Batch (nd) 0.23, 0.18, and 0.05, respectively	−20 °C for 52 wk <0.4 log CFU/g for <i>L. acidophilus</i> <0.2 log CFU/g for <i>B. animalis</i> <0.3 log CFU/g for <i>P. jensenii</i>	Nd	Ranadheera and others (2013)

MRS = de Man, Rogosa, Sharpe agar, FOS = fructo-oligosaccharides, EA = ellagic acid, GA = gallic acid, GSE = grape seed extract, PEO = peppermint essential oil, PPE = pomegranate peel extract, Nd = not determined (no data), MSNF = milk solids nonfat.

small-size ice crystals (both in intra- and extracellular domains), minimizing mechanical rupture of the cell membranes by interpenetrating ice crystals (Bremer and Ridley 2004). Moreover, ice cream hardening (the rapid cooling process at the end prior to storage) had a minor impact on bacterial cell lethality, ranging from 10% to 15% regardless of the freezing process (continuous or batch) implemented (Sheu and others 1993).

The developing shearing forces during the freezing process may also lead to sublethal injuries of probiotics due to mechanical rupture of the cell membranes, particularly in processes enabling very high shear stress, as with low temperature extrusion (LTE) freezing. Recently, the high-pressure low-temperature (HPLT) technique has been successfully applied for the production of ice cream with good air cell and ice crystal recovery, thus improving sensory properties (Volkert and others 2012). The method has also been previously applied in the production of anhydrobiotics, such as freeze-dried skim milk matrices containing *L. rhamnosus* GG (Volkert and others 2008). This may be an emerging method for the development of functional frozen dairy desserts with maximal viability of living cells.

Compositional features of ice cream mix. Ice cream components able to modify the colligative properties of the ice cream mixture (sugars, polyols, oligosaccharides, and proteins) can directly or indirectly affect the cryopreservation of probiotics. Nevertheless, these ingredients are deemed as cryoprotectants—many of which are used extensively for cell lyophilization—and it should be noted that their combined effect on ice crystallization, freeze concentration, and ice recrystallization has an impact on cryoprotection (Santivarangkna and others 2008). For instance, though polyols are regarded as efficient compounds for minimizing cellular injury during ice crystallization, their performance could be restricted during frozen storage due to their poor ability to control ice recrystallization. Similarly, ingredients that elevated the glass transition temperature (polysaccharides/high-molecular-weight sugars ensuring maximal stability of the probiotics in the rubbery/glassy matrix during frozen storage) might also induce high osmotic stress due to freeze concentration of the serum phase (Pehkonen and others 2008).

The majority of studies investigating survival of probiotic bacteria (in real or model ice cream systems) have revealed that *Bifidobacteria* are more freeze-resistant than *Lactobacilli* or *Saccharomyces* (Hekmat and McMahon 1992; Başığit and others 2006; Akin and others 2007; Akalin and Erişir 2008; Homayouni and others 2008b). Homayouni and others (2008b) reported that storing probiotic ice cream at -20°C for 6 mo was associated with higher inactivation rates of *L. casei* compared to *Bifidobacterium lactis*. Magariños and others (2007) demonstrated that freezing (scraped-surface freezer, -10°C draw temperature) resulted in lower viability of *L. casei* compared to *B. animalis*, whereas no significant changes on the inactivation rates of both species upon frozen storage were observed.

Increasing sugar or fat content in order to enhance texture or structural features has been reported to moderately impact probiotic cell viability. Alamprese and others (2002) did not observe any significant effect of sugar/fat content on the viability of *L. johnsonii* La1 in ice cream stored for 180 d at -28°C . Similarly, the presence of sucrose or aspartame in acidified ice cream did not affect viable counts of a symbiotic/probiotic blend composed of *L. acidophilus* and *L. rhamnosus* during 180 d at -20°C (Başığit and others 2006). On the other hand, Akin and others

(2007) found that 18% w/w of sucrose was required to ensure maximal survival of mixed strains of LAB and bifidobacteria, regardless of the species/strain. In model ice cream systems, Homayouni and others (2008b) demonstrated that probiotic osmotolerance in the presence of sucrose up to 25% w/w is strain- and type-dependent, with *Lactobacilli* (*L. casei*, *L. acidophilus*-La5) exhibiting a higher sugar tolerance than *Bifidobacteria* (*B. bifidum* and *B. longum*). However, including a very strong osmolytic agent (2% w/w glycerol) has been reported to have a negligible effect on the cryopreservation of mixed probiotic bacteria (*L. acidophilus* and *B. bifidum*) during freezing and storage (-20°C , 52 wk), while remarkably higher amounts of the cryoprotectant were required to achieve a significant cryoprotective effect (Hagen and Narvhus 1999).

Prebiotic fiber such as fructooligosaccharides (FOS), galactooligosaccharides (GOS), and resistant starch have also been tested as potential cryoprotective agents and as probiotic growth and activity stimulants in the gut environment for ice cream systems (Akin 2005; Akin and others 2007; Akalin and Erişir 2008). Inulin and oligofructose are the most common prebiotics implemented in probiotic ice cream production due to their established physiological aspects (modulation of gut flora bifidogenicity, suitability for diabetics, potential cancer prevention, improved mineral absorption, and improved lipid metabolism) and specific technological properties allowing their use as fat mimetics, sugar replacers, texturizers, mouthfeel improvers, and stabilizers (Coussement and Franck 2001; Franck 2002). The literature on FOS functionality in probiotic ice cream is rather contradictory; Akalin and Erişir (2008) reported a significant improvement of the viability of *B. animalis* Bb-12 and *L. acidophilus* La-5 during 90 d of storage at -18°C in the presence of 4% oligofructose, while no remarkable changes in viable counts of probiotic microflora were observed when inulin (4%) was added.

Akin and others (2007) revealed a strain dependency of the impact of inulin on the viability of the ice cream microflora. While the viability of yogurt culture starters (*L. delbrueckii* spp. *bulgaricus* and *S. thermophilus*) was not significantly improved by inulin (up to 2% w/w), a noticeable improvement was detected for *L. acidophilus* and *B. lactis*. On the other hand, in studies by Di Criscio and others (2010) and Ahmadi and others (2012), supplementation of probiotic ice cream with prebiotic fiber (inulin) did not exhibit any stimulating activity for *L. acidophilus*-La5, *L. rhamnosus*, and *L. casei*.

pH and ionic strength. Modulating the pH of the ice cream mix is another important factor that influences the survival of probiotics, particularly for hardened products. The decrease of pH is due to the metabolic activity of the microflora (producing lactic and/or acetic acid via the Emden–Meyerhof–Parnas pathway) or due to inclusion of acidic preparations (fruit juices, purees, or syrups) (Marshall and others 2003). Generally, based on manufacturing practices, the pH of probiotic ice cream might vary from 4.5 to 6.3 for fully fermented and unfermented formulations, respectively (Soukoulis and Tzia 2008). It is well established that *Lactobacilli* are more acid-tolerant than *Bifidobacteria*; consequently, a pH of 5.0 or higher, or 5.5 and higher, is usually recommended for probiotic ice creams, respectively (Laroia and Martin 1991). On the other hand, it has been shown that using symbiotic cultures (mixtures of LAB and probiotic bacteria) instead of single probiotic starter cultures can surmount the adverse results of acidic environmental conditions on probiotics (Davidson and others 2000).

Favaro-Trindade and others (2007) deduced that the freeze-induced injuries of *L. acidophilus* (74-2 and LAC4) were minimized in low-pH ice cream formulations containing added yellow mombin pulp (4.5 compared with 5.0), whereas the pH did not modify bacterial cell inactivation rates upon storage at -18°C for 105 d. In a previous study, the same research group had demonstrated an improved viability of *B. longum* and *B. lactis* in low-pH (4.5) ice creams containing acerola pulp. However, it should be noted that in both studies, the initial load of viable bacteria in the fully fermented systems (pH 4.5) was higher due to a prolonged incubation time, a factor well known to influence cell injuries during freezing (Santivangkna and others 2008). The incorporation of symbiotic cultures (*L. acidophilus*, *B. longum*, and *S. thermophilus*) in ice cream has been reported as a good strategy for reducing the lethal effects of frozen storage on probiotics as measured by the β -galactosidase enzyme activity (Kailasapathy and Sultana 2003). Davidson and others (2000) also demonstrated the synergistic action between LAB and probiotic bacteria by means of freeze tolerance although the authors did not observe a significant impact of the pH on bacterial cell recovery over 11-wk storage at -20°C .

Presence of oxygen. The presence of oxygen primarily due to air incorporation during the freezing-whipping step and secondarily due to oxygen dissolution in the ice cream mix during processing, including blending, homogenization, pumping, and oxygen permeation through the packaging material is another well-established route inducing bacterial death (Talwalkar and Kailasapathy 2004). Although the amount of air incorporated (also referred as overrun) varies according to the adopted manufacturing practices and the required structurization/texturization, it usually ranges from 20% for artisanal ice cream (gelato) to 50% to 100% for industrial frozen dairy desserts (Marshall and others 2003; Goff 2008). The lack of oxygen-scavenging cellular functionality in probiotic bacteria explains the toxic effect of oxygenic metabolites, such as superoxide anion (O_2^-), hydroxyl radicals (OH^-), and hydrogen peroxide (H_2O_2) in the cells (Mills and others 2011). In a recent study by Ferraz and others (2012), it was demonstrated that an overrun of 45% or less does not modify significantly the load of viable *L. acidophilus* DOWARU[®] cells for a storage period of 60 d at -18°C . A further increase of the overrun (60% to 90%) induced a significant reduction of microbial cells ranging from 1.02 to 1.98 log CFU/g after 60 d of storage at -18°C . Generally, the strictly anaerobic *Bifidobacteria* are pronouncedly more vulnerable to oxygen than *Lactobacilli*, which was also confirmed for aerated dairy desserts including ice cream (Homayouni and others 2008b), although lethality of the latter depended on the employed strain. It was shown that *L. rhamnosus* and *L. casei* exhibited a higher oxygen tolerance than *L. acidophilus* in unfermented ice creams (Homayouni and others 2008b; Abghari and others 2011). However, the presence of oxygen scavenging compounds (for example, L-cysteine, L-ascorbic acid, whey proteins, acid casein hydrolysate, catechins, and polyphenols) is regarded as effective for ensuring the survival of probiotic bacteria in dairy matrices including ice cream (Ravula and Shah 1998; Homayouni and others 2008b; Sagdic and others 2011; Gaudreau and others 2013; Pandiyan and others 2012c; Soukoulis and others 2014). Microencapsulation of probiotics also showed to hinder the formation of oxygenic metabolites in dependence of the encapsulant and the probiotic species/strain. For example, Shah and Ravula (2000), Homayouni and others (2008b), and Ahmadi and others (2012) found that the survival rates of microencapsulated probiotic cells in matrices composed of sodium alginate or bi-

nary blends of sodium alginate with either resistant starch or FOS significantly improved compared to unprotected bacterial cells. In addition, Godward and Kailasapathy (2003) demonstrated that symbiotic cultures instead of single or mixed probiotic strains can provide better cryoprotection and metabolic activity preservation compared to both free and encapsulated systems.

Impact of probiotics on quality characteristics of ice cream

Quantity and quality of ice cream mix components, together with the structure development process during the freezing-whipping step, are the most crucial factors that control the quality characteristics of ice cream (Marshall and others 2003). In this context, adding probiotics without any preacidification prior to freezing would not be expected to impact flavor-taste characteristics of the final product, as probiotic cells would not exert any remarkable metabolic activity leading to the formation of volatile and nonvolatile flavor compounds. On the other hand, incorporating probiotics into partially/fully pre fermented ice cream base, or immobilized within biopolymers, can improve the textural and sensory properties of ice cream. Specifically, increase of mix viscosity, enhancement of melting resistance, development of peculiar organoleptic properties such as refreshing and pleasantly sour flavor, improved body and controlled iciness, and ice crystal-induced grittiness have been reported (Christiansen and others 1996; Christiansen and others 1999; Aryana and Summers 2006; Salem and others 2006).

Unfermented probiotic ice creams. It is generally accepted that the addition of probiotics to plain ice cream mixes does not change composition, the viscosity, and physicochemical characteristics of the ice cream mixes, or the overrun values and instrumental texture profile of the finished frozen products (Alamprese and others 2002; Godward and Kailasapathy 2003; Alamprese and others 2005; Homayouni and others 2008a; Di Criscio and others 2010; Pandiyan and others 2012b). Minor or nonperceivable sensory changes (appearance, texture, body, flavor, and aroma) have been reported in a series of studies conducted using unfermented ice creams (Alamprese and others 2002; Di Criscio and others 2010; Ferraz and others 2012; Homayouni and others 2012; Pandiyan and others 2012b). Apparently, the low bacterial volume is insufficient to cause any significant change in the structure of the ice cream.

However, according to the studies of Di Criscio and others (2010) and Pandiyan and others (2012b), probiotic ice cream was rated as less appealing in terms of taste and aroma. In addition, Di Criscio and others (2010) demonstrated that the sensory profile of probiotic ice cream was strain-dependent, with the *L. casei* formulation was better perceived in terms of taste compared to *L. rhamnosus* GG. Triangle test sensory analysis of ice creams with or without added *L. rhamnosus* GG showed that probiotic ice cream was significantly different than plain ice cream although no perceivable "probiotic" off-flavors were reported (Alamprese and others 2005).

Fermented probiotic ice creams. Contrarily to nonacidified probiotic ice cream, the fermentation of the ice cream mix either by yogurt or symbiotic culture starters is associated with important colloidal, physicochemical, textural, flavor, and taste changes (Tamime and Robinson 2007). Thus, physicochemical phenomena such as gelation, casein micelle destabilization, formation of protein aggregates, protein-polysaccharide crosslinking, milk-protein interactions at the water-oil interface, and formation of fermentation metabolites such as exopolysaccharides (EPS),

lactic acid, and volatile organic compounds, have been reported to occur throughout fermented ice cream production (Christiansen and others 1996; Christiansen and others 1999; Salem and others 2006).

Parameters such as inoculum size, fermentation conditions, and pH endpoint, and the use of ropy (EPS-producing) strains have been reported as having a significant effect on the viscosity development of ice cream mix. Aryana and Summers (2006), investigating the impact of inoculum (0%, 0.002%, 0.02%, and 0.2% v/v) on several physicochemical properties of probiotic ice cream, observed a significant decline of viscosity when 0.2% or 0.02% v/v of a symbiotic culture (*L. acidophilus*, *Bifidobacterium*, and *L. casei*) was used. Experimental data dealing with the effect of acidification (evaluated by endpoint pH and titratable acidity) on the macroviscosity of probiotic ice cream mixes are contradictory. Typically, similar or lower viscosity, compared to the noninoculated samples, was found in fermented ice cream mixes using non-EPS-producing probiotic bacterial strains (Aryana and Summers 2006; Guner and others 2007). Contrarily, Salem and others (2006) demonstrated that blending ice cream mixes with fortified single-strain probiotic milk can result in a significant increase of the viscosity in the aged ice cream mixes. The authors found that the impact on viscosity was strain-dependent, with *L. gasseri*, *L. rhamnosus*, and *L. reuteri* imparting the largest thickening effect compared to *L. acidophilus* and *B. bifidum*.

On the other hand, the *in situ* use of ropy probiotic cultures has also been pinpointed as an effective strategy to enhance macroviscosity of probiotic ice cream mixes (Christiansen and others 1999; Goh and others 2008). Christiansen and others (1999) showed that the use of EPS-containing milk synthesized by *Lactococcus lactis* ssp. *cremoris* can successfully replace stabilizers when blended with plain (nonstabilized) ice cream mix at a ratio of 1:4. Ropy milk presence induced a significant viscosity increase of the final mix, mimicking the action of polysaccharides. Similarly, Goh and others (2008) reported a thickening effect on ice cream mixes by *L. delbrueckii*, comparable to that of stabilized (0.15% locust bean gum, 0.09% guar gum, and 0.01% w/w κ -carrageenan) ice cream.

In addition, the use of EPS-producing probiotic culture starters has been reported to impact the whipping ability of the ice cream mixes, overrun, and melting behavior of the finished frozen products (Christiansen and others 1999). Blending of ice cream mix (0.5% commercial stabilizer), with ropy milk at the ratio of 1:4 resulted in ice creams with acceptable body and texture, slightly perceivable iciness, high shape retention and melting resistance (17.5% compared with 10.3%), though the molten ropy milk-based systems had a slightly curdled appearance (Christiansen and others 1999). In a previous study, Christiansen and others (1996) reported that ice cream mixes containing up to 50% of probiotic ropy milk were associated with reduced air incorporation ability that was attributed to protein aggregation due to their lower pH value. Contrarily, Goh and others (2008) reported that fermented ice creams produced using different types of EPS-producing LAB had significantly higher overrun values than the control ones. Investigating the interrelationship between the viscoelastic profile of the ice cream mixes and the air–water interface interactions, the authors suggested that the molecular properties of the EPS formed by the ropy strains can influence significantly the air cells development and stability during the freezing process. The presence of EPS in the unfrozen serum phase was associated with foam colloids of high elastic modulus (G') enabling the stabilization of the formed fine air cells (Goh and others 2008). However, it should be noted that in the case of acidified ice creams produced using

a nonropy culture starters, air incorporation appears to be hampered proportionally to the extent of protein aggregation due to pH lowering (Goh and others 2008; Soukoulis and Tzia 2008).

Stability of probiotic bacteria during ice cream digestion as investigated by *in vitro* methods

An important aspect of the probiotic concept is the delivery of a large number of viable bacteria to the colon, thereby demonstrating cell survival upon during oral, gastric juice and intestinal phase transit. Despite the large number of studies investigating probiotic survival throughout ice cream manufacture, literature data on the ability of ice cream to deliver probiotic efficacy under simulated gastrointestinal tract (GIT) conditions (an accepted model to estimate survival rates of bacteria *in vivo*) (Cook and others 2012) are rather scarce. Maintaining probiotic viability under GIT conditions can be as challenging as cryopreservation, depending on the severity of the changes taking place following ingestion and particularly after the dissolution/disintegration of the matrix, exposing the released bacterial cells to the comparably harsh GIT conditions.

Oral processing leads to meltdown of the ice cream and its dilution with saliva, partial α -amylase-facilitated breakdown of complex carbohydrates and fat droplet coalescence takes place (Stokes and others 2013). This implies (except for probiotic ice cream systems where the bacterial cells are microencapsulated) that living cells are immediately exposed to GIT conditions.

It has been demonstrated that both gastric juice and bile salts during small intestinal digestion might induce a significant reduction of viable bacteria. However, it has been widely attested that the passage through the stomach is the most critical factor for delivering probiotic efficacy to the human host (Cook and others 2012). The varying harshness of gastric conditions including transit time (several minutes for liquid meals to ca. 2 to 3 h for solid meals) and pH (about 1.6 in fasting and pH 5 during the fed state) modulate the survival of probiotics, whereas their embedding in acid-resistant biopolymer matrices such as microcapsules and uncoated or lipid bilayered dry microparticles can provide protection against matrix disintegration during gastric passage (Cook and others 2012).

After passage through the stomach, probiotic cells enter the small intestine where pH and transit time are also variable, although pH conditions are considered less harsh than those of the gastric phase, ranging between 6.8 (duodenum) and 7.5 (ileum) (Cook and others 2012). In the small intestine, the most toxic parameter is bile salts originating from the gall bladder. Due to their interfacial properties, bile salts can alter the conformation of cell membrane proteins (inducing protein misfolding or denaturation) and interact with membrane lipids, modifying the structural integrity and permeability of the cell membranes (Li 2012). Moreover, bile salts have been reported to generate oxygen-free radicals, thereby altering RNA secondary structure and inducing DNA damage (Begley and others 2005).

dos Santos Leandro and others (2013) tested the viability of *L. delbrueckii* UFV H2b20 incubated under simulated GIT conditions (3 h in gastric juice, pH = 3.0) followed by a 12 h incubation in MRS broth (containing 0.3% w/w bile salts) and reported very good strain resistance in both cases (8.80 to 8.90 logCFU/g and 9.1 to 9.2 logCFU/g, respectively). The authors also reported that the fat content of the product did not significantly affect the strain's survivability.

In a comprehensive study, Ranadheera and others (2012) evaluated the impact of gastric pH (2.0, 3.0, and 4.0), bile salt

concentration (0% and 0.3%, w/w), and matrix system (ice cream and plain and fruit yogurt) on the viability of probiotic bacteria (*L. acidophilus* La-5, *B. animalis* Bb-12, and *Propionibacterium jensenii* 702). The authors demonstrated that the bacterial viability was largely affected (viability loss >4 logCFU/g) only at low pH (2.0), with *B. animalis* Bb-12 and *P. jensenii* 702 showing good acidic tolerance. This was mainly attributed to either the intrinsic resistance of probiotics at moderately acidic environments or the lower activity of pepsin at higher pH values. In the presence of bile salts, a viability loss up to 2 logCFU/g was noted, with *B. animalis* Bb-12 being the least vulnerable. With respect to the food matrix, it was deduced that ice cream generally provides good protection against GIT conditions due to the peculiar compositional features of ice cream, namely, high milk fat content, presence of polysaccharides, and cocoa powder in the case of chocolate flavor formulations. In a series of studies, Alamprese and others (2002, 2005) investigated several growth inhibitors known to influence gut colonization by probiotics, including bile salts, antibiotics, and sodium dodecyl sulfate (SDS) on the survival of *L. johnsonii* La1 and *L. rhamnosus* GG. Both strains showed that high resistance to most of the tested stress factors, and only at very low pH values (pH = 1.5) was a detectable decrease in survival observed (approximately 4 logCFU/g).

Ranadheera and others (2012) observed a strain and matrix dependency when scrutinizing the effect of dairy matrix on the adhesion properties of *L. acidophilus* La-5, *B. animalis* Bb-12, and *P. jensenii* 702 to Caco-2 cells. The probiotic bacteria used in probiotic ice cream exerted satisfactory adhesion ability (Figure 1) that was slightly lower than that of fruit stirred yogurt but higher than that of plain yogurt. Although it was not fully elucidated how ice cream improves the adhesion properties of probiotic cells, it was highlighted that prolonged frozen storage did not adversely impact adhesion ability. In another study, Deepika and others (2011) investigated the adhesion ability of *L. rhamnosus* GG to hexadecane (a marker of the bacterial cell surface hydrophobicity) and Caco-2 cells as influenced by the matrix (yogurt and ice cream with different fat and sugar contents) and storage time. The hydrophobicity of the bacterial cell surface in both systems increased during the first 3 d of storage, whereas a significant reduction was observed after the 1st wk of storage. No significant impact of matrix components (fat and sugars) was detected except for sugar-free and fat-free systems (lower surface hydrophobicity). The adhesion ability of *L. rhamnosus* GG to Caco-2 cells reached the highest level after 3 d of storage, while no impact of the fat or sugar content was detected. Yogurt is generally a better substrate for enhancing the adhesion properties of *L. rhamnosus* GG mainly because of the lower surface charge of the cells under acidic conditions (assessed by ζ -potential measurements) prevailing in yogurt, thus favoring adhesion primarily via hydrophobic and less via electrostatic interactions.

Pandiyan and others (2012c) investigated the survivability of *L. acidophilus* NCDC 14 and its impact on the fecal microflora in the GIT tract of healthy human volunteers receiving synbiotic ice cream for 15 d (supplemented with FOS and whey proteins). The pH of the collected stool samples was significantly reduced throughout the ice cream ingestion period, reaching its lowest value on day 15 due to the formation of short-chain fatty acids by the colonic microbiota. Microbiological counts of the fecal samples after 15 d of ice cream consumption revealed a 1 log CFU/g increase of *L. acidophilus* NCDC 14 in the gut epithelium. Finally, synbiotic ice cream consumption contributed to a reduced coliform load, attributed to the antagonistic adhesion

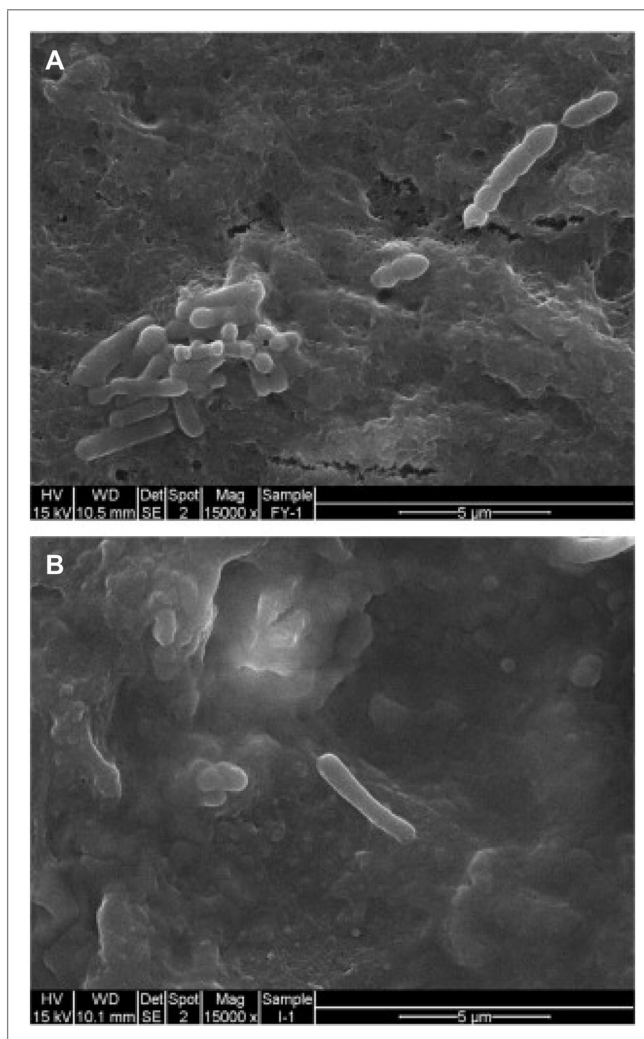


Figure 1—Adhesion of probiotic bacteria cells onto Caco-2 cells in stirred fruit yogurt (A) and plain ice cream (B) made from goat milk. (Source: Ranadheera and others 2012; image reproduced with the permission of Elsevier).

between probiotic and pathogenic bacteria and the production of bacteriocins by *L. acidophilus* (Pandiyan and others 2012c).

Prebiotics Technological and health aspects

According to Roberfroid (2007), the term prebiotics is used to describe “selectively fermented ingredients resulting in specific changes both in the composition and activity of the GIT microbiota that confer benefits upon the host’s well-being and health.” Most prebiotics are primarily composed of oligosaccharides or more rarely may contain also polysaccharides, such as inulin (Saad and others 2013). Prebiotics offer a considerably broad spectrum of technofunctional, physiological, and nutritional aspects. Depending on the type, degree of polymerization (DP), and degree of branching (DB), prebiotics can exert significant texturizing properties, as they may retain water, interact with milk proteins, and form aggregates composed of hydrated microcrystals (Schaller-Povolny and Smith 2002; Meyer and others 2011a). Moreover, the ability of several prebiotic fibers including inulin, polydextrose, oligofructose, wheat dextrin to promote emulsion, and foam stability, to enhance viscosity, to induce gelation, and to

mimic fat by reducing friction and imparting tongue lubrication during consumption has been reported (Fagan and others 2006; Meyer and others 2011b; Tárrega and others 2011; Soukoulis and Fisk in press). From a physiological point of view, the majority of the health claims for prebiotics are associated with their ability to modulate the colonic function by stimulating the growth of beneficial probiotic bacteria and suppressing the viability of pathogenic microorganisms such as *Escherichia coli*, *Campylobacter jejuni*, *Enterobacterium* spp. *Salmonella enteritidis*, and others (Saad and others 2013). In addition, ingestion of prebiotics has been associated with immunomodulatory effects, prevention of colorectal cancer, regulation of serum cholesterol and triglyceride levels, improvement of mineral absorption and bone mineralization, reduced plasma glucose levels, and anti-inflammatory and anticariogenic properties (Saad and others 2013).

The prebiotic benefits related to health and technological aspects have been exploited in a series of food products, including milk and related beverages, semisolid dairy products, such as yogurt, custards, and spreads and aerated dairy desserts, also o/w and w/o emulsions such as salad dressings and mayonnaise, as well as bakery and pasta products, extruded foods, and breakfast cereals (Brennan and others 2004; Homayouni and others 2008a; Debon and others 2010; Tárrega and others 2011; Mantzouridou and others 2012). The synbiotic concept has often been adopted during ice cream prototyping, in order to combine fair product texture and stability with enhanced survival of probiotics during GIT digestion.

Prebiotic functionality in ice cream systems

Many food applications of prebiotics refer to synbiotic ice cream prototyping due to their ability to stimulate probiotic growth in the digestive tract, much research (Table 2) has been conducted for exploiting the technological properties of FOS, polydextrose, and resistant starch (as fat replacers for mouthfeel and as body enhancers, and also as cryoprotectants, thickeners, and foaming agents) (Karaca and others 2009; Soukoulis and others 2010b). Due to their lower molecular weight compared to polysaccharides, starch hydrolysates (such as maltodextrins), corn syrup solids, and prebiotics exert a fair impact on colligative food properties, depending on their DP and DB. More specifically, the presence of inulin and oligofructose in model sucrose solutions and ice cream systems at 1.6% to 4.8% w/w was reported to affect their thermophysical properties such as freezing point, unfrozen and freezable water content, and glass transition temperature (Soukoulis and others 2009; Soukoulis and others 2010b).

The increase of carbon chain length and the branching degree appear to be related to the impact on freezing point temperature (FPT) and glass transition phenomena, with the higher DP inulin exhibiting a better cryoprotective potential compared to oligofructose (Soukoulis and others 2010a). In addition, long-chain inulin as a replacer of corn syrup solids (42 DE) led to significant reduction of mean ice crystal size and FPT depression of low-fat ice cream, supporting its cryoprotective role (Schaller-Povolny and Smith 2001). Although the mechanisms of the cryoprotective activity of FOS in ice creams are not fully explored, it has been attributed to synergistic effects in retaining and binding water, to reduced amounts of freezable water, to increased microviscosity, to controlled water diffusion in the freeze-concentrated serum phase, and to ice crystal size distribution (Schaller-Povolny and Smith 2001; Soukoulis and others 2009). Polydextrose has been used extensively as a cryoprotective bulking agent in low-fat or fat-free ice cream formulations due to its ability to reduce the FPT of ice cream mixes compared to sucrose (Baer and Baldwin 1984;

Alvarez and others 2005; Whelan and others 2008b). Moreover, adding polydextrose to ice cream can increase the uniformity of ice crystals, leading to enhanced creaminess and mouthfeel perception (Alvarez and others 2005).

The ability of FOS to improve the viscosity of dairy o/w emulsions (for example, dairy desserts and ice creams) is well established (Schaller-Povolny and Smith 2001; El-Nagar and others 2002; Karaca and others 2009; Debon and others 2010; Isik and others 2011; Tárrega and others 2011). Parameters such as DP, DB, and FOS concentration have been reported to impact the viscosity of semisolid dairy dessert systems (Tárrega and others 2011). Apart from the contribution of viscosity to the freezing-whipping process (a minimum viscosity is required for inducing fat destabilization and air cell stabilization), viscosity has been reported to interrelate with texture perception (creaminess, wateriness, mouth coating, tongue lubrication, friction, and roughness) during consumption (Soukoulis and others 2008; González-Tomás and others 2009; Soukoulis and others 2010b; Bayarri and others 2011).

Inulin and oligofructose have also been employed in ice creams as sugar replacers (for sucrose or corn syrup solids), exerting a significant enhancement of macroviscosity. This is likely not only due to an increase of total solids, but also due to the ability of inulin to hydrate and bind water (Schaller-Povolny and Smith 2001; El-Nagar and others 2002; Soukoulis and others 2009; Soukoulis and others 2010b). Karaca and others (2009) deduced that polydextrose did not impart viscous characteristics to regular, low-fat, and nonfat ice creams, while modified starch and inulin induced a prominent increase of macroviscosity. Contrary to other studies, the authors also observed that the thickening ability of inulin was strongly dependent on the milk fat content of ice cream, with nonfat formulations promoting the highest viscosity. In line with the former observations, no significant effects of adding polydextrose on the rheological profile of aged ice cream were reported by Whelan and others (2008a), Alvarez and others (2005), and Roland and others (1999). The feasibility of microfluidized and heat-treated resistant starch as thickening agents in ice cream was studied by Augustin and others (2007). The investigators reported a pronounced viscosity increase (7.6 cP) in the presence of resistant starch compared to native high-amylose starch and no-added starch (2.4 and 1.3 cP, respectively).

Prebiotics may also facilitate air incorporation and foam stabilization by increasing the viscosity of the aqueous phase (by increasing solute concentration or gelation) surrounding the air cell interface, raising the physical barrier against air cell destabilization via Ostwald ripening (Franck 2002; Herceg and others 2007). Although there is a lack of studies regarding the functionality of prebiotics such as FOS or resistant starch to act as colloids, most ice cream studies demonstrated that inulin, oligofructose, or resistant starch substantially enhanced air incorporation (overrun) and related properties such as melting resistance and shape retention (El-Nagar and others 2002; Augustin and others 2007; Akalin and Erişir 2008; Aykan and others 2008; Karaca and others 2009; Soukoulis and others 2010b).

Increasing the fat and sugar contents of frozen dairy desserts is further associated with superior texture and flavor characteristics such as enhanced mouth coating, tongue lubrication, creaminess, flavor release, reduced iciness, coarseness, friction, and wateriness (Koeferli and others 1996; Prindiville and others 1999). Ice cream reformulation on a customized dietary basis, for example, low-fat or low-sugar content, limited use of food additives, and so on, is challenging as in most cases texture, flavor and taste deterioration is experienced. The ability of many prebiotics to mimic fat

Table 2—Overview of studies investigating the impact of dietary fiber (excluding polysaccharides) on the technological and sensory properties of ice cream.

Dietary fiber	Functionality	Ice cream system	Rheological/mechanical properties	Thermophysical properties	Overrun and melting behavior	Sensory profile	Reference
Inulin	Fat replacer	Low-fat (5%) yoghurt-ice cream with 5%, 7%, and 9% inulin	Increase of consistency coefficient Increased stickiness Firmness reduction	Nd	Improvement of melting resistance	Improvement of smoothness Reduction of grittiness, iciness, coarseness, and hardness	El-Nagar and others (2002)
Inulin	Fat replacer	Low-fat (3%) ice cream	Nd	Nd	Nd	Impairment of perceived texture No impact on flavor, appearance, and color	Devereux and others (2003)
Inulin	Fat replacer	Low-fat ice cream (0%, 2%, and 4% fat) with 8%, 6%, and 4% inulin	No impact on hardness at 4% and 6% inulin, increased hardness at 8% Increase of consistency coefficient and pseudoplasticity at 8%	Nd	Increased overrun at 6% and 8% inulin Decrease of melting rate at 6% and 8% inulin	Flavor and taste score reduction (4% and 8%) Reduction of body and texture scores (8%)	Karaca and others (2009)
Inulin	Fat replacer	Low and reduced fat ice cream (3% and 6%) with 4% inulin	Reduction of consistency coefficient, apparent viscosity, and pseudoplasticity Increase of hardness	Nd	Overrun reduction by fat lowering Improvement of melting resistance	Nd	Akalin and others (2008)
Inulin	Fat replacer	Low (1.6%) and nonfat ice cream (5.7% and 6.2% inulin)	Viscosity enhancement	Nd	Decrease of overrun No impact on melting rate Improvement of shape retention	No impact on appearance, color, and flavor attributes Improvement of body and texture	Aykan and others (2008)
Inulin	Corn syrup solids (CSS) replacer	Reduced fat ice cream CSS (6.7%) were partially (50%) or fully replaced by inulin	Nd	Nd	Nd	Reduced iciness and increased chewiness in fresh and thermally abused samples Reduction of flavor and sweetness intensity of fresh and heat shocked samples	Schaller-Povolny and Smith (1999)
Inulin	Sugar replacer	Full-fat ice cream 10% and 30% sucrose replacement	Increase of consistency coefficient, apparent viscosity (at 50 s ⁻¹), and pseudoplasticity Increase of instrumental hardness	Elevation of T_g , increase of EMW, nonfreezable water amount, and ice crystals homogeneity	Improved overrun at 30% of sucrose replacement Reduction of air cells mean size Improvement of melting resistance	Reduction of perceived vanilla flavor, iciness, coarseness, brittleness, wateriness Increase of hardness, gumminess, and creaminess	Soukoulis and others (2010a)

(Continued)

Table 2—Continued.

Dietary fiber	Functionality	Ice cream system	Rheological/mechanical properties	Thermophysical properties	Overrun and melting behavior	Sensory profile	Reference
Inulin	Dietary fiber fortification	Full-fat ice cream mixes with 2% or 4% inulin	Viscosity and consistency coefficient enhancement at 4% Increase of shear thinning behavior at 4% Increased hardness at 5% and 10%	Elevation of T_f and EMW at 4% Elevation of T_g at both concentrations	Nd	Nd	Soukoulis and others (2009)
Oligofructose	Sugar replacement	Full-fat ice cream 10% and 30% sucrose replacement by oligofructose	Minor impact on apparent viscosity (50 s^{-1}) and pseudoplasticity Enhancement of consistency coefficient	Elevation of T_g , increase of EMW, nonfreezable water amount, and ice crystal homogeneity	Improved overrun at 30% of sucrose replacement Reduction of air cell mean size Improvement of melting resistance	Reduction of perceived iciness, coarseness, brittleness, wateriness Increase of hardness, gumminess, and creaminess More intense vanilla flavor and sweet taste	Soukoulis and others (2010a)
Oligofructose Polydextrose	Fat replacer	Low-fat (4%) ice cream	Nd	Nd	Nd	Lower coldness intensity No impact on creaminess, mouthcoating, and smoothness	Liou and Grün (2007)
Polydextrose	Sugar replacer—GI index reduction	Low GI ice cream Blends of polydextrose with tagatose, fructose, and erythritol	No impact on apparent viscosity (30 s^{-1}) Reduction of hardness	No impact on freezing point depression	Improved fat droplet destabilization Fat droplets mean size reduction dependent on blend composition	Primary sensory attributes (sweetness intensity, vanilla flavor, and dairy flavor) were impaired Good consumers' acceptability	Whelan and others (2008a)
Wheat dextrin	Sugar replacer	Full-fat ice cream 10% and 30% sucrose replacement					Soukoulis and others (2010a)

(Continued)

Table 2–Continued.

Dietary fiber	Functionality	Ice cream system	Rheological/mechanical properties	Thermophysical properties	Overrun and melting behavior	Sensory profile	Reference
Oat fiber	Dietary fiber fortifier	Ice cream mixes with 2% or 4% of oat fiber	Viscosity and shear thinning behavior enhancement at 4%	Depression of T_f at both concentrations Decrease of EMW at 2% Elevation of T_g at 4%	Nd	Nd	Soukoulis and others (2009)
Wheat fiber	Dietary fiber fortifier	Ice cream mixes with 2% or 4% of wheat fiber	Viscosity and shear thinning behavior enhancement at 4%	Depression of T_f at both concentrations Decrease of EMW at 2% Elevation of T_g at 4%	Nd	Nd	Soukoulis and others (2009)
Apple fiber	Dietary fiber fortifier	Ice cream mixes with 2% or 4% of apple fiber	Viscosity and shear thinning behavior enhancement at both concentrations Viscosity reduction	Elevation of T_f , EMW and T_g at both concentrations	Nd	Nd	Soukoulis and others (2009)
Citrus fiber	Stabilizing agent replacer	Ice cream without (0.4%, 0.8%, and 1.2% citrus fiber) or with added stabilizers (0.4% stabilizer and 0.4%, 0.8%, and 1.2% citrus fiber)		Nd	Improvement of melting resistance when combined with stabilizers	Nonperceivable differences in flavor, body, and texture of fully or partially stabilized systems	Dervisoglu and Yazici (2006)
	Fat replacer	Nonfat ice cream with 0.74% of orange peel fiber or orange peel-bagasse-seed fiber	Nd	Nd	Nd	Adverse effects on organoleptic quality in the individual citrus fiber systems Texture and color similar to full fat formulation Adverse impact on odor, flavor, and aftertaste	De Moraes Crizel and others (2013)
Soluble soybean polysaccharides (SSPS)	Fiber fortification	Nonfat ice cream 0% to 5% SSPS	Increase of consistency index No effect on pseudoplasticity Increase of firmness	Nd		No impact on appearance Decrease of flavor and sweetness intensity Adverse impact on texture	Chen and others (2010)

Nd = not determined (no data). EMW = effective molecular weight, T_f = freezing point temperature, T_g = glass transition temperature, GI = glycemic index.

technofunctionality has been extensively exploited in the manufacture of health-related food products such as low-fat or nonfat and low GI bakery and dairy food items (Franck 2002). Especially inulin has a dominating role on the development of low-fat and/or low-sugar ice cream due to the ability of its native or long-chain-like structure to create gels of various firmness via microcrystal aggregation and agglomeration (Meyer and others 2011a). In general, inulin has been reported to reduce iciness and to enhance chewiness of thermally abused ice creams via its ability to impact colligative properties. Similarly, partial substitution of sucrose by inulin or oligofructose improved texture and sensory parameters such as iciness, coarseness, wateriness, and brittleness. The ability of FOS to change ice crystallization/recrystallization processes via elevating the freezing point and reducing unfrozen water mobility was suggested as governing factors for these effects (Soukoulis and others 2010b).

Dietary Fiber

Definitions, technological, and physiological aspects

According to the Codex Alimentarius Commission, DF includes naturally occurring and enzymatically, chemically, or physically isolated and synthetic, edible carbohydrate polymers with 10 or more monomeric units, which are not hydrolyzed by endogenous enzymes in the human small intestine (Codex 2009). DF can be further classified into soluble and insoluble DF, high-molecular-weight (polysaccharides), and low-molecular-weight (oligosaccharides) DF (Westenbrink and others 2013), as well as into fermentable (by colonic bacteria) and nonfermentable fiber such as cellulose and lignin. While prebiotics are also fermentable DF, the focus in this section is on fibers that have not shown benefits on the colonic microbiota.

DF in food products exerts a wide range of technological aspects, including water-holding (WHC), water-swelling (WSC), and water-retention (WRC) capacity, influencing water solubility, oil holding capacity (OHC), viscosity, texture and texture stability, gel-forming capacity, and even antioxidant capacity (Elleuch and others 2011). Pectins, gums, and mucilages are generally characterized as water soluble, while cellulose, most types of hemicellulose, and lignin comprise the insoluble fraction of DF (Chawla and Patil 2010). Both water solubility and hydration properties are influenced by the fiber structure, presence of hydrophilic functions such as $-\text{OH}$, $-\text{COOH}$ or $-\text{SO}_4^{2-}$, temperature, and ionic strength (Elleuch and others 2011). On the other hand, viscosity is mainly affected by the type and concentration of soluble DF, while the impact of the insoluble DF fraction on the rheological behavior of aqueous systems is rather limited. In addition, depending on concentration (below or above the critical concentration c^*), pH, ionic strength, and temperature, soluble DF macromolecular conformation can vary from the dilute to the overlapping (entangled) state (Foster and Wolf 2011). Many random-coil soluble DFs such as alginates, carrageenans, gellan, or locust bean gum have the ability to form cross-linked gel networks that can be pH- or temperature-reversible. The latter has found many applications in the structuring and texturizing of semisolid and solid food systems.

DF incorporation into food products does not only aim to customize structure and texture, but also to provide specific dietary and physiological benefits. The physiological role of DFs is mainly attributed to their ability to enhance water binding, and viscosity/gel formation in the GIT tract, thus increasing fecal bulk, promoting peristalsis, and modifying colloidal elements of the food matrix, their biochemical aspects, and their impact on the

large bowel microbiota diversity and metabolic activity (Brownlee 2011). Adoption of a fiber (especially insoluble)-rich diet has been reported to be associated with reduced gut transit time, facilitating frequent defecation. Animal studies have also shown that the ingestion of fermentable DF, such as FOS or gum arabic, can affect the T-cell composition of gut-associated lymphoid tissue (GALT) compared to nonfermentable DF. On the other hand, it has been shown that both insoluble and soluble DF fractions can exert a protective effect on the colonic mucus barrier (Brownlee 2011). Ingestion of fair amounts of nonfermentable DF (about 30 g/d) is known to contribute to the balance of the colonic microbiota. Generally, the presence of nonfermentable DF in the digesta favors the growth of the native microbiota such as *Bifidobacteria* or *Lactobacilli*, resulting in the formation of short-chain fatty acids, in particularly butyric acid. The latter is believed to play an important role in modulating the health of the colonic epithelium (Topping and Clifton 2001), which is likely to be due to the anti-inflammatory properties of short-chain fatty acids (Vinolo and others 2011).

Additional health benefits conferred by consuming DF-rich food include lowered blood cholesterol levels and reduced postprandial serum glucose response, reducing the risk of developing obesity, cardiovascular disease, and type 2 diabetes (Hodge and others 2004; Brownlee 2011; Mudgil and Barak 2013). It is generally considered that lowering total and low-density lipoprotein cholesterol (LDL-C) takes place via altering cholesterol (re-)absorption, hepatic metabolism, and plasma clearance of lipoproteins (Mudgil and Barak 2013). DFs that promote viscosity enhancement or gelation (for example, oat- β -glucans, pectins, and guar gum) have been reported to constitute superior modulators of blood lipids (Mudgil and Barak 2013). In a recent randomized cross-over clinical study, it has been shown that the inclusion of high-viscosity fiber in the diet was able to reduce the plasma LDL-C by 9% and 17% compared to medium or low-viscosity DF-rich diets, respectively (Vuksan and others 2011). In addition, it has been reported that the ingestion of highly viscous DF was strongly correlated with reduced postprandial plasma glucose levels (Chutkan and others 2012). Several viscosity-related mechanisms, such as increased chyme viscosity, a decrease of the GI of ingested food, reduced glucose absorption, and reduced starch degradation in the small intestine in conjunction with improved hormonal responses to nutrients, may explain the impact of highly viscous DF on postprandial plasma glucose response (Chutkan and others 2012).

Despite these physiological aspects, polysaccharides are used in ice cream in rather limited amounts, usually not exceeding 0.5% to 0.6% w/w, and thus other food products can be considered better sources of DF. Except for prebiotics (FOS, inulin, resistant starch, and polydextrose) that can be added in remarkably higher amounts (up to 4% w/w) due to their limited thickening effect, attempting to further increase the amount of soluble fiber will lead to undesirable effects such as overstabilization and poor processing properties and handling. Chen and others (2010) used water-soluble soybean polysaccharide (SSPS) as an alternative to increase DF (from 1% to 4% w/w) in dairy desserts, including low-fat ice cream. The increased SSPS concentration induced a significant viscosity increase compared to conventionally stabilized analogs, although the changes in hardness and melting rates were rather minor. Mapping the consumers' willingness to purchase DF-rich ice cream formulations revealed a "moderate likely" intent to consume the SSPS-fortified products containing 2% w/w of SSPS, one of the highest levels for nonfat ice creams (Chen and others 2010).

Enrichment of ice cream with insoluble fiber

Enrichment of ice cream with insoluble fiber has been investigated in a series of studies to attempt improving its rheological properties, storage stability, and melting resistance, but also in order to provide health benefits (Dervisoglu 2006; Dervisoglu and Yazici 2006; Soukoulis and others 2009; de Moraes Crizel and others 2013).

The addition of DF (oat, wheat, and apple) with a high content of insoluble matter (45% to 93% w/w) into ice cream enhanced macroviscosity and induced a significant elevation of the glass transition and melting point of the frozen systems (Soukoulis and others 2009). The cryoprotective effects of oat and wheat fiber were attributed to their ability to retain high amounts of water, leading to hindered mobility of the water molecules in the freeze-concentrated serum. On the other hand, the presence of pectin, for example, in the case of apple fiber, can also contribute to cryopreservation of ice cream by controlling water molecule mobility in the unfrozen aqueous phase due to its thermodynamic incompatibility with the proteins present, exerting phase separation (Soukoulis and others 2009).

In a recent study, de Moraes Crizel and others (2013) have investigated the use of orange peel and orange peel-pulp-seed-isolated fiber (rich in soluble and insoluble fiber and total phenolics) as a potential fat replacer. The authors reported that incorporating both orange fiber types exerted a fat-mimetic function, allowing a 70% fat reduction (from 18 to 5 g/100 g in the finished product), with no significant modification of color, odor, and texture despite a slight impairment of the flavor and overall acceptability of the low-fat formulation. The study by de Moraes Crizel and others (2013) showed that the consumers' purchase intent did not decrease dramatically toward low-fat formulations (74% compared to 94% for regular ice cream), which appeared to be associated with a growing number of health aware and functional-food-conscious consumers.

In a series of studies, Dervisoglu (2006) evaluated the feasibility of using food industry by-products rich in insoluble DF, such as citrus fiber, hazelnut flour, and hazelnut kernel skin. Citrus fiber as an individual stabilizing agent (0.4% to 1.2% w/w) did not significantly impact viscosity development and air incorporation. However, increased citrus fiber content was associated with a remarkable improvement of ice cream melting resistance (Dervisoglu and Yazici 2006). Fortification of nonfat ice cream with hazelnut flour (up to 3.0% w/w) significantly improved overrun, viscosity, and melting resistance of ice creams without impairing appearance, flavor, body, and texture. On the other hand, hazelnut kernel skins, apart from its good foam formation and stabilizing ability, induced a significant deterioration of textural and sensory properties of the finished products (Dervisoglu 2006).

Low Glycemic Index Sweeteners

Overview

The GI concept of foods has been introduced in order to provide a quantifiable marker of the response of a specified amount of ingested carbohydrate compared to a standard food reference, for example, glucose or white wheat bread (Martin and others 2008). A high-GI food is expected to cause a faster and higher rise of blood glucose levels than a food with a low GI, given the same carbohydrate content. Factors such as DF content, the nature of starch and monosaccharide components, viscosity, particle size, food processing, ripeness, storage practices, the presence of alpha-amylase inhibitors, nutrient-starch interactions, and so on, have been reported to influence the GI (Agustin and others 2002).

Lowering the GI of food products has been associated with several health benefits, including decreased incidence of obesity, reduction of diabetes type 2 and coronary heart disease risk, and the prevention of hyperinsulinemia/insulin resistance-induced cancer such as colorectal, breast, and prostate cancer (Agustin and others 2002). Popovich and others (1996), studying the GI of several common food products, reported that a typical vanilla ice cream can be considered as a low-GI (42 ± 5 using dextrose as a reference) food. In a later study, Foster-Powell and others (2002) reported that the GI of ice cream depended on its formulation, with gelato and chocolate-frozen dairy desserts being characterized by the highest GI values, namely, 57 to 80.

Over the last decade, a constant increase of diabetes (the joint occurrence of diabetes type 2 and obesity) incidence has been recorded; the prevalence of diabetes from 2000 to 2030 is expected to rise from 171 to 366 million people worldwide (Ben-Avraham and others 2009). This has also impacted the ice cream industry toward producing products of reduced GI, targeting mainly consumers suffering from diabetes.

Reduction of GI in ice cream cannot be carried out by simply removing sucrose, as the resulting product would lack adequate scoopability, mouthfeel, and sweetness (Marshall and others 2003). Rather, the partial or total substitution of high-GI sweeteners (sucrose, dextrose, and corn syrup solids), by low-GI sweeteners (polyols, fructose, and the noncaloric sweeteners such as aspartame, cyclamate, or saccharin) is common practice to reduce the GI (Abril and others 1982; Goff and Jordan 1984; Özdemir and others 2003; Maia and others 2008; Ozdemir and others 2008; Whelan and others 2008a, 2008b; Soukoulis and others 2010a; Soukoulis and Tzia 2010). In most cases, the GI reduction is accompanied by a significant change of the colligative properties of ice cream (FPT, fraction of frozen water, and glass transition temperature), as well as growth and morphological aspects of ice crystals impacting texture and storage stability (Hagiwara and Hartel 1996; Miller-Livney and Hartel 1997; Ablett and others 2002; Muse and Hartel 2004).

Caloric sweeteners

Fructose has a GI of 14 to 23, is naturally occurring in fruits and vegetables, and is of higher sweetening power (1.2 to 1.7 times) than sucrose (O'Brien-Nabors 2001). In addition, its metabolism is insulin-independent. Fructose can be incorporated into ice cream either in pure crystalline form or as corn starch hydrolyzates that is high fructose corn syrups (HFCSs). However, it should be noted that the GI of the latter is pronouncedly higher than that of fruit- and vegetable-derived fructose, due to the presence of additional sugars. Studies on healthy and diabetic (insulin- and noninsulin-dependent) subjects have demonstrated that fructose intake produces a lower postprandial rise of glucose and insulin (Bantle 2005). Moreover, the passive absorption of fructose from the intestine and its high postprandial thermogenic response and hepatic oxidation has been reported to be associated with improved satiety (Melanson and others 2008). On the other hand, recent studies have shown that adopting diets high in fructose may lead to obesity, possibly due to inducing a rather lipogenic response. Furthermore, the strong impact of fructose on the colligative properties of ice cream may impair its texture (development of iciness and coarseness) and storage stability due to acceleration of ice recrystallization phenomena. For this reason, fructose is only used as a co-component of the bulk sweetening agents for improving mouthfeel, scoopability, and enhancing sweetness. Abril and others (1982) have demonstrated that combining fructose with low

DE maltodextrin and xylitol can impart texturizing properties and increase consumer acceptability. Partial substitution of sucrose by fructose has been reported to depress the freezing point and glass transition temperature of ice creams, reducing the instrumental hardness and melting resistance and increasing overrun (Soukoulis and others 2010b; Silva Junior and Lannes 2011). Despite the adverse impact of fructose on icy, coarse, and watery mouthfeel, it was shown that sucrose substitution by fructose (up to 30% w/w) can significantly promote flavor release (Soukoulis and others 2010b).

Polyols (polyhydric alcohols) are poly-hydroxyl compounds originating from their parent reducing sugars (O'Brien-Nabors 2001). Polyols are hygroscopic, heat- and acid-stable compounds that do not undergo Maillard reactions. They have a strong impact on colligative properties of aqueous solutions (many of them are widely used for cryopreservation of biological tissues) and they are characterized by a lower GI and lower to equal sweetening power (0.3 to 1.0) compared to sucrose (O'Brien-Nabors 2001). In addition, depending on the polyol type, several physiological features have been described, namely, low caloric load, anticariogenic properties, and low absorption in the digestive tract. On the other hand, excessive use of polyols (> 10 to 20 g for mannitol and xylitol, 23 g for sorbitol, and 29 g for isomalt) may lead to laxative effects and impair flavor and taste characteristics, resulting in cooling effects, and formation of unnatural or metallic off-flavors.

Polyols such as maltitol, sorbitol, xylitol, mannitol, and isomalt have been implemented in the manufacture of low or nonsugar ice creams (Özdemir and others 2003; Bordi and others 2004; Whelan and others 2008a; Soukoulis and others 2010a). In a series of studies, maltitol has been identified as a very efficient ingredient for lowering the GI and the sugar content of ice cream without compromising texture, flavor, or taste. Özdemir and others (2003) developed an ice cream product suitable for diabetic patients based exclusively on sorbitol or maltitol. The authors demonstrated that blood glucose concentration was reduced from 381 mg/100 g (sucrose-based product) to 104 and 108 mg/100 g in the case of sorbitol- or maltitol-based analogs, respectively.

Carlson and others (2011) developed a method to produce LGSs by the reaction of sucrose with an acceptor (sugars or sugar alcohols having free hydroxyl groups at 1 or more carbon position numbers, namely, 2, 3, or 6). The reaction was catalyzed by glucan-sucrase and is based on the glucose transfer from sucrose to the acceptor, releasing fructose and glucooligosaccharides. The resulting product has a GI lower than that of sucrose (17 to 38) and technofunctional properties similar to those of corn syrups. The incorporation of LGS in sugar-free ice cream allowed the efficient reduction of their GI without modifying significantly their colligative properties and mouthfeel.

Johannsen and others (2007) patented a method for producing low-GI ice cream formulations based on the technofunctional synergism of polyols, nondigestible nonstarch complex carbohydrates, whey proteins, and starch derivatives. The inventors reported that the developed sugar-free ice creams exerted a GI ranging from 20 to 37 and were organoleptically acceptable. Nonetheless, products were firmer and less sweet than conventional ice creams. In the same study, it was reported that the increase of lactose in ice cream does not only increase the occurrence of malabsorption-related disorders, but that it could also lead to an increased GI of the finished product.

In a similar approach, Anfinson and Tunland (2006) demonstrated that incorporating a blend of low-molecular weight (90 to 190 Da) low-digestible sweeteners (mannitol, maltitol, sorbitol,

lactitol, erythritol, xylitol, tagatose, fructose, and others) and fermentable fibers (inulin, oligofructose, resistant starch, and so on) can promote ice cream functionality and reduce its GI value without altering its colligative and organoleptic properties. According to the investigators, the low-digestible sweeteners tended to retain the original freezing point depression (FPD), sweetness, flavor intensity, texture, tongue lubrication, and mouthcoating. The need to control the FPD upon GI reduction in ice cream has also been highlighted by Whelan and others (2008b) who investigated the impact of different low-GI sweetening blends (polyols and prebiotics) on freezing characteristics, viscosity, overrun, and texture of ice cream. The authors deduced that reformulation of low GI frozen dairy desserts should target and match the FPD curves of the low-GI systems to those of conventional ice cream.

Noncaloric sweeteners

Over the last several years, a strong interest in plant-derived noncaloric sweeteners has arisen sweet diterpenoid glycosides such as ent-kaurene glycosides, sweet triterpenoid glycosides such as cycloartane, oleanane, and curcubitane glycosides, and sweet-tasting proteins or sweetness-inducing proteins (including thaumatin, monellin, and mabillin) did arise (Faus and Sisniega 2004; Pawar and others 2013). These noncaloric sweeteners are of negligible caloric content, but often possess sweetness many times higher than that of glucose. Therefore, small amounts in the final product often suffice. The introduction of sweet glycosides such as stevioside, rebaudioside (*Stevia rebaudiana* leaf extracts), and mogroside V (*Siraitia grosvenorii* fruit extracts) has become very popular in recent years, especially their use as sweeteners (or dietary supplements) for producing soft drinks and energy drinks, tabletop sweeteners, dairy products, fruit/vegetable products, chewing gum, soups, and sauces (Lemus-Mondaca and others 2012).

Apart from their low GI, stevioside-based sweeteners, allowed in the EU since 2011, are also characterized by good heat stability (up to 140 °C) and low degradation in both acidic and alkaline environments (pH range 2 to 10), and they act protectively against the degradation of water-soluble vitamins such as ascorbic acid (Kroyer 2010), present in ice creams containing fruit pulp rich in ascorbic acid. Jookan and others (2012), investigating the stability of steviol glycosides in different foods, including ice cream, reported no detectable degradation of steviol glycosides in ice creams stored at −18 °C for 12 wk. Nevertheless, steviol glycosides cannot totally replace sucrose (due to adverse effects on texture, flavor, and taste features), though they allow to reduce sugar content by up to 30% (from 22.8% to 16% w/w). Blending steviol glycosides with polyols (maltitol and erythritol) has been a good strategy for ice cream reformulation without impairing mouthfeel, taste, body, and texture (Jookan and others 2012). Giri and others (2013) reported that producing low-sugar-content ice cream (6.5% w/w) was feasible by adding steviol glycosides (0.05% w/w), while adverse effects of reducing the sugar content (crystallization and recrystallization phenomena, melting rate, hardness, and sensory traits) can be modulated by adding whey protein concentrate.

Ice cream reformulation using a patented low-GI blend (Trutina Dulcem®) composed of a fruit (kiwi), glycosides (steviosides, steviol glycosides, rebaudiosides, glycyrrhizin, mogroside V, and so on), and a low GI carbohydrate (fructose and polyols) has been successfully demonstrated (de Wees Allen 2008; de Wees Allen 2009). Postprandial *in-vivo* testing of sweet terpenoid glycoside ice creams in adults with or without diabetes symptoms revealed in both cases very low GI and glycemic load (GL) values, varying

from 21 to 24 and 1.8 to 2.1, respectively (serving size 55 g). Moreover, the inventor reported that the presence of Trutina Dulcem in the ice cream did not modify significantly its sensory profile, exhibiting good consumer acceptability, not stimulating fat storage in adipose tissue in both groups.

Ryu and others (2002) developed a method to produce a pulverized mulberry leaf extract product rich in bioactive compounds. The product, having a bland taste and flavor, was incorporated in ice creams at concentrations varying from 0.5% to 5% w/w. Sensory profiling of the mulberry-leaf-containing ice cream showed improved primary organoleptic features, taste, mouthfeel, and overall acceptability (when added at 0.5% to 2.0% w/w). Postprandial monitoring of the plasma glucose levels of 10 healthy subjects revealed a lower glucose blood appearance 45 min after ingestion in the case of ice creams enriched with mulberry leaf extract. It can be assumed that part of the activity was due to the presence of polyphenols, inhibiting glucose uptake from the gut (Ryu and others 2002).

Natural Antioxidants

Overview

Vegetable- and fruit-based food products and their by-products, cereals, pseudocereals, pulses, flours, tea and herbal teas, herb or spice extracts, cold-pressed oils, and so on, are some of the most common food items containing high amounts of antioxidant compounds such as polyphenols, carotenoids, tocopherols, tocotrienols, glutathione, ascorbic acid, and enzymes with antioxidant activity (Boskou 2006; Oreopoulou and Tzia 2006; Pellegrini and others 2008; Ogunlade and others 2009). It is well appreciated that adoption of a diet rich in natural antioxidants appears to be inversely associated with degenerative chronic diseases such as cardiovascular disease, cancer, obesity, and diabetes, as well as with the prevention of general inflammatory health implications and improvements of blood lipids such as cholesterol (Wilcox and others 2004; Bohn 2008).

Although interest in exploiting frozen dairy desserts as a carrier for these bioactive plant compounds against oxidative stress- and inflammation-related diseases has been increasing over the last few years, processing and food quality challenges must be overcome. For instance, several antioxidant compounds are considered as nonnutritive or even antinutritive, susceptible, or unstable under common processing practices such as for ice creams (heat treatment, aeration, and frozen storage), leading not only to partial loss of their physiological activity but also to food discoloration, development of off-flavor and aftertaste, and triggering lipid oxidation reactions. Thus, selection of antioxidants for ice cream prototyping is considered a rather challenging and laborious process.

Vanilla-based extracts

Undoubtedly, vanilla extract is the most common product with potential antioxidant activity used for ice cream production (Marshall and others 2003; Tai and others 2011). Typically, the vanillin amounts in ice cream products range from 5 to 100 ppm (Burdock 2005). Thus, vanilla, apart from its dominating role as flavoring in frozen dairy desserts (Cadena and others 2012), may also exert a fair antioxidant activity, enhancing ice cream resistance against light- or oxygen-induced oxidation that may result in flavor and aroma defects such as metallic, rancid, and cardboard off-notes (Shiota and others 2002). Vanilla extract components such as p-hydroxybenzoic acid, p-hydroxybenzaldehyde, vanillic acid, and vanillin have been held responsible for the potent antioxidant role of vanilla (Charles 2013). According to Kamat and others (2000),

vanillin can provide sufficient protection against protein oxidation and lipid peroxidation occurring in rat liver mitochondria. Dietary vanillin (1.25 to 50 ppm) also exerted a significant hypotriglyceridemic effect in high-fat fed animals at all levels tested, and at the highest concentration a remarkable reduction of body weight and perirenal adipose tissue was detected (Srinivasan and others 2008). Tai and others (2011) demonstrated that vanillin (20 μ M) exhibits a stronger antioxidant capacity than ascorbic acid (20 mM) using different multiple assays such as 2,2-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid) diammonium salt (ABTS^{•+}) scavenging, ORAC, and oxidative hemolysis inhibition (OxH-LIA). Moreover, vanillin (100 ppm) oral administration to mice resulted in detectable amounts of vanillin and its metabolites in plasma and a high antioxidant activity in the ORAC plasma assay. Recently, the ability of vanilla pod extracts (using supercritical fluids) to regulate cholesterol metabolism via upregulating the low-density lipoprotein receptor (LDLR) gene and downregulating the 3-hydroxy-3-methylglutaryl-coenzyme A reductase (HMGCR) gene in liver hepatocellular carcinoma (HepG2) cell lines was demonstrated (Al-Naqeb and others 2010). In addition, Makni and others (2011) studied the antioxidant, anti-inflammatory, and hepatoprotective properties of vanillin in carbon-tetrachloride-treated rats. It was found that vanillin pretreatment of rats prior to CCl₄ administration inhibited hepatic lipid peroxidation and protein carbonyl formation in the liver. Moreover, based on liver histopathology results, vanillin attenuated the expression of proinflammatory cytokines such as tumor necrosis factor α (TNF- α), interleukin-1 β (IL-1 β), and interleukin-6 (IL-6), preventing hepatic cell alteration and necrosis (Makni and others 2011). Nonetheless, the potential health benefits associated with ingesting vanillin in ice cream need to be clinically proven.

Vanillin can also promote free radical scavenging action in food products, providing an effective strategy to control quality loss due to lipid oxidation. Based on early considerations, the addition of pure or concentrated vanilla pod extracts, or synthetic vanilla flavorings was able to hinder oxygen uptake from unsaturated fatty acids and phospholipids after 12 mo of storage of spray-dried ice cream powders (Pyenson and Tracy 1950). Similarly, Gassenmeier (2003) reported that vanillin addition to ice cream mixes after heat treatment improved their stability against autooxidation, responsible for the development of cardboard off-flavors. This was mainly attributed to the inactivation of xanthine oxidase, known to catalyze the oxidation of vanillin to vanillic acid.

Fruits and fruit-based products

Incorporation of fruit preparations (fruit juices, purees, or concentrates) into frozen dairy desserts has been verified as an alternative strategy to increase their phenolic content (Table 3). Addition of frozen wild blueberry puree (5.33% w/w) and juice concentrate (4.92% to 5.33% w/w) was reported as an adequate means to deliver antioxidant functionality in soy-milk-based ice cream without affecting consumer acceptability (Camire and others 2006). Favaro-Trindade and others (2006) demonstrated that ice cream could be an excellent vehicle for delivering combined probiotic-antioxidant functionality by blending acerola fruit juice into ice cream mix fermented by *Bifidobacteria*. This allowed retaining living cell counts and ascorbic acid levels under prolonged frozen storage without altering flavor, taste, and texture. Sun-Waterhouse and others (2013) studied the effects of 3 different varieties of kiwifruit (green, yellow, and red flesh) added to regular ice cream, and they claimed potent health benefits due to the presence of several bioactive compounds, including ascorbic acid,

Table 3—Impact of ingredients containing natural antioxidants on antioxidant activity, technological, and organoleptic properties of ice cream.

Ingredient with antioxidant action	Ice cream system	DPPH inhibition percentage—antioxidant capacity	Total phenolics	Physicochemical properties	Texture and structure aspects	Sensory profile	Reference
Pomegranate peel extract (PPE)	Ice cream with 0.1% or 0.4% w/w PPE	4.9% in control 52.5% and 99% for ice cream with PPE	n.d. in control 73 and 363 mg/100 g for ice cream with 0.1% and 0.4% PPE	Decrease of pH	Nd	Impart sourness, astringency and unnatural flavor No impact on textural aspects Decrease of palatability and sweetness	Çam and others (2013)
Pomegranate peel extract	Ice cream with 0.5% or 1.0% w/w of microencapsulated PPE	n.d. in control 243 and 133 (EC_{50} , g of ice cream/g DPPH)	n.d. in control 327 and 893 mg/100 g for ice cream with 0.5% and 1% PPE	Nd	Nd	No impact on consumers' acceptability No perceivable defects relating to PPE addition	Çam and others (2014)
Pomegranate peel extract	Probiotic ice cream with 0.4% w/w PPE	approximately 55% for control approximately 80% for ice cream with PPE	approximately 30 mg/g for control approximately 540 mg/g for ice cream with PPE	Nd	Nd	No impact on tested sensory aspects (color, texture, flavor, taste, and overall acceptability)	Sagdic and others (2012)
Grape seed extract (GSE)	Probiotic ice cream with 0.4% w/w GSE	approximately 55% for control approximately 68% for ice cream with GSE	approximately 30 mg/g for control approximately 370 mg/g for ice cream with GSE	Nd	Nd	No impact on tested sensory aspects	Sagdic and others (2012)
Pomegranate seed oil (PSO)	Ice cream with 2% or 4% w/w PSO	4.9% in control 5.8% and 8.2% for ice cream with PSO	n.d. in control 69.4 and 66.3 mg/100 g for ice cream with 0.1% and 0.4% PSO	No impact on pH	Nd	Impart astringency, oxidized, and unnatural off-flavors Decrease of palatability	Çam and others (2013)
Peppermint essential oil (PEO)	Probiotic ice cream with 0.03% w/w PEO	Approximately 55% for control approximately 52% for ice cream with PEO	Approximately 30 mg/g for control approximately 50 mg/g for ice cream with PEO	Nd	Nd	No impact on tested sensory aspects	Sagdic and others (2012)

(Continued)

Table 3—Continued.

Ingredient with antioxidant action	Ice cream system	DPPH inhibition percentage—antioxidant capacity	Total phenolics	Physicochemical properties	Texture and structure aspects	Sensory profile	Reference
Herbal tea	Ice cream containing 2.5% or 5% w/w black tea, sage, linden, or chamomile extracts (water extraction at 40 or 80 °C)	Not reported	Approximately 100 mg/kg for control approximately 175 to 415 mg/kg for black tea approximately 165 to 260 mg/kg for sage approximately 180 to 340 mg/kg for linden approximately 120 to 185 mg/kg for chamomile	No significant impact on pH and acidity Decrease of pseudoplasticity and consistency coefficient	Reduced air incorporation in herbal tea extract containing samples	Acceptable taste, odour, color, and appearance Characteristics in ice creams with linden or chamomile Adverse impact on organoleptic quality on black tea or sage addition	Karaman and Kayacier (2012)
Grape wine lees (GWL)	Ice cream with 0.5%, 1%, 2%, or 4% w/w of GWL	1.59 mg/g for control 1.65 to 1.95 mg/g for systems with GWL	0.018 mg/g for control 0.11 to 0.40 mg/g for systems with GWL	Increase of acidity Improved melting resistance	Nd	Flavor, taste, texture, and body scores were maximized at 2% GWL	Sharma and others (2013)
Grape wine lees	Ice cream with 5%, 10%, and 15% w/w of GWL	18% for control 46% to 68% for systems with GWL	1.5 mg/mL for control 2.3 to 3.3 for system with GWL	Reduction of overrun Improvement of viscosity Reduction of pH Reduction of ice fusion enthalpy	Increase of fat destabilization index—fat droplets mean size Reduction of firmness and cohesiveness	Nd	Hwang and others (2009)
Kiwifruit juice	Ice cream containing 49% v/v green, gold, or red flesh kiwifruit juice	0.136, 0.208, and 0.338 Trolox eq. mg/g for ice cream with green, gold, and red flesh and red flesh kiwifruit juice	193, 232, and 399 mg/kg for ice cream with green, gold, and red flesh kiwifruit juice	Improved melting resistance and higher overrun in ice cream with red flesh kiwifruit juice	Storage modulus (<i>G'</i>) and fat destabilization index were higher at red-gold kiwifruit juice ice cream	Nd	Sun-Waterhouse and others (2013)
Persimmon puree	Ice cream containing 8%, 16%, 24%, 32%, or 40% w/w persimmon puree	20% for control 40% to 65% for persimmon puree based systems	Approximately 750 mg/kg for control approximately 975 to 1650 mg/kg for persimmon puree based systems	Enhanced melting resistance No impact on pH Reduction of mixes' viscosity	Lower hardness, work of penetration, and stickiness of persimmon puree based systems	Improvement of melting quality Adverse effect on taste on overall acceptability and taste (24% the most acceptable) No impact on odour, color, and structure aspects	Karaman and others (2014)

DPPH = di(phenyl)(2,4,6-trinitrophenyl)iminoazanium, Nd = not determined (no data).

caffeic acid, catechins, carotenoids (lutein and beta-carotene), salicylic acid, and o-coumaric acid. Moreover, favorable effects were also obtained in terms of rheological behavior, melting resistance, and overrun of the final product, with the red-flesh kiwifruit-supplemented food systems considered as the most acceptable. Recently, it was demonstrated that adding persimmon fruit puree to ice cream contributed to significant improvement of its total phenolic content and 2,2-diphenyl-1-picrylhydrazyl (DPPH) scavenging activity (Karaman and others 2014). In the same study, implementing the technique for order of preference by similarity to ideal solution (TOPSIS) approach, it was shown that sensory properties accounted for 80% of consumer preference response compared to 20% in the case of the health-promoting claims. It was deduced that 24% w/w ice cream enrichment with persimmon puree can satisfactorily meet both the criteria of acceptable organoleptic characteristics and antioxidant activity. On the other hand, Teh and others (2005) demonstrated that reducing the fat content of ice cream supplemented with blueberry concentrate had no significant effect on the amount of anthocyanins, ferrulic acid, and phenolic compounds, which depended exclusively on the amount of the added fruit preparation and thus highlighting that combining fruit preparations with ice cream constitutes a suitable carrier for health beneficial ingredients.

Agricultural waste products

Waste products of the agri-food industry such as molasses or pomace are generally regarded as good sources for bioactive compounds, including natural antioxidants (Ayala-Zavala and others 2011). Despite technological obstacles, in order to overcome exploiting food waste (mainly costs of drying, storage, and transport), their often high content of a broad range of bioactives and micronutrients renders their exploitation for food production purposes attractive (Oreopoulou and Tzia 2006). Important benefits have been revealed by the supplementation of ice cream with grape wine lees, a common winery waste rich in polyphenols and DF that is easily fermented by human intestinal host bacteria (Topping and Clifton 2001; Hwang and others 2009). Hwang and others (2009) evaluated the impact of wine grape lees solids addition (5%, 10%, or 15% w/w) to ice creams, concluding that 5% w/w of wine grape lees may improve rheological characteristics, reduce the freezable content and melting rate without influencing fat destabilization and air incorporation phenomena, while delivering important amounts of polyphenolics such as anthocyanins.

Recently, the feasibility to use pomegranate (*Punica granatum* L.) peel by-products (PBP) rich in phenolic compounds and PUFAs for producing functional ice cream was investigated (Çam and others 2013; Çam and others 2014). According to the findings of Çam and others (2013) adding PBP (at 0.1% and 0.4% w/w) was accompanied by a significant increase of phenolic compounds (mainly punicalagin, ellagic acid, and their derivatives), which was associated with higher DPPH-scavenging activity and more effective inhibition of α -glucosidase activity, the latter reducing the GI of prepared formulations. Similarly, enhanced antioxidant activity was also found following the addition of pomegranate seed oil (PSO) to ice cream (Çam and others 2013). In a subsequent study, Çam and others (2014) indicated that the antioxidant and α -glucosidase-inhibiting action of PBP-entrapped in maltodextrin-spray-dried matrices, compared to native PBP, were not significantly affected. In addition, microencapsulation of PBP appeared to be a good strategy for overcoming restrictions related to adverse organoleptic properties resulting from the presence of

phenolics, such as astringency, unsatisfactory tongue lubrication, and oral mucosa puckering (Çam and others 2014).

The addition of whey protein isolate (WPI) glycosylated with the sugars D-allose and D-psicose into ice cream has been referred to as a novel way for delivering excellent antioxidant properties together with improving foaming and emulsifying capacity (Puangmanee and others 2008). The authors suggested that free radical scavenging activity for the specific formulations was associated with the conjugation of Maillard reaction products found in WPI with the C-3 hydroxyl group of these sugars.

Herbal extracts

Herbal or green tea extracts were found to constitute suitable ingredients to increase the phenolic content of ice creams. In a study by Karaman and Kayacier (2012), the use of brewed black and herbal teas (chamomile, linden, and sage) at 2 temperatures for the production of ice cream suggested that antioxidant activity was both dependent on tea type and brewing temperature, with black tea or linden brewed at 80 °C resulting in the highest antioxidant activity of ice cream. The same authors also highlighted that the used herbal tea extracts may also confer important antimicrobial effects (against *Listeria monocytogenes*) and viscosity enhancements, but in most cases, they were accompanied with moderate deterioration of flavor (astringency, sharpness, and bitterness) and color attributes of the final products (Ozturk and others 2010; Karaman and Kayacier 2012).

Using finely ground green tea (macha) in ice cream has also been utilized for increasing the antioxidant capacity of the final product, but a negative impact on textural organoleptic properties such as astringency, bitterness, formation of clumps, powdery texture, and insufficient air incorporation has been reported. In a recent attempt to overcome these drawbacks, Fukuda (2012) developed a novel ice cream formulation by incorporating nonpolymer catechins and caffeine at usage rate ranging from 0.001 to 0.18. These systems exhibited very good air incorporation capacity and excellent texture and shape retention characteristics without adversely affecting the flavor of the product.

Omega-3,6 Fatty Acids and Polyunsaturated Fatty Acids

Omega-3 and ω -6 fatty acids comprise 2 important classes of PUFAs that are most commonly represented by alpha-linolenic acid and linoleic acid, respectively. Over the last several years, great interest has been built up to incorporate natural food ingredients rich in these fatty acids into other food products to improve their nutritional profile. The major health benefits associated with adopting a diet rich in PUFAs are primarily associated with the prevention of cardiovascular disease, rheumatoid arthritis, cancer, and inflammation, lowering of blood pressure, reinforcement of the immune system, and contributions to brain and nervous systems (Simopoulos 1991; Arab-Tehrani and others 2012). Huang (2011) demonstrated that incorporating omega 3/6 fatty acid extracts from fish or oil from algae in dairy matrices can be used to prevent several oral infections or inflammatory diseases such as caries, periodontitis, gingivitis, mouth ulcers, and halitosis, due to their antimicrobial activity against many oral pathogens.

Recently, several applications of omega-3-rich-oils such as fish, flaxseed, rice bran, and algal oil in frozen dairy desserts were launched (Gonzalez and others 2003; Chee and others 2007; Huang 2011; Song and others 2011). Gonzalez and others (2003) demonstrated that ice cream produced with alpha-linolenic acid fortified milk resulted in low-viscosity ice cream mixes and softer

frozen finished products without impacting oxidative alterations during storage. Similar results were also reported by Goh and others (2006); and (Méndez-Velasco and Goff 2012) who observed that a reduced milk fat to flaxseed oil ratio coincides with reduced product stiffness and melting resistance. This was mainly due to impaired structuring, that is, low partial fat coalescence during the freezing-whipping. The latter drawback may be surpassed by intelligent structure design of ice cream emulsions (ratio of saturated to unsaturated fatty acids, saturation level of the emulsifying agents), high-pressure low-temperature freezing, or the implementation of the 2-phase process that is based on combining an emulsion (fat, some protein, and water) with an aqueous solution composed of sugar solids, stabilizers, and remaining protein (Segall and Goff 2002; Sung and Goff 2010; Song and others 2011; Méndez-Velasco and Goff 2012).

Oleogel technology (for example, use of organogelating agents to provide liquid edible oils with solid-like properties) is a technique that has recently been successfully applied for saturated fatty acid replacement by polyunsaturated ones (Zulim Botega and others 2013a, 2013b). According to Zulim Botega and others (2013a), rice bran wax (RBW) oleogels were effectively emulsified into ice cream mixes, promoting the formation of small gelled fat droplets, as well as showing sufficient protein absorption on oil-water interfaces and successive displacement by competitive surface-active agents. In addition, RBW oleogels enhanced air incorporation and structure formation upon freezing, behaving more like crystallized than liquid lipids (Zulim Botega and others 2013a). However, a proper design of the emulsifying system is required to prevent structural collapse during melting (Zulim Botega and others 2013b). Naturally extracted lipid-rich organelles from oil seeds such as sunflower (*Helianthus annuus*) (oil bodies) with high oxidation stability (Fisk and others 2008) also have shown to be promising sources of PUFAs and as ice cream ingredients, potentially offering a source of oxidation-stable unsaturated fatty acids and additional surface active materials to the ice cream mix (Berry and others 2006).

Impairment organoleptic properties can be a restrictive parameter for their direct incorporation in frozen dairy desserts. Chee and others (2007) showed that consumer preference of omega 3/6-enriched regular ice creams is predominantly influenced by the flavoring system. It has been shown that strawberry flavoring can mask more effectively any fishy-like off-notes than vanilla flavoring (Chee and others 2007). In a recent study, Song and others (2011) incorporated omega-3 lipids in full-fat chocolate probiotic ice cream, demonstrating a perceivable deterioration of product acceptability, mainly due to the development of fishy off-flavors. However, it has also been shown that the presence of omega-3/6 fatty acids can promote survival of probiotic cells when subjected to freeze-induced or mechanical stress during freezing and storage.

The potential to deliver omega-3/6 fatty acids indirectly in dairy products, that is through modification of the diet of cows, has also been reported (Egger and others 2007). Using milk from cows fed with fodder supplemented with seed rich in PUFAs such as linseed, canola, cottonseed, and soybean has been proposed for producing smooth ice cream with improved nutritional value and nonperceivable flavor defects such as oxidized or rancid (Chen and others 2004). Feed fortification for milk enrichment with valuable fatty acids has also been proposed as a practice for dairy product enrichment including ice cream with conjugated linolenic acid (CLA) (Gonzalez and others 2003). According to recent considerations, the most representative isomers of CLA (9-cis, 10-trans, 11-trans, and 12-cis C18:2) appear to exert potent physiologi-

cal actions such as being anticarcinogenic, antiobese, antidiabetic, and antihypertensive (Koba and Yanagita 2013), although results remain controversial. Systematic studies of CLA on structural, textural, and sensory properties of ice cream are scarce, it has been shown that CLA-rich milk fat can provide adequate texture and body qualities (similar to CLA-free milk-fat-based ice cream), despite reduced macroviscosity of the ice cream due to the solid to liquid fat ratio reduction (Gonzalez and others 2003). In the same study, it was pointed out that using CLA-rich milk fat did not appear to impact the lipid autooxidation reactions occurring during ice cream storage. In addition, it has been evidenced that the type of dairy matrix (milk, butter, or yogurt) can critically affect the perception thresholds of CLA-associated defects such as aftertaste, rancidity, acid, and nutty off-flavors (Jimenez and others 2008).

Mineral and Trace Element Fortification

As for milk and milk products, frozen dairy desserts are regarded as good sources of inorganic microconstituents and water-soluble vitamins (Marshall and others 2003). Most of the minerals found in ice cream are present in form of salts in the freeze-concentrated serum phase, except for calcium and phosphorus that exceed their solubility and associate with casein micelles. In addition, some of the trace elements may also be present as complexes with proteins, for example, lactoferrin (Fe), or enzymes, like xanthine oxidoreductase (Fe, Mo) (Walstra and others 2006).

Liposoluble vitamins (vitamin K, D, E, and A) are found at very low amounts in ice cream, while they are absent in low-fat and nonfat formulations (Marshall and others 2003). Thus, mineral and trace element ice cream fortification aims both toward improving the nutritional value and to compensate for losses occurring during processing and storage, particularly due to heat treatment and skimming (Marshall and others 2003). Calcium and iron ice cream fortification is very common although there is also a growing interest in fortification with other micronutrients such as magnesium, zinc, manganese, cobalt, nickel, copper, and provitamin A carotenoids (Noël and others 2012).

Ice cream is considered a good source of calcium (836 mg/kg), but with ranges much lower than that of other dairy products such as milk, cheese, or yogurt (Chekri and others 2012), due to the lower amount of milk in the final product. Nevertheless, ice cream is regarded as an excellent vehicle for calcium administration to different target groups. In a recent study, Ferrar and others (2011) have shown that consumption of calcium-fortified low-fat ice cream by a target population of young women with habitually low calcium intake significantly reduced bone calcium resorption over a 28-d period, without any increase in body weight. The addition of dairy minerals for calcium fortification purposes in ice creams has further been reported as an efficient strategy for improving the bioavailability of calcium, regardless of the compositional diversity of the formulations. Van der Hee and others (2009) reported a similar fractional calcium absorption from butterfat- or coconut oil-standardized ice creams compared to low-fat milk in young adults (25 to 45 y), about 26±8% compared with 31±8%, respectively. It was deduced that changes in the long-chain saturated fatty acid (palmitic, oleic, and lauric) did not affect the intestinal absorption of calcium or the bone mineral mass.

Direct standardization of ice cream mixes prior to or during freezing-whipping has been proposed for mineral and water-soluble vitamin fortification of ice cream (Wendel 2010). Often, however, mineral standardization of ice cream using salts leads to significant technological and nutritional drawbacks, such as protein

and mineral precipitation, color and flavor deterioration, development of mouthfeel defects (chalky and gritty texture), and reduction of heat stability and bioavailability (Augustin 2003). Several strategies to overcome this hurdle have been proposed, with microencapsulation of minerals in hydrocolloid-based carriers or liposomes or complexation with anionic polysaccharides (such as carrageenans, pectins, or alginates) being the most effective strategies (Augustin 2003). In addition, food matrices rich in bioactive compounds including minerals such as fruit juices or concentrates have been proposed as alternatives to increase mineral content. However, changes in organoleptic profile and colloidal aspects of the finished products may be encountered (Soukoulis and Tzia 2010; Dagdemir 2011; Erkaya and others 2012; Karaman and others 2014).

Jacobson and others (2002) postulated that complexing calcium salts with hydrolyzed anionic polysaccharides (including pectins, cellulose, and carrageenans) with a DP ranging from 50 to 80 contributes to the efficient control of protein destabilization, that is, coagulation and precipitation, due to the presence of free calcium ions. Calcium-polysaccharide complexes can be used to stabilize calcium-fortified ice cream mixes without imparting undesirable high viscosity, flavor, and palatability defects. On the other hand, Wedral and others (1998) demonstrated that complexation of soluble calcium salts (calcium gluconate, lactate, or chloride) with alkali metal citrate at a ratio of 1:4 to 4:1 can be used for ice cream fortification, retaining its colloidal stability. Augustin and Williams (2001) exploited the ability of calcium and/or other minerals (iron, zinc, magnesium, and manganese) to migrate into the casein micelles following the addition of an effective amount of phosphate. The method has been claimed to more efficiently bind minerals in the micellar state compared to chelating agents (that is, citrates and polyphosphates), which at high concentrations destabilize the casein micelles, thereby releasing soluble calcium into the serum phase. The authors revealed that ice cream made from fortified milk using the phosphate-induced mineral complexation approach exerted similar melting behavior and textural profile than that composed of nonfortified milk, while only minor differences in the perceived flavor traits were observed.

Incorporation of raisin and grape juice concentrates or sugar cane molasses has been reported as an efficient way to administer iron in regular chocolate ice cream without imparting textural defects or off-flavors (Soukoulis and Tzia 2010). Adding 5% to 15% (w/w) dried Cape gooseberry pulp in conventional ice creams may increase the concentration of several minor abundant minerals (Fe, Mn, Zn, and S) without affecting or only slightly decreasing major abundant minerals, Ca, P, Mg, K, and Na (Erkaya and others 2012). Similarly, the addition of 5% to 20% (w/w) vegetable marrow (*Cucurbita pepo* L.) pulp in ice cream was associated with a significant increase of Zn, K, and Fe content, though it induced a significant reduction of the concentration of major minerals, that is, Ca and P (Dagdemir 2011). Moreover, in both cases, the incorporation of Cape gooseberry or vegetable marrow was accompanied by a remarkable improvement of the viscosity, melting, and sensory quality (Erkaya and others 2012; Dagdemir 2011). Karaman and others (2014) studying the impact of persimmon puree on the mineral content of ice cream revealed a significant increase of K content, though a significant reduction of Ca and Mg concentration was also observed. The authors deduced that persimmon enriched ice cream could be a nutritious dairy-based analog for people suffering from hypertension.

Conclusions

Over the last decade, the sector of functional frozen dairy desserts has experienced a remarkable growth, driven primarily by the increasing consumer awareness for healthy, nutritious, and additive-free products without compromising distinctive quality features such as palatability, flavor, and texture. Due to its structural complexity and the moderately low processing intensity, ice cream is generally considered as a good substrate for incorporating a broad spectrum of bioactive compounds, including living organisms. On the other hand, the direct relationship between structural elements and markers of quality and storage stability may pose obstacles on convenient functional ice cream prototyping. Consequently, incorporating functional ingredients and bioactive compounds must be realized without adversely affecting the colloidal and structural integrity of ice cream (sufficient air incorporation and fat droplet destabilization, minimal impact on crystallization and recrystallization phenomena, control of unfrozen water molecular mobility and diffusivity, and compatibility of the added bioactive compounds with major compositional elements), flavor, and taste attributes, ensuring adequate stability of the bioactive compounds. Probiotics can generally be easily incorporated in ice cream due to their relatively minor impact on ice cream texture and storage stability. On the other hand, incorporating prebiotics, DF, and LGSs in ice cream is a challenging task due to their ability to influence the colligative and rheological properties of the mixes, and consequently, the structure and texture development as well as the colloidal changes during storage.

Employing naturally occurring antioxidants such as polyphenols and carotenoids from by-products or in form of fruit preparations appears an effective strategy to promote ice cream quality and to drastically improve antioxidant (free radical scavenging, potentially reducing chronic health complications related to oxidative stress and inflammation) capacity of frozen dairy desserts. In addition, incorporating fruit preparations in ice cream can also ameliorate the mineral/trace element profile (for example, Ca, P, Mg, K, Na, Fe, and Zn) and provide an all-natural flavor and color.

Despite the conceptualization and development of several types of functional ice cream products, the disclosure of specific health benefits and physiological actions for marketing purposes still remains very challenging due to legislation restrictions. In most cases, food bioactives incorporated into ice cream should be judged for their health benefits by government agencies, for example, EFSA, FDA, and so on, and evidence of the health benefits should originate from clinical trials. From this perspective, developing a holistic approach in the field of functional ice cream that will also include findings from *in vivo* and clinical studies apart from technological profile characterization appears to be a growing future trend.

Author Contributions

Soukoulis has researched prior studies, interpreted results, drafted, and prepared manuscript. Fisk contributed to the manuscript; proofread and critically reviewed the manuscript. Bohn contributed with several additional ideas, wrote parts of the manuscript, and aided in the finalization of the manuscript.

References

- Abghari A, Sheikh-Zeinoddin M, Soleimanian-Zad S. 2011. Nonfermented ice cream as a carrier for *Lactobacillus acidophilus* and *Lactobacillus rhamnosus*. *Int J Food Sci Technol* 46:84–92.

- Ablett S, Clarke CJ, Izzard MJ, Martin DR. 2002. Relationship between ice recrystallisation rates and the glass transition in frozen sugar solutions. *J Sci Food Agric* 82:1855–9.
- Abril JR, Stull JW, Taylor RR, Angus RC, Daniel TC. 1982. Characteristics of frozen desserts sweetened with xylitol and fructose. *J Food Sci* 47:472–5.
- Agustin LS, Franceschi S, Jenkins JDA, Kendall CWC, La Vecchia C. 2002. Glycemic index in chronic disease: a review. *Eur J Clin Nutr* 56:1049–71.
- Ahmadi A, Milani E, Madadlou A, Mortazavi SA, Mokarram RR, Salarbashi D. 2012. Synbiotic yogurt-ice cream produced via incorporation of microencapsulated *Lactobacillus acidophilus* (la-5) and fructooligosaccharide. *J Food Sci Technol*. doi:10.1007/s13197-012-0679-y.
- Akalin AS, Erişir D. 2008. Effects of inulin and oligofructose on the rheological characteristics and probiotic culture survival in low-fat probiotic ice cream. *J Food Sci* 73:M184–8.
- Akalin AS, Karagözlü C, Ünal G. 2008. Rheological properties of reduced-fat and low-fat ice cream containing whey protein isolate and inulin. *European Food Res Technol* 227:889–95.
- Akin MB, Akin MS, Kirmaci Z. 2007. Effects of inulin and sugar levels on the viability of yogurt and probiotic bacteria and the physical and sensory characteristics in probiotic ice-cream. *Food Chem* 104:93–9.
- Akin MS. 2005. Effects of inulin and different sugar levels on viability of probiotic bacteria and the physical and sensory characteristics of probiotic fermented ice-cream. *Milchwissenschaft* 60:297–301.
- Al-Naqeb G, Maznah I, Gururaj B, Hadiza AA. 2010. Vanillin rich fraction regulates LDLR and HMGCR gene expression in HepG2 cells. *Food Res Intl* 43:2437–43.
- Alamprese C, Foschino R, Rossi M, Pompei C, Savani L. 2002. Survival of *Lactobacillus johnsonii* La1 and influence of its addition in retail-manufactured ice cream produced with different sugar and fat concentrations. *Intl Dairy J* 12:201–8.
- Alamprese C, Foschino R, Rossi M, Pompei C, Corti S. 2005. Effects of *Lactobacillus rhamnosus* GG addition in ice cream. *Intl J Dairy Technol* 58:200–6.
- Alvarez VB, Wolters CL, Vodovotz Y, Ji T. 2005. Physical properties of ice cream containing milk protein concentrates. *J Dairy Sci* 88:862–71.
- Alves LL, Richards NSPS, Mattanna P, Andrade DF, Rezer S, Milani LIG, Cruz AG, Faria JAF. 2013. Cream cheese as a symbiotic food carrier using *Bifidobacterium animalis* Bb-12 and *Lactobacillus acidophilus* La-5 and inulin. *Intl J Dairy Technol* 66:63–9.
- Anfinsen JR, Tungland BC. 2006. Reduced carbohydrate and nutritionally-enhanced frozen desserts and other food products. US20060286248 A1.
- Antunes AEC, Liserre AM, Coelho ALA, Menezes CR, Moreno I, Yotsuyanagi K, Azambuja NC. 2013. Acerola nectar with added microencapsulated probiotic. *LWT – Food Sci Technol* 54:125–31.
- Arab-Tehrani E, Jacquot M, Gaiani C, Imran M, Desobry S, Linder M. 2012. Beneficial effects and oxidative stability of omega-3 long-chain polyunsaturated fatty acids. *Trends Food Sci Technol* 25:24–33.
- Arihara K, Nakashima Y, Ishikawa S, Itoh M. 2004. Antihypertensive activities generated from porcine skeletal muscle proteins by lactic acid bacteria [abstract]. Abstracts of 50th International Congress of Meat Science and Technology, Aug 8–13 2004. Helsinki, Finland: Elsevier Ltd. p 236.
- Aryana KJ, Summers M. 2006. Probiotic, fat-free, no sugar-added ice cream. *Milchwissenschaft* 61:184–7.
- Augustin MA, Williams RPW. 2001. Nutritional mineral fortification of milk. WO2001072135 A1.
- Augustin MA. 2003. The role of microencapsulation in the development of functional dairy foods. *Aust J Dairy Technol* 58:156–60.
- Augustin MA, Htoon A, Sanguansri L. 2007. Starch treatment process. WO2005105851 A1.
- Ayala-Zavala JF, Vega-Vega V, Rosas-Domínguez C, Palafox-Carlos H, Villa-Rodríguez JA, Wasim-Siddiqui MD, Dávila-Aviña JE, González-Aguilar GA. 2011. Agro-industrial potential of exotic fruit byproducts as a source of food additives. *Food Res Intl* 44:1866–74.
- Aykan V, Sezgin E, Guzel-Seydim ZB. 2008. Use of fat replacers in the production of reduced-calorie vanilla ice cream. *Eur J Lipid Sci Technol* 110:516–20.
- Baer RJ, Baldwin KA. 1984. Freezing points of bulking agents used in manufacture of low-calorie frozen desserts. *J Dairy Sci* 67:2860–2.
- Bantle J. 2005. Is fructose the optimal low glycemic index sweetener? In: Bantle J, Slama G, editors. Nutritional management of diabetes mellitus and dysmetabolic syndrome. Basel, Switzerland: KargerAG, p 83–96.
- Başığit G, Kuleşan H, Karahan AG. 2006. Viability of human-derived probiotic lactobacilli in ice cream produced with sucrose and aspartame. *J Ind Microbiol Biotechnol* 33:796–800.
- Bayarri S, González-Tomás L, Hernando I, Lluch AM, Costell E. 2011. Texture perceived on inulin-enriched low-fat semisolid dairy desserts, rheological and structural basis. *J Texture Stud* 42:174–84.
- Begley M, Gahan CGM, Hill C. 2005. The interaction between bacteria and bile. *FEMS Microbiol Rev* 29:625–51.
- Ben-Avraham S, Harman-Boehm I, Schwarzfuchs D, Shai I. 2009. Dietary strategies for patients with type 2 diabetes in the era of multi-approaches; review and results from the Dietary Intervention Randomized Controlled Trial (DIRECT). *Diab Res Clin Pract* 86(Supplement 1):S41–8.
- Berry MJ, Quail PJ, Wilkinson JE. 2006. Ice cream confection. WO 2006042608 A1.
- Bigliardi B, Galati F. 2013. Innovation trends in the food industry: the case of functional foods. *Trends Food Sci Technol* 31:118–29.
- Bohn T. 2008. Bioavailability of non-provitamin A carotenoids. *Curr Nutr Food Sci* 4:240–58.
- Bolliger S, Wildmoser H, Goff HD, Tharp BW. 2000. Relationships between ice cream mix viscoelasticity and ice crystal growth in ice cream. *Intl Dairy J* 10:791–7.
- Bordi P, Cranage D, Stokols J, Palchak T, Powell L. 2004. Effect of polyols versus sugar on the acceptability of ice cream among a student and adult population. *Foodservice Res Int* 15:41–50.
- Boskou D. 2006. Sources of natural phenolic antioxidants. *Trends Food Sci Technol* 17:505–12.
- Bremer PJ, Ridley SC. 2004. Safety of frozen foods. In: Hui YH, Cornillon P, Guerrero Legaretta I, Lim MH, Murrel KD, Nip WP, editors. Handbook of frozen foods. New York: Marcell Dekker Inc.
- Brennan CS, Kuri V, Tudorica CM. 2004. Inulin-enriched pasta: effects on textural properties and starch degradation. *Food Chem* 86:189–93.
- Brownlee IA. 2011. The physiological roles of dietary fibre. *Food Hydrocolloids* 25:238–50.
- Burdock GA. 2005. Fenaroli's handbook of flavour ingredients. 5th ed. Boca Raton, Fla.: CRC Press.
- Burgain J, Gaiani C, Linder M, Scher J. 2011. Encapsulation of probiotic living cells: from laboratory scale to industrial applications. *J Food Eng* 104:467–83.
- Cadena RS, Cruz AG, Faria JAF, Bolini HMA. 2012. Reduced fat and sugar vanilla ice creams: sensory profiling and external preference mapping. *J Dairy Sci* 95:4842–50.
- Çam M, Erdoğan F, Aslan D, Dinç M. 2013. Enrichment of functional properties of ice cream with pomegranate by-products. *J Food Sci* 78:C1543–C50.
- Çam M, İçyer NC, Erdoğan F. 2014. Pomegranate peel phenolics: microencapsulation, storage stability and potential ingredient for functional food development. *LWT – Food Sci Technol* 55:117–23.
- Camire ME, Dougherty MP, Teh YH. 2006. Frozen wild blueberry-tofu-soymilk desserts. *J Food Sci* 71:S119–S23.
- Carlson TL, McElwee D, Remmert MA, Kim CHP, Walbert S, Woo A. 2011. Low glycemic sweeteners and products made using the same. EP 1729596 A2.
- Castro WF, Cruz AG, Bisinotto MS, Guerreiro LMR, Faria JAF, Bolini HMA, Cunha RL, Deliza R. 2013. Development of probiotic dairy beverages: rheological properties and application of mathematical models in sensory evaluation. *J Dairy Sci* 96:16–25.
- Champagne CP, Fustier P. 2007. Microencapsulation for the improved delivery of bioactive compounds into foods. *Curr Opin Biotechnol* 18:184–90.
- Chang Y, Hartel RW. 2002. Stability of air cells in ice cream during hardening and storage. *J Food Eng* 55:59–70.
- Charles DJ. 2013. Vanilla. In: Charles DJ, editor. Antioxidant properties of spices, herbs, and other sources. New York: Springer Science+Business Media. p 581–7.
- Chawla R, Patel GR. 2010. Soluble dietary fiber. *Compr Rev Food Sci Food Saf* 9:178–96.
- Chee CP, Djordjevic D, Faraji H, Decker EA, Hollender R, McClements DJ, Peterson DG, Roberts RF, Coupland JN. 2007. Sensory properties of vanilla and strawberry flavoured ice cream supplemented with omega-3 fatty acids. *Milchwissenschaft* 62:66–9.
- Chekri R, Noel L, Millour S, Vastel C, Kadar A, Sirot V, Leblanc JC, Guerin T. 2012. Calcium, magnesium, sodium and potassium levels in foodstuffs

- from the second French Total Diet Study. *J Food Compos Anal* 25: 97–107.
- Chen S, Bobe G, Zimmerman S, Hammond EG, Luhman CM, Boylston TD, Freeman AE, Beitz DC. 2004. Physical and sensory properties of dairy products from cows with various milk fatty acid compositions. *J Agric Food Chem* 52:3422–8.
- Chen W, Duizer L, Corredig M, Goff HD. 2010. Addition of soluble soybean polysaccharides to dairy products as a source of dietary fiber. *J Food Sci* 75:C478–C84.
- Christiansen PS, Edelsten D, Kristiansen JR, Nielsen EW. 1996. Some properties of ice cream containing *Bifidobacterium bifidum* and *Lactobacillus acidophilus*. *Milchwissenschaft* 51:502–4.
- Christiansen PS, Madeira AIMR, Edelsten D. 1999. The use of ropy milk as stabilizer in the manufacture of ice cream. *Milchwissenschaft* 54: 138–40.
- Chutkan R, Fahey G, Wright WL, McRorie J. 2012. Viscous versus nonviscous soluble fiber supplements: mechanisms and evidence for fiber-specific health benefits. *Am Acad Nurse Pract* 24:476–87.
- Codex. 2009. Report on the 30th session of the Codex Committee on Nutrition and Foods for Special Dietary Uses. ALINORM 09/32/26. (Appendix II). Rome.
- Cook KKL, Hartel RW. 2011. Effect of freezing temperature and warming rate on dendrite break-up when freezing ice cream mix. *Intl Dairy J* 21:447–53.
- Cook MT, Tzortzis G, Charalampopoulos D, Khutoryanskiy VV. 2012. Microencapsulation of probiotics for gastrointestinal delivery. *J Control Release* 162:56–67.
- Coussement P, Franck A. 2001. Inulin and oligofructose. In: Cho SS, Dreher ML, editors. *Handbook of dietary fiber*. New York: Marcel Dekker Inc. p 721–36.
- Cruz AG, Antunes AEC, Sousa ALOP, Faria JAF, Saad SMI. 2009. Ice-cream as a probiotic food carrier. *Food Res Intl* 42:1233–9.
- Cruz AG, Castro WF, Faria JAF, Lollo PCB, Maya-Farfán J, Freitas MQ, Rodrigues D, Oliveira CAF, Godoy HT. 2012. Probiotic yogurts manufactured with increased glucose oxidase levels: postacidification, proteolytic patterns, survival of probiotic microorganisms, production of organic acid and aroma compounds. *J Dairy Sci* 95:2261–9.
- Cruz AG, Cavalcanti RN, Guerreiro LMR, Sant'Ana AS, Nogueira LC, Oliveira CAF, Deliza R, Cunha RL, Faria JAF, Bolini HMA. 2013. Developing a prebiotic yogurt: rheological, physico-chemical and microbiological aspects and adequacy of survival analysis methodology. *J Food Eng* 114:323–30.
- Dağdemir E. 2011. Effect of vegetable marrow (*Cucurbita pepo* L.) on ice cream quality and nutritive value. *Asian J Chem* 23:4684–8.
- Davidson RH, Duncan SE, Hackney CR, Eigel WN, Boling JW. 2000. Probiotic culture survival and implications in fermented frozen yogurt characteristics. *J Dairy Sci* 83:666–73.
- de Moraes Crizel T, Jablonski A, de Oliveira Rios A, Rech R, Flôres SH. 2013. Dietary fiber from orange byproducts as a potential fat replacer. *LWT – Food Sci Technol* 53:9–14.
- de Wees Allen A. 2008. Novel sweetener compositions and methods of use. US 2008/0317901 A1
- de Wees Allen A. 2009. Low glycemic index confection. US 2009019695 A1
- Debon J, Prudêncio ES, Cunha Petrus JC. 2010. Rheological and physico-chemical characterization of prebiotic microfiltered fermented milk. *J Food Eng* 99:128–35.
- Deepika G, Rastall RA, Charalampopoulos D. 2011. Effect of food models and low-temperature storage on the adhesion of *Lactobacillus rhamnosus* GG to Caco-2 cells. *J Agric Food Chem* 59:8661–6.
- Dervisoglu M. 2006. Influence of hazelnut flour and skin addition on the physical, chemical and sensory properties of vanilla ice cream. *Intl J Food Sci Technol* 41:657–61.
- Dervisoglu M, Yazici F. 2006. Note. The effect of citrus fibre on the physical, chemical and sensory properties of ice cream. *Food Sci Technol Intl* 12:159–64.
- Devereux HM, Jones GP, McCormack L, Hunter WC. 2003. Consumer acceptability of low fat foods containing inulin and oligofructose. *J Food Sci* 68:1850–4.
- Di Criscio T, Fratianni A, Mignogna R, Cinquanta L, Coppola R, Sorrentino E, Panfili G. 2010. Production of functional probiotic, prebiotic, and synbiotic ice creams. *J Dairy Sci* 93:4555–64.
- dos Santos Leandro E, Andrade de Araujo E, Lopes da Conceição JP, Alencar de Moraes C, Fernandes de Carvalho A. 2013. Survival of *Lactobacillus delbrueckii* UFV H2b20 in ice cream produced with different fat levels and after submission to stress acid and bile salts. *J Funct Foods* 5: 503–7.
- Egger P, Holzer G, Segato S, Werth E, Schwenbacher F, Peratoner G, Andigetto I, Kasal A. 2007. Effects of oilseed supplements on milk production and quality in dairy cows fed a hay-based diet. *Ital J Anim Sci* 6:395–405.
- El-Nagar G, Clowes G, Tudoric CM, Kuri V, Brennan CS. 2002. Rheological quality and stability of yog-ice cream with added inulin. *Intl J Dairy Technol* 55:89–93.
- Elleuch M, Bedigian D, Roiseux O, Besbes S, Blecker C, Attia H. 2011. Dietary fibre and fibre-rich by-products of food processing: characterisation, technological functionality and commercial applications: a review. *Food Chem* 124:411–21.
- Erkaya T, Dağdemir E, Sengül M. 2012. Influence of Cape gooseberry (*Physalis peruviana* L.) addition on the chemical and sensory characteristics and mineral concentrations of ice cream. *Food Res Intl* 45:331–5.
- Esmerino EA, Cruz AG, Pereira EPR, Rodrigues JB, Faria JAF, Bolini HMA. 2013. The influence of sweeteners in probiotic Petit Suisse cheese in concentrations equivalent to that of sucrose. *J Dairy Sci* 96:5512–21.
- Euromonitor. 2012. Pre & Probiotics across the globe. Pre- and probiotics webinar. Available from: <http://richmedia.decisionnewsmedia.com/media/webinar/editorial/EditorialWebinar1/index.htm>
- Fagan CC, O'Donnell CP, Cullen PJ, Brennan CS. 2006. The effect of dietary fibre inclusion on milk coagulation kinetics. *J Food Eng* 77:261–8.
- FAO/WHO (2002). Guidelines for the evaluation of probiotics in food, Joint FAO/WHO working group report on drafting guidelines for the evaluation of probiotics in food. Available from: <http://ftp.fao.org/es/esn/food/wgreport2.pdf>
- Faus I, Sismiega H. 2004. Sweet tasting proteins. In: Fahnstock S, Steinbüchel A, editors. *Polyamides and complex proteinaceous materials II*. Weinheim: Wiley VHC. p 203–20.
- Favaro-Trindade CS, Bernardi S, Bodini RB, De Carvalho Balieiro JC, De Almeida E. 2006. Sensory acceptability and stability of probiotic microorganisms and vitamin C in fermented acerola (*Malpighia emarginata* DC.) ice cream. *J Food Sci* 71:S492–5.
- Favaro-Trindade CS, De Carvalho Balieiro JC, Dias PF, Sanino FA, Boschini C. 2007. Effects of culture, pH and fat concentration on melting rate and sensory characteristics of probiotic fermented yellow mombin (*Spondias mombin* L.) ice creams. *Food Sci Technol Intl* 13:285–91.
- Ferrar L, Van Der Hee RM, Berry M, Watson C, Miret S, Wilkinson J, Bradburn M, Eastell R. 2011. Effects of calcium-fortified ice cream on markers of bone health. *Osteoporosis Intl* 22:2721–31.
- Ferraz JL, Cruz AG, Cadena RS, Freitas MQ, Pinto UM, Carvalho CC, Faria JA, Bolini HM. 2012. Sensory acceptance and survival of probiotic bacteria in ice cream produced with different overrun levels. *J Food Sci* 77:S24–S8.
- Fisk I, White D, Lad M, Gray D. 2008. Oxidative stability of sunflower oil bodies. *Eur J Lipid Sci Technol* 110:962–8.
- Fogliano V, Vitaglione P. 2005. Functional foods: planning and development. *Mol Nutr Food Res* 49:256–62.
- Foster-Powell K, Holt SH, Brand-Miller JC. 2002. International table of glycemic index and glycemic load values: 2002. *Am J Clin Nutr* 76:5–56.
- Foster T, Wolf B. 2011. Hydrocolloid gums – their role and interactions in food. In: NIT, Spyropoulos F, Cox P, editors. *Practical food rheology an interpretive approach*. Oxford, UK: Wiley-Blackwell Ltd. p 61–79.
- Franck A. 2002. Technological functionality of inulin and oligofructose. *Brit J Nutr* 87:S287–S91.
- Fu N, Chen XD. 2011. Towards a maximal cell survival in convective thermal drying processes. *Food Res Intl* 44:1127–49.
- Fukuda M. 2012. Ice cream. EP 2491793 A1.
- Gassenmeier K. 2003. Vanillin and xanthine oxidase – key factors for the generation of a cardboard Off-note in vanilla ice cream. *LWT – Food Sci Technol* 36:99–103.
- Gaudreau H, Champagne CP, Remondetto GE, Bazinet L, Subirade M. 2013. Effect of catechins on the growth of oxygen-sensitive probiotic bacteria. *Food Res Intl* 53:751–7.
- Giri A, Ramachandra Rao HG, Ramesh V. 2013. Effect of incorporating whey protein concentrate into stevia-sweetened Kulfi on physicochemical and sensory properties. *Intl J Dairy Technol* 66:286–90.
- Godward G, Kailasapathy K. 2003. Viability and survival of free, encapsulated and co-encapsulated probiotic bacteria in ice cream. *Milchwissenschaft* 58:161–4.

- Goff DH, Jordan WK. 1984. Aspartame and polydextrose in a calorie-reduced frozen dairy dessert. *J Food Sci* 49:306–7.
- Goff HD. 1997. Instability and partial coalescence in whippable dairy emulsions. *J Dairy Sci* 80:2620–30.
- Goff HD. 2008. 65 Years of ice cream science. *Intl Dairy J* 18:754–8.
- Goh KKT, Nair RS, Matia-Merino L. 2008. Exploiting the functionality of lactic acid bacteria in ice cream. *Food Biophys* 3:295–304.
- Goh KKT, Ye A, Dale N. 2006. Characterisation of ice cream containing flaxseed oil. *Intl J Food Sci Technol* 41:946–53.
- Gomes AA, Braga SP, Cruz AG, Cadena RS, Lollo PCB, Carvalho C, Maya-Farfán J, Faria JAF, Bolini HMA. 2011. Effect of the inoculation level of *Lactobacillus acidophilus* in probiotic cheese on the physicochemical features and sensory performance compared with commercial cheeses. *J Dairy Sci* 94:4777–86.
- González-Tomás L, Bayarri S, Costell E. 2009. Inulin-enriched dairy desserts: physicochemical and sensory aspects. *J Dairy Sci* 92:4188–99.
- Gonzalez S, Duncan SE, O'Keefe SF, Sumner SS, Herbein JH. 2003. Oxidation and textural characteristics of butter and ice cream with modified fatty acid profiles. *J Dairy Sci* 86:70–7.
- Granato D, Branco GF, Cruz AG, Faria JDAF, Shah NP. 2010a. Probiotic dairy products as functional foods. *Compr Rev Food Sci Food Saf* 9:455–70.
- Granato D, Branco GF, Nazzaro F, Cruz AG, Faria JAF. 2010b. Functional foods and nondairy probiotic food development: trends, concepts, and products. *Compr Rev Food Sci Food Saf* 9:292–302.
- Guner A, Ardic M, Keles A, Dogruer Y. 2007. Production of yogurt ice cream at different acidity. *Intl J Food Sci Technol* 42:948–52.
- Hagen M, Narvhus JA. 1999. Production of ice cream containing probiotic bacteria. *Milchwissenschaft* 54:265–8.
- Hagiwara T, Hartel RW. 1996. Effect of sweetener, stabilizer, and storage temperature on ice recrystallization in ice cream. *J Dairy Sci* 79:735–44.
- Hasler C. 2002. Functional food: benefits, concerns and challenges. *J Nutr* 132:3772–81.
- Haynes IN, Playne MJ. 2002. Survival of probiotic cultures in low-fat ice-cream. *Aust J Dairy Technol* 57:10–4.
- Hekmat S, McMahon DJ. 1992. Survival of *Lactobacillus acidophilus* and *Bifidobacterium bifidum* in ice cream for use as a probiotic food. *J Dairy Sci* 75:1415–22.
- Herceg Z, Režek A, Lelas V, Krešić G, Franetović M. 2007. Effect of carbohydrates on the emulsifying, foaming and freezing properties of whey protein suspensions. *J Food Eng* 79:279–86.
- Hodge AM, O'Dea K, English DR, Giles GG. 2004. Glycemic index and dietary fiber and the risk of type 2 diabetes. *Diabetes Care* 27:2701–6.
- Holcomb JE, Frank J, McGregor JU. 1991. Viability of *Lactobacillus acidophilus* and *Bifidobacterium bifidum* in soft-serve frozen yogurt. *Cultured Dairy Prod J* 26:4–5.
- Homayouni A, Azizi A, Ehsani MR, Yarmand MS, Razavi SH. 2008a. Effect of microencapsulation and resistant starch on the probiotic survival and sensory properties of synbiotic ice cream. *Food Chem* 111:50–5.
- Homayouni A, Ehsani MR, Azizi A, Razavi SH, Yarmand MS. 2008b. Growth and survival of some probiotic strains in simulated ice cream conditions. *J Appl Sci* 8:379–82.
- Homayouni A, Azizi A, Javadi M, Mahdipour S, Ejtahed H. 2012. Factors influencing probiotic survival. *Int J Dairy Sci* 7:1–10.
- Huang C. 2011. Compositions and methods of omega polyunsaturated fatty acids for the treatment of oral diseases. *US 20120213713 A1*
- Hwang JY, Shyu YS, Hsu CK. 2009. Grape wine lees improves the rheological and adds antioxidant properties to ice cream. *LWT – Food Sci Technol* 42:312–8.
- Inoue K, Shiota K, Ito T. 1998. Preparation and properties of ice cream type frozen yogurt. *Intl J Dairy Technol* 51:44–56.
- Isik U, Boyacioglu D, Capanoglu E, Nilufer Erdil D. 2011. Frozen yogurt with added inulin and isomalt. *J Dairy Sci* 94:1647–56.
- Jacobson MR, Reddy S, Vadehra D, Wedral ER, Sher A. 2002. Calcium fortified foodstuff. *US6342257 B1*
- Jimenez M, Garcia HS, Beristain CI. 2008. Sensory evaluation of dairy products supplemented with microencapsulated conjugated linoleic acid (CLA). *LWT – Food Sci Technol* 41:1047–52.
- Johannsen L, Helmutraut A, Bedford G. 2007. Frozen confections. *EPI925210*.
- Jookan E, Amery R, Struyf T, Duquenne B, Geuns J, Meesschaert B. 2012. Stability of steviol glycosides in several food matrices. *J Agric Food Chem* 60:10606–12.
- Kailasapathy K, Sultana K. 2003. Survival and β -D-galactosidase activity of encapsulated and free *Lactobacillus acidophilus* and *Bifidobacterium lactis* in ice-cream. *Aust J Dairy Technol* 58:223–7.
- Kamat JP, Ghosh A, Devasagayam TPA. 2000. Vanillin as an antioxidant in rat liver mitochondria: inhibition of protein oxidation and lipid peroxidation induced by photosensitization. *Mol Cell Biol* 20:47–53.
- Karaca OB, Güven M, Yasar K, Kaya S, Kahyaoglu T. 2009. The functional, rheological and sensory characteristics of ice creams with various fat replacers. *Intl J Dairy Technol* 62:93–9.
- Karaman S, Kayacier A. 2012. Rheology of ice cream mix flavoured with black tea or herbal teas and effect of flavouring on the sensory properties of ice cream. *Food Bioprocess Technol* 5:3159–69.
- Karaman S, Tokar ÖS, Yüksel F, Çam M, Kayacier A, Dogan M. 2014. Physicochemical, bioactive and sensory properties of persimmon-based ice cream: technique for order preference by similarity to ideal solution to determine optimum concentration. *J Dairy Sci* 97:97–110.
- Kebary KMK. 1996. Viability of *Bifidobacterium bifidum* and its effect on quality of frozen Zabady. *Food Res Intl* 29:431–7.
- Khan MI, Arshad MS, Anjum FM, Sameen A, Aneeq UR, Gill WT. 2011. Meat as a functional food with special reference to probiotic sausages. *Food Res Intl* 44: 3125–33.
- Khan RS, Grigor J, Winger R, Win A. 2013. Functional food product development: opportunities and challenges for food manufacturers. *Trends Food Sci Technol* 30:27–37.
- Koba K, Yanagita T. 2013. Health benefits of conjugated linoleic acid (CLA). *Obes Res Clin Pract*. In press. Available from: <http://dx.doi.org/10.1016/j.orcp.2013.10.001>.
- Koeferli CRS, Piccinalli P, Sigrist S. 1996. The influence of fat, sugar and non-fat milk solids on selected taste, flavour and texture parameters of a vanilla ice-cream. *Food Qual Prefer* 7:69–79.
- Kroyer G. 2010. Stevioside and Stevia-sweetener in food: application, stability and interaction with food ingredients. *J Consum Prot Food Saf* 5:1–5.
- Laroia S, Martin JH. 1991. Effect of pH on survival of *Bifidobacterium bifidum* and *Lactobacillus acidophilus* in frozen fermented desserts. *Cultured Dairy Prod J* 11:13–21.
- Lemus-Mondaca R, Vega-Gálvez A, Zura-Bravo L, Ah-Hen K. 2012. Stevia rebaudiana Bertoni, source of a high-potency natural sweetener: a comprehensive review on the biochemical, nutritional and functional aspects. *Food Chem* 132:1121–32.
- Li G. 2012. Intestinal probiotics: interaction with bile salts and reduction of cholesterol. *Procedia Environ Sci* 12:1180–6.
- Liou BK, Grün IU. 2007. Effect of fat level on the perception of five flavor chemicals in ice cream with or without fat mimetics by using a descriptive test. *J Food Sci* 72:S595–604.
- Maganha LC, Rosim RE, Corassin CH, Cruz AG, Faria JAF, Oliveira CAF. 2014. Viability of probiotic bacteria in fermented skim milk produced with different levels of milk powder and sugar. *Intl J Dairy Technol* 67:89–94.
- Magariños H, Selaive S, Costa M, Flores M, Pizarro O. 2007. Viability of probiotic micro-organisms (*Lactobacillus acidophilus* La-5 and *Bifidobacterium animalis* subsp. *lactis* Bb-12) in ice cream. *Intl J Dairy Technol* 60:128–34.
- Maia MCA, Galvão APGLK, Modesta RCD, Pereira Jr N. 2008. Consumer evaluation of ice cream with xylitol. *Avaliação do consumidor sobre sorvetes com xilitol* 28:341–7.
- Makni M, Chtourou Y, Fetoui H, Garoui EM, Boudawara T, Zeghal N. 2011. Evaluation of the antioxidant, anti-inflammatory, and hemato-protective properties of vanillin in carbon tetrachloride-treated rats. *Eur J Pharmacol* 668:133–9.
- Malmo C, Stora A, Mauriello G. 2013. Microencapsulation of *Lactobacillus reuteri* DSM 17938 cells coated in alginate beads with chitosan by spray drying to use as a probiotic cell in a chocolate soufflé. *Food Bioprocess Technol* 6:795–805.
- Mantzouridou F, Spanou A, Kiosseoglou V. 2012. An inulin-based dressing emulsion as a potential probiotic food carrier. *Food Res Intl* 46:260–9.
- Marshall RT, Goff HD, Hartel RW. 2003. Ice cream. 3rd ed. New York: Aspen Publishers.
- Martin CL, Murphy SP, Au DLM. 2008. Compiling glycemic index and glycemic load values for addition to a food composition database. *J Food Compos Anal* 21:469–73.

- Melanson KJ, Angelopoulos TJ, Nguyen V, Zukley L, Lowndes J, Rippe JM. 2008. High-fructose corn syrup, energy intake, and appetite regulation. *Am J Clinical Nutr* 88:17385–44S.
- Méndez-Velasco C, Goff HD. 2012. Fat structure in ice cream: a study on the types of fat interactions. *Food Hydrocolloids* 29:152–9.
- Meyer D, Bayarri S, Tárrega A, Costell E. 2011a. Inulin as texture modifier in dairy products. *Food Hydrocolloids* 25:1881–90.
- Meyer D, Vermulst J, Tromp RH, De Hoog EHA. 2011b. The effect of inulin on tribology and sensory profiles of skimmed milk. *J Texture Stud* 42:387–93.
- Miller-Livney T, Hartel RW. 1997. Ice recrystallization in ice cream: interactions between sweeteners and stabilizers. *J Dairy Sci* 80:447–56.
- Mills S, Stanton C, Fitzgerald GF, Ross RP. 2011. Enhancing the stress responses of probiotics for a lifestyle from gut to product and back again. *Microb Cell Fact* 10:S19.
- Mohammadi R, Mortazavian AM, Khosrokhavar R, Da Cruz AG. 2011. Probiotic ice cream: viability of probiotic bacteria and sensory properties. *Ann Microbiol* 61:411–24.
- Mudgil D, Barak S. 2013. Composition, properties and health benefits of indigestible carbohydrate polymers as dietary fibre: a review. *Intl J Biol Macromol* 61:1–6.
- Muse MR, Hartel RW. 2004. Ice cream structural elements that affect melting rate and hardness. *J Dairy Sci* 87:1–10.
- Noël L, Chekri R, Millour S, Vastel C, Kadar A, Sirot V, Leblanc JC, Guérin T. 2012. Li, Cr, Mn, Co, Ni, Cu, Zn, Se and Mo levels in foodstuffs from the Second French TDS. *Food Chem* 132:1502–13.
- Nousia FG, Androutsakis PI, Fletouris DJ. 2011. Survival of *Lactobacillus acidophilus* LMGP-21381 in probiotic ice cream and its influence on sensory acceptability. *Int J Dairy Technol* 64:130–6.
- O'Brien-Nabors L. 2001. Alternative sweeteners. 3rd ed. New York: Marcel Dekker Inc.
- Ogunlade I, Tucker G, Fisk ID, Ogunlade A. 2009. Evaluation of antioxidant activity and vitamin E profile of some selected indigenous vegetables in Nigerian Diet. *J Food Agric Environ* 7:143–5.
- Oreopoulou V, Tzia C. 2006. Utilization of plant by-products for the recovery of proteins, dietary fibers, antioxidants and colorants. In: Oreopoulou V, Russ W, editors. Utilization of by-products and treatment of waste in the food industry. New York, USA: Springer Science & Business Media Inc. p 209–32.
- Özdemir C, Dagdemir E, Özdemir S, Sagdic O. 2008. The effects of using alternative sweeteners to sucrose on ice cream quality. *J Food Qual* 31:415–28.
- Özdemir C, Dagdemir E, Celik S, Özdemir S. 2003. An alternative ice cream production for diabetic patients. *Milchwissenschaft* 58:164–6.
- Ozturk I, Golec A, Karaman S, Sagdic O, Kayacier A. 2010. Evaluation of *Listeria monocytogenes* survival in ice cream mixes flavored with herbal tea using Taguchi method. *Foodborne Pathogen Dis* 7:1263–7.
- Pandiyar C, Annal Villi R, Kumaresan G, Murugan B, Gopalakrishnamurthy TR. 2012a. Development of synbiotic ice cream incorporating *Lactobacillus acidophilus* and *Saccharomyces boulardii*. *Intl Food Res J* 19:1233–9.
- Pandiyar C, Annal Villi R, Kumaresan G, Murugan B, Rajarajan G. 2012b. Effect of incorporation of inulin on the survivability of *Lactobacillus acidophilus* in synbiotic ice cream. *Intl Food Res J* 19:1729–32.
- Pandiyar C, Villy A, Kumaresan G, Murugan B, Gopalakrishnamurthy TR. 2012c. In vivo and in vitro effect of *Lactobacillus acidophilus* in synbiotic ice cream enriched with whey protein concentrate. *Intl Food Res J* 19:441–6.
- Patmore JV, Goff HD, Fernandes S. 2003. Cryo-gelation of galactomannans in ice cream model systems. *Food Hydrocolloids* 17:161–9.
- Pawar RS, Krynsky AJ, Rader JI. 2013. Sweeteners from plants – with emphasis on *Stevia rebaudiana* (Bertoni) and *Siraitia grosvenorii* (Swingle). *Anal Bioanal Chem* 405:4397–407.
- Pehkonen KS, Roos YH, Miao S, Ross RP, Stanton C. 2008. State transitions and physicochemical aspects of cryoprotection and stabilization in freeze-drying of *Lactobacillus rhamnosus* GG (LGG). *J Appl Microbiol* 104:1732–43.
- Pellegrini N, Serafini M, Salvatore S, Del Rio D, Bianchi M, Brighenti F. 2008. Total antioxidant capacity of spices, dried fruits, nuts, pulses, cereals and sweets consumed in Italy assessed by three different in vitro assays. *Mol Nutr Food Res* 50:1030–8.
- Pimentel TC, Cruz AG, Prudencio SH. 2013. Short communication: influence of long-chain inulin and *Lactobacillus paracasei* subspecies *paracasei* on the sensory profile and acceptance of a traditional yogurt. *J Dairy Sci* 96:6233–41.
- Popovich DG, Kendall CWC, Ransom TPP, Wolever TMS, Tariq N, Rosenberg RS, Jenkins AL, Jenkins DJA. 1996. Glycemic index of commonly consumed foods. *FASEB J* 10(3):A788.
- Prindiville EA, Marshall RT, Heymann H. 1999. Effect of milk fat on the sensory properties of chocolate ice cream. *J Dairy Sci* 82:1425–32.
- Puangmanee S, Shigeru H, Yuanxia S, Masahiro O. 2008. Application of whey protein isolate glycosylated with rare sugars to ice cream. *Food Sci Technol Res* 14:457–66.
- Pyenson H, Tracy PH. 1950. Vanillas as antioxidants in powdered ice cream mixes. *J Dairy Sci* 33:815–9.
- Ranadheera CS, Evans CA, Adams MC, Baines SK. 2012. In vitro analysis of gastrointestinal tolerance and intestinal cell adhesion of probiotics in goat's milk ice cream and yogurt. *Food Res Intl* 49:619–25.
- Ravula RR, Shah NP. 1998. Effect of acid casein hydrolysate and cysteine on the viability of yogurt and probiotic bacteria in fermented frozen dairy desserts. *Aust J Dairy Technol* 53:175–9.
- Regand A, Goff HD. 2003. Structure and ice recrystallization in frozen stabilized ice cream model systems. *Food Hydrocolloids* 17:95–102.
- Roberfroid MB. 2000. Concepts and strategy of functional food science: the European perspective. *Am J Clin Nutr* 71:1660–64.
- Roberfroid M. 2007. Prebiotics: the concept revisited. *J Nutr* 137:830–7.
- Roland AM, Phillips LG, Boor KJ. 1999. Effects of fat replacers on the sensory properties, color, melting, and hardness of ice cream. *J Dairy Sci* 82:2094–100.
- Ryu KS, Kim HB, Lee HS, Lee YK, Lee YW, Lim SH, Choung WY. 2002. Method for preparation of mulberry leaf powder and ice cream containing thereof. US6,376,002.
- Saad N, Delattre C, Urdaci M, Schmitter JM, Bressollier P. 2013. An overview of the last advances in probiotic and prebiotic field. *LWT – Food Sci Technol* 50:1–16.
- Saarela M, Virkajärvi I, Alakomi HL, Sigvart-Mattila P, Mättö J. 2006a. Stability and functionality of freeze-dried probiotic *Bifidobacterium* cells during storage in juice and milk. *Intl Dairy J* 16:1477–82.
- Saarela M, Virkajärvi I, Nohynek L, Vaari A, Mättö J. 2006b. Fibres as carriers for *Lactobacillus rhamnosus* during freeze-drying and storage in apple juice and chocolate-coated breakfast cereals. *Intl J Food Microbiol* 112:171–8.
- Sagdic O, Ozturk I, Cankurt H, Tornuk F. 2012. Interaction between some phenolic compounds and probiotic bacterium in functional ice Cream production. *Food Bioprocess Technol* 5:2964–71.
- Salem MME, Fathi FA, Awad RA. 2006. Production of functional ice cream. *Deutsche Lebensmittel-Rundschau* 102:326–9.
- Santivarangkna C, Kulosik U, Foerst P. 2008. Inactivation mechanisms of lactic acid starter cultures preserved by drying processes. *J Appl Microbiol* 105:1–13.
- Schaller-Povolny LA, Smith DE. 1999. Sensory attributes and storage life of reduced fat ice cream as related to inulin content. *J Food Sci* 64:555–9.
- Schaller-Povolny LA, Smith DE. 2001. Viscosity and freezing point of a reduced fat ice cream as related to inulin content. *Milchwissenschaft* 56:25–9.
- Schaller-Povolny LA, Smith DE. 2002. Interaction of milk proteins with inulin. *Milchwissenschaft* 57:494–7.
- Segall KI, Goff HD. 2002. A modified ice cream processing routine that promotes fat destabilization in the absence of added emulsifier. *Intl Dairy J* 12:1013–8.
- Senaka Ranadheera C, Evans CA, Adams MC, Baines SK. 2013. Production of probiotic ice cream from goat's milk and effect of packaging materials on product quality. *Small Ruminant Res* 112:174–80.
- Shah NP, Ravula RR. 2000. Microencapsulation of probiotic bacteria and their survival in frozen fermented dairy desserts. *Australian J Dairy Technol* 55:139–44.
- Sharma A, Kumar R, Azad ZRA, Adsule PG. 2013. Use of fine wine lees for value addition in ice cream. *J Food Sci Technol*. doi: 10.1007/s13197-013-1007-x
- Sheu TY, Marshall RT, Heymann H. 1993. Improving survival of culture bacteria in frozen desserts by microencapsulation. *J Dairy Sci* 76:1902–7.
- Shiota M, Ikeda N, Konishi H, Yoshioka T. 2002. Photooxidative stability of ice cream prepared from milk fat. *J Food Sci* 67:1200–7.

- Siaterlis A, Deepika G, Charalampopoulos D. 2009. Effect of culture medium and cryoprotectants on the growth and survival of probiotic lactobacilli during freeze drying. *Lett Appl Microbiol* 48:295–301.
- Silva Junior E, Lannes SCDS. 2011. Effect of different sweetener blends and fat types on ice cream properties. *Ciencia Tecnol Alimentaria* 31:217–20.
- Simopoulos AP. 1991. Omega-3 fatty acids in health and disease and in growth and development. *Am J Clin Nutr* 54:438–63.
- Song D, Khouryieh H, Abughazaleh AA, Salem MME, Hassan O, Ibrahim SA. 2011. Sensory properties and viability of probiotic microorganisms in chocolate ice cream supplemented with omega-3 fatty acids. *Milchwissenschaft* 66:172–5.
- Soukoulis C, Chandrinis I, Tzia C. 2008. Study of the functionality of selected hydrocolloids and their blends with κ -carrageenan on storage quality of vanilla ice cream. *LWT – Food Sci Technol* 41:1816–27.
- Soukoulis C, Lebesi D, Tzia C. 2009. Enrichment of ice cream with dietary fibre: effects on rheological properties, ice crystallisation and glass transition phenomena. *Food Chem* 115:665–71.
- Soukoulis C, Lyroni E, Tzia C. 2010a. Sensory profiling and hedonic judgement of probiotic ice cream as a function of hydrocolloids, yogurt and milk fat content. *LWT-Food Sci Technol* 43:1351–8.
- Soukoulis C, Rontogianni E, Tzia C. 2010b. Contribution of thermal, rheological and physical measurements to the determination of sensorially perceived quality of ice cream containing bulk sweeteners. *J Food Eng* 100:634–41.
- Soukoulis C, Fisk I. in press. Innovative ingredients and emerging technologies for controlling ice recrystallisation, texture and structure stability in frozen dairy desserts: a review. *Crit Rev Fd Sci Nutr*. doi: 10.1080/10408398.2013.876385.
- Soukoulis C, Tzia C. 2008. Impact of the acidification process, hydrocolloids and protein fortifiers on the physical and sensory properties of frozen yogurt. *Intl J Dairy Technol* 61:170–7.
- Soukoulis C, Tzia C. 2010. Response surface mapping of the sensory characteristics and acceptability of chocolate ice cream containing alternate sweetening agents. *J Sens Stud* 25:50–75.
- Soukoulis C, Yonekura L, Gan HH, Behboudi-Jobbehdar S, Parmenter C, Fisk I. 2014. Probiotic edible films as a new strategy for developing functional bakery products: the case of pan bread. *Food Hydrocolloids* 39:231–42.
- Srinivasan K, Platel K, Rao MVL. 2008. Hypotriglyceridemic effect of dietary vanillin in experimental rats. *European Food Res Technol* 228:103–8.
- Stokes JR, Boehm MW, Baier SK. 2013. Oral processing, texture, and mouthfeel: from rheology to tribology and beyond. *Curr Opin Colloid Surf Sci* 18:349–59.
- Sun-Waterhouse D, Edmonds L, Wadhwa SS, Wibisono R. 2013. Producing ice cream using a substantial amount of juice from kiwifruit with green, gold or red flesh. *Food Res Intl* 50:647–56.
- Sung KK, Goff HD. 2010. Effect of solid fat content on structure in ice creams containing palm kernel oil and high-oleic sunflower oil. *J Food Sci* 75:C274–C9.
- Tai A, Sawano T, Yazama F, Ito H. 2011. Evaluation of antioxidant activity of vanillin by using multiple antioxidant assays. *Biochim Biophys Acta* 1810:170–7.
- Talwalkar A, Kailasapathy K. 2004. A review of oxygen toxicity in probiotic yogurts: influence on the survival of probiotic bacteria and protective techniques. *Compr Rev Food Sci Food Saf* 3:117–24.
- Tamime AY, Robinson RK. 2007. *Tamime and Robinson's Yogurt Science and Technology*. 3rd ed. Cambridge, England: Woodhead Publishing Ltd. and CRC Press LCC
- Tárrega A, Torres JD, Costell E. 2011. Influence of the chain-length distribution of inulin on the rheology and microstructure of prebiotic dairy desserts. *J Food Eng* 104:356–63.
- Teh YH, Dougherty MP, Camire ME. 2005. Frozen blueberry-soy dessert quality. *J Food Sci* 70:S119–S22.
- Topping DL, Clifton PM. 2001. Short-chain fatty acids and human colonic function: roles of resistant starch and nonstarch polysaccharides. *Physiol Rev* 81:1031–64.
- Turgut T, Cakmakci S. 2009. Investigation of the possible use of probiotics in ice cream manufacture. *Int J Dairy Technol* 62:444–51.
- van der Hee RM, Miret S, Slettenaar M, Duchateau GSMJE, Rietveld AG, Wilkinson JE, Quail PJ, Berry MJ, Dainty JR, Teucher B, Fairweather-Tait SJ. 2009. Calcium absorption from fortified ice cream formulations compared with calcium absorption from milk. *J Am Diet Assoc* 109: 830–5.
- Varela P, Pintor A, Fiszman S. 2014. How hydrocolloids affect the temporal oral perception of ice cream. *Food Hydrocolloids* 36:220–8.
- Vinolo MAR, Rodrigues HG, Nachbar RT, Curi R. 2011. Regulation of inflammation by short chain fatty acids. *Nutrients* 3:858–76.
- Volkert M, Ananta E, Luscher CM, Knorr D. 2008. Effect of air freezing, spray freezing and pressure shift freezing on membrane integrity and viability of *Lactobacillus rhamnosus* GG. *J Food Eng* 87:532–40.
- Volkert M, Puaud M, Wille H-J, Knorr D. 2012. Effects of high pressure–low temperature treatment on freezing behavior, sensorial properties and air cell distribution in sugar rich dairy based frozen food foam and emulsions. *Innov Food Sci Emerg Technol* 13:75–85.
- Vuksan V, Jenkins AL, Rogovik AL, Fairgrieve CD, Jovanovski E, Leiter LA. 2011. Viscosity rather than quantity of dietary fibre predicts cholesterol-lowering effect in healthy individuals. *Brit J Nutr* 106:1349–52.
- Walstra P, Wouters JTM, Geurts TJ. 2006. Dairy science and technology. In: Walstra P, Wouters JTM, Geurts TJ, editors. 2nd ed. Boca Raton, Fla.: CRC Press Inc.
- Wendel HJ. 2010. Ice cream, useful for exclusive nutrition, comprises carbohydrate, protein and fat, which is free of trace elements, mineral substances, vitamins and optionally fibers and/or proteins. *DE 102008030964 A1*
- Westenbrink S, Brunt K, van der Kamp J-W. 2013. Dietary fibre: challenges in production and use of food composition data. *Food Chem* 562–7.
- Westerbeek JMM. 1995. Fermented ice cream products. A healthy challenge to the ice cream industry. *Scand Dairy Inf* 4:29–31.
- Whelan AP, Regand A, Vega C, Kerry JP, Goff HD. 2008a. Effect of trehalose on the glass transition and ice crystal growth in ice cream. *Intl J Food Sci Technol* 43:510–6.
- Whelan AP, Vega C, Kerry JP, Goff HD. 2008b. Physicochemical and sensory optimisation of a low glycemic index ice cream formulation. *Intl J Food Sci Technol* 43:1520–7.
- Wilcox JK, Ash SL, Catignani GL. 2004. Antioxidants and prevention of chronic disease. *Crit Rev Food Sci Nutr* 44:275–95.
- Zulim Botega DC, Marangoni A, Smith AK, Goff HD. 2013a. The potential application of rice bran wax oleogel to replace solid fat and enhance unsaturated fat content in ice cream. *J Food Sci* 78:C1334–C9.
- Zulim Botega DC, Marangoni A, Smith AK, Goff HD. 2013b. Development of formulations and processes to incorporate wax oleogels in ice cream. *J Food Sci* 78:C1845–51.