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Title: USE OF VERY LONG CHAIN INULIN FOR INHIBITING THE GROWTH OF ICE CRYSTALS IN A FROZEN FOODSTUFF

Abstract: The present invention is directed to the use of a very long chain inulin with a weight average degree of polymerization of between 50 and 103 for inhibiting the growth of ice crystals in a frozen foodstuff, preferably a frozen confection. The invention also relates to a frozen foodstuff, preferably a frozen confection, with inhibited ice crystal growth and to a method for producing a frozen confection with inhibited ice crystal growth.
Use of a very long chain inulin for inhibiting the growth of ice crystals in a frozen foodstuff

The present invention is directed to the use of a very long chain inulin for inhibiting the growth of ice crystals in a frozen foodstuff. The invention also relates to a method for inhibiting ice crystal growth in a frozen foodstuff and a frozen foodstuff with inhibited ice crystal growth.

Ice crystal formation is a known problem in frozen foodstuffs. When there is a gap between the time the frozen product is taken from the deep frozen section and placed in a domestic freezer, a partial thawing of the frozen product occurs before it is refrozen. Such cycles of temperature variation, which is known as "heat shock" cause the formation and growth of ice crystals in the product. The formation and growth of ice crystals in the product cause an undesirable crystallized texture in the frozen foodstuff. The crystallized texture and the icy mouth feel, as well as the impaired appearance, reduces the overall quality of the frozen foodstuff as perceived by the consumer.

Frozen confections are particularly appreciated for their creamy and smooth characteristics. In order to preserve the optimum organoleptic characteristics of smoothness, however, the frozen confections need to be stored and handled with care as even small temperature variations can be observed during storage, distribution, or handling. This is particularly the case when a consumer purchases a frozen confectionery and does not consume it quickly.

Various gums and emulsifiers have been used as additives to the frozen confection with the aim of improving the stability, smoothness, and the heat shock resistance of aerated frozen confections. For example, U.S. Pat. No. 4,500,553 to Liggett discloses that a gum stabilizer in combination with a monosaccharide and disaccharide sweeteners contributes to limiting the growth of ice crystals upon freezing and storage of frozen confection. In addition, U.S. Pat. No. 4, 542,035 to Huang discloses that adding to the product an edible multivalent cationic electrolyte such as calcium, magnesium, aluminum, iron, manganese and the like improve stability of the frozen product.

US patent application publication US20050123666A1 discloses an aerated frozen confection having reduced ice crystal growth after heat shock, and being prepared from a composition of
ingredients suitable for forming the confection and at least one emulsifier for reducing ice crystal growth after heat shock. The emulsifier is a polyol ester of a fatty acid which facilitates formation and stabilization of fat alpha crystals, the confection having an overrun of about 20% to about 250%, uniformly distributed small air cells having an average size of less than about 50 microns, ice crystals having a size of 30 microns or less after heat shock, and a smooth texture.

The problem addressed by the invention was to find further means for inhibiting the growth of ice crystals in a frozen foodstuff, particularly in a frozen confection.

In WO2006/108697 a very long chain inulin with a weight average degree of polymerization DPw of between 54 and 61 is described as a food additive that imparts beneficial usage properties to several foodstuffs as for example viscosity building and prebiotic effect. Ice cream is mentioned as an exemplary foodstuff containing said inulin. US20050123666A1 teaches that inulin can be added to the frozen confection as a freezing point depressant.

Surprisingly it was found out by the present inventors that inulin with a weight average degree of polymerization of between 50 and 103 is suitable for inhibiting the growth of ice crystals in a frozen foodstuff. Thus, the present invention relates to the use of an inulin with a weight average degree of polymerization of between 50 and 103 for inhibiting the growth of ice crystals in a frozen foodstuff.

Further objects of the present invention are a method for inhibiting the growth of ice crystals in a frozen foodstuff, a frozen foodstuff with inhibited ice crystal growth comprising an inulin with a weight average degree of polymerization of between 50 and 103, and a method for producing a frozen confection with inhibited ice crystal growth. These objects are defined in the appended claims and explained in more detail in the following description of the invention.

A further benefit of the inventive solutions is that the taste of the frozen product has a smooth and favorable texture, increased viscosity and body. Besides the inhibition of growth of ice crystals in a frozen foodstuff, very long chain inulin can at the same time act as a fat substitute in low caloric foodstuffs and as a viscosity building agent in certain foodstuffs, as for example ice cream yogurt, other dairy products and sauces.
For sake of brevity, inulin with a weight average degree of polymerization of between 50 and 103 and any other inulins which are described in the present description are hereinafter also referred to as "Very long chain inulin" or "VLCI". The term "inulin" is intended to mean in connection with the present invention a polyfructan which consists of a beta-2-1-linked chain of fructose molecules. This chain preferably, but not necessarily, has at its end a reducing alpha-D-glucose unit.

The terms weight average degree of polymerization (DPw) and number average degree of polymerization (DPn) used in the present disclosure are defined as in application WO 2006/108697 and measured by GPC-RI-MALLS, the method also disclosed in WO 2006/108697. In this connection and in connection with the present invention, the term "between" is also intended to include the respectively indicated numerical limits.

As far as DPw is concerned, very long chain inulin as used in the present invention has in a preferred embodiment an average degree of polymerization DPw of between 54 and 61, preferably between 55 and 60, particularly preferably between 56 and 57. In a another preferred embodiment, very long chain inulin which is used in the present invention has an average degree of polymerization DPw of between 65 and 81, preferably between 65 and 79, even more preferably between 66 and 78, very particularly even more preferably between 66 and 76, yet more preferably between 66 and 74 and most preferably between 66 and 73. In a still another preferred embodiment, very long chain inulin which is used in the present invention has an average degree of polymerization DPw of between 83 and 103, preferably between 84 and 100, more preferably between 83 and 98, even more preferably between 85 and 98, yet more preferably 85 and 95, still more preferably between 86 and 97 and most preferably between 86 and 94.

As far as DPn is concerned, very long chain inulin as used in the present invention has in a preferred embodiment an average degree of polymerization of between 44 and 48, preferably between 45 and 48, particularly preferably between 46 and 48. In a another preferred embodiment, very long chain inulin which is used in the present invention has an average degree of polymerization DPn of between 54 and 75, preferably between 54 and 72, even more preferably between 57 and 71, particularly preferably between 60 and 71. In a still another preferred embodiment, very long chain inulin which is used in the present invention has an average degree of polymerization DPn of between 66 and 89, preferably between 68
and 85, particularly preferably between 70 and 85 and even more preferably between 72 and 84.

Especially preferred according to the present invention are very long chain inulins with following characteristics:

5  (1) DPw of between 50-103 and DPn of between 41-89

(2) DPw of between 55-100 and DPn of between 45-85

(3) DPw of between 60-95 and DPn of between 50-81

(4) DPw of between 54-61 and DPn of between 44-48

(5) DPw of between 55-60 and DPn of between 45-48

10 (6) DPw of between 56-57 and DPn of between 46-48

(7) DPw of between 65-81 and DPn of between 54-75

(8) DPw of between 65-79 and DPn of between 54-72

(9) DPw of between 66-78 and DPn of between 57-71

(10) DPw of between 66-76 and DPn of between 60-71

15 (11) DPw of between 83-103 and DPn of between 66-89

(12) DPw of between 84-100 and DPn of between 68-85

(13) DPw of between 83-98 and DPn of between 70-85

(14) DPw of between 85-98 and DPn of between 72-84

(15) DPw of between 85-95 and DPn of between 72-81

20 An embodiment of very long chain inulin for use in the present invention which is particularly advantageous in its properties and which may include the characteristics of the previously described very long chain inulins has a particularly narrow molecular weight distribution expressed by the quotient between the weight average degree of polymerization and the
number average degree of polymerization DPw/DPn. This quantity is also referred to as polydispersity index. In a preferred embodiment, the quotient DPw/DPn is less than 1.25, in a more preferred embodiment is less than 1.20, in an even more preferred embodiment is less than 1.15 and in the most preferred embodiment is less than 1.10. The values for DPw and DPn are in this connection measured by the method of "gel permeation chromatography with light scattering and refractive index detection (GPC-RI-MALLS system)" described in WO 2006/108697. The molecular weight of a monomer for conversion calculations is set equal to 162 g/mol.

In a beneficial embodiment very long chain inulin which is used to the present invention has a molecular weight distribution of from 1620 g/mol to 40 000 g/mol, preferably from 2268 g/mol to 32 000 g/mol, particularly preferably from 2592 g/mol to 29 160 g/mol. In another preferred embodiment, very long chain inulin which is used in the present invention has a molecular weight distribution in the range from 650 to 48 000, more preferably 970 to 40 000 g/mol, even more preferably 1300 g/mol to 34 000 g/mol and most preferably from 4000 g/mol to 26 800 g/mol. In still another preferred embodiment, very long chain inulin for use in the present invention has a molecular weight distribution in the range from 650 to 48 000, more preferably 970 to 40 000 g/mol, even more preferably 1300 g/mol to 34 000 g/mol and most preferably from 4000 g/mol to 26 800 g/mol.

In one embodiment of the very long chain inulin the degree of branching is 0.5–2.0 mol%, more preferably 0.7–2.0 mol%, even more preferably 0.9 to 2.0 mol% and most preferably 1.1 to 2.0 mol%. The degree of branching is defined herein as the percentage number of beta-2-1-linked fructose monomers with additional branch point at position 6 of the fructose monomer (also abbreviated to "2-1,6-" hereinafter) based on the total number of all inulin monomers measured in a sample of the very long chain inulin with randomly distributed molecular weights. At its position 6, a "2-1,6-" fructose monomer within a polyfructose chain is linked to another polyfructose chain, consisting of at least two beta-2-1-linked fructose monomers, or to a single fructose monomer. The term “branch point” designates a position of a fructose monomer, within a polyfructose chain, to which another polyfructose chain consisting of at least two beta-2-1-linked fructose monomers, or a single fructose monomer is linked. The degree of branching is measured by the method of standard methylation analysis or alternatively by the method of reductive degradation after methylation. Both methods are described in detail in copending applications PCT/EP2007/004028, PCT/EP2007/004029, US 60/796,818, US 60/796,819.
Especially preferred in connection with the present invention are very long chain inulins as disclosed in WO 2006/108697, WO/2007/128559, WO/2007/128560, US 60/796,818 and US 60/796,819, which are incorporated by reference in their entirety.

The afore described very long chain inulins of the present invention can be prepared from artichoke (Cynara scolymus) roots as disclosed in WO 2006/108697, WO/2007/128559, WO/2007/128560, US 60/796,818 and US 60/796,819, which are incorporated by reference herein in their entirety.

A process for preparing very long chain inulin from artichoke roots is characterized by following steps:

a) artichoke (Cynara scolymus) roots are comminuted
b) an extract is obtained by treating the comminuted roots with water,
c) coloring constituents are removed from the extract obtained,
d) inulin is precipitated from the extract,
e) the inulin is reprecipitated at least once.

Artichoke roots are used as starting material, but the process is not restricted to a particular variety. The comminution is advantageously preceded by removing any adherent contaminants from the roots, e.g. by vigorous washing with water with a high-pressure cleaner. It is advantageously possible to wash the roots in the deep-frozen state in order to minimize the loss of mass of root material.

If necessary, the roots are initially comminuted coarsely, e.g. by chopping. Shredders are preferred for the further comminution. The product obtained is comminuted root material in the form of fibrous chips.

After the comminution, the comminuted root material is extracted with water, preferably at a temperature of 60°C to 95°C, most preferably 80–95°C. The extraction preferably takes place in the neutral to slightly alkaline pH range. A temperature of at least 60°C at pH 7–9 is advantageous because in this case enzymatic and acidic hydrolysis are suppressed. The concentration of comminuted root material in the water is preferably 10-40 % by weight, more preferably 20 – 30 % by weight, measured as fresh weight of roots based on the total weight of the extraction mixture.
Preferably a ratio between the dry matter of the shredded material used and the water as extraction medium is established which leads to a dry matter content in the extract of 8 - 12 % by weight and an inulin content of more than 6 % by weight, preferably 6 - 8 % by weight, based on the weight of the extract. A correspondingly suitable choice of extraction conditions, such as the ratio of water to root weight, can lead to a transfer of 80 - 90 % by weight of the inulin present in the roots into the extract. The aforementioned conditions are suitable to achieve a favorable crystallization and a high yield of the inulin from the extract, based on the observation that the high molecular weight inulin crystallizes from the extract even at a concentration as low as 5% by weight, based on the weight of the extract.

There is no special restriction on the extraction equipment, and conventional extraction techniques for plant material can be applied. It is most preferred for the extraction to take place in a jacket-heated extractor with agitator. In another highly preferred embodiment a heatable lauter tun is used as stirred extractor. Thus, the extraction of the inulin from the roots is combined with the separation of the extract from the spent chips by filtration, as described below. The extraction time after equilibration of the root/water mixture is preferably 30 min - 4 hours, preferably 1-2 hours. After this time, the extract is separated from the spent chips, e.g. by pumping off or straining off or filtration.

After separation of the extract from the spent chips, where appropriate, fibrous materials and plant fragments may remain as suspended materials in the extract. If present, these suspended materials are likewise removed from the extract. In this variant of the process, step b) of the process is thus followed, before step c), by a step in which suspended materials, mainly consisting of fibers, are removed from the extract. The acceptable amount of suspended materials and whether removal is to take place will be decided by the skilled worker from case to case. Removal of the suspended materials can take place by conventional separation techniques, as centrifugation or filtration. A desludging separator has proved particularly suitable. A screen or filter with appropriate fineness can also be used.

In a highly preferred embodiment, the suspended material can be filtered off by using the spent chips as a filter material. In this embodiment the spent chips are precipitated at the bottom of the extraction vessel equipped with a sieve at the bottom, like a lauter tun. The sieve is preferably a slit sieve. The precipitated spent chips are used as a filtration bed through which the extract flows. By using this technique a nearly quantitative removal of suspended
material is possible without using further filtration steps before further refining or brightening the extract or crystallizing the inulin.

The extracts are colored owing to their content of coloring constituents and colloidal suspended colorized matter. The coloring constituents consist, inter alia, of tannins and flavanoids and usually confer a yellow or brownish yellow and/or dark brownish color on the extract. The inulins which can be obtained directly from such extracts do not comply with the desired requirements concerning a neutral color. It is therefore necessary to remove the coloring constituents from the extract in step c) of the process. Process step c) for removing coloring constituents from plant extracts is generally also referred to as decolorization, clarification or "brightening" of plant extracts. These terms are equivalent in the context of the present invention.

The brightening can take place by adding lime and subsequent carbonation (CO₂ addition). The process of lime addition is known from the prior art and is used for example in obtaining sucrose from sugar beet. In an alternative brightening process, the interfering constituents are removed using an ion exchanger. Another useful process for removing coloring constituents by admixing magnesium ions (Mg²⁺) and an alkaline component to the plant extract is disclosed in co-pending applications PCT/EP2007/004028, PCT/EP2007/004029, US 60/796,818 and US 60/796,819. The whole of step c) may if required also be carried out more than once.

After step c), inulin is precipitated from the extract in step d). The precipitation can be effected for example by adding alcohols such as ethanol, methanol or isopropanol. In this case, depending on the amount of alcohol added or adjusted polarity of the liquid phase, initially high molecular weight inulin fractions are precipitated, so that it is possible to influence, via the amount of alcohol added, how quantitatively the inulin present in the extract is precipitated and which molecular weight fractions are predominantly obtained. Besides alcohol, it is also possible to employ other nonpolar organic liquids which are miscible with water.

For this purpose, in a particularly advantageous embodiment of this process step, to limit the use of alcohol, especially ethanol and isopropanol, the prepared extract is initially concentrated, preferably to one fourth to one fifth of its initial volume. The concentration can take place by evaporation or membrane filtration and a combination of both processes. Care must be taken in this case that the concentrate is kept hot during the concentration, preferably
at 60-95°C, in order to avoid precipitation of the inulin. An advantage of membrane filtration is the depletion, associated therewith, in low molecular weight substances accompanying the inulin. The subsequent precipitation of the inulin from the concentrate can be managed by the choice of increasing alcohol concentration so that the inulin is fractionated according to molecular size ranges which are characterized for example by the weight average degree of polymerization (DPw). Depending on the choice of the precipitation conditions, the result is fractions which have the DPw as described above.

It is more preferred to obtain inulin by cooling the extract than by alcoholic precipitation. The preferred conditions are such that the extract is cooled to a temperature of 2 - 10°C, more preferably 2 – 8°C, and kept at this temperature over a period of from 6 to 140 h, preferably 6 to 48 h, during which the inulin precipitates. The cooling rate and temperature, and the duration of the cooling influences the precipitation of the inulin from the extract and the breadth of the molecular weight distribution and thus at the same time the quantity. Choice of a longer period and lower temperature results in precipitation of more of low molecular weight inulins and a broader molecular weight distribution and thus a lower average molecular weight of the precipitated fraction. The precipitated inulin is separated from the liquid phase by conventional separation techniques such as, for example, centrifugation, decantation, and filtration.

In a preferred embodiment, inulin is crystallized for the first time after the extraction step b) and before step c) of the above described process. Such crystallisation is preferably done as described previously. Crystallization before step c) leads to an increase in the yield of high molecular weight inulin compared with direct brightening of the extract, and economizes the use of the brightening agents, i.e. magnesium compound and the alkaline component. It is advantageous to brighten the extract after the first crystallization of the inulin as in this case only the coloring constituents bound to the inulin crystals have to be removed, which leads to a similarly smaller amount of inulin bound to the brightening sludge.

A first precipitation and removal of the precipitated inulin can be followed by renewed cooling of the extract or addition of alcohol in order to obtain any inulin fractions which are still dissolved. A decision about repetition is made from case to case according to how quantitatively the inulin is to be obtained from the plants and what molecular weight distribution in the final product is desired.
The inulin concentration in the extract depends substantially on the inulin content of the roots and the concentration of the comminuted roots in the extract and is a further variable which has an effect on the precipitation of the inulin by cooling the extract. The dependence of the precipitation on the concentration can therefore be utilized in order to concentrate the liquid phase after the first precipitation, e.g. by evaporation, in order also to precipitate the low molecular weight fractions if this is desired.

In the last process step e), the precipitated inulin is reprecipitated. "Reprecipitation" means that the solid inulin, resulting from the previous process step, is redissolved and then precipitated and/or crystallized out of the solution again. Thus, process step e) can also be worded as: the inulin is dissolved and precipitated and/or crystallized again, wherein this step is done at least once. The crystallization differs from the precipitation in that predominantly crystalline structures are obtained.

The inulin is preferably dissolved under the influence of heat and preferably in water. Water with a temperature of 70-100°C, in particular 90-100°C, is particularly suitable.

The precipitation in step e) can take place by alcoholic precipitation as previously described. However, the inulin is preferably obtained by cooling the solution to 2 – 10°C, more preferably 2–8°C, over a period of 6 to 140 h, preferably 6 to 48 h.

The precipitation of the inulin dissolved in step e) can be repeated in order to obtain the inulin still remaining in the liquid phase. A decision about repetition is to be made from case to case according to how quantitatively the inulin is to be obtained from the plants and what molecular weight distribution in the final product is desired. The liquid phase can be concentrated in order to simplify the precipitation.

After reprecipitation, the resulting inulin solid is separated from the liquid phase by conventional separation techniques such as, for example, centrifugation, decantation, filtration.

In order to influence the molecular mass distribution and purity of the resulting inulin product, process step e) can be carried out more than once. It has emerged that the averages of the molecular weight and the averages of the degree of polymerization are shifted to higher values on repetition of the reprecipitation step e). It is thus possible to set various averages of
the molecular weight/degree of polymerization of the very long chain inulin as described
above within the described range.

If fine-particle impurities are still present, it is advantageous to insert one or more filtration
steps into the process. Any fine-particle impurities present are removed in the filtration. The
fineness of the filter is chosen by the skilled worker depending on the particle size of the
impurity.

The filtration step(s) can be inserted anywhere in the process after obtaining the extract. A
filtration step directly after obtaining the extract in step b) for example is advantageous. The
filtration step is to be distinguished from the removal of suspended materials as described
previously, because the particles removed by the filtration are finer than the suspended
materials, which consist mainly of fibers. In a further preferred embodiment, the filtration step
is carried out before step d).

The filtration step is preferably combined with a reprecipitation as described for process step
e). This entails the inulin being dissolved as previously described for step e), and the solution
then being filtered. After the filtration, the inulin is precipitated or crystallized out of the
filtered solution. The solid inulin resulting after the precipitation or crystallization can be
separated from the liquid phase by conventional separation techniques, such as, for example,
centrifugation, decantation and filtration.

In some cases the resulting inulin can be discolored by substances which can not be removed
by filtration. In such cases it is preferred to remove the coloring impurities by a treatment
with activated carbon. In one embodiment active charcoal is suspended in water and added to
an inulin solution at a temperature of above 80°C, preferably above 90°C. In case of a 20 %
by weight inulin solution the amount of active carbon is preferably in a range of 1 – 10 % by
weight, preferably 2 – 6 % by weight, more preferably 2 – 3 % by weight, based on the
weight of the inulin solution. After adsorption of the coloring impurities, the activated carbon
is removed by centrifugation and/or filtration. The activated-carbon suspension can be
preclarified by centrifugal separation of the activated-carbon sludge and then clarified by two-
stage filtration, for example with a combination of a kieselguhr precoat filter and a sheet filter.
It is important that during the separation of the active charcoal from the inulin solution the
temperature is maintained above 80°C, preferably above 90°C, in order to keep the inulin in
solution. After removal of the active charcoal, the inulin can be precipitated or crystallized
and separated from the liquid phase as described above.
After separation from the liquid phase, the final product can be washed again with water or a water/alcohol mixture. Washing with cold water at a temperature of 2-10°C is preferred. For this purpose, the inulin precipitate is slurried in water and the inulin is then sedimented again.

The resulting inulin is preferably dried in a further, last process step. The drying can take place by freeze drying, spray drying or drum drying. In a preferred embodiment, the inulin is in spray-dried form. It is self evident that in case of a spray drying process a precipitated or crystallized inulin must be brought into suspension (in water below about 80°C) or into solution (in water above about 80°C) again. Alternatively, a last precipitation or crystallization step, as described above, can be omitted and the suspended or dissolved inulin from the process can directly be spray dried. It is possible by adding spray-dried inulins to liquid prepared food products for the viscosity to be increased particularly effectively.

It has surprisingly been found that a frozen foodstuff with inhibited ice crystal growth can be produced by adding a very long chain inulin as previously described to the ingredients for forming a frozen foodstuff.

By applying the teaching of the invention ice crystal growth in a frozen foodstuff can be inhibited without unduly affecting the texture of the frozen foodstuff. In case of frozen confections, especially in case of ice cream, no “chewiness” or “gumminess” of the confection is observed when applying the teaching of the invention.

The terms “inhibited growth of ice crystals” or “inhibition of ice crystal growth” of the present invention in one sense mean that the ice crystal growth in a frozen foodstuff comprising very long chain inulin is retarded in comparison to a frozen foodstuff not comprising very long chain inulin when both foodstuffs are treated in the same manner and/or stored under similar conditions for the same period of time. In this sense, said terms mean that the ice crystal size in a frozen foodstuff comprising very long chain inulin is reduced in comparison to a comparative frozen foodstuff not comprising very long chain inulin when both foodstuffs are treated in the same manner and/or stored under similar conditions for the same period of time. A possible treatment is a temperature fluctuation (heat shock) process as described below.

The term “ice crystals” designates crystals comprising frozen water as the predominant or only component.
The inhibition of ice crystal growth occurs in a frozen foodstuff comprising very long chain inulin during shelf life of said foodstuff. Particularly, ice crystal growth is inhibited when said foodstuff is subjected during its shelf life to temperature fluctuations related to the storage and transportation of frozen foodstuff, particularly ice cream, ice milk, yogurt, or water ice product. Said temperature fluctuations can be simulated by a temperature cycling process, comprising: holding the foodstuff, especially a confection like ice cream, ice milk, yogurt, or water ice, at -25°C for 24 h, transferring the foodstuff to -15°C and holding for additional 24 h, returning to -25°C, wherein this sequence of steps is repeated four times. Any other testing method commonly used in the industry for evaluating the effect of temperature fluctuations on foodstuffs may also be employed. Above mentioned temperature fluctuations or temperature cycling process are commonly designated as “heat shock” or “temperature abuse”.

In one aspect, the present invention relates to a method for inhibiting the growth of ice crystals in a frozen foodstuff, the method comprising adding to the ingredients for forming a frozen foodstuff a very long chain inulin with a weight average degree of polymerization of between 50 and 103, or preferably any other very long chain inulin as previously described.

The very long chain inulin is added in an amount that is sufficient to inhibit crystal growth. Preferably, the very long chain inulin is added in an amount of 0.1 % to 8 % by weight, more preferably 0.1 % to 5 % by weight, most preferably 0.5 % to 4 % by weight of the total composition for forming the frozen foodstuff. Another preferred range is 1.5 % to 2.6 %. The term “total composition” means the total composition of ingredients for forming the frozen foodstuff.

In a further aspect, the present invention relates to a frozen foodstuff with inhibited ice crystal growth comprising a very long chain inulin with a weight average degree of polymerization of between 50 and 103, or preferably any other very long chain inulin as previously described.

Preferred frozen foodstuffs according to the invention are selected from confections, bakery, frozen dough, prebaked or parbaked frozen dough, meat, sausages, sauces, and frozen convenient foods, all aforementioned foodstuffs being frozen.

Highly preferred according to the invention are frozen confections as for example ice cream, such as milk ice cream, fruit ice cream, low fat ice cream or soft (serve) ice cream, water ice, frozen yogurt, sorbet, ice milk, frozen fruit juice.
A preferred ingredient composition for forming a frozen confection comprises a fat present in an amount of 0.1 % to 18 % by weight, more preferably 1 % to 10 % by weight of the total composition; a sweetener present in an amount of 2 % to 25 % by weight, more preferably 5 % to 20 % by weight of the total composition; and non-fat milk solids present in an amount from 6 % to 15 % by weight, more preferably 8 % to 12 % by weight of the total composition. Very long chain inulin is added in an amount as already described above.

Another preferred ingredient composition for forming a frozen confection comprises a fat present in an amount of 0.1 % to 18 % by weight, more preferably 1 % to 10 % by weight of the total composition; a sugar, preferably glucose or sucrose, present in an amount of 5 % to 15 % by weight of the total composition; an alternative sweetener other than glucose or sucrose present in an amount of 3% to 8% by weight of the total composition and non-fat milk solids present in an amount from 6 % to 15 % by weight, more preferably 8 % to 12 % by weight of the total composition. Very long chain inulin is added in an amount as already described above.

Still another preferred ingredient composition for forming a frozen confection, which is preferably a low fat ice cream, comprises 0.2 % – 5 % by weight, preferably 0.5 % – 4 % by weight fat, 40 % - 85 % by weight, preferably 50 % - 80 % by weight milk, preferably whole milk, 0 %– 8 % by weight, preferably 0 % - 7 % by weight non-fat dry milk, 0 % – 25 % by weight, preferably 0 % - 20 % by weight condensed skim, 2 % - 20 % by weight, preferably 5 % - 15 % by weight sugar, 0.1 % – 5 % by weight, preferably 0.5 - 4 % by weight very long chain inulin as described above.

The fat used may be a dairy fat, a non-dairy fat, or a mixture of both. When the fat is a dairy fat, it may be for instance, any milk fat source such as butter oil, butter, real cream, or a mixture thereof. When the fat is a non-dairy fat it may be, for instance, an edible oil or fat, preferably a vegetable oil such as coconut oil, palm kernel oil, palm oil, cotton oil, peanut oil, olive oil, soy bean oil, rapeseed, olive, hydrogenated coconut oil, hydrogenated soybean oil, palm olein etc., or mixtures thereof.

The non-fat milk solids may be of whey origin, for example powdered or concentrated defatted sweet whey or powdered or concentrated skim milk. The non-fat milk solids may also be derived from a commercial mixture of milk powder and whey proteins, whose functionality has been modified by specific denaturation treatments.
The sugar used may be sucrose, glucose, fructose, lactose, dextrose, trehalose, invert sugar either crystalline or liquid syrup form, or mixtures thereof.

The sweetener may be a corn sweetener in either a crystalline form of refined corn sugar (dextrose and fructose), a syrup, as glucose syrup, maltose syrup, or a dried corn syrup (corn syrup solids), a liquid corn syrup, a maltodextrin, glucose, or a mixture thereof.

An emulsifier can be present as a further ingredient in a frozen confection. For example, the emulsifiers include but are not limited to saturated mono-diglyceride, sorbitan tristearate ("STS"), sorbate, lactylated monoglycerides, acetylated monoglycerides, unsaturated monoglycerides, including monoglycerides with oleic acid, linoleic acid, linolenic acid, or other commonly available higher unsaturated fatty acids. The emulsifier blend should be present in an amount of about 0.01% to about 5%, preferably of about 0.04% to about 1%, and more preferably of about 0.2% to about 0.5% by weight of the total composition for forming the frozen confection.

A stabilizer can be present as a further ingredient in a frozen confection. The stabilizer may be, for instance, a hydrocolloid such as agar, gelatin, gum acacia, guar gum, locust bean gum, gum tragacanth, carrageenan and its salts, carboxymethyl cellulose, sodium alginate or propylene glycol alginate, or any mixture of hydrocolloids, carob flour, guar flour, alginates xanthan, starches. Preferably the stabilizer is present in an amount of about 0.1% to about 0.5% by weight of the total composition for forming the frozen confection.

In another aspect of the invention is provided a soft frozen confection. The term "soft" as used herein means that the frozen confection is scoopable at a temperature of -18°C or less. The term "scoopable" is used in its ordinary meaning in that the ice cream is sufficiently soft at freezer temperatures so that it can be scooped from a container by hand or extracted from a cartridge at that temperature for placement in a dish or cone as contrasted to a solid frozen block from which the confection is difficult to separate and remove. Thus, the invention provides a frozen confection which is easily scoopable after being taken from a home freezer by a consumer. In this regard, the frozen confection comprises at least one freezing point depressant. Suitable freezing point depressants include polyols such as sorbitol, mannitol, lactitol, xylitol, maltitol, glycerol, ethanol, and mixtures thereof. Preferably, the freezing point depressant is glycerol or ethanol. Other freezing point depressants include sucrose, dextrose, tagatose, fructose, glucose syrup, polydextrose, preferably corn syrup of a dextrose equivalent (DE)>40, or a mixture thereof. The freezing point depressant softens the frozen confection
thereby making it sufficiently soft to scoop from a container or receptacle at a temperature of about -18°C or less. Preferably, the freezing point depressant is present in an amount of about 0.5 to about 5% by weight of the frozen confection. Thus, a soft, scoopable frozen confection is with inhibited ice crystal growth is provided in which the freezing point has been reduced by the addition of freezing point depressants. Furthermore, as an added benefit of using a freezing point depressant is that a frozen confection may be produced having less fat, which enables manufacture of a less expensive confection such as ice cream, or alternatively, manufacture of an ice cream with reduced calories but which has the same softness and texture as a conventional high-fat ice cream.

The composition for forming a frozen confection may further contain ingredients which are selected from one or more members of the group consisting of colorants, mineral salts, flavorings, flavor modifiers, flavor enhancers, thickening agents, egg yolk, egg white, fruit juices and fruit syrups, as well as inclusions, such as but not limited to fruit or fruit pieces, for example, nuts, hazelnuts, whole or in pieces, or pieces of chocolate.

As stated above, the term “frozen confection” as used herein, unless otherwise indicated, encompasses ice cream, such as milk ice cream, fruit ice cream, low fat ice cream or soft (serve) ice cream, water ice, frozen yogurt, sorbet, ice milk, frozen fruit juice etc. The mixture suitable for a frozen product may be any conventional mix such as an ice cream mix, a frozen yogurt mix, a water ice mix, a fruit juice mix, a sorbet mix, or a combination thereof containing very long chain inulin as described above. An ice cream mix may contain fat, non-fat milk solids, carbohydrates, or stabilizers together with water and, if desired, other conventional ingredients, such as mineral salts, colorants, flavorings, inclusions, etc. A water ice mix comprises fruit juices, sugar, stabilizer, and small amounts of milkfat and non-fat milk solids.

In another aspect of the invention, the frozen confection is a light frozen confection. The term “light frozen confection” as used herein is intended to mean a frozen confection having a reduced calorie value. Advantageously, the light frozen confection of the present invention shows inhibited ice crystal growth after being exposed to temperature fluctuations, but also presents a smooth, and creamy, texture, a feature often lacking from conventional low calorie frozen desserts. This is due to the fact that very long chain inulin functions not only as an inhibitor of the growth of ice crystals but also acts as a fat or oil substitute with improved organoleptic properties (mouthfeel) and as a texturizing agent in the frozen confection. A
preferred frozen confection having a reduced calorie value is a low fat ice cream, preferably with a fat content of equal or lower than 5% by weight, more preferably of equal or lower than 3% by weight of the total composition. In the same manner, the use of very long chain inulin for inhibiting the growth of ice crystals in a frozen confection relates in a preferred embodiment to a low fat ice cream as defined above. The method for inhibiting the growth of ice crystals and the method for producing a frozen confection according to the appended claims in a preferred embodiment also relate to a frozen confection which is a low fat ice cream as defined above.

Very long chain inulin (VLCI) produces ice cream mixes with increased viscosity and body. Ice cream resulting from these mixes can resemble higher fat ice creams depending on fat content, solids and accompanying carbohydrates. VLCI is particularly effective in achieving a fat-like quality in ice cream. Where 3-4 % by weight chicory inulin with lower DP than VLCI can be required to act as a fat mimetic, VLCI is effective at levels between 1.5-2.6 % by weight, based on the total composition. Low fat ice cream with VLCI at levels of 1.5-2.6 % by weight in the mix is smoother and more like ice cream with 10 % fat. In fact VLCI levels of less than 2.6 % by weight might be optimal for low fat ice cream depending on solids and sugar levels and the amount of time the mix is allowed to age prior to freezing.

Additionally to very long chain inulin as a fat substitute, a frozen confection of the invention may also comprise a fat that is rich in polyunsaturated fatty acid. Suitable oils may be non-hydrogenated vegetable fats such as those high in oleic oils, or those having blends containing long-chain polyunsaturated fatty acids such as gammalinolenic, docosahexaenoic, and eicosapentaenoic acids. Thus, the low saturated fat may be obtained from sources such as seeds, sunflower, canola, safflower, rapeseed, soy, rice, borage, nuts including walnuts, almonds, macadamia, coconut, palm, palm kernal, apricot kernal, and other plants including olive, and peanut, or fish or microbial oils.

In a further aspect, the present invention relates to a method for producing a frozen confection with inhibited ice crystal growth during shelf life and/or temperature abuse, the method comprising: adding to the ingredients for forming a frozen confection a very long chain inulin with a weight average degree of polymerization of between 50 and 103, or preferably any other very long chain inulin as previously described; combining the very long chain inulin and the other ingredients with shear mixing to form a composition; pasteurizing and homogenizing the composition; and freezing the composition.
The composition of a frozen confection, which is produced by such method, the kind of ingredients, the amounts of ingredients, including very long chain inulin, are preferably chosen as already described above with respect to the frozen confection.

In one embodiment of the method, the preparation of frozen product of the invention can be carried out using conventional equipment for ice cream manufacture. The first step comprises combining the ingredients and the very long chain inulin under shear mixing to disperse and/or solubilize the ingredients to form a composition which is preferably an already homogeneous mass. One of ordinary skill in the art with little or no experimentation can determine mixing time and conditions to obtain the desired homogeneous mass.

In the next steps, the composition is pasteurized and homogenized. Pasteurization can be carried out first, followed by homogenization, or vice versa. It is more preferred to do the pasteurization step first.

Pasteurization of the composition is conducted under conditions commonly used in the industry. The pasteurization step is conducted at a temperature of about 60°C to about 100°C, preferably at about 75°C to about 85°C, for a time of about 10 seconds to about 30 minutes, preferably for at least about 25 seconds. Preferably, pasteurization is conducted by either high temperature short time (HTST) or low temperature long time (LTLT) processing.

The composition is conventionally homogenized, e.g., in a two stage homogenizer. The first stage is conducted at a pressure of about 70 bar to about 250 bar, preferably of about 100 bar to about 150 bar, more preferably about 140 bar. The second stage is conducted at a pressure of about 0 bar to about 50 bar, preferably of about 20 bar to about 35 bar. Homogenization is done at a temperature of about 62°C to about 75°C.

If homogenization is conducted before pasteurization, the composition is preheated after shear mixing to a temperature of about 62°C to about 75°C. If homogenization is conducted after pasteurization, the composition is cooled down to said temperature after pasteurization.

A composition pasteurized and homogenized under standard processing conditions results in a smooth viscous blend. Homogenization not only reduces the size of hydrated VLCI particles but also contributes to hydration itself.

The pasteurization and homogenization step is followed by cooling the composition to a temperature of about 0°C to about 10°C, and preferably at about 4°C. The mix is preferably
aged by allowing to stand at a temperature of about 0°C to about 6°C, preferably of about 1°C to about 5°C and for a time of about 1 hour to about 24 hours, preferably of about 2 hours to about 18 hours and more preferably of about 4 hours to about 12 hours. The mix is then colored and flavored as needed.

Optionally the composition may be aerated to create fine air cells within the composition. Aeration can be done as disclosed in US patent application publication US20050123666 A1. If aerated, the frozen product shows preferably an overrun of about 20% to about 120% for ice cream products and an overrun of about 5% to about 50% for water ice products. Overrun means the % increase in volume of frozen product greater than the volume of mix/composition used to produce that frozen product.

Subsequently, the frozen product is packaged into bulk containers, extruded for bars or cones, or packaged into small containers. Bulk containers include container sizes of 11.36 l (3 gallons) to 0.5 l, and small containers include container sizes of 250 ml to 50 ml.

The frozen products may afterwards be stored at a freezing temperature, usually at a temperature in the range of about -25°C to about -35°C, preferably of about -28°C to about -32°C, and more preferably at about -30°C.

If desired, the product can be repackaged before shipping. Also for individual sized portions the frozen products may be coated, for instance with chocolate or a compound coating. Compound coatings include coatings which do not contain 100% cocoa fat and coatings that contain any vegetable oil, such as canola oil, corn oil, soy oil, coconut oil, etc., or mixtures thereof. These coatings may also contain inclusions such as nut pieces, fruit pieces, rice crisps, or other additives therein. Furthermore, the frozen product may be placed between cookies, or other edible substrates to form ice cream sandwiches or the like. The final frozen products are then packaged and stored at a freezing temperature.
DESCRIPTION OF THE FIGURES:

Figure 1: Ice crystals in temperature abused, 3% fat ice cream with and without VLCI

Figure 2: Ice crystal growth in the three ice cream variables stored at -25°C (8 weeks) and -17°C (4 weeks) and samples cycled between -25°C and -15°C.

EXAMPLES

Hereinafter, the present invention is illustrated by working examples, which are intended to show the general idea of the invention, but which are not to be understood to restrict the scope of protection as defined by the appended claims.

Example 1 – Low Fat Ice Cream Mix with Very Long Chain Inulin (VLCI) – Batch process

Low fat ice cream mixes were produced in a batch process format using a 5 liter jacketed kettle. Formulas compared viscosity of mixes containing two different inulin types in ice cream mixes containing 3% butter fat. Formulas are listed in Table 1.
Table 1: Formulas for ice cream mixes containing inulin

<table>
<thead>
<tr>
<th></th>
<th>Full Fat Control</th>
<th>Low Fat Control</th>
<th>Low Fat with 4% VLCI</th>
<th>Low Fat with 4% Chicory Inulin</th>
<th>Low Fat with 3% VLCI</th>
<th>Low Fat with 4% VLCI (sheared)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cream 40%</td>
<td>24.78%</td>
<td>0.22%</td>
<td>0.67%</td>
<td>0.67%</td>
<td>0.56%</td>
<td>0.67%</td>
</tr>
<tr>
<td>Whole Milk</td>
<td>56.39%</td>
<td>78.68%</td>
<td>73.87%</td>
<td>73.87%</td>
<td>75.07%</td>
<td>73.87%</td>
</tr>
<tr>
<td>Non-fat Dry Milk</td>
<td>4.65%</td>
<td>6.92%</td>
<td>7.28%</td>
<td>7.28%</td>
<td>7.19%</td>
<td>7.28%</td>
</tr>
<tr>
<td>Sucrose</td>
<td>14.00%</td>
<td>14.00%</td>
<td>14.00%</td>
<td>14.00%</td>
<td>14.00%</td>
<td>14.00%</td>
</tr>
<tr>
<td>VLCI 1)</td>
<td>--</td>
<td>--</td>
<td>4.00%</td>
<td>--</td>
<td>3.00%</td>
<td>4.00%</td>
</tr>
<tr>
<td>Chicory inulin (DPw 33)</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>4.00%</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>Stabilizer/emulsifier 3)</td>
<td>0.18%</td>
<td>0.18%</td>
<td>0.18%</td>
<td>0.18%</td>
<td>0.18%</td>
<td>0.18%</td>
</tr>
</tbody>
</table>

1) very long chain inulin, DPw 75-81 (determined by GPC RI-MALLS)
2) Keystone 9185, Mainstreet Ingredients: guar gum, xanthan gum, carrageenan, polysorbate 80, mono and diglycerides

Ingredients were added to the kettle under agitation (300 rpm) and heated to 82°C for one minute, cooled rapidly to 30°C and transferred to one liter jars and stored at 4°C for 24 hours. One low fat batch containing 4% VLCI was further processed with additional shear (1200 rpm for 60 seconds) after pasteurization.

Table 2

<table>
<thead>
<tr>
<th></th>
<th>VISCOSITY cP</th>
<th>SENSORY DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full Fat Control</td>
<td>91.0</td>
<td>Smooth and creamy / Well rounded vanilla flavor</td>
</tr>
<tr>
<td>Low Fat Control</td>
<td>64.0</td>
<td>Very thin mouthfeel / Sharp vanilla flavor</td>
</tr>
<tr>
<td>Low Fat with 4% VLCI</td>
<td>240.0</td>
<td>Very thick, viscous texture / mild vanilla flavor</td>
</tr>
<tr>
<td>Low Fat with 4% Chicory Inulin</td>
<td>98.0</td>
<td>Thicker than low fat control / Sharp vanilla and caramel flavor</td>
</tr>
<tr>
<td>Low Fat with 3% VLCI</td>
<td>142.0</td>
<td>Smooth mouthfeel, not as thick as 4% VLCI / mild vanilla flavor</td>
</tr>
<tr>
<td>Low Fat with 4% VLCI (sheared)</td>
<td>189.0</td>
<td>Not as viscous as 4% VLCI without shear, chewy texture / mild vanilla flavor</td>
</tr>
</tbody>
</table>

Viscosity of mixes, measured in a Brookfield RVT Viscometer using a RV-1 spindle, are reported in Table 2. Each sample tasted and evaluated for comparative sensory properties.
Low fat mixes containing 3% or 4% VLCI had highest viscosities, considerably higher than mixes with long chain chicory inulin or 12% fat. Shearing the mix with 4% VLCI after pasteurization reduced the viscosity by approximately 25%.

Mouthfeel differences trended with the viscosity data. 3% VLCI was more similar to the mouthfeel of 12% fat mixes than the other inulin containing mixes.

Example 2 – Effects of Very Long Chain Inulin on Low Fat Ice Cream

Methods

As a reference, a typical “economy” ice cream mix was produced to the following specifications: 10% butterfat, 11% non-fat dairy solids, and 36% total solids. Very Long Chain Inulin (DPw 60, GPC-MALLS) was used at a concentration of 2.6% w/w in the mix.

Formulas for full fat economy ice cream, 3% (low fat) ice cream and 3% (low fat) ice cream with VLCI are listed in Table 3. To maintain similar total solids concentration in the low fat ice creams, corn syrup solids were used to replace part of the butter fat and sucrose used the in 10% fat formula.

Table 3

<table>
<thead>
<tr>
<th>INGREDIENT LIST (%)</th>
<th>10% Fat Control</th>
<th>3% Fat Control</th>
<th>VLCI&lt;sup&gt;1)&lt;/sup&gt; 3% Fat</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cream</td>
<td>21.56</td>
<td>2.31</td>
<td>2.54</td>
</tr>
<tr>
<td>Whole milk</td>
<td>48.87</td>
<td>57.34</td>
<td>55.00</td>
</tr>
<tr>
<td>Condensed Skim</td>
<td>14.91</td>
<td>21.19</td>
<td>21.70</td>
</tr>
<tr>
<td>Sucrose</td>
<td>14.50</td>
<td>10.00</td>
<td>10.00</td>
</tr>
<tr>
<td>Corn Syrup Solids (36 DE)</td>
<td>-</td>
<td>9.00</td>
<td>8.00</td>
</tr>
<tr>
<td>VLCI</td>
<td>-</td>
<td>-</td>
<td>2.60</td>
</tr>
<tr>
<td>Stabilizer/Emulsifier</td>
<td>0.16</td>
<td>0.16</td>
<td>0.16</td>
</tr>
<tr>
<td>Vanilla</td>
<td>0.8</td>
<td>0.8</td>
<td>0.8</td>
</tr>
</tbody>
</table>

<sup>1)</sup> very long chain inulin, DPw 59 (determined by GPC RI-MALLS)
Total non-fat dairy solids (serum solids) level is routinely 1-2% higher in low fat ice cream formulas to accommodate the loss of butter fat. A commercial stabilizer blend (Keystone 9185, Mainstreet Ingredients, guar gum, xanthan gum, carrageenan, polysorbate 80, mono and diglycerides) was used at a range recommended by the manufacturer for 10% butter fat ice creams. Stabilizer concentration was held constant for all three formulas.

154 kgs (70 pounds) of mix was produced for each variable by blending the dry ingredients (sucrose, corn syrup solids, VLCI and stabilizer) and separately combining the condensed skim, whole milk and cream. The dry blend was then slowly added to the milk blend with agitation using a lightning mixer. Mixes were pasteurized at 82°C for 20 seconds in a Microthermixin™ fluid processing system, cooled immediately to 65°C and homogenized in a two stage homogenizer (138/34.5 bar). Homogenized mixes were cooled to 4°C and aged for 4 hours prior to freezing.

Before freezing double fold, natural vanilla was added to each mix at a rate of 4ml/kg. Mixes were frozen in a Crepaco, single barrel, continuous ice cream freezer. During freezing over run was targeted for 80% (0.62 g/ml). Exit temperature for all variables was -5°C. Ice cream was packaged into pint containers and “hardened” at -27°C.

Ice cream mix viscosity was measured using a Brookfield RTV viscometer with an RV-1 spindle at 2.5 to 20 RPMs. Ice cream was evaluated for sensory attributes after one day, one week and one month of storage followed by monthly intervals thereafter. At one week melt characteristics were measured by placing 250g of ice cream on a sieve at room temperature and measuring the amount of melted product over time. Geometry of the melted product was also observed.

Sensitivity to ice crystal growth during shelf life was determined by cycling the products between two different temperatures to accelerate growth rates and comparing them to control samples held a constant temperature. Temperature stressed ice cream was held at -25°C for 24 hours then transferred to -15°C and held for an additional 24 hours before returning to -25°C. This cycling process was repeated four times. Temperature cycled ice cream was compared to
control samples stored at -27°C for 8 weeks and -15°C for 4 weeks. Ice crystals were imaged by cold stage microscopy. Size distribution was determined by digital image analysis of ice crystals in the micrographs.

5 Results:

VLCI, like chicory inulin (DPw 33), produces ice cream mixes with increased viscosity and body. Ice cream resulting from these mixes can resemble higher fat ice creams depending on fat content, solids and accompanying carbohydrates. VLCI is particularly effective in achieving a fat-like quality in ice cream. Where 4% chicory inulin can be required to act as a fat mimetic, VLCI is effective at levels between 1.75 -2.6%.

Mix pasteurized and homogenized under standard processing conditions results in a smooth viscous blend. Homogenization not only reduces the size of hydrated VLCI particles but also contributes to hydration itself.

Low fat ice cream with VLCI at levels of 2.6% in the mix is smoother and more like ice cream with 10% fat. In fact VLCI levels of less than 2.6% might be optimal for low fat ice cream depending on solids and sugar levels and the amount of time the mix is allowed cure prior to freezing.

Mixes containing VLCI produced on the Microthermix™ were smooth tasting. The texture of the mix with VLCI tasted creamier and was more “mouth coating” than its low fat counter part without VLCI. While it did not have the unctuousness of full fat mix, it approximated the 10% fat mix more than the 3% fat mix.

The viscosity of low fat mix with VLCI was higher than either the low fat or full fat mixes. In samples measured 10 days after preparation (Table 4), viscosity of the VLCI containing mix was nearly double that of the 10% fat or the 3% fat mixes.
Table 4

<table>
<thead>
<tr>
<th></th>
<th>Mix Viscosity cP (day 10)</th>
<th>Percent Over Run</th>
<th>Percent Melt</th>
</tr>
</thead>
<tbody>
<tr>
<td>10% Fat</td>
<td>97</td>
<td>75</td>
<td>20</td>
</tr>
<tr>
<td>3% Fat</td>
<td>91</td>
<td>70</td>
<td>30</td>
</tr>
<tr>
<td>3% Fat with VLCl</td>
<td>178</td>
<td>69</td>
<td>30</td>
</tr>
</tbody>
</table>

Overrun for all samples was targeted at ~80% and was measured after hardening at 71%. When measured after one week there was no significant difference between the three formulas (Table 4). There was no evidence of shrink or loss of volume during storage. Melt profiles followed a pattern where the 3% fat ice creams “melted” more rapidly which is typical of low fat products. The method actually measures melt as well as flow or separation from the body of the ice cream so differences are a combination of actual melting rates and structure which can result from churned fat during freezing or from over stabilization of the ice cream mix. Melt characteristics did not suggest either defect in any of the samples.

Ice crystal size in low fat ice cream (3%) doubled from 31 μm to 65 μm during temperature cycling (Table 5). Ice crystal size in 3% fat ice cream with VLCl did not change during temperature abuse suggesting that VLCl retards the growth of ice crystals in low fat ice cream (Figures 1 and 2). There was no significant crystal growth in full fat ice cream or ice cream with 3% fat plus VLCl for any temperature storage condition (Figure 2). Crystals in full fat ice cream appeared to grow from an average of 34 μm to 41 μm after temperature cycling. However, while there was apparent crystal growth trend in the 10% fat ice cream, the change was not significant.
Table 5 - Ice crystal size distribution for ice creams containing 10% fat, 3% fat and 3% fat with VLCI after temperature cycling from -25°C to -15°C (all sizes in μm)

<table>
<thead>
<tr>
<th></th>
<th>Initial Crystal Size</th>
<th>Crystal Size After Temperature Abuse</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10% Fat</td>
<td>3% Fat</td>
</tr>
<tr>
<td>Mean size</td>
<td>34,1</td>
<td>31,1</td>
</tr>
<tr>
<td>Std. Dev.</td>
<td>13,6</td>
<td>12,2</td>
</tr>
<tr>
<td>Minimum</td>
<td>9,0</td>
<td>3,1</td>
</tr>
<tr>
<td>Maximum</td>
<td>80,5</td>
<td>74,3</td>
</tr>
<tr>
<td>C.V. (%)</td>
<td>39,9</td>
<td>39,2</td>
</tr>
<tr>
<td>Sample size</td>
<td>559,0</td>
<td>457,0</td>
</tr>
</tbody>
</table>

Day One. From a sensory perspective VLCI containing ice cream was noticeably different from low fat ice cream without VLCI when tasted as soft ice cream (exiting the freezer at -5°C) and after hardening at -27°C. Low fat ice cream was “thin”, lacked body and melting rapidly in the mouth while low fat ice cream with VLCI had more body and was retained longer in the mouth. As soft ice creams both had strong almost objectionable vanilla flavor profile common to low fat ice creams using natural vanilla. After hardening the harsh vanilla flavor was reduced in both ice creams, but to a greater extent in the ice cream with VLCI. Hardened VLCI ice cream developed even more body, retention in the mouth and creaminess.

One Week. There was little change from the sensory characteristics observed in the hardened ice creams at day one. The body and mouthfeel of the VLCI containing low fat ice cream was more like that of the 10% fat than the 3% fat.

One Month. All ice creams had noticeable ice crystals, but in a set of temperature abused samples, VLCI clearly had the least noticeable amount of ice crystals while the 3% fat sample without VLCI had the most. 10% fat was between the extremes.
Summary

Very long chain inulin changed the texture of 3% fat ice cream, making it smoother, longer lasting in the mouth and creamier than 3% fat alone. In minimally stabilized products the increase in cohesiveness was not perceived as chewy or gummy, a negative often associated with stabilizers and high levels of inulin. In temperature stressed samples, VLCI retarded ice crystal growth when compared to low fat ice cream without VLCI.
Claims

1. The use of an inulin with a weight average degree of polymerization of between 50 and 103 for inhibiting the growth of ice crystals in a frozen foodstuff.

2. The use of claim 1, wherein the inulin is added to the ingredient mix for forming a frozen foodstuff.

3. The use of claim 1 or 2, wherein the inulin is added in an amount of 0.1 % to 8 % by weight of the total composition for forming the frozen foodstuff.

4. The use of one of claims 1-3, wherein the frozen foodstuff is a frozen confection.

5. The use of claim 4, wherein the frozen confection has a reduced calorie value.

6. The use of claim 4 or 5, wherein the frozen confection is an ice cream, a water ice, a frozen fruit juice, a frozen yogurt, a sorbet, an ice milk or a mixture thereof.

7. The use of one of claims 4-6, wherein the composition for forming the frozen confection comprises a fat present in an amount of 0.1 % to 18 % by weight of the total composition; a sweetener present in an amount of 2 % to 25 % by weight of the total composition; and non-fat milk solids present in an amount from 6 % to 15 % by weight of the total composition.

8. The use of claim 7, wherein the non-fat milk solids are of whey origin.

9. The use of one of claims 4-8, wherein the frozen confection is a frozen confection with inhibited ice crystal growth in a temperature cycling process comprising:
    holding the confection at -25°C for 24 h,
    transferring the confection to -15°C and holding for additional 24 h,
    returning to -25°C, wherein this sequence of steps is repeated four times.

10. A frozen foodstuff with inhibited ice crystal growth comprising an inulin with a weight average degree of polymerization of between 50 and 103.
11. The frozen foodstuff of claim 10, which is selected from frozen bakery, frozen dough, frozen meat, frozen sausages, and frozen sauces.

12. The frozen foodstuff of claim 10, which is a frozen confection.

13. The frozen foodstuff of claim 12, which is selected from milk ice cream, soft ice cream, water ice, frozen fruit juice, frozen yogurt, sorbet, ice milk or a mixture thereof.

14. The frozen foodstuff of claim 12 or 13, wherein the frozen confection has a reduced calorie value.

15. The frozen foodstuff of one of claims 12-14, wherein the total composition for forming the frozen confection comprises a fat present in an amount of 0.1 % to 18 % by weight of the total composition; a sweetener present in an amount of 2 % to 25 % by weight of the total composition; and non-fat milk solids present in an amount from 6 % to 15 % by weight of the total composition.

16. The frozen foodstuff of claim 15, wherein the non-fat milk solids are of whey origin.

17. The frozen foodstuff of one of claims 10-16, including inulin in an amount of 0.1 % to 8 % by weight of the total composition for forming the frozen foodstuff.

18. A method for producing a frozen confection with inhibited ice crystal growth, the method comprising:

   adding to the ingredients for forming a frozen confection an inulin with a weight average degree of polymerization of between 50 and 103;

   combining the inulin and the other ingredients with shear mixing to form a composition;

   pasteurizing and homogenizing the composition;

   freezing the composition.

19. The method of claim 18, wherein the inulin is added in an amount of 0.1 % to 8 % by weight of the total composition.

20. The method of claims 18 or 19, wherein the composition comprises a fat present in an amount of 0.1 % to 18 % by weight of the total composition; a sweetener present in an amount of 2 % to 25 % by weight of the total composition; and non-fat milk solids present in an amount from 6 % to 15 % by weight of the total composition.
Figure 2

Ice crystal growth in low fat ice cream with VLCI
# INTERNATIONAL SEARCH REPORT

**International application No**

PCT/EP2008/005216

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## A. CLASSIFICATION OF SUBJECT MATTER

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<th>INV.</th>
<th>A23G9/34</th>
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According to International Patent Classification (IPC) or to both national classification and IPC

## B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

A23G A21D A23L

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched:

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, FSTA, WPI Data

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## C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
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<td>X</td>
<td>WO 2006/108697 A (BAYER CROPSCIENCE GMBH [DE]; HELLWEGE ELKE [DE]; PEETERS ROGER [NL]; P) 19 October 2006 (2006-10-19) cited in the application page 2, line 1 - last line page 8, paragraph 1; claim 14</td>
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<td>X</td>
<td>US 5 169 671 A (HARADA TSUTOMU [JP] ET AL) 8 December 1992 (1992-12-08) column 2, line 49 - line 65; claims 1,4; example 26</td>
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<td>X</td>
<td>US H2 095 H1 (YOUNG TIMOTHY J [US]) 6 January 2004 (2004-01-06) claims 1,4; examples 10,11</td>
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**X** See patent family annex.

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Further documents are listed in the continuation of Box C.

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**X** See patent family annex.

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**"&" document member of the same patent family**

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**Date of the actual completion of the international search**

19 September 2008

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**Date of mailing of the international search report**

07/10/2008

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**Name and mailing address of the ISA/ European Patent Office, P.B. 5618 Patentlaan 2 NL - 2280 HV RIJWijk Tel. (+31-70) 340-2049, Rx 31 651 epo ni. Fax: (+31-70) 340-2016**

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**Authorized officer**

Rinaldi, Francesco

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* Form PCT/ISA/210 (second sheet) (April 2005)
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<td>SCHALLER-POVOLNY L A ET AL: &quot;Sensory attributes and storage life of reduced fat ice cream as related to inulin content.&quot; JOURNAL OF FOOD SCIENCE, vol. 64, no. 3, 1999, pages 555-559, XP002488984 abstract Conclusions</td>
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<td>WOUTERS R: &quot;Technological and nutritional advances in ice cream production.&quot; LATTE, vol. 23, no. 8, 1998, pages 24-26, XP009103438 page 24, left-hand column</td>
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<td>DE 38 15 950 A1 (JUchem FRANZ GMBH &amp; CO KG [DE]) 23 November 1989 (1989-11-23) column 1, line 59 - line 60; claims</td>
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