Commission Implementing Regulation (EU) 2017/2470 of 20 December 2017 establishing the Union list of novel foods in accordance with Regulation (EU) 2015/2283 of the European Parliament and of the Council on novel foods (Text with EEA relevance)

[F1ANNEX

F2... LIST OF NOVEL FOODS

Textual Amendments

- Substituted by Commission Implementing Regulation (EU) 2018/1023 of 23 July 2018 correcting Implementing Regulation (EU) 2017/2470 establishing the Union list of novel foods (Text with EEA
- Word in Annex heading omitted (31.12.2020) by virtue of The Novel Food (Amendment) (EU Exit) Regulations 2019 (S.I. 2019/702), regs. 1, 64(a); 2020 c. 1, Sch. 5 para. 1(1)

Content of the list

1. The F3... list shall consist of Tables 1 and 2.

Textual Amendments

Word in Annex para. 1 omitted (31.12.2020) by virtue of The Novel Food (Amendment) (EU Exit) Regulations 2019 (S.I. 2019/702), regs. 1, **64(b)**; 2020 c. 1, Sch. 5 para. 1(1)

2. Table 1 includes the authorised novel foods and contains the following information:

Column 1 : Authorised novel food

Conditions under which the novel food may be used. This column is Column 2

further subdivided into two: Specified food category and Maximum

Additional specific labelling requirements Column 3

Column 4 Other requirements

3. Table 2 includes the specifications on novel foods and contains the following information:

Authorised novel food Column 1

Column 2 **Specifications**

TABLE 1: AUTHORISED NOVEL FOODS

Authorised novel food	Conditions under which the novel food may be used		Additional specific labelling requirements	Other requirements	[^{F8} Data Protection]
N - Acetyl-D- neuraminic acid	Specified food category Infant and follow-on formulae as defined by Regulation (EU) No 609/2013 a	Maximum levels 0,05 g/L of reconstituted formula	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'N-acetyl-D-neuraminic acid'		

Processed cereal-based foods and baby foods for infants and young children as defined by Regulation (EU) No 609/2013	0,05 g/kg for solid foods	Food supplements containing N -acetyl-D-neuraminic acid shall bear a statement that the food supplement should not be given
Foods for special medical purposes for infants and young children as defined by Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the infants and young children for whom the products are intended but in any case not higher than the maximum levels specified for the category mentioned in the table corresponding to the products.	to infants, young children and children under 10 years of age where they consume breast milk or other foods with added N-acetyl-D-neuraminic acid within the same twenty four hour period.
Total diet replacement foods for weight control as defined by Regulation (EU) No 609/2013	0,2 g/L (drinks) 1,7 g/kg (bars)	
Foods bearing statements on the absence or reduced presence of gluten in accordance with the	1,25 g/kg	

requirements of Commission Implementing Regulation (EU) No 828/2014 b	
Unflavoured pasteurised and sterilised (including UHT) milk-based products	0,05 g/L
Unflavoured fermented milk-based products, heat treated after fermentation, flavoured fermented milk products including heat-treated products	0,05 g/L (beverages) 0,4 g/kg (solids)
Dairy analogues, including beverage whiteners	0,05 g/L (beverages) 0,25 g/kg (solids)
Cereal bars	0,5 g/kg
Table top sweeteners	8,3 g/kg
Fruit and vegetable-based drinks	0,05 g/L
Flavoured drinks	0,05 g/L
Speciality coffee, tea, herbal and fruit infusions, chicory; tea, herbal and fruit infusions and chicory	0,2 g/kg

	extracts; tea, plant, fruit and cereal preparations for infusions Food Supplements as defined in Directive 2002/46/EC c	300 mg/day for general population older than 10 years 55 mg/day for infants 130 mg/day for young children 250 mg/day for children between 3 to 10 years of age		
Adansonia digitata (Baobab) dried fruit pulp	Not specified		The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Baobab fruit pulp '	
Ajuga reptans extract from cell cultures	Specified food category Food Supplements	Maximum levels In line with normal use		
	as defined in Directive 2002/46/EC	in food supplements of a similar extract of the flowering aerial parts of <i>Ajuga reptans</i>		
L-Alanyl-L- Glutamine	Specified food category	Maximum levels		
	Food Supplements as defined in Directive 2002/46/EC			

Algal oil	Foods for special medical purposes as defined in Regulation (EU) No 609/2013 excluding foods for infants and young children Drinks intended to meet the expenditure of intense muscular effort especially for sportsmen Specified	Maximum	The			
from the microalgae	food category	levels of DHA	designation of the novel			
Ulkenia sp.	Bakery products (breads, rolls and sweet biscuits)	200 mg/100 g	food on the labelling of the foodstuffs containing it shall be	od on the belling of e foodstuffs ntaining shall be		
	Cereal bars	500 mg/100 g	micro-algae			
	Non- alcoholic beverages (including milk based beverages)	60 mg/100 ml	Ulkenia sp. '			
[F9 Allanblacki seed oil	a Specified food category	Maximum levels	The designation of the novel			
	Yellow fat spreads and cream based spreads	30 g/100 g	food on the labelling of the foodstuffs containing it shall be '	ood on the abelling of ne foodstuffs ontaining		
	Mixtures of vegetable oils (*) and milk (falling under	30 g/100 g	Allanblackia seed oil'			

	and o oils a in Par Anne Regu	pt olive oils olive pomace s defined rt VIII of x VII of lation (EU) 308/2013.]			
Aloe macroclada Baker leaf extract	Specified food category Food Supplements	Maximum levels In line with normal use			
	as defined in Directive 2002/46/EC	in food supplements of the similar gel derived from Aloe vera (L.) Burm.			
Antarctic Krill oil from Euphausia superba	Specified food category	Maximum levels of combined DHA and EPA	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Lipid extract from the crustacean Antarctic Krill (Euphausia superba)'		
	Dairy products except milk- based drinks	200 mg/100 g or for cheese products 600 mg/100 g			
	Dairy analogues except drinks	200 mg/100 g or for analogues to cheese products 600 mg/100 g			
	Non- alcoholic beverages Milk-based drinks Dairy analogue drinks	80 mg/100 ml			

Spreadable fat and dressings	600 mg/100 g
Cooking fats	360 mg/100 ml
Breakfast cereals	500 mg/100 g
Bakery products (breads, rolls and sweet biscuits)	200 mg/100 g
Nutrition bars/cereal bars	500 mg/100 g
Food Supplements as defined in Directive 2002/46/EC	3 000 mg/ day for the general population 450 mg/day for pregnant and lactating women
Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended
Total diet replacement for weight control as defined in Regulation (EU) No 609/2013 and meal replacements for weight control	250 mg/meal
Processed cereal-based food and baby food	200 mg/100 ml

	intended for infants and young children covered by Regulation (EU) No 609/2013 Foods intended to meet the expenditure of intense muscular effort, especially for sportsmen			
	Foods bearing statements on the absence or reduced presence of gluten in accordance with the requirements of Commission Implementing Regulation (EU) No 828/2014			
Antarctic Krill oil rich in phospholipids from Euphausia superba	Specified food category Dairy products	Maximum levels of combined DHA and EPA 200 mg/100 g or for cheese	The designation of the novel food on the labelling of the foodstuffs containing	
superou	except milk- based drinks	products 600 mg/100 g	it shall be 'Lipid extract	
	Dairy analogues except drinks	200 mg/100 g or for analogues to cheese products 600 mg/100 g	from the crustacean Antarctic Krill (Euphausia superba)'	
	Non- alcoholic beverages	80 mg/100 ml		

Milk-based drinks Dairy analogue drinks	
Spreadable fat and dressings	600 mg/100 g
Cooking fats	360 mg/100 ml
Breakfast cereals	500 mg/100 g
Bakery products (breads, rolls and sweet biscuits)	200 mg/100 g
Nutrition bars/cereal bars	500 mg/100 g
Food Supplements as defined in Directive 2002/46/EC	3 000 mg/ day for the general population 450 mg/day for pregnant and lactating women
Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended
Total diet replacement for weight control as defined in Regulation (EU) No 609/2013 and meal replacements	250 mg/meal

	for weight control			
	Processed cereal-based food and baby food intended for infants and young children covered by Regulation (EU) No 609/2013	200 mg/100 ml		
	Foods intended to meet the expenditure of intense muscular effort, especially for sportsmen			
	Foods bearing statements on the absence or reduced presence of gluten in accordance with the requirements of Commission Implementing Regulation (EU) No 828/2014			
Arachidonic acid-rich oil from	Specified food category	Maximum levels	The designation of the novel	
the fungus Mortierella alpina	Infant formula and follow-on formula as defined in Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013	food on the labelling of the foodstuffs containing it shall be 'Oil from Mortierella alpina' or 'Mortierella alpina oil'	

	Foods for special medical purposes for premature infants as defined in Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013		
Argan oil from Argania spinosa	Specified food category As seasonings Food Supplements as defined in Directive 2002/46/EC	Maximum levels Not specified In line with normal food use of vegetable oils	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Argan oil' and if used as seasoning 'Vegetable oil only for seasoning 'shall be mentioned on the label	
Astaxanthin- rich oleoresin from Haematococcu pluvialis algae	Specified food category Food Supplements as defined in Directive 2002/46/EC	Maximum levels 40-80 mg/day of oleoresin, resulting in ≤ 8 mg astaxanthin per day	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Astaxanthin'	
Basil seeds (Ocimum basilicum)	Specified food category Fruit juice and fruit/vegetable blend beverages	Maximum levels 3 g/200 ml for addition of whole basil seeds (Ocimum basilicum)		
[^{F4} Betaine	Specified food category Drink powders,	Maximum levels ^g 60 mg/100 g	The designation of the novel food on the labelling of	Authorised on 22 August 2019. This inclusion is based on

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	isotonic and energy drinks intended for sportsmen		the foodstuffs containing it shall be 'betaine'.
	Protein and cereal bars intended for sportsmen	500 mg/100 g	The labelling of foods containing betaine shall bear
	Meal replacements intended for sportsmen	20 mg/100 g	a statement that the foods should not be used if food
	Total diet replacement for weight control as defined under Regulation (EU) No 609/2013	500 mg/100 g (bar) 136 mg/100 g (soup) 188 mg/100 g (porridge) 60 mg/100 g (beverages)	supplements containing betaine are consumed the same day.
	Foods for Special Medical Purposes as defined under Regulation (EU) No 609/2013 for adults	400 mg/day]	

proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: DuPont Nutrition Biosciences ApS, Langebrogade Copenhagen K, DK-1411, Denmark. During the period of data protection, the novel food betaine is authorised for placing on the market within the Union only by DuPont Nutrition Biosciences ApS unless a subsequent applicant obtainsauthorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation

> (EU) 2015/2283

				or with the agreement of DuPont Nutrition Biosciences ApS, End date of the data protection: 22 August 2024.
Fermented black bean extract	Specified food category	Maximum levels	The designation of the novel	
	Food Supplements as defined in Directive 2002/46/EC	4,5 g/day	food on the labelling of the foodstuffs containing it shall be ' Fermented black bean (Soya) extract ' 'or ' Fermented Soya extract '	
Bovine lactoferrin	Specified food category	Maximum levels	The designation of the novel	
	Infant formula and follow-on formula as defined in Regulation (EU) No 609/2013 (ready to drink)	100 mg/100 ml	food on the labelling of the foodstuffs containing it shall be 'Lactoferrin from cows' milk'	
	Foods on dairy basis intended for young children (ready to eat/ drink)	200 mg/100 g		
	Processed cereal food (solid)	670 mg/100 g		
	Foods for special medical	Depending on the needs of		

	purposes as defined in Regulation (EU) No 609/2013	the individual up to 3 g/day		
	Beverages based on milk	200 mg/100 g		
	Powdered drink mixes based on milk (ready to drink)	330 mg/100 g		
	Beverages based on fermented milk (including yoghurt drinks)	50 mg/100 g		
	Non- alcoholic drinks	120 mg/100 g		
	Products based on yoghurt	80 mg/100 g		
	Products based on cheese	2 000 mg/100 g		
	Ice cream	130 mg/100 g		
	Cakes and pastries	1 000 mg/100 g		
	Candies	750 mg/100 g		
	Chewing gum	3 000 mg/100 g		
[F10Bovine milk basic whey protein	Specified food category	Maximum levels	The designation of the novel	Authorised on 20 November 2018 This
isolate	Infant formulae as defined in Regulation (EU) No 609/2013 Follow-on formulae as defined in	30 mg/100 g (powder) 3,9 mg/100 mL (reconstituted) 30 mg/100 g (powder)	food on the labelling of the foodstuffs containing it shall be 'Milk whey protein isolate'. Food supplements containing	2018 . This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article

Regulation (EU) No 609/2013 Total diet replacement foods for weight control as defined by Regulation (EU) No 609/2013 Foods for special medical purposes as defined in Regulation (EU) No 609/2013 Food **Supplements** as defined in Directive 2002/46/EC

4,2 mg/100 mL(reconstituted) 300 mg/day 30 mg/100 g (powder formula for infants during the first months of life until the introduction of appropriate | adolescents complementar under the age feeding) 3,9 mg/100 mL (reconstituted formula for infants during the first months of life until the introduction of appropriate complementary feeding) 30 mg/100g (powder formula for infants when appropriate complementary feeding is introduced) 4.2 mg/100mL (reconstituted formula for infants when appropriate complementary feeding is

introduced)

58 mg/day for young

children

380 mg/

day for

children and

adolescents from 3 to 18 years of age

bovine milk basic whey protein isolate shall bear the following statement: 'This food supplement should not be consumed by infants/ children/ of one/three/ eighteen (*) years' (*) Depending on the age group the food supplement is intended for.

26 of Regulation (EU) 2015/2283. Applicant: Armor Protéines S.A.S., 19 bis, rue de la Libération 35460 Saint-Brice-en-Coglès, France. During the period of data protection the novel food bovine milk basic whey protein isolate is authorised for placing on the market within the Union only by Armor Protéines S.A.S. unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement

		610 mg/day for adults 25 mg/day for infants 58 mg/day for young children 250 mg/ day for children and adolescents from 3 to 18 years of age 610 mg/day for adults]		of Armor Protéines S.A.S. End date of the data protection: 20 November 2023 .
Buglossoides arvensis seed oil	Specified food category	Maximum levels of stearidonic acid (STA)	The designation of the novel food on the	
	Dairy	250 mg/100 g	labelling of	
	products and analogues	75 mg/100 g for drinks	the foodstuffs containing it shall be 'Refined Buglossoides oil'	
	Cheese and cheese products	750 mg/100 g		
	Butter and other fat and oil emulsions including spreads (not for cooking or frying purposes)	750 mg/100 g		
	Breakfast cereals	625 mg/100 g		
	Food supplements as defined in Directive 2002/46/EC, excluding food supplements for infants and young children	500 mg/day		
	Foods for special medical	In accordance with the particular		

	purposes as defined in Regulation (EU) No 609/2013, excluding foods for special medical purposes intended for infants and young children	nutritional requirements of the persons for whom the products are intended		
	Total diet replacement for weight control as defined in Regulation (EU) No 609/2013 and meal replacements for weight control	250 mg/meal		
Calanus finmarchicus oil	Food supplements as defined in Directive	Maximum levels 2,3 g/day	The designation of the novel food on the labelling of the foodstuffs	
	2002/46/EC		containing it shall be 'oil from Calanus finmarchicus (crustacean)'	

			homopolymer, maleated, esters with polyethylene glycol mono- Me ether) ' or ' Gum base (including CAS No: 1246080-53-4)	
Chewing gum base (Methyl	Specified food category	Maximum levels	The designation of the novel	
vinyl ether- maleic anhydride copolymer)	Chewing gum	2 %	food on the labelling of the foodstuffs containing it shall be 'Gum base (including methyl vinyl ether-maleic anhydride copolymer) 'or 'Gum base (including CAS No 9011-16-9) '	
Chia oil from Salvia hispanica	Specified food category	Maximum levels	The designation of the novel	
pc.u	Fats and oils	10 %	food on the labelling of the foodstuffs containing it shall be 'Chia oil (Salvia hispanica)'	
	Pure chia oil	2 g/day		
	Food Supplements as defined in Directive 2002/46/EC	2 g/day		
[^{F11} Chia seeds (<i>Salvia</i>	Specified food category	Maximum levels	The designation of the novel	
hispanica)	Bread products	5 % (whole or ground chia seeds)	food on the labelling of the foodstuffs	
	Baked products	10 % whole chia seeds	containing it shall be 'Chia seeds	
	Breakfast cereals	10 % whole chia seeds	(Salvia hispanica)'	

Sterilised ready to eat meals based on cereal grains, pseudocereals grains and/or pulses Fruit, nut and seed mixes Pre-packaged Chia seed as such Confectionery (including chocolate and chocolate and chocolate products), excluding chewing gums Dairy products (including yoghurt) and analogues Edible ices Fruit and vegetables products (including fruit spreads, compotes with/without cereals, fruit-preparations to underlay or to be mixed with dairy products, fruit desserts, mixed fruits with coconut milk for a twin pot)
Pre-packaged Chia seed as such Confectionery (including chocolate and chocolate products), excluding chewing gums Dairy products (including yoghurt) and analogues Edible ices Fruit and vegetables products (including fruit spreads, compotes with/without cereals, fruit-preparations to underlay or to be mixed with dairy products, fruit desserts, mixed fruits with coconut milk for a
Chia seed as such Confectionery (including chocolate and chocolate products), excluding chewing gums Dairy products (including yoghurt) and analogues Edible ices Fruit and vegetables products (including fruit spreads, compotes with/without cereals, fruit-preparations to underlay or to be mixed with dairy products, fruit desserts, mixed fruits with coconut milk for a
(including chocolate and chocolate and chocolate products), excluding chewing gums Dairy products (including yoghurt) and analogues Edible ices Fruit and vegetables products (including fruit spreads, compotes with/without cereals, fruit-preparations to underlay or to be mixed with dairy products, fruit desserts, mixed fruits with coconut milk for a
products (including yoghurt) and analogues Edible ices Fruit and vegetables products (including fruit spreads, compotes with/without cereals, fruit- preparations to underlay or to be mixed with dairy products, fruit desserts, mixed fruits with coconut milk for a
Fruit and vegetables products (including fruit spreads, compotes with/without cereals, fruit-preparations to underlay or to be mixed with dairy products, fruit desserts, mixed fruits with coconut milk for a
vegetables products (including fruit spreads, compotes with/without cereals, fruit- preparations to underlay or to be mixed with dairy products, fruit desserts, mixed fruits with coconut milk for a
Non-

	fruit juice and fruit/ vegetable blend beverages) Puddings that do not require heat treatment at or above 120 °C in their manufacture, processing or preparation]			
Chitin- glucan from Aspergillus niger	Specified food category Food	Maximum levels 5 g/day	The designation of the novel food on the	
	Supplements as defined in Directive 2002/46/EC		labelling of the foodstuffs containing it shall be 'Chitin- glucan from Aspergillus niger'	
Chitin- glucan complex from Fomes fomentarius	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Chitinglucan from Fomes fomentarius'	
	Food Supplements as defined in Directive 2002/46/EC	5 g/day		
Chitosan extract from fungi (Agaricus bisporus ; Aspergillus niger)	Specified food category	Maximum levels	The designation of the novel	
	Food Supplements as defined in Directive 2002/46/EC	In line with normal use in food supplements of chitosan from crustaceans	food on the labelling of the foodstuffs containing it shall be 'Chitosan extract from Agaricus bisporus' or 'Chitosan	

			extract from Aspergillus niger'		
Chondroitin sulphate	Specified food category Food supplements	Maximum levels 1 200 mg/day	The designation of the novel food on the labelling of		
	as defined in Directive 2002/46/ EC for adult population, excluding pregnant and lactating women		the foodstuffs containing it shall be ' Chondroitin sulphate derived from microbial fermentation and sulphation '		
Chromium Picolinate	Specified food category	Maximum levels of total chromium	The designation of the novel food on the		
	Foods covered by Regulation (EU) No 609/2013	250 μg/day	labelling of the foodstuffs containing it shall be ' Chromium		
	Foods fortified in accordance with Regulation (EC) No 1925/2006 d		Picolinate '		
Cistus incanus L. Pandalis	Specified food category	Maximum levels	The designation of the novel		
herb	Herbal infusions	Intended daily intake: 3 g herbs/day (2 cups/day)	food on the labelling of the foodstuffs containing it shall be 'Cistus incanus L. Pandalis herb'		
Citicoline	Specified food category	Maximum levels	1. The desig of the	nation	

	as defined in Directive 2002/46/EC Foods for special medical purposes as defined in Regulation (EU) No 609/2013	250 mg per serving and a maximum daily consumption level of 1 000 mg	on the labelling of the foodstuffs containing it shall be 'Citicoline' 2. The labelling of foods containing citicoline shall bear a statement that the product is not intended to be consumed by children	
Clostridium butyricum	Specified food category Food Supplements as defined in Directive 2002/46/EC	Maximum levels 1,35 × 10 ⁸ CFU/day	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Clostridium butyricum MIYAIRI 588 (CBM 588)' or 'Clostridium butyricum (CBM 588)'	

[F8D-ribose	Specified food category	Maximum levels	The designation of the novel
	Cereal bars	0,20 g/100 g	food on the
	Fine bakery wares	0,31 g/100 g	labelling of the foodstuffs containing it
	Chocolate confectionery (excluding chocolate bars)	0,17 g/100 g	shall be 'D- ribose'. The labelling of foods containing D-ribose
	Milk- based drinks (excluding malts and shakes)	0,08 g/100 g	shall bear a statement that the foods should not be used if food
	Drinks intended to meet the expenditure of intense muscular effort especially for sportsmen, isotonic and energy drinks	0,80 g/100 g	supplements containing D-ribose are consumed the same day.
	Bars intended to meet the expenditure of intense muscular effort especially for sportsmen	3,3 g/100 g	
	Meal replacement for weight control (as drinks)'	0,13 g/100 g	
	Meal replacement for weight control (as bars)	3,30 g/100 g	
	Confectionery	0,20 g/100 g	

Authorised on 16 April 2019. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Bioenergy Life Science, Inc., 13840 Johnson St. NE, Minneapolis, Minnesota, 55304, USA. During the period of data protection, the novel food D-ribose is authorised for placing on the market within the Union only by Bioenergy Life Science, Inc. unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article

	Tea and infusions (in powder form to be reconstituted)	0,23 g/100 g]		26 of Regulation (EU) 2015/2283 or with the agreement of Bioenergy Life Science, Inc. End date of the data protection: 16 April 2024 (5 years).
Extract of defatted cocoa	Specified food category	Maximum levels	Consumers shall be instructed not	
powder	Nutrition bars	1 g/day and	to consume more than	
	Milk based beverages	300 mg polyphenols corresponding	600 mg polyphenols	
	Any other foods (including food supplements as defined in Directive 2002/46/ EC) which have become established vehicles for functional ingredients and which are typically positioned for consumption by health conscious adults	to not more than 550 mg of extract of defatted cocoa powder in one portion of food (or food supplement)	corresponding to 1,1 g of extract of defatted cocoa powder per day	
Low fat cocoa extract	Specified food category	Maximum levels	Consumers shall be instructed not	
	Foods including food supplements as defined	730 mg per serving and around 1,2 g/ day	to consume more than 600 mg of cocoa	

	in Directive 2002/46/EC		flavanols per day	
Coriander seed oil from Coriandrum	Specified food category	Maximum levels	The designation of the novel	
sativum	Food Supplements as defined in Directive 2002/46/EC	600 mg/day	food on the labelling of the foodstuffs containing it shall be ' Coriander seed oil '	
[F12Cranberry extract powder	Specified food category	Maximum levels	The designation of the novel	Authorised on 20 November 2018. This
powder	Food Supplements as defined in Directive 2002/46/EC for the adult population	350 mg/day]	food on the labelling of the foodstuffs containing it shall be 'cranberry extract powder'	inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Ocean Spray Cranberries Inc. One Ocean Spray Drive Lakeville- Middleboro, MA, 02349, USA. During the period of data protection the novel food, cranberry extract powder, is authorised for placing on the market within the Union only by Ocean Spray Cranberries

				Inc. unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Ocean Spray Cranberries Inc. End date of the data protection: 20 November 2023.
Crataegus pinnatifida dried fruit	Specified food category	Maximum levels	The designation of the novel	
	Herbal infusions	In line with normal	food on the labelling of	
	Jams and jellies in accordance with Directive 2001/113/EC	food use of Crataegus laevigata	the foodstuffs containing it shall be ' Crataegus pinnatifida dried fruit'	
α-	Not specified		The	
cyclodextrin			designation of the novel food on the labelling of the foodstuffs containing	

γ- cyclodextrin	Not specified		it shall be 'Alpha-cyclodextrin' or 'α-cyclodextrin'. The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Gamma-Cyclodextrin' or 'γ-Cyclodextrin'.	
[F13]Decorticate grains of Digitaria exilis (Kippist) Stapf (Traditional food from a third country)	dNot specified		The designation of the novel food on the labelling of the foodstuffs containing it shall be 'decorticated fonio (Digitaria exilis) grains']	
Dextran preparation	Specified food	Maximum levels	The designation	
produced by Leuconostoc mesenteroides	nostoc Bakery 5 % food on the			
Diacylglycerol oil of plant	food	Maximum levels	The designation	
origin	Cooking oils		of the novel food on the	
	Fat spreads		labelling of the foodstuffs containing it shall be '	
	Salad dressings			
	Mayonnaise		Diacylglycerol oil of plant origin (at	

	Meal replacement for weight control (as drinks) Bakery products Yoghurt type products		least 80 diacylgly)	
Dihydrocapsia (DHC)	food category Cereal bars Biscuits, cookies and	Maximum levels 9 mg/100 g 9 mg/100 g	1.	The design of the novel food on	nation	
	Rice based snacks	12 mg/100 g		the labell of the foods conta it shall be	stuffs ining ydrocapsiate' lements ining letic drocapsiate	
	Carbonated drinks, dilutable drinks, fruit juice based beverages	1,5 mg/100 ml				
	Vegetable drinks	2 mg/100 ml	2.	Food suppl		
	Coffee based drinks, tea based drinks	1,5 mg/100 ml		synth		
	Flavoured water — still	1 mg/100 ml		be labell as		
	Precooked oatmeal cereal	2,5 mg/100 g		'not intend for		
	Other cereals	4,5 mg/100 g		childi up	en	
	Ice cream, dairy desserts	4 mg/100 g		to 4.5	,	
	Pudding mixes (ready to eat)	2 mg/100 g		years	5	
	Products based on yoghurt	2 mg/100 g				

	Chocolate confectionery	7,5 mg/100 g		
	Hard candy	27 mg/100 g		
	Sugar-free gum	115 mg/100 g		
	Whitener/ creamer	40 mg/100 g		
	Sweeteners	200 mg/100 g		
	Soup (ready to eat)	1,1 mg/100 g		
	Salad dressing	16 mg/100 g		
	Vegetable protein	5 mg/100 g		
	Ready to eat meals	3 mg/meal		
	Meal replacements for weight control	3 mg/meal		
	Meal replacement for weight control (as drinks)	1 mg/100 ml		
	Food Supplements as defined in Directive 2002/46/EC	3 mg/single intake 9 mg/day		
	Non- alcoholic powdered drink mixes	14,5 mg/kg equivalent to 1,5 mg/100 ml		
F14Dried nerial parts of <i>Hoodia</i>	Specified food category	Maximum levels	The designation of the novel	Authorised on 3 September 2018. This
parviflora	Food Supplements as defined in Directive 2002/46/ EC for adult population	9,4 mg/day]	food on the labelling of the foodstuffs containing it shall be 'dried aerial parts of Hoodia parviflora'	inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article

ANNEX

Document Generated: 2024-04-16

Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

26 of Regulation (EU) 2015/2283. Applicant: Desert Labs, Ltd Kibbutz Yotvata, 88820 Israel. During the period of data protection the novel food dried aerial parts of Hoodia parviflora is authorised for placing on the market within the Union only by Desert Labs, Ltd unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Desert Labs, Ltd. End date of the data protection: 3 September 2023.

Dried extract of Lippia citriodora from cell cultures	Specified food category Food Supplements as defined in Directive 2002/46/EC	In line with normal use in food supplements of a similar extract from the leaves of Lippia citriodora	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'dried extract of <i>Lippia citriodora</i> from cell cultures HTN ® Vb'	
Echinacea angustifolia extract from cell cultures	Specified food category Food Supplements as defined in Directive 2002/46/EC	In line with normal use in food supplements of a similar extract from the root of Echinacea angustifolia		
[F15 Echinacea purpurea extract from cell cultures	Specified food category Food Supplements as defined in Directive 2002/46/EC	In line with normal use in food supplements of a similar extract from florets within the flower head of Echinacea purpurea]	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'dried extract of <i>Echinacea purpurea</i> from cell cultures EchiPure-PCTM'	
Echium plantagineum oil	Specified food category Milk-based products and drinkable yoghurt products	Maximum levels of stearidonic acid (STA) 250 mg/100 g; 75 mg/100 g for drinks	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Refined echium oil'	

	delivered in a single dose Cheese preparations Spreadable fat and	750 mg/100 g 750 mg/100 g			
	dressings Breakfast cereals	625 mg/100 g			
	Food supplements as defined in Directive 2002/46/EC	500 mg/day			
	Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended			
	Total diet replacement for weight control as defined in Regulation (EU) No 609/2013 and meal replacements for weight control	250 mg/meal			
[F16 Ecklonia cava phlorotannins	Food supplements as defined in Directive 2002/46/EC intended for the general population, excluding children	Maximum levels 163 mg/ day for adolescents from 12 to 14 years of age 230 mg/ day for adolescents above 14 years of age	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Ecklonia cava Phlorotannins' Food supplements containing		

under the age of 12 years	263 mg/day for adults]	Ecklonia phlorotar shall bea following statemen (a)	nnins r the g
		(b)	years. This food supplement should not be consumed by persons with thyroid disease or by persons who are aware of or have been identified as being at risk of developing

			dis (c) The food sup sho not be con if oth food sup con iod are also con (*) De on the age gro the food sup con iod are also con the age gro the food sup is	d oplement ould oplement ould oplements of oplements of oplements of oplements of oplement op	
[^{F17} Egg membrane	Specified food	Maximum levels	The designation		Authorised on 25 November
hydrolysate	Food Supplements as defined in Directive 2002/46/ EC intended for the general adult population	450 mg/day]	of the novel food on the labelling of the foodstuff containing it shall be 'egg membrane hydrolysate'.		2018. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Biova, LLC., 5800 Merle Hay Rd, Suite 14 PO Box 394 Johnston 50131,

				Iowa USA. During the period of data protection the novel food egg membrane hydrolysate is authorised for placing on the market within the Union only by Biova, LLC. unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Biova, LLC. End date of the data protection: 25 November 2023
Epigallocatech gallate as	in pecified food category	Maximum levels	The labelling shall bear a	
	Foods	150 mg of	statement that consumers	
	including food	extract in one portion of	should not consume	
	supplements	food or food	more than	
sinensis)	as defined	supplement	300 mg of	
	in Directive 2002/46/EC		extract per day	
	7111177/76/17			

[^{F18} L- ergothioneine	Specified food category	Maximum levels	The designation	
	Alcohol-free beverages	0,025 g/kg	of the novel food on the labelling of	
	drinks containing it	the foodstuffs containing it		
	' Fresh ' milk products(*)	0,040 g/kg	shall be 'L- ergothioneine	
	Cereal bars	0,2 g/kg		
	Chocolate confectionery	0,25 g/kg		
	Food supplements as defined in Directive 2002/46/EC	30 mg/day for general population (excluding pregnant and lactating women) 20 mg/day for children older than 3 years		
	(*) When used in milk products L-ergothioneine may not replace in whole or in part, any milk constituent]			
[F16Extract of three herbal	Specified food category	Maximum levels	The designation	
roots (Cynanchum wilfordii Hemsley, Phlomis umbrosa Turcz. and Angelica gigas Nakai)	Food supplements as defined in Directive 2002/46/ EC for adult population	175 mg/day]	of the novel food on the labelling of the foodstuffs containing it shall be 'extract of three herbal roots (Cynanchum wilfordii Hemsley, Phlomis umbrosa Turcz. and Angelica gigas Nakai)'.	

			The labelling of food supplements containing the extract of mixture of the three herbal roots shall bear a statement in close proximity to the list of ingredients indicating that it should not be consumed by individuals with known celery allergy.	
Ferric Sodium	Specified food	Maximum levels	The designation	
EDTA	category	(expressed as	of the novel food on the	
		anhydrous EDTA)	labelling of the foodstuffs	
	Food supplements as defined in Directive 2002/46/EC	18 mg/day for children 75 mg/day for adults	containing it shall be 'Ferric Sodium EDTA'	
	Foods covered by Regulation (EU) No 609/2013	12 mg/100 g		
	Foods fortified in accordance with Regulation (EC) No 1925/2006			
Ferrous ammonium	Specified food	Maximum levels	The designation	
phosphate	Food supplements as defined	To be used in compliance with	of the novel food on the labelling of the foodstuffs containing	

	in Directive 2002/46/EC Foods covered by Regulation (EU) No 609/2013 Foods fortified in accordance with Regulation (EC) No 1925/2006	Directive 2002/46/EC, Regulation (EU) No 609/2013 and/or Regulation (EC) No 1925/2006	it shall be 'Ferrous ammonium phosphate'		
Fish peptides from Sardinops	Specified food category	Maximum levels fish peptide product	The designation of the novel food on the		
sagax	Foods based on yoghurt, yoghurt drinks, fermented milk products, and powdered milk	0,48 g/100 g (ready to eat/ drink)	labelling of the foodstuffs containing it shall be 'Fish (Sardinops sagax) peptides'		
	Flavoured water, and vegetable- based drinks	0,3 g/100 g (ready to drink)			
	Breakfast cereals	2 g/100 g			
	Soups, stews and soup powders	0,3 g/100 g (ready to eat)			
Flavonoids from Glycyrrhiza glabra	Specified food category	Maximum levels of flavonoids from Glycyrrhiza glabra	of the	Beverages national ining flavonoids shall be presented to the final	
	Beverages based on milk	120 mg/day	on the	consumer as single i ng rtions.	
	Beverages based on yoghurt		of the foods		

Beverages			it	
based on fruit			shall	
or vegetables			be	
or vegetables			'Flav	onoids
Food	120 mg/day		from	
Supplements				rrhiza
as defined			glabr	
in Directive			L.	
2002/46/EC		2.	The	
		۷.	labell	ing
Total diet	120 mg/day		of	mg
replacement			the	
for weight				
control as			foods	
defined in			where	
Regulation			the	
(EU) No			produ	ict
609/2013			was	
F 1 C	120 /1		addec	1
Foods for	120 mg/day		as a	
special			novel	
medical			food	
purposes as			ingre	dient
defined in			shall	
Regulation			bear	
(EU) No			a	
609/2013			stater	nent
			that:	
			(a)	the
			` ′	product
				should
				not
				be
				consumed
				by
				pregnant
				and
				breast
				feeding
				women,
				children
				and
				young
				adolescents;
			(1.)	and
			(b)	people
				taking
				prescription
				drugs
				should
				only
				consume
				the

			3.	The amou of flavor in the final food shall be indica on the labell of the food conta it.	noids ated ing	
[F19Fruit pulp, pulp juice, concentrated pulp juice from Theobroma cacao L. (Traditional food from a third country)	Not specified	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'cocoa (Theobroma cacao L.) pulp', 'cocoa (Theobroma cacao L.) pulp juice' or 'cocoa (Theobroma				

Fucoidan extract from the seaweed Fucus	Specified food category Foods	cacao L.) concentrated pulp juice' depending on the form used.] Maximum levels	The designation of the novel food on the		
vesiculosus	including food the food supplements as defined in Directive 2002/46/ EC for the general labelling the food containing it shall be seawed Fucus	labelling of the foodstuffs containing it shall be 'Fucoidan extract from seaweed	labelling of the foodstuffs containing it shall be 'Fucoidan extract from seaweed Fucus		
Fucoidan extract from the seaweed	Specified food category	Maximum levels	The designation		
Undaria pinnatifida	Foods including food supplements as defined in Directive 2002/46/ EC for the general population	250 mg/day	of the novel food on the labelling of the foodstuffs containing it shall be 'Fucoidan extract from seaweed Undaria pinnatifida'		
2'- Fucosyllactoso	Specified food category	Maximum levels	1. The design of	nation	
	Unflavoured pasteurised and sterilised (including UHT) milk-based products	1,2 g/l	the novel food on the labell of		
	Unflavoured fermented	1,2 g/l beverages	the foods conta		
	milk-based products 19,2 g/kg products other than beverages	it shall be '2'- fucosyllactose'.			

Flavoured fermented milk-based products including heat-treated products Dairy analogues, including beverage whiteners	1,2 g/l beverages 19,2 g/kg products other than beverages 1,2 g/l beverages 12 g/kg for products other than beverages	2.	The labelling of food supplements containing 2'-fucosyllactose shall bear a statement that the supplements
	400 g/kg for whitener		should not
Cereal bars Table-top sweeteners	12 g/kg 200 g/kg		be used if other
Infant formula as defined in Regulation (EU) No 609/2013	1,2 g/l alone or in combination with up to 0,6 g/l of lacto- N - neotetraose at a ratio of 2:1 in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer	3.	foods with added 2'- fucosyllactose are consumed the same day. The labelling of food supplements containing 2'- fucosyllactose
Follow-on formula as defined in Regulation (EU) No 609/2013	1,2 g/l alone or in combination with up to 0,6 g/l of lacto- N - neotetraose at a ratio of 2:1 in the final product ready for use, marketed as such or reconstituted		intended for young children shall bear a statement that the supplements should not

Processed cereal-based food and baby food for infants and young children as defined in Regulation (EU) No 609/2013	as instructed by the manufacturer 12 g/kg for products other than beverages 1,2 g/l for liquid food ready for use, marketed as such or reconstituted as instructed by the manufacturer	be used if breast milk or other foods with added 2'-fucosy are consuthe same day.	yllactose
Milk-based drinks and similar products intended for young children	1,2 g/l for milk-based drinks and similar products added alone or in combination with up to 0,6 g/l lacto- N -neotetraose, at a ratio of 2:1 in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer		
Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended		
Total diet replacement for weight control as	4,8 g/l for drinks 40 g/kg for bars		

defined in	
Regulation (EU) No	
Bread and pasta products bearing statements on the absence or reduced presence of gluten in accordance with the requirements of Commission Implementing Regulation (EU) No 828/2014	60 g/kg
Flavoured drinks	1,2 g/l
Coffee, tea (excluding black tea), herbal and fruit infusions, chicory; tea, herbal and fruit infusions and chicory extracts; tea, plant, fruit and cereal preparations for infusions, as well as mixes and instant mixes of these products	9,6 g/l — the maximum level refers to the products ready to use
Food supplements as defined in Directive 2002/46/EC, excluding food	3,0 g/day for general population 1,2 g/day for young children

	supplements for infants				
[F202'- Fucosyllactose Difucosyllacto	Specified food seategory	Maximum levels	The designation of the novel		Authorised on 19.12.2019. This inclusion
mixture ('2'-FL/DFL ') (microbial source)	Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products	2,0 g/L	Difucosyllactor mixture shall bear a statement that they should not be used if breast milk or other foods containing added 2'- Fucosyllactose and/or	proprietary scientific evidence an scientific da protected in accordance with Article	scientific evidence and scientific data protected in
	Unflavoured fermented milk-based products	2,0 g/L (beverages) 20 g/kg (products other than beverages)		supplements containing the 2'- Fucosyllactose/ Difucosyllactose	Regulation (EU) 2015/2283. Applicant: Glycom A/ S, Kogle Allé 4, DK-2970 Hørsholm, Denmark. During the period of data protection, the novel food 2'-
	Flavoured fermented milk-based products including heat-treated products	2,0 g/L (beverages) 20 g/kg (products other than beverages)			
	Beverages (flavoured drinks)	2,0 g/L			Fucosyllactose/ Difucosyllactose mixture is
	Cereal bars	20 g/kg	Difucosyllactor are consumed	se	authorised for placing
	Infant formula as defined under Regulation (EU) No 609/2013	1,6 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer	the same day.		on the market within the Union only by Glycom A/S, unless a subsequent applicant obtains authorisation
	Follow-on formula as defined under Regulation (EU) No 609/2013	1,2 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			food refere to the propr scient evide scient protes

Processed cereal-based food and baby food for infants and young children as defined under Regulation (EU) No 609/2013	1,2 g/L (beverages) in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer 10 g/kg for products other than beverages
Total diet replacement foods for weight control as defined under Regulation (EU) No 609/2013	4,0 g/L (beverages) 40 g/kg (products other than beverages)
Food for special medical purposes as defined under Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended
Food Supplements as defined in Directive 2002/46/EC intended for the general population excluding infants	4,0 g/day]
[F21Milk-based drinks and similar products intended	I ^{F22} 1.2 g/L in the final product ready for use, marketed as such or

with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Glycom A/S. End date of the data protection: 19.12.2024.

ı		ma a a m a t : t = t = .1	I	l	1
	for young children	reconstituted as instructed			
		by the			
		manufacturer]			
Galacto- oligosacchario	category	Maximum levels (expressed as ratio kg galacto- oligosacchari kg final food)	de/		
	Food Supplements as defined in Directive 2002/46/EC	0,333			
	Milk	0,02			
	Milk drinks	0,03			
	Meal replacement for weight control (as drinks)	0,02			
	Dairy analogue drinks	0,02			
	Yoghurt	0,033			
	Dairy based deserts	0,043			
	Frozen dairy deserts	0,043			
	Fruit drinks and energy drinks	0,021			
	Infant meal replacement drinks	0,012			
	Baby juice	0,025			
	Baby yogurt drink	0,024			
	Baby desert	0,027			
	Baby snack	0,143			
	Baby cereals	0,027			

	Drinks intended to meet the expenditure of intense muscular effort especially for sportsmen Juice Fruit pie fillings	0,013 0,021 0,059
	Fruit preparations	0,125
	Bars	0,125
	Cereals	0,125
	Infant formula and follow-on formula as defined in Regulation (EU) No 609/2013	0,008
Glucosamine HCl	Specified food category	Maximum levels
	Food Supplements as defined in Directive 2002/46/EC	In line with normal food use of glucosamine from shell
	Foods covered by Regulation (EU) No 609/2013	fish
	Meal replacement for weight	
	control Foods	_

	of intense muscular effort, especially for sportsmen Foods bearing statements on the absence or reduced presence of gluten in accordance with the requirements of Commission Implementing Regulation (EU) No 828/2014				
Glucosamine sulphate KCl	Specified food	Maximum levels			
	category				
	Food Supplements as defined in Directive 2002/46/EC	In line with normal food use of glucosamine from shell fish			
Glucosamine sulphate NaCl	Specified food category	Maximum levels			
	Food Supplements as defined in Directive 2002/46/EC	In line with normal food use of glucosamine from shell fish			
Guar Gum	Specified food category	Maximum levels	The design	nation	
	Fresh dairy products such as yogurts, fermented milks, fresh cheeses and other dairy-based desserts.	1,5 g/100 g	the novel food on the labell of the foods	ing	

Fruit or vegetable- based liquid foodstuffs (of the ' smoothie ' variety)	1,8 g/100 g 3,25 g/100 g	2.	containing it shall be 'Guar Gum'. A specific mention
vegetable- based compotes			of the possible
Cereals accompanied by a dairy product, in packaging containing two compartments	10 g/100 g in the cereals None in the accompanying dairy product 1 g/100 g in the product when ready to eat		risks of digestive discomfort linked to the exposure of children aged under 8 to guar gum must be visible on the label of any foodstuffs containing it. For example, 'Excessive consumption of these products may cause digestive discomfort, especially for children under

8 years of age'. 3. In the case of products with two compartments containing dairy and cereal products respectively, the instructions for use must clearly specify the need to mix the cereal and the dairy products respectively. The containing dairy and cereal products respectively, the instructions for use must clearly specify the cereal and the dairy productbefore consumption, in order to take into account the potential risk of gastro-intestinal obstruction. Heat-treated milk Specified Maximum levels Ma						
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of age*. 3. In the case of products with two compartments containing dairy and cereal products respectively, the instructions for use must clearly specify the need to mix the cereal and the dairy productbefore consumption, in order to take into account the potential risk of gastro-intestinal obstruction. Heat- Specified Maximum					years	
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11000					oosu uction.	
treated milk food levels		Specified				
	treated milk	food	levels			
products <i>category</i>	products	category				

fermented with Bacteroides xylanisolvens	Fermented milk products (in liquid, semi-liquid and spray- dried powder forms)				
Hydroxytyros	forms)	Maximum levels 0,215 g/kg 0,175 g/kg	The designation of the novel food on the labelling of the food products containing it shall be 'hydroxytyroso The labelling of the food products containing hydroxytyroso shall bear the following statements: (a) This food products consumption of the food products containing hydroxytyroso shall bear the following statements: (a) This food products consumption of the food products containing hydroxytyroso shall bear the following statements: (b) This food products containing hydroxytyroso shall bear the following statements: (a) This food products containing hydroxytyroso shall bear the following statements: (b) This food products containing hydroxytyroso shall bear the following statements: (a) This food products containing hydroxytyroso shall bear the following statements: (b) This food products containing hydroxytyroso shall bear the following statements: (a) This food products consumption hydroxytyroso shall bear the following statements: (b) This food products containing hydroxytyroso shall bear the following statements: (b) This food products containing hydroxytyroso shall bear the following statements: (b) This food products containing hydroxytyroso shall bear the following statements: (c) This food products containing hydroxytyroso shall bear the following statements: (b) This food products containing hydroxytyroso shall bear the following statements:	ict d imed ren ten, ing	
			shoul not be used	d	

Ice Structuring Protein type III HPLC 12	Specified food category Edible ices	Maximum levels	for cooking, baking or frying' The designation of the novel food on the labelling of the foodstuffs containing it shall be ' Ice Structuring	
Aqueous extracts of dried	Specified food category	Maximum levels	Protein ' The designation of the novel	
leaves of Ilex guayusa	Herbal infusions	In line with normal use	food on the labelling of	
	Food Supplements as defined in Directive 2002/46/EC	in herbal infusions and food supplements of a similar aqueous extract of dried leaves of Ilex paraguariensis	the foodstuffs containing it shall be 'Extracts of dried leaves of <i>Ilex guayusa</i> '	
[^{F23} Infusion from coffee	Specified food	Maximum levels	The designation	
leaves of	<i>category</i> Herbal		of the novel food on the	
Coffea arabica	infusions]		labelling of	
L. and/ or <i>Coffea</i>			the foodstuffs containing	
canephora			it shall be 'Infusion	
Pierre ex A. Froehner			from coffee	
(Traditional food from			leaves of <i>Coffea</i>	
a third			arabica and/	
country)			or Coffea canephora '.	
Isomalto- oligosacchario	Specified lefood	Maximum levels	1. The designation	
JIISUSACCIIAI IC	category		of	
			the novel	
			food	

	Energy- Reduced Soft Drinks Energy	6,5 % 5,0 %		on the labell of the	stuffs aining maltooligosacchari ds aining	
	Foods intended to meet the expenditure of intense muscular efforts, especially for sportsmen (including isotonic drinks)	6,5 %	2.	conta it shall be 'Isom Food conta the novel		ride'.
	Processed Vegetables and Vegetable Juices	5 %		as 'a sourc of gluco		
	Other Soft Drinks	5 %	-			
	Cereals Bars	10 %				
	Cookies, Biscuits	20 %	_			
	Breakfast Cereal Bars	25 %				
	Hard Candies	97 %	-			
	Soft Candies/ Chocolate Bars	25 %				
	Meal replacement for weight control (as bars or milk based)	20 %				
Isomaltulose	Not specified		1.	The desig of the novel food on	nation	

			the labelling of the foodstuffs containing it shall be 'Isomaltulose'. 2. The designation of the novel food on the labelling shall be accompanied by indication that the 'Isomaltulose is a source of glucose and fructose'.
[F24Lactitol	Specified food category Food Supplements as defined in Directive 2002/46/EC (capsules, tablets or powder) intended for the adult population	Maximum levels 20 g/day]	The designation of the novel food on the labelling of the food supplements containing it shall be 'Lactitol'
Lacto- N - neotetraose	Specified food category	Maximum levels	1. The designation of the novel

Unflavoured pasteurised and sterilised (including UHT) milk-based products	0,6 g/l	food on the labelling of the foodstuffs containing
Unflavoured fermented milk-based products	0,6 g/l for beverages 9,6 g/kg for products other than beverages	it shall be 'lacto- N - neotetraose'.
Flavoured fermented milk-based products including heat-treated products	0,6 g/l for beverages 9,6 g/kg for products other than beverages	2. The labelling of food supplements containing lacto- N -
Dairy analogues, including beverage whiteners	0,6 g/l for beverages 6 g/kg for products other than beverages 200 g/kg for whitener	neotetraose shall bear a statement that the supplements
Cereal bars	6 g/kg	should not
Table-top sweeteners	100 g/kg	be used
Infant formula as defined in Regulation (EU) No 609/2013	0,6 g/l in combination with up to 1,2 g/l of 2'-fucosyllactose at a ratio of 1:2 in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer	if other foods with added lacto- N- neotetraose are consumed the same day. 3. The labelling of
Follow-on formula as defined in	0,6 g/l in combination with up to	food

Regulation (EU) No 609/2013	1,2 g/l of 2'- fucosyllactose at a ratio of 1:2 in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer	supplements containing lacto- N - neotetraose intended for young children shall bear a
Processed cereal-based food and baby food for infants and young children as defined in Regulation (EU) No 609/2013	6 g/kg for products other than beverages 0,6 g/l for liquid food ready for use, marketed as such or reconstituted as instructed by the manufacturer	statement that the supplements should not be used if breast milk or other foods
Milk-based drinks and similar products intended for young children	0,6 g/l for milk-based drinks and similar products added alone or in combination with 2'-fucosyllactose, at a ratio of 1:2 in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer	with added lacto- N - neotetraose are consumed the same day.
Foods for special medical purposes as defined in Regulation	In accordance with the particular nutritional requirements of the persons	

(EU) No 609/2013	for whom the products are intended
Total diet replacement for weight control as defined in Regulation (EU) No 609/2013	2,4 g/l for drinks 20 g/kg for bars
Bread and pasta products bearing statements on the absence or reduced presence of gluten in accordance with the requirements of Commission Implementing Regulation (EU) No 828/2014	30 g/kg
Flavoured drinks	0,6 g/l
Coffee, tea (excluding black tea), herbal and fruit infusions, chicory; tea, herbal and fruit infusions and chicory extracts; tea, plant, fruit and cereal preparations for infusions, as well as mixes and instant mixes	4,8 g/l — the maximum level refers to the products ready to use

	of these products Food supplements as defined in Directive 2002/46/EC, excluding food supplements for infants	1,5 g/day for general population 0,6 g/day for young children		
[F25Lacto- N-tetraose ('LNT') (microbial source)	Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products	Maximum levels 1,0 g/l	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'lacto- N-tetraose'.	Authorised on 23.4.2020. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance
	Unflavoured fermented milk-based products	1,0 g/l (beverages) 10 g/kg (products other than beverages)	The labelling of food supplements containing lacto- <i>N</i> -tetraose shall bear a statement that they should not be used if breast milk or other foods containing added lacto- <i>N</i> - tetraose are consumed the same day.	with Article 26 of Regulation (EU) 2015/2283. Applicant: Glycom A/
	Flavoured fermented milk-based products including heat-treated products	1,0 g/l (beverages) 10 g/kg (products other than beverages)		S, Kogle Allé 4, DK-2970 Hørsholm, Denmark. During the period of data protection, the novel
	Beverages (flavoured drinks)	1,0 g/l		food lacto- N- tetraose is authorised
	Cereal bars Infant formula as defined under Regulation (EU) No 609/2013	10 g/kg 0,8 g/l in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer		for placing on the market within the Union only by Glycom A/S, unless a subsequent applicant obtains authorisation for the novel food without reference

Follow-on formula as defined under Regulation (EU) No 609/2013	0,6 g/l in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer
Processed cereal-based food, baby food for infants and young children as defined under Regulation (EU) No 609/2013	0,6 g/l (beverages) in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer 5 g/kg for products other than beverages
Milk based drinks and similar products intended for young children	0,6 g/l (beverages) in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer 5 g/kg for products other than beverages
Total diet replacement foods for weight control as defined under Regulation (EU) No 609/2013	2,0 g/l (beverages) 20 g/kg (products other than beverages)
Food for special	In accordance

to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Glycom A/S. End date of the data protection: 23.4.2025.

	medical purposes as defined under Regulation (EU) No 609/2013	with the particular nutritional requirements of the persons for whom the products are intended		
	Food Supplements as defined in Directive 2002/46/EC, excluding infants	2,0 g/day for young children, children, adolescents, and adults]		
[F26]Lonicera caerulea L. berries (haskap) (Traditional food from a third country)	Not specified		The designation of the novel food on the labelling of the foodstuffs containing it shall be 'haskap (Lonicera caerulea) berries']	
Lucerne leaf extract from <i>Medicago</i>	Specified food category	Maximum levels	The designation of the novel	
sativa	Food supplements as defined in Directive 2002/46/EC	10 g/day	food on the labelling of the foodstuffs containing it shall be 'Lucerne (
			Medicago sativa) protein' or 'Alfalfa (Medicago sativa) protein'.	
Lycopene	Specified food category	Maximum levels	sativa) protein' or 'Alfalfa (Medicago sativa)	

(including concentrates)		it shall be 'Lycopene'	
Drinks intended to meet the expenditure of intense muscular effort especially for sportsmen	2,5 mg/100 g		
Total diet replacement for weight control as defined in Regulation (EU) No 609/2013 and meal replacements for weight control	8 mg/meal		
Breakfast cereals	5 mg/100 g		
Fats and dressings	10 mg/100 g		
Soups other than tomato soups	1 mg/100 g		
Bread (including crispy breads)	3 mg/100 g		
Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended		
Food supplements as defined in Directive 2002/46/EC	15 mg/day		

Lycopene from <i>Blakeslea</i>	Specified food category	Maximum levels	The designation of the novel		
trispora	Fruit/ vegetable juice-based drinks (including concentrates)	2,5 mg/100 g	food on the labelling of the foodstuffs containing it shall be 'Lycopene'	the foodstuffs containing it shall be '	
	Drinks intended to meet the expenditure of intense muscular effort especially for sportsmen	2,5 mg/100 g			
	Total diet replacement for weight control as defined in Regulation (EU) No 609/2013 and meal replacements for weight control	8 mg/meal			
	Breakfast cereals	5 mg/100 g			
	Fats and dressings	10 mg/100 g			
	Soups other than tomato soups	1 mg/100 g			
	Bread (including crispy breads)	3 mg/100 g			
	Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the			

Lycopene	Food supplements as defined in Directive 2002/46/EC	products are intended 15 mg/day Maximum	The		
from tomatoes	food category	levels	designation of the novel		
	Fruit/ vegetable juice-based drinks (including concentrates)	2,5 mg/100 g	food on the labelling of the foodstuffs containing it shall be ' Lycopene'	labelling of the foodstuffs containing it shall be '	
	Drinks intended to meet the expenditure of intense muscular effort especially for sportsmen	2,5 mg/100 g			
	Total diet replacement for weight control as defined in Regulation (EU) No 609/2013 and meal replacements for weight control	8 mg/meal			
	Breakfast cereals	5 mg/100 g			
	Fats and dressings	10 mg/100 g			
	Soups other than tomato soups	1 mg/100 g			
	Bread (including crispy breads)	3 mg/100 g			

	Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended		
	Food supplements as defined in Directive 2002/46/EC	15 mg/day		
Lycopene oleoresin from	Specified food category	Maximum levels of lycopene	The designation of the novel	
tomatoes	Fruit/ vegetable juice-based drinks (including concentrates)	2,5 mg/100 g	food on the labelling of the foodstuffs containing it shall be ' Lycopene	
	Drinks intended to meet the expenditure of intense muscular effort especially for sportsmen	2,5 mg/100 g	oleoresin from tomatoes '	
	Total diet replacement for weight control covered by Regulation (EU) No 609/2013 and meal replacements for weight control	8 mg/meal		
	Breakfast cereals	5 mg/100 g		
	Fats and dressings	10 mg/100 g		

[F16Hen egg white lysozyme hydrolysate	Soups other than tomato soups Bread (including crispy breads) Foods for special medical purposes as defined in Regulation (EU) No 609/2013 Specified food category Food supplements as defined in Directive 2002/46/ EC intended	1 mg/100 g 3 mg/100 g In accordance with the particular nutritional requirements of the persons for whom the products are intended Maximum levels 1000 mg/day]	The designation of the novel food on the labelling of food supplements containing it shall be 'Hen	
	for adult population	Maximum	egg white lysozyme hydrolysate	
Magnesium citrate malate	Specified food category Food Supplements as defined in Directive 2002/46/EC	levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Magnesium citrate malate'	
Magnolia Bark Extract	Specified food category Mints (confectionary products) Chewing gum	Maximum levels 0,2 % for breath freshening purposes. Based on a 0,2 % maximum incorporation	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Magnolia Bark Extract'	

		level and a maximum gum/mint size of 1,5 g each, each gum or mint serving will contain no more than 3 mg of magnolia bark extract.			
Maize-germ oil high in	Specified food	Maximum levels	The designation		
unsaponifiabl matter	Food Supplements as defined in Directive 2002/46/EC Chewing gum	2 g/day 2 %	of the novel food on the labelling of the foodstuffs containing it shall be ' Maize-germ oil extract'		
Methylcellulo	food	Maximum levels	The designation	Methylcellulos is not to be	e
	Edible ices	2 %	of the novel food on the	used in foods specially	
	Edible ices Flavoured drinks Flavoured or unflavoured fermented	2 /0	labelling of the foodstuffs containing it shall be ' Methylcellulos	prepared for young children	
	milk products Cold desserts (dairy, fat, fruit, cereal, egg-based products)				
	Fruit preparations (pulps, purees or compotes)				
	Soups and broths				
[F271- Methylnicotin chloride	Specified affilde category	Maximum levels	The designation of the novel		Authorised on 2 September 2018. This
- 2-	Food Supplements as defined	58 mg/day]	food on the labelling of the foodstuffs		inclusion is based on proprietary

in Directive containing it scientific 2002/46/EC shall be '1evidence and Methylnicotinamide for the adult scientific data chloride'. population protected in excluding Food accordance pregnant supplements with Article and lactating containing 1-26 of Methylnicotinamide Regulation women shall bear the (EU) following 2015/2283. statement: Applicant: This food Pharmena supplement SA, Wolczanska should be 178, 90 530 consumed by adults only Lodz, Poland. excluding During the period of data pregnant and lactating protection women thenovel food 1methylnicotinamide chloride is authorised for placing on the market within the Union only by Pharmena S.A. unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Pharmena

S.A.

					End date of the data protection: 2 September 2023
(6S)-5- methyltetrahy acid, glucosamine salt	Specified d f016 lic category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be '(6S)-5-methyltetrahydacid, glucosamine salt 'or '5MTHF-glucosamine'	rofolic	
	Food Supplements as defined in Directive 2002/46/EC as a source of folate				
Monomethyls (Organic Silicon)	i Spacif iyd food category	Maximum levels of silicon	The designation of the novel		
Sincony	Food Supplements as defined in Directive 2002/46/ EC for adult population (in liquid form)	10,40 mg/day	food on the labelling of the food supplements containing it shall be 'Organic silicon (monomethylsi,'	lanetriol)	
Mycelial extract from	Specified food category	Maximum levels	The designation of the novel		
Shiitake mushroom (Lentinula	Bread products	2 ml/100 g	food on the labelling of		
edodes)	Soft drinks	0,5 ml/100 ml	the foodstuffs containing		
	Ready prepared meals	2,5 ml per meal	it shall be 'extract from the mushroom		

	Foods based on yoghurt Food supplements as defined in Directive 2002/46/EC	1,5 ml/100 ml 2,5 ml per day dose	Lentinula edodes' or 'extract from Shiitake mushroom'	
[F28Nicotinami riboside chloride	Food Supplements as defined in Directive 2002/46/EC	Maximum levels 300 mg/day for the general adult population, excluding pregnant and lactating women 230 mg/day for pregnant and lactating women]	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Nicotinamide riboside chloride'	Authorised on 20 February 2020. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: ChromaDex Inc., 10900 Wilshire Boulevard Suite 600, Los Angeles, CA 90024 USA. During the period of data protection, the novel food is authorised for placing on the market within the Union only by ChromaDex Inc. unless a subsequent applicant obtains authorisation for that novel food without reference

				to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of ChromaDex Inc. End date of the data protection: 20 February 2025.
Noni fruit juice (Morinda	Specified food category	Maximum levels	The designation of the novel	
citrifolia)	Pasteurised fruit and fruit nectar based drinks	30 ml with one serving (up to 100 % noni juice) or 20 ml twice a day, not more than 40 ml per day	food on the labelling of the foodstuffs containing it shall be 'Noni juice' or 'Juice of Morinda citrifolia'	
Noni fruit juice powder (Morinda citrifolia)	Food supplements as defined in Directive 2002/46/EC	6,6 g/day (equivalent to 30 ml of noni juice)	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Noni juice powder' or 'Juice powder of Morinda citrifolia'	
Noni fruit puree and concentrate	Specified food category	Maximum levels	The designation of the novel	
(Morinda citrifolia)		Fruit puree	food on the labelling of	

Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

Candy/ confectionery	45 g/100 g
Cereal bars	53 g/100 g
Powdered nutritional drink mixes (dry weight)	53 g/100 g
Carbonated beverages	11 g/100 g
Ice cream & sorbet	31 g/100 g
Yoghurt	12 g/100 g
Biscuits	53 g/100 g
Buns, cakes and pastries	53 g/100 g
Breakfast cereals (wholegrain)	88 g/100 g
Jams and jellies in accordance with Directive 2001/113/EC	133 g/100 g Based on pre- processing quantity to produce final 100 g product
Sweet spreads, fillings and icings	31 g/100 g
Savoury sauces, pickles, gravies and condiments	88 g/100 g
Food Supplements as defined in Directive 2002/46/EC	26 g/day
	Fruit concentrate
Candy/ Confectionery	10 g/100 g
Cereal bars	12 g/100 g

the foodstuffs containing it shall be: For fruit puree: ` Morinda citrifolia fruit puree' or 'Noni fruit puree' For fruit concentrate: *' Morinda* citrifolia fruit concentrate' or 'Noni fruit concentrate'

	For the preparation of infusions	A cup of infusion to be consumed shall not be prepared with more than 1 g of dried and roasted leaves	the novel food on the labell of the foods	ing	
Noni leaves (<i>Morinda</i> citrifolia)	Specified food category	Maximum levels	1. The design of	nation	
	Food Supplements as defined in Directive 2002/46/EC	6 g/day			
	Savoury sauces, pickles, gravies and condiments	20 g/100 g			
	Sweet spreads, fillings and icings	7 g/100 g			
	Jams and jellies in accordance with Directive 2001/113/EC	30 g/100 g			
	Breakfast cereals (wholegrain)	20 g/100 g			
	Buns, cakes and pastries	12 g/100 g			
	Biscuits	12 g/100 g			
	Ice cream & sorbet Yoghurt	7 g/100 g 3 g/100 g			
	Carbonated beverages	3 g/100 g			
	Powdered nutritional drink mixes (dry weight)	12 g/100 g			

		of Morinda citrifolia	containir it shall be 'Noni leaves' or 'leaves of Morinda citrifolia '. 2. Instruction shall be given to the consume that a cup of infusion should not be prepared with more than 1 g of dried and roasted leaves of Morinda citrifolia	ons
Noni fruit powder (<i>Morinda</i>	Specified food category	Maximum levels	The designation of the novel	
citrifolia)	Food Supplements as defined in Directive 2002/46/EC	2,4 g per/day	food on the labelling of the foodstuffs containing it shall be 'Morinda citrifolia fruit powder' or	

			' Noni fruit powder '	
Odontella aurita microalgae	Specified food category	Maximum levels	The designation of the novel	
-	Flavoured pasta	1,5 %	food on the labelling of	
	Fish soups	1 %	the foodstuffs containing	
	Marine terrines	0,5 %	it shall be ' Odontella	
	Broth preparations	1 %	aurita microalgae'	
	Crackers	1,5 %		
	Frozen breaded fish	1,5 %		
Oil enriched with phytosterols/phytostanols	Specified food category	Maximum levels of phytosterols/ phytostanols	In accordance with Annex III.5 to Regulation	
	Spreadable fats as defined in Annex VII, Part VII and Appendix II, points B and C of Regulation (EU) No 1308/2013, and excluding cooking and frying fats and spreads based on butter or other animal fat Milk based products, such as products based on semiskimmed and skimmed milk products,	1. The productonta the novel food ingreshall be prese in such a mann that they can be easily dividinto portion that conta either a maxim of 3 g (in case	(EÜ) No ldts69/2011 ining dient ted ons in	

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	or cereals,	per			
	products	day)			
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	fermented	maxi	mum		
	milk such as	of 1			
	yoghurt and	g (in			
	cheese based	case			
	products (fat	of			
	content ≤ 12	three			
	g per 100	portio)fis		
	g), where	per			
	possibly the milk fat has	day)			
	been reduced	of added	4		
	and the fat or				
			sterols/		
	protein has	2. The	stanols.		
	been partly or	amou	nt		
	fully replaced by vegetable	of	1111		
	fat or protein		sterols/		
			stanols		
	Soya drinks	added			
	Salad	to a	.		
	dressings,	conta	iner		
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	sauces	shall	-5°5		
	Sauces	not			
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Oil	Specified	Maximum	The		
extracted	food	levels	designation		
from squids	category	of DHA	of the novel		
nom squius		and EPA	food on the		
		combined	labelling of		
	Dairy	200 mg/100 g	the foodstuffs		
	products	or for cheese	containing		
	products	of for cheese	Containing		

except milk-based beverages	products 600 mg/100 g	it shall be ' Squid oil '.
Dairy analogues except drinks	200 mg/100 g or for analogues to cheese products 600 mg/100 g	
Spreadable fat and dressings	600 mg/100 g	
Breakfast cereals	500 mg/100 g	
Bakery products (breads and bread rolls)	200 mg/100 g	
Cereal bars	500 mg/100 g	
Non- alcoholic beverages (including milk-based beverages)	60 mg/100 ml	
Food Supplements as defined in Directive 2002/46/EC	3 000 mg/day for general population 450 mg/day for pregnant and lactating women	
Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products intended	
Total diet replacement for weight control defined in Regulation (EU) No	200 mg/meal	

[F5Partially defatted chia seed (Salvia hispanica)	and meal replacements for weight control Specified food category Powder with his content	Maximum levels	The designation of the novel food on the		
powders	Unflavoured fermented milk products, including natural unflavoured buttermilk (excluding sterilised buttermilk) non-heat-treated after fermentation	0,7 %	labelling of the foodstuffs containing it shall be 'Partially defatted chia seed (Salvia hispanica) powder'	the foodstuffs containing it shall be 'Partially defatted chia seed (Salvia hispanica)	
	Unflavoured fermented milk products, heat- treated after fermentation	0,7 %			
	Flavoured fermented milk products including heat-treated products	0,7 %			
	Confectionery	10 %			
	Fruit juices as defined by Directive 2001/112/ EC h and vegetable juices	2,5 %			
	Fruit nectars as defined by Directive 2001/112/EC	2,5 %			

and vegetable nectars and similar products	
Flavoured drinks	3 %
Food supplements as defined in Directive 2002/46/EC excluding food supplements for infants and young children	7,5 g/day
Powder with hi content	gh fibre
Confectionery	4 %
Fruit juices as defined by Directive 2001/112/EC and vegetable juices	2,5 %
Fruit nectars as defined by Directive 2001/112/EC and vegetable nectars and similar products	4 %
Flavoured drinks	4 %
Food supplements as defined in Directive 2002/46/EC excluding food supplements for infants and young children	12 g/day]

ъ .	Cnacified	Manisses			
Pasteurised	Specified	Maximum	The wording		
fruit-based	food	levels	' pasteurised		
preparations	category		by high-		
produced	Types of		pressure		
using high-	fruit:		treatment		
pressure	apple, apricot,		' shall be		
treatment	banana,		displayed		
***************************************	blackberry,		next to the		
	•		name of		
	blueberry,		the fruit		
	cherry,				
	coconut,		preparations		
	fig, grape,		as such and in		
	grapefruit,		any product		
	mandarin,		in which it is		
	mango,		used		
	melon,				
	peach, pear,				
	pineapple,				
	prune,				
	raspberry,				
	rhubarb,				
	· ·				
	strawberry				
[F29Phenylcaps	Spacified	Maximum	The		Authorised on
1 Henyicap	food category	levels	designation		19 December
			of the novel		2019. This
	Foods for	2,5 mg/day	food on the		inclusion
	special				
			i ianeiling of		is hased on
	medical		labelling of		is based on
			the foodstuffs		proprietary
	medical		the foodstuffs containing		proprietary scientific
	medical purposes as defined under		the foodstuffs containing it shall be '		proprietary scientific evidence and
	medical purposes as defined under Regulation		the foodstuffs containing	n	proprietary scientific evidence and scientific data
	medical purposes as defined under Regulation (EU) No		the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in
	medical purposes as defined under Regulation (EU) No 609/2013		the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance
	medical purposes as defined under Regulation (EU) No 609/2013 excluding		the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for		the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants,		the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young		the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children		the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU)
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children		the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283.
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age		the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant:
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children		the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: aXichem AB,
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age of 11 years	2.5 mg/davil	the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: aXichem AB, Södergatan
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age of 11 years Food	2,5 mg/day]	the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: aXichem AB, Södergatan 26, SE 211
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age of 11 years Food supplements	2,5 mg/day]	the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: aXichem AB, Södergatan 26, SE 211 34, Malmö
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age of 11 years Food supplements as defined	2,5 mg/day]	the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: aXichem AB, Södergatan 26, SE 211 34, Malmö Sweden.
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age of 11 years Food supplements as defined in Directive	2,5 mg/day]	the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: aXichem AB, Södergatan 26, SE 211 34, Malmö Sweden. During the
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age of 11 years Food supplements as defined in Directive 2002/46/EC	2,5 mg/day]	the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: aXichem AB, Södergatan 26, SE 211 34, Malmö Sweden. During the period of data
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age of 11 years Food supplements as defined in Directive	2,5 mg/day]	the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: aXichem AB, Södergatan 26, SE 211 34, Malmö Sweden. During the period of data protection,
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age of 11 years Food supplements as defined in Directive 2002/46/EC intended for	2,5 mg/day]	the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: aXichem AB, Södergatan 26, SE 211 34, Malmö Sweden. During the period of data
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age of 11 years Food supplements as defined in Directive 2002/46/EC intended for the general	2,5 mg/day]	the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: aXichem AB, Södergatan 26, SE 211 34, Malmö Sweden. During the period of data protection,
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age of 11 years Food supplements as defined in Directive 2002/46/EC intended for the general population,	2,5 mg/day]	the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: aXichem AB, Södergatan 26, SE 211 34, Malmö Sweden. During the period of data protection, the novel food
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age of 11 years Food supplements as defined in Directive 2002/46/EC intended for the general population, excluding	2,5 mg/day]	the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: aXichem AB, Södergatan 26, SE 211 34, Malmö Sweden. During the period of data protection, the novel food phenylcapsaicin
	medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age of 11 years Food supplements as defined in Directive 2002/46/EC intended for the general population,	2,5 mg/day]	the foodstuffs containing it shall be '	n	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: aXichem AB, Södergatan 26, SE 211 34, Malmö Sweden. During the period of data protection, the novel food

	under the age of 11 years				on the market within the Union only by aXichem AB, unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of aXichem AB.
Phosphated maize starch	Specified food category	Maximum levels	The designation of the novel		
	Baked bakery products	15 %	food on the labelling of		
	Pasta		the foodstuffs containing		
	Breakfast cereals		it shall be 'Phosphated maize starch'		
	Cereal bars		maize staten		
Phosphatidyls from fish phospholipids	food	Maximum levels of phosphatidyls	The designation Win Re novel		
Beverages based on yoghurt Powders	Beverages based on	50 mg/100 ml 3 500 mg/100	food on the labelling of the foodstuffs containing it shall be 'Fish		
	based on milk			rine	
	Foods based on yoghurt	80 mg/100 g			

	Cereal bars	350 mg/100 g			
	Chocolate based confectionary	200 mg/100 g			
	Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In compliance with Regulation (EU) No 609/2013			
	Food supplements as defined in Directive 2002/46/EC	300 mg/day			
Phosphatidyls from soya	food	Maximum levels of phosphatidyls	The designation		
phospholipids	Beverages based on yoghurt	50 mg/100 ml	food on the labelling of the foodstuffs		
	Powders based on milk powder	3,5 g/100 g (equivalent to 40 mg/100 ml ready to drink)	containing it shall be 'Soya phosphatidylse	rine	
	Foods based on yoghurt	80 mg/100 g			
	Cereal bars	350 mg/100 g			
	Chocolate based confectionary	200 mg/100 g			
	Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In compliance with Regulation (EU) No 609/2013			
Phospholipid product containing	Specified food category	Maximum levels of phosphatidyls	The designation evinus	The product is not intended to	
equal amounts of	Breakfast cereals	80 mg/100 g	food on the labelling of	be marketed to pregnant	

phosphatidyls and	ecineal bars	350 mg/100 g	the foodstuffs	or breast-	
phosphatidic acid	Foods based on yogurt	80 mg/100 g	containing shall be 'Soy phosphatidylse	feeding women rine	
	Soy-based yogurt-like products	80 mg/100 g	and phosphatidic acid '		
	Yogurt based- drinks	50 mg/100 g			
	Soy-based yogurt-like drinks	50 mg/100 g			
	Powders based on milk powder	3,5 g/100 g (equivalent to 40 mg/100 ml ready-to drink)			
	Food Supplements as defined in Directive 2002/46/EC	800 mg/day			
	Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In compliance with Regulation (EU) No 609/2013			
Phospholipide from egg yolk	_S Specified food category	Maximum levels			
·	Not specified				
Phytoglycoger	Specified food category	Maximum levels	The designation of the novel		
	Processed foods	25 %	food on the labelling of the foodstuffs containing it shall be ' Phytoglycogen'		
Phytosterols/ phytostanols	Specified food category	Maximum levels	In accordance with Annex III.5 of Regulation		

Rice drinks	(EU) No 1. They 1169/2011
Rye bread	shall
with flour	be
containing	presented
≥ 50 % rye	in
(wholemeal	such
rye flour,	
whole or	a
cracked rye	manner
	that
kernels and	they
rye flakes)	can
and $\leq 30 \%$	be
wheat; and	easily
with $\leq 4 \%$	divided
added sugar	into
but no fat	portions
added.	that
Salad	contain
dressings,	either
mayonnaise	a
	maximum
and spicy	of 3
sauces.	g (in
Soya drink	case
•	of 1
Milk type	portion/
products,	day)
such as semi-	or a
skimmed and	maximum
skimmed	of 1
milk type	g (in
products,	case
possibly with	of 3
the addition	portions/
of fruits and/	day)
or cereals,	of
where	added
possibly the	phytosterols/
milk fat has	phytosterois/ phytostanols.
been reduced,	The phytostanois.
or where milk	I I
fat and/or	amount of
protein has	phytosterols/
been partly or	phytostanols
fully replaced	added to a
by vegetable	container of
fat and/or	beverages
protein.	shall not
_	exceed 3 g.
Products	Salad
based on	dressings,
fermented	mayonnaise
milk such	and spicy
	·

	as yoghurt and cheese type products (fat content < 12 % per 100 g), where possibly the milk fat has been reduced, or where milk fat and/or protein has been partly or fully replaced by vegetable fat and/or protein	sauces shall be packed as single portions		
	Spreadable fats as defined in Annex VII, Part VII and Appendix II, points B and C of Regulation (EU) No 1308/2013, and excluding cooking and frying fats and spreads based on butter or other animal fat.			
	Food Supplements as defined in Directive 2002/46/EC	3 g/day		
Plum kernel oil	Specified food	Maximum levels		
	For frying and as seasoning	In line with normal food use of vegetable oils		
Potato proteins (coagulated)	Not specified		The designation of the novel	

and hydrolysates thereof			food on the labelling of the foodstuffs containing it shall be ' Potato protein	
Prolyl oligopeptidase (enzyme preparation)	Food Supplements as defined in Directive 2002/46/ EC for general adult population	Maximum levels 120 PPU/ day (2,7 g of enzyme preparation/ day) (2 × 10 6 PPI/day) PPU – Prolyl Peptidase Units or Proline Protease Units PPI – Protease Picomole International	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Prolyl oligopeptidase,'	
[F30Protein extract from	Specified food category	Maximum levels		
	Food Supplements as defined in Directive 2002/46/EC	3 capsules or 3 tablets/day; equalising 12,6 mg pig kidney		
	Food for special medical purposes as defined in Regulation (EU) No 609/2013]	extract a day Diamine oxidase (DAO) content: 0,9 mg/day (3 capsules or 3 tablets with a content of DAO of 0,3 mg/capsule or 0,3 mg/tablet)		
[F31Pyrroloqui quinone	nothicified food category	Maximum levels	The designation of the novel	Authorised on 2 September 2018. This
disodium salt	Food Supplements	20 mg/day]	food on the labelling of	inclusion is based on proprie

scientific

Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

as defined in Directive 2002/46/ EC intended for the adult population, excluding pregnant and lactating women the foodstuffs containing it shall be 'Pyrrologuinoline quinone disodium salt'. Food supplements containing Pyrroloquinoline quinone disodium salt shall bear the following statement: This food supplement should be consumed by adults only excluding pregnant and lactating

women

evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Mitsubishi Gas Chemical Company, Inc., Mitsubishi Building 5-2 Marunouchi 2-chome, Chiyodaku, Tokyo 100-8324, Japan. During the period of data protection the novel food Pyrroloquinoline quinone disodium salt is authorised for placing on the market within the Union only by Mitsubishi Gas Chemical Company, Inc., unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance

					with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Mitsubishi Gas Chemical Company, Inc. End date of the data protection: 2 september 2023
Rapeseed oil high in	Specified food	Maximum levels	The designation		
unsaponifiable		1.5.	of the novel		
matter	Food Supplements as defined in Directive 2002/46/EC	1,5 g per portion recommended for daily consumption	food on the labelling of the foodstuffs containing it shall be ' Rapeseed oil extract '		
Rapeseed Protein	As a vegetable protein source in foods except in infant formula and follow-on formula		of the novel food on the labell of the foods conta it shall be 'Rape prote 2. Any foods	ing stuffs ining eseed in'. stuff ining seed	

			statement that this ingredient may cause allergic reaction to consumers who are allergic to mustard and products thereof. Where relevant, this statement shall appear in close proximity to the list of	
[F32Refined shrimp peptide concentrate	Specified food category Food Supplements as defined in Directive 2002/46/EC for the adult population	Maximum levels 1 200 mg/day]	ingredients. The designation of the novel food on the labelling of the foodstuffs containing it shall be 'refined shrimp peptide concentrate'.	Authorised on 20 November 2018. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Marealis AS.,

			End date of the data protection: 20 November 2023.
			Hus, 2nd floor, N-9008 Tromsø Postal address: P.O. Box 1065, 9261 Tromsø, Norway. During the period of data protection the novel food refined shrimp peptideconcentrate is authorised for placing on the market within the Union only by Marealis AS unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Marealis AS.

	Food Supplements as defined in Directive 2002/46/ EC for adult population (capsule or tablet form)	150 mg/day		the novel food on the labelling of the food supplements containing it shall be ' Trans
Trans-	Specified	Maximum	2.	resveratrol'. The labelling of food supplements containing trans- resveratrol shall bear a statement that people using medicines should only consume the product under medical supervision.
Trans- resveratrol (microbial source)	Food supplements as defined in Directive 2002/46/EC	In line with normal use in food supplements of resveratrol extracted from Japanese	1.	The designation of the novel food on the labelling of the

		knotweed (Fallopia japonica)	food supplements containing it shall be ' Trans - resveratrol'. 2. The labelling of food supplements containing trans-resveratrol shall bear a statement that people using medicines should only consume the product under medical supervision.
Rooster comb extract	Specified food category	Maximum levels	The designation of the novel
	Milk-based drinks	40 mg/100 g or mg/100 ml	food on the labelling of
	Milk based fermented drinks	80 mg/100 g or mg/100 ml	the foodstuffs containing it shall be ' Rooster comb
	Yoghurt-type products	65 mg/100 g or mg/100 ml	extract ' or ' Cockerel comb extract
	Fromage frais	110 mg/100 g or mg/100 ml	comb extract
Sacha inchi oil from Plukenetia volubilis	Specified food category As for linseed oil	Maximum levels In line with normal food	The designation of the novel food on the labelling of

		use of linseed oil	the foodstuffs containing it shall be 'Sacha inchi oil (Plukenetia volubilis)'	
Salatrims	Specified food category	Maximum levels	1. The desig	nation
	Bakery products and confectionary		the nove food on the label of the foods contait shall be 'redu energ fat (salar 2. There shall be a state that excest consumay lead to gastrintes	ling stuffs sining ced gy trims)'. e ment ssive amption o- tinal rbance. e ment

			by	ron	
Schizochytrium sp. oil rich in DHA and EPA	m Specified food category	Maximum levels of DHA and EPA combined:	The designation of the novel food on the labelling of	en.	
	Food Supplements as defined in Directive 2002/46/ EC for adult population excluding pregnant and lactating women	3 000 mg/day	the foodstuffs containing it shall be 'DHA and EPA-rich oil from the microalgae Schizochytrium sp.'		
	Food Supplements as defined in Directive 2002/46/EC for pregnant and lactating women	450 mg/day			
	Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended			
	Total diet replacement for weight control as defined in Regulation (EU) No 609/2013 and meal replacements for weight control	250 mg/meal			
	Milk-based drinks and similar	200 mg/100 g			

products intended for young children Processed cereal based food and baby food for infants and young children as defined in Regulation (EU) No 609/2013	
Foods intended to meet the expenditure of intense muscular effort, especially for sportsmen	
Foods bearing statements on the absence or reduced presence of gluten in accordance with the requirements of Commission Implementing Regulation (EU) No 828/2014	
Bakery Products (Breads, Rolls and Sweet Biscuits)	200 mg/100 g
Breakfast Cereals	500 mg/100 g
Cooking Fats	360 mg/100 g

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	Dairy Analogues except drinks	600 mg/100 g for cheese; 200 mg/100 g for soy and imitation milk products (excluding drinks)			
	Dairy Products except milk- based drinks	600 mg/100 g for cheese; 200 mg/100 g for milk products (including milk, fromage frais and yoghurt products; excluding drinks)			
	Non- alcoholic Beverages (including dairy analogue and milk-based drinks)	80 mg/100 g			
	Cereal/ Nutrition Bars	500 mg/100 g			
	Spreadable Fats and Dressings	600 mg/100 g			
[F33Schizochyte sp. (ATCC PTA-9695)	ritim food category	Maximum levels of DHA	The designation of the novel		
oil	Dairy products except milk- based drinks	200 mg/100 g or for cheese products 600 mg/100 g	food on the labelling of the foodstuffs containing it shall be		
	Dairy analogues except drinks	200 mg/100 g or for analogues to cheese products 600 mg/100 g	'Oil from the microalgae Schizochytrium sp.'	,	

Spreadable fats and dressings	600 mg/100 g
Breakfast cereals	500 mg/100 g
Food Supplements as defined in Directive	250 mg DHA/day for general population
2002/46/EC	450 mg DHA/day for pregnant and lactating women
Total diet replacement for weight control as defined in Regulation (EU) No 609/2013 and meal replacements for weight control	250 mg/meal
Milk-based drinks and similar products intended for young children	200 mg/100 g
Foods intended to meet the expenditure of intense muscular effort, especially for sportsmen	
Foods bearing statements on the absence or reduced presence of gluten in	

accordance with the requirements of Commission Implementing Regulation (EU) No 828/2014		
Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended	
Bakery products (breads, rolls, and sweet biscuits)	200 mg/100 g	
Cereal bars	500 mg/100 g	
Cooking fats	360 mg/100 g	
Non- alcoholic beverages (including dairy analogue and milk-based drinks)	80 mg/100 ml	
Infant formula and follow-on formula as defined in Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013	
Processed cereal-based foods and baby foods for infants	200 mg/100 g	

	and young children as defined in Regulation (EU) No 609/2013 Fruit/ vegetable puree	100 mg/100 gl			
	iSpacified food	Maximum			
sp. strain (FCC-3204) oil	Food supplements as defined in the Food Supplements (England) Regulations 2003, excluding food supplements for infants and children under the age of 3.	levels of DHA 1000mg/day	The designation of the novel food on the labelling of the foodstuffs containing it is 'Oil from the microalgae Schizochytriun sp.'.	1	
	Infant formula and follow-on formula as defined in Regulation 609/2013	In accordance with Regulation 609/2013.	The labelling of food supplements containing Schizochytrium sp. strain (FCC-3204) oil must bear a statement that they should not be consumed by infants and children under the age of 3.]	ı	
[F35Schizochyttsp.(FCC-3204	rispacified food category	Maximum levels of DHA			
oil	Food supplements as defined in the Food Supplements (Scotland)	1 g/day	The designation of the novel food on the labelling of the foodstuffs		

	Regulations 2003 excluding food supplements for infants and children under 3 years of age		containing it is "Oil from the microalgae <i>Schizochytrium</i> sp.".	
	Infant formula and follow-on formula as defined in Regulation (EU) 609/2013	In accordance with Regulation (EU) 609/2013	The labelling of food supplements containing <i>Schizochytrium</i> sp. (FCC-3204) oil must bear a statement that they should not be consumed by infants and children under 3 years of age.]	
[F36Schizochyttsp.(FCC-3204] oil	Food Supplements as defined in the Food Supplements (Wales) Regulations 2003, excluding food supplements for infants and children under 3 years of age	Maximum levels of DHA 1g/day	The designation of the novel food on the labelling of the foodstuffs containing it is "Oil from the microalgae Schizochytrium sp.". The labelling of food supplements containing Schizochytrium sp.	
	Infant formula and follow-on formula as defined in Regulation	In accordance with Regulation (EU) No 609/2013	oil must bear a statement that they should not be consumed	

			under 3 years of age.]		
Spron Con Dan property of the control of the contro	tritimecified food category	Maximum levels of DHA	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Oil from the microalgae Schizochytrium sp.'		
	Dairy products except milk- based drinks	200 mg/100 g or for cheese products 600 mg/100 g		labelling of the foodstuffs containing it shall be 'Oil from the microalgae Schizochytrium	
	Dairy analogues except drinks	200 mg/100 g or for analogues to cheese products 600 mg/100 g			
	Spreadable fat and dressings	600 mg/100 g			
	Breakfast cereals	500 mg/100 g			
	Food Supplements as defined in Directive	250 mg DHA/day for general population			
	2002/46/EC	450 mg DHA/day for pregnant and lactating women			
	Total diet replacement for weight control as defined in Regulation (EU) No 609/2013 and meal replacements for weight control	250 mg/meal			
	Milk-based drinks and similar products intended	200 mg/100 g			

for young children Processed		
cereal-based foods and baby foods for infants and young children as defined in Regulation (EU) No 609/2013		
Foods intended to meet the expenditure of intense muscular effort, especially for sportsmen		
Foods bearing statements on the absence or reduced presence of gluten in accordance with the requirements of Implementing Regulation (EU) No 828/2014		
Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended	
Bakery products (breads, rolls,	200 mg/100 g	

	and, sweet biscuits) Cereal bars Cooking fats Non-alcoholic beverages (including dairy analogue and milk-based drinks)	500 mg/100 g 360 mg/100 g 80 mg/100 ml			
	Fruit/ vegetable puree	100 mg/100 gl			
[F18 Schizochyttsp. (T18) oil	iSpa cified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Oil from the microalgae Schizochytrium sp.'.		
	Dairy products except milk- based drinks	200 mg/100 g or for cheese products 600 mg/100 g			
	Dairy analogues except drinks	200 mg/100 g or for analogues to cheese products 600 mg/100 g		microalgae Schizochytrium	
	Spreadable fats and dressings	600 mg/100 g			
	Breakfast cereals	500 mg/100 g			
	Food supplements as defined in Directive	250 mg DHA/day for general population			
	2002/46/EC	450 mg DHA/day for pregnant and lactating women			
	Total diet replacement	250 mg/meal			

for weight control as defined in Regulation (EU) No 609/2013 and meal replacements for weight control	
Milk-based drinks and similar products intended for young children	200 mg/100 g
Foods intended to meet the expenditure of intense muscular effort, especially for sportsmen	
Foods bearing statements on the absence or reduced presence of gluten in accordance with the requirements of Commission Implementing Regulation (EU) No 828/2014	
Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the

	I	I 1 .	I	
		products are intended		
	Bakery products (breads, rolls and, sweet biscuits)	200 mg/100 g		
	Cereal bars	500 mg/100g		
Cooking fats	360 mg/100 g			
	Non- alcoholic beverages (including dairy analogue and milk-based drinks)	80 mg/100 ml		
	Infant formula and follow-on formula as defined in Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013		
	Processed cereal-based foods and baby foods for infants and young children as defined in Regulation (EU) No 609/2013	200 mg/100 g		
	Fruit/ vegetable puree	100 mg/100 gl		
р.	isna cified food category	Maximum levels of DHA		
WZU477) iil	Infant formula and follow-on formula as defined in	In accordance with Regulation 609/2013.	The designation of the novel food on the labelling of the foodstuffs	

Regulation 609/2013	containing it is 'Oil from the microalgae Schizochytriun sp.'.	evidence and scientific data protected in accordance with Article 26 of Regulation 2015/2283. Applicant: Progress Biotech BV of Canaalstaete, Kanaalweg 33, 2903LR Capelle aan den Ijssel, The Netherlands. During the period of data protection, Schizochytrium sp. (WZU477) oil is authorised for placing on the market within England only by Progress Biotech BV unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of
		protected in accordance with Article

				of Progress Biotech BV. The data protection will expire at the end of 29th June 2027.]
[F39Schizochytr (WZU477) oil	formula and follow- on formula as defined in Regulation (EU) 609/2013			
			During the period of data protection, the novel food Schizochytrium sp. (WZU477) oil is authorised for placing on the market within Scotland only by Progress Biotech BV unless a subsequent	

				applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Progress Biotech BV.
				will expire at the end of 29 June 2027.]
[^{F40} Schizochyti sp. (WZU477) oil	Infant formula and follow-on formula as defined inRegulation (EU) No 609/2013	Maximum levels of DHA In accordance withRegulation (EU) No 609/2013	The designation of the novel food on the labelling of the foodstuffs containing it is "Oil from the microalgae Schizochytrium sp."	Included in the list on 30 June 2022. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Progress Biotech BV, Canaalstaete, Kanaalweg 33, 2903LR Capelle aan den Ijssel, the Netherlands.

			During the period of data protection, Schizochytrium sp. (WZU477) oil is authorised for placing on the market within Wales only by Progress Biotech BV unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Progress Biotech BV. The data protection ends at the end of 29 June 2027.]
[F41Syrup from Sorghum bicolor (L.) Moench (Traditional food from a third country)	Not specified	This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of	

			Regulation (EU) 2015/2283.	
Fermented soybean extract	Specified food category Food Supplements as defined in Directive 2002/46/EC (capsules, tablets or powder form) intended for the adult population, excluding pregnant and lactating women	Maximum levels 100 mg/day	Applicant: Progress Biotech BV of Canaalstaete, Kanaalweg 33, 2903LR Capelle aan den Ijssel, the Netherlands.	
Spermidine- rich wheat germ extract (Triticum aestivum)	Specified food category Food Supplements as defined in Directive 2002/46/ EC intended for the adult population, excluding pregnant and lactating women	Maximum levels Equivalent of max. 6 mg/day spermidine	The data protection will expire at the end of 29 June 2027.	
Sucromalt	Specified food category Not specified	Maximum levels	of the novel food on the labell of the foods	ing tuffs ining

			2. The design of the nove food on the label shall be according to the	lling mpanied eation uct ce	
Sugar cane fibre	Specified food category	Maximum levels			
	Bread	8 %			
	Bakery goods	5 %			
	Meat and muscle products	3 %			
	Seasonings and spices	3 %			
	Grated cheeses	2 %			
	Special diet foods	5 %			
	Sauces	2 %			
	Beverages	5 %			
[F42Sugars obtained from cocoa (Theobroma cacao L.) pulp	Not specified		The designation of the novel food on the labelling of the foodstuffs		

			containing it shall be 'sugars obtained from cocoa (Theobroma cacao L.) pulp', 'Glucose obtained from cocoa (Theobroma cacao L.) pulp' or 'Fructose obtained from cocoa (Theobroma cacao L.) pulp' or 'Indiana cacao L.) pulp' or 'Indiana cacao L.) pulp', depending on the form used.]	
Sunflower oil extract	Specified food	Maximum levels	The designation	
	Food Supplements as defined in Directive 2002/46/EC	1,1 g/day	of the novel food on the labelling of the foodstuffs containing it shall be ' Sunflower oil extract '	
Dried Tetraselmis chuii	Specified food category	Maximum levels	The designation of the novel	
microalgae	Sauces	20 % or 250mg/day	food on the labelling of the foodstuffs	
	Special salts	1 %	containing	
	Condiment	250 mg/day	it shall be 'Dried	
	Food Supplements as defined in Directive 2002/46/EC	250 mg/day	microalgae Tetraselmis chuii ' or 'Dried microalgae T. chuii ' Food supplements containing dried	

			microalgae Tetraselmis chuii shall bear the following statement: 'Contains negligible amounts of iodine'		
Therapon barcoo / Scortum	Intended use identical to that of the salmon, namely the preparation of culinary fish products and dishes, including cooked, raw, smoked and baked fish products				
D-Tagatose	Specified food category Not specified	Maximum levels	of the novel food on the labell of the foods	ing tuffs ining ose'. ling oct e	

			greate than 1 % D- Tagat (as consu shall bear a stater 'exce	ining er cose amed) ment ssive amption ace ve	
[^{F18} Taxifolin- rich extract	Specified food category Yogurt plain/ Yogurt with	Maximum levels 0,020 g/kg	The designation of the novel food on the labelling of		
	fruits (*)		the foodstuffs		
	Kephir (*)	0,008 g/kg	containing it shall be '		
	Buttermilk (*)	0,005 g/kg	taxifolin-rich extract '		
	Milk powder	0,052 g/kg			
	Cream (*)	0,070 g/kg			
	Sour cream	0,050 g/kg			
	Cheese (*)	0,090 g/kg			
	Butter (*)	0,164 g/kg			
	Chocolate confectionery	0,070 g/kg			
	Non- alcoholic beverages	0,020 g/L			
	Food supplements as defined in Directive 2002/46/EC intended for	100 mg/day			

	produ rich e replac in par	n used in milk acts Taxifolin- extract may not bee in whole or rt, any milk ituent]				
Trehalose	Specified food category Not specified	Maximum levels	1.	of the novel food on the labell of the foods contait shall be 'Treh and shall be displation the labell of the product as such or in the list of	ing tuffs ining alose' nyed ing act	

			of the novel food on the labell shall be accor by indicathat the	nation ing mpanied ation alose e	
[^{F43} UV- treated baker's yeast	Specified food category	Maximum level of vitamin D ₂	The designation of the novel	The novel food must be inactivated	
(Saccharomyce cerevisiae)	Syeast- leavened breads and rolls	5 μg/100 g	food on the labelling of food containing it is "vitamin	for use in infant formula, follow-on formula,	
	Yeast- leavened fine bakery wares	5 μg/100 g	D yeast" or "vitamin D ₂ yeast".	processed cereal-based food and food	
	Food supplements as defined in the Food Supplements (England) Regulations 2003	In accordance with any relevant requirements contained in regulations applying in relation to England and made under regulation 4 of the Nutrition (Amendment etc.) (EU Exit)		for special medical purposes.]	

	Regulations 2019	
Pre-packed fresh or dry yeast for home baking	45 μg/100 g for fresh yeast, 200μg/100 g for dry yeast	The designation of the novel food on the labelling of food containing it is "vitamin D yeast" or "vitamin D ₂ yeast". The labelling of the novel food must bear a statement that the food is only intended for baking and it should not be eaten raw. The labelling of the novel food must bear instructions for use for the final consumer to ensure a maximum concentration of 5 µg/100 g of vitamin D ₂ in the final home-baked product is not exceeded.
Dishes, including ready-to- eat meals (excluding soups and salads)	3 μg/100 g	The designation of the novel food on the labelling of food containing it
Soups and salads	5 μg/100 g	is "vitamin D yeast" or

Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

Fried or extruded cereal, seed or root-based products	5 μg/100 g
Infant formula and follow-on formula as defined in Regulation (EU) No. 609/2013 ^j	In accordance with Regulation (EU) No. 609/2013
Processed cereal-based food as defined in Regulation (EU) No. 609/2013	In accordance with Regulation (EU) No. 609/2013
Processed fruit products	1.5 μg/100 g
Processed vegetables	2 μg/100 g
Bread and similar products	5 μg/100 g
Breakfast cereals	4 μg/100 g
Pasta, doughs and similar products	5 μg/100 g
Other cereal- based products	3 μg/100 g
Spices, seasonings, condiments, sauce ingredients, dessert sauces/ toppings	10 μg/100 g
Protein products	10 μg/100 g
Cheese	2 μg/100 g

 $\text{``vitamin}\ D_2$ yeast".

	Dairy desserts and similar products Fermented milk or fermented cream Dairy powders and concentrates Milk based products,	2 μg/100 g 1.5 μg/100 g 25 μg/100 g 0.5 μg/100 g			
	whey and cream Meat and dairy analogues	2.5 μg/100 g			
	Total diet replacement for weight control as defined in Regulation (EU) No. 609/2013	5 μg/100 g			
	Meal replacement for weight control	5 μg/100 g			
	Food for special medical purposes as defined in Regulation (EU) No. 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended.			
[F44UV- treated baker's yeast (Saccharomyc cerevisiae)	Specified food category esyeast-leavened breads and rolls	Maximum levels of Vitamin D# 5 μg/100 g	The designation of the novel food on the labelling of food containing it is "vitamin D yeast" or	The novel food must be inactivated for use in infant formula, follow-on formula, processed cereal-based	

Yeast- leavened fine bakery wares	5 μg/100 g	"vitamin D# yeast".	food and food for special medical
Food supplements as defined in the Food Supplements (Scotland) Regulations 2003	In accordance with any relevant requirements contained in regulations applying in relation to Scotland and made under regulation 4 of the Nutrition (Amendment etc.) (EU Exit) Regulations 2019		purposes.]
Pre-packed fresh or dry yeast for home baking	45 μg/100 g for fresh yeast 200 μg/100 g for dry yeast	The designation of the novel food on the labelling of food containing it is "vitamin D yeast" or "vitamin D# yeast". The labelling of the novel food must bear a statement that the food is only intended for baking and it should not be eaten raw. The labelling of the novel food must bear instructions for use for the final consumer	

Dishes, including ready-to-eat meals (excluding soups and salads)	3 μg/100 g	to ensure a maximum concentration of 5µg/100g of vitamin D# in the final home-baked product is not exceeded. The designation of the novel food on the labelling of food containing it
Soups and salads	5 μg/100 g	is "vitamin D yeast" or "vitamin D#
Fried or extruded cereal, seed or root-based products	5 μg/100 g	yeast".
Infant formula and follow-on formula as defined in Regulation (EU) No. 609/2013	In accordance with Regulation (EU) No. 609/2013	
Processed cereal-based food as defined in Regulation (EU) No. 609/2013	In accordance with Regulation (EU) No. 609/2013	
Processed fruit products	1.5 μg/100 g	
Processed vegetables	2 μg/100 g	
Bread and similar products	5 μg/100 g	
Breakfast cereals	4 μg/100 g	

Pasta, doughs and similar products	5 μg/100 g
Other cereal- based products	3 μg/100 g
Spices, seasonings, condiments, sauce ingredients, dessert sauces/ toppings	10 μg/100 g
Protein products	10 μg/100 g
Cheese	2 μg/100 g
Dairy desserts and similar products	2 μg/100 g
Fermented milk or fermented cream	1.5 μg/100 g
Dairy powders and concentrates	25 μg/100 g
Milk-based products, whey and cream	0.5 μg/100 g
Meat and dairy analogues	2.5 μg/100 g
Total diet replacement for weight control as defined by Regulation (EU) No. 609/2013	5 μg/100 g
Meal replacement for weight control	5 μg/100 g

	Food for special medical purposes as defined by Regulation (EU) No. 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended			
[F45UV- treated baker's yeast (Saccharomyco cerevisiae)	Specified food category Yeast-leavened breads and	Maximum levels of vitamin D ₂ 5 μg/100 g	The designation of the novel food on the labelling of food	The novel food must be inactivated for use in infant formula, follow-on	
	Yeast- leavened fine bakery wares	5 μg/100 g	containing it is "vitamin D yeast" or "vitamin D ₂ yeast".	formula, processed cereal-based food and food for special medical purposes.]	
	Food supplements as defined in the Food Supplements (Wales) Regulations 2003	In accordance with any relevant requirements contained in regulations applying in relation to Wales and made under regulation 4 of the Nutrition (Amendment etc.) (EU Exit) Regulations 2019			
	Pre-packed fresh or dry yeast for home baking	45 μg/100 g for fresh yeast 200 μg/100 g for dry yeast	The designation of the novel food on the labelling of food containing it is "vitamin D yeast" or "vitamin D ₂ yeast". The labelling of the novel		

			food must bear a statement that the food is only intended for baking and should not be eaten raw. The labelling of the novel food must bear instructions for use for the final consumer to ensure a maximum concentration of 5µg/100g of vitamin D ₂ in the final home-baked product is not	
			product is not exceeded.	
UV-treated bread	Dishes, including ready-to-eat meals (excluding soups and salads) Soups and salads Fried or extruded cereal, seed or root-based products Infant	3 μg/100 g 5 μg/100 g 5 μg/100 g	The designation of the novel food on the labelling of food containing it is "vitamin D yeast" or "vitamin D ₂ yeast".	
	formula and follow-on formula as defined in Regulation (EU) No 609/2013	accordance with Regulation (EU) No 609/2013		
	Processed cereal-	In accordance		

based	with
food as	Regulation
defined in	(EU) No
Regulation	609/2013
(EU) No	
609/2013	
Processed	1.5 μg/100 g
fruit	
products	
Processed	2 μg/100 g
vegetables	
Bread and	5 μg/100 g
similar	
products	
Breakfast	4 μg/100 g
cereals	
Pasta,	5 μg/100 g
doughs	
and similar	
products	
Other	3 μg/100 g
cereal-	
based	
products	
Spices,	10 μg/100 g
seasonings,	
condiments,	
sauce	
ingredients,	
dessert	
sauces/	
toppings	
Protein	10 μg/100 g
products	, 0
Cheese	2 μg/100 g
Dairy	2 μg/100 g
desserts	F-8' 8
and similar	
products	
Fermented	1.5 µg/100 g
milk or	F-8 8
fermented	
cream	
Dairy	25 μg/100 g
powders	- 1-8-1-4-8
and	
concentrates	
Milk-based	0.5 μg/100 g
products,	
whey and	
cream	

food category levels of vitamin D 3 defectory Pasteurised whole milk as defined in Regulation (EU) No 1308/2013 to be consumed as such 5-32 μg/kg for general population excluding infants th Pasteurised semissimmed milk as defined in		Meat and dairy analogues Total diet replacement for weight control as defined in Regulation (EU) No 609/2013 Meal replacement for weight	2.5 μg/100 g 5 μg/100 g 5 μg/100 g
food category levels of vitamin D 3 design on the category Pasteurised whole milk as defined in Regulation (EU) No 1308/2013 to be consumed as such 5-32 μg/kg for general population excluding infants the label		Food for special medical purposes as defined in Regulation (EU) No	with the particular nutritional requirements of the persons for whom the products are
1308/2013 to be consumed as such Pasteurised semi- for general skimmed population milk as defined in infants food shall be 'UV- treate 2. Whe treate the formula with the food shall be 'UV- treate 2. Whe for general treate 3. Whe milk with the food shall be 'UV- treate 4. The food shall be 'UV-	JV-treated nilk	food category Pasteurised whole milk as defined in Regulation	Maximum levels of vitamin D 3 5-32 μg/kg for general population excluding
(EU) No		1308/2013 to be consumed as such Pasteurised semi- skimmed milk as defined in Regulation	1-15 µg/kg for general population excluding

			accordance with Point 2 of	
			Part A of Annex	
			XIII	
			Regulation (EU)	
			No 1169/2011	
			of the	
			European	
			Parliament and	
			of	
			the Council,	
			the	
			designation for	
			the labelling	
			shall	
			be accompanied	
			by	
			'contains vitamin	
			D	
			produced	
			by UV-	
			treatment'	
			ʻmilk	
			containing vitamin	
			D	
			resulting from	
			UV-	
-F7- va	Specified	Maximum	treatment'.	A 2241: 1
[F7Vitamin D 2 mushroom	food	levels of	The designation	Authorised on 27 August
powder	category	vitamin D	of the novel food on the	2020. This inclusion
		2	labelling of	is based on
	1		the foodstuffs	proprietary

Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

Breakfast cereals	2,25 μg of vitamin D ₂ /100 g
Yeast- leavened bread and pastries	2,25 μg of vitamin D ₂ /100 g
Grain products and pastas	2,25 μg of vitamin D ₂ /100 g
Fruit juice and fruit/ vegetable blend beverages	1,125 µg of vitamin D ₂ /100 mL
Milk and dairy products (excluding fluid milks)	2,25 μg of vitamin D ₂ /100 g/1,125 μg of vitamin D ₂ /100 mL (beverages)
Cheese (excluding cottage cheese, ricotta cheese, and hard-grating cheeses)	$^{2,25}\mu g$ of vitamin D $_2$ /100 g
Meal replacement bars and beverages	2,25 μg of vitamin D ₂ /100 g/1,125 μg of vitamin D ₂ /100 mL (beverages)
Dairy analogues	2,25 μg of vitamin D ₂ /100 g/1,125 μg of vitamin D ₂ /100 mL (beverages)
Meat analogues	2,25 μg of vitamin D ₂ /100 g
Soups and broths	2,25 μg of vitamin D ₂ /100 g

containing it shall be 'UV-treated mushroom powder containing vitamin D' or 'UV-treated mushroom powder containing vitamin D₂' The labelling of food supplements containing vitamin D₂ mushroom powder shall bear a statement that they should not be consumed by infants

scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Oakshire Naturals, LP., PO Box 388 Kennett Square, Pennsylvania 19348, United States. During the period of data protection, the novel food vitamin D₂ mushroom powder is authorised for placing on the market within the Union only by Oakshire Naturals, LP., unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of

	Extruded vegetable snacks Foods for Special Medical Purposes as defined under Regulation (EU) No 609/2013 excluding those intended for infants	2,25 μg of vitamin D ₂ /100 g 15 μg/day		Regulation (EU) 2015/2283 or with the agreement of Oakshire Naturals, LP. End date of the data protection: 27 August 2025.
	Food supplements as defined in Directive 2002/46/EC intended for the general population excluding infants	15 μg/day]		
[F46Vitamin D ₂ mushroom powder	Specified food category	Maximum levels of vitamin D ₂	The designation of the novel	Included in the list on 15
	Produfact	_	food on the	May 2023. This inclusion
	Breakfast cereals	2.1 μg/100 g	food on the labelling of food	This inclusion is based on proprietary
		_	food on the labelling of food containing it is "UV- treated mushroom powder	This inclusion is based on proprietary scientific evidence and scientific data protected in accordance
	Yeast leavened bread and similar	2.1 μg/100 g	food on the labelling of food containing it is "UV-treated mushroom	This inclusion is based on proprietary scientific evidence and scientific data protected in

Dairy products and analogues other than beverages	2.1 µg/100 g (marketed as such or reconstituted as instructed by the manufacturer)	must bear a statement that they should not be consumed by infants and children under 3 years
Dairy products and analogues as beverages	1.1 µg/100 ml (marketed as such or reconstituted as instructed by the manufacturer)	of age.]
Milk and dairy powders	21.3 µg/100 g (marketed as such or reconstituted as instructed by the manufacturer)	
Meat analogues	2.1 μg/100 g	
Soups	2.1 µg/100 ml (marketed as such or reconstituted as instructed by the manufacturer)	
Extruded vegetable snacks	2.1 μg/100 g	
Meal replacement for weight control	2.1 μg/100 g	
Food for special medical purposes as defined in Regulation (EU) No. 609/2013 excluding those	In accordance with the particular nutritional requ of the persons for whom the products are intended.	irements

protection, Vitamin D₂ mushroom powder is authorised for placing on the market, within England, only by MBio, Monaghan Mushrooms unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation 2015/2283 or with the agreement of MBio, Monaghan Mushrooms.

	intended for infants				
	Food supplements as defined in the Food Supplements (England) Regulations 2003 excluding food supplements for infants and children under 3 years of age.	15 μg of vitamin D ₂ /day			
^{F47} Vitamin D# mushroom	Specified food category	Maximum levels of vitamin D#	The designation of the novel		Included in the list on 15 May 2023.
I F	Breakfast cereals	2.1 μg/100 g	food on the labelling of food containing it is "UV-treated mushroom powder containing vitamin D#". The labelling of food supplements, as defined by the Food Supplements (Scotland) Regulations 2003, containing vitamin D# mushroom powder must bear a statement that they should not be consumed by infants and children under 3 years of age.		This inclusion is based on proprietary
	Yeast leavened bread and similar pastries	2.1 μg/100 g			scientific evidence and scientific data protected in accordance
	Grain products and pasta and similar products	2.1 μg/100 g		#". ling nts,	with Article 26 of Regulation (EU) 2015/2283. Applicant: MBio, Monaghan Mushrooms, Tullygony, Tyholland, Co. Monaghan, Ireland, H18
	Fruit/ vegetable juices and nectars	1.1 µg/100 ml (marketed as such or reconstituted as instructed by the manufacturer)		Supplements (Scotland) Regulations 2003, containing vitamin D# mushroom powder must bear a statement that they should not be consumed	
	Dairy products and analogues other than beverages	2.1 µg/100 g (marketed as such or reconstituted as instructed by the manufacturer)			
	Dairy products and	1.1 µg/100 ml (marketed			authorised for placing on the market,

analogues as beverages Milk and dairy powders	as such or reconstituted as instructed by the manufacturer) 21.3 µg/100 g (marketed as such or reconstituted as instructed by the
Meat analogues	manufacturer) 2.1 μg/100 g
Soups	2.1 µg/100 ml (marketed as such or reconstituted as instructed by the manufacturer)
Extruded vegetable snack	2.1 μg/100 g
Meal replacement for weight control	2.1 μg/100 g
Food for special medical purposes as defined in Regulation (EU) 609/2013 excluding those intended for infants	In accordance with the particular nutritional requirements of the persons for whom the products are intended
Food supplements as defined in the Food Supplements (Scotland) Regulations 2003 excluding	15 μg of vitamin D#/ day

within Scotland, only by MBio, Monaghan Mushrooms unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of MBio, Monaghan Mushrooms. The data protection will expire at the end of 14 May 2028.]

rF48x7*.	food supplements for infants and children under 3 years of age	Maximum	The		Included in	
[F48Vitamin D ₂ mushroom powder	Specified food category	Maximum levels of vitamin D 2	designation of the novel food on the labelling of food		Included in the list on 15 May 2023.	
	Breakfast cereals	2.1 μg/100 g		This inclusion is based on proprietary		
	Yeast- leavened bread and similar pastries	2.1 μg/100 g	containing it is "UV- treated mushroom powder		scientific evidence and scientific data protected in accordance	
	Grain products and pasta and similar products	2.1 μg/100 g	containing vitamin D ₂ ". The labelling of food supplements, as defined in the Food Supplements (Wales) Regulations 2003, containing vitamin D ₂ mushroom powder must bear a statement that they should not be consumed by infants and children under 3 years of age.	vitamin D ₂ ". The labelling of food supplements,		with Article 26 of Regulation (EU) 2015/2283. Applicant:
	Fruit / vegetable juices and nectars	1.1 µg/100 ml (marketed as such or reconstituted as instructed by the manufacturer)			MBio, Monaghan Mushrooms, Tullygony, Tyholland, Co. Monaghan, Ireland, H18 FW95. During the period of data protection, vitamin D ₂ mushroom powder is	
	Dairy products and analogues other than beverages	2.1 µg/100 g (marketed as such or reconstituted as instructed by the manufacturer)				
	Dairy products and analogues as beverages	1.1 µg/100 ml (marketed as such or reconstituted as instructed by the manufacturer)		under 3 years of age.		authorised for placing on the market, within Wales, only by MBio, Monaghan Mushrooms
	Milk and dairy powders	21.3 µg/100 g (marketed as such or reconstituted as instructed			unless a subsequent applicant obtains authorisation for the novel	

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	Meat analogues Soups Extruded vegetable snack Meal replacement for weight control Food for special medical purposes as defined in Regulation (EU) No 609/2013 excluding those intended for	by the manufacturer) 2.1 µg/100 g 2.1 µg/100 ml (marketed as such or reconstituted as instructed by the manufacturer) 2.1 µg/100 g 2.1 µg/100 g In accordance with the particular nutritional requirements of the persons for whom the products are intended		food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of MBio, Monaghan Mushrooms. The data protection will expire at the end of 14 May 2028.]
Vitamin K 2 (menaquinone	Food supplements as defined in the Food Supplements (Wales) Regulations 2003 excluding food supplements for infants and children under 3 years of age To be used in complete the complements of the complements of the complete t	2002/46/	The designation of the novel	
Vitamin K 2 (menaquinone	with Directive	2002/46/	designation	

	609/2013 and/o (EC) No 1925/		labelling of the foodstuffs containing it shall be 'Menaquinone or 'Vitamin K		
Wheat bran extract	Specified food category	Maximum levels	The designation of the novel	The 'Wheat Bran Extract 'may not be	
	Beer and substitutes	0,4 g/100 g	food on the labelling of the foodstuffs containing it shall be '	introduced onto the	
	Ready to eat cereals	9 g/100 g		market as a food supplement	
	Dairy products	2,4 g/100 g	Wheat bran extract '	or food supplement ingredient. Nor may it be added to infant formula.	
_	Fruit and vegetable juices	0,6 g/100 g			
	Soft drinks	0,6 g/100 g			
	Meat preparations	2 g/100 g			
[^{F49} Xylo- oligosaccharic	Specified lefood category	Maximum levels ^j	The designation of the novel		
	White bread	14 g/kg	food on the		
	Wholemeal bread	14 g/kg	labelling of the foodstuffs containing		
	Breakfast cereals	14 g/kg	it shall be 'Xylo-		
	Biscuits	14 g/kg	oligosaccharid	58	
	Soy drink	3,5 g/kg			
	Yoghurt i	3,5 g/kg			
	Fruit spreads	30 g/kg			
	Chocolate confectionery	30 g/kg			
	Food supplements as defined in Directive 2002/46/ EC for the general adult population	2 g/day]			

[F50 Yarrowia lipolytica yeast biomass	Specified food category Food Supplements as defined in Directive 2002/46/EC, excluding food supplements for infants and young children	Maximum levels 6 g/day for children from 10 years of age, adolescents and general adult population 3 g/day for children from 3 to 9 years of age]	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Yarrowia lipolytica yeast heat-killed biomass'		
[F513'- Sialyllactose (3'-SL) sodium salt (microbial source)	Specified food category Unflavoured pasteurised and sterilised (including UHT) milk products	Maximum levels (expressed as 3'- Sialyllactose) 0.25 g/L l unflavoured	the foodstuffs containing it is '3'- Sialyllactose sodium salt'. The labelling	Included in the list on 30th June 2022. This inclusion is based on proprietary scientific evidence and scientific data protected in	
	Flavoured fermented milk-based products including heat-treated products Unflavoured	0.25 g/L (beverages) 2.5g/kg (products other than beverages)	of food supplements containing 3'- Sialyllactose sodium salt must bear a statement that they should not	supplements containing 3'- Sialyllactose sodium salt must bear a statement that they should not	accordance with Article 26 of Regulation 2015/2283. Applicant: Glycom A/S of Kogle Allé 4, DK-2970 Hørsholm,
	fermented milk-based products	(beverages) 0.5g/ kg (products other than beverages)	be consumed: (a) if foods containing added 3'- Sialyllactose sodium	Denmark. During the period of data protection, 3'-Sialyllactose	
	Beverages (flavoured drinks, excluding drinks with a pH less than 5)	0.25 g/L	salt are consumed the same day (b) by infants and young children.	sodium salt is authorised for placing on the market within England only by Glycom A/S unless a	
	Cereal bars Infant formula as	2.5g/kg 0.2 g/L in the final product		subsequent applicant obtains authorisation	

defined in Regulation 609/2013	ready for use, marketed as such or reconstituted as instructed by the manufacturer
Follow-on formula as defined in Regulation 609/2013	0.15 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer
Processed cereal-based food and baby food for infants and young children as defined in Regulation 609/2013	0.15 g/L (beverages) in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer. 1.25 g/kg for products other than beverages
Milk-based drinks and similar products intended for young children	0.15 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer
Total diet replacement fo weight control as defined in Regulation 609/2013	0.5 g/L o(theforages) 5g/ kg (products other than beverages)

for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation 2015/2283 or with the agreement of Glycom A/S. The data protection will expire at the end of 29th June 2027.]

	Food for special medical purposes as defined in Regulation 609/2013 Food Supplements as defined the Food Supplements (England) Regulations 2003, excluding food supplements for infants and young children				
[F523'- Sialyllactose (3'-SL) sodium salt (microbial source)	Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products Flavoured fermented milk-based products including heat-treated products Unflavoured fermented milk-based products Beverages	Maximum levels (expressed as 3'-Sialyllactose) 0.25 g/L 0.25 g/L (beverages) 2.5 g/kg (products other than beverages) 0.25 g/L	The designation of the novel food on the labelling of the foodstuffs containing it is "3'-Sialyllactose sodium salt". The labelling of food supplements containing 3'-Sialyllactose so salt must bear a statement that they should not be consumed: a) if foods containing added 3'-Sialyllactose sodium salt are consumed the same day, b) by infants	odium	Included in the list on 30 June 2022. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Glycom A/S, Kogle Allé 4, DK-2970 Hørsholm, Denmark. During the period of data protection, the novel food 3'-sialyllactose
	(flavoured drinks, excluding	(beverages)	and young children.		sodium salt is authorised for

drinks with a pH less than 5)	
Cereal bars	0.5 g/kg (products other than beverages)
Infant formula as defined in Regulation (EU) 609/2013	0.25 g/L
Follow-on formula as defined in Regulation (EU) 609/2013	2.5 g/kg
Processed cereal-based food and baby food for infants and young children as defined in Regulation (EU) 609/2013	0.2 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer
Milk-based drinks and similar products intended for young children	0.15 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer
Total diet replacement foods for weight control as defined in Regulation (EU) 609/2013	0.15 g/L (beverages) in the final product ready for use, marketed as such or reconstituted as instructed

placing on the market within Scotland only by Glycom A/S, unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Glycom A/S. The data protection will expire at the end of 29 June 2027.]

	Food for special medical purposes as defined in Regulation (EU) 609/2013	by the manufacturer 1.25 g/kg for products other than beverages		
	Food supplements as defined in the Food Supplements (Scotland) Regulations 2003, excluding food supplements for infants and young children	0.15 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer		
[F533'- Sialyllactose (3'-SL) sodium salt (microbial source)	Unflavoured pasteurised and sterilised (including UHT) milk products	Maximum levels (expressed as 3'- Sialyllactose) 0.25 g/L l unflavoured	The designation of the novel food on the labelling of the foodstuffs containing it is "3'-Sialyllactose sodium salt". The labelling of food	Included in the list on 30 June 2022. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article
	Flavoured fermented milk-based products including heat-treated products	0.25 g/L (beverages) 2.5g/kg (products other than beverages)	supplements containing 3'- Sialyllactose sodium salt must bear a statement that they	26 of Regulation (EU) 2015/2283. Applicant: Glycom A/ S, Kogle Alle
	Unflavoured fermented milk-based products	0.25 g/L (beverages) 0.5 g/kg (products other than beverages)	should not be consumed: a) if foods containing added 3'- Sialyllactose sodium salt are	4, DK-2970 Horsholm, Denmark. During the period of data protection, 3'- Sialyllactose sodium salt is

Beverages (flavoured drinks, excluding drinks with a pH less than 5)	0.25 g/L	consumed the same day; b) by infants and young children.
Cereal bars	2.5 g/kg	
Infant formula as defined in Regulation (EU) No 609/2013	0.2 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer	
Follow-on formula as defined in Regulation (EU) No 609/2013	0.15 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer	
Processed cereal-based food and baby food for infants and young children as defined in Regulation (EU) No 609/2013	0.15 g/L (beverages) in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer. 1.25 g/kg (products other than beverages)	
Milk-based drinks and similar products intended for young children	0.15 g/L in the final product ready for use, marketed as such or reconstituted as instructed	

authorised for placing on the market within Wales only by Glycom A/S unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Glycom A/S. The data protection ends at the end of 29 June 2027.]

	Total diet replacement foods for weight control as defined in Regulation (EU) No 609/2013	by the manufacturer 0.5 g/L (beverages) 5g/kg (products other than beverages)		
	Food for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended		
	Food supplements as defined in the Food Supplements (Wales) Regulations 2003, excluding food supplements for infants and young children	0.5 g/day		
[F516'- Sialyllactose (6'-SL) sodium salt (microbial	Specified food category	Maximum levels (expressed as 6'- Sialyllactose)	The designation of the novel food on the labelling of	Included in the list on 30th June 2022. This inclusion
source)	Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products Unflavoured fermented	0.5 g/L (beverages)	the foodstuffs containing it is '6'- Sialyllactose sodium salt'. The labelling of food supplements containing 6'- Sialyllactose sodium salt must bear	is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation 2015/2283.

Flavoured fermented milk-based products including heat-treated products	2.5g/kg (products other than beverages) 0.5 g/L (beverages) 5.0 g/kg (products other than beverages) 0.5 g/L	a sta that show be co (a) in cont adde Sialy sodir salt a cons same (b) b
(flavoured drinks, excluding drinks with a PH less than 5)	0.0 g 2	and y
Cereal bars	5.0 g/kg	
Infant formula as defined in Regulation 609/2013	0.4 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer	
Follow-on formula as defined in Regulation 609/2013	0.3 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer	
Processed cereal-based food and baby food for infants and young children as defined in Regulation 609/2013	0.3 g/L (beverages) in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer 2.5 g/kg for products	

a statement that they should not be consumed:
(a) if foods containing added 6'-Sialyllactose sodium salt are consumed the same day
(b) by infants and young children.

Applicant: Glycom A/S of Kogle Allé 4, DK-2970 Hørsholm. Denmark. During the period of data protection, 6'-Sialyllactose sodium salt is authorised for placing on the market within England only by Glycom A/S unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation 2015/2283 or with the agreement of Glycom A/S. The data protection will expire at the end of 29th June 2027.]

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	Milk based drinks and similar products intended for young children	other than beverages 0.3 g/L (beverages) in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer		
	Total diet replacement foods for weight control as defined in Regulation 609/2013	1.0 g/L (beverages) 10.0 g/kg (products other than beverages)		
	Food for special medical purposes as defined in Regulation 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended		
	Food Supplements as defined in the Food Supplements (England) Regulations 2003, excluding food supplements for infants and young children	1.0 g/day		
[F546'- Sialyllactose (6'-SL) sodium salt (microbial source)	Specified food category	Maximum levels (expressed as 6'- Sialyllactose)	The designation of the novel food on the labelling of the foodstuffs	Included in the list on 30 June 2022. This inclusion is based on proprietary

Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products	0.5 g/L
Unflavoured fermented milk-based products	0.5 g/L (beverages)
Flavoured fermented milk-based products including heat-treated products	2.5 g/kg (products other than beverages)
Beverages (flavoured drinks, excluding drinks with a pH less than 5)	0.5 g/L (beverages)
Cereal bars	5.0 g/kg (products other than beverages)
Infant formula as defined in Regulation (EU) 609/2013	0.5 g/L
Follow-on formula as defined in Regulation (EU) 609/2013	5.0 g/kg
Processed cereal-based food and baby food for infants and young children as	0.4 g/L in the final product ready for use, marketed as such or reconstituted as instructed

containing it is '6'-Sialyllactose sodium salt'. The labelling of food supplements containing 6'-Sialyllactose (6'-SL) sodium salt must bear a statement that they should not be consumed: (a) if foods containing added 6'-Sialyllactose sodium salt are consumed on the same day, (b) by infants and young children.

scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Glycom A/ S, Kogle Allé 4, DK-2970 Hørsholm. Denmark. During the period of data protection, the novel food 6'sialyllactose sodium salt is authorised for placing on the market within Scotland only by Glycom A/S, unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Glycom A/S.

	defined in Regulation (EU) 609/2013	by the manufacturer		The data protection will expire at the end of
	Milk based drinks and similar products intended for young children	0.3 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer		29June 2027.]
	Total diet replacement foods for weight control as defined in Regulation (EU) 609/2013	0.3 g/L (beverages) in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer		
	Food for special medical purposes as defined in Regulation (EU) 609/2013	2.5 g/kg for products other than beverages		
	Food supplements as defined in the Food Supplements (Scotland) Regulations 2003, excluding food supplements for infants and young children	0.3 g/L (beverages) in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer		
F ⁵⁵ "6'- Sialyllactose (6'-SL) sodium salt	Specified food category	Maximum levels (expressed	The designation of the novel food on the	Included in the list on 30 June 2022.

(microbial source)

	as 6'- Sialyllactose)
Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products	0.5 g/L
Unflavoured fermented milk-based products	0.5 g/L (beverages) 2.5 g/kg (products other than beverages)
Flavoured fermented milk-based products including heat-treated products	0.5 g/L (beverages) 5.0 g/kg (products other than beverages)
Beverages (flavoured drinks, excluding drinks with a PH less than 5)	0.5 g/L
Cereal bars	5.0 g/kg
Infant formula as defined in Regulation (EU) No 609/2013	0.4 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer
Follow-on formula as defined in Regulation (EU) No 609/2013	0.3 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer

labelling of the foodstuffs containing it is "6'-Sialyllactose sodium salt". The labelling of food supplements containing 6'-Sialyllactose sodium salt must bear a statement that they should not be consumed: a) if foods containing added 6'-Sialyllactose sodium salt are consumed the same day; b) by infants and young children.

This inclusion is authorised based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Glycom A/ S, Kogle Alle 4, DK-2970 Horsholm. Denmark. During the period of data protection, 6'-Sialyllactose sodium salt is authorised for placing on the market within Wales only by Glycom A/S unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Glycom A/S.

Processed cereal-based food and baby food for infants and young children as defined in Regulation (EU) No 609/2013	0.3 g/L (beverages) in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer 2.5 g/kg (products other than beverages)
Milk based drinks and similar products intended for young children	0.3 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer
Total diet replacement foods for weight control as defined in Regulation (EU) No 609/2013	1.0 g/L (beverages) 10.0 g/kg (products other than beverages)
Food for special medical purposes as defined under Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended
Food supplements as defined in the Food Supplements (Wales) Regulations 2003,	1.0 g/day

The data protection ends at the end of 29 June 2027.]

Yeast beta- glucans	excluding food supplements for infants and young children Specified food category	1.0 g/L (beverages)	The designation of the novel	
	Food supplements as defined in Directive 2002/46/EC, excluding food supplements for infants and young children	10.0 g/kg (products other than beverages)	food on the labelling of the foodstuffs containing it shall be 'Yeast (Saccharomyces cerevisiae) beta-glucans'	
	Total diet replacement for weight control as defined in Regulation (EU) No 609/2013	In accordance with the particular nutritional requirements of the persons for whom the products are intended		
	Food for special medical purposes as defined in Regulation (EU) No 609/2013, excluding food for special medical purposes intended for infants and young children	1.0 g/day.		
	Beverages based on fruit and/or vegetable juices	1,3 g/kg		

including concentrate and dehydrated juices	
Fruit- flavoured drinks	0,8 g/kg
Cocoa beverages preparation powder	38,3 g/kg (powder)
Other beverages	0,8 g/kg (ready to drink)
	(powder)
Cereal bars	6 g/kg
Breakfast cereals	15,3 g/kg
Wholegrain and high fibre instant hot breakfast cereals	1,5 g/kg
Cookie-type biscuits	6,7 g/kg
Cracker-type biscuits	6,7 g/kg
Milk based beverages	3,8 g/kg
Fermented milk products	3,8 g/kg
Milk product analogues	3,8 g/kg
Dried milk/ milk powder	25,5 g/kg
Soups and soup mixes	0,9 g/kg (ready to eat)
	1,8 g/kg (condensed)
	6,3 g/kg (powder)

F ⁵⁶ Zeaxanthir	Chocolate and confectionery Protein bars and powders Jam, marmalade and other fruit spreads Specified	4 g/kg 19,1 g/kg 11,3 g/kg Maximum	The	
	Food Supplements as defined in Directive 2002/46/EC	levels 2 mg/day]	designation of the novel food on the labelling of the foodstuffs containing it shall be 'Zeaxanthin'.	
Zinc L-pidolate	Specified food category Foods covered by Regulation (EU) No 609/2013 Milk based drinks and similar products intended for young children Meal replacement for weight control Foods intended to meet the expenditure of intense muscular effort, especially for sportsmen Food bearing statement on	Maximum levels 3 g/day	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Zinc L-pidolate'	

ANNEX

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Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

the absence or reduced presence of gluten in accordance with the requirements of Commission Implementing Regulation (EU) No 828/2014		
Food Supplements as defined in Directive 2002/46/EC]		

- a Regulation (EU) No 609/2013 of the European Parliament and of the Council of 12 June 2013 on food intended for infants and young children, food for special medical purposes, and total diet replacement for weight control and repealing Council Directive 92/52/EEC, Commission Directives 96/8/EC, 1999/21/EC, 2006/125/EC and 2006/141/EC, Directive 2009/39/EC of the European Parliament and of the Council and Commission Regulations (EC) No 41/2009 and (EC) No 953/2009 (OJ L 181, 29.6.2013, p. 35).
- b Commission Implementing Regulation (EU) No 828/2014 of 30 July 2014 on the requirements for the provision of information to consumers on the absence or reduced presence of gluten in food (OJ L 228, 31.7.2014, p. 5).
- c Directive 2002/46/EC of the European Parliament and of the Council of 10 June 2002 on the approximation of the laws of the Member States relating to food supplements (OJ L 183, 12.7.2002, p. 51).
- d Regulation (EC) No 1925/2006 of the European Parliament and of the Council of 20 December 2006 on the addition of vitamins and minerals and of certain other substances to foods (OJ L 404, 30.12.2006, p. 26).
- e Council Directive 2001/113/EC of 20 December 2001 relating to fruit jams, jellies and marmalades and sweetened chestnut purée intended for human consumption (OJ L 10, 12.1.2002, p. 67).
- f Regulation (EU) No 1308/2013 of the European Parliament and of the Council of 17 December 2013 establishing a common organisation of the markets in agricultural products and repealing Council Regulation (EEC) No 922/72, (EEC) No 234/79, (EC) No 1037/2001 and (EC) 1234/2007 (OJ L 347, 20.12.2013, p. 671).
- g [F4Maximum use levels in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer.]
- h [FSCouncil Directive 2001/112/EC of 20 December 2001 relating to fruit juices and certain similar products intended for human consumption (OJ L 10, 12.1.2002, p. 58).]
- i [F6When used in milk products xylo-oligosaccharides shall not replace, in whole or in part, any milk constituent.
- j Maximum levels calculated on the basis of the specifications of Powder form 1.]
- k [F7The minimum specification for vitamin D content in vitamin D $_2$ mushroom powder of 1 000 μ g vitamin D $_2$ /gram of mushroom powder is used.]

Textual Amendments

- **F4** Inserted by Commission Implementing Regulation (EU) 2019/1294 of 1 August 2019 authorising the placing on the market of betaine as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- F5 Inserted by Commission Implementing Regulation (EU) 2020/500 of 6 April 2020 authorising the placing on the market of partially defatted chia seed (Salvia hispanica) powders as novel foods under

- Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F6** Inserted by Commission Implementing Regulation (EU) 2020/916 of 1 July 2020 authorising the extension of use of xylo-oligosaccharides as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- F7 Inserted by Commission Implementing Regulation (EU) 2020/1163 of 6 August 2020 authorising the placing on the market of vitamin D2 mushroom powder as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F8** Inserted by Commission Implementing Regulation (EU) 2019/506 of 26 March 2019 authorising the placing on the market of D-ribose as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council, and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F9** Substituted by Commission Implementing Regulation (EU) 2019/110 of 24 January 2019 authorising an extension of use of Allanblackia seed oil as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F10** Substituted by Commission Implementing Regulation (EU) 2019/1686 of 8 October 2019 authorising the extension of use of bovine milk basic whey protein isolate as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- F11 Substituted by Commission Implementing Regulation (EU) 2020/24 of 13 January 2020 authorising an extension of use of chia seeds (Salvia hispanica) as a novel food and the change of the conditions of use and the specific labelling requirements of chia seeds (Salvia hispanica) under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- F12 Inserted by Commission Implementing Regulation (EU) 2018/1631 of 30 October 2018 authorising the placing on the market of cranberry extract powder as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F13** Inserted by Commission Implementing Regulation (EU) 2018/2016 of 18 December 2018 authorising the placing on the market of decorticated grains of Digitaria exilis as a traditional food from a third country under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F14** Inserted by Commission Implementing Regulation (EU) 2018/1133 of 13 August 2018 authorising the placing on the market of dried aerial parts of Hoodia parviflora as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- F15 Substituted by Commission Implementing Regulation (EU) 2019/1272 of 29 July 2019 correcting Implementing Regulation (EU) 2017/2470 establishing the Union list of novel foods and Implementing Decision (EU) 2017/2078 authorising an extension of use of yeast beta-glucans as a novel food ingredient under Regulation (EC) No 258/97 of the European Parliament and of the Council (Text with EEA relevance).
- F16 Inserted by Commission Implementing Regulation (EU) 2020/1559 of 26 October 2020 amending Implementing Regulation (EU) 2017/2470 establishing the Union list of novel foods (Text with EEA relevance).
- F17 Inserted by Commission Implementing Regulation (EU) 2018/1647 of 31 October 2018 authorising the placing on the market of egg membrane hydrolysate as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).

- F18 Substituted by Commission Implementing Regulation (EU) 2020/1559 of 26 October 2020 amending Implementing Regulation (EU) 2017/2470 establishing the Union list of novel foods (Text with EEA relevance).
- F19 Inserted by Commission Implementing Regulation (EU) 2020/206 of 14 February 2020 authorising the placing on the market of fruit pulp, pulp juice, concentrated pulp juice from Theobroma cacao L. as a traditional food from a third country under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F20** Inserted by Commission implementing Regulation (EU) 2019/1979 of 26 November 2019 authorising the placing on the market of 2'-Fucosyllactose/Difucosyllactose mixture as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- F21 Words in Annex Table 1 inserted (E.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (England) Regulations 2022 (S.I. 2022/560), regs. 1(3), 2(2)(a)(i); words inserted (S.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Scotland) Regulations 2022 (S.S.I. 2022/168), reg. 1(3), sch. 1 para. 1; and words inserted (W.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Wales) Regulations 2022 (S.I. 2022/575), reg. 1(4), Sch. 1 para. 1
- F22 Words in Annex Table 1 inserted (E.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (England) Regulations 2022 (S.I. 2022/560), regs. 1(3), 2(2)(a)(ii); words inserted (S.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Scotland) Regulations 2022 (S.S.I. 2022/168), reg. 1(3), sch. 1 para. 1; and words inserted (W.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Wales) Regulations 2022 (S.I. 2022/575), reg. 1(4), Sch. 1 para. 1
- F23 Inserted by Commission Implementing Regulation (EU) 2020/917 of 1 July 2020 authorising the placing on the market of infusion from coffee leaves of Coffea arabica L. and/or Coffea canephora Pierre ex A. Froehner as a traditional food from a third country under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F24** Substituted by Commission Implementing Regulation (EU) 2018/1293 of 26 September 2018 amending Implementing Regulation (EU) 2017/2470 as regards the conditions of use of the novel food lactitol (Text with EEA relevance).
- F25 Inserted by Commission Implementing Regulation (EU) 2020/484 of 2 April 2020 authorising the placing on the market of lacto-N-tetraose as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- F26 Inserted by Commission Implementing Regulation (EU) 2018/1991 of 13 December 2018 authorising the placing on the market of berries of Lonicera caerulea L. as a traditional food from a third country under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F27** Inserted by Commission Implementing Regulation (EU) 2018/1123 of 10 August 2018 authorising the placing on the market of 1-methylnicotinamide chloride as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F28** Inserted by Commission Implementing Regulation (EU) 2020/16 of 10 January 2020 authorising the placing on the market of nicotinamide riboside chloride as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F29** Inserted by Commission Implementing Regulation (EU) 2019/1976 of 25 November 2019 authorising the placing on the market of Phenylcapsaicin as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).

- **F30** Substituted by Commission Implementing Regulation (EU) 2020/973 of 6 July 2020 authorising a change of the conditions of use of the novel food 'protein extract from pig kidneys' and amending Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F31** Inserted by Commission Implementing Regulation (EU) 2018/1122 of 10 August 2018 authorising the placing on the market of pyrroloquinoline quinone disodium salt as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- F32 Inserted by Commission Implementing Regulation (EU) 2018/1633 of 30 October 2018 authorising the placing on the market of refined shrimp peptide concentrate as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F33** Substituted by Commission Implementing Regulation (EU) 2019/387 of 11 March 2019 authorising an extension of use of Schizochytrium sp. (ATCC PTA-9695) oil as a novel food and the change of the designation and of the specific labelling requirement of Schizochytrium sp. (ATCC PTA-9695) oil under Regulation (EU) 2015/2283 of the European Parliament and of the Council, and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F34** Words in Annex Table 1 inserted (E.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (England) Regulations 2022 (S.I. 2022/560), reg. 1(3), **Sch. 1**
- **F35** Words in Annex Table 1 inserted (S.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Scotland) Regulations 2022 (S.S.I. 2022/168), reg. 1(3), sch. 2 para. 1
- **F36** Words in Annex Table 1 inserted (W.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Wales) Regulations 2022 (S.I. 2022/575), reg. 1(4), **Sch. 2 para. 1**
- **F37** Substituted by Commission Implementing Regulation (EU) 2019/109 of 24 January 2019 authorising an extension of use of Schizochytrium sp. oil as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council, and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- F38 Words in Annex Table 1 inserted (E.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (England) Regulations 2022 (S.I. 2022/560), reg. 1(3), Sch. 2
- **F39** Words in Annex Table 1 inserted (S.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Scotland) Regulations 2022 (S.S.I. 2022/168), reg. 1(3), sch. 3 para. 1
- **F40** Words in Annex Table 1 inserted (W.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Wales) Regulations 2022 (S.I. 2022/575), reg. 1(4), **Sch. 3 para. 1**
- **F41** Inserted by Commission Implementing Regulation (EU) 2018/2017 of 18 December 2018 authorising the placing on the market of syrup from Sorghum bicolor (L.) Moench as a traditional food from a third country under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F42** Inserted by Commission Implementing Regulation (EU) 2020/1634 of 4 November 2020 authorising the placing on the market of sugars obtained from cocoa (Theobroma cacao L.) pulp as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F43** Words in Annex Table 1 substituted (E.) (15.5.2023) by The Food Additives, Food Flavourings and Novel Foods (Authorisations) (England) Regulations 2023 (S.I. 2023/334), reg. 1(1), **Sch. 4** (with reg. 4)
- F44 Words in Annex Table 1 substituted (S.) (15.5.2023) by The Food Additives, Food Flavourings and Novel Foods (Authorisations) (Scotland) Regulations 2023 (S.S.I. 2023/78), reg. 1(1), sch. 4 para. 1 (with reg. 5)

Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

- F45 Words in Annex Table 1 substituted (W.) (15.5.2023) by The Food Additives, Food Flavourings, and Novel Foods (Authorisations) and Food and Feed (Miscellaneous Amendments) (Wales) Regulations 2023 (S.I. 2023/343), reg. 1(4), Sch. 4 para. 2 (with reg. 4)
- **F46** Words in Annex Table 1 inserted (E.) (15.5.2023) by The Food Additives, Food Flavourings and Novel Foods (Authorisations) (England) Regulations 2023 (S.I. 2023/334), reg. 1(1), **Sch. 5** (with reg. 4)
- F47 Words in Annex Table 1 inserted (S.) (15.5.2023) by The Food Additives, Food Flavourings and Novel Foods (Authorisations) (Scotland) Regulations 2023 (S.S.I. 2023/78), reg. 1(1), sch. 5 para. 1 (with reg. 5)
- **F48** Words in Annex Table 1 inserted (W.) (15.5.2023) by The Food Additives, Food Flavourings, and Novel Foods (Authorisations) and Food and Feed (Miscellaneous Amendments) (Wales) Regulations 2023 (S.I. 2023/343), reg. 1(4), **Sch. 5 para. 2** (with reg. 4)
- **F49** Substituted by Commission Implementing Regulation (EU) 2020/916 of 1 July 2020 authorising the extension of use of xylo-oligosaccharides as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F50** Inserted by Commission Implementing Regulation (EU) 2019/760 of 13 May 2019 authorising the placing on the market of Yarrowia lipolytica yeast biomass as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F51** Words in Annex Table 1 inserted (E.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (England) Regulations 2022 (S.I. 2022/560), reg. 1(3), **Sch. 3** (as amended by S.I. 2022/619, regs. 1(1), **2(2)**)
- **F52** Words in Annex Table 1 inserted (S.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Scotland) Regulations 2022 (S.S.I. 2022/168), reg. 1(3), sch. 4 para. 1
- F53 Words in Annex Table 1 inserted (W.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Wales) Regulations 2022 (S.I. 2022/575), reg. 1(4), Sch. 4 para. 1
- **F54** Words in Annex Table 1 inserted (S.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Scotland) Regulations 2022 (S.S.I. 2022/168), reg. 1(3), sch. 5 para. 1
- F55 Words in Annex Table 1 inserted (W.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Wales) Regulations 2022 (S.I. 2022/575), reg. 1(4), Sch. 5 para. 1
- F56 Substituted by Commission Implementing Regulation (EU) 2018/1132 of 13 August 2018 authorising the change of the designation and specific labelling requirement of the novel food synthetic zeaxanthin under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).

TABLE 2: SPECIFICATIONS

Authorised Novel Food	Specifications
N -Acetyl-D-	Description:
neuraminic acid	
	Definition:
	Chemical name:
	IUPAC names:
	N -Acetyl-D-neuraminic acid (dihydrate)
	5-Acetamido-3,5-dideoxy-D-glycero-D-galacto-non-2-ulopyranosonic
	acid (dihydrate)
	Synonyms:

Sialic acid (dihydrate)

Chemical formula:

C₁₁ H₁₉ NO₉ (acid)

C₁₁ H₂₃ NO₁₁ (C₁₁ H₁₉ NO₉ * 2H₂O) (dihydrate)

Molecular mass:

309,3 Da (acid)

345,3 (309,3 + 36,0) (dihydrate)

CAS No.:

131-48-6 (free acid)

50795-27-2 (dihydrate)

Specifications:

Description: white to off-white crystalline powder

pH (20 °C, 5 % solution): 1,7 – 2,5

N-Acetyl-D-neuraminic acid (dihydrate): > 97.0 %

Water (dihydrate calculates to 10.4 %): $\leq 12.5 \%$ (w/w)

Ash, sulphated: < 0.2 % (w/w)

Acetic acid (as free acid and/or sodium acetate): < 0.5 % (w/w)

Heavy Metals:

Iron: < 20.0 mg/kg

Lead: < 0.1 mg/kg

Residual proteins: < 0.01 % (w/w) **Residual solvents:**

2-Propanol: < 0.1 % (w/w)

Acetone: < 0.1 % (w/w)

Ethyl acetate: < 0.1 % (w/w)

Microbiological criteria:

Salmonella: Absence in 25 g

Aerobic mesophilic total count: < 500 CFU/g

Enterobacteriaceae: Absence in 10 g

Cronobacter (Enterobacter) sakazakii: Absence in 10 g

Listeria monocytogenes: Absence in 25 g

Bacillus cereus: < 50 CFU/g

Yeasts: < 10 CFU/g Moulds: < 10 CFU/g

Residual endotoxins: < 10 EU/mg

CFU: Colony Forming Units; EU: Endotoxin Units.

Adansonia digitata (Baobab) dried fruit pulp

Description/Definition:

The Baobab (*Adansonia digitata*) fruits are harvested from trees. The hard shells are cracked open and the pulp is separated from the seeds and the shell. This is milled, separated into coarse and fine lots (particle size 3 to 600 µ) and then packaged.

Typical nutritional components:

Moisture (loss on drying) (g/100 g): 4,5-13,7

Protein (g/100 g): 1,8-9,3

Fat (g/100 g): 0-1,6

Total carbohydrate (g/100 g): 76,3-89,5

Total sugars (as glucose): 15,2-36,5

Sodium (mg/100 g): 0,1-25,2

Analytical specifications:

Foreign matter: Not more than 0,2 %

Moisture (loss on drying) (g/100 g): 4,5-13,7

Ash (g/100 g): 3,8-6,6

Ajuga reptans extract from cell cultures	Description/Definition: Hydroalcoholic extract from <i>Ajuga reptans</i> L. tissue cultures which is substantially equivalent to extracts from flowering aerial parts of <i>Ajuga reptans</i> obtained by traditional cultures.
L-Alanyl-L-Glutamine	Description/Definition: L-Alanyl-L-Glutamine is produced by fermentation with a genetically modified strain of <i>Escherichia coli</i> . During the fermentation process, the ingredient is secreted into the growth medium from which it is subsequently separated and purified to a concentration of > 98 %. Appearance: White crystalline powder Purity: > 98 % Infrared spectroscopy: Conformity with ref. standard Appearance of solution: Colourless and clear Assay (dry basis): 98-102 % Related substances (each): ≤ 0,2 % Residue on ignition: ≤ 0,1 % Loss on drying: ≤ 0,5 % Optical rotation: +9,0 - +11,0 ° pH (1 %; H $_2$ O): 5,0-6,0 Ammonium (NH $_4$): ≤ 0,020 % Chloride (Cl): ≤ 0,020 % Sulphate (SO $_4$): ≤ 0,020 % Microbiological criteria: <i>Escherichia coli</i> : Absence/g
Algal oil from the microalgae <i>Ulkenia</i> sp.	Description/Definition: Oil from the micro-algae <i>Ulkenia</i> sp. Acid value: ≤ 0.5 mg KOH/g Peroxide value (PV): ≤ 5.0 meq/kg oil Moisture and volatiles: ≤ 0.05 % Unsaponifiables: ≤ 4.5 % Trans-fatty acids: ≤ 1.0 % DHA content: ≥ 32 %
[^{F9} Allanblackia seed oil	Description/Definition: Allanblackia seed oil is obtained from the seeds of the allanblackia species: A. floribunda (synonymous with A. parviflora) and A. stuhlmannii. Composition of fatty acids (as a % of the total fatty acids): Lauric acid — Myristic acid — Palmitic acid (C12:0 – C14:0 – C16:0): sum of these acids < 4,0 % Stearic acid (C18:0): 45-58 % Oleic acid (C18:1): 40-51 % Poly unsaturated fatty acids (PUFA): < 2 % Characteristics: Free fatty acids: max 0,1 % of total fatty acids Trans fatty acids: max 1,0 % of total fatty acids Peroxide value: max 1,0 meq/kg Unsaponifiable matter: max 1,0 % (w/w) of the oil Saponification value: 185-198 mg KOH/g]
Aloe macroclada	Description/Definition:

Baker leaf extract

Powdered gel extract derived from the leaves of *Aloe macroclada* Baker which is substantially equivalent to the same gel derived from *Aloe vera*

(L.) Burm.f. leaves.

Ash: 25 %

Dietary fibres: 28,6 %

Fat: 2,7 %

Moisture: 4,7 %

Polysaccharides: 9,5 % Protein: 1,63 %

Protein: 1,63 % Glucose: 8,9 %

[F58] Antarctic Krill oil from Euphausia superba

Description/Definition:

To produce lipid extract from Antarctic Krill (*Euphausia superba*) deepfrozen crushed krill or dried krill meal is subjected to lipid extraction with an approved extraction solvent (under Directive 2009/32/EC). Proteins and krill material are removed from the lipid extract by filtration. The extraction solvents and residual water are removed by evaporation.

Saponification value: $\leq 230 \text{ mg KOH/g}$ Peroxide value (PV): $\leq 3 \text{ meq O}_2/\text{kg oil}$

Oxidative stability: All food products containing Antarctic Krill oil from *Euphausia superba* should demonstrate oxidative stability by appropriate and recognised national/international test methodology (e.g. AOAC). Moisture and volatiles: ≤ 3 % or 0,6 expressed as water activity at 25 °C

Phospholipids: $\geq 35 \%$ to < 60 %

Trans-fatty acids: $\leq 1 \%$

EPA (eicosapentaenoic acid): $\geq 9 \%$ DHA (docosahexaenoic acid): $\geq 5 \%$]

Antarctic Krill oil rich in phospholipids from Euphausia superba

Description/Definition:

Oil rich in phospholipids is produced from Antarctic krill (*Euphausia superba*) by repeated solvent washings with an approved solvent (under Directive 2009/32/EC) to increase phospholipid content of the oil. Solvents are removed from the final product by evaporation.

Saponification value: $\leq 230 \text{ mg KOH/g}$ Peroxide value (PV): $\leq 3 \text{ meq O}_2/\text{kg oil}$

Moisture and volatiles: ≤ 3 % or 0,6 expressed as water activity at 25 °C

Phospholipids: $\geq 60 \%$ Trans-fatty acids: $\leq 1 \%$

EPA (eicosapentaenoic acid): $\geq 9 \%$ DHA (docosahexaenoic acid): $\geq 5 \%$

Arachidonic acid-rich oil from the fungus *Mortierella alpina*

Description/Definition:

The clear yellow arachidonic acid-rich oil is obtained by fermentation of the non-genetically modified strains IS-4, I49-N18, FJRK-MA01 and CBS 210.32 of the fungus *Mortierella alpina* using a suitable liquid. The oil is then extracted from the biomass and purified.

Arachidonic acid: ≥ 40 % by weight of the total fatty acid content

Free fatty acids: ≤ 0.45 % of the total fatty acid content

Trans fatty acids: $\leq 0.43\%$ of the total fatty acid content

Unsaponifiable matter: ≤ 1.5 % Peroxide value (PV): ≤ 5 meq/kg

Anisidin value: ≤ 20 Acid value: $\leq 1,0$ KOH/g Moisture: $\leq 0,5$ %

Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

Argan oil from Argania spinosa

Description/Definition:

Argan oil is the oil obtained by cold pressing of the almond like kernels of the fruits of *Argania spinosa* (L.) Skeels. Kernels may be roasted prior to pressing, but with no direct contact with a flame.

Composition:

Palmitic acid (C16:0): 12-15 % Stearic acid (C18:0): 5-7 % Oleic acid (C18:1): 43-50 % Linoleic acid (C18:2): 29-36 % Unsaponifiable matter: 0,3-2 % Total sterols: 100-500 mg/100 g Total tocopherols: 16-90 mg/100 g

Oleic acidity: 0,2-1,5 %

Peroxide value (PV): $\leq 10 \text{ meq O}_2/\text{kg}$

Astaxanthinrich oleoresin from *Haematococcus* pluvialis algae

Description/Definition:

Astaxanthin is a carotenoid produced by *Haematococcus pluvialis* algae. Production methods for the growth of the algae are variable; using either closed systems exposed to sunlight or strictly controlled illuminated light; alternatively open ponds may be used. The algal cells are harvested and dried; the oleoresin is extracted using either super critical CO ₂ or a solvent (ethyl acetate). The Astaxanthin is diluted and standardized to 2,5 %, 5,0 %, 7,0 %, 10 %, 15 % or 20 % using olive oil, safflower oil, Sunflower oil or MCT (Medium Chain Triglycerides).

Composition of the Oleoresin:

Fat: 42,2- 99 % Protein: 0,3-4,4 % Carbohydrate: 0-52,8 %

Fibre: < 1,0 % Ash: 0,0-4,2 %

Specification of Carotenoids w/w%

Total Astaxanthins: 2,9-11,1 % 9-cis-astaxanthin: 0,3-17,3 %

13-cis-astaxanthin: 0,2-7,0 %

Astaxanthin monoesters: 79,8-91,5 % Astaxanthin diesters: 0,16-19,0 %

B-Carotene: 0,01-0,3 %

Lutein: 0-1.8 %

Canthaxanthin: 0-1,30 % **Microbiological criteria:**

Total aerobic bacteria: < 3 000 CFU/g Yeast and Moulds: < 100 CFU/g

Coliforms: < 10 CFU/g E. coli: Negative Salmonella: Negative Staphylococcus: Negative

Basil seeds (Ocimum basilicum)

Description/Definition:

Basil (*Ocimum basilicum* L.) belongs to the family ' *Lamiaceae*' within the order 'Lamiales'. Post-harvest the seeds are cleaned mechanically. Flowers, leaves and other parts of the plant are removed. Highest level of purity of Basil seeds has to be ensured by filtering (optical, mechanical). Production process of fruit juice and fruit/vegetable blend beverages containing Basil seeds (*Ocimum basilicum* L.) includes seed

pre-hydration and pasteurisation steps. Microbiological controls and

monitoring systems are in place.

Dry Matter: 94,1 % Protein: 20,7 % Fat: 24,4 %

Carbohydrate: 1,7 %

Dietary Fibre: 40,5 % (Method: AOAC 958,29)

Ash: 6,78 %

[F4Betaine

Description/Definition:

Betaine (N,N,N-trimethylglycine or carboxy-N,N,N-

trimethylmethanaminium), in anhydrous (CH₃)₃N⁺CH₂COO⁻ (CAS

No: 107-43-7) and monohydrate (CH₃)3N⁺CH₂COO⁻.H₂O (CAS No: 590-47-6) forms is obtained from processing of sugar beets (i.e. molasses, vinasses or betaine-glycerol).

Characteristics/Composition

Appearance: Free-flowing white crystals Betaine: \geq 99,0 % (w/w on dry weight basis)

Moisture: $\leq 2.0 \%$ (anhydrous); $\leq 15.0 \%$ (monohydrate)

Ash: $\leq 0.1 \%$ pH: 5,0-7,0

Residual protein: $\leq 1.0 \text{ mg/g}$

Heavy metals:

Arsenic: < 0,1 mg/kg Mercury: < 0,005 mg/kg Cadmium: < 0,01 mg/kg Lead: < 0,05 mg/kg **Microbiological criteria:**

Total viable count: $\leq 100 \text{ CFU/g}$

Coliforms: Negative/10 g
Salmonella sp.: Negative/25 g

Yeast: $\leq 10 \text{ CFU/g}$ Mould: $\leq 10 \text{ CFU/g}$

CFU: Colony Forming Units.]

Fermented black bean extract

Description/Definition:

Fermented black bean extract (Touchi extract) is a fine light-brown protein-rich powder obtained by water extraction of small soybeans (*Glycine max (L.) Merr.*) fermented with *Aspergillus oryzae*. The extract contains an α -glucosidase inhibitor.

Characteristics:

Fat: $\leq 1.0 \%$

Protein: $\geq 55 \%$ Water: $\leq 7.0 \%$

Ash: $\leq 10 \%$

Carbohydrate: ≥ 20 %

α-glucosidase inhibitory activity: IC50 min 0,025 mg/ml

Soy isoflavone: $\leq 0.3 \text{ g/}100 \text{ g}$

Bovine lactoferrin

Description/Definition:

Bovine lactoferrin is a protein that occurs naturally in cows' milk. It is an iron-binding glycoprotein of approximately 77 kDa and consists of a single polypeptide chain of 689 amino acids.

Production process: Bovine lactoferrin is isolated from skimmed milk or cheese whey via ion exchange and subsequent ultra-filtration steps.

Finally, it is dried by freeze drying or spraying and the large particles are sieved out. It is a virtually odourless, light pinkish powder.

Physical-Chemical properties of Bovine lactoferrin:

Moisture: < 4,5 % Ash: < 1,5 %

Arsenic: < 2,0 mg/kg Iron: < 350 mg/kg Protein: > 93 %

of which bovine lactoferrin: > 95 % of which other proteins: < 5,0 % pH (2 % solution, 20 °C): 5,2-7,2

Solubility (2 % solution, 20 °C): complete

[F10Bovine milk basic whey protein isolate

Description

Bovine milk basic whey protein isolate is a yellowish grey powder obtained from bovine skimmed milk via a series of isolation and purification steps.

Characteristics/Composition

Total protein (w/weight of product): ≥ 90 % Lactoferrin (w/weight of product): 25-75 % Lactoperoxidase (w/weight of product): 10-40 % Other proteins (w/weight of product): ≤ 30 %

TGF-β2: 12-18 mg/100 g

Moisture: $\leq 6.0 \%$

pH (5 % solution w/v): 5,5 - 7,6

Lactose: ≤ 3.0 % Fat: ≤ 4.5 % Ash: ≤ 3.5 % Iron: ≤ 25 mg/100 g Heavy Metals

Lead: < 0,1 mg/kg Cadmium: < 0,2 mg/kg Mercury: < 0,6 mg/kg Arsenic: < 0,1 mg/kg Microbiological criteria:

Aerobic mesophilic count: ≤ 10 000 CFU/g

Enterobacteriaceae : ≤ 10 CFU/g *Escherichia coli* : Negative/g

Coagulase positive *Staphylococci*: Negative/g

Salmonella : Negative/25 g Listeria : Negative/25 g

Cronobacter spp.: Negative/25 g

Moulds: $\leq 50 \, \hat{CFU/g}$ Yeasts: $\leq 50 \, CFU/g$

CFU: Colony Forming Units

Buglossoides arvensis seed oil

Description/Definition:

Refined Buglossoides oil is extracted from the seeds of *Buglossoides* arvensis (L.) I.M.Johnst

Alpha-linolenic acid: ≥ 35 % w/w of total fatty acids Stearidonic acid: ≥ 15 % w/w of total fatty acids Linoleic acid: $\geq 8,0$ % w/w of total fatty acids Trans fatty acids: $\leq 2,0$ % w/w of total fatty acids

Acid value: $\leq 0.6 \text{ mg KOH/g}$

Peroxide value (PV): $\leq 5.0 \text{ meq O}_2/\text{kg}$ Unsaponifiable content: $\leq 2.0 \%$

Protein content (total nitrogen): ≤ 10 µg/ml

Pyrrolizidine alkaloids: Not detectable with a detection limit of 4,0 µg/kg

Calanus finmarchicus oil

Description/Definition:

The novel food is ruby coloured, slightly viscous oil with a slight shellfish odour extracted from the crustacean (marine zooplankton) Calanus finmarchicus. The ingredient consists primarily of wax esters (> 85 %) with minor amounts of triglycerides and other neutral lipids.

Specifications:

Water: < 1,0 % Wax esters: > 85 % Total fatty acids: > 46 %

Eicosapentaenoic acid (EPA): > 3,0 % Docosahexaenoic acid (DHA): > 4,0 %

Total fatty alcohols: > 28 % C20:1 n-9 fatty alcohol: > 9,0 % C22:1 n-11 fatty alcohol: > 12 % Trans fatty acids: < 1,0 %

Astaxanthinesters: < 0,1 %

Peroxide value (PV): < 3.0 meq. O₂/kg

Chewing

Description/Definition: The novel food ingredient is a synthetic polymer (Patent gum base

glycol)

(monomethoxypolyethlyde) (2006016179). It consists of branched polymers of monomethoxypolyethylene glycol (MPEG) grafted onto polyisoprenegraft-maleic anhydride (PIP-g-MA), and unreacted MPEG (less than 35

> % by weight). White to off-white colour.

CAS No.: 1246080-53-4 **Characteristics:** Moisture: < 5,0 %

Aluminium: < 3.0 mg/kgLithium: < 0,5 mg/kg Nickel: < 0.5 mg/kg

Residual anhydride: < 15 µmol/g Polydispersity index: < 1,4 Isoprene: < 0.05 mg/kgEthylene oxide: < 0.2 mg/kg

Free maleic anhydride: < 0,1 %Total oligomeres (less than 1 000 Dalton):

 $\leq 50 \text{ mg/kg}$

Ethylene glycol: < 200 mg/kg Diethylene glycol: < 30 mg/kg

Monoethylene glycol methyl ether: < 3,0 mg/kg Diethylene glycol methyl ether: < 4,0 mg/kg Triethylene glycol methyl ether: < 7,0 mg/kg

1,4-Dioxane: < 2,0 mg/kgFormaldehyde: < 10 mg/kg

Chewing gum base (Methyl vinyl ether-maleic

Description/Definition:

Methyl vinyl ether-maleic anhydride copolymer is an anhydrous copolymer of methyl vinyl ether and maleic anhydride.

Free-flowing, white to white-off powder

anhydride copolymer)

CAS No: 9011-16-9

Purity:

Assay value: At least 99,5 % in dry matter Specific viscosity (1 % MEK): 2-10 Residual methyl vinyl ether: ≤ 150 ppm Residual maleic anhydride: ≤ 250 ppm

Acetaldehyde: ≤ 500 ppm Methanol: ≤ 500 ppm Dilauroyl peroxide: ≤ 15 ppm Total heavy metals: ≤ 10 ppm **Microbiological criteria:**

Total aerobic plate count: $\leq 500 \text{ CFU/g}$

Mould/yeast: ≤ 500 CFU/g Escherichia coli : Negative to test Salmonella : Negative to test

Staphylococcus aureus: Negative to test Pseudomonas aeruginosa: Negative to test

Chia oil from Salvia hispanica

Description/Definition:

Chia oil is produced from Chia (Salvia hispanica L.) seeds (99,9 % pure) by cold pressing. No solvents are used and, once pressed, the oil is held in decantation tanks and a filtration process employed to remove impurities. It can also be produced by extraction with supercritical CO $_2$.

Production process:

Produced by cold pressing. No solvents are used and, once pressed, the oil is held in decantation tanks and a filtration process employed to remove impurities.

Acidity expressed as oleic acid: \leq 2,0 % Peroxide value (PV): \leq 10 meq/kg Insoluble impurities: \leq 0,05 % Alpha linolenic acid: \geq 60 % Linoleic acid: 15-20 %

Chia seeds (Salvia hispanica

Description/Definition:

Chia (*Salvia hispanica* L.) is a summer annual herbaceous plant belonging to the *Labiatae* family. Post-harvest the seeds are cleaned mechanically. Flowers, leaves and other parts of the plant are removed.

Dry matter: 90-97 % Protein: 15-26 % Fat: 18-39 %

Carbohydrate (*): 18-43 % Crude Fibre(**): 18-43 %

Ash: 3-7 %

- (*) Carbohydrates include the fibre value
- (**) Crude fibre is the part of fibre made mainly of indigestible cellulose, pentosans and lignin

Production process:

Production process of fruit juices and fruit juice blends beverages, containing Chia seeds, includes seed pre-hydration and pasteurisation steps. Microbiological controls and monitoring systems are in place.

Chitinglucan from

Description/Definition:

Aspergillus niger

Chitin-glucan is obtained from the mycelium of *Aspergillus niger*; it is a slightly yellow, odourless, free-flowing powder. It has a dry matter content of 90 % or more.

Chitin-glucan is composed largely of two polysaccharides:

chitin, composed of repeating units of N-acetyl-D-glucosamine

(CAS No: 1398-61-4),

 beta (1, 3)-glucan, composed of repeating units of D-glucose (CAS No: 9041-22-9).

Loss on drying: $\leq 10 \%$

Chitin-glucan: $\geq 90 \%$

Ratio of chitin to glucan: 30:70 to 60:40

Ash: $\leq 3.0 \%$ Lipids: $\leq 1.0 \%$ Proteins: $\leq 6.0 \%$

Chitin-glucan complex from Fomes fomentarius

Description/Definition:

Chitin-glucan complex is obtained from the cell walls of the fruit bodies of the fungus *Fomes fomentarius*. It consists primarily of two polysaccharides:

Chitin, composed of repeating units of N-acetyl-D-glucosamine

(CAS No: 1398-61-4);

Beta-(1,3)(1,6)-D-glucan, composed of repeating units of D-

glucose (CAS No: 9041-22-9).

The production process consists of several steps, including: cleaning, reduction in size and grinding, softening in water and heating in an alkaline solution, washing, drying. No hydrolysis is applied during the production process.

Appearance: Powder, odourless, flavourless, brown

Purity:

Moisture: $\leq 15 \%$ Ash: $\leq 3.0 \%$

Chitin-glucan: ≥ 90 %

Ratio of chitin to glucan: 70:20

Total carbohydrates, excluding glucans: $\leq 0.1 \%$

Proteins: ≤ 2,0 % Lipids: ≤ 1,0 % Melanins: ≤ 8,3 % Additives: None pH: 6,7-7,5

Heavy metals:

Lead (ppm): $\leq 1,00$ Cadmium (ppm): $\leq 1,00$ Mercury (ppm): $\leq 0,03$

Arsenic (ppm): ≤ 0.20 Microbiological criteria:

Total mesophilic bacteria: $\leq 10^3 / g$

Yeast and moulds: $\leq 10^3 / g$ Coliforms at 30 °C: $\leq 10^3 / g$

 $E. coli : \leq 10/g$

Salmonella and other pathogenic bacteria: Absence/25 g

Chitosan extract from fungi

Description/Definition:

(Agaricus bisporus ; Aspergillus niger)

The chitosan extract (containing mainly poly(D-glucosamine)) is obtained from stems of *Agaricus bisporus* or from the mycelium of *Aspergillus niger*.

The patented production process consists of several steps, including: extraction and deacetylation (hydrolysis) in alkaline medium, solubilisation in acidic medium, precipitation in alkaline medium, washing and drying.

Synonym: Poly(D-glucosamine) Chitosan CAS number: 9012-76-4 Chitosan formula: (C₆H₁₁NO₄)_n Appearance: fine free-flowing powder Aspect: Off –white to slightly brownish

Odour: Odourless

Purity:

Chitosan content (% w/w dry weight):≥ 85 Glucan content (% w/w dry weight): ≤ 15 Loss on drying (% w/w dry weight): ≤ 10 Viscosity (1 % in 1 % acetic acid): 1-15

Degree of acetylation (in % mol/wet weight): 0-30

Viscosity (1 % in 1 % acetic acid) (mPa.s): 1-14 for chitosan from

Aspergillus niger; 12-25 for chitin from *Agaricus bisporus*

Ash (% w/w dry weight): $\leq 3,0$ Proteins (% w/w dry weight): $\leq 2,0$

Particle size: > 100 nm

Tapped density (g/cm^3) : 0,7-1,0

Fat binding capacity $800 \times (\text{w/w wet weight})$: pass **Heavy metals:**

Mercury (ppm): $\leq 0,1$ Lead (ppm): $\leq 1,0$ Arsenic (ppm): $\leq 1,0$ Cadmium (ppm): $\leq 0,5$ Microbiological criteria:

Aerobic count (CFU/g): $\leq 10^3$

recoole count (Cr 6/g). = 10

Yeast and mould count (CFU/g): $\leq 10^3$ Escherichia coli (CFU/g): ≤ 10 Enterobacteriaceae (CFU/g): ≤ 10

Salmonella: Absence/25g

Listeria monocytogenes: Absence/25g

Chondroitin sulphate

Description/Definition:

Chondroitin sulphate (sodium salt) is a biosynthetic product. It is obtained by chemical sulphation of chondroitin derived from fermentation by the bacterium *Escherichia coli* O5:K4:H4 strain U1-41 (ATCC 23502).

Chondroitin sulphate (sodium salt) (% dry basis): 95-105

MWw (weight avg.) (kDa): 5-12 MWn (number avg.) (kDa): 4-11 Dispersity ($w_h/w_{0.05}$): ≤ 0.7

Sulphation pattern ($\Delta \text{Di-6S}$) (%): ≤ 85

Loss on drying (%) (105 °C to constant weight): ≤ 10.0

Residue on ignition (% dry basis): 20-30

Protein (% dry basis): ≤ 0.5 Endotoxins (EU/mg): ≤ 100

Total organic impurities (mg/kg): ≤ 50

Chromium Picolinate

Description/Definition:

Chromium picolinate is a reddish free-flowing powder, slightly soluble in water at pH 7. The salt is also soluble in polar organic solvents. Chemical name: tris(2pyridinecarboxylato-N,O)chromium(III) or 2-

pyridinecarboxylic acid chromium(III) salt

CAS No.: 14639-25-9Chemical formula: Cr(C₆H₄NO₂)₃

Chemical characteristics: Chromium Picolinate: ≥ 95 % Chromium (III): 12-13 % Chromium (VI): not detected

Water: $\leq 4.0 \%$

Cistus incanus L. Pandalis

herb

Description:

Cistus incanus L. Pandalis herb; species belonging to the Cistaceae family and native to the Mediterranean region, Chalkidiki Peninsula.

Composition:

Moisture: 9–10 g/100 g herbs Protein: 6,1 g/100 g herbs Fat: 1,6 g/100 g herbs

Carbohydrates: 50,1 g/100 g herbs

Fiber: 27,1 g/100 g herbs Minerals: 4,4 g/100 g herbs

Sodium: 0,18 g Potassium: 0,75 g Magnesium: 0,24 g Calcium: 1,0 g Iron: 65 mg

Vitamin B $_1$: 3,0 μg Vitamin B $_2$: 30 μg Vitamin B $_6$: 54 μg Vitamin C: 28 m g

Vitamin A: less than 0,1 mg Vitamin E: 40–50 mg Alpha-Tocopherol: 20–50 mg

Beta and Gamma-Tocopherols: 2–15 mg

Delta-Tocopherol: 0,1–2 mg

Citicoline

Description/Definition:

Citicoline is produced by a microbial process.

Citicoline is composed of cytosine, ribose, pyrophosphate and choline.

White crystalline powder

Chemical name: Choline cytidine 5'-pyrophosphate, Cytidine 5'-

(trihydrogen diphosphate) P'-[2-(trimethylammonio)ethyl]ester inner salt

Chemical formula: C₁₄ H₂₆ N₄ O₁₁ P₂ Molecular weight: 488,32 g/mol

CAS No.: 987-78-0

pH (sample solution of 1 %): 2,5-3,5

Purity:

Assay value: \geq 98 % of dry matter

Loss on drying (100 °C for 4 hours): $\leq 5.0 \%$

Ammonium: $\leq 0.05 \%$

Arsenic: Not more than 2 ppm

Free phosphoric acids: $\leq 0.1 \%$ 5'-Cytidylic acid: $\leq 1.0 \%$ Microbiological criteria: Total plate count: $\leq 10^3$ CFU/g

Yeast and moulds: $\leq 10^2$ CFU/g Escherichia coli: Absence in 1 g

Clostridium butyricum

Description/Definition:

Clostridium butyricum (CBM-588) is a Gram-positive, spore-forming, obligate anaerobic, non-pathogenic, non-genetically modified bacterium.

Depository number FERM BP-2789

Microbiological criteria:

Total viable aerobic count: $\leq 10^3$ CFU/g

Escherichia coli: Not detected in 1 g Staphylococcus aureus: Not

detected in 1 g

Pseudomonas aeruginosa: Not detected in 1 g

Yeast and moulds: $\leq 10^2$ CFU/g

[F8D-ribose

Description

D-ribose is an aldopentose monosaccharide which is produced by fermentation using a transketolase-deficient strain of Bacillus subtilis.

Chemical formula: C₅H₁₀O₅

CAS No: 50-69-1

Molecular mass: 150.13 Da Characteristics/Composition

Appearance: Dry with powdery texture, white to slightly yellow in colour

Specific rotation $[\alpha]_D^{25}$: -19.0° to -21.0°

D-ribose purity (% dry basis): -HPLC/RI^h Method 98,0-102,0 %

Ash: < 0.2 %

Loss on drying (moisture): < 0.5 %Clarity on solution: ≥ 95 % transmittance

Heavy metals Lead: $\leq 0.1 \text{ mg/kg}$

Arsenic: $\leq 0.1 \text{ mg/kg}$ Cadmium: $\leq 0.1 \text{ mg/kg}$ Mercury: $\leq 0.1 \text{ mg/kg}$

Microbiological criteria

Total plate count: ≤ 100 CFUⁱ/g

Yeast: $\leq 100 \text{ CFU/g}$ Moulds: $\leq 100 \text{ CFU/g}$ Coliforms: ≤ 10 CFU/g

Salmonella sp: Negative/25 g]

Extract of defatted cocoa powder

Cocoa (Theobroma cacao L.) Extract

Appearance: Dark brown powder free of visible impurities

Physical and chemical properties: Polyphenol content: Min 55,0 % GAE Theobromine content: Max 10,0 %

Ash content: Max 5.0 % Moisture content: Max 8,0 % Bulk density: 0,40-0,55 g/cm³

	pH: 5,0-6,5 Residual solvent: Max 500 ppm
Low fat cocoa extract	Low fat Cocoa (<i>Theobroma cacao</i> L.) extract Appearance: Dark red to purple powder Cocoa extract, concentrate: Min 99 % Silicon dioxide (technological aid): Max 1,0 % Cocoa flavanols: Min. 300 mg/g — Epicatechin: Min. 45 mg/g Loss on drying: Max. 5,0 %
[F59]Coriander seed oil from Coriandrum sativum	Description/Definition: Coriander seed oil is an oil containing glycerides of fatty acids that is produced from the seeds of the coriander plant <i>Coriandrum sativum</i> L. Slight yellow colour, bland taste CAS No: 8008-52-4 Composition of fatty acids: Palmitic acid (C16:0): 2-5 % Stearic acid (C18:0): < 1,5 % Petroselinic acid (cis-C18:1(n-12)): 60-75 % Oleic acid (cis-C18:1 (n-9)): 7-15 % Linoleic acid (C18:2): 12-19 % α-Linolenic acid (C18:3): < 1,0 % Trans fatty acids: ≤ 1,0 % Purity: Refractive index (20 °C): 1,466-1,474 Acid value: ≤ 2,5 mg KOH/g Peroxide value (PV): ≤ 5,0 meq/kg Iodine value: 88-110 units Saponification value: 179-200 mg KOH/g Unsaponifiable matter: ≤ 15 g/kg]
[F12Cranberry extract powder	Description/Definition: Cranberry extract powder is a water-soluble phenolic-rich powder extract prepared through an ethanolic extraction from the juice concentrate of sound, mature berries of the cranberry cultivar Vaccinium macrocarpon. Characteristics/Composition Moisture (% w/w): ≤ 4 Proanthocyanidins — PACs (% w/w dry weight) — OSC-DMAC method ^{ce} : 55.0-60.0 or — BL-DMAC method ^{ce} : 15.0-18.0 Total phenolics (GAE ^f , % w/w dry weight) ^e — Folin-Ciocalteau method: > 46.2 Solubility (water): 100 %, with no visible insoluble particles Ethanol Content (mg/kg): ≤ 100 Screen Analysis: 100 % through 30 mesh screen Appearance and aroma, as powder: Free-flowing, deep red colour. Earthy aroma with no burnt character. Heavy metals: Arsenic (ppm): < 3 Microbiological criteria: Yeast: < 100 CFU/g Mould: < 100 CFU/g Aerobic plate count: < 1 000 CFU/g

Coliforms: < 10 CFU/g *Escherichia coli* : < 10 CFU/g Salmonella: Absent in 375 gl

Crataegus pinnatifida dried fruit

Description/Definition:

Dried fruits of Crataegus pinnatifida species belonging to the Rosaceae family and native to north China and Korea.

Composition: Dry matter: 80 %

Carbohydrates: 55 g/kg fresh weight

Fructose: 26,5–29,3 g/100 g Glucose: 25,5–28,1 g/100 g

Vitamin C: 29,1 mg/100 g fresh weight Sodium: 2,9 g/100 g fresh weight

Compotes are products obtained by thermal processing of the edible part of one or several species of fruits, whole or in pieces, sieved or not, without significant concentration. Sugars, water, cider, spices and lemon juice may be used.

α-cyclodextrin

Description/Definition:

A non-reducing cyclic saccharide consisting of six α -1,4-linked D-glucopyranosyl units produced by the action of cyclodextrin glucosyltransferase (CGTase, EC 2.4.1.19) on hydrolyzed starch. Recovery and purification of α -cyclodextrin may be carried out using one of the following procedures: precipitation of a complex of α -cyclodextrin with 1-decanol, dissolution in water at elevated temperature and reprecipitation, steam-stripping of the complexant, and crystallisation of αcyclodextrin from the solution; or chromatography with ion-exchange or gel filtration followed by crystallisation of α -cyclodextrin from the purified mother liquor; or membrane separation methods such as ultrafiltration and reverse osmosis: Description: Virtually odourless, white or almost white crystalline solid.

Synonyms: α -cyclodextrin, α -dextrin, cyclohexaamylose,

cyclomaltohexaose, α-cycloamylase

Chemical name: CyclohexaamyloseCAS No.: 10016-20-3

Chemical formula: (C₆H₁₀O₅)₆

Formula weight: 972,85 Assay: \geq 98 % (dry basis)

Identification:

Melting range: Decomposes above 278 °C

Solubility: Freely soluble in water; very slightly soluble in ethanol Specific rotation: $[\alpha]_D^{25}$: Between +145° and +151° (1% solution) Chromatography: The retention time for the major peak in a liquid chromatogram of the sample corresponds to that for α -cyclodextrin in a chromatogram of reference α-cyclodextrin (available from *Consortium* für Elektrochemische Industrie GmbH, München, Germany or Wacker Biochem Group, Adrian, MI, USA) using the conditions described in the METHOD OF ASSAY

Purity:

Water: ≤ 11 % (Karl Fischer Method) Residual complexant: $\leq 20 \text{ mg/kg}$

(1-decanol)

Reducing substances: $\leq 0.5 \%$ (as glucose)

Sulphated ash: $\leq 0.1 \%$ Lead: $\leq 0.5 \text{ mg/kg}$ **Method of assay:**

Determine by liquid chromatography using the following conditions: Sample solution: Weigh accurately about 100 mg of test sample into a 10 ml volumetric flask and add 8 ml of deionised water. Dissolve the sample completely using an ultra-sonification bath (10-15 min) and dilute to the mark with purified deionised water. Filter through a 0,45-micrometer filter

Reference solution: Weigh accurately about 100 mg of α -cyclodextrin into a 10 ml volumetric flask and add 8 ml of deionised water. Dissolve the sample completely using an ultra-sonification bath and dilute to the mark with purified deionised water.

Chromatography: Liquid chromatograph equipped with a refractive index detector and an integrating recorder.

Column and packing: Nucleosil-100-NH₂ (10 µm) (Macherey & Nagel

Co. Düren, Germany) or similar

Length: 250 mm Diameter: 4 mm Temperature: 40 °C

Mobile phase: acetonitrile/water (67/33, v/v)

Flow rate: 2,0 ml/min

Injection volume: 10 μ lProcedure: Inject the sample solution into the chromatograph, record the chromatogram, and measure the area of the α -CD peak. Calculate the percentage of α -cyclodextrin in the test sample as follows:

% α -cyclodextrin (dry basis) = $100 \times (A_S/A_R) (W_R/W_S)$ where

 A_S and A_R are the areas of the peaks due to α -cyclodextrin for the sample solution and reference solution, respectively.

 W_S and W_R are the weights (mg) of the test sample and reference α -cyclodextrin, respectively, after correcting for water content.

γ-cyclodextrin

Description/Definition:

A non-reducing cyclic saccharide consisting of eight α -1,4-linked D-glucopyranosyl units produced by the action of cyclodextrin glucosyltransferase (CGTase, EC 2.4.1.19) on hydrolysed starch. Recovery and purification of γ -cyclodextrin may be carried out by precipitation of a complex of γ -cyclodextrin with 8-cyclohexadecen-1-one, dissolution of the complex with water and n-decane, steam-stripping of the aqueous phase and recovery of gamma-CD from the solution by crystallisation.

Virtually odourless, white or almost white crystalline solid Synonyms: γ-cyclodextrin, γ-dextrin, cyclooctaamylose,

cyclomaltooctaose, γ-cycloamylase Chemical name: Cyclooctaamylose

CAS number: 17465-86-0Chemical formula: $(C_6H_{10}O_5)_8$

Assay: \geq 98 % (dry basis)

Identification:

Melting range: Decomposes above 285 °C

Solubility: Freely soluble in water; very slightly soluble in ethanol Specific rotation: $[\alpha]_D^{25}$: between + 174° and + 180° (1% solution)

Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

Purity:

Water: ≤ 11 %

Residual complexant (8-cyclohexadecen-1-one (CHDC)): ≤ 4 mg/kg

Residual solvent (n-decane): < 6mg/kg Reducing substances: $\leq 0.5 \%$ (as glucose)

Sulphated ash: $\leq 0.1 \%$

[F13Decorticated grains of

Digitaria exilis (Kippist) Stapf (fonio) (Traditional food

from a third

country)

Description/Definition

The traditional food is the decorticated grain (bran removed) of Digitaria exilis (Kippist) Stapf.

Digitaria exilis (Kippist) Stapf) is an annual herbaceous plant belonging to the Poaceae family.

Typical nutritional components of decorticated grain of fonio

Carbohydrates: 76,1 g/100 g of fonio

Water: 12,4 g/100 g of fonio Protein: 6,9 g/100 g of fonio Fat: 1,2 g/100 g of fonio Fibre: 2,2 g/100 g of fonio Ash: 1,2 g/100 g of fonio Phytate content: $\leq 2.1 \text{ mg/g}$

Dextran preparation produced by Leuconostoc mesenteroides

2.

1. Powdered form:

Carbohydrates: 60 % with: (Dextran: 50 %, Mannitol: 0,5 %,

Fructose: 0,3 %, Leucrose: 9,2 %)

Protein: 6.5 % Lipid: 0.5 % Lactic acid: 10 % Ethanol: traces Ash: 13 % Moisture: 10 % Liquid form:

Carbohydrates: 12 % with: (Dextran: 6,9 %, Mannitol: 1,1 %,

Fructose: 1,9 %, Leucrose: 2,2 %)

Protein: 2,0 % Lipid: 0,1 % Lactic acid: 2,0 % Ethanol: 0.5 % Ash: 3,4 % Moisture: 80 %

Diacylglycerol oil of plant origin

Description/Definition:

Manufactured from glycerol and fatty acids derived from edible vegetable oils, in particular from sovbean oil (Glycine max) or rapeseed oil (Brassica campestris, Brassica napus) using a specific enzyme.

Acylglycerol Distribution:

Diacylglycerols (DAG): $\geq 80 \%$

1,3-Diacylglycerols (1,3-DAG): \geq 50 %

Triacylglycerols (TAG): $\leq 20 \%$ Monoacylglycerols (MAG): ≤ 5,0 %

Fatty Acid Composition (MAG, DAG, TAG):

Oleic acid (C18:1): 20-65 % Linoleic acid (C18:2): 15-65 % Linolenic acid (C18:3): $\leq 15 \%$ Saturated fatty acids: $\leq 10 \%$

Others:

Acid value: $\leq 0.5 \text{ mg KOH/g}$ Moisture and volatile: $\leq 0.1 \%$ Peroxide value (PV): $\leq 1.0 \text{ meg/kg}$

Unsaponifiables: $\leq 2.0 \%$ Trans fatty acids≤ 1,0 %

MAG = monoacylglycerols, DAG = diacylglycerols, TAG =

triacylglycerols

Dihydrocapsiate (DHC)

Description/Definition:

Dihydrocapsiate is synthesised by enzyme-catalysed esterification of vanillyl alcohol and 8-methylnonanoic acid. Following the esterification dihydrocapsiate is extracted with n-hexane.

Viscous to colourless to yellow liquid Chemical formula: C₁₈ H₂₈ O₄

CAS No: 205687-03-2

Physical-chemical properties:

Dihydrocapsiate: > 94 % 8-Methylnonanoic acid: < 6.0 %

Vanillyl acohol: < 1,0 %

Other synthesis related substances: < 2,0 %

[F14Dried aerial parts of *Hoodia* parviflora

Description/Definition:

It is the whole dried aerial parts of *Hoodia parviflora* N.E.Br., (family Apocynaceae)

Characteristics/Composition

Plant material: Aerial parts of at least 3-year-old plants

Appearance: Light green to tan fine powder

Solubility (water): > 25 mg/mL

Moisture: < 5,5 %

 A_{w} : < 0,3

pH: < 5.0

Protein: < 4.5 g/100 g

Fat: < 3 g/100 g

Carbohydrate (including dietary fibre): < 80 g/100 g

Dietary fibre: < 55 g/100 g Total sugars: < 10.5 g/100 g

Ash: < 20 %

Hoodigosides P57: 5-50 mg/kg

L: 1 000–6 000 mg/kg O: 500-5 000 mg/kg

Total: 1 500-11 000 mg/kg

Heavy metals:

Arsenic: < 1,00 mg/kgMercury: < 0.1 mg/kgCadmium: < 0.1 mg/kgLead: < 0.5 mg/kg

Microbiological criteria:

Aerobic plate count: < 10⁵ CFU/g Escherichia coli : < 10 CFU/g *Staphylococcus aureus* : < 50 CFU/g Total coliforms: < 10 CFU/g

Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

Yeast: $\leq 100 \text{ CFU/g}$ Mould: $\leq 100 \text{ CFU/g}$

Salmonella species: Negative/25 g Listeria monocytogenes : Negative/25 g

CFU: Colony Forming Units

Dried extract of *Lippia* citriodora from cell cultures

Description/Definition:

Dried extract of Lippia citriodora (Palau) Kunth from cell cultures HTN

® Vb.

Echinacea angustifolia extract from cell cultures

Description/Definition:

Extract of the roots of *Echinacea angustifolia* obtained from plant tissue culture which is substantially equivalent to a root extract from *Echinacea angustifolia* obtained in ethanol-water titrated to 4 % echinacoside.

[F15 Echinacea purpurea extract from cell cultures

Description/Definition:

Dried extract of *Echinacea purpurea* from cell cultures EchiPure-PCTM]

Echium plantagineum oil

Description/Definition:

Echium oil is the pale yellow product obtained by refining oil extracted from the seeds of *Echium plantagineum* L. Stearidonic acid: $\geq 10 \%$ w/

w of total fatty acids

Trans fatty acids: ≤ 2.0 % (w/w of total fatty acids)

Acid value: $\leq 0.6 \text{ mg KOH/g}$

Peroxide value (PV): ≤ 5.0 meq O₂/kg

Unsaponifiable content: ≤ 2,0 %

Protein content (total nitrogen): $\leq 20 \mu g/ml$

Pyrrolizidine alkaloids: Not detectable with a detection limit 4,0 μg/kg

[F16Ecklonia cava phlorotannins

Description/Definition

Ecklonia cava phlorotannins are obtained via alcohol extraction from the edible marine alga *Ecklonia cava*. The extract is a dark brown powder, rich in phlorotannins, polyphenolic compounds found as secondary metabolites in certain brown algae species.

Characteristics/Composition

Phlorotannin content: 90 ± 5 % Antioxidant activity: > 85 %

Moisture: < 5 % Ash: < 5 %

Microbiological criteria

Total viable cell count: < 3 000 CFU/g

Mould/yeast: < 300 CFU/g Coliforms: Negative to test Salmonella spp.: Negative to test Staphylococcus aureus: Negative to test

Heavy metals and Halogens

Lead: < 3,0 mg/kg Mercury: < 0,1 mg/kg Cadmium: < 3,0 mg/kg Arsenic: < 25,0 mg/kg

Inorganic Arsenic: < 0,5 mg/kg Iodine: 150,0 – 650,0 mg/kg

CFU: Colony Forming Units]

[F17Egg membrane hydrolysate

Description

The egg membrane hydrolysate is derived from the eggshell membranes of chicken eggs. The eggshells undergo hydro-mechanical separation in order to obtain the egg membranes, which are then further processed using a patented solubilisation method. Following the solubilisation process, the solution is filtered, concentrated, spray-dried and packaged.

Characteristics/Composition

Chemical parameters	Methods
Total nitrogen- containing compounds (% w/w): ≥ 88	Combustion according to AOAC 990.03 and AOAC 992.15
Collagen (% w/ w): ≥ 15	Sircol TM Soluble Collagen Assay
Elastin (% w/ w): ≥ 20	Fastin TM Elastin Assay
Total glycosaminoglyca (% w/w): ≥ 5	USP26 (chondroitin sulphate K0032 method)
Calcium: ≤ 1 %	

Physical parameters

pH: 6.5 - 7.6Ash (% w/w): ≤ 8 Moisture (% w/w): ≤ 9 Water activity: ≤ 0.3

Solubility (in water): soluble Bulk density: ≥ 0.6 g/cc

Heavy metals Arsenic $\leq 0.5 \text{ mg/kg}$

Microbiological criteria

Aerobic plate count: \leq 2 500 CFU/g Escherichia coli : \leq 5 MPN/g Salmonella : Negative (in 25 g)

Coliforms: $\leq 10 \text{ MPN/g}$

Staphylococcus aureus : ≤ 10 CFU/g Mesophilic spore count: ≤ 25 CFU/g Thermophilic spore count: ≤ 10 CFU/10 g

Yeast: $\leq 10 \text{ CFU/g}$ Mould: $\leq 200 \text{ CFU/g}$

CFU: Colony Forming Units; MPN = Most Probable Number; USP:

United States Pharmacopeia.]

Epigallocateching allate as a purified extract from green tea leaves

Epigallocatechin Description/Definition:

A highly purified extract from the leaves of green tea (*Camellia sinensis (L.) Kuntze*) in the form of a fine, off-white to pale pink powder. It is composed of a minimum of 90 % epigallo-catechin gallate (EGCG), and has a melting point between approx. 210 and 215 °C

(Camellia sinensis)

Appearance: off-white to pale pink powder

Chemical name: polyphenol (-) epigallocatechin-3-gallate

Synonyms: epigallocatechin gallate (EGCG)

CAS No.: 989-51-5

INCI name: epigallocatechin gallate Molecular mass: 458,4 g/mol Loss on drying: max 5,0 %

Heavy metals: Arsenic: max 3,0 ppm Lead: max 5,0 ppm

Assay:

Min. 94 % EGCG (on dry material)

max. 0,1 % caffeine

Solubility: EGCG is fairly soluble in water, ethanol, methanol and

acetone

Lergothioneine

Definition

Chemical name (IUPAC): (2S)-3-(2-thioxo-2,3-dihydro-1 H -imidazol-4-

yl)-2-(trimethylammonio)-Propanoate Chemical formula: $C_9 H_{15} N_3 O_2 S$

Molecular mass: 229,3 Da CAS No.: 497-30-3

Parameter	Specification	Method		
Appearance	White powder	Visual		
Optical rotation	$[\alpha]_D \ge (+) 122^{\circ}$ $(c = 1, H_2 O)^{a)}$	Polarimetry		
Chemical purity	≥ 99,5 % ≥ 99,0 %	HPLC [Eur. Ph. 2,2.29] 1H-NMR		
Identification	Compliant with the structure C: $47,14 \pm 0,4 \%$ H: $6,59 \pm 0,4 \%$ N: $18,32 \pm 0,4 \%$	1H-NMR Elemental analysis		
Total residual solvents (methanol, ethyl acetate, isopropanol, ethanol)	[Eur. Ph. 01/2008:50400] < 1 000 ppm	Gas chromatography [Eur. Ph. 01/2008:20424]		
Loss on drying	Internal standard < 0,5 %	[Eur. Ph. 01/2008:20232]		
Impurities	< 0,8 %	HPLC/GPC or 1H-NMR		
Heavy metals b)c)				
Lead	< 3,0 ppm	ICP/AES		
Cadmium	< 1,0 ppm	(Pb, Cd)		
Mercury	< 0,1 ppm	Atomic fluorescence (Hg)		

Microbiological specifications b)					
Total viable aerobic count (TVAC)	≤ 1 x 10 ³ CFU/g	[Eur. Ph. 01/2011:50104]			
Total yeast and mould count (TYMC)	≤ 1 x 10 ² CFU/g				
Escherichia coli	Absence in 1 g				

Eur. Ph.: European Pharmacopoeia; 1H-NMR: proton nuclear magnetic resonance; HPLC: high-performance liquid chromatography; GPC: gel permeation chromatography; ICP/AES: Inductively coupled plasma atomic emission spectroscopy;

CFU: colony-forming units.

- a) Lit. $[\alpha]_D = (+) 126.6^{\circ} (c = 1, H_2 O)$
- b) Analyses conducted on each batch
- c) Maximum levels in accordance with Regulation (EC) No 1881/2006

[F16Extract of three herbal roots (Cynanchum wilfordii Hemsley, Phlomis umbrosa Turcz. and Angelica gigas Nakai)

Description/Definition

The mixture of the three herbal roots is yellowish brown fine powder produced by hot-water extraction, concentration by evaporation, and spray drying

Composition of the extract of mixture of the 3 herbal roots

Cynanchum wilfordii root: 32,5 % (w/w) Phlomis umbrosa root: 32,5 % (w/w) Angelica gigas root: 35,0 % (w/w)

Specifications

Loss on drying: NMT 100 mg/g

Assay

Cinnamic acid: 0.012 - 0.039 mg/g

Shanzhiside methyl ester: 0.20 - 1.55 mg/g

Nodakenin: 3,35 – 10,61 mg/g Methoxsalen: < 3 mg/g Phenols: 13,0 – 40,0 mg/g Coumarins: 13,0 – 40,0 mg/g Iridoids: 13,0 – 39,0 mg/g Saponins: 5,0 – 15,5 mg/g

Nutritive components

Carbohydrates: 600 - 880 mg/g

Proteins: 70 - 170 mg/g

Fats: < 4 mg/g

Microbiological parameters

Total viable plate count: < 5000 CFU/g Total mold and yeast: < 100 CFU/g Coliform bacteria: < 10 CFU/g Salmonella: Negative/25 g Escherichia coli: Negative/25 g Staphylococcus aureus: Negative/25 g

Heavy metals Lead: < 0,65 mg/kg

Arsenic: < 3,0 mg/kg Mercury: < 0,1 mg/kg Cadmium: < 1,0 mg/kg CFU: Colony Forming Units]

Ferric Sodium EDTA

Description/Definition:

Ferric Sodium EDTA (ethylenediaminetetraacetic acid) is an odourless free-flowing, yellow to brown powder with a chemical purity of more

than 99 % (w/w). It is freely soluble in water. Chemical formula: $C_{10}H_{12}$ FeN $_2$ NaO $_8$ * $3H_2$ O

Chemical characteristics: pH of 1 % solution: 3,5-5,5

Iron: 12,5-13,5 % Sodium: 5,5 % Water: 12,8 %

Organic matter (CHNO): 68,4 %

EDTA: 65,5-70,5 %

Water insoluble matter: $\leq 0.1 \%$ Nitrilo-triacetic acid: $\leq 0.1 \%$

Ferrous ammonium phosphate

Description/Definition:

Ferrous ammonium phosphate is a grey/green fine powder, practically insoluble in water and soluble in dilute mineral acids.

CAS No.: 10101-60-7

Chemical formula: FeNH 4 PO 4

Chemical characteristics:

pH of 5 % suspension in water: 6,8-7,8

Iron (total): $\geq 28 \%$ Iron (II): 22-30 % (w/w) Iron (III): $\leq 7,0 \%$ (w/w) Ammonia: 5-9 % (w/w) Water: $\leq 3,0 \%$

Fish peptides from Sardinops sagax

Description/Definition:

The novel food ingredient is a peptide mixture, which is obtained by an alkaline protease-catalysed hydrolysis of fish (*Sardinops sagax*) muscle, subsequent isolation of the peptide fraction by column chromatography, concentration under vacuum and spray drying.

Yellowish white powderPeptides (1) (short chain peptides, dipeptides and tripeptides with a molecular weight of less than 2 kDa): $\geq 85 \text{ g}/100 \text{ g}$

Val-Tyr (dipeptide): 0,1-0,16 g/100 g

Ash: $\leq 10 \text{ g}/100 \text{ g}$ Moisture: $\leq 8 \text{ g}/100 \text{ g}$

(1) Kjeldahl method

Flavonoids from Glycyrrhiza glabra

Description/Definition:

Flavonoids derived from the roots or rootstock of *Glycyrrhiza glabra* L. are extracted with ethanol followed by further extraction of this ethanolic extract with medium-chain triglycerides. It is a dark-brown coloured liquid, containing 2,5 % to 3,5 % of glabridin.

Moisture: < 0,5 % Ash: < 0,1 %

Peroxide value (PV): < 0.5 meq/kg

Glabridin: 2,5-3,5 % of fat Glycyrrhizinic acid: < 0,005 %

Fat including polyphenol-type substances: $\geq 99 \%$

Protein: < 0.1 %

Carbohydrates: not detectable

[F19Fruit pulp, pulp juice, concentrated pulp juice from *Theobroma cacao* L. (Traditional food from a third country)

Description/Definition

The traditional food is the fruit pulp from the cocoa (*Theobroma cacao* L) plant, which is the 'aqueous, mucilaginous and acidic substance in which the seeds are embedded'.

Cocoa fruit pulp is obtained by splitting cocoa pods followed by separation from husks and beans; the pulp is then subject to pasteurisation and freezing. Cocoa pulp juice and/or cocoa concentrated pulp juice are produced following processing (enzymatic treatment, pasteurization, filtration, and concentration).

Typical compositional data of cocoa fruit pulp, pulp juice, concentrated pulp juice

Protein (g/100 g): 0,0 to 2,0 Total fat (g/100 g): 0,0 to 0,2 Total sugars (g/100 g): > 11,0 Brix level (° Brix): \geq 14 pH: 3,3 to 4,0

Microbiological criteria

Total Plate Count (aerobic): < 10 000 cfuⁱ/g

Enterobacteriaceae: ≤ 10 cfu/g *Salmonella*: Absence in 25 g]

Fucoidan extract from the seaweed Fucus vesiculosus

Description/Definition:

Fucoidan from the seaweed *Fucus vesiculosus* is extracted using aqueous extraction in acidic solution and filtration processes without the use of organic solvents. The resulting extract is concentrated and dried to yield the fucoidan extract with the following specifications:

Off-white to brown powder

Odour and Taste: Bland odour and taste Moisture: < 10 % (105 °C for 2 hours) pH value: 4,0-7,0 (1 % suspension at 25 °C)

Heavy metals:

Arsenic (inorganic): < 1,0 ppm

Cadmium: < 3,0 ppm Lead: < 2,0 ppm

Mercury: < 1,0 ppm **Microbiological criteria:** Total aerobic microbial count: < 10 000 CFU/g

Yeast and mould count: < 100 CFU/g Total enterobacteria count: Absence/g

Escherichia coli : Absence/g
Salmonella : Absence/10 g
Staphylococcus aureus : Absence/g

Composition of the two permitted types of extracts, based on the level of

fucoidan: Extract 1:

Fucoidan: 75-95 % Alginate: 2,0-5,5 %

Polyphloroglucinol: 0,5-15 %

Mannitol: 1-5 %

Natural salts/Free Minerals: 0,5-2,5 % Other carbohydrates: 0,5-1,0 %

Protein: 2,0-2,5 %

Extract 2:

Fucoidan: 60-65 % Alginate: 3,0-6,0 %

Polyphloroglucinol: 20-30 %

Mannitol: < 1,0 %

Natural salts/Free Minerals: 0,5-2,0 % Other carbohydrates: 0,5-2,0 %

Protein: 2,0-2,5 %

Fucoidan extract from the seaweed *Undaria* pinnatifida

Description/Definition:

Fucoidan from seaweed *Undaria pinnatifida* is extracted using aqueous extraction in acidic solution and filtration processes without the use of organic solvents. The resulting extract is concentrated and dried to yield the fucoidan extract with the following specifications:

Off-white to brown powder

Odour and Taste: Bland odour and tasteMoisture: < 10 % (105 °C for 2

hours)

pH value: 4,0-7,0 (1 % suspension at 25 °C)

Heavy metals:

Arsenic (inorganic): < 1,0 ppm

Cadmium: < 3,0 ppm Lead: < 2,0 ppm Mercury: < 1,0 ppm **Microbiology:**

Total aerobic microbial count: < 10 000 CFU/g

Yeast and mould count: < 100 CFU/g Total enterobacteria count: Absence/g

Escherichia coli : Absence/g *Salmonella* : Absence/10 g

Staphylococcus aureus: Absence/g

Composition of the two permitted types of extracts, based on the level of

fucoidan: Extract 1:

Fucoidan: 75-95 % Alginate: 2,0-6,5 %

Polyphloroglucinol: 0,5-3,0 %

Mannitol: 1-10 %

Natural salts/Free Minerals: 0,5-1,0 % Other carbohydrates: 0,5-2,0 %

Protein: 2,0-2,5 %

Extract 2:

Fucoidan: 50-55 % Alginate: 2,0-4,0 %

Polyphloroglucinol: 1,0-3,0 %

Mannitol: 25-35 %

Natural salts/Free Minerals: 8-10 % Other carbohydrates: 0,5-2,0 %

Protein: 1,0-1,5 %

2′-

Fucosyllactose

Definition:

(synthetic)

Chemical name: α -L-Fucopyranosyl- $(1\rightarrow 2)$ - β -D-galactopyranosyl-

 $(1\rightarrow 4)$ - D-glucopyranose

Chemical formula: C₁₈ H₃₂ O₁₅

CAS No: 41263-94-9

Molecular weight: 488,44 g/mol

Description:

2'-fucosyllactose is a white to off-white powder that is produced by a chemical synthesis process.

Purity:

2'-Fucosyllactose: ≥ 95 % D-Lactose: ≤ 1,0 w/w % L-Fucose: ≤ 1,0 w/w %

Difucosyl- D-lactose isomers: \leq 1,0 w/w % 2'-Fucosyl- D-lactulose: \leq 0,6 w/w % pH (20 °C, 5 % solution): 3,2-7,0

Water (%): \leq 9,0 % Ash, sulphated: \leq 0,2 % Acetic acid: \leq 0,3 %

Residual solvents (methanol, 2-propanol, methyl acetate, acetone): ≤ 50.0

mg/kg singly, ≤ 200.0 mg/kg in combination

Residual proteins: $\leq 0.01 \%$

Heavy Metals:

Palladium: ≤ 0,1 mg/kg Nickel: ≤ 3,0 mg/kg Microbiological criteria:

Aerobic mesophilic bacteria total count: ≤ 500 CFU/g

Yeasts and Moulds: $\leq 10 \text{ CFU/g}$ Residual endotoxins: $\leq 10 \text{ EU/mg}$

2'-Fucosyllactose (microbial source)

[F60 Definition:

Chemical name: α -L-Fucopyranosyl- $(1\rightarrow 2)$ - β -D-galactopyranosyl-

 $(1\rightarrow 4)$ -D-glucopyranose

Chemical formula: C₁₈ H₃₂ O₁₅

CAS No: 41263-94-9

Molecular weight: 488,44 g/mol

Source:

Genetically modified strain of

Escherichia coli K-12

Source:

Genetically modified strain of

Escherichia coli BL21

Description:

2'-Fucosyllactose is a white to offwhite powder that is produced by a microbial process.

Purity:

2'-Fucosyllactose: ≥ 83 % D-Lactose: ≤ 10,0 % L-Fucose: ≤ 2,0 %

Difucosyl-D-lactose: ≤ 5,0 % 2'-Fucosyl-D-lactulose: ≤ 1,5 % Sum of saccharides (2'-Fucosyllactose, D-Lactose, L-Fucose, Difucosyl-D-lactose, 2'-

Fucosyl-D-lactulose): $\geq 90 \%$

Description:

2'-Fucosyllactose is a white to off white powder and the liquid concentrate ($45\% \pm 5\%$ w/v) aqueous solution is a colourless to slight yellow clear aqueous solution. 2'-Fucosyllactose is produced by a microbiological process.

Purity:

2'-Fucosyllactose: ≥ 90 %

Lactose: $\leq 5.0 \%$ Fucose: $\leq 3.0 \%$

3-Fucosyllactose: $\leq 5.0 \%$

pH (20 C, 5 % solution): 3,0-7,5

Water: $\leq 9.0 \%$

Sulphated ash: $\leq 2.0 \%$ Acetic acid: $\leq 1.0 \%$ Residual proteins: $\leq 0.01 \%$

Microbiological criteria:

Aerobic mesophilic bacteria total

count: ≤ 3 000 CFU/g Yeasts: ≤ 100 CFU/g Moulds: ≤ 100 CFU/g Endotoxins: ≤ 10 EU/mg Fucosylgalactose: ≤ 3,0 % Difucosyllactose: ≤ 5,0 %

Glucose: $\leq 3.0 \%$ Galactose: $\leq 3.0 \%$ Water: $\leq 9.0 \%$ (powder)

Ash, sulphated: $\leq 0.5 \%$ (powder

and liquid)

Residual proteins: $\leq 0.01 \%$ (powder and liquid)

Heavy Metals:

Lead: $\leq 0.02 \text{ mg/kg}$ (powder and

liquid)

Arsenic: ≤ 0.2 mg/kg (powder and

liquid)

Cadmium: $\leq 0.1 \text{ mg/kg}$ (powder

and liquid)

Mercury: ≤ 0.5 mg/kg (powder and

liquid)

Microbiological criteria:

Total plate count: $\leq 10^4$ CFU/g (powder), $\leq 5\,000$ CFU/g (liquid) Yeasts and Moulds: ≤ 100 CFU/g (powder); ≤ 50 CFU/g (liquid) Enterobacteriaceae/Coliforms: absence in 11 g (powder and liquid) Salmonella: negative/100 g (powder), negative/200 ml (liquid) Cronobacter: negative/100 g (powder), negative/200 ml (liquid) Endotoxins: ≤ 100 EU/g (powder), ≤ 100 EU/ml (liquid) Aflatoxin M1: $\leq 0,025$ µg/kg (powder and liquid)]

[F612'-Fucosyllactose/ Difucosyllactose mixture ('2'-FL/DFL') (microbial source)

Description / Definition:

2'-Fucosyllactose/Difucosyllactose mixture is a purified, white to off-white [^{F62}powder or agglomerates thereof that is produced by a microbial process].

Source: Genetically modified strain of Escherichia coli strain K-12 DH1

Characteristics/Composition

Appearance: White to off white powder or agglomerates

Sum of 2'-Fucosyllactose, Difucosyllactose, [F63D-Lactose, L-Fucose, and

3-Fucosyllactose] (% of dry matter): \geq 92,0 % (w/w)

Sum of 2'-fucosyllactose and difucosyllactose (% of dry matter): $\geq 85,0$ % (w/w)

2'-Fucosyllactose (% of dry matter): \geq 75,0 % (w/w) Difucosyllactose (% of dry matter): \geq 5,0 % (w/w)

D-Lactose: ≤ 10,0 % (w/w) L-Fucose: ≤ 1,0 % (w/w)

2'-Fucosyl-D-lactulose: \leq 2,0 % (w/w) Sum of other carbohydratesl: \leq 6,0 % (w/w)

Moisture: $\leq 6.0 \%$ (w/w) Ash, sulfated: $\leq 0.8 \%$ (w/w)

pH (20 °C, 5 % solution): 4,0-6,0 Residual protein: \leq 0,01 % (w/w)

Microbiological criteria:

Aerobic mesophilic bacteria total plate count: ≤ 1000 CFU/g

Enterobacteriaceae : ≤ 10 CFU/g Salmonella sp.: Negative/25 g

Yeast: $\leq 100 \text{ CFU/g}$ Mould: $\leq 100 \text{ CFU/g}$

Residual endotoxins: ≤ 10 EU/mg

CFU: Colony Forming Units; EU: Endotoxin Units]

[F642'-

source)

Fucosyllactose/ Difucosyllactose mixture ('2'-FL/DFL') (microbial

Description:

2'-Fucosyllactose/Difucosyllactose mixture is a purified, white to offwhite powder or agglomerate thereof that is produced by a microbial process.

Source:

Genetically modified strain of Escherichia coli K-12 DH1

Characteristics/Composition:

Appearance: White to off white powder or agglomerates

Sum of 2'-Fucosyllactose, Difucosyllactose, D-Lactose, L-Fucose, and 3-

Fucosyllactose (% of dry matter): \geq 92.0 % (w/w)

Sum of 2'-Fucosyllactose and Difucosyllactose (% of dry matter): ≥ 85.0

% (w/w)

2'-Fucosyllactose (% of dry matter): \geq 75.0 % (w/w) Difucosyllactose (% of dry matter): \geq 5.0 % (w/w)

D-Lactose: $\leq 10.0 \%$ (w/w) L-Fucose: $\leq 1.0 \%$ (w/w)

2'-Fucosyl-D-lactulose: $\leq 2.0 \text{ (w/w)}$

Sum of other carbohydrates (11): $\leq 6.0 \%$ (w/w)

Moisture: \leq 6.0 % (w/w) Ash, sulfated: \leq 0.8 % (w/w) pH (20 °C, 5 % solution): 4.0 -6.0 Residual protein: \leq 0.01 % (w/w)

Microbiological criteria:

Aerobic mesophilic total plate count: ≤ 1000 CFU/g Enterobacteriaceae:

 $\leq 10 \text{ CFU/g}$

Salmonella sp.: Negative/25 g

Yeast: $\leq 100 \text{ CFU/g}$ Mould: $\leq 100 \text{ CFU/g}$

Residual endotoxins: ≤ 10 EU/mg

CFU: Colony Forming Units; EU: Endotoxin Units]

[F652'-

Fucosyllactose / Difucosyllactose ('2'-FL/DFL') (microbial source)

Description/Definition:

2'-Fucosyllactose/Difucosyllactose mixture is a purified, white to offwhite powder or agglomerates thereof that is produced by a microbial process.

Source: Genetically modified strain of *Escherichia coli* strain K-12 DH1

Characteristics/Composition:

Appearance: White to off-white powder or agglomerates

Sum of 2'-Fucosyllactose, Difucosyllactose, D-Lactose, L-Fucose, and

3-Fucosyllactose (% of dry matter): \geq 92.0 % (w/w)

Sum of 2'-fucosyllactose and difucosyllactose (% of dry matter): ≥ 85.0

% (w/w)

2'-Fucosyllactose (% of dry matter): ≥ 75.0 % (w/w) Difucosyllactose (% of dry matter): ≥ 5.0 % (w/w) D-Lactose: $\leq 10.0 \%$ (w/w) L-Fucose: $\leq 1.0 \%$ (w/w) 2'-Fucosyl-D-lactulose: $\leq 2.0 \%$ (w/w) Sum of other carbohydrates : $\leq 6.0 \%$ (w/w) Moisture: $\leq 6.0 \%$ (w/w) Ash, sulfated: $\leq 0.8 \%$ (w/w) pH (20 °C, 5 % solution): 4.0-6.0 Residual protein: $\leq 0.01 \%$ (w/w) Microbiological criteria: Aerobic mesophilic bacteria total plate count: ≤ 1000 CFU/g Enterobacteriaceae: ≤ 10 CFU/g Salmonella sp.: Negative/25 g Yeast: $\leq 100 \text{ CFU/g}$ Mould: $\leq 100 \text{ CFU/g}$ Residual endotoxins: ≤ 10 EU/mg CFU: Colony Forming Units; EU: Endotoxin Units] Galacto-**Description/Definition:** oligosaccharide Galacto-oligosaccharide is produced from milk lactose by an enzymatic process using β-galactosidases from Aspergillus oryzae, Bifidobacterium bifidum, Pichia pastoris, Sporobolomyces singularis, Kluyveromyces lactis, Bacillus circulans, and Papiliotrema terrestris. GOS: min 46 % Dry Matter (DM) Lactose: max 40 % DM Glucose: max 27 % DM Galactose: min 0.8 % DM Ash: max 4,0 % DM Protein: max 4,5 % DM Nitrite: max. 2 mg/kg Glucosamine White crystalline odourless powder HCl from Molecular formula: C₆H₁₃NO₅ · HCl Aspergillus Relative molecular mass: 215,63 g/mol D-Glucosamine HCl 98,0-102,0 % of reference standard (HPLC) niger and genetically Specific rotation $+70.0^{\circ} - +73.0^{\circ}$ modified strain of E. coli K-12 Glucosamine White crystalline odourless powder sulphate Molecular formula: $(C_6H_{14}NO_5)_2SO_4 \cdot 2KCl$ KCl from Relative molecular mass: 605,52 g/mol Aspergillus D-Glucosamine Sulphate 2KCl 98,0-102,0 % of reference standard niger and (HPLC) genetically Specific Rotation +50.0° to +52.0° modified strain of E. coli K-12 Glucosamine White crystalline odourless powder sulphate Molecular formula: (C₆H₁₄NO₅)₂SO₄ · 2NaCl NaCl from Relative molecular mass: 573,31 g/mol Aspergillus D-Glucosamine HCl: 98-102 % of reference standard (HPLC) niger and

genetically modified strain of *E. coli* K-12

Specific Optical Rotation: +52° - +54°

Guar Gum

Description/Definition:

Native guar gum is the ground endosperm of seeds from natural strains of guar *Cyamopsis tetragonolobus* L. Taub. (*Leguminosae* family). It consists of a high molecular weight polysaccharide, primarily composed of galactopyranose and mannopyranose units combined through glycosidic linkages, which may be described chemically as a galactomannan (galactomannan content not less than 75 %).

Appearance: White to yellowish powder

Molecular weight: Between 50 000 – 8 000 000 Daltons

CAS number: 9000-30-0 Einecs Number: 232-536-8

Purity: As specified by Commission Regulation (EU) No 231/2012 laying down specifications for food additives listed in Annexes II and III to Regulation (EC) No 1333/2008 of the European Parliament and of the Council^a & by Commission Implementing Regulation (EU) 2015/175 of 5 February 2015 laying down special conditions applicable to the import of guar gum originating in or consigned from India due to contamination risks by pentachlorophenol and dioxins^b.

Physico-chemical properties:

Powder

Shelf-life: 2 years Colour: White Odour: Light

Average diameter of particles: 60-70µm

Moisture: Max 15 %

Viscosity * at 1 hour — Viscosity * at 2 hours: Min 3 600 mPa.s

Viscosity * at 24 hours: Min 4 000 mPa.s Solubility: Soluble in hot and cold water

pH for 10g/L, at 25 °C - 6-7,5

Flakes

Useful life: 1 year

Colour: White/off white with absence or minimal presence of black spots

Odour: Light

Average diameter of particles: 1-10 mm

Moisture: Max 15 %

Viscosity * at 1 hour: Min 3 000 mPa.s

Viscosity * at 2 hours — Viscosity * at 24 hours —

Solubility — Soluble in hot and cold water

pH for 10g/L, at 25 °C - 5-7,5

(*) The measurements of viscosity are carried out under the following conditions: 1 %, 25 °C, 20 rpm

Heat-treated milk products fermented with Bacteroides xylanisolvens

Description/Definition:

Heat-treated fermented milk products are produced with *Bacteroides xylanisolvens* (DSM 23964) as starter culture.

Semi-skimmed milk (between 1,5 % and 1,8 % fat) or skimmed milk (0,5 % fat or less) is pasteurised or ultra-heat-treated before starting the fermentation with *Bacteroides xylanisolvens* (DSM 23964). The

resulting fermented milk product is homogenised and then heat-treated to inactivate *Bacteroides xylanisolvens* (DSM 23964). The final product does not contain viable cells of *Bacteroides xylanisolvens* (DSM 23964) (1).

(1) Modified DIN EN ISO 21528-2.

Hydroxytyrosol

Description/Definition:

Hydroxytyrosol is a pale yellow viscous liquid obtained by chemical

synthesis

Molecular formula: C₈H₁₀O₃ Molecular weight: 154,6 g/mol

CAS No: 10597-60-1 Moisture ≤ 0,4 %

Odour: CharacteristicTaste: Slightly bitter Solubility (water): Miscible with water

pH: 3,5-4,5

Refractive Index: 1,571-1,575

Purity:

Hydroxytyrosol: \geq 99 % Acetic acid: \leq 0,4 %

Hydroxytyrosol acetate: ≤ 0,3 %

Sum of homovanillic acid, iso-homovanilic acid, and 3-

methoxy-4hydroxyphenylglycol: ≤ 0,3 %

Heavy Metals

Lead: ≤ 0,03 mg/kg Cadmium: ≤ 0,01 mg/kg Mercury: ≤ 0,01 mg/kg **Residual Solvents**

Ethyl acetate: \leq 25,0 mg/kg Isopropanol: \leq 2,50 mg/kg Methanol: \leq 2,00 mg/kg Tetrahydrofuran: \leq 0,01 mg/kg

Ice Structuring Protein type III HPLC 12

Description/Definition:

The Ice Structuring Protein (ISP) preparation is a light-brown liquid produced by submerged fermentation of a genetically-modified strain of food-grade baker's yeast (*Saccharomyces cerevisiae*) in which a synthetic gene for the ISP has been inserted into the yeast's genome. The protein is expressed and secreted into the growth medium where it is separated from the yeast cells by micro-filtration and concentrated by ultra-filtration. As a result, the yeast cells are not transferred into the ISP preparation as such or under an altered form. The ISP preparation consists of native ISP, glycosylated ISP and proteins and peptides from the yeast and sugars as well as acids and salts commonly found in food. The concentrate is stabilised with 10 mM citric acid buffer.

Assay: ≥ 5 g/l active ISP

pH: 2,5-3,5 Ash: ≤ 2,0 % DNA: Not detectable

Aqueous extract of dried

Description/Definition:

Dark brown liquid. Aqueous extracts of dried leaves of *Ilex guayusa*. **Composition:**

leaves of *Ilex* guayusa

Protein: < 0,1 g/100 ml Fat: < 0,1 g/100 ml

Carbohydrate: 0,2–0,3 g/100 ml Total sugars: < 0,2 g/100 ml Caffeine: 19,8–57,7 mg/100 ml Theobromine: 0,14–2,0 mg/100 ml Chlorogenic acids: 9,9–72,4 mg/100ml

[F23] Infusion from coffee leaves of Coffea arabica L. and/or Coffea canephora Pierre ex A. Froehner (Traditional food from a third country)

Description/Definition:

The traditional food consists of an infusion of leaves from *Coffea* arabica L. and/or *Coffea* canephora Pierre ex A.Froehner (family:

Rubiaceae).

The traditional food is prepared by mixing a maximum of 20 g of dried leaves from *Coffea arabica* L. and/or *Coffea canephora* Pierre ex A.Froehner with 1 L of hot water. Leaves are removed and the infusion is then subjected to pasteurization (at least 71 °C for 15 seconds).

Composition:

Visual: Brown green liquid Odour and taste: Characteristic

Chlorogenic acid (5-CQA): < 100 mg/L

Caffeine: < 80 mg/L

Epigallocatechin gallate (EGCG): < 700 mg/L

Microbiological criteria: Total plate count: < 500 CFU/g

Total yeast and mould count: < 100 CFU/g

Total coliforms: < 100 CFU/g Escherichia coli: Absence in 1 g Salmonella: Absence in 25 g

Heavy metals:

Lead (Pb): < 3,0 mg/L Arsenic (As): < 2,0 mg/L Cadmium (Cd): < 1,0 mg/L CFU: Colony Forming Units]

Isomaltooligosaccharide

Powder:

Solubility (water) (%): > 99 Glucose (% dry basis): ≤ 5.0

Isomaltose + DP3 to DP9 (% dry basis): \geq 90

Moisture (%): ≤ 4.0

Sulphated ash(g/100 g): ≤ 0.3

Heavy metals:

Lead (mg/kg): ≤ 0.5 Arsenic (mg/kg): ≤ 0.5

Syrup:

Dried solids (g/100 g): > 75Glucose (% dry basis): ≤ 5.0

Isomaltose + DP3 to DP9 (% dry basis): \geq 90

pH: 4 - 6

Sulphated ash(g/100 g): ≤ 0.3

Heavy metals:

Lead (mg/kg): ≤ 0.5 Arsenic (mg/kg): ≤ 0.5

Isomaltulose

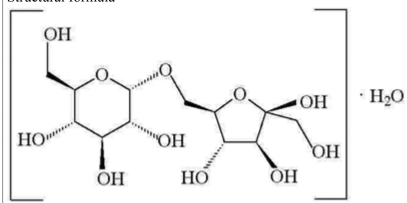
Description/Definition:

A reducing disaccharide that consists of one glucose and one fructose moiety linked by an alpha-1,6-glycosidic bond. It is obtained from sucrose by an enzymatic process. The commercial product is the monohydrate. Appearance: Virtually odourless, white or almost white crystals with a sweet tasteChemical name: $6-O-\alpha-D$ -glucopyranosyl-D-fructofuranose, monohydrate

CAS No.: 13718-94-0

Chemical formula: C₁₂ H₂₂ O₁₁ · H₂ O

Structural formula



Formula weight: 360,3 (monohydrate)

Purity:

Assay: \geq 98 % on the dry basis

Loss on drying: $\leq 6.5 \%$ (60 °C, 5 hours)

Heavy metals: Lead: $\leq 0.1 \text{ mg/kg}$

Determine using an atomic absorption technique appropriate to the specified level. The selection of sample size and method of sample preparation may be based on the principles of the method described in FNP 5(¹), 'Instrumental methods'

Food and Nutrition Paper 5 Rev. 2 — Guide to specifications for general notices, general analytical techniques, identification tests, test solutions and other reference materials (JECFA), 1991, 322 pp., English, ISBN 92-5-102991-1.

Lactitol Description/Definition:

Crystalline powder or colourless solution manufactured via catalytic hydrogenation of lactose. Crystalline products occur in anhydrous, monohydrate and dihydrate forms. Nickel is used as a catalyst.

Chemical name: 4-O-β-D-Galactopyranosyl-D-glucitol

Chemical formula: C₁₂ H₂₄ O₁₁ Molecular weight: 344,31 g/mol CAS No: 585-86-4 **Purity:**

Solubility (in water): Very soluble in water Specific rotation $[\alpha]_D^{20} = +13^\circ \text{ to } +16^\circ$

Assay: ≥ 95 % d.b (d.b — expressed on the dry weight basis)

Water: $\le 10.5 \%$

Other polyols: ≤ 2.5 % d.b Reducing sugars: ≤ 0.2 % d.b Chlorides: ≤ 100 mg/kg d.b

Sulphates: $\leq 200 \text{ mg/kg d.b}$ Sulphated ash: $\leq 0,1 \% \text{ d.b}$ Nickel: $\leq 2,0 \text{ mg/kg d.b}$ Arsenic: $\leq 3,0 \text{ mg/kg d.b}$ Lead: $\leq 1,0 \text{ mg/kg d.b}$

Lacto- N - neotetraose (synthetic)

Definition:

Chemical name: β -D-Galactopyranosyl- $(1\rightarrow 4)$ -2-acetamido-2-deoxy- β -D-glucopyranosyl- $(1\rightarrow 3)$ - β -D-galactopyranosyl- $(1\rightarrow 4)$ - D-

glucopyranose

Chemical formula: C₂₆ H₄₅ NO₂₁

CAS No: 13007-32-4

Molecular weight: 707,63 g/mol

Description:

Lacto-N-neotetraose is a white to off-white powder. Produced by a chemical synthesis process and is isolated by crystallisation.

Purity:

Assay (water free): \geq 96 %

D-Lactose: ≤ 1,0 %

Lacto-N-triose II: $\leq 0.3 \%$

Lacto-N-neotetraose fructose isomer: ≤ 0,6 %

pH (20 °C, 5 % solution): 5,0-7,0

Water: $\leq 9.0 \%$

Ash, sulphated: $\leq 0.4 \%$

Acetic acid: ≤ 0.3 %Residual solvents (methanol, 2-propanol, methyl acetate, acetone): ≤ 50 mg/kg singly, ≤ 200 mg/kg in combination

Residual proteins: ≤ 0,01 % Palladium: ≤ 0,1 mg/kg Nickel: ≤ 3,0 mg/kg Microbiological criteria:

Aerobic mesophilic bacteria total count: ≤ 500 CFU/g

Yeasts: $\leq 10 \text{ CFU/g}$ Moulds: $\leq 10 \text{ CFU/g}$

Residual endotoxins: ≤ 10 EU/mg

[F66Lacto N neotetraose (microbial source)

Definition:

Chemical name: β -D-Galactopyranosyl- $(1\rightarrow 4)$ -2-acetamido-2-deoxy- β -D-glucopyranosyl- $(1\rightarrow 3)$ - β -D-galactopyranosyl- $(1\rightarrow 4)$ -D-glucopyranose

Chemical formula: C₂₆ H₄₅ NO₂₁

CAS No: 13007-32-4

Molecular weight: 707,63 g/mol

Source:

Genetically modified strain of Escherichia coli K-12

Description:

Lacto- *N* -neotetraose is a white to off-white powder that is produced by a microbiological process.

Purity:

Assay (water free): $\geq 80 \%$

D-Lactose: $\leq 10,0 \%$ Lacto- N -triose II: $\leq 3,0 \%$

para -Lacto- N -neohexaose: $\leq 5.0 \%$

Lacto- N -neotetraose fructose isomer: $\leq 1.0 \%$

Sum of saccharides (Lacto- N -neotetraose, D-Lactose, Lacto- N -triose II, para -Lacto- N -neotetraose, Lacto- N -neotetraose fructose isomer): $\geq 92\%$

pH (20 C, 5 % solution): 4,0-7,0

Water: $\leq 9.0 \%$

Ash, sulphated: $\leq 0.4 \%$

Residual solvents (methanol): $\leq 100 \text{ mg/kg}$

Residual proteins: $\leq 0.01 \%$ Microbiological criteria:

Aerobic mesophilic bacteria total count: $\leq 500 \text{ CFU/g}$

Yeasts: $\leq 10 \text{ CFU/g}$ Moulds: $\leq 10 \text{ CFU/g}$

Residual endotoxins: ≤ 10 EU/mg

CFU: Colony Forming Units; EU: Endotoxin Units.]

[F25Lacto- N - tetraose ('LNT') (microbial source)

Definition:

Chemical formula: C₂₆ H₄₅ O₂₁

Chemical name: β -D-Galactopyranosyl- $(1\rightarrow 3)$ -2-acetamido-2-deoxy- β -D-glucopyranosyl- $(1\rightarrow 3)$ - β -D-galactopyranosyl- $(1\rightarrow 4)$ -D-glucopyranose

Molecular mass: 707,63 Da CAS No 14116-68-8

Description:

Lacto-N-tetraose is a purified, white to off-white amorphous powder that is produced by a microbial process.

Source: Genetically modified strain of *Escherichia coli* strain K-12

DH1

Characteristics/Composition:

Appearance: White to off-white powder

Sum of lacto- N -tetraose, D-Lactose and lacto- N -tetraose II (% of dry

matter): $\geq 90.0 \% (w/w)$

Lacto- N -tetraose (% of dry matter): ≥ 70.0 % (w/w)

D-Lactose: $\leq 12,0 \% \text{ (w/w)}$

Lacto- N -tetraose II: $\leq 10.0 \%$ (w/w)

Para -lacto- N -hexaose-2: $\leq 3.5 \%$ (w/w)

Lacto- N -tetraose fructose isomer: $\leq 1,0 \%$ (w/w)

Sum of other carbohydrates: $\leq 5.0 \%$ (w/w)

Moisture: $\leq 6.0 \%$ (w/w) Ash, sulfated: $\leq 0.5 \%$ (w/w) pH (20 °C, 5 % solution): 4,0–6,0 Residual protein: $\leq 0.01 \%$ (w/w)

Microbiological criteria:

Aerobic mesophilic bacteria total plate count: ≤ 1 000 CFU/g

Enterobacteriaceae : ≤ 10 CFU/g *Salmonella* sp.: Negative/25 g

Yeast: $\leq 100 \text{ CFU/g}$ Mould: $\leq 100 \text{ CFU/g}$

Residual endotoxins: ≤ 10 EU/mg

CFU: Colony Forming Units; EU: Endotoxin Units.]

I^{F26}*Lonicera caerulea* L. berries (haskap)

Description/Definition:

The traditional food are fresh and frozen berries from *Lonicera caerulea* var. edulis.

Lonicera caerulea L. is a deciduous shrub belonging to the Caprifoliaceae family.

(Traditional food from a third country)

(Traditional food | Typical nutritional components of haskap berries (given in fresh

berries):

Carbohydrates: 12,8 %

Fibre: 2,1 % Lipids: 0,6 % Proteins: 0,7 % Ash: 0,4 % Water: 85,5 %]

Lucerne leaf extract from *Medicago sativa*

Description/Definition:

The Lucerne (*Medicago sativa* L.) is processed within 2 hours after harvest. It is chopped and crushed. By passing through an oleaginous-type press, the Lucerne provides a fibrous residue and press juice (10% of dry matter). The dry matter of this juice contains about 35% of crude protein. The press juice (pH 5,8-6,2) is neutralised. Preheating and vapour injection allows coagulation of proteins associated with carotenoid and chlorophyll pigments. The protein precipitate is separated by centrifugation and thereafter dried. After adding ascorbic acid the Lucerne protein concentrate is granulated and stored in inert gas or in cold storage.

Composition:

Protein: 45-60 % Fat: 9-11 %

Free carbohydrates (soluble fibre): 1-2 % Polysaccharides (insoluble fibre): 11-15 %

including cellulose: 2-3 %

Minerals: 8-13 % Saponins: ≤ 1,4 % Isoflavones: ≤ 350 mg/kg Coumestrol: ≤ 100 mg/kg Phytates: ≤ 200 mg/kg L-canavanine: ≤ 4,5 mg/kg

Lycopene

Description/Definition:

Synthetic lycopene is produced by the Wittig condensation of synthetic intermediates commonly used in the production of other carotenoids used in food. Synthetic lycopene consists of ≥ 96 % lycopene and minor quantities of other related carotenoid components. Lycopene is presented either as a powder in a suitable matrix or an oily dispersion. The colour is dark red or red-violet. Antioxidative protection has to be assured.

Chemical name: Lycopene

CAS No.: 502-65-8 (all -trans lycopene)

Chemical formula: C ₄₀ H ₅₆ Formula weight: 536,85 Da

Lycopene from *Blakeslea trispora*

Description/Definition:

The purified lycopene from *Blakeslea trispora* consists of \geq 95 % lycopene and \leq 5 % other carotenoids. It is presented either as a powder in a suitable matrix or an oily dispersion. The colour is dark red or redviolet. Anti-oxidative protection has to be assured.

Chemical name: Lycopene

CAS No.: 502-65-8 (all trans lycopene)

Chemical formula: C₄₀H₅₆ Formula weight: 536,85 Da Document Generated: 2024-04-16

Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

Lycopene from tomatoes

Description/Definition:

The purified lycopene from tomatoes (*Lycopersicon esculantum* L.) consists of ≥ 95 % lycopene and ≤ 5 % other carotenoids. It is presented either as a powder in a suitable matrix or an oily dispersion. The colour is dark red or red-violet. Anti-oxidative protection has to be assured.

Chemical name: Lycopene

CAS No.: 502-65-8 (all trans lycopene)

Chemical formula: C₄₀H₅₆ Formula weight: 536,85 Da

Lycopene oleoresin from tomatoes

Description/Definition:

Lycopene oleoresin from tomatoes is obtained by solvent extraction of ripe tomatoes (*Lycopersicon esculentum Mill.*) with subsequent removal of the solvent. It is a red to dark brown viscous, clear liquid.

Total lycopene: 5-15 %

Thereof trans-lycopene: 90-95 %

Total carotenoids (calculated as lycopene): 6,5-16,5 %

Other carotenoids: 1,75 %

(Phytoene/phytofluene/β-carotene): (0,5-0,75/0,4-0,65/0,2-0,35 %)

Total tocopherols: 1,5-3,0 % Unsaponifiable matter: 13-20 % Total fatty acids: 60-75 % Water (Karl Fischer): ≤ 0.5 %

[F16Hen egg white lysozyme hydrolysate

Description/Definition

Hen egg white lysozyme hydrolysate is obtained from hen egg white lysozyme by an enzymatic process, using subtilisin from *Bacillus licheniformis*.

The product is a white to light yellow powder.

Specification

Protein (TN(*) x 5,30): 80-90 %

Tryptophan: 5-7 %

Moisture: < 5 %

Ratio Tryptophan/LNAA(**): 0,18-0.25

Degree of hydrolysis: 19-25 %

Ash: < 10 %
Sodium: < 6 %
Heavy metals
Arsenic: < 1 ppm
Lead: < 1 ppm
Cadmium: < 0,5 ppm
Mercury: < 0,1 ppm
Microbiological criteria

Total aerobic count: < 10³ CFU/g

Total combined yeasts/moulds count: $< 10^2$ CFU/g

Enterobacteria: < 10 CFU/g

Salmonella spp: Absence in 25 g

Escherichia coli: Absence in 10 g

Staphylococcus aureus: Absence in 10 g

Pseudomonas aeruginosa: Absence in 10 g

* TN: total nitrogen

** LNAA: large neutral amino acids]

Magnesium citrate malate

Description/Definition:

Magnesium citrate malate is a white to yellowish-white, amorphous powder. Chemical formula: Mg₅(C₆H₅O₇)₂(C₄H₄O₅)₂

Chemical name: Pentamagnesium di-(2-hydroxybutanedioate)-di-(2-

hydroxypropane-1,2,3-tricarboxylate)

CAS No.: 1259381-40-2

Molecular weight: 763,99 Daltons (anhydrous)

Solubility: Freely soluble in water (about 20 g in 100 ml) Description of the physical state: Amorphous powder

Assay magnesium: 12,0-15,0 %

Loss on drying (120 °C/4 hours): \leq 15 % Colour (solid): White to yellowish-white

Colour (20 % aqueous solution): Colourless to yellowish Appearance (20 % aqueous solution): Clear solution

pH (20 % aqueous solution): Approx. 6,0

Impurities:

Chloride: $\leq 0.05 \%$ Sulphate: $\leq 0.05 \%$ Arsenic: ≤ 3.0 ppm Lead: ≤ 2.0 ppm Cadmium: ≤ 1 ppm Mercury: ≤ 0.1 ppm

Magnolia Bark Extract

Description/Definition:

Magnolia bark extract is obtained from the bark of the plant *Magnolia officinalis* L. and produced with supercritical carbon dioxide. The bark is washed and oven dried to reduce moisture content before being crushed and extracted with supercritical carbon dioxide. The extract is dissolved in medical-grade ethanol and re-crystallised to yield magnolia bark extract.

Magnolia bark extract is mainly composed of two phenolic compounds, magnolol and honokiol.

Appearance: Light brownish powder

Purity:

Magnolol: \geq 85,2 % Honokiol: \geq 0,5 %

Magnolol & Honokiol: ≥ 94 %

Total Eudesmol: $\leq 2 \%$

Moisture: 0,50 % **Heavy metals:**

Arsenic (ppm): ≤ 0.5 Lead (ppm): ≤ 0.5

Methyl eugenol (ppm): ≤ 10 Tubocurarine (ppm): ≤ 2.0 Total Alkaloid (ppm): ≤ 100

Maize-germ oil high in unsaponifiable matter

Description/Definition:

Maize-germ oil high in unsaponifiable matter is produced by vacuum distillation and it is different from refined maize-germ oil in the concentration of the unsaponifiable fraction (1,2 g in refined maize-germ oil and 10 g in 'maize-germ oil high in unsaponifiable matter').

Purity:

Unsaponifiable matter: > 9,0 g/100 g

Tocopherols: $\geq 1.3 \text{ g/}100 \text{ g}$ α-tocopherol (%): 10-25 % β -tocopherol (%): < 3.0 % γ-tocopherol (%): 68-89 % δ -tocopherol (%): < 7.0 % Sterols, triterpenic alcohols, methylsterols: > 6,5 g/100 g Fatty acids in triglycerides: palmitic acid: 10,0-20.0 % stearic acid: < 3,3 % oleic acid: 20,0-42,2 % linoleic acid: 34,0-65,6 % linolenic acid: < 2,0 % Acid value: $\leq 6.0 \text{ mg KOH/g}$ Peroxide value (PV): $\leq 10 \text{ mEq O}_2/\text{kg}$ **Heavy metals:** Iron (Fe): $< 1500 \,\mu g/kg$ Copper (Cu): $< 100 \mu g/kg$

Impurities:

Polycyclic aromatic hydrocarbons (PAH) Benzo(a)pyrene: $< 2 \mu g/kg$ Treatment with active carbon is required to ensure that polycyclic aromatic hydrocarbons (PAH) are not enriched in the production of 'maize-germ oil high in unsaponifiable matter'

Methylcellulose

Description/Definition:

Methyl cellulose is cellulose obtained directly from natural strains of fibrous plant material and partially etherified with methyl groups.

Chemical name: Methyl ether of cellulose

Chemical formula: The polymers contain substituted anhydroglucose units with the following general formula:

C6H7O2(OR1)(OR2)(OR3) where R1, R2, R3 each may be one of the following:

— Н — СН₃ or — СН₂СН₃

Molecular weight: Macromolecules: from about 20 000 (n about 100) up to about 380 000 g/mol (n about 2 000)

Assay: Content not less than 25 % and not more than 33 % of methoxyl groups (-OCH 3) and not more than 5 % of hydroxyethoxyl groups (-OCH 2 CH 2 OH)

Slightly hygroscopic white or slightly yellowish or greyish odourless and tasteless, granular or fibrous powder.

Solubility: Swelling in water, producing a clear to opalescent, viscous, colloidal solution. Insoluble in ethanol, ether and chloroform. Soluble in glacial acetic acid.

Purity:

Loss on drying: $\leq 10 \%$ (105 °C, 3 hours)

Sulphated Ash: ≤ 1.5 % determined at 800 ± 25 °C

pH: \geq 5,0 and \leq 8,0 (1 % colloidal solution)

Heavy metals:

Arsenic: ≤ 3,0 mg/kg Lead: ≤ 2,0 mg/kg Mercury: ≤ 1,0 mg/kg Cadmium: ≤ 1,0 mg/kg

[F27]1- Definition:

Methylnicotinami@emical name: 3-carbamoyl-1-methyl-pyridinium chloride

chloride Chemical formula: C₇H₉N₂OCl

CAS No: 1005-24-9 Molecular weight: 172,61 Da

Description

1-Methylnicotinamide chloride is white or off-white, crystalline solid

produced by a chemical synthesis process.

Characteristics/Composition

Appearance: White – off-white, crystalline solid

Purity: \geq 98,5 % Trigonelline: \leq 0,05 % Nicotinic Acid: \leq 0,10 % Nicotinamide: \leq 0,10 %

Largest unknown impurity: $\leq 0.05 \%$ Sum of unknown impurities: $\leq 0.20 \%$ Sum of all impurities: $\leq 0.50 \%$

Solubility: soluble in water and methanol. Practically insoluble in 2-

propanol and dichloromethane

Moisture: $\leq 0.3 \%$ Loss on drying: $\leq 1.0 \%$ Residue on ignition: $\leq 0.1 \%$

Residual Solvents and Heavy Metals

Methanol: $\leq 0.3 \%$ Heavy metals: $\leq 0.002 \%$ **Microbiological criteria:**

Total aerobic microbial count: ≤ 100 CFU/g

Mould/yeast: ≤ 10 CFU/g

Enterobacteriaceae: absence in 1 g *Pseudomonas aeruginosa*: absence in 1 g *Staphylococcus aureus*: absent in 1 g

CFU: Colony Forming Units]

(6S)-5- Description/Definition:

methyltetrahydrofoliemical name: N-[4-[[[(6S)-2-amino-1,4,5,6,7,8-hexahydro-5-methyl-4-acid, oxo-6-pteridinyl]methyl]amino]benzoyl]-L-glutamic acid, glucosamine

glucosamine salt

salt Chemical formula: C₃₂ H₅₁ N₉ O₁₆

Molecular weight: 817,80 g/mol (anhydrous)

CAS No.: 1181972-37-1

Appearance: Creamy to light-brown powder **Purity:**

Diastereoisomeric purity: At least 99 % of (6S)-5-methyltetrahydrofolic

acid

Glucosamine assay: 34-46 % in dry basis

5-Methyltetrahydrofolic acid assay: 54-59 % in dry basis

Water: $\leq 8.0 \%$ **Heavy metals:** Lead: $\leq 2.0 \text{ ppm}$ Cadmium: $\leq 1.0 \text{ ppm}$ Mercury: $\leq 0.1 \text{ ppm}$ Arsenic: $\leq 2.0 \text{ ppm}$ Boron: $\leq 10 \text{ ppm}$

Microbiological criteria:

Total aerobic microbial count: ≤ 100 CFU/g

Yeasts and moulds: ≤ 100 CFU/g *Escherichia coli*: Absence in 10g

Monomethylsilan Deixeription/Definition:

(Organic Silicon)

Chemical name: Silanetriol, 1-methyl-Chemical formula: CH₆O₃Si

Molecular weight: 94,14 g/mol

CAS No: 2445-53-6

Purity:

Organic Silicon (monomethylsilanetriol) preparation (aqueous solution):

Acidity (pH): 6,4-6,8 Silicon: 100-150 mg Si/l

Heavy metals: Lead: $\leq 1,0~\mu g/l$ Mercury: $\leq 1,0~\mu g/l$ Cadmium: $\leq 1,0~\mu g/l$ Arsenic: $\leq 3,0~\mu g/l$

Solvents:

Methanol: $\leq 5.0 \text{ mg/kg}$ (residual presence)

Mycelial extract from Shiitake mushroom (Lentinula edodes)

Description/Definition:

The novel food ingredient is a sterile aqueous extract obtained from the mycelium of *Lentinula edodes* cultivated in a submerged fermentation. It is a light brown, slightly turbid liquid.

Lentinan is a β -(1-3) $\bar{\beta}$ -(1-6)-D-glucan which has a molecular weight of approximately 5×10^5 Daltons, a degree of branching of 2/5 and a triple helical tertiary structure.

Purity/Composition of the mycelial extract from Lentinula edodes:

Moisture: 98 % Dry matter: 2 %

Free glucose: < 20 mg/ml Total protein(¹): < 0,1 mg/ml

N-containing constituents (2): < 10 mg/ml

Lentinan: 0.8 - 1.2 mg/ml

(1) Bradford method

(2) Kjeldahl method

[F28Nicotinamide riboside chloride

Description/Definition:

The novel food is a synthetic form of nicotinamide riboside. The novel food contains ≥ 90 % nicotinamide riboside chloride, predominantly in its β form, the remaining components being residual solvents, reaction by-products and degradation products.

Nicotinamide riboside chloride: CAS number: 23111-00-4 EC number: 807-820-5

IUPAC name: 1-[(2R,3R,4S,5R)-3,4-dihydroxy-5-

(hydroxymethyl)oxolan-2-yl]pyridin-1-ium-3-carboxamide;chloride

Chemical formula: C₁₁ H₁₅ N₂O₅Cl Molecular weight: 290,7 g/mol **Characteristics/Composition:** Colour: White to light brown

Form: Powder

Identification: Conforms by NMR (nuclear magnetic resonance)

Nicotinamide riboside chloride: ≥ 90 %

Water content: ≤ 2 % **Residual solvents:** Acetone: $\leq 5~000$ mg/kg Methanol: $\leq 1~000$ mg/kg Acetonitrile: ≤ 50 mg/kg

Methyl tert-butyl ether: $\leq 500 \text{ mg/kg}$

Reaction by-products:
Methyl acetate: ≤ 1 000 mg/kg
Acetamide: ≤ 27 mg/kg

Acetamide: $\leq 27 \text{ mg/kg}$ Acetic acid: $\leq 5 000 \text{ mg/kg}$ **Heavy metals:**

Arsenic: ≤ 1 mg/kg
Microbiological criteria:

Total Plate Count: ≤ 1 000 CFU/g Yeast and Mould: ≤ 100 CFU/g Escherichia coli: Absence in 10 gl

Noni fruit juice (*Morinda citrifolia*)

Description/Definition:

Noni fruits (fruits of *Morinda citrifolia* L.) are pressed. The obtained juice is pasteurised. An optional fermentation step before or after the

pressing may occur. Rubiadin: ≤ 10 μg/kg Lucidin: ≤ 10 μg/kg

Noni fruit juice powder (*Morinda* citrifolia)

Description/Definition:

Seeds and skin of the sun-dried fruits of *Morinda citrifolia* are separated. The obtained pulp is filtered to separate juice from the flesh. Desiccation of the produced juice occurs in one or two ways: Either by atomisation using maize maltodextrins, this mixture is obtained by keeping the rates of inflow of the juice and maltodextrins constant Or by zeodratation or drying and then mixing with an excipient, this process allows the juice to be dried initially and then mixed with

maltodextrins (same amount as used in atomisation).

Noni fruit puree and concentrate (*Morinda* citrifolia)

Description/Definition:

The fruits of *Morinda citrifolia* are harvested by hand. Seeds and skin may be separated mechanically from the pureed fruits. After pasteurisation, the puree is packaged in aseptic containers and stored under cold conditions.no_br *Morinda citrifolia* concentrate is prepared from *M. citrifolia* puree by treatment with pectinolytic enzymes (50–60 °C for 1-2 h). Then the puree is heated to inactivate the pectinases and then immediately cooled. The juice is separated in a decanter centrifuge. Afterwards the juice is collected and pasteurised, prior to being concentrated in a vacuum evaporator from a brix of 6 to 8 to a brix of 49 to 51 in the final concentrate.

Composition:

Puree:

Moisture: 89-93 % Protein: < 0,6 g/100 g Fat: ≤ 0,4 g/100 g Ash: < 1,0 g/100 g

Total carbohydrates: 5-10 g/100 g

Fructose: 0,5-3,82 g/100 g Glucose: 0,5-3,14 g/100 g Dietary fibre: < 0,5-3 g/100 g

5,15-dimethylmorindol (1): $\leq 0.254 \,\mu\text{g/ml}$

Lucidin (1): Not detectable Alizarin (1): Not detectable Rubiadin (1): Not detectable

Concentrate:

Moisture: 48-53 % Protein: 3-3,5 g/100 g Fat: < 0,04 g/100 g Ash: 4,5-5,0 g/100 g

Total carbohydrates: 37-45 g/100 g

Fructose: 9-11 g/100 g Glucose: 9-11 g/100 g Dietary fibre: 1,5-5,0 g/100 g

5,15-dimethylmorindol (1): $\leq 0,254 \,\mu\text{g/ml}$

(1) By an HPLC-UV method developed and validated for the analysis of anthraquinones in Morinda citrifolia puree and concentrate. Limits of detection: 2,5 ng/ml (5,15 dimethylmorindol); 50,0 ng/ml (lucidin); 6,3 ng/ml (alizarin) and 62,5 ng/ml (rubiadin).

Noni leaves (*Morinda citrifolia*)

Description/Definition:

After cutting, the leaves of *Morinda citrifolia* are subject to drying and roasting steps. The product has a particle size ranging from broken leaves to coarse powder with fines. It is of greenish brown to brown colour.

Purity/Composition:

Moisture: < 5,2 % Protein: 17- 20 % Carbohydrate: 55-65 %

Ash: 10-13 % Fat: 4-9 %

Oxalic acid: < 0,14 % Tannic acid: < 2,7 %

5,15-dimethylmorindol: < 47 mg/kgRubiadin: non detectable, $\le 10 \mu\text{g/kg}$ Lucidin: non detectable, $\le 10 \mu\text{g/kg}$

Noni fruit powder (Morinda citrifolia)

Description/Definition:

Noni fruit powder is made from pulped noni (*Morinda citrifolia L.*) fruits by freeze-drying. Fruits are pulped and seeds are removed. After freeze-drying during which water is removed from noni fruits, the remaining noni pulp is milled to a powder and encapsulated.

Purity/Composition

Moisture: 5,3-9 % Protein: 3,8-4,8 g/100 g Fat: 1-2 g/100 g

Ash: 4,6-5,7 g/100 g

Total carbohydrates: 80-85 g/100 g Fructose: 20,4-22,5 g/100 g Glucose: 22-25 g/100 g

	Dietary fibre: 15,4-24,5 g/100 g 5,15-dimethylmorindol (1): \leq 2,0 μ g/ml
	(1) By an HPLC-UV method developed and validated for the analysis of anthraquinones in Morinda citrifolia fruit powder. Limits of detection: 2,5 ng/ml (5,15 dimethylmorindol)
Odontella aurita microalgae	Silicon: 3,3 % Crystalline silica: max 0,1-0,3 % as impurity
Oil enriched with phytosterols/phytostanols	Description/Definition: Oil enriched with phytosterols/phytostanols is composed of an oil fraction and a phytosterol fraction. Acylglycerol Distribution: Free fatty acids (expressed as oleic acid): $\leq 2,0$ % Monoacylglycerols (MAG): ≤ 10 % Diacylglycerols (DAG): ≤ 25 % Triacylglycerols (TAG): Making up the balance Phytosterol fraction: β-sitosterol: ≤ 80 % β-sitostanol: ≤ 15 % campesterol: ≤ 40 % campestanol: $\leq 5,0$ % stigmasterol: ≤ 30 % brassicasterol $\leq 3,0$ % other sterols/stanols: $\leq 3,0$ % Others: Moisture and volatile: $\leq 0,5$ % Peroxide value (PV): $< 5,0$ meq/kg Trans fatty acids: ≤ 1 % Contamination/Purity (GC-FID or equivalent method) of phytosterols/phytostanols: Phytosterols and phytostanols extracted from sources other than vegetable oil suitable for food have to be free of contaminants, best ensured by a purity of more than 99 %.
Oil extracted from squids	Acid value: ≤ 0.5 KOH/g oil Peroxide value (PV): ≤ 5 meq O $_2$ /kg oil p-Anisidine value: ≤ 20 Cold test at 0 °C: ≤ 3 hours Moisture: ≤ 0.1 % (w/w) Unsaponifiable matter: ≤ 5.0 %Trans fatty acids: ≤ 1.0 % Docosahexaeonic acid: ≥ 20 % Eicosapentaenoic acid: ≥ 10 %
[FSPartially defatted chia seed (Salvia hispanica) powders	Description/Definition: The novel foods are partially defatted chia seed (Salvia hispanica) powders obtained by pressing and grinding of the whole seeds of Salvia hispanica L. Physical—sensorial: Foreign matter: 0,1 %

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Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

	Powder with high protein content	Powder with high fibre content	
Particle size	≤ 130 μm	≤ 400 μm	
Chemical compo	sition:		
	Salvia hispanica powder with high protein content	Salvia hispanica powder with high fibre content	
Moisture	≤ 9,0 %	≤ 9,0 %	
Protein	≥ 40,0 %	≥ 24,0 %	
Fat	≤ 17 %	≤ 12 %	
Fibre	≤ 30 %	≥ 50 %	

Microbiological criteria:

Total plate count: ≤ 10 000 CFU/g

Yeasts: ≤ 500 CFU/g Moulds: ≤ 500 CFU/g

 $Staphylococcus\ aureus: \le 10\ CFU/g$

Coliforms: < 100 MPN/g

Enterobacteriaceae: ≤ 100 CFU/g
Bacillus cereus: ≤ 50 CFU/g
Escherichia coli: < 10 MPN/g
Listeria monocytogenes: Absence/g
Salmonella spp.: Absence in 25 g
Contaminants:

Arsenic: $\leq 0,1$ ppm Cadmium: $\leq 0,1$ ppm Lead: $\leq 0,1$ ppm Mercury: $\leq 0,1$ ppm Total aflatoxins: ≤ 4 ppb Ochratoxin A: ≤ 1 ppb]

Pasteurised fruit-based preparations produced using high-pressure treatment

_ FF-1			
Parameter	Target	Comments	
Fruit storage before high- pressure treatment	Minimum 15 days at – 20 °C	Fruit harvested and stored in conjunction with good/hygienic agricultural and manufacturing practices	
Fruit added	40 % to 60 % of thawed fruit	Fruit homogenised and added to other ingredients	
pН	3,2 to 4,2		
° Brix	7 to 42	Assured by added sugars	
a w	< 0,95	Assured by added sugars	
Final storage	60 days maximum at + 5 °C maximum	Equivalent to storage regimen for conventionally processed product	

[F29PhenylcapsaiciPescription/Definition:

Phenylcapsaicin (N-[(4-hydroxy-3-methoxyphenyl)methyl]-7-phenylhept-6-ynamide, C $_{21}$ H $_{23}$ NO $_3$, CAS no: 848127-67-3), is synthesized chemically via a two step synthesis process involving in a first step the production of the acetylenic acid intermediate through a reaction of phenyl acetylene with a carboxylic acid derivative, and in a second step a series of reactions of the acetylenic acid intermediate with vanillylamine derivative to produce phenylcapsaicin.

Characteristics/Composition:

Purity (% of dry matter): \geq 98 %

Moisture: $\leq 0.5 \%$

Total synthesis related production by-products: $\leq 1.0 \%$

N,N -dimethyl formamide: $\leq 880 \text{ mg/kg}$

Dichloromethane: ≤ 600 mg/kg Dimethoxyethane: ≤ 100 mg/kg

Ethyl acetate: $\leq 0.5 \%$ Other solvents: $\leq 0.5 \%$

Heavy metals:

Lead: ≤ 1.0 mg/kg Cadmium: ≤ 1.0 mg/kg Mercury: ≤ 0.1 mg/kg Arsenic: ≤ 1.0 mg/kg **Microbiological criteria:**

Total plate count: ≤ 10 CFU/g

Coliforms: $\leq 10 \text{ CFU/g}$

Escherichia coli : Negative/10 g Salmonella sp.: Negative/10 g Yeast and mould: ≤ 10 CFU/g CFU: Colony Forming Units]

Phosphated maize starch

Description/Definition:

Phosphated maize starch (phosphated distarch phosphate) is a chemically modified resistant starch derived from high amylose starch by combining chemical treatments to create phosphate cross-links between carbohydrate residues and esterified hydroxyl groups.

The novel food ingredient is a white or nearly white powder.

CAS No: 11120-02-8

Chemical formula: $(C_6H_{10}O_5)_n$ [$(C_6H_9O_5)_2PO_2H$]x [$(C_6H_9O_5)_2PO_2H$]x

 PO_3H_2

n = number of glucose units; x, y = degrees of substitutionThe chemical characteristics of phosphated distarch phosphate:

Loss on drying: 10-14 %

pH: 4,5-7,5

Dietary fibre: $\geq 70 \%$ Starch: 7-14 % Protein: $\leq 0.8 \%$ Lipids: $\leq 0.8 \%$

Residual bound phosphorus: ≤ 0.4 % (as phosphorus) 'high amylose

maize' as source

Phosphatidylserin Description/Definition:

from fish phospholipids

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Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

The novel food ingredient is yellow to brown powder.

Phosphatidylserine is obtained from fish phospholipids by an enzymatic transphosphorylation with the amino acid L-serine.

Specification of the phosphatidylserine product manufactured from fish phospholipids:

Moisture: < 5,0 % Phospholipids: $\geq 75 \%$ Phosphatidylserine: $\geq 35 \%$ Glycerides: < 4,0 % Free L-serine: < 1,0 %

Tocopherols: $< 0.5 \% (^1)$

Peroxide value (PV): $< 5.0 \text{ meq O}_2/\text{kg}$

Tocopherols may be added as antioxidants according to $(^1)$ Commission Regulation (EU) No 1129/2011

PhosphatidylserinDescription/Definition:

from soya phospholipids

The novel food ingredient is off-white to light yellow powder. It is also available in liquid form with a clear brown to orange colour. The liquid form contains medium chain triacylglycerides (MCT) as a carrier. It contains lower levels of Phosphatidylserine due to the fact that it includes significant amounts of oil (MCT).

Phosphatidylserine from soya phospholipids is obtained through enzymatic transphosphatidylation of high-phosphatidylcholine soybean lecithin with the amino acid L-serine. Phosphatidylserine consists of a glycerophosphate skeleton conjugated with two fatty acids and L-serine via a phosphodiester linkage.

Characteristics of Phosphatidylserine from soya phospholipids: Powder form:

Moisture: < 2.0 %Phospholipids: $\geq 85 \%$ Phosphatidylserine: ≥ 61 %

Glycerides: < 2.0 % free L-serine: < 1.0 % Tocopherols: < 0,3 % Phytosterols: < 0,2 %

Liquid form: Moisture: < 2,0 %

Phospholipids: ≥ 25 %Phosphatidylserine: ≥ 20 %

Glycerides: not applicable free L-serine: < 1.0 % Tocopherols: < 0,3 % Phytosterols: < 0,2 %

Phospholipid product containing

egual amounts of

Description/Definition:

The product is manufactured through enzymatic conversion of soy lecithin. The phospholipid product is a highly concentrated, yellowbrown powder form of phosphatidylserine and phosphatidic acid at an equal level.

phosphatidylserin specification of the product:

and phosphatidic acid

Moisture: $\leq 2.0 \%$

Total phospholipids: $\geq 70 \%$ Phosphatidylserine: $\geq 20 \%$ Phosphatidic acid: ≥ 20 %

N. J. P. J.	Glycerides: ≤ 1,0 % Free L-serine: ≤ 1,0 % Tocopherols: ≤ 0,3 % Phytosterols: ≤ 2,0 % Silicon dioxide is used with a maximum content of 1,0 %
Phospholipides from egg yolk	85 % and 100 % pure Phospholipides from egg yolk
Phytoglycogen	Description: White to off-white powder which is an odourless, colourless, flavourless polysaccharide derived from non-GM sweet corn using conventional food processing techniques Definition: Glucose polymer ($C_6H_{12}O_6$)n with linear linkages of $\alpha(1-4)$ glycosidic bonds branched every 8 to 12 glucose units by $\alpha(1-6)$ glycosidic bonds Specifications: Carbohydrates: 97 % Sugars: 0,5 % Fibre: 0,8 % Fat: 0,2 % Protein: 0,6 %
Phytosterols/phytostanols	Phytosterols and phytostanols are sterols and stanols that are extracted from plants and may be presented as free sterols and stanols or esterified with food grade fatty acids. Composition (with GC-FID or equivalent method): β-sitosterol: < 81 % β-sitostanol: < 35 % campesterol: < 40 % campestanol: < 15 % stigmasterol: < 30 % brassicasterol: < 3,0 % other sterols/stanols: < 3,0 % Contamination/Purity (GC-FID or equivalent method): Phytosterols and phytostanols extracted from sources other than vegetable oil suitable for food have to be free of contaminants, best ensured by a purity of more than 99 % of the phytosterol/phytostanol ingredient.
Plum kernel oil	Description/Definition: Plum kernel oil is a vegetable oil obtained by cold pressing of plum (Prunus domestica) kernels. Composition: Oleic acid (C18:1): 68 % Linoleic acid (C18:2): 23 % γ-Tocopherol:80 % of total tocopherols β-Sitosterol: 80-90 % of total sterols Triolein: 40-55 % of triglycerides Cyanhydric acid: maximum 5 mg/kg oil
Potato proteins (coagulated) and	Dry substance: $\geq 800 \text{ mg/g}$ Protein (N * 6,25): $\geq 600 \text{ mg/g}$ (dry substance) Ash: $\leq 400 \text{ mg/g}$ (dry substance) Glycoalkaloid (total): $\leq 150 \text{ mg/kg}$

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thereof Prolvl oligopeptidase

preparation)

(enzyme

hydrolysates

Lysinoalanine (total): $\leq 500 \text{ mg/kg}$ Lysinoalanine (free): $\leq 10 \text{ mg/kg}$

Specification of the enzyme: Systematic name: Prolyl oligopeptidase

Synonyms: Prolyl endopeptidase, proline-specific endopeptidase,

endoprolylpeptidase Molecular weight: 66 kDa

Enzyme Commission number: EC 3.4.21.26

CAS number: 72162-84-6

Source: A genetically modified strain of Aspergillus niger (GEP-44)

Description: Prolyl oligopeptidase is available as an enzyme preparation containing approximately 30 % maltodextrin.

Specifications of the enzyme preparation of prolyl oligopeptidase:

Activity: $> 580\ 000\ PPI(^1)/g (> 34.8\ PPU(^2)/g)$

Appearance: Microgranulate

Colour: Off-white to orange yellowish. The colour may change from

batch to batch Dry Matter: > 94 % Gluten: < 20 ppm **Heavy metals:** Lead: $\leq 1.0 \text{ mg/kg}$ Arsenic: $\leq 1.0 \text{ mg/kg}$

Cadmium: $\leq 0.5 \text{ mg/kg}$ Mercury: $\leq 0.1 \text{ mg/kg}$ Microbiological criteria:

Total aerobic plate count: $\leq 10^3$ CFU/g Total yeasts and moulds: $\leq 10^2$ CFU/g Sulphite reducing anaerobes: ≤ 30 CFU/g

Enterobacteriaceae : < 10 CFU/g Salmonella: Absence in 25 g Escherichia coli: Absence in 25 g Staphylococcus aureus: Absence in 10 g Pseudomonas aeruginosa: Absence in 10 g Listeria monocytogenes: Absence in 25 g

Antimicrobial activity: AbsentMycotoxins: Below limits of detection: Aflatoxin B1, B2, G1, G2 ($< 0.25 \mu g/kg$), total Aflatoxins ($< 2.0 \mu g/kg$), Ochratoxin A ($< 0.20 \mu g/kg$), T-2 Toxin ($< 5 \mu g/kg$), Zearalenone ($< 2.5 \mu g/kg$) $\mu g/kg$), Fumonisin B1 and B2 (< 2,5 $\mu g/kg$)

- $(^1)$ PPI - Protease Picomole International
- PPU Prolyl Peptidase Units or Proline Protease Units $(^2)$

[F30Protein extract from pig kidneys

Description/Definition:

The protein extract is obtained from homogenised pig kidneys through a combination of salt precipitation and high speed centrifugation. The obtained precipitate contains essentially proteins with 7 % of the enzyme diamine oxidase (enzyme nomenclature E.C. 1.4.3.22) and is resuspended in a physiologic buffer system. The obtained pig kidney extract is formulated as encapsulated enteric coated pellets or enteric coated tablets to reach the active sites of digestion.

Basic Product:

Specification: pig kidney protein excerpt with natural content of Diamine

oxidase (DAO):

Physical condition: liquid

Colour: brownish

Appearance: slightly turbid solution

pH value: 6,4-6,8

Enzymatic activity: > 2 677 kHDU DAO/ml (DAO REA (DAO

Radioextractionassay)) **Microbiological criteria:**

Brachyspira spp.: negative (Real Time PCR)

Listeria monocytogenes : negative (Real Time PCR)

Staphylococcus aureus : < 100 CFU/g

Influenza A: negative (Reverse Transcription Real Time PCR)

Escherichia coli : < 10 CFU/g

Total aerobic microbiological count: < 10⁵ CFU/g

Yeasts/moulds count: < 10⁵ CFU/g

Salmonella: Absence/10g

Bile salt resistant enterobacteriaceae: < 10⁴ CFU/g

Final product:

Specification pig kidney protein excerpt with natural content of DAO

(E.C. 1.4.3.22) in an enteric coated formulation:

Physical condition: solid Colour: yellow grey

Appearance: micropellets or tablets

Enzymatic activity: 110-220 kHDU DAO/g pellet or g tablet (DAO REA

(DAO Radioextractionassay))

Acid stability 15 min 0,1M HCl followed by 60 min Borat pH = 9,0: > 68 kHDU DAO/g pellet or g tablet (DAO REA (DAO

Radioextractionassay)) Humidity: < 10 %

Staphylococcus aureus: < 100 CFU/g

Escherichia coli : < 10 CFU/g

Total aerobic microbiological count: < 10⁴ CFU/g Total combined yeasts/moulds count: < 10³ CFU/g

Salmonella: Absence/10g

Bile salt resistant enterobacteriaceae: < 10² CFU/g]

[F31Pyrrologuinolinefinition:

quinone disodium salt Chemical name: disodium 9-carboxy-4,5-dioxo-1 H-pyrrolo[5,4-

f]quinoline-2,7-dicarboxylate

Chemical formula: C₁₄H₄N₂Na₂O₈

CAS No: 122628-50-6 Molecular weight: 374,17 Da

Description

Pyrroloquinoline quinone disodium salt is a reddish-brown powder produced by the non-genetically modified bacterium *Hyphomicrobium denitrificans* strain CK-275.

Characteristics/Composition

Appearance: Reddish-brown powder

Purity: ≥ 99,0 % (dry weight)

UV absorbance (A322/A259): 0.56 ± 0.03 UV absorbance (A233/A259): 0.90 ± 0.09

Moisture: ≤ 12,0 %

Residual Solvent

Ethanol: ≤ 0,05 %

Heavy metals

Lead: < 3 mg/kg

Arsenic: < 2 mg/kg

Microbiological criteria:

Total viable cell count: ≤ 300 CFU/g

Mould/yeast: ≤ 12 CFU/g Coliforms: absent in 1 g

Hyphomicrobium denitrificans : ≤ 25 CFU/g

CFU: Colony Forming Units]

Rapeseed oil high in unsaponifiable matter

Description/Definition:

Rapeseed oil high in unsaponifiable matter' is produced by vacuum distillation and it is different from refined rapeseed oil in the concentration of the unsaponifiable fraction (1 g in refined rapeseed oil and 9 g in 'rapeseed oil high in unsaponifiable matter'). There is a minor reduction of triglycerides containing monounsaturated and polyunsaturated fatty acids.

Purity:

Unsaponifiable matter: > 7,0 g/100 g

Tocopherols: > 0.8 g/100 g α-tocopherol (%): 30-50 % γ-tocopherol (%): 50-70 % δ-tocopherol (%): < 6.0 %

Sterols, triterpenic alcohols, methylsterols: > 5,0 g/100 g

Fatty acids in triglycerides:

palmitic acid: 3-8 % stearic acid: 0,8-2,5 % oleic acid: 50-70 % linoleic acid: 15-28 % linolenic acid: 6-14 % erucic acid: < 2,0 %

Acid value: $\leq 6.0 \text{ mg KOH/g}$

Peroxide value (PV): $\leq 10 \text{ mEq O}_2/\text{kg}$

Heavy metals:

Iron (Fe): < 1 000 μg/kg Copper (Cu): < 100 μg/kg

Impurities:

Polycyclic aromatic hydrocarbons (PAH) Benzo(a)pyrene: $< 2 \mu g/kg$ Treatment with active carbon is required to ensure that polycyclic aromatic hydrocarbons (PAH) are not enriched in the production of 'rapeseed oil high in unsaponifiable matter.

Rapeseed Protein

Definition

Rapeseed protein is an aqueous protein-rich extract from rapeseed press cake originating from non-genetically modified *Brassica napus* L. and *Brassica rapa* L.

Description:

White to off-white, spray dried powder

Total protein: $\geq 90 \%$ Soluble protein: $\geq 85 \%$ Moisture: $\leq 7,0 \%$

Carbohydrates: ≤ 7,0 %

Fat: ≤ 2,0 % Ash: ≤ 4,0 % Fibre: ≤ 0,5 %

Total glucosinolates: ≤ 1 mmol/kg

Purity:

Total phytate: $\leq 1,5 \%$ Lead: $\leq 0,5 \text{ mg/kg}$ Microbiological criteria:

Yeast and mould count: ≤ 100 CFU/g Aerobic bacteria count: ≤ 10 000 CFU/g

Total coliform count: ≤ 10 CFU/g Escherichia coli: Absence in 10 g Salmonella: Absence in 25 g

[F32Refined shrimp peptide concentrate

Description

Refined shrimp peptide concentrate is a peptide mixture obtained from northern shrimp (*Pandalus borealis*) shells and heads via a series of purification steps following enzymatic proteolysis using a protease from *Bacillus licheniformis* and/or *Bacillus amyloliquefaciens*.

Characteristics/Composition

Total Dry matter (%): \geq 95,0 %

Peptides (w/weight dry matter): ≥ 87.0 % of which peptides with

molecular weight $< 2 \text{ kDa} : \ge 99.9 \%$

Fat (w/w): $\leq 1.0 \%$

Carbohydrates (w/w): $\leq 1.0 \%$

Ash (w/w): $\leq 15,0 \%$ Calcium: $\leq 2,0 \%$ Potassium: $\leq 0,15 \%$ Sodium: $\leq 3,5 \%$

Heavy Metals

Arsenic (inorganic): $\leq 0,22$ mg/kg Arsenic (organic): $\leq 51,0$ mg/kg

Cadmium: $\leq 0.09 \text{ mg/kg}$ Lead: $\leq 0.18 \text{ mg/kg}$

Total mercury: ≤ 0,03 mg/kg

Microbiological criteria:

Total viable cell count: ≤ 20 000 CFU/g

Salmonella: ND/25g

Listeria monocytogenes : ND/25g *Escherichia coli* : ≤ 20 CFU/g

Coagulase positive *Staphylococcus aureus*: ≤ 200 CFU/g

Pseudomonas aeruginosa : ND/25g

Mould/yeast: $\leq 20 \text{ CFU/g}$

CFU : Colony Forming Units ND : Not Detectable

Transresveratrol

Description/Definition:

Synthetic *Trans* -resveratrol is off-white to beige crystals.

Chemical name: 5-[(E)-2-(4-hydroxyphenyl)ethenyl]benzene-1,3-diol

Chemical formula: C₁₄ H₁₂ O₃ Molecular weight: 228,25 Da

CAS No: 501-36-0

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Purity:

Trans -resveratrol: ≥ 98 %-99 %

Total by-products (related substances): $\leq 0.5 \%$

Any single related substance: $\leq 0.1 \%$

Sulphated ash: $\leq 0.1 \%$ Loss on drying: $\leq 0.5 \%$

Heavy metals: Lead: ≤ 1,0 ppm Mercury: ≤ 0,1 ppm Arsenic: ≤ 1,0 ppm

Impurities:

Diisopropylamine: ≤ 50 mg/kg

Microbial source: A genetically modified strain of Saccharomyces

cerevisiae

Appearance: Off-white to slight yellow powder

Particle size: 100 % less than 62,23 µm

Trans-resveratrol content: Min. 98 % w/w (dry weight basis)

Ash: Max. 0,5 % w/w Moisture: Max. 3 % w/w

Rooster comb extract

Description/Definition:

Rooster comb extract is obtained from *Gallus gallus* by enzymatic hydrolysis of rooster comb and by subsequent filtration, concentration and precipitation steps. The principal constituents of rooster comb extract are the glycosaminoglycans hyaluronic acid, chondroitin sulphate A and dermatan sulphate (chondroitin sulphate B). White or almost white hygroscopic powder.

Hyaluronic acid: 60-80 %

Chondroitin sulphate A: ≤ 5.0 %Dermatan sulphate (chondroitin sulphate

B): ≤ 25 % pH: 5,0-8,5 **Purity:**

Chlorides: $\leq 1.0 \%$ Nitrogen: $\leq 8.0 \%$

Loss on drying: (105 °C for 6 hours): \leq 10 %

Heavy metals: Mercury: ≤ 0,1 mg/kg Arsenic: ≤ 1,0 mg/kg Cadmium: ≤ 1,0 mg/kg Chromium: ≤ 10 mg/kg Lead: ≤ 0,5 mg/kg Microbiological criteria:

Total viable aerobic count: $\leq 10^2$ CFU/g

Escherichia coli : Absence in 1 g Salmonella : Absence in 1 g

Staphylococcus aureus : Absence in 1 g Pseudomonas aeruginosa : Absence in 1g

Sacha Inchi oil from Plukenetia volubilis

Description/Definition:

Sacha inchi oil is a 100 % cold pressed vegetable oil obtained from the seeds of *Plukenetia volubiis* L. It is a transparent, fluid (liquid) and shiny oil at room temperature. It has a fruity, light, green vegetable taste without undesirable flavours.

Aspect, limpidity, shine, colour: Fluid at room temperature, clean, shiny yellow gold

Odour and taste: Fruity, vegetable without non acceptable taste or odour

Purity:

Water and Volatiles: < 0,2 g/100 g

Impurities insoluble in hexane: < 0,05 g/100 g

Oleic acidity: < 2.0 g/100 g

Peroxide value (PV): $< 15 \text{ meq O}_2/\text{kg}$

Trans fatty acids: < 1.0 g/100 g

Total unsaturated fatty acids: > 90 %Omega 3 alpha linolenic acid

(ALA): > 45 %

Saturated fatty acids: < 10 % No trans fatty acids (< 0,5 %) No erucic acid (< 0,2 %)

More than 50 % of tri-linolenin and di-linolenin-triglycerides

Phytosterols composition and level No cholesterol (< 5,0 mg/100 g)

Salatrims

Description/Definition:

Salatrim is the internationally recognised acronym for (short and long chain acyl triglyceride molecules). Salatrim is prepared by non-enzymatic inter-esterification of triacetin, tripropionin, tributyrin, or their mixtures with hydrogenated canola, soybean, cottonseed, or sunflower oil.

Description: Clear slightly amber liquid to a light coloured ways solid.

Description: Clear, slightly amber liquid to a light coloured waxy solid at room temperature. Free of particulate matter and of foreign or rancid odour.

Glycerol ester disribution: Triacylglycerols: > 87 %

Diacylglycerols: ≤ 10 % Monoacylglycerols: ≤ 2,0 %

Fatty acid composition:

MOLE % LCFA (long chain fatty acids): 33-70 % MOLE % SCFA (short chain fatty acids): 30-67 % Saturated long chain fatty acids: < 70 % by weight

Trans fatty acids: $\leq 1.0 \%$

Free fatty acids as oleic acid: $\leq 0.5 \%$

Triacylglycerol profile:

Triesters (short/long of 0,5 to 2,0): \geq 90 %

Triesters (short/long = 0): \leq 10 % Unsaponifiable material: \leq 1,0 %

Moisture: $\leq 0.3 \%$ Ash: $\leq 0.1 \%$

Colour: $\leq 3.5 \text{ Red (Lovibond)}$ Peroxide value (PV): $\leq 2.0 \text{ Meq/Kg}$

Schizochytrium sp. oil rich in DHA and EPA

Acid value: $\leq 0.5 \text{ mg KOH/g}$

Peroxide value (PV): ≤ 5.0 meg/kg oil

Oxidative stability: All food products containing *Schizochytrium sp.* oil rich in DHA and EPA should demonstrate oxidative stability by appropriate and recognised national/international test methodology (e.g.

AOAC)

Moisture and volatiles: $\leq 0.05 \%$

Unsaponifiables: $\leq 4.5 \%$ Trans-fatty acids: $\leq 1 \%$

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	DHA content: ≥ 22,5 % EPA content: ≥ 10 %
[F33Schizochytrium sp. (ATCC PTA-9695) oil	The novel food is obtained from the strain ATCC PTA-9695 of the microalgae <i>Schizochytrium</i> sp. Peroxide value (PV): ≤ 5.0 meq/kg oil Unsaponifiables: ≤ 3.5 % Trans-fatty acids: ≤ 2.0 % Free fatty acids: ≤ 0.4 % Docosapentaenoic acid (DPA) n-6: ≤ 7.5 % DHA content: ≥ 35 %]
f ^{F67} Schizochytrium sp. strain (FCC-3204) oil	Description/Definition: The novel food is an oil produced from the strain FCC-3204 of the microalgae Schizochytrium sp. Composition: Acid value: ≤ 0.5 mg KOH (potassium hydroxide)/g Peroxide value (PV): ≤ 5.0 meq (milliequivalent)/kg oil Moisture and volatiles: ≤ 0.05 % Unsaponifiables: ≤ 4.5 % Trans-fatty acids: ≤ 1.0 % Docosahexaenoic acid (DHA): ≥ 32.0 % P-anisidine value: ≤ 10]
Schizochytrium sp. oil	Acid value: ≤ 0.5 mg KOH/g Peroxide value (PV): ≤ 5.0 meq/kg oil Moisture and volatiles: ≤ 0.05 % Unsaponifiables: ≤ 4.5 % Trans-fatty acids: ≤ 1.0 % DHA content: ≥ 32.0 %
[F68Schizochytrium sp. (T18) oil	Acid value: ≤ 0.8 mg KOH/g Peroxide value (PV): ≤ 5.0 meq/kg oil Moisture and volatiles: ≤ 0.05 % Unsaponifiables: ≤ 3.5 % Trans-fatty acids: ≤ 2.0 % Free fatty acids: ≤ 0.4 % DHA content: ≥ 35 %]
[F69Schizochytrium sp. (WZU477) oil	Description/Definition: The novel food is an oil produced from the strain WZU477 of the microalgae Schizochytrium sp. Composition: Acid value: ≤ 0.5 mg KOH (potassium hydroxide)/g Peroxide value (PV): ≤ 5.0 meq (milliequivalent)/kg oil Moisture and volatiles: ≤ 0.05 % Unsaponifiables: ≤ 4.5 % Trans-fatty acids: ≤ 1.0 % Docosahexaenoic acid (DHA): ≥ 32.0 % P-anisidine value: ≤ 10]
[F41Syrup from Sorghum bicolor (L.) Moench.	Description/Definition The traditional food is syrup from <i>Sorghum bicolor</i> (L.) Moench (genus, <i>Sorghum</i> ; family, <i>Poaceae</i> (alt. <i>Gramineae</i>)).

from a third country)

(Traditional food | The syrup is obtained from stalks of S. bicolor, after applying production processes such as crushing, extraction, and evaporation including a heat treatment in order to obtain a minimum of 74 °Brix syrup

Compositional data of syrup from *Sorghum bicolor* (L.) Moench

Water: 22,7 g/100 g

Ash: 2,4

Sugars, total: > 74.0 g/100 g

Fermented sovbean extract

Description/Definition:

Fermented soybean extract is an odourless milk-white coloured powder. It is comprised of 30 % fermented soybean extract powder and 70 % resistant dextrin (as carrier) from corn-starch, which is added during the processing. Vitamin K₂ is removed during the manufacturing process. Fermented soybean extract contains nattokinase isolated from natto, a foodstuff produced by the fermentation of non-genetically modified soybeans (Glycine max (L.)) with a selected strain of Bacillus subtilis var. natto.

Nattokinase activity: 20 000 -28 000 Fibrin degradation unit/g(1)

Identity: Confirmable

Condition: No offensive taste or smell

Loss on drying: $\leq 10 \%$ Vitamin K₂: ≤ 0.1 mg/kg

Heavy metals:

Lead: $\leq 5.0 \text{ mg/kg}$ Arsenic: $\leq 3.0 \text{ mg/kg}$ Microbiological criteria:

Total viable aerobic count: $\leq 10^3$ CFU(3)/g

Yeast and mould: $\leq 10^2$ CFU/g

Coliforms: ≤ 30 CFU/g

Spore-forming bacteria: ≤ 10 CFU/g Escherichia coli: Absence/25 g Salmonella: Absence/25 g Listeria: Absence/25 g

 $(^1)$ Assay method as described by Takaoka et al. (2010).

[F70Spermidinerich wheat germ extract (Triticum aestivum)

Description/Definition:

Spermidine-rich wheat germ extract is obtained from non-fermented, non-sprouting wheat germs (Triticum aestivum) by the process of solidliquid extraction targeting specifically, but not exclusively polyamines. Spermidine:(N-(3-aminopropyl)butane-1,4-diamine):0,8-2,4 mg/g

Spermine: 0,4-1,2 mg/g

Spermidine trichloride $< 0.1 \mu g/g$ Putrescine: < 0.3 mg/g

Cadaverine: $\leq 16.0 \, \mu g/g$

Mycotoxins:

Aflatoxins (total): $< 0.4 \mu g/kg$ Microbiological criteria:

Total aerobic bacteria: < 10 000 CFU/g Yeast and moulds: < 100 CFU/g *Escherichia coli:* < 10 CFU/g Salmonella: Absence/25g

Listeria monocytogenes: Absence/25g]

ANNEX

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Sucromalt

Description/Definition:

Sucromalt is a complex mixture of saccharides which is produced from sucrose and a starch hydrolysate by means of an enzymatic reaction. In this process, glucose units are attached to saccharides from the starch hydrolysate by means of an enzyme produced by the bacterium Leuconostoc citreum or by means of a recombinant strain of the production organism *Bacillus licheniformis*. The resulting oligosaccharides are characterised by the presence of α -(1 \rightarrow 6) and α - $(1\rightarrow 3)$ glycosidic compounds. The overall product is syrup, in addition to these oligosaccharides, contains mainly fructose but also the disaccharide leucrose and other disaccharides.

Total solids: 75-80 % Moisture: 20-25 % Sulphatase: Max 0.05 %

pH: 3,5-6,0

Conductivity < 200 (30 %)

Nitrogen < 10 ppm Fructose: 35-45 % d.w. Leucrose: 7-15 % d.w.

Other disaccharides: Max 3 % Higher saccharides: 40-60 % d.w

Sugar cane fibre

Description/Definition:

Sugar Cane Fibre is derived from the dry cell wall or fibrous residue remaining after expression or extraction of sugar juice from sugar cane, of the Saccharum genotype. It consists primarily of cellulose and hemicellulose.

The production process consists of several steps, including: chipping, alkaline digestion, removal of lignins and other noncellulosic components, bleaching of purified fibres, acid washing and neutralization. Moisture: $\leq 7.0 \%$

Ash: $\leq 0.3 \%$

Total Dietary Fibre (AOAC) dry basis (all insoluble): $\geq 95 \%$ of which: Hemicellulose (20-25 %) and cellulose (70-75 %)

Silica (ppm): ≤ 200 Protein: 0,0 % Fat: Trace pH: 4-7

Heavy metals:

Mercury (ppm): ≤ 0.1 Lead (ppm): ≤ 1.0 Arsenic (ppm): ≤ 1.0 Cadmium (ppm): ≤ 0.1 Microbiological criteria:

Yeast and moulds (CFU/g): $\leq 1~000$

Salmonella: Absence

Listeria monocytogenes: Absence

IF42Sugars obtained from cocoa (Theobroma cacao L.) pulp

Description/Definition:

Sugars are obtained from the concentrated cocoa pulp (*Theobroma* cacao L.) juice either via a drying process or via a purification process to produce high purity glucose or fructose.

Sugars produced by a drying process

Nutritional composition:

Total sugars (g/100g): > 80

Moisture (%): < 5

Microbiological criteria:

Total Plate Count (aerobic) (cfu/g): < 10⁴

Moulds and Yeasts (cfu/g): < 50

Enterobacteriaceae (cfu/g): < 10

Salmonella spp.: Absence in 25 g

Alicyclobacillus: Absence in 50 g

Thermo-acidophilic bacteria: Absence in 50 g Sugars produced by a purification process

Nutritional composition of Glucose obtained from cocoa (*Theobroma*

cacao L.) pulp:

Glucose content (%): > 93

Ash (%): < 0.2

Moisture (%): < 1.0

Nutritional composition of Fructose obtained from cocoa (Theobroma

cacao L.) pulp:

Fructose content (%): > 98

Glucose content (%): < 0.5 %

Ash (%): < 0.2

Moisture (%):< 0.5

Microbiological criteria for glucose and fructose obtained from cocoa (

Theobroma cacao L.) pulp:

Total Plate Count (aerobic) (cfu/g): $< 10^4$

Salmonella spp.: Absence in 25 g]

Sunflower oil extract

Description/Definition:

The sunflower extract is obtained by a concentration factor of 10 of the unsaponifiable fraction of refined sunflower oil extracted from the seeds of the sunflower, *Helianthus Annuus* L.

Composition:

Oleic acid (C18:1): 20 % Linoleic acid (C18:2): 70 % Unsaponifiable matter: 8,0 %

Phytosterols: 5,5 %

Tocopherols: 1,1 %

Dried Tetraselmis chuii microalgae

Description/Definition:

The dried product is obtained from the marine microalgae *Tetraselmis chuii*, belonging to the *Chlorodendraceae* family, cultivated in sterile sea water in closed photobioreactors insulated from the outside air.

Purity/Composition:

Identified by means of nuclear marker rDNA 18 S (sequence analysed no less than 1 600 base pairs) in the National Centre for Biotechnology information (NCBI) database: Not less than 99,9 %

Humidity: ≤ 7,0 % Proteins: 35-40 % Ashes: 14-16 %

Carbohydrates: 30-32 %

Fibre: 2-3 % Fat: 5-8 %

Saturated fatty acids: 29-31 % of total fatty acids

	Monounsaturated fatty acids: 21-24 % of total fatty acids Polyunsaturated fatty acids: 44-49 % of total fatty acids Iodine: ≤ 15 mg/kg
Therapon barcoo / Scortum	Description/Definition: Scortum/ Therapon barcoo is a species of fish in the family Terapontidae. It is an endemic fresh water species from Australia. It is now reared in fish farms. Taxonomic Identification: Class: Actinopterygii > order: Perciformes > family: Terapontidae > genus: Therapon or Scortum barcoo Composition of fish flesh: Protein (%): 18-25 Moisture (%): 65-75 Ash (%): 0,5-2,0 Energy (KJ/Kg): 6000-11500 Carbohydrates (%): 0,0 Fat (%): 5-15 Fatty acids (mg FA/g fillet): Σ PUFA n-3: 1,2-20,0 Σ PUFA n-6: 0,3-2,0 PUFA n-6: 0,3-2,0 Total omega 3 acids: 1,6-40,0 Total omega 6 acids: 2,6-10,0
D-Tagatose	Description/Definition: Tagatose is produced by isomerization of galactose by means of chemical or enzymatic conversion, or by epimerization of fructose by means of enzymatic conversion. These are single-step conversions. Appearance: White or almost white crystals Chemical name: D-tagatose Synonym: D- <i>lyxo</i> -Hexulose CAS number: 87-81-0 Chemical formula: C $_6$ H $_{12}$ O $_6$ Formula weight: 180,16 (g/mol) Purity: Assay: ≥ 98 % on a dry weight basis Loss on drying: ≤ 0,5 % (102 °C, 2 hours) Specific Rotation: [α] $_D$ 20 : − 4 to − 5,6 ° (1 % aqueous solution)(1) Melting range: 133−137 °C Heavy metals: Lead: ≤ 1,0 mg/kg(*)
	(*) Determine using an atomic absorption technique appropriate to the specified level. The selection of sample size and method of sample preparation may be based on the principles of the method described in FNP 5. 'Instrumental methods'(1).
	Food and nutrition paper 5 Rev 2 – Guide to specifications for general notices, general analytical techniques, identification tests, test solutions and other reference materials (JECFA) 1991, 307 p.; English – ISBN 92-5-102991-1
[^{F18} Taxifolin- rich extract]	Description:

Taxifolin-rich extract from the wood of Dahurian Larch (*Larix gmelinii* (Rupr.) Rupr) is a white to pale-yellow powder that crystallizes from hot aqueous solutions.

[F18 Definition:

Chemical name: [(2R,3R)-2-(3,4 dihydroxyphenyl)-3,5,7-trihydroxy-2,3-dihydrochromen-4-one, also called (+) trans (2R,3R)- dihydroquercetin] and with no more than 2 % of the cis-form

Specifications:

Physical parameter

Moisture: $\leq 10 \%$ *Compound analysis* Taxifolin (m/m): $\geq 90,0 \%$ of the dry weight

Heavy Metals, Pesticide

Lead: ≤ 0.5 mg/kg Arsenic: ≤ 0.02 mg/kg Cadmium: ≤ 0.5 mg/kg Mercury: ≤ 0.1 mg/kg

Dichlorodiphenyltrichloroethane (DDT): ≤ 0.05 mg/kg

Residual solvents Ethanol: < 5 000 mg/kg **Microbiological criteria**

Total Plate Count (TPC): $\leq 10^4$ CFU/g

Enterobacteria: ≤ 100/g

Yeast and Mould: ≤ 100 CFU/g Escherichia coli : Absence/1 g Salmonella : Absence/10 g

Staphylococcus aureus: Absence/1 g

Pseudomonas: Absence/1g

Usual range of components of the Taxifolin-rich extract (as per dry substance)

Extract component	Content, usual observed range (%)
Taxifolin	90 – 93
Aromadendrin	2,5 – 3,5
Eriodictyol	0.1 - 0.3
Quercetin	0,3 – 0,5
Naringenin	0.2 - 0.3
Kaempferol	0,01 – 0,1
Pinocembrin	0,05 - 0,12
Unidentified flavonoids	1 – 3
Water(*)	1,5

(*) Taxifolin in its hydrated form and during the drying process is a crystal. This results on the inclusion of water of crystallisation in a quantity of 1,5 %.

Trehalose

Description/Definition:

A non-reducing disaccharide that consists of two glucose moieties linkes by an α -1,1-glucosidic bond. It is obtained from liquefied starch or from sucrose by a multistep enzymatic process. The commercial product is the dihydrate. Virtually odourless, white or almost white crystals with a sweet taste

Synonyms: α,α -trehalose

Chemical name: α-D-glucopyranosyl-α-D-glucopyranoside, dihydrate

CAS No.: 6138-23-4 (dihydrate)

Chemical formula: C₁₂H₂₂O₁₁ · 2H₂O (dihydrate)

Formula weight: 378,33 (dihydrate) Assay: ≥ 98 % on the dry basis

Determine using an atomic absorption technique appropriate to the specified level. The selection of sample size and method of sample preparation may be based on the principles of the method described in FNP 5 (1), 'Instrumental methods'

Method of assay:

Principle: trehalose is identified by liquid chromatography and quantified by comparison to a reference standard containing standard trehalose Preparation of sample solution: weigh accurately about 3 g of dry sample into a 100 ml volumetric flask and add about 80 ml of purified, deionised water. Bring sample to complete dissolution and dilute to mark with purified deionised water. Filter through a 0,45 micron filter Preparation of standard solution: dissolve accurately weighed quantities of dry standard reference trehalose in water to obtain a solution having known concentration of about 30 mg of trehalose per ml.

Apparatus: liquid chromatography equipped with a refractive index detector and integrating recorder

Conditions:

Column: Shodex Ionpack KS-801 (Showa Denko Co.) or equivalent

length: 300 mmdiameter: 10 mmtemperature: 50 °C

Mobile phase: water flow rate: 0,4 ml/min Injection volume: 8 µl

Procedure: inject separately equal volumes of the sample solution and the standard solution into the chromatograph.

Record the chromatograms and measure the size of response of the trehalose peak

Calculate the quantity, in mg, of trehalose in 1 ml of the sample solution by the following formula:% trehalose = $100 \times (R_U/R_S) (W_S/W_U)$ where

R S = peak area of trehalose in the standard preparation
R U = peak area of trehalose in the sample preparation
W S = weight in mg of trehalose in the standard preparation
W S = weight of dry sample in mg

 W_{U} = weight of dry sample in mg

Characteristics:

Identification:

Solubility: Freely soluble in water, very slightly soluble in ethanol Specific rotation: $[\alpha]_D^{20} = +179^{\circ}$ (5 % aqueous solution, dihydrate), +199° (5 % aqueous solution, anhydrous substance)

Melting point: 97 °C (dihydrate)

Purity:

Loss on drying: $\leq 1.5 \%$ (60 °C, 5h)

Total ash: $\leq 0.05 \%$ **Heavy metals:** Lead: $\leq 1.0 \text{ mg/kg}$

[F18UV-treated mushrooms (Agaricus bisporus)

Description/Definition

Commercially grown *Agaricus bisporus* to which UV light treatment is applied to harvested mushrooms.

UV radiation: a process of radiation in ultraviolet light within the wavelength of 200-800 nm.

Vitamin D 2

Chemical name: (3\beta,5\,Z,7\,E,22\,E)-9,10-secoergosta-5,7,10(19),22-

tetraen-3-ol

Synonym: Ergocalciferol

CAS No: 50-14-6

Molecular weight: 396,65 g/mol

Contents

Vitamin D₂ in the final product: 5-20 μg/100 g fresh weight at the

expiration of shelf life.]

[F71UV-treated baker's yeast (Saccharomyces cerevisiae)

Description/Definition

Baker's yeast (Saccharomyces cerevisiae) is treated with ultraviolet light to induce the conversion of ergosterol to vitamin D# (ergocalciferol). Vitamin D# content in the yeast concentrate varies between 800,000 - 3,500,000 IU vitamin D/100g ($200-875 \mu g/g$). The yeast is inactivated for use in infant formula, follow-on formula, processed cereal-based food, and food for special medical purposes as defined by Regulation (EU) No. 609/2013. The yeast can be active or inactive for use in other foods. The yeast concentrate is blended with regular baker's yeast in order not to exceed the maximum level in the pre-packed fresh or dry yeast for home baking.

Tan-coloured, free-flowing granules.

Vitamin D#

Chemical name: (5Z,7E,22E)-(3S)-9,10-secoergosta-5,7,10(19),22-

tetraen-3-ol

Synonym: Ergocalciferol CAS No.: 50-14-6

Molecular weight: 396.65 g/mol

Microbiological criteria for the yeast concentrate

Coliforms: $\leq 10^3$ CFU/g Escherichia coli: ≤ 10 CFU/g Salmonella spp: Absence in 25 g CFU: Colony Forming Units.]

UV-treated bread

Description/Definition:

UV-treated bread is yeast leavened bread and rolls (without toppings) to which a treatment with ultraviolet radiation is applied after baking in order to convert ergosterol to vitamin D₂ (ergocalciferol).

UV radiation: A process of radiation in ultraviolet light within the wavelength of 240-315 nm for maximum of 5 seconds with energy input of 10-50 mJ/cm².

Vitamin D $_2$:

Chemical name: (5Z,7E,22E)-3S-9,10-secoergosta-5,7,10(19),22-

tetraen-3-ol

Synonym: Ergocalciferol

CAS No: 50-14-6

Molecular weight: 396,65 g/mol

Contents:

Vitamin D₂ (ergocalciferol) in the final product: $0.75-3 \mu g/100 g(^{1})$

Yeast in dough: $1-5 \text{ g}/100 \text{ g} (^2)$

(1) EN 12821, 2009, European Standard.

(2) Recipe calculation.

UV-treated milk

Description/Definition:

UV-treated milk is cow's milk (whole and semi-skimmed) to which a treatment with ultraviolet (UV) radiation via turbulent flow is applied after pasteurisation. The treatment of the pasteurised milk with UV radiation results in an increase in the vitamin D $_3$ (cholecalciferol) concentrations by conversion of 7-dehydrocholesterol to vitamin D $_3$. UV radiation: A process of radiation in ultraviolet light within the wavelength of 200-310 nm with energy input of 1 045 J/l.

Vitamin D 3:

 $\label{lem:chemical name: continuous} Chemical name: (1S,3Z)-3-[(2E)-2-[(1R,3aS,7aR)-7a-methyl-1-[(2R)-6-methylheptan-2-yl]-2,3,3a,5,6,7-hexahydro-1H-inden-4-ylidene]-4-methylidenecyclohexan-1-ol$

Synonym: Cholecalciferol

CAS No: 67-97-0

Molecular weight: 384,6377 g/mol

Contents:

Vitamin D $_3$ in the final product:

Whole milk(1)0,5-3,2 µg/100 g(2)

Semi-skimmed milk(1): $0.1-1.5 \mu g/100 g(^2)$

- (1) As defined by Regulation (EU) No 1308/2013 of the European Parliament and of the Council of 17 December 2013 establishing a common organisation of the markets in agricultural products and repealing Council Regulations (EEC) No 922/72, (EEC) No 234/79, (EC) No 1037/2001 and (EC) No 1234/2007 (OJ L 347, 20.12.2013, p. 671).
- $(^2)$ HPLC

[F7Vitamin D 2 mushroom powder

Description/Definition

Vitamin D₂ mushroom powder is a granular powder made from homogenised *Agaricus bisporus* mushrooms that have been exposed to UV light.

The mushrooms are washed, homogenised and suspended in water to produce a mushroom slurry. The mushroom slurry is passed under a UV lamp. The slurry is then filtered, dried and ground, producing vitamin D $_{\rm 2}$ mushroom powder.

UV radiation: A process of radiation in ultraviolet light within a range of wavelength similar to those UV-treated novel foods authorised under the novel food regulation.

Characteristics/Composition

Vitamin D₂ content: 1 000–1 300 μg/g of mushroom powder¹

Moisture: ≤ 10,0 % Ash: ≤ 13,5 % **Heavy Metals**

Lead (as Pb): ≤ 0.5 mg/kg Cadmium: ≤ 0.5 mg/kg Mercury: ≤ 0.1 mg/kg Arsenic: ≤ 0.3 mg/kg

Mycotoxins

Aflatoxins (sum of B1+B2+G1+G2): $< 4 \mu g/kg$

Microbiological criteria:

Total plate count: ≤ 5 000 CFU^g/g Yeast and mould: ≤ 100 CFU/g Salmonella sp.: Absent in 25 g Staphylococcus aureus : ≤ 10 CFU/g Escherichia coli : ≤ 10 CFU/g

Coliforms: $\leq 10 \text{ CFU/g}$ *Enterobacteriaceae* : $\leq 10 \text{ CFU/g}$

Listeria monocytogenes: Absent in 25 g]

[^{F72}Vitamin D# mushroom powder

Description/Definition

The novel food is mushroom powder produced from dried whole *Agaricus bisporus* mushrooms. The process includes drying, milling and the controlled exposure of the mushroom powder to ultraviolet light.

Characteristics/Composition

Vitamin D# content: 580-595 μg/g of mushroom powder

Ash: $\leq 13.5\%$ Water activity: < 0.5Moisture content: $\leq 7.5\%$ Carbohydrates: $\leq 35\%$ Total dietary fibre: $\geq 15\%$

Crude protein (N x 6.25): \geq 22%

Fat: ≤ 4.5% **Heavy metals**

Lead: ≤ 0.5 mg/kg Cadmium: ≤ 0.5 mg/kg Mercury: ≤ 0.1 mg/kg Arsenic: ≤ 0.3 mg/kg

Mycotoxins

Aflatoxin B1: $\leq 0.1 \,\mu\text{g/kg}$

Aflatoxins (sum of B1 + B2 + G1 + G2): $< 4 \mu g/kg$

Microbiological criteria

Total plate count: ≤ 5000 CFU 14

Total yeast and mould count: ≤ 100 CFU/g

Escherichia coli: < 10 CFU/g
Salmonella spp.: Absence in 25 g
Staphylococcus aureus: ≤ 10 CFU/g

Coliforms: ≤ 10 CFU/g Listeria spp.: Absence in 25 g

Enterobacteriaceae: < 10 CFU/g CFU: Colony Forming Units.]

Vitamin K₂ (menaquinone)

This novel food is produced by a synthetic or microbiological process. Vitamin K ₂ (2-methyl-3-all-trans-polyprenyl-1,4-naphthoquinones), or the menaquinone series, is a group of prenylated naphthoquinone derivatives. The number of isoprene residues, where 1 isoprene unit consists of 5 carbons comprising the side chain, is used to characterise the menaquinone homologues containing primarily MK-7 and to a smaller extent MK-6.

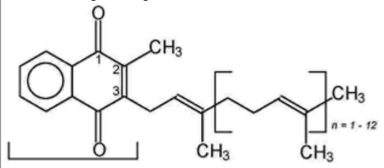
Vitamin K $_2$ (menaquinones) series with menaquinone-7 (MK-7)(n = 6) being C $_{46}$ H $_{64}$ O $_2$, menaquinone-6 (MK-6)(n = 5) being C $_{41}$ H $_{56}$ O $_2$ and menaquinone-4 (MK-4)(n = 3) being C $_{31}$ H $_{40}$ O $_2$.

Chemical Name: (all-E)-2-(3,7,11,15,19,23,27-

Heptamethyl-2, 6, 10, 14, 18, 22, 26-octacosa heptaenyl)-3-methyl-1, 4-methyl-1, 4-methy

naphtalenedione

CAS Number: 2124-57-4 Molecular formula: C₄₆ H₆₄ O₂ Molecular weight: 649 g/mol



2-methyl-1,4-naphthoquinone (menadione moiety)

Specification of synthetic Vitamin K_2 (menaquinone-7)

Appearance: Yellow powder

Purity: Max 6,0 % cis-isomer, max 2,0 % other impurities

Content: 97-102 % Menaquinone-7 (including at least 92 % all-trans

Menaquinone-7)

Specifications of microbiologically produced Vitamin K ₂ (menaquinone-7)

Source: Bacillus subtilis spp. natto and Bacillus licheniformis

Appearance: Yellow powder or oil suspension

Wheat bran extract

Description/Definition:

White crystalline powder obtained by enzymatic extraction from *Triticum aestivum* L. bran, rich in arabinoxylan oligosaccharides

Dry matter: Min. 94 %

Arabinoxylan oligosaccharides: Min 70 % of dry matter

Average degree of polymerisation of arabinoxylan oligosaccharides: 3-8 Ferulic acid (bound to arabinoxylan oligosaccharides): 1-3 % of dry

matter

Total poly/oligosaccharides: Min 90 %

Protein: Max 2 % of dry matter

Ash: Max 2 % of dry matter **Microbiological parameters:**

Mesophilic bacteria – total count: Max 10 000/g

Yeasts: Max 100/g Fungi: Max 100/g

Salmonella: Absence in 25g Bacillus cereus: Max 1000/g

Clostridium perfringens: Max 1000/g

[F73Xylooligosaccharides

Description:

The novel food is a mixture of xylo-oligosaccharides (XOS) which are obtained from corncobs (*Zea mays* subsp. *mays*) via hydrolysis by a xylanase from *Trichoderma reesei* followed by a purification process.

Characteristics/Composition

Parameter	Powder form 1	Powder form 2	Syrup form
Moisture (%)	≤ 5,0	≤ 5,0	70-75
Protein (g/100 g)	< 0,2		
Ash (%)	≤ 0,3		
рН	3,5-5,0		
Total carbohydrate content (g/100 g)	≥ 97	≥ 95	≥ 70
XOS content (dry basis) (g/100 g)	≥ 95	≥ 70	≥ 70
Other carbohydrates (g/100 g) (a)	2,5-7,5	2-16	1,5-31,5
Monosaccharides total (g/100 g)	0-4,5	0-13	0-29
Glucose (g/100 g)	0-2	0-5	0-4
Arabinose (g/100 g)	0-1,5	0-3	0-10
Xylose (g/100 g)	0-1,0	0-5	0-15
Disaccharides total (g/100 g)	27,5-48	25-43	26,5-42,5
Xylobiose (XOS DP2) (g/100 g)	25-45	23-40	25-40
Cellobiose (g/100 g)	2,5-3	2-3	1,5-2,5

Oligosaccharides total (g/100 g)	41-77	36-72	32-71
xylotriose (XOS DP3) (g/100 g)	27-35	18-30	18-30
xylotetraose (XOS DP4) (g/100 g)	10-20	10-20	8-20
xylopentaose (XOS DP5) (g/100 g)	3-10	5-10	3-10
xylohexaose (XOS DP6) (g/100 g)	1-5	1-5	1-5
Xyloheptaose (XOS DP7) (g/100 g)	0-7	2-7	2-6
Maltodextrin (g/100 g) (b)	0	20-25	0
Copper (mg/kg)	< 5,0		
Lead (mg/kg)	< 0,5		
Arsenic (mg/kg)	< 0,3		
Salmonella (CFU (°)/25 g)	Negative		
E, coli (MPN (Negative		
Yeast (CFU/g)	< 10		
Mould (CFU/g)	< 10		
DP :	Degree of poly	merization	

DP : Degree of polymerization

- (a) Other carbohydrates include monosaccharides (glucose, xylose and arabinose) and cellobiose.
- (b) Maltodextrin content is calculated according to the amount added in the process.
- (°) CFU: Colony Forming Units.
- (d) MPN: Most Probable Number.]

I^{F50}Yarrowia lipolytica yeast biomass

Description/Definition:

The novel food is the dried and heat-killed biomass of the yeast *Yarrowia lipolytica* .

Characteristics/Composition:

Protein: 45-55 g/100 g Dietary fibre: 24-30 g/100 g Sugars: < 1,0 g/100 g

Fat: 7-10 g/100 g Total ash: \leq 12 % Water content: \leq 5 % Dry matter content: \geq 95 % **Microbiological criteria:**

Total Aerobic Microbial Count: $\leq 5 \times 10^3$ CFU/g Total Yeast and Mould Count: $\leq 10^2$ CFU/g

Viable Yarrowia lipolytica cells^j: < 10 CFU/g (i.e. limit of detection)

Coliforms: $\leq 10 \text{ CFU/g}$

Salmonella spp.: Absence in 25 g]

[^{F74}3'-Sialyllactose (3'-SL) sodium salt (microbial source)

Description:

3'-Sialyllactose (3'-SL) sodium salt is a purified, white to off-white powder or agglomerate that is produced by a microbial process and contains limited levels of lactose, 3'-sialyl-lactulose, and sialic acid **Source:**

Genetically modified strain of Escherichia coli K-12 DH1

Definition:

Chemical formula: C₂₃H₃₈NO₁₉Na

Chemical name: N-Acetyl- α -D-neuraminyl- $(2\rightarrow 3)$ - β -D-galactopyranosyl-

(1→4)-D-glucose, sodium salt Molecular mass: 655.53 Da CAS No 128596-80-5

Characteristics/Composition:

Appearance: White to off-white powder or agglomerate

Sum of 3'-Sialyllactose sodium salt, D-Lactose, and Sialic acid (% of dry

matter): $\geq 90.0\%$ (w/w)

3'-Sialyllactose sodium salt (% of dry matter): \geq 88.0 % (w/w)

D-Lactose: $\leq 5.0 \%$ (w/w) Sialic acid: $\leq 1.5 \%$ (w/w)

3'-Sialyl-lactulose: $\leq 5.0 \%$ (w/w)

Sum of other carbohydrates: $\leq 3.0 \%$ (w/w)

Moisture: $\leq 8.0 \%$ (w/w) Sodium: 2.5 - 4.5 % (w/w) Chloride: $\leq 1.0 \%$ (w/w)

pH (20 °C, 5 % solution): 4.5 -6.0 Residual protein: \leq 0.01 % (w/w)

Microbiological criteria:

Aerobic mesophilic bacteria total plate count: ≤ 1000 CFU/g

Enterobacteriaceae: ≤ 10 CFU/g Salmonella sp.: Absence in 25 g

Yeast: $\leq 100 \text{ CFU/g}$ Mould: $\leq 100 \text{ CFU/g}$

Residual endotoxins: ≤ 10 EU/mg

CFU: Colony Forming Units; EU: Endotoxin Units]

[^{F75}6'-Sialyllactose (6'-SL) sodium salt (microbial source)

Description:

6'-Sialyllactose (6'-SL) sodium salt is a purified, white to off-white powder or agglomerate that is produced by a microbial process and contains limited levels of lactose, 6'-sialyl-lactulose, and sialic acid

Source

Genetically modified strain of Escherichia coli K-12 DH1

Definition:

Document Generated: 2024-04-16

Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

Chemical formula: C23H38NO19Na

Chemical name: N-Acetyl- α -D-neuraminyl- $(2\rightarrow 6)$ - β -D-

galactopyranosyl- $(1\rightarrow 4)$ -D-glucose, sodium salt

Molecular mass: 655.53 Da CAS No 157574-76-0

Characteristics/Composition:

Appearance: White to off-white powder or agglomerate

Sum of 6'-Sialyllactose sodium salt, D-Lactose and Sialic acid (% of dry

matter): $\geq 94.0 \% (w/w)$

6'-Sialyllactose sodium salt (% of dry matter): \geq 90.0 % (w/w)

D-Lactose: $\leq 5.0 \%$ (w/w) Sialic acid: $\leq 2.0 \%$ (w/w)

6'-Sialyl-lactulose: $\leq 3.0 \%$ (w/w)

Sum of other carbohydrates: $\leq 3.0 \%$ (w/w)

Moisture: $\leq 6.0 \%$ (w/w) Sodium: 2.5-4.5 % (w/w) Chloride: $\leq 1.0 \%$ (w/w)

pH (20 °C, 5 % solution): 4.5-6.0 Residual protein: ≤ 0.01 % (w/w)

Microbiological criteria:

Aerobic mesophilic bacteria total plate count: ≤ 1 000 CFU/g

Enterobacteriaceae: ≤ 10 CFU/g Salmonella sp.: Absence in 25 g

Yeast: $\leq 100 \text{ CFU/g}$ Mould: $\leq 100 \text{ CFU/g}$

Residual endotoxins: ≤ 10 EU/mg

CFU: Colony Forming Units; EU: Endotoxin Units]

Yeast betaglucans

Description/Definition:

Beta-glucans are complex, high molecular mass (100–200 kDa) polysaccharides, found in the cell wall of many yeasts and cereals. The chemical name for 'yeast beta-glucans' is (1-3),(1-6)- β -D-glucans. Beta-glucans consist of a backbone of β -1-3-linked glucose residues that are branched by β -1-6-linkages, to which chitin and mannoproteins are linked by β -1-4-bonds.

Beta-glucans are isolated from yeast Saccharomyces cerevisiae.

The tertiary structure of the glucan cell wall of *Saccharomyces cerevisiae* consists of chains of β -1,3-linked glucose residues, branched by β -1,6-linkages, forming a backbone to which are linked chitin via β -1,4- bonds, β -1,6-glucans and some mannoproteins.

This novel food is available in three different forms: soluble, insoluble and insoluble in water, but dispersible in many liquid matrices.

Chemical characteristics yeast (Saccharomyces cerevisiae) betaglucans:

Soluble form:

Total carbohydrates: > 75 % Beta-glucans (1,3/1,6): > 75 %

Ash: < 4,0 % Moisture: < 8,0 % Protein: < 3,5 % Fat: < 10 % **Insoluble form:**

Total carbohydrates: > 70 %

Beta-glucans (1,3/1,6): > 70 %

Ash: ≤ 12 % Moisture: < 8,0 % Protein: < 10 % Fat: < 20 %

Insoluble in water, but dispersible in many liquid matrices:

(1,3)-(1,6)- β -D-Glucans: > 80 %

Ash: < 2,0 % Moisture: < 6,0 % Protein: < 4,0 % Total fat: < 3,0 %

Microbiological data for insoluble in water, but dispersible in many

liquid matrices:

Total plate count: < 1 000 CFU/g Enterobacteriaceae: < 100 CFU/g Total coliforms: < 10 CFU/g

Yeast: < 25 CFU/g Mould: < 25 CFU/g

Salmonella: Absence in 25 g Escherichia coli: Absence in 1 g Bacillus cereus: < 100 CFU/g

Staphylococcus aureus: Absence in 1 g

Heavy metals for insoluble in water, but dispersible in many liquid

matrices:

[FIS Lead: < 0,2 mg/kg Arsenic: < 0,2 mg/kg Mercury: < 0,1 mg/kg Cadmium: < 0,1 mg/kg]

Zeaxanthin

Description/Definition:

Zeaxanthin is a naturally occurring xanthophyll pigment, it is an oxygenated carotenoid.

The synthetic zeaxanthin is presented either as a spray-dried powder of gelatin or starch base ('beadlets') with added α -tocopherol and ascorbyl palmitate or as a corn oil suspension with added α -tocopherol. Synthetic zeaxanthin is produced by a multi-step chemical synthesis from smaller molecules.

Orange-red crystalline powder with little or no odour.

Chemical formula: C₄₀H₅₆O₂

CAS No: 144-68-3

Molecular weight: 568,9 daltons **Physical-chemical properties:**

Loss on drying: < 0,2 %

All -trans zeaxanthin: > 96 %

Cis-zeaxanthin: < 2,0 %

Other carotenoids: < 1,5 %

Triphenylphosphine oxid (CAS No 791-28-6): < 50 mg/kg

Zinc L-pidolate

Description/Definition:

Zinc L-pidolate is a white to off-white powder, with characteristic odour. International non-proprietary name (INN): L-pyroglutamic acid, Zinc salt Synonyms: Zinc 5-oxoproline, Zinc pyroglutamate, Zinc pyrrolidone carboxylate, Zinc PCA, L-Zinc pidolate

CAS No.: 15454-75-8

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Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

Molecular formula: (C₅ H₆ NO₃)₂ Zn Relative anhydrous molecular mass: 321,4 Appearance: White to slightly white powder

Purity:

Zinc L-pidolate (purity): \geq 98 % pH (10 % aqueous sol.): 5,0-6,0 Specific rotation: 19,6 ° - 22,8 °

Water: $\le 10,0 \%$

Glutamic acid: < 2,0 %

Heavy metals: Lead: ≤ 3,0 ppm Arsenic: ≤ 2,0 ppm Cadmium: ≤ 1,0 ppm Mercury: ≤ 0,1 ppm Microbiological criteria:

Total viable mesophilic count: ≤ 1 000 CFU/g

Yeasts and moulds: ≤ 100 CFU/g

Pathogen: Absence

- a Commission Regulation (EU) No 231/2012 of 9 March 2012 laying down specifications for food additives listed in Annexes II and III to Regulation (EC) No 1333/2008 of the European Parliament and of the Council (OJ L 83, 22.3.2012, p. 1).
- b Commission Implementing Regulation (EU) 2015/175 of 5 February 2015 laying down special conditions applicable to the import of guar gum originating in or consigned from India due to contamination risks by pentachlorophenol and dioxins (OJ L 30, 6.2.2015, p. 10).
- c [F12OSC-DMAC (4-dimethylaminocinnamaldehyde) method (Ocean Spray Cranberries, Inc) Martin MA, Ramos S, Mateos R, Marais JPJ, Bravo-Clemente, L, Khoo C and Goya L. Food Res Intl 2015 71: 68-82. Modified from Cunningham DG, Vannozzi S, O'Shea E, Turk R (2002) In: Ho C-T, Zheng QY (eds) Quality Management of Nutraceuticals ACS Symposium series 803, Washington DC. Quantitation of PACs by DMAC Color Reaction pp 151-166.
- d BL-DMAC 4-dimethylaminocinnamaldehyde) method (Brunswick Lab) Multi-laboratory validation of a standard method for quantifying proanthocyanidins in cranberry powders. Prior RL, Fan E, Ji H, Howell A, Nio C, Payne MJ, Reed J. J Sci Food Agric. 2010 Jul;90(9):1473-8.
- e The different values for these three parameters are due to the different methods used.
- f GAE: Gallic Acid Equivalents.
- **g** [F57CFU: Colony Forming Units.]]
- h [F8HPLC/RI: High-performance liquid chromatography coupled with refractive index detection.
- i CFU: Colony-forming unit.]
- **j** [F50To be tested immediately after the heat-treatment step. Measures have to be in place to prevent cross-contamination with viable *Yarrowia lipolytica* cells during packaging and/or storage of the NF.]
- k [F203'-Fucosyllactose, 2'-Fucosyl-galactose, Glucose, Galactose, Mannitol, Sorbitol, Galactitol, Trihexose, Allo-lactose and other structurally related carbohydrates.]
- I [F7Converted from International Units (IU) using the conversion factor of 0,025 μ g = 1 IU.]]

Textual Amendments

F57 Substituted by Commission Implementing Regulation (EU) 2020/1163 of 6 August 2020 authorising the placing on the market of vitamin D2 mushroom powder as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).

- **F58** Substituted by Commission Implementing Regulation (EU) 2019/108 of 24 January 2019 authorising the change of specifications of the novel food ingredient lipid extract from Antarctic Krill (Euphausia superba) under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F59** Substituted by Commission Implementing Regulation (EU) 2019/2165 of 17 December 2019 authorising the change of the specifications of the novel food coriander seed oil from Coriandrum sativum under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F60** Substituted by Commission Implementing Regulation (EU) 2019/388 of 11 March 2019 authorising the change of the specifications of the novel food 2'-fucosyllactose produced with Escherichia coli K-12 under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F61** Inserted by Commission implementing Regulation (EU) 2019/1979 of 26 November 2019 authorising the placing on the market of 2'-Fucosyllactose/Difucosyllactose mixture as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- **F62** Words in Annex Table 2 substituted (W.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Wales) Regulations 2022 (S.I. 2022/575), reg. 1(4), **Sch. 1 para. 2(a)**
- **F63** Words in Annex Table 2 substituted (W.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Wales) Regulations 2022 (S.I. 2022/575), reg. 1(4), **Sch. 1 para. 2(b)**
- **F64** Words in Annex Table 2 substituted (E.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (England) Regulations 2022 (S.I. 2022/560), regs. 1(3), **Sch. 4**
- **F65** Words in Annex Table 2 substituted (S.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Scotland) Regulations 2022 (S.S.I. 2022/168), reg. 1(3), sch. 1 para. 2
- **F66** Substituted by Commission Implementing Regulation (EU) 2019/1314 of 2 August 2019 authorising the change of the specifications of the novel food Lacto-N-neotetraose produced with Escherichia coli K-12 under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- F67 Words in Annex Table 2 inserted (E.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (England) Regulations 2022 (S.I. 2022/560), regs. 1(3), Sch. 5; inserted (S.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Scotland) Regulations 2022 (S.S.I. 2022/168), reg. 1(3), sch. 2 para. 2; and inserted (W.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Wales) Regulations 2022 (S.I. 2022/575), reg. 1(4), Sch. 2 para. 2
- **F68** Substituted by Commission Implementing Regulation (EU) 2020/478 of 1 April 2020 correcting Implementing Regulation (EU) 2017/2470 establishing the Union list of novel foods (Text with EEA relevance).
- F69 Words in Annex Table 2 inserted (E.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (England) Regulations 2022 (S.I. 2022/560), regs. 1(3), Sch. 6; inserted (S.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Scotland) Regulations 2022 (S.S.I. 2022/168), reg. 1(3), sch. 3 para. 2; and inserted (W.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Wales) Regulations 2022 (S.I. 2022/575), reg. 1(4), Sch. 3 para. 2
- **F70** Substituted by Commission Implementing Regulation (EU) 2020/443 of 25 March 2020 authorising the change of the specifications of the novel food spermidine-rich wheat germ extract (Triticum aestivum) under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- F71 Words in Annex Table 2 substituted (E.) (15.5.2023) by The Food Additives, Food Flavourings and Novel Foods (Authorisations) (England) Regulations 2023 (S.I. 2023/334), reg. 1(1), Sch. 6 (with reg. 4);

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Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX. (See end of Document for details)

- substituted (S.) (15.5.2023) by The Food Additives, Food Flavourings and Novel Foods (Authorisations) (Scotland) Regulations 2023 (S.S.I. 2023/78), reg. 1(1), sch. 4 para. 2 (with reg. 5); and substituted (W.) (15.5.2023) by The Food Additives, Food Flavourings, and Novel Foods (Authorisations) and Food and Feed (Miscellaneous Amendments) (Wales) Regulations 2023 (S.I. 2023/343), reg. 1(4), Sch. 4 para. 3 (with reg. 4)
- F72 Words in Annex Table 2 inserted (E.) (15.5.2023) by The Food Additives, Food Flavourings and Novel Foods (Authorisations) (England) Regulations 2023 (S.I. 2023/334), reg. 1(1), Sch. 7 (with reg. 4); inserted (S.) (15.5.2023) by The Food Additives, Food Flavourings and Novel Foods (Authorisations) (Scotland) Regulations 2023 (S.S.I. 2023/78), reg. 1(1), sch. 5 para. 2 (with reg. 5); and inserted (W.) (15.5.2023) by The Food Additives, Food Flavourings, and Novel Foods (Authorisations) and Food and Feed (Miscellaneous Amendments) (Wales) Regulations 2023 (S.I. 2023/343), reg. 1(4), Sch. 5 para. 3 (with reg. 4
- **F73** Inserted by Commission Implementing Regulation (EU) 2018/1648 of 29 October 2018 authorising the placing on the market of xylo-oligosaccharides as a novel food under Regulation (EU) 2015/2283 of the European Parliament and of the Council and amending Commission Implementing Regulation (EU) 2017/2470 (Text with EEA relevance).
- F74 Words in Annex Table 2 inserted (E.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (England) Regulations 2022 (S.I. 2022/560), regs. 1(3), Sch. 7; inserted (S.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Scotland) Regulations 2022 (S.S.I. 2022/168), reg. 1(3), sch. 4 para. 2; and inserted (W.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Wales) Regulations 2022 (S.I. 2022/575), reg. 1(4), Sch. 4 para. 2
- F75 Words in Annex Table 2 inserted (E.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (England) Regulations 2022 (S.I. 2022/560), regs. 1(3), Sch. 7 (as amended by S.I. 2022/619, regs. 1(1), 2(3)); inserted (S.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Scotland) Regulations 2022 (S.S.I. 2022/168), reg. 1(3), sch. 5 para. 2; and inserted (W.) (30.6.2022) by The Novel Foods (Authorisations) and Smoke Flavourings (Modification of Authorisations) (Wales) Regulations 2022 (S.I. 2022/575), reg. 1(4), Sch. 5 para. 2

Changes to legislation:

There are currently no known outstanding effects for the Commission Implementing Regulation (EU) 2017/2470, ANNEX.