This text is meant purely as a documentation tool and has no legal effect. The Union's institutions do not assume any liability for its contents. The authentic versions of the relevant acts, including their preambles, are those published in the Official Journal of the European Union and available in EUR-Lex. Those official texts are directly accessible through the links embedded in this document

$ightharpoonup \underline{B}$ 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)

(OJ L 351, 30.12.2017, p. 72)

Amended by:

		(Official Jou	rnal
		No	page	date
► <u>M1</u>	Commission Implementing Regulation (EU) 2018/460 of 20 March 2018	L 78	2	21.3.2018
<u>M2</u>	Commission Implementing Regulation (EU) 2018/461 of 20 March 2018	L 78	7	21.3.2018
► <u>M3</u>	Commission Implementing Regulation (EU) 2018/462 of 20 March 2018	L 78	11	21.3.2018
► <u>M4</u>	Commission Implementing Regulation (EU) 2018/469 of 21 March 2018	L 79	11	22.3.2018
► <u>M5</u>	Commission Implementing Regulation (EU) 2018/991 of 12 July 2018	L 177	9	13.7.2018
<u>M6</u>	Commission Implementing Regulation (EU) 2018/1011 of 17 July 2018	L 181	4	18.7.2018
<u>M7</u>	Commission Implementing Regulation (EU) 2018/1018 of 18 July 2018	L 183	9	19.7.2018
<u>M8</u>	Commission Implementing Regulation (EU) 2018/1032 of 20 July 2018	L 185	9	23.7.2018
► <u>M9</u>	Commission Implementing Regulation (EU) 2018/1023 of 23 July 2018	L 187	1	24.7.2018
► <u>M10</u>	Commission Implementing Regulation (EU) 2018/1122 of 10 August 2018	L 204	36	13.8.2018
► <u>M11</u>	Commission Implementing Regulation (EU) 2018/1123 of 10 August 2018	L 204	41	13.8.2018
► <u>M12</u>	Commission Implementing Regulation (EU) 2018/1132 of 13 August 2018	L 205	15	14.8.2018
► <u>M13</u>	Commission Implementing Regulation (EU) 2018/1133 of 13 August 2018	L 205	18	14.8.2018
► <u>M14</u>	Commission Implementing Regulation (EU) 2018/1293 of 26 September 2018	L 243	2	27.9.2018
► <u>M15</u>	Commission Implementing Regulation (EU) 2018/1631 of 30 October 2018	L 272	17	31.10.2018
► <u>M16</u>	Commission Implementing Regulation (EU) 2018/1632 of 30 October 2018	L 272	23	31.10.2018

► <u>M17</u>	Commission Implementing Regulation (EU) 2018/1633 of 30 October 2018	L 272	29	31.10.2018
► <u>M18</u>	Commission Implementing Regulation (EU) 2018/1647 of 31 October 2018	L 274	51	5.11.2018
► <u>M19</u>	Commission Implementing Regulation (EU) 2018/1648 of 29 October 2018	L 275	1	6.11.2018
► <u>M20</u>	amended by Commission Implementing Regulation (EU) 2023/65 of 6 January 2023	L 6	1	9.1.2023
► <u>M21</u>	Commission Implementing Regulation (EU) 2018/1991 of 13 December 2018	L 320	22	17.12.2018
► <u>M22</u>	Commission Implementing Regulation (EU) 2018/2016 of 18 December 2018	L 323	1	19.12.2018
► <u>M23</u>	Commission Implementing Regulation (EU) 2018/2017 of 18 December 2018	L 323	4	19.12.2018
► <u>M24</u>	Commission Implementing Regulation (EU) 2019/108 of 24 January 2019	L 23	4	25.1.2019
► <u>M25</u>	Commission Implementing Regulation (EU) 2019/109 of 24 January 2019	L 23	7	25.1.2019
► <u>M26</u>	Commission Implementing Regulation (EU) 2019/110 of 24 January 2019	L 23	11	25.1.2019
► <u>M27</u>	Commission Implementing Regulation (EU) 2019/387 of 11 March 2019	L 70	17	12.3.2019
► <u>M28</u>	Commission Implementing Regulation (EU) 2019/388 of 11 March 2019	L 70	21	12.3.2019
► <u>M29</u>	Commission Implementing Regulation (EU) 2019/456 of 20 March 2019	L 79	13	21.3.2019
► <u>M30</u>	Commission Implementing Regulation (EU) 2019/506 of 26 March 2019	L 85	11	27.3.2019
►M31	Commission Implementing Regulation (EU) 2019/760 of 13 May 2019	L 125	13	14.5.2019
► M32	Commission Implementing Regulation (EU) 2019/1272 of 29 July 2019	L 201	3	30.7.2019
► M33	Commission Implementing Regulation (EU) 2019/12/2 of 27 July 2019	L 201	16	2.8.2019
<u> 14155</u>	2019	L 204	10	2.8.2019
► <u>M34</u>	Commission Implementing Regulation (EU) 2019/1314 of 2 August 2019	L 205	4	5.8.2019
► <u>M35</u>	Commission Implementing Regulation (EU) 2019/1686 of 8 October 2019	L 258	13	9.10.2019
► <u>M36</u>	amended by Commission Implementing Regulation (EU) 2023/65 of 6 January 2023	L 6	1	9.1.2023
► <u>M37</u>	Commission Implementing Regulation (EU) 2019/1976 of 25 November 2019	L 308	40	29.11.2019
► <u>M38</u>	Commission Implementing Regulation (EU) 2019/1979 of 26 November 2019	L 308	62	29.11.2019
► <u>M39</u>	Commission Implementing Regulation (EU) 2019/2165 of 17 December 2019	L 328	81	18.12.2019
► <u>M40</u>	Commission Implementing Regulation (EU) 2020/16 of 10 January 2020	L 7	6	13.1.2020
► <u>M41</u>	Commission Implementing Regulation (EU) 2020/24 of 13 January 2020	L 8	12	14.1.2020
► <u>M42</u>	Commission Implementing Regulation (EU) 2020/206 of 14 February 2020	L 43	66	17.2.2020
► <u>M43</u>	Commission Implementing Regulation (EU) 2020/443 of 25 March 2020	L 92	7	26.3.2020

► <u>M44</u>	Commission Implementing Regulation (EU) 2020/478 of 1 April 2020	L 102	1	2.4.2020
► <u>M45</u>	Commission Implementing Regulation (EU) 2020/484 of 2 April 2020	L 103	3	3.4.2020
► <u>M46</u>	amended by Commission Implementing Regulation (EU) 2021/1318 of 9 August 2021	L 286	5	10.8.2021
► <u>M47</u>	Commission Implementing Regulation (EU) 2020/500 of 6 April 2020	L 109	2	7.4.2020
► <u>M48</u>	Commission Implementing Regulation (EU) 2020/916 of 1 July 2020	L 209	6	2.7.2020
► <u>M49</u>	Commission Implementing Regulation (EU) 2020/917 of 1 July 2020	L 209	10	2.7.2020
► <u>M50</u>	Commission Implementing Regulation (EU) 2020/973 of 6 July 2020	L 215	7	7.7.2020
► <u>M51</u>	Commission Implementing Regulation (EU) 2020/1163 of 6 August 2020	L 258	1	7.8.2020
► <u>M52</u>	Commission Implementing Regulation (EU) 2020/1559 of 26 October 2020	L 357	7	27.10.2020
► <u>M53</u>	Commission Implementing Regulation (EU) 2020/1634 of 4 November 2020	L 367	39	5.11.2020
► <u>M54</u>	Commission Implementing Regulation (EU) 2020/1820 of 2 December 2020	L 406	29	3.12.2020
► <u>M55</u>	Commission Implementing Regulation (EU) 2020/1821 of 2 December 2020	L 406	34	3.12.2020
► <u>M56</u>	Commission Implementing Regulation (EU) 2020/1822 of 2 December 2020	L 406	39	3.12.2020
► <u>M57</u>	Commission Implementing Regulation (EU) 2020/1993 of 4 December 2020	L 410	62	7.12.2020
► <u>M58</u>	Commission Implementing Regulation (EU) 2021/50 of 22 January 2021	L 23	7	25.1.2021
► <u>M59</u>	Commission Implementing Regulation (EU) 2021/51 of 22 January 2021	L 23	10	25.1.2021
► <u>M60</u>	Commission Implementing Regulation (EU) 2021/82 of 27 January 2021	L 29	16	28.1.2021
► <u>M61</u>	Commission Implementing Regulation (EU) 2021/96 of 28 January 2021	L 31	201	29.1.2021
► <u>M62</u>	amended by Commission Implementing Regulation (EU) 2023/65 of 6 January 2023 $$	L 6	1	9.1.2023
► <u>M63</u>	Commission Implementing Regulation (EU) 2021/120 of 2 February 2021	L 37	1	3.2.2021
► <u>M64</u>	Commission Implementing Regulation (EU) 2021/668 of 23 April 2021	L 141	3	26.4.2021
► <u>M65</u>	Commission Implementing Regulation (EU) 2021/670 of 23 April 2021	L 141	14	26.4.2021
► <u>M66</u>	Commission Implementing Regulation (EU) 2021/882 of 1 June 2021	L 194	16	2.6.2021
► <u>M67</u>	Commission Implementing Regulation (EU) 2021/900 of 3 June 2021	L 197	71	4.6.2021
► <u>M68</u>	Commission Implementing Regulation (EU) 2021/912 of 4 June 2021	L 199	10	7.6.2021
► <u>M69</u>	Commission Implementing Regulation (EU) 2021/1318 of 9 August 2021	L 286	5	10.8.2021
► <u>M70</u>	Commission Implementing Regulation (EU) 2021/1319 of 9 August 2021	L 286	12	10.8.2021
► <u>M71</u>	Commission Implementing Regulation (EU) 2021/1326 of 10 August 2021	L 288	24	11.8.2021
► <u>M72</u>	Commission Implementing Regulation (EU) 2021/1377 of 19 August 2021	L 297	20	20.8.2021
► <u>M73</u>	Commission Implementing Regulation (EU) 2021/1974 of 12 November 2021	L 402	5	15.11.2021
► <u>M74</u>	Commission Implementing Regulation (EU) 2021/1975 of 12 November 2021	L 402	10	15.11.2021
► <u>M75</u>	Commission Implementing Regulation (EU) 2021/2029 of 19 November 2021	L 415	9	22.11.2021
► <u>M76</u>	Commission Implementing Regulation (EU) 2021/2079 of 26 November 2021	L 426	16	29.11.2021

► <u>M77</u>	Commission Implementing Regulation (EU) 2021/2129 of 2 December 2021	L 432	13	3.12.2021
► <u>M78</u>	Commission Implementing Regulation (EU) 2021/2191 of 10 December 2021	L 445	1	13.12.2021
► <u>M79</u>	Commission Implementing Regulation (EU) 2022/47 of 13 January 2022	L 9	29	14.1.2022
► <u>M80</u>	Commission Implementing Regulation (EU) 2022/168 of 8 February 2022	L 28	5	9.2.2022
► <u>M81</u>	Commission Implementing Regulation (EU) 2022/169 of 8 February 2022	L 28	10	9.2.2022
► <u>M82</u>	Commission Implementing Regulation (EU) 2022/187 of 10 February 2022	L 30	102	11.2.2022
► <u>M83</u>	Commission Implementing Regulation (EU) 2022/188 of 10 February 2022	L 30	108	11.2.2022
► <u>M84</u>	Commission Implementing Regulation (EU) 2022/196 of 11 February 2022	L 31	46	14.2.2022
► <u>M85</u>	Commission Implementing Regulation (EU) 2022/202 of 14 February 2022	L 33	41	15.2.2022
► <u>M86</u>	Commission Implementing Regulation (EU) 2022/672 of 22 April 2022	L 122	24	25.4.2022
► <u>M87</u>	Commission Implementing Regulation (EU) 2022/673 of 22 April 2022	L 122	27	25.4.2022
► <u>M88</u>	Commission Implementing Regulation (EU) 2022/684 of 28 April 2022	L 126	10	29.4.2022
► <u>M89</u>	Commission Implementing Regulation (EU) 2022/961 of 20 June 2022	L 165	41	21.6.2022
► <u>M90</u>	Commission Implementing Regulation (EU) 2022/965 of 21 June 2022	L 166	118	22.6.2022
► <u>M91</u>	Commission Implementing Regulation (EU) 2022/966 of 21 June 2022	L 166	125	22.6.2022
► <u>M92</u>	Commission Implementing Regulation (EU) 2022/1160 of 5 July 2022	L 179	25	6.7.2022
► <u>M93</u>	Commission Implementing Regulation (EU) 2022/1365 of 4 August 2022	L 205	230	5.8.2022
► <u>M94</u>	Commission Implementing Regulation (EU) 2022/1373 of 5 August 2022	L 206	28	8.8.2022
► <u>M95</u>	Commission Implementing Regulation (EU) 2022/1381 of 8 August 2022	L 207	12	9.8.2022
► <u>M96</u>	Commission Implementing Regulation (EU) 2022/2534 of 21 December 2022	L 328	85	22.12.2022
► <u>M97</u>	Commission Implementing Regulation (EU) 2022/2535 of 21 December 2022	L 328	91	22.12.2022
► <u>M98</u>	Commission Implementing Regulation (EU) 2023/4 of 3 January 2023	L 2	3	4.1.2023
► <u>M99</u>	Commission Implementing Regulation (EU) 2023/5 of 3 January 2023	L 2	9	4.1.2023
► <u>M100</u>	Commission Implementing Regulation (EU) 2023/6 of 3 January 2023	L 2	16	4.1.2023
► <u>M101</u>	Commission Implementing Regulation (EU) 2023/7 of 3 January 2023	L 2	21	4.1.2023
► <u>M102</u>	Commission Implementing Regulation (EU) 2023/52 of 4 January 2023	L 3	1	5.1.2023
► <u>M103</u>	Commission Implementing Regulation (EU) 2023/58 of 5 January 2023	L 5	10	6.1.2023
► <u>M104</u>	Commission Implementing Regulation (EU) 2023/65 of 6 January 2023	L 6	1	9.1.2023
► <u>M105</u>	Commission Implementing Regulation (EU) 2023/113 of 16 January 2023	L 15	1	17.1.2023

► <u>M106</u>	Commission Implementing Regulation (EU) 2023/267 of 8 February 2023	L 39	1	9.2.2023
► <u>M107</u>	Commission Implementing Regulation (EU) 2023/463 of 3 March 2023	L 68	32	6.3.2023
► <u>M108</u>	Commission Implementing Regulation (EU) 2023/652 of 20 March 2023	L 81	23	21.3.2023
► <u>M109</u>	Commission Implementing Regulation (EU) 2023/667 of 22 March 2023	L 84	3	23.3.2023
► <u>M110</u>	Commission Implementing Regulation (EU) 2023/859 of 25 April 2023	L 111	17	26.4.2023
► <u>M111</u>	Commission Implementing Regulation (EU) 2023/931 of 8 May 2023	L 124	1	10.5.2023
► <u>M112</u>	Commission Implementing Regulation (EU) 2023/937 of 10 May 2023	L 125	12	11.5.2023
► <u>M113</u>	Commission Implementing Regulation (EU) 2023/938 of 10 May 2023	L 125	16	11.5.2023
► <u>M114</u>	Commission Implementing Regulation (EU) 2023/943 of 11 May 2023	L 126	41	12.5.2023
► <u>M115</u>	Commission Implementing Regulation (EU) 2023/948 of 12 May 2023	L 128	52	15.5.2023
► <u>M116</u>	Commission Implementing Regulation (EU) 2023/949 of 12 May 2023	L 128	60	15.5.2023
► <u>M117</u>	Commission Implementing Regulation (EU) 2023/950 of 12 May 2023	L 128	68	15.5.2023
► <u>M118</u>	Commission Implementing Regulation (EU) 2023/951 of 12 May 2023	L 128	73	15.5.2023
► <u>M119</u>	Commission Implementing Regulation (EU) 2023/961 of 12 May 2023	L 129	3	16.5.2023
► <u>M120</u>	Commission Implementing Regulation (EU) 2023/972 of 10 May 2023	L 132	46	17.5.2023
► <u>M121</u>	Commission Implementing Regulation (EU) 2023/1581 of 1 August 2023	L 194	4	2.8.2023
► <u>M122</u>	Commission Implementing Regulation (EU) 2023/1582 of 1 August 2023	L 194	8	2.8.2023
► <u>M123</u>	Commission Implementing Regulation (EU) 2023/1583 of 1 August 2023	L 194	13	2.8.2023
► <u>M124</u>	Commission Implementing Regulation (EU) 2023/2145 of 16 October 2023	L 2145	1	17.10.2023
► <u>M125</u>	Commission Implementing Regulation (EU) 2023/2210 of 20 October 2023	L 2210	1	23.10.2023
► <u>M126</u>	Commission Implementing Regulation (EU) 2023/2214 of 23 October 2023	L 2214	1	24.10.2023
► <u>M127</u>	Commission Implementing Regulation (EU) 2023/2215 of 23 October 2023	L 2215	1	24.10.2023
► <u>M128</u>	Commission Implementing Regulation (EU) 2023/2847 of 20 December 2023	L 2847	1	21.12.2023
► <u>M129</u>	Commission Implementing Regulation (EU) 2023/2851 of 20 December 2023	L 2851	1	21.12.2023
► <u>M130</u>	Commission Implementing Regulation (EU) 2024/1023 of 8 April 2024	L 1023	1	9.4.2024
► <u>M131</u>	Commission Implementing Regulation (EU) 2024/1026 of 8 April 2024	L 1026	1	9.4.2024
► <u>M132</u>	Commission Implementing Regulation (EU) 2024/1027 of 8 April 2024	L 1027	1	9.4.2024
► <u>M133</u>	Commission Implementing Regulation (EU) 2024/1037 of 9 April 2024	L 1037	1	10.4.2024
► <u>M134</u>	Commission Implementing Regulation (EU) 2024/1046 of 9 April 2024	L 1046	1	10.4.2024

	02017R2470 —	EN — 01.05	.2024 —	<u> - 046.001 — 6</u>
► <u>M135</u>	Commission Implementing Regulation (EU) 2024/1047 of 9 April 2024	L 1047	1	10.4.2024
► <u>M136</u>	Commission Implementing Regulation (EU) 2024/1048 of 9 April 2024	L 1048	1	10.4.2024

11.4.2024

► M137 Commission Implementing Regulation (EU) 2024/1052 of 10 April L 1052 2024

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)

Article 1

Union list of authorised novel foods

The Union list of novel foods authorised to be placed on the market within the Union as referred to in Article 6(1) of Regulation (EU) 2015/2283 is hereby established and set out in the Annex to this Regulation.

Article 2

This Regulation shall enter into force on the twentieth day following that of its publication in the Official Journal of the European Union.

This Regulation shall be binding in its entirety and directly applicable in all Member States.

ANNEX

UNION LIST OF NOVEL FOODS

Content of the list

- 1. The Union list shall consist of Tables 1 and 2.
- 2. Table 1 includes the authorised novel foods and contains the following information:
 - Column 1: Authorised novel food
 - Column 2: Conditions under which the novel food may be used. This column is further subdivided into two: Specified food category and Maximum levels
 - Column 3: Additional specific labelling requirements
 - Column 4: Other requirements
- 3. Table 2 includes the specifications on novel foods and contains the following information:
 - Column 1: Authorised novel food
 - Column 2: Specifications

Table 1: Authorised novel foods

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀				
N-Acetyl-D-neuraminic acid	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs						
	Infant and follow-on formulae as defined by Regulation (EU) No 609/2013 (1)	0,05 g/L of reconstituted formula	neuraminic acid' Food supplements containing N- acetyl-D-neuraminic acid shall bear a statement that the food supplement should not be given 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.	neuraminic acid' Food supplements containing N- acetyl-D-neuraminic acid shall bear a statement that the food supplement should not be given 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	Food supplements containing <i>N</i> -acetyl-D-neuraminic acid shall bear a statement that the food supplement should not be given to infants, young children and children under 10 years of age where they consume breast milk or other foods with added <i>N</i> -acetyl-D-neuraminic	neuraminic acid' Food supplements containing <i>N</i> - acetyl-D-neuraminic acid shall bear a statement that the food supplement should not be given to infants, young children and children under 10 years of age where they consume breast milk or other foods with added <i>N</i> -acetyl-D-neuraminic	neuraminic acid' Food supplements containing <i>N</i> -acetyl-D-neuraminic acid shall bear		
	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							
	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.							
	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 requirements of Commission Implementing Regulation (EU) No 828/2014 (²)	1,25 g/kg							
	Unflavoured pasteurised and sterilised (including UHT) milk-based products	0,05 g/L							

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	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 extracts; tea, plant, fruit and cereal preparations for infusions	0,2 g/kg			
	Food Supplements as defined in Directive 2002/46/EC (³)	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			

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
▼ <u>M99</u>						
	Acheta domesticus (house cricket) partially defatted powder	ouse cricket) artially defatted	Maximum levels (g/100 g) 1. (marketed as such or reconstituted according to the instructions)	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Acheta'		Authorised on 24.1.2023 This inclusion is based o proprietary scientifi evidence and scientific dat
	powder	Multigrain bread and rolls; crackers and breadsticks	2	domesticus (house cricket) partially defatted powder'. 2. The labelling of the foodstuffs		protected in accordance with Article 26 of Regulation (EU
		Cereal bars	3	containing Acheta domesticus		2015/2283.
		Pre-mixes for baked products (dry)	3	(house cricket) partially defatted powder shall bear a statement that this ingredient may cause allergic reactions to consumers with known allergies to crus- taceans, molluscs, and products		Applicant: 'Cricket One Co Ltd', 383/3/51 Quang Trun
		Biscuits	1,5			street, Ward 10, Go Va
		Pasta-based products (dry)	0,25			district, Ho Chi Minh Cit
		Stuffed pasta-based products (dry)	3			Vietnam.
		Sauces	1	thereof, and to dust mites.		During the period of day protection, the novel foo
		Processed potato products, legume- and vegetable-based dishes, pizza, pasta-based dishes	1	This statement shall appear in close proximity to the list of ingredients.		Acheta domesticus (he cricket) partially defa powder is authorised
		Whey powder	3	1		placing on the marked within the Union only b
		Meat analogues	5	1		'Cricket One Co. Ltd', 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)
		Soups and soup concentrates or powders	1	1		
		Maize flour based snacks	4	1		
		Beer-like beverages	0,1	1		
		Chocolate confectionary	2	1		
		Nuts and oilseeds	2	1		
		Snacks other than chips	5	1		
		Meat preparations	2			2015/2283, or with the agreement of 'Cricket Or
						Co. Ltd'. End date of the da protection: 24.1.2028.
▼ <u>M9</u>	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'		

02017R2470 - EN - 01.05.2024 - 046.001 - 11

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀		
03							
Frozen, paste, dried and powder forms of <i>Alphitobius</i>	Specified food category Cereal bars	Maximum levels (g/100g) 25 (Dried form) 25 (Powder form)	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Frozen/		Authorised on 26.1.20 This inclusion is based proprietary scientific of		
diaperinus larvae (lesser mealworm)	Bread and rolls	20 (Powder form)	paste Alphitobius diaperinus larvae (lesser mealworm)' or		protected in accordance of Article 26 of Regulation (
	Processed and breakfast cereals	10 (Dried form) 10 (Powder form)	'Dried/powder Alphitobius diaperinus larvae (lesser mealworm)' depending on the		2015/2283. Applicant: Ynsect NL I Harderwijkerweg 14		
	Porridge	15 (Powder form)	form used.		3852 AB Ermelo,		
	Pre-mixes (dry) for baked products	10 (Powder form)	2. The labelling of food supplements containing the		Netherlands.		
	Dried pasta-based products	10 (Powder form)	novel food shall bear a		During the period of data protection, the novel food i		
	Stuffed pasta-based products	28 (Frozen or paste form) 10 (Powder form)	statement that those food supplements should not be consumed by persons under 18 years of age. 3. The labelling of the foodstuffs containing frozen, paste, dried or powder forms of Alphitobius diaperinus larvae (lesser mealworm) shall bear a statement that this ingredient may cause allergic reactions to consumers with known allergies to crustaceans, and products thereof, and to dust mites. This statement shall appear in close proximity to the list of ingredients.	supplements should not be	supplements should not be	authorised for placing on the market within the Union onleby Ynsect NL B.V., unless	
	Whey powder	35 (Powder form)			subsequent applicant obtain authorisation for that nove food without reference to the proprietary scientifi		
	Soups	15 (Powder form)					
	Cereal-, pasta-based dishes	5 (Powder form)					
	Pizza-based dishes	5 (Dried form) 5 (Powder form)		mealworm) shall bear a	with	data protected in accordance with Article 26 of Regulation (EU) 2015/2283, or	
	Noodles	10 (Powder form)			with the agreement		
	Snacks other than chips	10 (Dried form) 10 (Powder form)		to crustaceans, and products		Ynsect NL B.V. End date of the protection: 26.1.2028.	
	Chips/crisps	10 (Powder form)			protection: 20.1.2026.		
	Crackers and bread sticks	10 (Powder form)					
	Peanut butter	15 (Powder form)	ingredients.				
	Ready-to-eat savoury based sandwich	20 (Powder form)					
	Meat preparations	14 (Frozen or paste form) 5 (Powder form)					
	Meat analogues	40 (Frozen or paste form) 15 (Powder form)					
	Milk and dairy analogues	10 (Powder form)					
	Chocolate confectionary	5 (Powder form)					
	Food supplements as defined in Directive 2002/46/EC for the adult population	4 g/day (Powder form)	-				

V 1V12						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Ajuga reptans extract from cell	Specified food category	Maximum levels			
	cultures	Food Supplements as defined in Directive 2002/46/EC	In line with normal use in food supplements of a similar extract of the flowering aerial parts of <i>Ajuga reptans</i>			
▼ <u>M80</u>	<u>.</u>					
	Akkermansia muci- niphila (pasteurised)	Foods for special medical purposes as defined under Regulation (EU) No 609/2013 for the adult population, excluding pregnant and lactating women	3.4×10^{10} cells/day	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'pasteurised Akkermansia muciniphila'.		Authorised on 1 March 2022. This inclusion is based on proprietary scientific evidence and scientific data protected in
		Food supplements as defined in Directive 2002/46/EC for the adult population, excluding pregnant and lactating women	3.4×10^{10} cells/day	The labelling of food supplements containing pasteurised Akkermansia muciniphila shall bear a statement that they should be consumed by adults only, excluding pregnant and lactating women.		accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: A-Mansia Biotech S.A., rue Granbonpré, 11, Bâtiment H, 1435 Mont-Saint-Guibert. Belgium. During the period of data protection, the novel food pasteurised Akkermansia muciniphila is authorised for placing on the market within the Union only by A-Mansia Biotech 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 Mansia Biotech S.A End date of the data protection: 1 March 2027.

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
L-Alanyl-L- Glutamine	Specified food category	Maximum levels			
	Food Supplements as defined in Directive 2002/46/EC				
	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				
Algal oil from the microalgae <i>Ulkenia</i> sp.	Specified 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		
	Bakery products (breads, rolls and sweet biscuits)	200 mg/100 g	micro-algae Ulkenia sp.'		
	Cereal bars	500 mg/100 g			
	Non-alcoholic beverages (including milk based beverages)	60 mg/100 ml			

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
<u>M26</u>						
	Allanblackia seed oil	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
		Yellow fat spreads and cream based spreads	30 g/100 g	containing it shall be 'Allanblackia seed oil'		
		Mixtures of vegetable oils (*) and milk (falling under the food category: Dairy analogues, including beverage whiteners)	30 g/100 g			
		(*) Except olive oils and olive pomace oils as Regulation (EU) No 1308/2013.	defined in Part VIII of Annex VII of			
<u>M9</u>						
	Aloe macroclada Baker leaf extract	Specified food category	Maximum levels			
		Food Supplements as defined in Directive 2002/46/EC	In line with normal use 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		
		Dairy products except milk-based drinks	200 mg/100 g or for cheese products 600 mg/100 g	from the crustacean Antarctic Krill (Euphausia superba)'		
		Dairy analogues except drinks	200 mg/100 g or for analogues to cheese products 600 mg/			

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	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			

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	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				
Antarctic Krill oil rich in phosp- holipids from Euphausia superba	Specified food category	Maximum levels of combined DHA and EPA	(Eupnausia superva)		
	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/			
	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			

Auth	horised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		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 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				
	phorata mycelia	Specified food category	Maximum levels	The designation of the novel food on the labelling of food		
powd	der	Food supplements as defined in Directive 2002/46/EC, excluding infants, children, and adolescents younger than 14 years of age	990 mg/day	supplements containing it shall be 'Antrodia camphorata mycelia powder'. 2. The labelling of the food supplements containing Antrodia camphorata mycelia powder shall bear a statement that this food supplement should not be consumed by infants, children, and adolescents younger than 14 years of age.		

Aqueous ethanolic extract of Labisia pumila Food supplements as defined in Directive 2002/46/EC for the adult population, excluding pregnant and lactating women Specified food category Maximum levels 1. The designation of the novel food on the labelling of the foodstuffs containing it shall be 'aqueous ethanolic extract of Labisia pumila'. 2. The labelling of food supplements containing the novel food supplements containing the novel food shall bear a statement that they should only be consumed by persons above 18 years of age excluding pregnant and lactating women.	quirements ► M30 Data Protection
Aqueous ethanolic extract of Labisia pumila Food supplements as defined in Directive 2002/46/EC for the adult population, excluding pregnant and lactating women Specified food category Maximum levels 1. The designation of the novel food on the labelling of the foodstuffs containing it shall be 'aqueous ethanolic extract of Labisia pumila'. 2. The labelling of food supplements containing the novel food shall bear a statement that they should only be consumed by persons above 18 years of age excluding	
	Authorised on 6 June This inclusion is bas proprietary sci evidence and scientifi protected in accordance Article 26 of lation (EU) 2015/2283 Applicant: Medika Sdn. Bhd., No. 44B Bola Tampar 13/14 States 13, 40100 Shah Selangor, Malaysia. If the period of data protected in accordance than olic extract of Inpumila is authorised placing on the within the Union on Medika Natura Sdn. unless a substantial applicant obtains au ation for the novel without reference to proprietary sci evidence or scientific protected in accordance Article 26 of lation (EU) 2015/22

				T	ı ı	
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
7 <u>M128</u>	1					
	Apple fruit cell culture biomass	Specified food category	Maximum levels			
		Food supplements as defined in Directive 2002/46/EC for the adult population	0,15 mg/day	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'apple fruit cell culture biomass'. The labelling of food supplements containing the novel food shall bear a statement that they should only be consumed by persons above 18 years of age.		
<u>M69</u>	Arachidonic	Specified food category	Maximum levels	The designation of the novel food		
	Aracildonic acid-rich oil from 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	on the labelling of the foodstuffs containing it shall be 'Oil from Mortierella alpina' or 'Mortierella alpina oil'		
		Foods for special medical purposes for infants as defined in Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013			

Authorised novel f	Conditions under which the n	ovel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Argan oil from Argania spinosa	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
	As seasonings	Not specified	containing it shall be 'Argan oil' and if used as seasoning 'Vegetable oil only for seasoning' shall be		
	Food Supplements as defined in Directive 2002/46/EC	In line with normal food use of vegetable oils	mentioned on the label		
<u>M121</u>					
Astaxanthin-rich oleoresin from	Specified food category	Maximum levels of astaxanthin	The designation of the novel food on the labelling of the foodstuffs		
Haematococcus pluvialis algae	Food supplements as defined in Directive 2002/46/EC excluding infants and young children	2,3 mg astaxanthin per day for children 3 to less than 10 years of age	containing it shall be 'Astaxanthin rich oleoresin from <i>Haematococcus pluvialis</i> algae' The labelling of food supplements containing Astaxanthin rich oleoresin from <i>Haematococcus pluvialis</i> algae shall bear a statement that they should not be consumed: (a) if other food supplements		
		5,7 mg astaxanthin per day for adolescents 10 to less than 14 years of age			
		8 mg astaxanthin per day for general population older than 14 years of age	under 3 years of age (c) by infants and children under 10 years of age (12)		
			(d) by infants, children and adolescents under 14 years of age (12).		

V 1V17						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
▼ <u>M12</u>	_					
	Partially hydrolysed protein from spent	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		Authorised on 10 January 2024. This inclusion is
	barley (Hordeum vulgare) and rice	Fried or extruded cereal, seed or root-based products	5 g/100 g	containing it shall be 'Partially hydrolysed protein from spent		based on proprietary scientific evidence and
	(Oryza sativa)	Confectionery including chocolate	5 g/100 g	barley and rice'.		scientific data protected in accordance with Article 26
		Breakfast cereals	5 g/100 g	In accordance with Article 21 to		of Regulation (EU) 2015/
		Pastas and rice (or other cereal)-based dishes	8 g/100 g	Regulation (EU) No 1169/2011.		2283.
		Soups (dry mixture)	50 g/100 g			Applicant: Evergrain LLC, 3205 S. 9th St, St. Louis,
		Soups (ready-to-eat)	5 g/100 g			Missouri, 63118 USA. During the period of data
		Sauces	10 g/100 g			protection, the novel food
		Dried sauce preparation	50 g/100 g			partially hydrolysed protein from spent barley (<i>Hordeum</i>
		Meat analogues	15 g/100 g			vulgare) and rice (Oryza sativa) is authorised for placing on the market within the Union only by Evergrain 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
		Cereal bars	30 g/100 g			
		Butter and margarine/oil blends	10 g/100 g			
		Milk analogues based ice creams	10 g/100 g			
		Milk analogues	5 g/100 ml			
		Nut/seeds paste/emulsion	15 g/100 g			
		Energy drinks	8 g/100 ml			
		Soft drinks marketed in relation to physical exercise	5 g/100 ml			Article 26 of Regulation (EU) 2015/2283 or with the agreement of
		Cola type drinks	5 g/100 g			Evergrain LLC.
		Powdered drink bases	90 g/100 g	7		End date of the date
		Beverages based on fruit and/or vegetable juices	5 g/100 ml			protection: 10 January 2029.
		Cream, cheese and yoghurt (non-soy) analogues	10 g/100 g			
		Hummus	10 g/100 g	7		
		Alcohol-free beer	5 g/100 ml			
		Meal replacement for weight control	30 g/100 g	7		

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Basil seeds (Ocimum basilicum)	Specified food category	Maximum levels			
	Fruit juice and fruit/vegetable blend beverages	3 g/200 ml for addition of whole basil seeds (<i>Ocimum basilicum</i>)			
<u>4</u>					
Beta-glucan from Euglena gracilis	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		Authorised on 30 A 2024. This inclusion
microalgae	Cereal bars	670 mg/100 g	containing it shall be 'beta-glucan from Euglena gracilis microalgae'.		based on proprie scientific evidence
	Total diet replacement for weight control as defined in Regulation (EU) No 609/2013	600 mg/day			scientific data protected accordance with Article 2 of Regulation (EU) 201 2283.
	Food supplements as defined in Directive 2002/46/EC, excluding food supplements for infants and young children	100 mg/day for children from 3 to 9 years of age 150 mg/day for children from 10 to 17 years of age 200 mg/day for adults	 The designation of the novel food on the labelling of the foodstuffs containing it shall be 'beta-glucan from Euglena gracilis microalgae'. The labelling of food supplements containing the novel food shall bear a statement that they should only be consumed by persons above 3 years of age/above 9 years of age/adults, depending on the age group the product is intended for 		Applicant: Kemin For L.C., 1900 Scott Ave Des Moines, IA 500 United States. During period of data protect the novel food beta-glu from Euglena gramicroalgae is authorised placing on the ma within the Union only Kemin Foods L.C., unlessubsequent applicant obt authorisation for the note of the proprietary scient evidence or scientific protected in accordance Article 26 of Rollation (EU) 2015/2283 with the agreement Kemin Foods L.C. End date of the protection 30 April 2029

food Conditions under which	the novel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Specified food category	Maximum levels (⁷)	The designation of the novel food on the labelling of the foodstuffs		Authorised on 22 Aug 2019. This inclusion
Drink powders, isotonic and energy of intended for sportsmen	drinks 60 mg/100 g	containing it shall be 'betaine'. The labelling of foods containing betaine shall bear a statement that the foods should not be used if		based on proprietary so tific evidence and scient data protected in accorda with Article 26 of Re
Protein and cereal bars intended sportsmen	for 500 mg/100 g	food supplements containing betaine are consumed the same day.		lation (EU) 2015/2283. Applicant: DuPont Nutri Biosciences ApS, Lar brogade 1 Copenhagen
Meal replacements intended for sportsn	en 20 mg/100 g		the the transfer of the transf	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 obtains authorisation for the novel food without reference to
				the proprietary scienevidence or scientific protected in accordance Article 26 of Regulation (2015/2283 or with agreement of Du Nutrition Biosciences Ap End date of the protection: 22 August 2
el	Specified food category Drink powders, isotonic and energy of intended for sportsmen Protein and cereal bars intended sportsmen Meal replacements intended for sportsm Total diet replacement for weight contradefined under Regulation (EU) No 609 Foods for Special Medical Purpose defined under Regulation (EU) No 609	Drink powders, isotonic and energy drinks intended for sportsmen Protein and cereal bars intended for sportsmen Meal replacements intended for sportsmen Total diet replacement for weight control as defined under Regulation (EU) No 609/2013 Foods for Special Medical Purposes as defined under Regulation (EU) No 609/2013 Maximum levels (7) 60 mg/100 g 500 mg/100 g 500 mg/100 g (bar) 136 mg/100 g (soup) 188 mg/100 g (porridge) 60 mg/100 g (beverages)	Specified food category Maximum levels (7) Drink powders, isotonic and energy drinks intended for sportsmen Protein and cereal bars intended for sportsmen Meal replacements intended for sportsmen The designation of the novel food on the labelling of the foodstuffs containing it shall be 'betaine'. The labelling of foods containing betaine shall bear a statement that the foods should not be used if food supplements containing betaine are consumed the same day. Meal replacements intended for sportsmen 20 mg/100 g 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) Foods for Special Medical Purposes as defined under Regulation (EU) No 609/2013	Specified food category Maximum levels (7) Drink powders, isotonic and energy drinks intended for sportsmen Protein and cereal bars intended for sportsmen Meal replacements intended for sportsmen Total diet replacement for weight control as defined under Regulation (EU) No 609/2013 Foods for Special Medical Purposes as defined under Regulation (EU) No 609/2013 The designation of the novel food on the labelling of the foodstuffs containing it shall be "betaine". The labelling of the foodstuffs containing betaine shall bear a statement that the foods should not be used if food supplements containing betaine are consumed the same day. Soo mg/100 g Total diet replacement for weight control as defined under Regulation (EU) No 609/2013 Total diet replacement for weight control as defined under Regulation (EU) No 609/2013 The designation of the novel food on the labelling of the foodstuffs containing it shall be "betaine". The designation of the novel food on the labelling of the foodstuffs containing the shall be "betaine". The designation of the novel food on the labelling of the foodstuffs containing the shall be "betaine". The designation of the novel food on the labelling of the foodstuffs containing the shall be "betaine". The designation of the novel food on the labelling of the foodstuffs containing betaine shall bear a statement that the foods should not be used if food supplements containing betaine shall bear a statement that the foods should not be used if food supplements containing betaine shall bear a statement that the foods should not be used if food supplements containing betaine shall bear a statement that the foods should not be used if food supplements containing betaine shall bear a statement that the foods should not be used if food supplements containing betaine shall bear a statement that the foods should not be used if food supplements containing betaine shall bear a statement that the foods should not be used if food supplements containing betaine shall bear a statement that

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Fermented black bean extract	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Fermented black bean (Soya) extract' or		
	Food Supplements as defined in Directive 2002/46/EC	4,5 g/day	'Fermented Soya extract'		
Bovine lactoferrin	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Lactoferrin from cows' milk'		
	Infant formula and follow-on formula as defined in Regulation (EU) No 609/2013 (ready to drink)	100 mg/100 ml			
	Foods on dairy basis intended for young children (ready to eat/drink)	200 mg/100 g			

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Processed cereal food (solid)	670 mg/100 g			
	Foods for special medical purposes as defined in Regulation (EU) No 609/2013	Depending on the needs of 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			

	Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
M35 M36		Specified food category Infant formulae as defined in Regulation (EU) No 609/2013	Maximum levels 30 mg/100 g (powder) 3,9 mg/100 mL (reconstituted)	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Milk whey		Authorised on 20 Novembe 2018. This inclusion is base on proprietary scientifications.
		Follow-on formulae as defined in Regulation (EU) No 609/2013 Total diet replacement foods for weight control as defined in Regulation (EU) No 609/2013 Foods for special medical purposes as defined in Regulation (EU) No 609/2013 Foods for special medical purposes as defined in Regulation (EU) No 609/2013 and a sum of the second of th	30 mg/100 g (powder) 4,2 mg/100 mL (reconstituted) 300 mg/day	(*) Depending on the age group the food supplement is intended for.	pr A 20 Pr de Sa Fr da fo w or m or S. ap at w pr ev pr A 20 ag Pr	evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Armon 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 Unior 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 of Armon Protéines S.A.S. End date of the data protection 20 November 2023.
			feeding) 30 mg/100 g (powder formula for			
			infants when appropriate complementary feeding is introduced) 4,2 mg/100 mL (reconstituted formula for infants when appropriate complementary feeding is introduced) 58 mg/day for young children 380 mg/day for children and			
		Food supplements as defined in Directive 2002/46/EC	adolescents from 3 to 18 years of age 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			

02017R2470 - EN - 01.05.2024 - 046.001 - 27

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
▼ <u>M96</u>						
	Bovine milk beta-lactoglobulin	Specified food category	Maximum levels (g NF/100 ml)	The designation of the novel food on the labelling of the foodstuffs		Authorised on 11 January 2023. This inclusion is
	(β-lactoglobulin)	Soft drinks marketed in relation to physical exercise	25	containing it shall be 'bovine milk beta-lactoglobulin' or 'bovine milk β -lactoglobulin'.	based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/	
		Whey powder (reconstituted)	8			2283. Applicant: Arla Foods
		Milk based drinks and similar products	12			Ingredients Group P/S, Sønderhøj 10-12, 8260 Viby J, Denmark. During the
		Foods for special medical purposes as defined in Regulation (EU) No 609/2013 intended for the general population older than 3 years of age, excluding pregnant and lactating women	In accordance with the particular nutritional requirements of the persons for whom the products are intended			period of data protection, the novel food beta-lactoglobulin (β-lactoglobulin) is authorised for placing on the market within the Union only by Arla Foods Ingredients Group P/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 Arla Foods Ingredients Group P/S. End date of the data protection: 11 January 2028.

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
<u>7</u>					
Bovine milk osteo- pontin	Specified food category Maximum levels The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Bovine milk Osteopontin'. Specified food category Maximum levels The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Bovine milk Osteopontin'.		Authorised on 26 M 2023. This inclusion based on propri scientific evidence scientific data protecte accordance with Article of Regulation (EU) 2 2283.		
	Follow-on formula as defined in Regulation (EU) No 609/2013 (13)		Application Ingredition Sønder J Desperiod novel for ponting placing the United Ingredition unless obtains novel for to the evidence protect. Article lation with the special special service of the ser	Applicant: Arla F Ingredients Group Sønderhøj 10-12 8260 J Denmark. During period of data protection	
	Milk-based drinks intended for young children	151 mg/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			novel food Bovine milk of pontin is authorised placing on the market with the Union only by Arla F. Ingredients Group unless a subsequent apply obtains authorisation for novel food without refet to the proprietary scie evidence or scientific protected in accordance Article 26 of F. lation (EU) 2015/228; with the agreement of Foods Ingredients Group
					End date of the protection: 26 March 200

Au	uthorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	glossoides vensis seed oil	Specified food category	Maximum levels of stearidonic acid (STA)	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Refined		
		Dairy products and analogues	250 mg/100 g	Buglossoides oil'		
			75 mg/100 g for drinks			
		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 purposes as defined in Regulation (EU) No 609/2013, excluding foods for special medical purposes intended for infants and young children	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			
91						
	lanus	Specified food category	Maximum levels	1. The designation of the novel food		
jini	finmarchicus oil	Food supplements as defined in Directive 2002/46/EC, excluding food supplements for infants and young children	1,0 g/day (< 0,1 % astaxanthin esters, resulting in < 1,0 mg astaxanthin per day) for the general population, excluding infants and young children 2,3 g/day (from 0,1 % to \leq 0,25 % astaxanthin esters, resulting in \leq 5,75 mg astaxanthin per day) for the general population older than 14 years of age	on the labelling of the foodstuffs containing it shall be 'oil from Calanus finmarchicus (crustacean)'. The labelling of food supplements containing Calanus finmarchicus oil shall bear a statement that those food supplements should not be consumed:		

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
				 a) if other food supplements containing astaxanthin esters are consumed on the same day. b) by infants and children younger than 3 years. c) by children younger than 14 years, if the ingredient contains ≥ 0,1 % astaxanthin. 		
▼ <u>M77</u>	Calcium fructoborate	Specified food category Food supplements as defined in Directive	Maximum levels 220 mg/day	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'calcium'		Authorised on 23 December 2021. This inclusion is based on proprietary scientific
		2002/46/EC for the adult population, excluding food supplements for pregnant and lactating women		fructoborate'. 2. The labelling of food supplements containing calcium fructoborate shall bear a statement that those food supplements should not be consumed by population under 18 years of age and by pregnant and lactating women.		evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: VDF Future-Ceuticals, Inc., 300 West 6th Street Momence, Illinois 60954, the United States. During the period of data protection, the novel food calcium fructoborate is authorised for placing on the market within the Union only by VDF Future-Ceuticals, 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 VDF Future-Ceuticals, Inc. End date of the date protection: 23 December 2026

	Authorised novel food	Conditions under which the no	val food may be used	Additional specific labelling requirements	Other requirements	►M30 Data Protection ◀
	Authorised novel lood	Conditions under which the no	ver root may be used	Additional specific labelling requirements	Other requirements	MISO Data Protection
▼ <u>M85</u>						
	Calcium L-Methyl- folate	Specified food category	Maximum levels (expressed as folic acid)	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Calcium L-Methylfolate'.		
		Foods for special medical purposes and total diet replacement for weight control as defined in Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013	2 Mean monaic .		
		Infant formulae and follow-on formula as defined by 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 by Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013			
		Food supplements as defined in Directive 2002/46/EC	In accordance with Directive 2002/ 46/EC			
		Food fortified in accordance with Regulation (EC) No 1925/2006	In accordance with Regulation (EC) No 1925/2006			
▼ <u>M137</u>						
	Calcidiol monohydrate	Specified food category	Maximum levels	1. The designation of the novel food on the labelling of the foodstuffs containing it shall be "calcidial		Authorised on 1 May 2024. This inclusion is based on proprietary scientific
		Food supplements as defined in Directive 2002/46/EC, excluding food supplements for infants and young children	10 μg/day for children from 11 years of age and adults 5 μg/day for children from 3 to 10 years of age	containing it shall be "calcidiol (calcifediol) monohydrate (vitamin D)".		evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283.

▼<u>M137</u>

Authorised novel food	Conditions under which the novel food may be used		Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
			2. The labelling of food supplements containing the novel food shall bear a statement that they should not be consumed by infants and children under 3 years of age/children under 11 years of age, depending on the age group the product is intended for.		Applicant: DSM Nutritional Products Ltd., Wurmisweg 576, 4303 Kaiseraugst, Switzerland. During the period of data protection, the novel food calcidiol monohydrate is authorised for placing on the market within the Union only by DSM Nutritional Products 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 DSM Nutritional Products Ltd. End date of the date protection: 1 May 2029.
Dried nuts of Canarium ovatum Engl.	Specified food category Not specified	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'nuts of <i>Canarium ovatum</i> ' and/or 'pilinuts' and/or 'pili (<i>Canarium ovatum</i>) nuts'.		

▼<u>M106</u>

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
			2. The labelling of the foodstuffs containing dried nuts of Canarium ovatum Engl. shall bear a statement that the dried nuts of Canarium ovatum Engl. may cause allergic reactions to consumers with known allergies to cashew and walnut. This statement shall appear in close proximity to the list of ingredients or, in the absence of a list of ingredients, in close proximity to the name of the food.		
Canarium indicum L. dried nuts (Kenari) (Traditional food from a third country)	Specified food category Not specified	Maximum levels (g/100 g)	1. The designation of the traditional food on the labelling of the foodstuffs containing it shall be 'dried kenari (Canarium indicum) nuts'. 2. The labelling of the foodstuffs containing dried nuts of Canarium indicum L. shall bear a statement that the nuts may cause allergic reactions to consumers with known allergies to hazel, cashew and pistachio. This statement shall appear in close proximity to the list of ingredients or, in the absence of a list of ingredient, in close proximity to the name of the food.		

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
4					
Cellobiose	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		Authorised on 1 June 2 This inclusion is based
	Food supplements as defined in Directive 2002/46/EC for the general population, excluding infants and young children	3 g/day	containing it shall be 'cellobiose'. 2. The labelling of food supplements containing cellobiose shall bear a statement		proprietary scie. evidence and scientific protected in accordance Article 26 of R lation (EU) 2015/2283.
	Dried, canned-tinned, raw cured (or seasoned), cooked cured (or seasoned) meat	2 g/100 g	that those food supplements should not be consumed by infants and young children.		Applicant: SAVA Ingredients GmbH, Dü Straße 67, 50189 Els
	Fresh raw, preserved or partly preserved sausages	2 g/100 g			Germany. During the p of data protection, the food cellobiose is author for placing on the m
	Meat based spreadable-textured specialties	2 g/100 g			within the Union only SAVANNA Ingred GmbH, unless a subse
	Liver based spreadable-textured specialties	2 g/100 g			applicant obtains auth ation for the novel without reference to
	Savoury sauce dry preparation	40 g/100 g			proprietary scient evidence or scientific
	Table-top sweeteners in powder form	60 g/100 g			protected in accordance Article 26 of I lation (EU) 2015/228
	Table-top sweeteners in tablets	60 g/100 g			with the agreement SAVANNA Ingred GmbH.
					End date of the protection: 1 June 2028

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Authorised novel food 82 Cetylated fatty acids	Specified food category Food supplements as defined in Directive 2002/46/EC for the adult population	Maximum levels	1. The designation of the novel food on the labelling of the food supplements containing it shall be 'cetylated fatty acids preparation'. 2. The labelling of food supplements containing the novel food shall bear a statement that those food supplements should not be consumed by persons under 18 years of age.	Other requirements	Authorised on 3 Mar 2022. This inclusion based on propriets scientific evidence a scientific data protected accordance with Article of Regulation (EU) 201 2283. Applicant: Pharmanu S.p.A., Via Delle Len 216/b, 56122 Pisa, Ita During the period of da protection, the novel fo cetylated fatty acids is autonated to the sum of the component of the science of the scien
					orised for placing on market within the Un only by Pharmanutra S.p. unless a subsequent applic obtains authorisation for novel food without refere to the proprietary scient evidence or scientific d protected in accordance w Article 26 of Regulation (F 2015/2283 or with agreement of Pharmanu S.p.A. End date of the d protection: 3 March 2027

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Chewing gum base (monomethoxypoly- ethylene glycol)	Specified food category Chewing gum	Maximum levels 8 %	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Gum base (including 1,3-butadiene, 2-methylhomopolymer, maleated, esters with polyethylene glycol mono-Me ether)' or 'Gum base (including CAS No: 1246080-53-4)'		
Chewing gum base (Methyl vinyl ether-maleic anhydride copolymer)	Specified food category Chewing gum	Maximum levels 2 %	The designation of the novel 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 <i>Salvia</i> hispanica	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Chia oil (Salvia hispanica)'		
	Fats and oils Pure chia oil	10 % 2 g/day	(Sarra Inspanca)		
	Food Supplements as defined in Directive 2002/46/EC	2 g/day			

WIJ						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
<u>M64</u>						
	Chia seeds (Salvia hispanica)	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
	mspunicu)	Bread products	5 % (whole or ground chia seeds)	containing it shall be 'Chia seeds (Salvia hispanica)'		
		Baked products	10 % whole chia seeds			
		Breakfast cereals	10 % whole chia seeds			
		Sterilised ready to eat meals based on cereal grains, pseudocereal grains and/or pulses	5 % whole chia seeds			
		Fruit, nut and seed mixes				
		Chia seeds 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 twin pot)				
		Non-alcoholic beverages (including 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				

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► M30 Data Protection
Chitin-glucan from Aspergillus niger	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
Aspergulus niger	Food Supplements as defined in Directive 2002/46/EC	5 g/day	containing it shall be 'Chitin-glucan from Aspergillus niger'		
Chitin-glucan	Specified food category	Maximum levels	The designation of the novel food		
complex from Fomes fomentarius	Food Supplements as defined in Directive 2002/46/EC	5 g/day	on the labelling of the foodstuffs containing it shall be 'Chitin-glucan from Fomes fomentarius'		
Chitosan extract	Specified food category	Maximum levels	The designation of the novel food on		
from fungi (Agaricus bisporus; Aspergillus niger)	Food Supplements as defined in Directive 2002/46/EC	In line with normal use in food supplements of chitosan from crustaceans	the labelling of the foodstuffs containing it shall be 'Chitosan extract from <i>Agaricus bisporus</i> ' or 'Chitosan extract from <i>Aspergillus niger</i> '		
Chondroitin sulphate	Specified food category	Maximum levels	The designation of the novel food		
	Food supplements as defined in Directive 2002/46/EC for adult population, excluding pregnant and lactating women	1 200 mg/day	on the labelling of the foodstuffs containing it shall be 'Chondroitin sulphate derived from microbial fermentation and sulphation'		
Chromium	Specified food category	Maximum levels of total chromium	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Chromium Picolinate'		
Picolinate	Foods covered by Regulation (EU) No 609/2013	0/ 250 μg/day c			
	Foods fortified in accordance with Regulation (EC) No 1925/2006 (4)				
Chromium-	Specified food category	Maximum levels	The designation of the novel food on		
containing yeast (Yarrowia lipolytica) biomass	Food supplements as defined in Directive 2002/46/EC, excluding food supplements for infants and young children	2 g/day for children from 3 to 9 years of age, resulting in 46 μg of chromium per day	the labelling of the foodstuffs containing it shall be 'chromium-containing yeast (<i>Yarrowia lipolytica</i>) biomass'		
		4 g/day for children from 10 years of age, adolescents and adults, resulting in 92 μg of chromium per day	The labelling of food supplements containing chromium-containing yeast (<i>Yarrowia lipolytica</i>) biomass shall bear a statement that the food supplements should not be consumed by infants and young children (children under 3 years of age)/children from 3 to 9 years of age (12).		

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Cistus incanus L. Pandalis herb	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Cistus		
	Herbal infusions	Intended daily intake: 3 g herbs/day (2 cups/day)	incanus L. Pandalis herb'		
Citicoline	Specified food category	Maximum levels	 The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Citicoline' The labelling of foods containing citicoline shall bear a statement that the product is not intended to be consumed by children 		
	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	250 mg per serving and a maximum daily consumption level of 1 000 mg			
Clostridium butyricum	Specified food category	Maximum levels	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)'		
	Food Supplements as defined in Directive 2002/46/EC	1,35 × 10 ⁸ CFU/day			

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Coffea arabica L. and/or Coffea	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
canephora Pierre ex A.Froehner dried cherry pulp and its infusion (Traditional food from a third country)	Coffee cherry pulp from <i>Coffea arabica</i> L. and/or <i>Coffea canephora</i> Pierre ex A.Froehner for the preparation of infusions		containing it shall be 'coffee cherry pulp' and/or 'cascara (coffee cherry pulp)', and/or 'coffee cherry pulp infusion' and/or 'coffee cherry pulp dried infusion'. If the product containing the novel food contains more than 150 mg/l of caffeine (as such or after reconstitution), it shall be labelled with the following indication: 'High caffeine content. Not recommended for children or pregnant or breast-feeding women' in the same field of vision as the name of the food,		
	Coffee, coffee and chicory extracts, instant coffee, tea, herbal- and fruit-infusions, coffee substitutes, coffee mixes and instant mixes for hot beverages (and their flavoured counterparts).				
	Flavoured and unflavoured non-alcoholic ready-to-drink beverages		followed by the caffeine content expressed in mg per 100 ml. Typical infusion preparations are prepared with up to 6 g of coffee cherry pulp per 100 ml of hot water (> 75 °C). For the coffee cherry pulp placed on the market as such for the preparation of infusions, instructions shall be given to the consumer on the preparation.		

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
<u>M30</u>						
	D-ribose	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		Authorised on 16 April 2019. This inclusion is
		Cereal bars	0,20 g/100 g	containing it shall be 'D-ribose'.		based on proprietary scientific evidence and
		Fine bakery wares	0,31 g/100 g	The labelling of foods containing D-ribose shall bear a statement that		scientific data protected in accordance with Article 26
		Chocolate confectionery (excluding chocolate bars)	0,17 g/100 g	the foods should not be used if food supplements containing D-ribose are consumed the same day.		of Regulation (EU) 2015/2283.
		Milk-based drinks (excluding malts and shakes)	0,08 g/100 g			Applicant: Bioenergy Life Science, Inc., 13840 Johnson St. NE,
		Drinks intended to meet the expenditure of intense muscular effort especially for sportsmen, isotonic and energy drinks			Minneapolis, Minnesota, 55304, USA. During the period of data protection, the novel food D-ribose is authorised for placing on	
		Bars intended to meet the expenditure of intense muscular effort especially for sportsmen	3,3 g/100 g			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 26 of Regulation (EU) 2015/2283 or with the
		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			
		Tea and infusions (in powder form to be reconstituted)	0,23 g/100 g			agreement of Bioenergy Life Science, Inc.
						End date of the data protection: 16 April 2024 (5 years).

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Extract of defatted cocoa powder	Specified food category	Maximum levels	Consumers shall be instructed not to consume more than 600 mg polyphenols corresponding to 1,1 g of		
	Nutrition bars	1 g/day and 300 mg polyphenols corresponding to not more than 550 mg of extract of defatted	extract of defatted cocoa powder per day		
	Milk based beverages cocoa powder in one portion of food (or food supplement)				
	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				
Low fat cocoa extract	Specified food category	Maximum levels	Consumers shall be instructed not to consume more than 600 mg of cocoa flavanols per day		
	Foods including food supplements as defined in Directive 2002/46/EC	730 mg per serving and around 1,2 g/day			
Coriander seed oil from <i>Coriandrum</i> sativum	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Coriander		
	Food Supplements as defined in Directive 2002/46/EC	600 mg/day	seed oil'		

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Cranberry extract powder	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		Authorised on 20 November 2018. This inclusion is base
	Food Supplements as defined in Directive 2002/46/EC for the adult population	350 mg/day	containing it shall be 'cranberry extract powder'		on proprietary scientific evidence and scientific dat protected in accordance wit 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 dat protection the novel food cranberry extract powder, authorised for placing of the market within the Unio only by Ocean Spra Cranberries Inc. unless subsequent applicant obtain authorisation for the nove food without reference to the proprietary scientific evidence or scientific dat protected in accordance with Article 26 of Regulation (EU 2015/2283 or with the agreement of Ocean Spra Cranberries Inc.
					End date of the dar protection: 20 November 2023.

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Crataegus pinna- tifida dried fruit	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
		Herbal infusions	In line with normal food use of Crataegus laevigata	containing it shall be 'Crataegus pinnatifida dried fruit'		
		Jams and jellies in accordance with Directive 2001/113/EC (5)				
		Compotes				
	α-cyclodextrin	Not specified		The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Alpha-cyclodextrin' or 'α-cyclodextrin'		
	γ-cyclodextrin	Not specified		The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Gamma-Cyclodextrin' or 'γ-Cyclodextrin'		
▼ <u>M22</u>	Decorticated grains of <i>Digitaria exilis</i> (Kippist) Stapf	Not specified		The designation of the novel food on the labelling of the foodstuffs containing it shall be 'decorticated fonio (<i>Digitaria exilis</i>) grains'		
	(Traditional food from a third country)			Tome (Digital tales) grams		
▼ <u>M9</u>						
	Dextran preparation produced	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
	by Leuconostoc mesenteroides	Bakery products	5 %	containing it shall be 'Dextran'		

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Diacylglycerol oil of plant origin	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
piant origin	Cooking oils		containing it shall be 'Diacylglycerol oil of plant origin (at least 80 %		
	Fat spreads		diacylglycerols)'		
	Salad dressings				
	Mayonnaise				
	Meal replacement for weight control (as drinks)				
	Bakery products				
	Yoghurt type products				
Dihydrocapsiate (DHC)	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
	Cereal bars	9 mg/100 g 9 mg/100 g	containing it shall be 'Dihydro-capsiate'		
	Biscuits, cookies and crackers		2. Food supplements containing		
	Rice based snacks	12 mg/100 g	synthetic dihydrocapsiate will be labelled as 'not intended for		
	Carbonated drinks, dilutable drinks, fruit juice based beverages	1,5 mg/100 ml	children up to 4.5 years'		
	Vegetable drinks	2 mg/100 ml			
	Coffee based drinks, tea based drinks	1,5 mg/100 ml			
	Flavoured water — still	1 mg/100 ml			
	Precooked oatmeal cereal	2,5 mg/100 g			
	Other cereals	4,5 mg/100 g			
	Ice cream, dairy desserts	4 mg/100 g			
	Pudding mixes (ready to eat)	2 mg/100 g			

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	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			
M54 Dried Euglena	Specified food category	Maximum levels	The designation of the novel food		Authorised on 23 December
gracilis	Breakfast cereal bars, granola bars and protein bars	630 mg/100 g	on the labelling of the foodstuffs containing it shall be 'dried		2020. This inclusion is based on proprietary scientific
	Yoghurt	150 mg/100 g	biomass of Euglena gracilis algae'.		evidence and scientific data protected in accordance with
	Yoghurt Beverages	95 mg/100 g	The labelling of food supplements		Article 26 of Regulation (EU)
	Fruit and vegetable juices, nectars, fruit/vegetable blend beverages	120 mg/100 g	containing dried Euglena gracilis shall bear a statement that those food supplements should not be		2015/2283. Applicant: Kemin Foods
	Fruit-Flavoured Drinks	40 mg/100 g	consumed by infants/children under		L.C., 2100 Maury Street
	Meal replacement beverages	75 mg/100 g	3 years of age/children under 10		Des Moines, IA 50317,
	Food supplements as defined in Directive	100 mg/day for young children	years of age/children and adolescents under 18 years of		USA.
	2002/46/EC, excluding food supplements for infants	150 mg/day for children from 3 to 9 years of age	age (12).		
		225 mg/day for children from 10 years of age and adolescents (to 17 years of age)			
		375 mg/day for adults			

▼<u>M54</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Total diet replacement for weight control as defined by Regulation (EU) No 609/2013	190 mg/meal			During the period of data protection, the novel food is authorised for placing on the market within the Union only by Kemin Foods L.C. 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 Kemin Foods L.C. End date of the data protection: 23 December
Dried aerial parts of Hoodia parviflora	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'dried aerial		Authorised on 3 September 2018. This inclusion is based on proprietary scientific
	Food Supplements as defined in Directive 2002/46/EC for adult population	9,4 mg/day	parts of <i>Hoodia parviflora</i> '		evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Desert Labs, Ltd Kibbutz Yotvata, 88820 Israel.

▼<u>M13</u>

	Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
						During the period of data protection the novel food dried aerial parts of <i>Hoodia parviflora</i> 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.
▼ <u>M9</u>	Dried extract of Lippia citriodora from cell cultures	Specified food category Food Supplements as defined in Directive 2002/46/EC	Maximum levels 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 angus- tifolia extract from	Specified food category	Maximum levels			
	cell cultures	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 <i>Echinacea angustifolia</i>			

<u>M9</u>						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
<u>M32</u>	Echinacea purpurea extract from cell cultures	Specified food category Food Supplements as defined in Directive 2002/46/EC	Maximum levels 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-PC TM '		
<u> 19</u>	Echium plan- tagineum oil	Specified food category	Maximum levels of stearidonic acid (STA)	The designation of the novel food on the labelling of the foodstuffs		
		Milk-based products and drinkable yoghurt products delivered in a single dose	250 mg/100 g; 75 mg/100 g for drinks	containing it shall be 'Refined echium oil'		
		Cheese preparations	750 mg/100 g			
		Spreadable fat and dressings	750 mg/100 g			
		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			

▼<u>M52</u>

_						
	Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	<i>Ecklonia cava</i> phlo- rotannins	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Ecklonia cava Phlorotannins'.		
		Food supplements as defined in Directive 2002/46/EC intended for the general population, excluding children under the age of 12 years	163 mg/day for adolescents from 12 to 14 years of age 230 mg/day for adolescents above	Ecklonia cava phlorotannins shall bear the following statement:		
			14 years of age 263 mg/day for adults	(a) This food supplement should not be consumed by children/adolescents under the age of twelve/fourteen/eighteen(*) years.		
				(b) 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 thyroid disease.		
				(c) This food supplement should not be consumed if other food supplements containing iodine are also consumed.		
				(*) Depending on the age group the food supplement is intended for.		

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
▼ <u>M18</u>						
	Egg membrane hydrolysate	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'egg membrane hydrolysate'.		Authorised on 25 Novembe 2018. This inclusion is based on proprietary scientific evidence and scientific data
		Food Supplements as defined in Directive 2002/46/EC intended for the general adult population	450 mg/day	memorane nyaronysate .		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

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Epigallocatechin gallate as a purified extract from green	Specified food category	Maximum levels	The labelling shall bear a statement that consumers should not consume more than 300 mg of extract per day		
	tea leaves (Camellia sinensis)	Foods including food supplements as defined in Directive 2002/46/EC	150 mg of extract in one portion of food or food supplement			
▼ <u>M52</u>						
	L-ergothioneine	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
		Alcohol-free beverages	0,025 g/kg	containing it shall be 'L-ergo-		
		Milk-based drinks	0,025 g/kg	thioneine'		
		'Fresh' milk products(*)	0,040 g/kg			
		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-ergothioneir any milk constituent	ne may not replace in whole or in part,			
▼ <u>M108</u>	3					
	Roasted and popped kernels	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
	from the seeds of Euryale ferox Salisb. (makhana) (Traditional food from a third country)	Processed nuts		containing it shall be 'roasted seeds of Euryale ferox' or 'makhana (Euryale ferox) roasted seeds'		

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
<u>M52</u>						
	Extract of three herbal roots	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
	(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	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.		
<u>M9</u>						
	Ferric Sodium EDTA	Specified food category	Maximum levels (expressed as anhydrous EDTA)	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Ferric		
		Food supplements as defined in Directive 2002/46/EC	18 mg/day for children	Sodium EDTA'		
			75 mg/day for adults			
		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 category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
	phosphate	Food supplements as defined in Directive 2002/46/EC	To be used in compliance with Directive 2002/46/EC, Regulation (EU) No 609/2013 and/or Regu-	containing it shall be 'Ferrous ammonium phosphate'		
		Foods covered by Regulation (EU) No 609/2013	lation (EC) No 1925/2006			
		Foods fortified in accordance with Regulation (EC) No 1925/2006				

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Fish peptides from Sardinops sagax	Specified food category	Maximum levels fish peptide product	The designation of the novel food on the labelling of the foodstuffs		
sarainops sagax	Foods based on yoghurt, yoghurt drinks, fermented milk products, and powdered milk	0,48 g/100 g (ready to eat/drink)	(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	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Flavonoids'	Beverages containing flavonoids shall	
	Beverages based on milk	120 mg/day	from Glycyrrhiza glabra L.'	be presented to the final	
	Beverages based on yoghurt			consumer as single portions.	
	Beverages based on fruit or vegetables				
	Food Supplements as defined in Directive 2002/46/EC	120 mg/day	(a) the product should not be consumed by pregnant and breast feeding women,		
	Total diet replacement for weight control as defined in Regulation (EU) No 609/2013	120 mg/day	children and young adolescents; and (b) people taking prescription		
	Foods for special medical purposes as defined in Regulation (EU) No 609/2013	120 mg/day	drugs should only consume the product under medical supervision;		
			(c) a maximum of 120 mg of flavonoids per day should be consumed.		
			3. The amount of flavonoids in the final food shall be indicated on the labelling of the food containing it.		

V 1V19						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Fruit 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 (<i>Theobroma cacao</i> L.) pulp', 'cocoa (<i>Theobroma cacao</i> L.) pulp juice' or 'cocoa (<i>Theobroma cacao</i> L.) concentrated pulp juice' depending on the form used.			
<u>M9</u>	Fucoidan extract from the seaweed Fucus vesiculosus	Specified food category Foods including food supplements as defined in Directive 2002/46/EC for the general	Maximum levels 250 mg/day	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Fucoidan extract from seaweed <i>Fucus vesiculosus</i> '.		
	Fucoidan extract from the seaweed Undaria pinnatifida	Specified food category Foods including food supplements as defined in Directive 2002/46/EC for the general	Maximum levels 250 mg/day	Culosus'. The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Fucoidan extract from seaweed Undaria pinna-		
<u>M117</u>	2'-Fucosyllactose	population Specified food category	Maximum levels	tifida' 1. The designation of the novel food		
		Unflavoured pasteurised and sterilised (including UHT) milk-based products		on the labelling of the foodstuffs containing it shall be '2'-fucosyllactose'.		
		Unflavoured fermented milk-based products	1,2 g/l for beverages	2. The labelling of food supplements containing 2'-fuco-		
			19,2 g/kg for products other than beverages	syllactose shall bear a statement that the supplements should not be used if other foods with added 2'-fucosyllactose are consumed the same day.		
		Flavoured fermented milk-based products including heat-treated products	1,2 g/l for beverages 19,2 g/kg for products other than beverages			

▼<u>M117</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► M30 Data Protection ◀
	Dairy analogues, including beverage whiteners	1,2 g/l for beverages	3. The labelling of food		
		12 g/kg for products other than beverages	supplements containing 2'-fuco- syllactose intended for young children shall bear a statement that the supplements should not		
		400 g/kg for whitener	be used if breast milk or other foods with added 2'-fucosyl- lactose are consumed the same		
	Cereal bars	12 g/kg	e,		
	Table-top sweeteners	200 g/kg			
	Infant 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			
	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			
	Processed cereal-based foods and baby foods for infants and young children as defined under Regulation (EU) No 609/2013	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			
	Milk based drinks and similar products intended for young children	1,2 g/l for milk-based drinks and similar products in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			

▼<u>M117</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Foods 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			
	Total diet replacement for weight control as defined in Regulation (EU) No 609/2013	4,8 g/l for drinks			
		40 g/kg for bars			
	Bread and pasta products bearing statements on the absence or reduced presence of gluten in accordance with the requirements of Im- plementing 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, for the general population, excluding infants	3,0 g/day for general population			
		1,2 g/day for young children			

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
▼ <u>M38</u>						
	2'-Fucosyllactose/ Difucosyllactose	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		Authorised on 19.12.2019. This inclusion is based on
	mixture ('2'-FL/ DFL')	Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products	2,0 g/L	containing it shall be '2'-Fucosyllactose/Difucosyllactose mixture'.		proprietary scientific evidence and scientific data protected in accordance with
	(microbial source)	Unflavoured fermented milk-based products	2,0 g/L (beverages)	The labelling of food supplements containing the 2'-Fucosyllactose/Difucosyllactose mixture shall bear		Article 26 of Regulation (EU) 2015/2283.
			20 g/kg (products other than beverages)	I		Applicant: Glycom A/S, Kogle Allé 4, DK-2970 Hørsholm, Denmark. During
		Flavoured fermented milk-based products including heat-treated products	2,0 g/L (beverages) 20 g/kg (products other than beverages)			the period of data protection, the novel food 2'-Fucosyl- lactose/Difucosyllactose mixture is authorised for placing on the market
		Beverages (flavoured drinks)	2,0 g/L		s a f t t e F F F F F F F F F F F F F F F F F	within the Union only by Glycom A/S, unless a subsequent applicant obtains
		Cereal bars	20 g/kg			authorisation for the novel food without reference to the proprietary scientific
		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	- - r -		evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Glycom A/S.
		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			End date of the data protection: 19.12.2024.
		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			

▼<u>M38</u>

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		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			
▼ <u>M58</u>		Milk-based drinks and similar products intended for young children	1,2 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			
▼ <u>M75</u>	3-Fucosyllactose (3-FL)	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		Authorised on 12 December 2021. This inclusion is based
	(microbial source)	Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products	0,85 g/L	containing it shall be '3-Fucosyllactose'.	or ev	on proprietary scientific evidence and scientific data protected in accordance with
		Unflavoured and flavoured fermented milk- based products including heat-treated products	0,5 g/L (beverages) 5,0 g/kg (products other than beverages)	The labelling of food supplements containing 3-Fucosyllactose (3-FL) shall bear a statement that they should not be consumed: a) if foods containing added 3-		Article 26 of Regulation (EU) 2015/2283.
		Dairy analogues	0,85 g/L (beverages) 8,5 g/kg (products other than beverages)	Fucosyllactose are consumed on the same day; b) by infants and children under 3 years of age.		

▼<u>M75</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Flavoured drinks, energy and sports drinks	1,0 g/L			Applicant: DuPont Nutrition & Biosciences ApS Lange-
	Cereal bars	30,0 g/kg			brogade 1, 1001 Copenhagen K, Denmark. During the
	Infant formula as defined under Regulation (EU) No 609/2013	0,85 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			period of data protection, the novel food 3-Fucosyl- lactose is authorised for placing on the market within the Union only by
	Follow-on formula as defined under Regulation (EU) No 609/2013	0,85 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer		cience subseq author food	DuPont Nutrition & Biosciences ApS, unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific
	Milk-based drinks and similar products intended for young children	0,85 g/L (beverages) in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the
	Processed cereal-based food and baby food for infants and young children as defined under 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		Nutrition & Bio ApS. End date of the	Nutrition & Biosciences ApS. End date of the data protection: 12 December
		3,0 g/kg for products other than beverages			2026.
	Total diet replacement foods for weight control as defined under Regulation (EU)	2,0 g/L (beverages)			
	No 609/2013	30,0 g/kg (products other than beverages)			
	Foods 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, excluding food supplements for infants and young children	5,0 g/day			

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
3-Fucosyllactose ('3-FL')	Specified food category Infant formula as defined under Regulation	Maximum levels 0,90 g/l in the final product ready	The designation of the novel food on the labelling of the foodstuffs containing it shall be '3-fucosyl-		Authorised on 25.1.20 This inclusion is based proprietary scient
(produced by a derivative strain of E. coli BL21(DE3))	(EU) No 609/2013	for use, marketed as such or reconstituted as instructed by the manufacturer facturer for use, marketed as such or reconlactorse'. lactose'. The labelling of food supplements	lactose'. The labelling of food supplements containing 3-Fucosyllactose (3-FL) shall bear a statement that (a) they should not be consumed by children under 3 years of age; (b) they should not be used if other foods containing added 3-Fucosyllactose are consumed on the same day.		evidence and scientific data protected in accordance with Article 26 of Regulation (EU 2015/2283. Applicant: 'Chr. Hansen A.S', Bøge Allé 10-12, 2970 Hørsholm, Denmark. During the period of data protection the novel food 3-Fucosyllactose is authorised for placing on the marker within the Union only by Chr. Hansen A/S unless a subsequent applicant obtains authorisation for the novel food without reference to
	Follow-on formula as defined under Regulation (EU) No 609/2013	1,20 g/l in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			
	Processed cereal-based foods for infants and young children and baby foods for infants and young children as defined under Regulation (EU) No 609/2013	1,20 g/l or 1,20 g/kg 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	1,20 g/l in the final product ready for use, marketed as such or recon- stituted as instructed by the manu- facturer			
	Foods for special medical purposes for infants and young children as defined under 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 0,9 g/l or 0,9 g/kg (if it is intended for infants from 0 until 6 months) and 1,2 g/l or 1,2 g/kg (if it is intended for infants of 6-12 months and/or for young children) in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer.			the proprietary sciene vidence or scientific protected in accordance Article 26 of Regulation 2015/2283 or with agreement of 'Chr. Ha A/S'. End date of the protection: 25.1.2028.
	Foods for special medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants and young children	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, for the general population, excluding infants and young children	3 g/day			

V <u>IVI</u>						
	Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	3-Fucosyllactose ('3-FL') (produced by	Specified food category	Maximum levels (expressed as 3-Fucosyllactose)	The designation of the novel food on the labelling of the foodstuffs containing it shall be '3-Fucosyl-		Authorised on 12 November 2023. This inclusion is based on proprietary scientific
	derivative strain of E. coli K-12 DH1)	Infant formula as defined under Regulation (EU) No 609/2013	1,75 g/L in the final product ready for use, marketed as such or recon- stituted as instructed by the manu- facturer	(b) they should not be used if other foods containing added 3-Fuco-syllactose are consumed on the same day.		evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: 'Glycom A/S', Kogle Allé 4, 2970
		Follow-on formula as defined under Regulation (EU) No 609/2013	1,75 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			Hørsholm, Denmark. During the period of data protection, the novel food 3-Fucosyllactose produced by derivative strain of <i>E. coli</i> K-12 DH1 is authorised for
		Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products	2,0 g/L			placing on the market within the Union 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'. End date of the data protection: 12 November
		Unflavoured fermented milk-based products	2,0 g/L (beverages)			
			4,0 g/kg (products other than beverages)			
		Flavoured fermented milk-based products including heat-treated products	2,0 g/L (beverages)			
			12,0 g/kg (products other than beverages)			2028.
		Cereal bars	25,0 g/kg			

▼<u>M125</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Milk based drinks and similar products	2,0 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			
		12,0 g/kg (products other than beverages)			
	Beverages (flavoured drinks, excluding drinks with a pH less than 5)	1,25 g/L			
	Total diet replacement foods for weight control as defined under Regulation (EU) No 609/2013	2,0 g/L (beverages)			
		25,0 g/kg (products other than beverages)			
	Foods for special medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants and young children	In accordance with the particular nutritional requirements of the persons for whom the products are intended but in any case not higher 4,0 g/L or 4,0 g/kg in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer.			
	Food Supplements as defined in Directive 2002/46/EC, for the general population, excluding infants and young children	4,0 g/day			

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection
Galacto-oligos- accharide	Specified food category	Maximum levels (expressed as ratio kg galacto-oligosaccharide/kg final food)			
	Food supplements as defined in Directive 2002/46/EC	0,333			
	Food supplements as defined in Directive 2002/46/EC, excluding infants and young children	0,450 (corresponding to 5,4 g galacto-oligosaccharide/serving; maximum 3 servings/day up to a maximum of 16,2 g/day)			
	Foods for special medical purposes as defined by Regulation (EU) No 609/2013, excluding infants and young children	In accordance with the particular nutritional requirements of the persons for whom the products are intended but not more than 0,128 (corresponding to a maximum of 8,25 g galacto-oligosaccharide/day)			
	Milk	0,020			
	Milk drinks	0,030			
	Meal replacement for weight control (as drinks)	0,020			
	Dairy analogue drinks	0,020			
	Yoghurt	0,033			
	Dairy based desserts	0,043			
	Frozen dairy desserts	0,043			
	Fruit drinks and energy drinks	0,021			
	Infant meal replacement drinks	0,012			
	Baby juice	0,025			
	Baby yogurt drink	0,024			

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		Baby dessert	0,027			
		Baby snack	0,143			
		Baby cereals	0,027			
		Drinks intended to meet the expenditure of intense muscular effort especially for sportsmen	0,013			
		Juice	0,021			
		Fruit pie filling	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			
		Dairy confectionery	0,05			
		Cheese and processed cheese	0,1			
		Butter and spreadable fats	0,1			
9						
	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 fish			
		Foods covered by Regulation (EU) No 609/2013				
		Meal replacement for weight control				

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	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				
Glucosamine	Specified food category	Maximum levels			
sulphate KCl	Food Supplements as defined in Directive 2002/46/EC	In line with normal food use of glucosamine from shell fish			
Glucosamine	Transfer de la constant de la consta				
sulphate NaCl	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 designation of the novel food on the labelling of the foodstuffs containing it shall be 'Guar Gum'. A specific mention of the possible risks of digestive discomfort linked to the exposure of children aged under		
	Fresh dairy products such as yogurts, fermented milks, fresh cheeses and other dairy-based desserts.	1,5 g/100 g			
	Fruit or vegetable-based liquid foodstuffs (of the 'smoothie' variety)	1,8 g/100 g			
	Fruit or vegetable-based compotes	3,25 g/100 g	8 to guar gum must be visible on		
	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	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 product		

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
			before consumption, in order to take into account the potential risk of gastro-intestinal obstruction.		
Heat-treated milk products fermented with <i>Bacteroides</i>	Specified food category	Maximum levels			
xylanisolvens	Fermented milk products (in liquid, semi-liquid and spray-dried powder forms)				
Hydroxytyrosol	Specified food category	Maximum levels	The designation of the novel food on the labelling of the food		
	Fish and vegetable oils, (except olive oils and olive pomace oils as defined in Part VIII of	0,215 g/kg	products containing it shall be 'hydroxytyrosol'.		
	Annex VII of Regulation (EU) No 1308/2013 (6)), placed as such on the market		The labelling of the food products containing hydroxytyrosol shall bear the following statements: (a) This food product should not be consumed by children under the age of three years, pregnant women, and lactating women;		
	Spreadable fats as defined in Part VII of Annex VII of Regulation (EU) No 1308/2013, placed as such on the market	0,175 g/kg			
			(b) This food product should not be used for cooking, baking or frying'		
Ice Structuring Protein type III	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
HPLC 12	Edible ices	0,01 %	containing it shall be 'Ice Structuring Protein'		
Aqueous extracts of dried leaves of <i>Ilex</i>	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
guayusa	Herbal infusions	In line with normal use in herbal infusions and food supplements of	containing it shall be 'Extracts of dried leaves of <i>Ilex guayusa</i> '		
	Food Supplements as defined in Directive 2002/46/EC	11			

V 1V12						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Infusion from coffee leaves of Coffea arabica L. and/or Coffea canephora Pierre ex A.	Specified food category Infusion from coffee leaves of Coffea arabica L. and/or Coffea canephora Pierre ex A. Froehner placed on the market as such	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Infusion from coffee leaves' or 'Dried infusion from coffee leaves', depending on the form to be marketed.		
	Froehner (Traditional food from a third country)	Flavoured and unflavoured non-alcoholic ready-to-drink beverages (14) Coffee, coffee and chicory extracts, instant coffee, tea, herbal- and fruit-infusions, coffee substitutes, coffee mixes and instant mixes for beverages (and their flavoured counterparts) (14)				
	Iron hydroxide adipate tartrate	Specified food category Food supplements as defined in Directive 2002/46/EC for the adult population Food supplements as defined in Directive 2002/46/EC for children and adolescents under 18 years of age, excluding children under 4 years of age	Maximum levels ≤ 100 mg/day (≤ 30 mg Fe/day) ≤ 50 mg/day (≤ 14 mg Fe/day)	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'iron hydroxide adipate tartrate (nano)'. The labelling of food supplements containing iron hydroxide adipate tartrate shall bear a statement that they should not be consumed by children and adolescents under the age of 18/children under 4 years of age (*) (*) Depending on the age group the food supplement is intended for.		Authorised on 28.8.2022. The inclusion is based of proprietary scientific data protected in accordance with Article 26 of Regulation (EU 2015/2283). Applicant: Nemysis Limited Suite 4.01 Ormond Buildin 31-36 Ormond Quay Uppe Arran Quay Dublin 7, DO F6DC, Dublin, Ireland During the period of data protection, the novel food in hydroxide adipate tartrate authorised for placing on the market within the Union only Nemysis Limited, unless subsequent applicant obtain authorisation for the novel food without reference to the proprietary scientific data protected in accordance with Article 26 of Regulation (EU 2015/2283) or with the agreement of Nemysis Limited End date of the data protection 28.8.2027.

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀	
Iron milk caseinate	Specified food category	Maximum levels	The designation of the novel food		Authorised on 4 June 2023. This inclusion is based or proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regu-	
	Milk and dairy powder products	500 mg/100 g (≤ 10 mg Fe/100 g)	containing it shall be 'iron milk caseinate'. The labelling of food supplements			
	Soft-drinks marketed in relation to physical exercise	85 mg/100 g (≤ 1,7 mg Fe/100 g)				
	Powder cocoa beverage preparations	400 mg/100 g (≤ 8 mg Fe/100 g)	bear a statement that		lation (EU) 2015/2283.	
	Powder or liquid malt-based coffee substitutes	1 050 mg/100 g (≤ 21 mg Fe/100 g)	(a) they should not be consumed by children under 3 years of age;(b) they should not be consumed if other foods containing iron milk caseinate and/or if other foods with added iron are consumed the same day.		Applicant: 'Société des Produits Nestlé S.A.' Avenue Nestlé 55	
	Cereal bars	350 mg/100 g (≤ 7 mg Fe/100 g)		lk ls	1800 Vevey, Switzerland. During the period of data protection, the iron milk caseinate is authorised for placing on the market within the Union only by 'Société des Produits Nestlé S.A.' unless a subsequent applicant obtains authoris-	
	Noodles other than glass noodles	75 mg/100 g (< 1.5 mg Fe/100 g)				
	•					
	Total diet replacement for weight control as defined under Regulation (EU) No 609/2013	235 mg/meal (≤ 4,7 mg Fe/meal) or 700 mg/day (≤ 14,0 mg/Fe/day)				ation for the novel food without reference to the proprietary scientific
	Foods for special medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants and young children	In accordance with the particular nutritional requirements of the persons for whom the products are intended			evidence or scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283 o with the agreement o 'Société des Produits Nestle S.A.'. End date of the data	
	Food supplements as defined in Directive 2002/46/EC, for the adult population	700 mg/day (≤ 14 mg Fe/day)				
	Food supplements as defined in Directive 2002/46/EC, for children and adolescents under 18 years of age, excluding infants and young children	350 mg/day (≤ 7 mg Fe/day)			protection: 4 June 2028.	
	Iron milk caseinate	Milk and dairy powder products Soft-drinks marketed in relation to physical exercise Powder cocoa beverage preparations Powder or liquid malt-based coffee substitutes Cereal bars Noodles other than glass noodles Stock cubes or granulates (bouillon base) Single meal replacements for weight control Total diet replacement for weight control as defined under Regulation (EU) No 609/2013 Foods for special medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants and young children Food supplements as defined in Directive 2002/46/EC, for the adult population Food supplements as defined in Directive 2002/46/EC, for children and adolescents under 18 years of age,	Milk and dairy powder products 500 mg/100 g (≤ 10 mg Fe/100 g)	Iron milk caseinate Specified food category	Specified food category Maximum levels Specified food category Maximum levels	

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection
Isomalto-oligos- accharide	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
	Energy-Reduced Soft Drinks	6,5 %	containing it shall be 'Isom- altooligosaccharide'.	containing it shall be 'Isom-	
	Energy Drinks	5,0 %	Foods containing the novel ingredient must be labelled as 'a source of glucose'.		
	Foods intended to meet the expenditure of intense muscular efforts, especially for sportsmen (including isotonic drinks)	6,5 %	Source of gamesse !		
	Fruit Juices	5 %			
	Processed Vegetables and Vegetable Juices	5 %			
	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		The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Isomaltulose'.		

▼M9

V 1V19						
	Authorised novel food	Conditions under which the nov	el food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
				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'.		
▼ <u>M90</u>						
	Jatropha curcas L.	Specified food category	Maximum levels (g/100g)	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'kernels from edible <i>Jatropha curcas</i> L.'		Authorised on 12 July 2022. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU)
	(edible variety) kernels	Kernels as such, candied or sugar preserved and as processed nuts			s from	
		Cereal bars	5			
		Breakfast cereals	5			2015/2283. Applicant: 'JatroSolution:
		Dried fruits	5			GmbH', Echterdinger Strasse 30, 70599 Stuttgart,
						Germany. During the period of data protection, the nove food kernels from the edibly variety of <i>Jatropha curcas</i> It is authorised for placing of the market within the Uniod only by 'JatroSolution GmbH', unless a subsequer applicant obtains authorisation for the novel foo 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 'JatroSolution GmbH'.
						End date of the dat protection: 12 July 2027.

Lactitol Specified food category Food supplements as defined in Directive 2002/46/EC intended for the adult population Maximum levels The designation of the novel food on the labelling of the food supplements containing it shall be 'Lactitol'	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	►M30 Data Protection ◀
Laction Specified food category	M130	Conditions under which the no	l look may be used	reductional specific labelling requirements	other requirements	But Hotelion 4
Dairy analogues, including beverage whiteners Dairy analogues, including beverages Dairy analogues, including beverage Dairy analogues, in		Food supplements as defined in		on the labelling of the food supplements containing it shall be		
Lacto-N-neotetraose Specified food category Maximum levels		population		Lactitoi		
Unflavoured pasteurised and sterilised (including UHT) milk-based products Unflavoured fermented milk-based products 9,6 g/kg for products other than beverages 9,6 g/kg for products other than beverages 10,6 g/l for beverages 9,6 g/kg for products other than beverages including heat-treated products Dairy analogues, including beverage Whiteners 0,6 g/l for beverages 0,6 g/l for beverages 0,6 g/l for beverages 1,6 g/kg for products other than beverages whiteners 0,6 g/l for beverages 0,6 g/l for beverages 1,6 g/kg for products other than beverages whiteners 0,6 g/l for beverages 0,6 g/l for beverages 1,6 g/kg for products other than beverages whiteners 0,6 g/l for beverages 0,6 g/l for beverages 1,6 g/kg for products other than beverages whiteners 0,6 g/l for beverages 1,6 g/kg for products other than beverages whiteners 1,6 g/kg for products other than beverages whiteners 1,0 g/kg for whitener 0,6 g/l for beverages 1,6 g/kg for products other than beverages whiteners 1,7 leabelling of food supplements containing lacto-N-neotetraose are consumed the same day. 2,7 leabelling of food supplements containing lacto-N-neotetraose are consumed the same day. 3,7 leabelling of food supplements containing lacto-N-neotetraose are consumed the same day. 4,8 g/kg for products other than beverages whiteners 6 g/kg for whitener 1,9 g/kg for whitener 1,0 g/kg for whitener 2,0 g/kg for whitener 3,0 g/kg for whitener 4,0 g/kg for	M119 Lacto-N-neotetraose	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs.		
Unflavoured fermented milk-based products 9,6 g/kg for products other than beverages Flavoured fermented milk-based products including heat-treated products Dairy analogues, including beverage Dairy analogues, including beverage Mitteners Dairy analogues, including beverage Of g/kg for products other than beverages Of g/kg for whitener Of g/kg Of g/kg for products other than beverages Of g/kg for whitener Of g/kg Of g/kg for products other than beverages Of g/kg for whitener Of g/kg Of g/kg for whitener Of g/kg Of g/kg for whitener			0,6 g/l	containing it shall be 'lacto-N-neotetraose'. 2. The labelling of food supplements containing lacto-N-neotetraose shall bear a statement that the supplements should not be used if other foods with added lacto-N-		
Solution		Unflavoured fermented milk-based products	0,6 g/l for beverages			
Flavoured fermented milk-based products including heat-treated products 9,6 g/kg for products other than beverages 9,6 g/kg for products other than beverages 0,6 g/l for beverages 0,6 g/l f						
9,6 g/kg for products other than beverages Dairy analogues, including beverage whiteners 0,6 g/k for beverages 0,6 g/l for beverages 0,6 g/l for beverages 0,6 g/l for beverages 0,6 g/l for beverages 6 g/kg for products other than beverages 200 g/kg for whitener Cereal bars 6 g/kg Table-top sweeteners 100 g/kg Infant formula as defined under Regulation (EU) No 609/2013 9,6 g/kg for products other than beverages 0,6 g/l for beverages 200 g/kg for whitener 6 g/kg 100 g/kg			0,6 g/l for beverages	same day.		
Dairy analogues, including beverage whiteners 0,6 g/l for beverages 6 g/kg for products other than beverages 200 g/kg for whitener Cereal bars 6 g/kg Table-top sweeteners 100 g/kg Infant formula as defined under Regulation (EU) No 609/2013 beverages 0,6 g/l for beverages that the supplements should not be used if breast milk or other foods with added lacto-N-neotetraose are consumed the same day.				supplements containing lacto-N-neotetraose intended for young		
6 g/kg for products other than beverages 200 g/kg for whitener Cereal bars 6 g/kg Table-top sweeteners 100 g/kg Infant formula as defined under Regulation (EU) No 609/2013 neotetraose are consumed the same day.			0,6 g/l for beverages	that the supplements should not be used if breast milk or other		
Cereal bars 6 g/kg Table-top sweeteners 100 g/kg Infant 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 manu-				neotetraose are consumed the		
Table-top sweeteners 100 g/kg Infant 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 manu-			200 g/kg for whitener			
Infant formula as defined under Regulation (EU) No 609/2013 Infant formula as defined under use, marketed as such or reconstituted as instructed by the manu-		Cereal bars	6 g/kg			
Regulation (EU) No 609/2013 use, marketed as such or reconstituted as instructed by the manu-		Table-top sweeteners	100 g/kg			
			use, marketed as such or reconstituted as instructed by the manu-			

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	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 foods and baby foods for infants and young children as defined under 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			
	Milk based drinks and similar products intended for young children	0,6 g/l for milk-based drinks and similar products in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			
	Foods 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			
	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			

_	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		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	4,8 g/l – the maximum level refers to the products ready to use			
		Food Supplements as defined in Directive 2002/46/EC, for the general population, excluding infants	1,5 g/day for general population 0,6 g/day for young children			
<u>45</u>						
	Lacto-N-tetraose ('LNT')	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		Authorised on 23.4.20 This inclusion is based
((microbial source)	Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products	1,0 g/l	containing it shall be 'lacto- <i>N</i> -tetraose'. The labelling of food supplements		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 lacto-Nettraose is authorised for placing on the marke within the Union only by Glycom A/S, unless a subsequent applicant obtains
		Unflavoured fermented milk-based products	1,0 g/l (beverages) 10 g/kg (products other than beverages)	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.		
		Flavoured fermented milk-based products including heat-treated products	1,0 g/l (beverages) 10 g/kg (products other than beverages)			
		Beverages (flavoured drinks)	1,0 g/l			authorisation for the n food without reference
		Cereal bars	10 g/kg			the proprietary scientific evidence or scientific data protected in accordance with
	Infant formula as defined under Regulation (EU) No 609/2013	0,8 g/l in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer	.		Article 26 of Regulate (EU) 2015/2283 or with agreement of Glycom A	

▼<u>M45</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	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			End date of the data protection: 23.4.2025.
	Processed cereal-based food, baby food for infants and young children as defined under Regulation (EU) No 609/2013				
	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 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, excluding infants	2,0 g/day for young children, children, adolescents, and adults			

-						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Lacto-N-tetraose ('LNT')	Specified food category	Maximum levels (expressed as lacto-N-tetraose)	The designation of the novel food on the labelling of the foodstuffs		Authorised on 24.1.2023. This inclusion is based on
(produced derivative	(produced by derivative strains of <i>E. coli</i> BL21(DE3))	Infant formula as defined under Regulation (EU) No 609/2013	1,82 g/L in the final product ready for use, marketed as such or recon- stituted as instructed by the manu- facturer	containing it shall be 'lacto-N-tetraose'. The labelling of food supplements containing lacto-N-tetraose (LNT) shall bear a statement that Y (a) they should not be consumed by children under 3 years of age; (b) they should not be used if other foods containing added lacto-N-tetraose are consumed the same day.		proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283.
		Follow-on formula as defined under Regulation (EU) No 609/2013	1,82 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			Applicant: 'Chr. Hansen A/S', Boege Allé 10-12, 2970 Hoersholm, Denmark. During the period of data protection, the novel food Lacto-N-tetraose is authorised for placing on the market within the Union only by 'Chr. Hansen A/S' unless a subsequent
		Processed cereal-based foods and baby foods for infants and young children as defined under Regulation (EU) No 609/2013	1,82 g/L or 1,82 g/kg 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 1,82 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer Foods for special medical purposes for infants and young children as defined under 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 1,82 g/L or 1,82 g/kg in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer.	for use, marketed as such or reconstituted as instructed by the manu-			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 'Chr. Hanser A/S'. End date of the data protection: 24.1.2028.	
		Foods for special medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants and young children	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, for the general population, excluding infants and young children	4,6 g/day			

MI						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	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'		
<u>M9</u>	Lucerne leaf extract	Specified food category	Maximum levels	The designation of the novel food		
	from Medicago sativa	Food supplements as defined in Directive 2002/46/EC	10 g/day	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	The designation of the novel food on the labelling of the foodstuffs		
		Fruit/vegetable juice-based drinks (including concentrates)	2,5 mg/100 g	containing 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			

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	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 Blakeslea trispora	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
•	Fruit/vegetable juice-based drinks (including concentrates)	2,5 mg/100 g	containing 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			

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Lycopene from tomatoes	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		_
	Fruit/vegetable juice-based drinks (including concentrates)	2,5 mg/100 g	containing 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	of the		
	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 tomatoes	Specified food category	Maximum levels of lycopene	The designation of the novel food on the labelling of the foodstuffs		
from tomatoes	Fruit/vegetable juice-based drinks (including concentrates)	2,5 mg/100 g	containing it shall be 'Lycopene oleoresin from tomatoes'		
	Drinks intended to meet the expenditure of intense muscular effort especially for sportsmen	2,5 mg/100 g			

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		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			
		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			
<u>M52</u>						
	Hen egg white	Specified food category	Maximum levels	The designation of the novel food		
	lysozyme hydrolysate	Food supplements as defined in Directive 2002/46/EC intended for adult population	1000 mg/day	on the labelling of food supplements containing it shall be 'Hen egg white lysozyme hydrolysate'.		
<u>M9</u>						
	Magnesium citrate	Specified food category	Maximum levels The designat	The designation of the novel food		
	malate	Food Supplements as defined in Directive 2002/46/EC		on the labelling of the foodstuffs containing it shall be 'Magnesium citrate malate'		
	Magnolia Bark	Specified food category	Maximum levels	The designation of the novel food		
	Extract	Mints (confectionary products)	0,2 % for breath freshening	on the labelling of the foodstuffs containing it shall be 'Magnolia		
		Chewing gum	purposes. Based on a 0,2 % maximum incorporation 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.	Bark Extract'		
	Maize-germ oil high in unsaponifiable matter Specified food category Food Supplements as defined in Directive 2002/46/EC	Maximum levels	The designation of the novel food			
		Food Supplements as defined in Directive 2002/46/EC	2 g/day	on the labelling of the foodstuffs containing it shall be 'Maize-germ oil extract'		
		Chewing gum	2 %			

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Methylcellulose	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs	Methylcellulose is not to be	
		Edible ices	2 %	containing it shall be 'Methylcel- lulose'	used in foods specially prepared for young children	
		Flavoured drinks				
		Flavoured or unflavoured fermented milk products				
		Cold desserts (dairy, fat, fruit, cereal, egg-based products)				
		Fruit preparations (pulps, purees or compotes)				
		Soups and broths				
▼ <u>M11</u>						
	1-Methylnicoti- namide chloride	Specified food category Food Supplements as defined in Directive 2002/46/EC for the adult population excluding pregnant and lactating women	Maximum levels 58 mg/day	The designation of the novel food on the labelling of the foodstuffs containing it shall be '1- Methylnicotinamide chloride'. Food supplements containing 1- Methylnicotinamide shall bear the following statement: This food supplement should be consumed by adults only excluding pregnant and lactating women		Authorised on 2 September 2018. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Pharmena SA, Wolczanska 178, 90 530 Lodz, Poland. During the period of data protection the

▼<u>M11</u>

	Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
						novel food 1-methylnicotinamide 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.
▼ <u>M9</u>	(6S)-5-methyltet- rahydrofolic acid, glucosamine salt	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be '(6S)-5-methyltetrahydrofolic acid, glucosamine salt' or '5MTHF-glucosamine'		protection: 2 September 2023
		Food Supplements as defined in Directive 2002/46/EC as a source of folate				
	Monomethylsil- anetriol (Organic	Specified food category	Maximum levels of silicon	The designation of the novel food on the labelling of the food supplements containing it shall be		
	Silicon)	Food Supplements as defined in Directive 2002/46/EC for adult population (in liquid form)	10,40 mg/day	'Organic silicon (monomethylsilanetriol)'		

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀	
Monosodium salt of	Specified food category	Maximum levels	The designation of the novel food		Authorised on 30 A	
L-5-methyltetrahy- drofolic acid	Food supplements as defined in Directive 2002/46/EC, excluding food supplements for infants and young children	(expressed as folic acid)	2. The labelling of food supplements containing monosodium salt of L-5-methyltetrahydrofolic acid shall bear a statement that the food supplements should not be consumed by infants and young children (children under 3 years of age)		2024. This inclusion based on proprie scientific evidence scientific data protected accordance with Article of Regulation (EU) 20 2283.	
	Infant formula and follow-on formula as defined in Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013		sodium salt of L-5-methyltetrahy- drofolic acid shall bear a statement that the food supplements should not be consumed by infants and young children (children under 3 years of age) Solitoria		Applicant: Merck & KmG, Im Laternenacker 8200 Schaffhausen, S
	Processed cereal-based foods and baby foods for infants and young children as defined in Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013			zerland. During the pe of data protection, the ne food monosodium salt L-5-methyltetrahydrofolic acid is authorised for place	
	Foods for special medical purposes as defined in Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013			on the market within Union only by Merck Cie KmG, unless subsequent applicant obt	
	Total diet replacement for weight control as defined in Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013			authorisation for the n food without reference the proprietary scien	
	Food fortified in accordance with Regulation (EC) No 1925/2006	In accordance with Regulation (EC) No 1925/2006			evidence or scientific protected in accordance Article 26 of R lation (EU) 2015/2283 with the agreement Merck & Cie KmG.	
					End date of the protection: 30 April 202	

▼M9

V IVI)						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
▼ <u>M87</u>						
	Mung bean (<i>Vigna</i> radiata) protein	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'mung bean protein from <i>Vigna radiata</i> '.		Authorised on 15 May 2022. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with
		Protein products	20 g/100 g			Article 26 of Regulation (EU) 2015/2283. Applicant: Eat Just, Inc., 2000 Folsom Street San Francisco, CA 94110 USA. During the period of data protection, the novel mung bean protein is authorised for placing on the market within the Union only by Eat Just, 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 Eat Just, Inc. End date of the date protection: 15 May 2027.

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Mycelial extract	Specified food category	Maximum levels	The designation of the novel food		
	from Shiitake mushroom	Bread products	2 ml/100 g	on the labelling of the foodstuffs containing it shall be 'extract from		
	(Lentinula edodes)	Soft drinks	0,5 ml/100 ml	the mushroom <i>Lentinula edodes</i> ' or 'extract from Shiitake mushroom'		
		Ready prepared meals	2,5 ml per meal	extract from Simtake musinoom		
		Foods based on yoghurt	1,5 ml/100 ml			
		Food supplements as defined in Directive 2002/46/EC	2,5 ml per day dose			
▼ <u>M92</u>	Nicotinamide riboside chloride	Specified food category	Maximum levels			Authorised on 20 February
	riboside chioride	Food supplements as defined in Directive 2002/46/EC	300 mg/day for the adult population, excluding pregnant and lactating women		containing it shall be 'nicotinamide riboside chloride'.	2020. This inclusion is based on proprietary scientific evidence and scientific data protected in
			230 mg/day for pregnant and lactating women			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.

	Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		Foods for special medical purposes as defined by Regulation (EU) No 609/2013 for the adult population, excluding pregnant and lactating women	In accordance with the particular nutritional requirements of the persons for whom the products are intended	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'nicotinamide riboside chloride' The labelling of foodstuffs		
		Total diet replacement for weight control as defined by Regulation (EU) No 609/2013 for the adult population, excluding pregnant and lactating women	500 mg/day	containing the novel food shall bear a statement that those foods should only be consumed by persons above 18 years of age excluding pregnant and lactating women.		
		Meal replacements for the adult population, excluding pregnant and lactating women	150 mg/meal (maximum 2 meals/day up to a maximum of 300 mg/day)			
▼ <u>M9</u>						
	Noni fruit juice (Morinda citrifolia)	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
		Pasteurised fruit and fruit nectar based drinks	30 ml with one serving (up to 100 % noni juice)	containing it shall be 'Noni juice' or 'Juice of Morinda citrifolia'		
			or 20 ml twice a day, not more than 40 ml per day			
	Noni fruit juice powder (<i>Morinda</i> 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'		

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Noni fruit puree and concentrate (Morinda citrifolia)	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be:		
,		Fruit puree	For fruit puree: 'Morinda citrifolia fruit puree' or 'Noni fruit puree'		
	Candy/confectionery	45 g/100 g	For fruit concentrate: 'Morinda citrifolia fruit concentrate' or 'Noni fruit concentrate'		
	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			

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	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			
	2001/113/EC	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			
	Powdered nutritional drink mixes (dry weight)	12 g/100 g			
	Carbonated beverages	3 g/100 g			
	Ice cream & sorbet	7 g/100 g			
	Yoghurt	3 g/100 g			
	Biscuits	12 g/100 g			
	Buns, cakes and pastries	12 g/100 g			
	Breakfast cereals (wholegrain)	20 g/100 g			
	Jams and jellies in accordance with Directive 2001/113/EC	30 g/100 g			
	Sweet spreads, fillings and icings	7 g/100 g			

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Savoury sauces, pickles, gravies and condiments	20 g/100 g			
	Food Supplements as defined in Directive 2002/46/EC	6 g/day			
Noni leaves (Morinda citrifolia)	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
(Aormai Cargona)	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 of Morinda citrifolia	containing it shall be 'Noni leaves' or 'leaves of <i>Morinda citrifolia</i> '. 2. Instructions shall be given to the consumer that a cup of infusion should not be prepared with more than 1 g of dried and roasted leaves of Morinda citrifolia.		
Noni fruit powder (Morinda citrifolia)	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Morinda citrifolia fruit powder' or 'Noni fruit powder'		
(Adriana Carifolia)	Food Supplements as defined in Directive 2002/46/EC	2,4 g per/day			
Odontella aurita microalgae	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
microaigae	Flavoured pasta	1,5 %	containing it shall be 'Odontella aurita microalgae'		
	Fish soups	1 %			
	Marine terrines	0,5 %			
	Broth preparations	1 %]		
	Crackers	1,5 %			
	Frozen breaded fish	1,5 %			

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Oil enriched with phytosterols/ phytostanols	Specified food category	Maximum levels of phytosterols/ phytostanols	In accordance with Annex III.5 to Regulation (EU) No 1169/2011		
	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	novel food ingredient shall be presented in such a manner that they can be easily divided into			
	Milk based products, such as products based on semi-skimmed and skimmed milk products, possibly with the addition of fruits and/or cereals, products based on fermented milk such as yoghurt and cheese based products (fat content ≤ 12 g per 100 g), where possibly the milk fat has been reduced and the fat or protein has been partly or fully replaced by vegetable fat or protein	per day) of added phytosterols/phytostanols. 2. The amount of phytosterols/phytostanols added to a container of beverages shall not exceed 3 g. 3. Salad dressings, mayonnaise and spicy sauces shall be packed as single portions.			
	Soya drinks				
	Salad dressings, mayonnaise and spicy sauces				

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Oil extracted from squids	Specified food category	Maximum levels of DHA and EPA combined	The designation of the novel food on the labelling of the foodstuffs		
	Dairy products except milk-based beverages	200 mg/100 g or for cheese products 600 mg/100 g	containing it shall be 'Squid oil'.		
	Dairy analogues except drinks	200 mg/100 g or for analogues to cheese products 600 mg/			
	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 609/2013 and meal replacements for weight control	200 mg/meal			
E-tour t form Pour	Specified food estagem	Maximum levels	The decimal of the second field		And wind an 22 December
Extract from Panax notoginseng and Astragalus membra- naceus	Specified food category Food supplements as defined in Directive 2002/46/EC for the general adult population, excluding food supplements for pregnant women	35 mg/day	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Extract from Panax notoginseng and Astragalus membranaceus' The labelling of food supplements containing extract from Panax notoginseng and Astragalus membranaceus shall bear a statement that those food supplements should not be consumed by the population under 18 years of age and by pregnant women.		Authorised on 23 December 2020. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283.

▼<u>M55</u>

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
						Applicant: NuLiv Science, 1050 W. Central Ave., Building C, Brea, CA 92821, USA.
						During the period of data protection, the novel food is authorised for placing on the market within the Union only by NuLiv Science, 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 NuLiv Science.
						End date of the data protection: 23 December 2025.
▼ <u>M126</u>	<u>5</u>					
	Partially defatted chia seed (Salvia	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
	hispanica L.) powders	Powder with high protein content		containing it shall be 'Partially defatted chia seed (Salvia hispanica)		
		Unflavoured fermented milk products, including natural unflavoured buttermilk (excluding sterilised buttermilk) non-heat-treated after fermentation	0,7 %	powder'		
		Unflavoured fermented milk products, heat-treated after fermentation	0,7 %	1		
		Flavoured fermented milk products including heat-treated products	0,7 %	_		
		Confectionery	10 %	1		

▼<u>M126</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Fruit juices as defined by Council Directive 2001/112/EC (8) and vegetable juices	2,5 %			
	Fruit nectars as defined by Directive 2001/112/EC and vegetable nectars and similar products	2,5 %			
	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 high fi	ibre content			Authorised for use in cakes and pastries, processed fruit
	Confectionery	4 %			and vegetables (including
	Fruit juices as defined by Directive 2001/112/ EC and vegetable juices	2,5 %			vegetable-based dishes), bread and rolls, pasta based products and protein products on 13 November 2023. This
	Fruit nectars as defined by Directive 2001/112/EC and vegetable nectars and similar products	4 %			inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of
	Flavoured drinks	4 %			Regulation (EU) 2015/2283.
	Food supplements as defined in Directive 2002/46/EC excluding food supplements for infants and young children	12 g/day			Applicant: Functional Products Trading Arica S.A./BENEXIA, Luis Pasteur 5850, Oficina 403, Quinto Piso.
	Cakes and pastries	5 g/100 g		During the period protection, partially chia seed (Salvia L.) powder with a h content for use in c pastries, processed vegetables (vegetable-based dish	Vitacura, Santiago – Chile. During the period of data
	Processed fruit and vegetables (including vegetable-based dishes)	10 g/100 g			protection, partially defatted chia seed (<i>Salvia hispanica</i> L.) powder with a high fibre
	Bread and rolls	10 g/100 g			content for use in cakes and pastries, processed fruit and
	Pasta based products	8 g/100 g			vegetable-based dishes), bread
	Protein products	10 g/100 g			and rolls, pasta based products and protein products is auth- orised for placing on the

▼<u>M126</u>

	Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
						market within the Union only by Functional Products Trading Arica S.A./BENEXIA, unless a subsequent applicant obtains authorisation for the same 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 Functional Products Trading Arica S.A./BENEXIA. End date of the data protection:
						13 November 2028.
▼ <u>M63</u>						
	Partially defatted rapeseed powder	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Partially defatted Rapeseed powder'. Any foodstuff containing 'Partially defatted Rapeseed powder' from		
	from Brassica rapa L. and	Cereal bars mixed	20 g/100 g			
	Brassica napus L.	Muesli and similar breakfast cereals	20 g/100 g			
		Extruded breakfast cereal products	20 g/100 g	Brassica rapa L. and Brassica napus L.' shall bear a statement that this ingredient may cause		
		Snacks (excluding potato crisps)	15 g/100 g	allergic reaction to consumers who are allergic to mustard and		
		Breads and rolls with added special ingredients (such as seeds, raisins, herbs)	7 g/100 g	products thereof. That statement shall appear in close proximity to the list of ingredients.		
		Brown breads bearing statements on the absence or reduced presence of gluten in accordance with the requirements of Commission Implementing Regulation (EU) No 828/2014	7 g/100 g			

_	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
_		Multigrain bread and rolls	7 g/100 g			
		Meat substitutes	10 g/100 g			
_		Meat balls	10 g/100 g			
▼ <u>M9</u>						
_	Pasteurised fruit-based prep-	Specified food category	Maximum levels	The wording 'pasteurised by		
a u	run-based prep- arations produced asing high-pressure reatment	Types of fruit: apple, apricot, banana, blackberry, blueberry, cherry, coconut, fig, grape, grapefruit, mandarin, mango, melon, peach, pear, pineapple, prune, raspberry, rhubarb, strawberry		high-pressure treatment' shall be displayed next to the name of the fruit preparations as such and in any product in which it is used		
▼ <u>M100</u>						
	Pea and rice protein	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Pea and rice protein fermented by Shiitake mushroom mycelia'.	Authorised on 24.1.2023. This inclusion is based or	
I	fermented by Lentinula edodes	Bakery wares, breads, rolls, croutons, pizza	5 g/100 g			proprietary scientific data
	Shiitake nushroom) mycelia	Breakfast cereals and cereal bars	33 g/100 g			protected in accordance with Article 26 of Regulation (EU)
		Fruit- and vegetable-based drinks	20 g/100 ml		A Ii S	2015/2283. Applicant: MycoTechnology Inc., 18250 E. 40th Avenue
		Ready-to-mix beverage powders	93 g/100 g			
		Cocoa and chocolate confectionary	7 g/100 g			Suite 50, Aurora, 80011
		Dairy analogues and non-dairy meal replacements for weight control	11 g/100 g			Colorado, United States. During the period of data protection, the novel food
		Fermented milk-based products	5 g/100 g			pea and rice protein fermented by Lentinula
		Pasta-based products	15 g/100 g			edodes (Shiitake mushroom)
		Meat preparations and meat products	14 g/100 g			mycelia is authorised for placing on the market
		Soups (ready-to-eat) and soup concentrates or powders	3 g/100 g			within the Union only by MycoTechnology, Inc.
		Salads	26 g/100 g			applicant obtains authoris-
		Meat analogues	40 g/100 g			ation for the novel food
		Milk-based drinks	1 g/100 g			
		Single meal replacements for weight control	1 g/100 g			

▼<u>M100</u>

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
						without reference to th proprietary scientific dat protected in accordance with Article 26 of Regulation (EU 2015/2283 or with th agreement of MycoTech nology, Inc.
- 3.535						protection: 24.1.2028.
▼ <u>M37</u>						
	Phenylcapsaicin	Foods for special medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants, young children and children under the age of 11 years	Maximum levels 2,5 mg/day	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'phenylcapsaicin'.		Authorised on 19 Decembe 2019. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU 2015/2283.
		Food supplements as defined in Directive 2002/46/EC intended for the general population, excluding children under the age of 11 years	2,5 mg/day			Applicant: aXichem AB Södergatan 26, SE 211 34 Malmö Sweden. During the period of data protection the novel food phenylcap saicin is authorised fo placing on the marke within the Union only by aXichem AB, unless a subsequent applicant obtain authorisation for the nove 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.

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Phosphated maize	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
	starch	Baked bakery products	15 %	containing it shall be 'Phosphated		
		Pasta		maize starch'		
		Breakfast cereals				
		Cereal bars				
▼ <u>M11</u>	2					
	Phosphated wheat	Specified food category	Maximum levels	The designation of the novel food		
	starch	Baked bakery products	15 %	on the labelling of the foodstuffs containing it shall be 'Phosphated wheat starch'.		
		Pasta				
		Breakfast cereals				
		Cereal bars				
▼ <u>M9</u>						
	Phosphatidylserine from fish phosp- holipids	Specified food category	Maximum levels of phosphati- dylserine	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Fish phos-		
	nonpius	Beverages based on yoghurt	50 mg/100 ml	phatidylserine'		
		Powders based on milk powders	3 500 mg/100 g (equivalent to 40 mg/100 ml ready to drink)			
		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			

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Phosphatidylserine from soya phosp- holipids	Specified food category	Maximum levels of phosphati- dylserine	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Soya phos-		
nonpius	Beverages based on yoghurt	50 mg/100 ml	phatidylserine'		
	Powders based on milk powder	3,5 g/100 g (equivalent to 40 mg/100 ml ready to drink)			
	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 equal amounts of	Specified food category	Maximum levels of phosphati- dylserine	The designation of the novel food on the labelling of the foodstuffs containing shall be 'Soy phosphatidylserine and phosphatidic acid'	The product is not intended to be marketed to	
phosphatidylserine and phosphatidic	Breakfast cereals	80 mg/100 g		pregnant or breast-feeding	
acid	Cereal bars	350 mg/100 g		women	
	Foods based on yogurt	80 mg/100 g			
	Soy-based yogurt-like products	80 mg/100 g			
	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			

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Phospholipides	Specified food category	Maximum levels			
from egg yolk	Not specified				
Phytoglycogen	Specified food category	Maximum levels	The designation of the novel food		
	Processed foods		on the labelling of the foodstuffs containing it shall be 'Phytog- lycogen'		
Phytosterols/	Specified food category	Maximum levels	In accordance with Annex III.5 of		
phytostanols	Rice drinks	1. They shall be presented in such a manner that they can be easily divided into portions that contain either a maximum of 3 g (in case of 1 portion/day) or a maximum of 1 g (in case of 3 portions/day) of added phytosterols/phytostanols. The amount of phytosterols/phytostanols added to a container of beverages shall not exceed 3 g. Salad dressings, mayonnaise and spicy sauces shall be packed as single portions			
	Rye bread with flour containing ≥ 50 % rye (wholemeal rye flour, whole or cracked rye kernels and rye flakes) and ≤ 30 % wheat; and with ≤ 4 % added sugar but no fat added.				
	Salad dressings, mayonnaise and spicy sauces.				
	Soya drink				
	Milk type products, such as semi-skimmed and skimmed milk type products, possibly with the addition of fruits and/or cereals, 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.				
	Products based on fermented milk such 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				
	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			

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Plum kernel oil	Specified food category	Maximum levels			
	For frying and as seasoning	In line with normal food use of vegetable oils			
Potato proteins (coagulated) and hydrolysates thereof	Not specified		The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Potato protein'		
Prolyl oligopep-	Specified food category	Maximum levels	The designation of the novel food		
tidase (enzyme preparation)	Food Supplements as defined in Directive 2002/46/EC for general adult population	120 PPU/day (2,7 g of enzyme preparation/day) (2 × 10 ⁶ PPI/day)	on the labelling of the foodstuffs containing it shall be 'Prolyl oligopeptidase'		
		PPU – Prolyl Peptidase Units or Proline Protease Units			
		PPI – Protease Picomole International			
M136			1 771 1 1 1 1 1 1 1 1		
Protein concentrate from <i>Lemna gibba</i>	Specified food category Cereal bars	Maximum levels 10 g/100 g	1. The designation of the novel food on the labelling of the foodstuffs		Authorised on 30 April 2024. This inclusion is
and Lemna minor	Prepacked bread and rolls	1,7 g/100 g	containing it shall be 'protein concentrate from the <i>Lemna</i>		based on proprietary scientific evidence and
	Powdered drink mixes	20 g/100 g	gibba and Lemna minor plants' or 'protein concentrate from the		scientific data protected in accordance with Article 26
	Noodles	6 g/100 g	Lemna gibba plant' depending on the presence of Lemna minor.		of Regulation (EU) 2015/ 2283.
			2. Where foods containing the novel food include an amount of vitamin K that is considered significant in accordance with point 2 of Part A of Annex XIII to Regulation (EU) No 1169/2011, the nutrition declaration shall indicate the amount of vitamin K.		

▼<u>M136</u>

_	Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		Food supplements as defined in Directive 2002/46/EC for the adult population	1 g/day	1. The designation of the novel food on the labelling of the foodstuffs containing it shall be 'protein concentrate from the <i>Lemna gibba</i> and <i>Lemna minor</i> plants' or 'protein concentrate from <i>Lemna gibba</i> plant' depending on the presence of <i>Lemna minor</i> .		Applicant: ABC Kroos BV Drosteweg 8, 8101 NE Raalte, NETHERLANDS During the period of date protection, the novel food protein concentrate from Lemna gibba and Lemna minor is authorised fo placing on the marke
				2. The labelling of food supplements containing the novel food shall bear a statement that they should only be consumed by adults.		within the Union only by ABC Kroos BV, unless a subsequent applicant obtain authorisation for the nove food without reference to the proprietary scientific evidence or scientific data
				3. Where food supplements containing the novel food include an amount of vitamin K that is considered significant in accordance with point 2 of Part A of Annex XIII to Regulation (EU) No 1169/2011 and Article 8 of Directive 2002/46/		protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of ABC Kroos BV.
				EC, the labelling of food supplements containing novel food shall indicate the amount of vitamin K.		protection: 30 April 2029.
M50						
	Protein extract rom pig kidneys	Specified food category	Maximum levels			
1.	rom pig kidneys	Food Supplements as defined in Directive 2002/46/EC	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)			

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
<u> 410</u>					
Pyrroloquinoline quinone disodium salt	Specified food category Food Supplements as defined in Directive 2002/46/EC intended for the adult population, excluding pregnant and lactating women	Maximum levels 20 mg/day	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Pyrroloquinoline 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		Authorised on 2 September 2018. This inclusion is based on proprietary scientific evidence are scientific data protected accordance with Article 2 of Regulation (EU) 2015 2283. Applicant: Mitsubishi Garchemical Company, Inc. Mitsubishi Building 5-Marunouchi 2-chom Chiyoda-ku, Tokyo 100 8324, Japan. During the period of data protection the novel food Pyrroloquinolin quinone disodium salt is authorised for placing on the market within the Union on by Mitsubishi Gas Chemic Company, Inc., unless subsequent applicant obtain authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected accordance with Article 26 Regulation (EU) 2015/228 or with the agreement of Mitsubishi Gas Chemic Company, Inc. End date of the da protection: 2 september 202

	Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Rapeseed oil high in unsaponifiable	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
	matter	Food Supplements as defined in Directive 2002/46/EC	1,5 g per portion recommended for daily consumption	containing it shall be 'Rapeseed oil extract'		
	Rapeseed Protein	As a vegetable protein source in foods except in infant formula and follow-on formula		 The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Rapeseed protein'. Any foodstuff containing 'rapeseed protein' shall bear a 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 ingredients. 		
▼ <u>M17</u>	Refined 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	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., Stortorget 1, Kystens 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 peptide

▼<u>M17</u>

	Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
						concentrate is authorised of placing on the market with the Union only by Marea AS unless a subseque applicant obtains author ation for the novel for without reference to the proprietary scientific day rotected in accordance what is a considered and accordance what is a greement of Marealis A End date of the day rotection: 20 November 2023.
<u>M59</u>						
	Trans-resveratrol	Specified food category	Maximum levels	The designation of the novel food on the labelling of the food		
		Food supplements as defined in Directive 2002/46/EC for the adult population	150 mg/day	supplements containing it shall be ' <i>Trans</i> -resveratrol'. 2. The labelling of food supplements containing <i>trans</i> -resveratrol shall bear a statement that people using medicines should only consume the product under medical supervision.		

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Trans-resveratrol	Specified food category	Maximum levels	1. The designation of the novel food		
(microbial source)	Food supplements as defined in Directive 2002/46/EC	In line with normal use in food supplements of resveratrol extracted from Japanese knotweed (Fallopia japonica)	on the labelling of the food supplements containing it shall be ' <i>Trans</i> -resveratrol'.		
			The labelling of for supplements containing trans-resveratrol shall bear statement that people using medicines should only consumate product under medicines supervision. The designation of the povel for		
Rooster comb	Specified food category	Maximum levels	The designation of the novel food		
extract	Milk-based drinks	40 mg/100 g or mg/100 ml	on the labelling of the foodstuffs containing it shall be 'Rooster		
	Milk based fermented drinks	80 mg/100 g or mg/100 ml	comb extract' or 'Cockerel comb extract'		
	Yoghurt-type products	65 mg/100 g or mg/100 ml			
	Fromage frais	110 mg/100 g or mg/100 ml			
Sacha inchi oil from	Specified food category	Maximum levels	The designation of the novel food		
Plukenetia volubilis	As for linseed oil	In line with normal food use of linseed oil	on the labelling of the foodstuffs containing it shall be 'Sacha inchi oil (Plukenetia volubilis)'		
Salatrims	Specified food category	Maximum levels	1. The designation of the novel food		
	Bakery products and confectionary		on the labelling of the foodstuffs containing it shall be 'reduced energy fat (salatrims)'.		
			2. There shall be a statement that excessive consumption may lead to gastro-intestinal disturbance.		
			3. There shall be a statement that the products are not intended for use by children.		

Cooking fats

VI						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Schizochytrium sp. oil rich in DHA and EPA	Specified food category	Maximum levels of DHA and EPA combined	The designation of the novel food on the labelling of the foodstuffs		
	EFA	Food supplements as defined in Directive 2002/46/EC for the adult population excluding pregnant and lactating women	3 000 mg/day	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 products intended for young children	200 mg/100 g			
		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)					
		Breakfast cereals	500 mg/100 g			
				1	1	

360 mg/100 g

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		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			
		Fish analogues	300 mg/100 g			
		Meat analogues	300 mg/100 g			
▼ <u>M27</u>						_
	Schizochytrium sp. (ATCC PTA-9695)	Specified 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 <i>Schizochytrium</i> sp.'		
	oil	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			
		Spreadable fats and dressings	600 mg/100 g			
		Breakfast cereals	500 mg/100 g			
		Food Supplements as defined in Directive 2002/46/EC	250 mg DHA/day for general population			
			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			

▼<u>M27</u>

Α	Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection <
		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 and young children as defined in Regulation (EU) No 609/2013	200 mg/100 g			
		Fruit/vegetable puree	100 mg/100 g			
71						
	chizochytrium sp.	Specified food category	Maximum levels of DHA	The designation of the novel food		
(F)	CC-3204) oil	Infant formula and follow-on formula as defined in Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013	on the labelling of the foodstuffs containing it shall be 'Oil from the microalgae <i>Schizochytrium</i> sp.'.		
		Food supplements as defined in Directive 2002/46/EC for the general population above 3 years of age	1 g/day	The labelling of food supplements containing <i>Schizochytrium</i> sp. (FCC-3204) oil shall bear a statement that they should not be consumed by infants and children under 3 years of age.		

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
125						
	Schizochytrium sp.	Specified food category	Maximum levels of DHA	The designation of the novel food on the labelling of the foodstuffs		
		Dairy products except milk-based drinks	200 mg/100 g or for cheese products 600 mg/100 g	containing it shall be 'Oil from the microalgae <i>Schizochytrium</i> sp.'		
		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 2002/46/EC	250 mg DHA/day for general population			
			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			
		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				
		L	1	1		

02017R2470 — EN — 01.05.2024 — 046.001 — 110

▼<u>M25</u>

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		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, 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			
		Fruit/vegetable puree	100 mg/100 g			
▼ <u>M52</u>						
	Schizochytrium sp. (T18) oil	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
	` '	Dairy products except milk-based drinks	200 mg/100 g or for cheese products 600 mg/100 g	containing it shall be 'Oil from the microalgae <i>Schizochytrium</i> sp.'.		
		Dairy analogues except drinks	200 mg/100 g or for analogues to cheese products 600 mg/100 g			
		Spreadable fats and dressings	600 mg/100 g			
		Breakfast cereals	500 mg/100 g			

▼<u>M52</u>

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Food supplements as defined in Directive 2002/46/EC	250 mg DHA/day for general population			
		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/100g			

▼<u>M52</u>

V 1V132						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		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 g			
▼ <u>M65</u>						
	Schizochytrium sp.	Specified food category	Maximum levels of DHA	The designation of the novel food on the labelling of the foodstuffs		Authorised on 16 May 2021. This inclusion is based on
	(WZU477) oil	Infant formula and follow-on formula as defined in Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013	containing it shall be 'Oil from the microalgae Schizochytrium sp.'		proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283.
						Applicant: Progress Biotech by, Canaalstaete, Kanaalweg 33, 2903LR Capelle aan den Ijssel, the Netherlands.
						During the period of data protection, the novel food is authorised for placing on the market within the Union only by Progress Biotech by unless a subsequent applicant obtains authorisation for that novel food without reference to the proprietary scientific evidence or scientific data

▼<u>M65</u>

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
						protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of Progress Biotech bv. End date of the data protection: 16 May 2026 (5 years).
▼ <u>M57</u>						
	Selenium-containing yeast (Yarrowia lipolytica) biomass	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'selenium-containing yeast (Yarrowia		
		Food supplements as defined in Directive 2002/46/EC (³), excluding food supplements for infants and children under 4 years of age	50 mg/day for children from 4 to 6 years of age, resulting in 10 μg of selenium per day 100 mg/day for children from 7 to 10 years of age, resulting in 20 μg of selenium per day 500 mg/day for adolescents from 11 to 17 years of age, resulting in 100 μg of selenium per day 800 mg/day for adults, resulting in 160 μg of selenium per day	lipolytica) biomass'. The labelling of food supplements containing selenium-containing yeast (Yarrowia lipolytica) biomass shall bear a statement that the food supplements should not be consumed by infants and children under 4 years of age/children under 7 years of age/children under 11 years of age/children and adolescents under 18 years of age (12).		

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
1 2 3'-Sialyllactose (3'-SL) sodium salt	Specified food category	Maximum levels (expressed as 3'-Sialyllactose)	The designation of the novel food on the labelling of the foodstuffs		Authorised on 18 February 2021. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283.
(microbial source)	Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products	0,25 g/L	containing it shall be '3'-Sialyl-lactose sodium salt'. The labelling of food supplements containing 3'-Sialyllactose sodium		
	Unflavoured fermented milk-based products	0,25 g/L (beverages)	salt shall 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 and young children		
		0,5 g/kg (products other than beverages)			Applicant: Glycom Kogle Allé 4, DK-2 Hørsholm, Denmark. Du the period of data protec the novel food 3'-si-
	Flavoured fermented milk-based products including heat-treated products	0,25 g/L (beverages)		lactose sodium salt is a orised for placing on market within the U	
		2,5 g/kg (products other than beverages)			only by Glycom A/S, un a subsequent appl obtains authorisation for novel food without refer
	Beverages (flavoured drinks, excluding drinks with a pH less than 5)	0,25 g/L			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.
	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			End date of the protection: 18 Febr 2026.
	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 recon- stituted as instructed by the manu- facturer	.		

▼<u>M62</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Processed cereal-based food and baby food for infants and young children as defined in Regulation (EU) No 609/2013				
		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 foods for weight control as defined in Regulation (EU) No 609/2013	0,5 g/L (beverages)			
		5 g/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 Directive 2002/46/EC, excluding food supplements for infants and young children	0,5 g/day			

1412					
Authorised novel	food Conditions under which the ne	ovel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
<u>M122</u>					
3'-Sialyllactose (SL') sodium sal		Maximum levels	sodium salt shall bear a statement that (a) they should not be consumed by children under 3 years of age; (b) they should not be used if other foods containing added 3'-sialyl-lactose sodium salt are consumed the same day.		Authorised on 6 February 2023. This inclusion is based on proprietary
(produced by derivative strain <i>E. coli</i> BL21(DI		0,28 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer		scientific eviden scientific data pro accordance with A of Regulation (E 2283. Applicant: 'Chr. H S', Boege Allé 10	scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/
	Follow-on formula as defined under Regulation (EU) No 609/2013	0,28 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer		During the period of data protection, the novel food 3'-Sialyllactose sodium salt is authorised for placing on the market within the Union only by Chr. Hansen A/S unless a subsequent applicant obtains authorised.	
	Processed cereal-based foods for infants and young children and baby foods for infants and young children as defined under Regulation (EU) No 609/2013	product ready for use, marketed as		ation 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 'Chr. Hansen A/S'.	
	Milk based drinks and similar products intended for young children	0,28 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			End date of the data protection: 6 February 2028.

▼<u>M122</u>

-	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		Foods for special medical purposes for infants and young children as defined under 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 0,28 g/L or 0,28 g/kg in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer.			
		Foods for special medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants and young children	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, for the general population, excluding infants and young children	0,7 g/day			
▼ <u>M135</u>						
	3'-Sialyllactose (3'- SL) sodium salt	Specified food category	Maximum levels (expressed as 3'-Sialyllactose)	The designation of the novel food on the labelling of the foodstuffs containing it shall be '3'-Sialyllactose sodium salt'. The labelling of food supplements		Authorised on 30 April 2024. This inclusion is based on proprietary scientific evidence and scientific data
i I	produced using a derivative strain of E. coli W (ATCC	Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products	0,25 g/L			
9	9637))	Unflavoured fermented milk-based products	0,25 g/L (beverages)	containing 3'-Sialyllactose (3'-SL) sodium salt shall bear a statement		protected in accordance with Article 26 of Regu-
			0,5 g/kg (products other than beverages)	that they should not be consumed: (a) if foods containing added 3'- Sialyllactose sodium salt are consumed on the same day; (b) by children under 3 years of age. Applic Bio Centra Nakan Tokyo During protect	lation (EU) 2015/2283. Applicant: Kyowa H	lation (EU) 2015/2283. Applicant: Kyowa Hakko Bio Co., Ltd, Nakano
		Flavoured fermented milk-based products	0,25 g/L (beverages)		Central Park South, Nakano 4-10-2, Nakano-ku	
		including heat-treated products	2,5 g/kg (products other than beverages)			Tokyo, 164-0001 Japan. During the period of data protection, the novel food
		Beverages (flavoured drinks, excluding drinks with a pH less than 5)	0,25 g/L			3'-sialyllactose sodium salt

▼<u>M135</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Cereal bars	2,5 g/kg			produced using a derivative
	Infant formula as defined under 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			strain of E. coli W (ATCC 9637) is authorised for placing on the market within the Union only by Kyowa Hakko Bio Co., Ltd, unless a subsequent applicant
	Follow-on formula as defined under 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			obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in accordance with
	Processed cereal-based food and baby food for infants and young children as defined under 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			Article 26 of Regulation (EU) 2015/2283 or with the agreement of Kyowa Hakko Bio Co., Ltd. End date of the data
		1,25 g/kg for products other than beverages			protection: 30 April 2029.
	Milk based drinks and similar products	0,15 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 under Regulation (EU)	0,5 g/L (beverages)			
	No 609/2013 5,0 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 Directive 2002/46/EC, excluding food supplements for infants and young children	1,0 g/day			

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
<u>M60</u>						
	6'-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 containing it shall be '6'-Sialyl-		Authorised on 17 February 2021. 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 6'-sialyllactose sodium salt is authorised 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 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.
	(Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products	0,5 g/L	lactose sodium salt'. The labelling of food supplements		
		Unflavoured fermented milk-based products	0,5 g/L (beverages)	containing 6'-Sialyllactose (6'-SL) sodium salt shall bear a statement		
			2,5 g/kg (products other than beverages)	a) if foods containing added 6'- Sialyllactose sodium salt are consumed on the same day. b) by infants and young children		
		Flavoured fermented milk-based products including heat-treated products	0,5 g/L (beverages)			
		including hear-treated products	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 under 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 under 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			End date of the data protection: 17 February 2026.
		Processed cereal-based food and baby food for infants and young children as defined under 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 for products other than beverages			

▼<u>M60</u>

Authorised novel f	conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Milk based drinks and similar products intended for young children	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 under Regulation (EU)	1,0 g/L (beverages)]		
	No 609/2013	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 Directive 2002/46/EC, excluding food supplements for infants and young children	1,0 g/day			
115					
6'-Sialyllactose ('6'-SL') sodium	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		Authorised on 4 June 2023. This inclusion is based on
(produced by derivative strains <i>E. coli</i> BL21(DE.	Infant formula as defined under Regulation (EU) No 609/2013	0,70 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer	containing it shall be '6'-Sialyl-lactose sodium salt'. The labelling of food supplements containing 6'-Sialyllactose (6'-SL) sodium salt shall bear a statement that (a) they should not be consumed by children under 3 years of age; (b) they should not be consumed if other foods containing added 6'-sialyllactose sodium salt are consumed the same day.	proprievide prote Artic (EU) t Appl S', I Hoer Durin	proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283.
	Follow-on formula as defined under Regulation (EU) No 609/2013	0,70 g/L in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			Applicant: 'Chr. Hansen A/S', Bøge Allé 10-12, 2970 Hoersholm, Denmark. During the period of data protection, the novel food
	Processed cereal-based foods for infants and young children and baby foods for infants and young children as defined under Regulation (EU) No 609/2013	0,70 g/L or 0,70 g/kg in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer			6'-Sialyllactose sodium salt is authorised for placing on the market within the Union only by Chr. Hansen A/S unless a subsequent applicant obtains authoris-
	Milk based drinks and similar products intended for young children	0,70 g/L in the final product ready for use, marketed as such or recon- stituted as instructed by the manu- facturer			ation for the novel food without reference to the proprietary scientific evidence or scientific data

▼<u>M115</u>

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		Foods for special medical purposes for infants and young children as defined under 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 0,70 g/L or 0,70 g/kg in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer.			protected in accordance with Article 26 of Regulation (EU) 2015/2283 or with the agreement of 'Chr. Hansen A/S'. End date of the data protection: 4 June 2028.
		Foods for special medical purposes as defined under Regulation (EU) No 609/2013 excluding foods for infants and young children	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, for the general population, excluding infants and young children	1,8 g/day			
▼ <u>M127</u>						
	6'-Sialyllactose (6'- SL) sodium salt	Specified food category	Maximum levels (expressed as 6'- Sialyllactose)	The designation of the novel food on the labelling of the foodstuffs containing it shall be '6'-Sialyl-		Authorised on 13.11.2023. This inclusion is based on proprietary scientific
	E. coli W (ATCC	Unflavoured pasteurised and unflavoured sterilised (including UHT) milk products	0,5 g/L	lactose sodium salt'. The labelling of food supplements containing 6'-Sialyllactose (6'-SL)	evidence and scientific data protected in accordance with Article 26 of Regu-	
	9637))	Unflavoured fermented milk-based products	0,5 g/L (beverages)	sodium salt shall bear a statement that they should not be consumed:		lation (EU) 2015/2283.
			2,5 g/kg (products other than beverages)	(a) if foods containing added 6'- Sialyllactose sodium salt are		Applicant: Kyowa Hakko Bio Co., Ltd, 1-9-2, Otemachi, Choyoda-ku Tokyo, 100-0004, Japan.
		Flavoured fermented milk-based products including heat-treated products	0,5 g/L (beverages)	consumed on the same day;		During the period of data protection, the novel food
		metating near-neared products	5,0 g/kg (products other than beverages)	(b) by children under 3 years of age.		6'-sialyllactose sodium salt produced by derivative strain of <i>E. coli</i> W (ATCC
		Beverages (flavoured drinks, excluding drinks with a pH less than 5)	0,5 g/L			9637) is authorised for placing on the market within
		Cereal bars	5,0 g/kg			

▼<u>M127</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Infant formula as defined under 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			the Union only by Kyowa Hakko Bio Co., Ltd, unless a subsequent applicant obtains authorisation for the
	Follow-on formula as defined under 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			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 Kyowa Hakko Bio Co., Ltd.
	Processed cereal-based food and baby food for infants and young children as defined under 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			End date of the data protection: 13.11.2028.
		2,5 g/kg for products other than beverages			
	Milk based drinks and similar products	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 under Regulation (EU) No 609/2013	1,0 g/L (beverages)			
		10,0 g/kg (products other than beverages)			

▼<u>M127</u>

Authorised novel food	Conditions under which the nov	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	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, excluding food supplements for infants and young children	1,0 g/day			
Syrup from Sorghum bicolor (L.) Moench (Traditional food from a third country)	Not specified		The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Sorghum (Sorghum bicolor) syrup'		
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	 The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Fermented soybean extract'. The labelling of food supplements containing fermented soybean extract shall bear a statement that persons taking medication should only consume the product under medical supervision. 		

▼<u>M9</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection <
Spermidine-rich wheat germ extract (Triticum aestivum)	Specified food category Food Supplements as defined in Directive 2002/46/EC intended for the adult population, and least time respect and least time respect to the second control of the second cont	Maximum levels Equivalent of max. 6 mg/day spermidine	The designation of the novel food on the labelling of the food supplements containing it shall be 'spermidine-rich wheat germ extract'		
Sucromalt	Specified food category Not specified	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Sucromalt'. The designation of the novel food on the labelling shall be accompanied by indication that the product is a source of glucose and fructose.		
Sugar cane fibre	Specified food category Bread Bakery goods Meat and muscle products	Maximum levels 8 % 5 % 3 %			
	Seasonings and spices Grated cheeses Special diet foods Sauces	3 % 2 % 5 % 2 %			
	Beverages	5 %			
Sugars 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 (<i>Theobroma cacao</i> L.) pulp', 'Glucose obtained from cocoa (<i>Theobroma cacao</i> L.) pulp' or 'Fructose obtained from cocoa (<i>Theobroma cacao</i> L.) pulp', depending on the form used.		

▼M9

V <u>IVI</u>						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Sunflower oil extract	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
		Food Supplements as defined in Directive 2002/46/EC	1,1 g/day	containing it shall be 'Sunflowe oil extract'		
▼ <u>M73</u>						
	Synsepalum dulcificum dried	Specified food category	Maximum levels	The designation of the novel food on the labelling of food		Authorised on 5 December 2021. This inclusion
	fruits	Food Supplements as defined in Directive 2002/46/EC for adult population excluding pregnant and lactating women	0,7 g/day	supplements containing it shall be 'dried Synsepalum dulcificum fruits' 2. The labelling of food supplements containing Synsepalum dulcificum dried fruits shall bear a statement that this food supplement should be consumed by adults only excluding pregnant and lactating women.	11 n n o- n ur d d	based on proprietar scientific evidence an scientific data protected is accordance with Article 2 of Regulation (EU) 2013 2283. Applicant: Medicinal Garder S.L. Marqués de Urquijo 4' 1° D, Office 1, Madrid 28008, Spain.
						During the period of da protection, the novel food authorised for placing on the market within the Union on by Medicinal Gardens S.J. unless a subsequent application obtains authorisation for the novel food without referent to the proprietary scientific evidence or scientific da protected in accordance with Article 26 of Regulation (EU 2015/2283) or with the agreement of Medicin Gardens S.L. End date of the data protection 5 December 2026.

▼M9

V <u>IVI</u>						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
▼ <u>M89</u>						
	Tetrahydrocur- cuminoids	Specified food category Food Supplements as defined in Directive 2002/46/EC for the adult population, excluding pregnant and lactating women	Maximum levels 140 mg/day	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'tetrahydrocurcuminoids'. The labelling of food supplements containing tetrahydrocurcuminoids shall bear a statement that a) they should be consumed by adults only, excluding pregnant and lactating women; b) they should not be consumed if other food supplements containing curcumin and/or curcuminoids are consumed on the same day.		Authorised on 11 July 2022. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: 'Sabinsa Europe GmbH', Monzastrasse 4, 63225 Langen, Germany. During the period of data protection, the novel food tetrahydrocurcuminoids is authorised for placing on the market within the Union only by 'Sabinsa Europe GmbH' 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 'Sabinsa Europe GmbH'. End date of the data protection: 11 July 2027.

▼<u>M9</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Dried Tenebrio molitor larva (yellow mealworm)	Specified food category Dried Tenebrio molitor larva, whole or in powder	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Dried <i>Tenebrio molitor</i> larva (yellow mealworm)'. The labelling of the foodstuffs containing dried <i>Tenebrio</i>	Other requirements	Authorised on 22 June 20 This inclusion is based proprietary scientific evide and scientific data protected accordance with Article 26 Regulation (EU) 2015/2283 Applicant: SAS EAP Gro
	Protein products 10 g/100 g	10 g/100 g	molitor larva (yellow mealworm) shall bear a statement that this ingredient may cause allergic		35 Boulevard du Li Échange, 31650 Saint-Ore de-Gameville, France.
	Biscuits	10 g/100 g	reactions to consumers with known allergies to crustaceans and products thereof, and to dust mites. This statement shall appear in close proximity to the list of ingredients.	During the period of oprotection, the novel food authorised for placing on	
	Pasta-based products	10 g/100 g 10 g/100 g		market within the Union of by SAS EAP Group, unles ubsequent applicant obtauthorisation for that not food without reference to proprietary scientific evide or scientific data protecte accordance with Article 2 Regulation (EU) 2015/22 or with the agreement of SEAP Group.	
					End date of the data protect 22 June 2026.

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
Dried Tetraselmis chuii microalgae	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
	Sauces	20 % or 250mg/day	containing it shall be 'Dried microalgae <i>Tetraselmis chuii</i> ' or 'Dried microalgae <i>T. chuii</i> '		
	Special salts	1 %	Food supplements containing dried microalgae <i>Tetraselmis chuii</i> shall bear the following statement:		
	Condiment	250 mg/day	'Contains negligible amounts of iodine'		
	Food Supplements as defined in Directive 2002/46/EC	250 mg/day			
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	Maximum levels	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'D-Tagatose'.		
	Not specified				
			2. The labelling of any product where the level of D-Tagatose exceeds 15 g per serving and all beverages containing greater than 1 % D-Tagatose (as consumed) shall bear a statement 'excessive consumption may produce laxative effects'.		
Taxifolin-rich	Specified food category	Maximum levels	The designation of the novel food		
extract	Yogurt plain/Yogurt with fruits ^(*)	0,020 g/kg	on the labelling of the foodstuffs containing it shall be 'taxifolin-rich		
	Kephir ^(*)	0,008 g/kg	extract'		
	Buttermilk ^(*)	0,005 g/kg			

▼ <u>M52</u>

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		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 the general population, excluding infants, young children, children and adolescents younger than 14 years	100 mg/day			
		(*) When used in milk products Taxifolin-rich part, any milk constituent	extract may not replace in whole or in			
▼ <u>M9</u>						
	Trehalose	Specified food category	Maximum levels	The designation of the novel food on the labelling of the foodstuffs		
		Not specified		containing it shall be 'Trehalose' and shall be displayed on the labelling of the product as such or in the list of ingredients of foodstuffs containing it. 2. The designation of the novel food on the labelling shall be accompanied by indication that the 'Trehalose is a source of glucose'.		

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection <
<u>152</u>						
	UV-treated mushrooms (Agaricus bisporus)	Specified food category	Maximum levels of vitamin D_2	The designation on the label of the novel food as such or of the foodstuffs containing it shall be		
	(Aguricus visporus)	Mushrooms (Agaricus bisporus)	20 μg of vitamin D ₂ /100 g fresh weight	'UV-treated mushrooms (Agaricus bisporus)'.		
				2. The designation on the label of the novel food as such or of the foodstuffs containing it shall be accompanied by indication that a 'controlled light treatment was used to increase vitamin D levels' or 'UV treatment was used to increase vitamin D ₂ levels'.		
[84						
	UV-treated baker's yeast (Sacchar-	Specified food category	Maximum levels of vitamin D_2	The designation of the novel food on the labelling of the foodstuffs		
	omyces cerevisiae)	Yeast-leavened breads and rolls	5 μg/100 g	containing it shall be 'vitamin D yeast' or 'vitamin D ₂ yeast'		
		Yeast-leavened fine bakery wares	5 μg/100 g			
		Food supplements as defined in Directive 2002/46/EC	In accordance with Directive 2002/ 46/EC			
		Pre-packed fresh or dry yeast for home baking	45 μg/100 g for fresh yeast 200 μg/100 g for dried yeast	The designation of the novel food on the labelling of the foodstuffs shall be 'vitamin D yeast' or 'vitamin D ₂ yeast'.		

02017R2470 - EN - 01.05.2024 - 046.001 - 131

▼<u>M84</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
			 The labelling of the novel food shall bear a statement that the foodstuff is only intended for baking and that it should not be eaten raw. The labelling of the novel food shall bear instructions for use for the final consumers so that a maximum concentration of 5 μg/100 g of vitamin D₂ in final home-baked products is not exceeded. 		
	Dishes, incl. ready-to-eat meals (excluding soups and salads)	3 μg/100 g	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'vitamin D yeast' or 'vitamin D ₂ yeast'		
	Soups and salads	5 μg/100 g	yeast of vitaliili D ₂ yeast		
	Fried or extruded cereal, seed or root-based products	5 μg/100 g			
	Infant formula and follow-on formula as defined by Regulation (EU) No 609/2013	In accordance with Regulation (EU) No 609/2013			
	Processed cereal-based food as defined by 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			

▼<u>M84</u>

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	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 dessert 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 replacements for weight control	5 μg/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			

Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
UV-treated bread	n 'c		The designation on the label of the novel food shall be accompanied by 'contains vitamin D produced by UV-treatment'		
	Yeast leavened bread and rolls (without toppings)	3 μg vitamin $D_2/100$ g	C V-deadlicht		
UV-treated milk	Specified food category	Maximum levels of vitamin D_3	The designation on the label of the novel food shall be 'UV-treated'.		
	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	2. Where UV-treated milk contains an amount of vitamin D that is considered significant in accordance with Point 2 of Part A of Annex XIII to 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' or 'milk containing vitamin D resulting from UV-treatment'.		
	Pasteurised semi-skimmed milk as defined in Regulation (EU) No 1308/2013 to be consumed as such	1-15 μg/kg for general population excluding infants			

tamin D ₂	Specified food category Breakfast cereals Yeast-leavened bread and pastries	wel food may be used	The designation of the novel food on the labelling of the foodstuffs	Other requirements	
_	Breakfast cereals	- ` ` /	on the labelling of the foodstuffs		
_	Breakfast cereals	- ` ` /	on the labelling of the foodstuffs		
isiirooni powdei		2,25 μg of vitamin $D_2/100$ g	f vitamin D ₂ /100 g on the labelling of the foodstuffs containing it shall be 'UV-treated mushroom powder containing vitamin D' or 'UV-treated		Authorised on 27 August 2020. This inclusion is based on proprietary 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 D ₂ 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 Regulation
	Yeast-leavened bread and pastries				
		2,25 μg of vitamin $D_2/100~g$			
Г	Grain products and pastas	2,25 μg of vitamin $D_2/100$ g	mushroom powder containing vitamin D ₂ '		
	Fruit juice and fruit/vegetable blend beverages	1,125 μg of vitamin $D_2/100$ mL	The labelling of food supplements containing vitamin D ₂ mushroom powder shall bear a statement that		
	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)	they should not be consumed by infants		
	Cheese (excluding cottage cheese, ricotta cheese, and hard-grating cheeses)	2,25 μg 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			(EU) 2015/2283 or with agreement of Oak
	Extruded vegetable snacks	2,25 μg of vitamin D ₂ /100 g			Naturals, LP.
	Foods for Special Medical Purposes as defined under Regulation (EU) No 609/2013 excluding those intended for infants	15 μg/day			End date of the da protection: 27 August 202
	Food supplements as defined in Directive 2002/46/EC intended for the general population excluding infants	15 μg/day			
		milks) Cheese (excluding cottage cheese, ricotta cheese, and hard-grating cheeses) Meal replacement bars and beverages Dairy analogues Meat analogues Soups and broths Extruded vegetable snacks Foods for Special Medical Purposes as defined under Regulation (EU) No 609/2013 excluding those intended for infants Food supplements as defined in Directive 2002/46/EC intended for the general popu-	milks) µg of vitamin D₂/100 mL (beverages) Cheese (excluding cottage cheese, ricotta cheese, and hard-grating cheeses) Meal replacement bars and beverages 2,25 μg of vitamin D₂/100 g/1,125 μg of vitamin D₂/100 g/1,125 μg of vitamin D₂/100 mL (beverages) Dairy analogues 2,25 μg of vitamin D₂/100 g/1,125 μg of vitamin D₂/100 mL (beverages) Meat analogues 2,25 μg of vitamin D₂/100 g Soups and broths 2,25 μg of vitamin D₂/100 g Extruded vegetable snacks 2,25 μg of vitamin D₂/100 g Extruded vegetable snacks 2,25 μg of vitamin D₂/100 g Extruded vegetable snacks 2,25 μg of vitamin D₂/100 g Foods for Special Medical Purposes as defined under Regulation (EU) No 609/2013 excluding those intended for infants Food supplements as defined in Directive 2002/46/EC intended for the general popu-	Milk and dairy products (excluding fluid milks) 2,25 μg of vitamin D ₂ /100 g/1,125 μg of vitamin D ₂ /100 g/1,125 μg of vitamin D ₂ /100 g/1,125 μg of vitamin D ₂ /100 g mL (beverages) Cheese (excluding cottage cheese, ricotta cheese, and hard-grating cheeses) 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 2,25 μg of vitamin D ₂ /100 g Extruded vegetable snacks 2,25 μg of vitamin D ₂ /100 g Extruded vegetable snacks 2,25 μg of vitamin D ₂ /100 g 15 μg/day 15 μg/day	Milk and dairy products (excluding fluid milks) 2,25 μg of vitamin D ₂ /100 g/1,125 μg of vitamin D ₂ /100 g mL (beverages) they should not be consumed by infants Cheese (excluding cottage cheese, ricotta cheese, and hard-grating cheeses) 2,25 μg of vitamin D ₂ /100 g/1,125 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 Extruded vegetable snacks 2,25 μg of vitamin D ₂ /100 g Foods for Special Medical Purposes as defined under Regulation (EU) No 609/2013 excluding those intended for infants 15 μg/day Food supplements as defined in Directive 2002/46/EC intended for the general popu- 15 μg/day

▼<u>M9</u>

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀	
▼ <u>M76</u>							
	Vitamin D ₂ mushroom powder	Specified food category	Maximum levels of vitamin D_2	1. The designation of the novel food on the labelling of the foodstuffs		Authorised on 19 December 2021. This inclusion is based	
	musiiroom powder	Breakfast cereals	2,1 μg/100 g	containing it shall be 'UV-treated		on proprietary scientific evidence and scientific data protected in accordance with	
		Yeast leavened bread and similar pastries	2,1 μg/100 g	mushroom powder containing vitamin D ₂ '			
		Grain products and pasta and similar products	2,1 μg/100 g	2. The labelling of food supplements containing vitamin		Article 26 of Regulation (EU) 2015/2283.	
		Fruit/vegetable juices and nectars	1,1 µg/100 ml (marketed as such or reconstituted as instructed by the manufacturer)	D ₂ mushroom powder shall bear a statement that they should not be consumed by infants and children under 3 years of age.		Applicant: MBio, Monaghan Mushrooms, Tullygony, Tyholland, Co. Monaghan, Ireland. During the period of data protection, the novel food vitamin D ₂ mushroom powder is authorised for placing on the market within the Union 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	
		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)				
		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)			Article 26 of Regulation (EU) 2015/2283 or with the agreement of MBio, Monaghan Mushrooms.	
		Extruded vegetable snack	2,1 μg/100 g			End date of the date protection: 19 December	
		Meal replacement for weight control	2,1 μg/100 g			2026.	
		Foods for Special Medical Purposes as defined under Regulation (EU) No 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 Directive 2002/46/EC excluding food supplements intended for infants and young children	15 μg of vitamin D ₂ /day				

WIS						
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Vitamin D ₂ mushroom powder	Specified food category	Maximum levels of vitamin D_2 (μ g/ 100 g or 100 ml)	The designation of the novel food on the labelling of the foodstuffs		Authorised on 24 January 2023. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Monterey Mushrooms Inc, 260 Westgate Drive Watsonville, CA
		Milk analogues	1,1	containing it shall be 'UV-treated mushroom powder containing		
		Dairy analogues other than milks	2,2	vitamin D ₂ ' 2. The labelling of food		
		Breakfast cereals and cereal bars	2,2	supplements containing vitamin D ₂ mushroom powder shall bear		
		Soups	2,2	a statement that they should not be consumed by infants and		
		Dried soups	22,5	children under 3 years of age.		95076, the United States.
		Whey powder	14,1			During the period of data protection, the novel food
		Fruit/vegetable juices and nectars	1,1			vitamin D ₂ mushroo powder is authorised to
		Fruit/vegetable juice powder	12,4			placing on the market within the Union only by Monterey Mushrooms Inc, unless a subsequent applicant obtains authorisation for the novel food without reference to the proprietary scientific evidence or scientific data protected in
		Fruit/vegetable juice concentrate (liquid)	3,4			
		Soft drinks marketed in relation to physical exercise and fermented non-alcoholic drinks (with exclusion of dairy fermented drinks)	1,1			
		Foods for Special Medical Purposes as defined under Regulation (EU) No 609/2013 excluding those intended for infants In accordance with the particular nutritional requirements of the persons for whom the products are intended but not higher than 15 μg/day Total diet replacement for weight control as defined in			accordance with Article 26 Regulation (EU) 2015/2283 with the agreement Monterey Mushrooms Inc. End date of the date protecti	
			15 μg/day			24 January 2028.
		Regulation (EU) No 609/2013				
		Meal replacements for weight control	5 μg/meal			
		Food supplements as defined in Directive 2002/46/EC, excluding food supplements for infants and young children	15 μg/day	1		

Autho	orised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► M30 Data Protection •
Vitam quino	nin K ₂ (mena- one)	To be used in compliance with Directive 2002/ and/or Regulation (EC) No 1925/2006	46/EC, Regulation (EU) No 609/2013	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'Menaquinone' or 'Vitamin K ₂ '		
Wheat	nt bran extract	Specified food category	Maximum levels	The designation of the novel food	The 'Wheat Bran Extract' may not be	
		Beer and substitutes	0,4 g/100 g	on the labelling of the foodstuffs containing it shall be 'Wheat bran		
		Ready to eat cereals	9 g/100 g	extract'	introduced onto	
		Dairy products	2,4 g/100 g]	the market as a food supple- ment or food supplement ingredient. Nor	
		Fruit and vegetable juices	0,6 g/100 g]		
		Soft drinks	0,6 g/100 g]		
		Meat preparations	2 g/100 g		may it be added to infant formula.	
_{'8} ——						
Wolffid	<i>ia arrhiza</i> and/	Specified food category	Maximum levels	The designation of the novel food		
fresh p	olffia globosa plants (Tradi- food from a country)	Wolffia arrhiza and/or Wolffia globosa fresh plants as such		on the labelling of the foodstuffs containing it shall be 'Wolffia arrhiza and Wolffia globosa' or 'Wolffia arrhiza' or 'Wolffia globosa' depending on the plant used.		
8						
Xylo-o acchai	oligos- crides	Specified food category	Maximum levels (10)	The designation of the novel food on the labelling of the foodstuffs		
acciiai	iriucs	White bread	14 g/kg	containing it shall be 'Xylo-oligos-		
		Wholemeal bread	14 g/kg	accharides'		
		Breakfast cereals	14 g/kg			
		Biscuits	14 g/kg			
		Soy drink	3,5 g/kg			
		Yoghurt (9)	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			

				1	1	
	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
1113	-					
	Yarrowia lipolytica	Specified food category	Maximum levels	1. The designation of the novel food		
	yeast biomass	Food supplements as defined in Directive 2002/46/EC, excluding food supplements for infants and young children	6 g/day for children from 10 years of age, adolescents and general adult population	on the labelling of the foodstuffs containing it shall be 'Yarrowia lipolytica yeast biomass'.		
			3 g/day for children from 3 to 9 years of age	2. The labelling of meal replacements for weight control containing <i>Yarrowia lipolytica</i>		
		Meal replacements for weight control for the adult population	3 g/meal (maximum 2 meals/day up to a maximum of 6 g/day)	yeast biomass shall bear a statement that they should only be used by persons above 18 years of age and should not be used if food supplements containing <i>Yarrowia lipolytica</i> yeast biomass are consumed on the same day.		
<u>19</u>	Yeast beta-glucans	Specified food category	Maximum levels of pure beta-glucans from yeast (Sacchar- omyces cervisiae)	containing it shall be 'Yeast		
		Food supplements as defined in Directive 2002/46/EC, excluding food supplements for infants and young children	1,275 g/day for children older than 12 years and general adult popu- lation	(Saccharomyces cerevisiae) beta- glucans'		
			0,675 g/day for children younger than 12 years			
		Total diet replacement for weight control as defined in Regulation (EU) No 609/2013	1,275 g/day			
		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	1,275 g/day			
		Beverages based on fruit and/or vegetable juices including concentrate and dehydrated juices	1,3 g/kg			
		Fruit-flavoured drinks	0,8 g/kg			
		Cocoa beverages preparation powder	38,3 g/kg (powder)			

02017R2470 - EN - 01.05.2024 - 046.001 - 139

	Authorised novel food	Conditions under which the no	vel food may be used	Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
		Other beverages	0,8 g/kg (ready to drink)			
			7 g/kg (powder)	7		
		Cereal bars	6 g/kg	7		
		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	1		
		Milk based beverages	3,8 g/kg	7		
		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)			
		Chocolate and confectionery	4 g/kg			
		Protein bars and powders	19,1 g/kg			
		Jam, marmalade and other fruit spreads	11,3 g/kg			
<u>M12</u>	Zeaxanthin	Specified food category	Maximum levels	The designation of the novel food		
		Food Supplements as defined in Directive 2002/46/EC	2 mg/day	on the labelling of the foodstuffs containing it shall be 'Zeaxanthin'.		
<u>M9</u>	Zinc L-pidolate	Specified food category	Maximum levels	The designation of the novel food		
		Foods covered by Regulation (EU) No 609/2013	3 g/day	on the labelling of the foodstuffs containing it shall be 'Zinc L-pidolate'		
		Milk based drinks and similar products intended for young children		L-pidoiate		
		Meal replacement for weight control				
		Foods intended to meet the expenditure of intense muscular effort, especially for sportsmen				

Authorised novel food	Conditions under which the novel food may be used		Additional specific labelling requirements	Other requirements	► <u>M30</u> Data Protection ◀
	Food bearing statement on 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				

- (¹) 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).
- (2) 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).
- (3) 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).
- (4) 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).
- (5) 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).
- (e) 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).
- ►M33 (7) Maximum use levels in the final product ready for use, marketed as such or reconstituted as instructed by the manufacturer. ◄
- ► M47 (8) Council 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).
- ▶ M48 (9) When used in milk products xylo-oligosaccharides shall not replace, in whole or in part, any milk constituent.
- (10) Maximum levels calculated on the basis of the specifications of Powder form 1. ◀
- ►M51 (11) The minimum specification for vitamin D content in vitamin D₂ mushroom powder of 1 000 µg vitamin D₂/gram of mushroom powder is used. ◄
- (12) Depending on the age group the food supplement is intended for.
- (13) Without prejudice to the requirements of Regulation (EU) No 609/2013 and Regulation (EU) 2016/127.
- (14) Not a traditional food use.

Authorised novel food	Conditions under which t	he novel food may be us	sed		Additional specific labelling requirements	Other requirements	Data protection
Frozen, dried and powder forms of Acheta domesticus	Specified food category	(marketed as suc	vels (g/100g) h or reconstituted he instructions)	1.	The designation of the novel food on the labelling of the foodstuffs containing it shall		Authorised on 3 March 2022. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Fair Insects BV, Industriestraat 3, 5107 NC
(house cricket)		Frozen	Dried or powder	1	be 'Frozen Acheta domesticus (house cricket)',		
	Frozen, dried, and powder forms of Acheta domesticus				'Dried/powdered Acheta domesticus (house cricket)'depending on the		
	Protein products other than meat analogues	40	20	2.	foodstuffs containing frozen, dried or powder forms of <i>Acheta domesticus</i> (house cricket) shall bear a statement that this ingredient may cause		Dongen, the Netherlands. During the period of data protection, the novel food is auth-
	Bread and rolls	30	10				orised for placing on the market within the Union only by Fair Insects BV, 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 Fair Insects BV.
	Bakery wares, cereal bars, and stuffed pasta products	30	15				
	Biscuits	30	8	allergic reactions consumers with known	allergic reactions to		
	Pasta-based products (dry)	3	1		allergies to crustaceans, molluscs and products thereof, and to dust mites. This statement shall appear		
	Soups and soup concentrates or powders	20	5				
	Processed potato products, legumes- and vegetable- based dishes, and pasta- or pizza-based products	15	5		in close proximity to the list of ingredients.		End date of the data protection: 3 March 2027.
	Corn flour based snacks	40	20				
	Beer-like beverages, alcoholic drink mixes	1	1]			
	Nuts, oilseeds and chickpeas	40	25				
	Sauces	30	10				
	Meat preparations	40	16				
	Meat analogues	80	50				
	Chocolate confectionary	30	10				
	Frozen fermented milk based products	15	5				

Authorised novel food	Conditions under which t	he novel food may be us	sed	Additional specific labelling requirements	Other requirements	Data protection
Frozen, dried and powder forms of <i>Locusta migratoria</i> (migratory locust)	Specified food category	(marketed as suc	vels (g/100 g) h or reconstituted he instructions)	The designation of the novel food on the labelling of the foodstuffs containing it shall be 'frozen Locusta migratoria'		Authorised on 5.12.2021. This inclusion is based on proprietary scientific evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: Fair Insects BV, Industriestraat 3, 5107 NC
	. 5 .	Frozen	Dried or Powder	(migratory locust)', 'dried/ powder Locusta migratoria (migratory locust)', 'Whole Locusta migratoria (migratory		
	Frozen, dried and powder forms of Locusta migratoria			locust) powder' depending on the form used. 2. The labelling of the foodstuffs containing frozen dried or		Dongen, the Netherlands. During the period of data protection, the novel food is authorised for placing on the market
	Processed potato products; legumes-based dishes and pasta-based products	15	5	powder forms of Locusta migratoria (migratory locust) shall bear a statement that this ingredient may cause allergic reactions to consumers with known allergies to crustaceans, molluscs and products thereof, and to mites. This statement shall appear in close proximity to the list of ingredients.		within the Union only by Fair Insects BV, 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 Fair Insects BV. End date of the data protection: 5.12.2026.
	Meat analogues	80	50			
	Soups and concentrated soups	15	5			
	Canned/jarred legumes and vegetables	20	15			
	Salads	15	5			
	Beer-like beverages, Alcoholic drink mixes	2	2			
	Chocolate confectionery	30	10			
	Nuts, oilseeds and chickpeas		20			
	Frozen fermented milk-based products	15	5			
	Sausages	30	10			

▼<u>M74</u>

	Authorised novel food	Conditions under which t	he novel food may be u	sed	Additional specific labelling requirements	Other requirements	Data protection
▼ <u>M78</u>	Frozen, dried and powder forms of	Specified food category	(marketed as suc	evels (g/100g) ch or reconstituted the instructions)	Depending on the form used, the designation of the novel food on the labelling of the		Authorised on 1 March 2022. This inclusion is based on proprietary scientific evidence
	yellow mealworm (Tenebrio molitor		Frozen	Dried or powder	foodstuffs containing it shall be 'frozen yellow mealworm		and scientific data protected in accordance with Article 26 of
	larva)	Frozen, dried and powder forms of yellow mealworm (<i>Tenebrio molitor</i> larva)			(Tenebrio molitor larva)', 'dried yellow mealworm (Tenebrio molitor larva)', or 'yellow mealworm (Tenebrio molitor larva) powder'. 2. The labelling of the foodstuffs containing frozen, dried and powder forms of yellow mealworm (Tenebrio molitor larva) shall bear a statement that this ingredient may cause allergic reactions to consumers with known allergies to crustaceans and products thereof and to dust mites. This statement shall appear in close proximity to the list of ingredients.		Regulation (EU) 2015/2283. Applicant: Fair Insects BV, Industriestraat 3, 5107 NC Dongen, the Netherlands. During the period of data protection, the novel food is authorised for placing on the market within the Union only by Fair Insects BV, unless a subsequent applicant obtains authorisation for that novel food without
		Multigrain bread and rolls; crackers and breadsticks	30	10			
		Cereal bars	30	15			
		Dried pasta based products; pasta based dishes (excluding dried puffed pasta); pizza and pizza-like dishes	15	10			
		Dried stuffed pasta based products	30	15			reference to the proprietary scientific evidence or scientific
		Pre-mixes (dry) for baked products	30	15			data protected in accordance with Article 26 of Regulation
		Sauces	30	10			(EU) 2015/2283, or with the
		Potato, legumes based dishes	15	10			agreement of Fair Insects BV. End date of the date protection:
		Whey powder	40	20			1 March 2027.
		Meat analogues	80	50			
		Soups and salads	20	5			
		Chips/crisps	40	20			
		Beer-like beverages; mixed alcoholic drinks; alcoholic drink mixes	1	1			
		Chocolate confectionary	30	10			
		Nuts, oilseeds and chickpeas	40	30			
		Frozen fermented milk-based products	15	5			
		Meat preparations	40	16			

N-Acetyl-D-neuraminic acid	Description: N-Acetyl-D-neuraminic acid is a white to off-white crystalline powder
·	
	N-Activi-D-neuralinine acturis a white to on-white crystamine powder
	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:
	$C_{11}H_{19}NO_9$ (acid)
	$C_{11}H_{23}NO_{11}$ ($C_{11}H_{19}NO_9 * 2H_2O$) (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: $< 0.1 \text{ mg/kg}$
	Residual proteins: < 0,01 % (w/w)

s, an mally der).	02017R2470 - EN - 01.05.2024 - 046.001 - 146

Authorised Novel Food	Specifications
	Residual solvents:
	2-Propanol: < 0,1 % (w/w)
	Acetone: $< 0.1 \% \text{ (w/w)}$
	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.

▼ M83

Frozen, dried and powder forms of *Acheta domesticus* (house cricket)

Description/Definition:

The novel food consists of the whole, frozen, dried and powder forms of the house cricket. The term 'house cricket' refers to the adult *Acheta domesticus*, an insect species that belongs to the Gryllidae family.

The novel food is intended to be marketed in three different forms, namely: (i) thermally processed and frozen whole *A. domesticus* (AD frozen); (ii) thermally processed and freeze-dried whole *A. domesticus* (AD dried), and (iii) thermally processed freeze-dried and ground whole *A. domesticus* (whole AD powder). A minimum 24 hours fasting period is required before killing the insects by freezing, to allow the adults to discard their bowel content.

Characteristics/Composition (AD frozen):

Ash (% w/w): 0,6–1,2 Moisture (% w/w): 76–82 Crude protein (N x 6,25) (% w/w): 12–21 Digestible Carbohydrates (% w/w): 0,1–2 Fat (% w/w): 3–12

of which saturated (% w/w): 36-45

Characteristics/Composition (AD dried or powder):

Ash (% w/w): 2,9–5,1

Moisture (% w/w): ≤ 5

Crude protein (N x 6,25) (% w/w): 55–65

Digestible Carbohydrates (% w/w): 1–4

Fat (% w/w): 29–35

of which saturated (% w/w): 36–45

▼<u>M83</u>

Authorised Novel Food	Specifications	
	Peroxide value (Meq O ₂ /kg fat): ≤ 5 Dietary fibre (% w/w): 0,8–3 (¹8)Chitin (% w/w): 0,7–3,0 Heavy metals: Lead: ≤ 0,05 mg/kg Cadmium: ≤ 0,06 mg/kg Mycotoxins: Aflatoxins (Sum of B1, B2, G1, G2): ≤ 4 μg/kg Aflatoxin B1 (μg/kg): ≤ 2 Deoxynivalenol: ≤ 200 μg/kg Ochratoxin A: ≤ 1 μg/kg Dioxins and dioxin like PCBs	Peroxide value (Meq O_2 /kg fat): ≤ 5 Dietary fibre (% w/w): 3–6 (18)Chitin (% w/w): 5,3-10,0 Heavy metals: Lead: $\leq 0,05$ mg/kg Cadmium: $\leq 0,06$ mg/kg Mycotoxins: Aflatoxins (Sum of B1, B2, G1, G2): ≤ 4 µg/kg Aflatoxin B1 (µg/kg): ≤ 2 Deoxynivalenol: ≤ 200 µg/kg Ochratoxin A: ≤ 1 µg/kg Dioxins and dioxin like PCBs
	Sum of dioxins and dioxin-like PCBs UB, ((¹¹)WHO₂₀₀₅ PCDD/F-PCB-TEQ): ≤ 1,25 pg/g fat Microbiological criteria: Total aerobic colony count: ≤ 10⁵ (²)CFU/g Yeasts and moulds: ≤ 100 CFU/g Escherichia coli: ≤ 50 CFU/g Salmonella spp.: Absence in 25 g Listeria monocytogenes: Absence in 25 g Sulfite-reducing Anaerobes: ≤ 30 CFU/g Bacillus cereus (presumptive): ≤ 100 CFU/g Enterobacteriaceae (presumptive): < 100 CFU/g Coagulase-positive staphylococci: ≤ 100 CFU/g	Sum of dioxins and dioxin-like PCBs UB, ((¹¹)WHO₂₀₀₅ PCDD/F-PCB-TEQ): ≤ 1,25 pg/g fat Microbiological criteria: Total aerobic colony count: ≤ 10⁵ CFU/g Yeasts and moulds: ≤ 100 CFU/g Escherichia coli: ≤ 50 CFU/g Salmonella spp.: Absence in 25 g Listeria monocytogenes: Absence in 25 g Sulfite-reducing Anaerobes: ≤ 30 CFU/g Bacillus cereus (presumptive): ≤ 100 CFU/g Enterobacteriaceae (presumptive): < 100 CFU/g Coagulase-positive staphylococci: ≤ 100 CFU/g

	Authorised Novel Food	Specifications
▼ M99		
	Acheta domesticus (house cricket) partially defatted powder	Description/Definition : The novel food is partially defatted powder obtained from whole <i>Acheta domesticus</i> (house cricket) following a series of steps involving a 24 hours fasting period of the insects to allow them to discard their bowel content, the sacrifice of the insects by freezing, washing, thermal processing, drying, oil extraction (mechanical extrusion), and grinding.
		Characteristics/Composition:
		Crude protein (N x 6,25) (% w/w): 74,0 - 78,0
		Fat (% w/w): 9,0 – 12,0
		Moisture (% w/w): $3.0 - 6.0$
		Crude fibre (% w/w): 8,0 – 10,0
		Chitin (22) (% w/w): 4,0-8,5
		Ash (% w/w): ≤ 5.6
		Peroxide value (Meq O_2/kg fat): ≤ 5.0
		Manganese: ≤ 100,0 mg/kg
		Cyanide: $\leq 5.0 \text{ mg/kg}$
		Heavy metals:
		Lead: $\leq 0.1 \text{ mg/kg}$
		Cadmium: $\leq 0.025 \text{ mg/kg}$
		Mycotoxins:
		Aflatoxins (Sum of B1, B2, G1, G2): $\leq 0.4 \mu g/kg$
		Deoxynivalenol: ≤ 200,0 μg/kg
		Ochratoxin A: ≤ 1,0 μg/kg
		Dioxins and dioxin like PCBs:
		Sum of dioxins and dioxin-like PCBs UB, ((23)WHO ₂₀₀₅ PCDD/F-PCB-TEF): ≤ 1,25 pg/g fat
		Microbiological criteria:
		Total aerobic microbial count: $\leq 10^5$ CFU/g
		Yeasts and moulds: ≤ 100 CFU/g
		Escherichia coli: ≤ 50 CFU/g
		Salmonella spp.: Not detected in 25 g
		Listeria monocytogenes: Not detected in 25 g
		Bacillus cereus (presumptive): ≤ 100 CFU/g
		Enterobacteriaceae (presumptive): < 100 CFU/g
		Coagulase-positive staphylococci: ≤ 100 CFU/g

	Adansonia digitata (Baobab) dried	Description/Definition:
	fruit pulp	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	Description/Definition:
	cultures	Hydroalcoholic extract from Ajuga reptans L. tissue cultures which is substantially equivalent to extracts from flowering aerial parts of Ajuga reptans obtained
		by traditional cultures.
▼ <u>M80</u>		
	Akkermansia muciniphila	Description:
	(pasteurised)	Pasteurised Akkermansia muciniphila (strain ATCC BAA-835, CIP 107961) is produced by anaerobic growth of the bacteria followed by pasteurisation,
		concentration of the cells, cryopreservation, and freeze drying.
		Characteristics/Composition:
		Total A. muciniphila cell count (cells/g): 2.5×10^{10} to 2.5×10^{12}
		Viable A. muciniphila cell count (CFU/g): < 10 (LoD)(*)
		Water activity: ≤ 0.43
		Moisture (%): $\leq 12,0$
		Protein (%): $\leq 35,0$
		Fat $(\%)$: $\leq 4,0$
		Crude ash (%): $\leq 21,0$
		Carbohydrates (%): 36,0 – 86,0
		Microbiological criteria: Aerobic mesophilic total count: ≤ 500 CFU(**)/g
		Sulphite reducing anaerobes: ≤ 50 CFU/g
		Coagulase ⁺ Staphylococci: ≤ 10 CFU/g
		Enterobacteriaceae: ≤ 10 CFU/g

Specifications

Authorised Novel Food	Specifications
	Yeast: ≤ 10 CFU/g
	Mould: $\leq 10 \text{ CFU/g}$
	Bacillus cereus: ≤ 100 CFU/g Listeria spp.: Absence in 25 g
	Salmonella spp.: Absence in 25 g
	Escherichia coli: Absence in 1 g
	(*) LoD: Limit of Detection;
	(**) Colony Forming Units.
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: $\leq 0.1 \%$
	Loss on drying: $\leq 0.5\%$ Optical rotation: $+9.0 - +11.0^{\circ}$
	pH (1 %; H ₂ O): 5,0-6,0
	Ammonium (NH ₄): $\leq 0.020 \%$
	Chloride (Cl): $\leq 0.020\%$
	Sulphate (SO ₄): $\leq 0.020 \%$
	Microbiological criteria:
	Escherichia coli: Absence/g
Algal oil from the microalgae	Description/Definition:
Ulkenia sp.	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: $\leq 0.05 \%$
	Unsaponifiables: ≤ 4,5 %
	Trans-fatty acids: $\leq 1.0 \%$
	DHA content: ≥ 32 %

02017F
7R2470 -
— EN
01.0
05.2024
1 - 04
6.001 -
-151

V 1/1/2		
	Authorised Novel Food	Specifications
▼ <u>M26</u>		
	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
▼ <u>M9</u>		
	Aloe macroclada Baker leaf extract	Description/Definition:
		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 %
		Glucose: 8,9 %
- 3.510		

▼M103

Frozen, paste, dried and powder forms of *Alphitobius diaperinus* larvae (lesser mealworm)

Description/Definition:

The novel food consists of the frozen, paste, dried, and powder forms of the whole lesser mealworm. The term 'lesser mealworm' refers to the larval form of *Alphitobius diaperinus*, an insect species that belongs to the family of *Tenebrionidae* (darkling beetles).

The entire lesser mealworms are meant for human consumption, no parts are removed.

The novel food is intended to be marketed in 4 different forms, namely: (i) whole blanched and frozen *A. diaperinus* larvae (ADL frozen), (ii) paste from whole blanched, ground, and frozen *A. diaperinus* larvae (ADL paste), (iii) whole blanched, and freeze-dried *A. diaperinus* larvae (ADL dried), and (iv) powder from whole blanched, freeze-dried and ground *A. diaperinus* larvae (ADL powder).

A minimum 24 hours fasting period is required to allow the larvae to discard their bowel content before killing the insects by a thermal treatment.

Authorised Novel Food	Specifications		
	Characteristics/Composition (ADL frozen or paste):	Characteristics/Composition (ADL dried or powder):	
	Ash (% w/w): ≤ 1.5	Ash (% w/w): ≤ 5	
	Moisture (% w/w): 65-80	Moisture (% w/w): 1-5	
	Crude protein (N × 6,25) (% w/w): 12-25	Crude protein (N × 6,25) (% w/w): 50-70	
	Digestible Carbohydrates (% w/w): 0,4-2	Digestible Carbohydrates (% w/w): 1,5-3,5	
	Fat (% w/w): 5-12	Fat (% w/w): 20-35	
	Peroxide value (Meq O_2/kg fat): ≤ 0.2	Peroxide value (Meq O_2/kg fat): ≤ 5	
	Dietary fibre (% w/w): 1-4	Dietary fibre (% w/w): 3-6	
	(27) Chitin (% w/w): 1,0-2,6	(27) Chitin (% w/w): 3,0-9,1	
	Heavy metals:	Heavy metals:	
	Lead: ≤ 0,1 mg/kg	Lead: ≤ 0,1 mg/kg	
	Cadmium: ≤ 0,05 mg/kg	Cadmium: ≤ 0,05 mg/kg	
	Mycotoxins:	Mycotoxins:	
	Aflatoxins (Sum of B1, B2, G1, G2): $\leq 4 \mu g/kg$	Aflatoxins (Sum of B1, B2, G1, G2): ≤ 4 μg/kg	
	Aflatoxin B1 (μ g/kg): ≤ 2	Aflatoxin B1 (μ g/kg): ≤ 2	
	Deoxynivalenol: ≤ 200 μg/kg	Deoxynivalenol: ≤ 200 μg/kg	
	Ochratoxin A: ≤ 1 µg/kg	Ochratoxin A: ≤ 1 µg/kg	
	Microbiological criteria:	Microbiological criteria:	
	Total aerobic colony count: $\leq 10^5$ (25) CFU/g	Total aerobic colony count: ≤ 10 ⁵ CFU/g	
	Yeasts and moulds: ≤ 100 CFU/g	Yeasts and moulds: ≤ 100 CFU/g	
	<i>Escherichia coli</i> : ≤ 50 CFU/g	Escherichia coli: ≤ 50 CFU/g	
	Salmonella spp.: Absence in 25 g	Salmonella spp.: Absence in 25 g	
	Listeria monocytogenes: Absence in 25 g	Listeria monocytogenes: Absence in 25 g	
	Sulphite-reducing Anaerobes: ≤ 30 CFU/g	Sulfite-reducing Anaerobes: ≤ 30 CFU/g	
	Bacillus cereus: ≤ 100 CFU/g	Bacillus cereus: ≤ 100 CFU/g	
	Enterobacteriaceae: ≤ 100 CFU/g	Enterobacteriaceae: ≤ 100 CFU/g	
	Coagulase-positive <i>staphylococci</i> : ≤ 100 CFU/g	Coagulase-positive staphylococci: ≤ 100 CFU/g	

	Authorised Novel Food	Specifications
<u>M24</u>		
	Antarctic Krill oil from Euphausia	Description/Definition:
	superba	To produce lipid extract from Antarctic Krill (<i>Euphausia superba</i>) deep-frozen crushed krill or dried krill meal is subjected to lipid extraction with a approved extraction solvent (under Directive 2009/32/EC). Proteins and krill material are removed from the lipid extract by filtration. The extraction solven and residual water are removed by evaporation.
		Saponification value: ≤ 230 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 <i>Euphausia superba</i> should demonstrate oxidative stability by appropriate as recognised national/international test methodology (e.g. AOAC).
		Moisture and volatiles: ≤ 3 % or 0,6 expressed as water activity at 25 °C
		Phospholipids: ≥ 35 % to < 60 %
		Trans-fatty acids: ≤ 1 %
		EPA (eicosapentaenoic acid): ≥ 9 %
		DHA (docosahexaenoic acid): ≥ 5 %
М9		
	Antarctic Krill oil rich in phosp-	Description/Definition:
	holipids from Euphausia superba	Oil rich in phospholipids is produced from Antarctic krill (<i>Euphausia superba</i>) by repeated solvent washings with an approved solvent (under Directive 200 32/EC) to increase phospholipid content of the oil. Solvents are removed from the final product by evaporation.
		Saponification value: ≤ 230 mg KOH/g
		Peroxide value (PV): $\leq 3 \text{ meq } O_2/kg \text{ oil}$
		Moisture and volatiles: ≤ 3 % or 0,6 expressed as water activity at 25 °C
		Phospholipids: ≥ 60 %
		Trans-fatty acids: ≤ 1 %
		EPA (eicosapentaenoic acid): ≥ 9 %
		DHA (docosahexaenoic acid): ≥ 5 %

	0
	2
	0
	1
	17
	4
	(3
	17R247
	-
	7
) —]
	E
	\neg
	-
	(
)]
	1.0
	.05.2
	5
	<i>i</i> ,
	\simeq
	5.2024
	7,
	_
	0
	4
	6.
	$\overline{}$
	00
)1
	1.
	ž
ı	4

▼ <u>M97</u>			
	Antrodia	camphorata	mvcelia

powder

Authorised Novel Food

Specifications

Description/Definition:

The novel food is the freeze-dried mycelia of the fungus *Antrodia camphorata* (strain BCRC 39106), which has been grown by solid-state cultivation. The freeze-dried mycelia are then milled into a powder. *Antrodia camphorata* is a synonym of *Taiwanofungus camphoratus* (family: Fomitopsidaceae).

Characteristics/Composition:

Loss on drying (Moisture): < 10 %

Carbohydrates: ≤ 80 g/100 g

Protein: $\leq 20 \text{ g}/100 \text{ g}$

 $Ash: \leq 6~g/100g$

Fat: \leq 6 g/100 g

Total triterpenoids: 1,0 - 10,0 g/100 g

Antroquinonol: 1,0 - 20,0 mg/g

Heavy metals:

Arsenic: < 0,5 mg/kg

Microbiological criteria:

Total aerobic microbial count: ≤ 10³ *CFU/g
Total yeast and mould count: ≤ 100 CFU/g
Escherichia coli: Not detected in 10 g
Salmonella spp.: Not detected in 25 g
Staphylococcus aureus: Not detected in 10 g

*CFU: Colony Forming Units

▼M120

Aqueous ethanolic extract of Labisia pumila

Description/Definition:

The novel food is a hydroalcoholic extract obtained from a dried whole plant of Labisia pumila (Blume) Fern.-Vill.

The production process of the novel food starts with washing, drying and grinding of the plant *Labisia pumila*. The ground plant material is then extracted twice with a mixture of water and ethanol (50/50 v/v). The liquid extract is then concentrated, mixed with maltodextrin (which is used as a drying aid) in a ratio of 2:1 and spray-dried.

▼<u>M120</u>

Aı	thorised Novel Food	Specifications
		Characteristics/composition (including maltodextrin):Particle size: > 90 % through 120 mesh (125 µm)
		Ash: < 10 %
		Acid-insoluble ash: < 1 %
		Moisture: < 8 %
		Ethanol: $< 1 \% (w/w)$
		Gallic acid: 2-10 % (w/w)
		Carbohydrate: 70-90 g/100 g
		Protein: $< 9 \% (w/w)$
		Total fat: $< 3 \% (w/w)$
		Saponin (as ardisiacripsin A): < 1,5 % (w/w)
		Microbiological criteria:
		Aerobic plate count: $< 1 \times 10^4 \text{ CFU/g}$
		Yeast and mould: $< 5 \times 10^2 \text{ CFU/g}$
		E. coli: not detected in 10 g
		S.aureus: not detected in 10 g
		Salmonella: not detected in 25 g
		P. aeruginosa: not detected in 10 g
		cfu: colony forming units
		w/w: weight per weight
▼ <u>M128</u>		
Annle frui	t cell culture biomass	Description/Definition:
прри пи	t cen culture bioliuss	The novel food is a biomass of cultivated and homogenised cells of the Swiss apple variety Uttwiler Spätlauber (Malus domestica Borkh.).
		The production process consists of collecting under sterile conditions specific sections of the apple, which are then placed on solid medium with the aim to induce the formation of a primary callus tissue comprised of dedifferentiated cells under sterile conditions. The callus cells are then cultivated in liquid medium and subsequently homogenised, heat treated and dried.

▼<u>M128</u>

	Authorised Novel Food	Specifications
		Characteristics/composition:
		Moisture: 10,9–15,5 g/100 g
		Ash: 11,8–20,8 g/100 g
		Proteins: 14,3–20,0 g/100 g
		Fats: 0,6–2,5 g/100 g
		Non-digestible carbohydrates: 17,1–25,2 g/100 g
		Other carbohydrates (calculated (29)): 21,9–38,9 g/100 g
		Total sugars: 17,1–32,6 g/100g
		Fructose: 10,8–20,2 g/100 g
		Glucose: 3,8–7,0 g/100 g
		Total phenols: 0,15-0,29 g/100 g
		Malic acid: 0,41-1,19 g/100 g
		Succinic acid: 0,14–0,26 g/100 g
▼ <u>M9</u>		
	Arachidonic acid-rich oil from the fungus <i>Mortierella alpina</i>	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 <i>Mortierella alpina</i> 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: ≤ 0,5 % of the total fatty acid content
		Unsaponifiable matter: ≤ 1,5 %
		Peroxide value (PV): ≤ 5 meq/kg
		Anisidin value: ≤ 20
		Acid value: ≤ 1,0 KOH/g
		Moisture: ≤ 0,5 %

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 <i>Argania spinosa</i> (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): < 10 meq O ₂ /kg
<u></u>	
Astaxanthin-rich oleoresin from	
	Description:
	Description: Astaxanthin is a carotenoid produced by <i>Haematococcus pluvialis</i> algae. Production methods for the growth of the algae are variable; using '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).
	Astaxanthin is a carotenoid produced by <i>Haematococcus pluvialis</i> algae. Production methods for the growth of the algae are variable; using '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 %
Astaxanthin-rich oleoresin from Haematococcus pluvialis algae	Astaxanthin is a carotenoid produced by <i>Haematococcus pluvialis</i> algae. Production methods for the growth of the algae are variable; using '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).
	Astaxanthin is a carotenoid produced by <i>Haematococcus pluvialis</i> algae. Production methods for the growth of the algae are variable; using '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). Characteristics/Composition:
	Astaxanthin is a carotenoid produced by <i>Haematococcus pluvialis</i> algae. Production methods for the growth of the algae are variable; using '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). Characteristics/Composition: Fat: 42,2-99 %
	Astaxanthin is a carotenoid produced by <i>Haematococcus pluvialis</i> algae. Production methods for the growth of the algae are variable; using '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 CO₂ 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). Characteristics/Composition: Fat: 42,2-99 % Protein: ≤ 4,4 %
	Astaxanthin is a carotenoid produced by <i>Haematococcus pluvialis</i> algae. Production methods for the growth of the algae are variable; using '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 or MCT (Medium Chain Triglycerides). Characteristics/Composition: Fat: 42,2-99 % Protein: $\leq 4,4$ % Carbohydrate: $\leq 52,8$ %
	Astaxanthin is a carotenoid produced by <i>Haematococcus pluvialis</i> algae. Production methods for the growth of the algae are variable; using '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). Characteristics/Composition: Fat: 42,2-99 % Protein: $\leq 4,4$ % Carbohydrate: $\leq 52,8$ % Fibre: $< 1,0$ %

9-cis-astaxanthin: 0,3-30,0 %

Specifications

▼<u>M131</u>

	Authorised Novel Food	Specifications
		13-cis-astaxanthin: 0,2-7,0 %
		Astaxanthin monoesters: 66,7-91,5 %
		Astaxanthin diesters: 0,16-32,5 %
		Beta-Carotene: 0,01-0,3 %
		Lutein: ≤ 1,8 %
		Canthaxanthin: ≤ 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
▼ <u>M12</u>	9	
	Partially hydrolysed protein from	Description/Definition:
	spent barley (Hordeum vulgare) and rice (Oryza sativa)	The novel food is partially hydrolysed protein from spent barley (<i>Hordeum vulgare</i>) and rice (<i>Oryza sativa</i>), residues obtained from the solid by-product of beer production that contains 45-70 % spent barley and 30-55 % spent rice.
		The novel food is produced by enzymatically treating the pasteurised spent barley and rice residues of the mash step of beer production. Several mechanical treatment steps of the partial hydrolysate are employed to obtain the final product.
		Characteristics/composition:
		Appearance: powder
		Degree of hydrolysis: 1-7 %
		Proteins (N x 6,25): 78-90 %
		Moisture: 2-8 %
		Carbohydrates: 2-10 %

▼ <u>M129</u>

Authorised Novel Food	Specifications
	Fat: 0-2 %
	Ash: 1-8 %
	Heavy metals:
	Arsenic (mg/kg): ≤ 0.2
	Cadmium (mg/kg): ≤ 0.1
	Lead (mg/kg): ≤ 0.2
	Mercury (mg/kg): ≤ 0.01
	Myaatayinga
	Mycotoxins: Aflatoxin B1: ≤ 2 μg/kg
	Sum of aflatoxins (B1, B2, G1, G2): $\leq 4 \mu g/kg$
	Deoxynivalenol: < 200 μg/kg
	Fumonisins (sum of B1, B2): $\leq 200 \mu g/kg$
	Ochratoxin A: $\leq 3 \mu g/kg$
	Zearalenone: $\leq 20 \mu g/kg$
	Patulin: ≤ 50 µg/kg
	Antinutritional factors:
	Phytic acid: < 0,25 %
	Microbiological criteria:
	Total aerobic microbial count (CFU/g): < 10 ⁴
	Coliforms (CFU/g): < 100
	Total yeast and mould count (CFU/g): < 100
	Salmonella spp.: Not detected in 25 g
	Escherichia coli (CFU/g): < 10
	Staphylococcus aureus (CFU/g): < 10
	Listeria monocytogenes: Not detected in 25 g
	Bacillus cereus (CFU/g): < 100 CFU: colony forming units
	CFO. Colony forming units

n-GM	
ted by ne pH	

Authorised Novel Food	Specifications
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 %

▼M134

Beta-glucan from *Euglena gracilis* microalgae

Description/Definition:

The novel food, beta-glucan from Euglena gracilis microalgae (paramylon), is a linear, unbranched beta-1,3-D-glucan polymer derived from the non-GM microalga Euglena gracilis.

The novel food is produced by fermentation, followed by pH adjustment and homogenization to release the beta-glucan granules. The granules are isolated by decanting and washing, and subsequently, acidified and filtered. After drying, the product is milled. The process includes conditions such as an alkaline pH and heat-killing step of the microalga to ensure the absence of viable *Euglena gracilis* cells in the novel food.

Characteristics/composition:

Appearance: cream white powder

Beta-glucan (30): (%) ≥ 95

Moisture (%): ≤ 6

Ash (%): ≤ 1

Heavy metals:

Lead (mg/kg): ≤ 0.5

Cadmium (mg/kg): ≤ 0.5

Mercury (mg/kg): ≤ 0.05

Arsenic (mg/kg): ≤ 0.02

▼M134

Authorised Novel Food	Specifications
	Microbiological criteria:
	Total aerobic microbial count (CFU/g): ≤ 3 000
	Total yeast and mould count (CFU/g): ≤ 100
	Coliforms (MPN/g): ≤ 30
	Escherichia coli: Not detected in 10 g
	Staphylococcus aureus: Not detected in 10 g
	Salmonella spp.: Not detected in 25 g
	Listeria monocytogenes: Not detected in 25 g
	CFU: colony forming units, MPN: most probable number.
3	

▼ M33

Betaine

Description/Definition:

Betaine (N,N,N-trimethylglycine or carboxy-N,N,N-trimethylmethanaminium), in anhydrous (CH_3)₃N⁺ CH_2COO^- (CAS No: 107-43-7) and monohydrate (CH_3)₃N⁺ CH_2COO^- . H_2O (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: ≥ 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: ≤ 1,0 mg/g

Heavy metals:

Arsenic: < 0,1 mg/kg

Mercury: < 0,005 mg/kg

Cadmium: < 0,01 mg/kg

Lead: < 0,05 mg/kg

▼<u>M33</u>

•	Authorised Novel Food	Specifications
·		Microbiological criteria:
		Total viable count: ≤ 100 CFU/g
		Coliforms: Negative/10 g
		Salmonella sp.: Negative/25 g
		Yeast: ≤ 10 CFU/g
		Mould: $\leq 10 \text{ CFU/g}$
		CFU: Colony Forming Units.
▼ <u>M9</u>		
]	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: ≤ 1,0 %
		Protein: ≥ 55 %
		Water: ≤ 7,0 %
		Ash: ≤ 10 %
		Carbohydrate: ≥ 20 %
		α-glucosidase inhibitory activity: IC50 min 0,025 mg/ml
		Soy isoflavone: $\leq 0.3 \text{ g/}100 \text{ g}$

Description/Definition:

Bovine lactoferrin

		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
▼ <u>M35</u>		
	Bovine milk basic whey protein	Description
	isolate	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

Specifications

▼<u>M35</u>

	Authorised Novel Food	Specifications
		Lactose: ≤ 3,0 %
		Fat: ≤ 4,5 %
		$Ash: \leq 3.5\%$
		Iron: $\leq 25 \text{ mg/}100 \text{ g}$
		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: ≤ 50 CFU/g
		Yeasts: $\leq 50 \text{ CFU/g}$
		CFU: Colony Forming Units
▼ <u>M96</u>		
	Bovine milk beta-lactoglobulin	Description:
	(β-lactoglobulin)	Beta-lactoglobulin (β-lactoglobulin) protein is a white to cream powder produced from bovine whey by a series of steps involving filtration, concentration,
		crystallisation, re-dissolution (in water), pH adjustment to acidic or neutral pH, re-concentration and drying.
		CAS number: 9045-23-2
		Molecular weight: 36,7 kDa (dimer); 18,3 kDa (monomer)

Authorised Novel Food	Specifications
	Characteristics/Composition:
	pH (10 % solution): 3,5-8,0
	Protein (N x 6,38) (%): \geq 86,0
	Beta-lactoglobulin (% of protein): ≥ 90,0
	Lactose (%): $\leq 1,0$
	Fat $(\%)$: $\leq 1,0$
	Ash (%): ≤ 5.0
	Moisture (%): ≤ 5.5
	Heavy Metals:
	Cadmium (mg/kg): < 0,2
	Lead (mg/kg): < 0,1
	Mercury (mg/kg): < 0,01
	Contaminants:
	Aflatoxin M1 (μg/kg): < 0,01
	Microbiological criteria:
	Total plate count: ≤ 5 000 CFU/g
	Total yeast/moulds count: ≤ 10 CFU/g
	Enterobacteriaceae: ≤ 10 CFU/g
	Salmonella spp.: Absent in 25 g
	Bacillus cereus: < 100 CFU/g
	Listeria monocytogenes: Absent in 25 g
	Staphylococcus aureus: < 10 CFU/g
	Sulfite-reducing clostridia: < 10 CFU/g
	CFU: Colony Forming Units; kDa: kiloDaltons
M107	
Bovine milk osteopontin	Description
•	Bovine milk osteopontin is isolated from pasteurised or microfiltered bovine whey or milk by ion exchange chromatography, ultrafiltration to remove low molecular weight components and spray drying. During this filtration steps lactose and whey proteins predominantly alpha-lactalbumin and beta lactoglobulin are removed.
	Characteristics/Composition
	Protein % as is $(N \times 6,38)$: $76,5-80,5$
	Bovine milk osteopontin (bmOPN) (% of protein): ≥ 84,5

▼<u>M107</u>

ull-length bmOPN (MW 33,9 kDa) (% of bmOPN): ≥ 15 I-terminal fragment bmOPN (MW 19,8 kDa) (% of bmOPN): ≥ 70 Other milk protein (% of protein): ≤ 14,5 Moisture: < 9,5 % actose: ≤ 1,0 % act: ≤ 1,0 % ach: ≤ 11 % ansolubility index (mL) ≤ 1,0 Meavy metals acad: < 0,05 mg/kg admium: < 0,05 mg/kg Mercury: < 0,05 mg/kg Arrecury: < 0,05 mg/kg
other milk protein (% of protein): $\leq 14,5$ Moisture: $< 9,5$ % $< actose: \leq 1,0 % < act: \leq 1,0 % < ac: \leq 1,0 % < ac:$
Moisture: < 9,5 % Moisture: < 1,0 % Moisture: ≤
cactose: $\leq 1,0\%$ cat: $\leq 1,0\%$ cash: $\leq 11\%$ cashsolubility index (mL) $\leq 1,0$ leavy metals cead: $< 0,05 \text{ mg/kg}$ cadmium: $< 0,05 \text{ mg/kg}$
at: ≤ 1,0 % ash: ≤ 11 % asolubility index (mL) ≤ 1,0 leavy metals lead: < 0,05 mg/kg Cadmium: < 0,05 mg/kg Mercury: < 0,05 mg/kg
ash: ≤ 11 % asolubility index (mL) $\leq 1,0$ Heavy metals acad: $< 0,05$ mg/kg Cadmium: $< 0,05$ mg/kg Mercury: $< 0,05$ mg/kg
nsolubility index (mL) \leq 1,0 Leavy metals Lead: < 0,05 mg/kg Cadmium: < 0,05 mg/kg Mercury: < 0,05 mg/kg
Heavy metals Lead: < 0,05 mg/kg Cadmium: < 0,05 mg/kg Mercury: < 0,05 mg/kg
Lead: < 0,05 mg/kg Cadmium: < 0,05 mg/kg Mercury: < 0,05 mg/kg
Cadmium: < 0,05 mg/kg Mercury: < 0,05 mg/kg
Mercury: < 0,05 mg/kg
vrsenic: < 0.5 mg/kg
flatoxin M1 $< 0,1 \mu g/kg$
Aicrobiological criteria
otal plate count (30 °C) (CFU/g): ≤ 5000
$fould/yeast (CFU/g): \le 100$
Pacillus cereus (CFU/g): < 50
ulfur-reducing Clostridia (CFU/g): < 10
taphylococcus aureus: Not detected in 1 g
interobacteriaceae (CFU/g): < 10
almonella spp.: Not detected in 25 g
FU: Colony Forming Units
Description/Definition:
defined Buglossoides oil is extracted from the seeds of <i>Buglossoides arvensis</i> (L.) I.M.Johnst
Alpha-linolenic acid: ≥ 35 % w/w of total fatty acids
tearidonic acid: ≥ 15 % w/w of total fatty acids
inoleic acid: ≥ 8,0 % w/w of total fatty acids
MOREC ACIO: < 0.0 % W/W OF IOIAL PARTY ACIOS
Tio Tot Mo Pac ultap Ent Tali CFU

	Authorised Novel Food	Specifications
		Acid value: ≤ 0,6 mg KOH/g
		Peroxide value (PV): ≤ 5.0 meq O_2/kg
		Unsaponifiable content: $\leq 2.0 \%$
		Protein content (total nitrogen): $\leq 10 \mu \text{g/ml}$
		Pyrrolizidine alkaloids: Not detectable with a detection limit of 4,0 µg/kg
▼ <u>M91</u>		
	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) <i>Calanus finmarchicus</i> . 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 %
		Astaxanthin esters: ≤ 0,25 %
		Peroxide value (PV): < 3.0 meq. O_2/kg
▼ <u>M77</u>		
	Calcium fructoborate	Description/Definition
		The novel food is calcium fructoborate, a calcium salt tetrahydrate of a bis(fructose) ester of boric acid in the form of a powder, represented by Ca[(C6H10O6)2B]2•4H2O, with a molecular mass of 846 Da.

Authorised Novel Food	Specifications
	The novel food is produced by chemical synthesis whereby fructose is combined with boric acid in water to produce a bis(fructose) ester of boric acid through various heating and mixing processes. Calcium carbonate is then added to produce a solution containing the calcium salt of fructoborate (tetrahydrate). The solution is freeze-dried, ground to produce the final powdered product, and then packaged and stored under representative storage conditions (22 ± 1°C RH 55-60 %).
	Characteristics/composition
	Free moisture: < 5,0 %
	Calcium: 4,5-5 %
	Boron: 2,5-2,9 %
	Fructose: 80-85 %
	Ash: 15-16 %
	Heavy metals
	Arsenic: ≤ 1 mg/kg
	Microbiological criteria
	Total plate count: $\leq 1~000~\mathrm{CFU/g}^{\mathrm{(a)}}$
	Yeast and mould: < 100 CFU/g
	Coliforms: ≤ 10 CFU/g
	Escherichia coli: < 10 CFU/g
	Salmonella spp.: Absence in 25 g
	Coagulase-positive staphylococci: Absence in 1 g
	(a) CFU: colony forming units
M85	
Calcium L-Methylfolate	Description:
	The novel food is produced by chemical synthesis starting from folic acid.
	It is a white to light yellowish, almost odourless, crystalline powder, sparingly soluble in water and very slightly soluble or insoluble in most organic solvents.
	Definition:
	Chemical formula: C ₂₀ H ₂₃ CaN ₇ O ₆

02
020
)1
7]
R
24
17
0
E
Z
0
01.05.2024
0
5.
2
)5.2024
4
0
4
046.001
00
1
1
69
_

Systematic name: N-{4-[[((6S)-2-amino-1,4,5,6,7,8-hexahydro-5-methyl-4-oxo-6-pteridinyl)methyl]amino]benzoyl}-L-glutamic acid, calcium salt.

CAS Numbers: 129025-21-4 (Calcium salt with an unspecified ratio of L-5-MTHF/Ca²⁺) and 151533-22-1 (Calcium salt with specified 1:1 ratio of L-5-MTHF/Ca²⁺).

Specifications

Molecular weight: 497,5 Daltons

Synonyms: L-methylfolate, calcium; L-5-methyltetrahydrofolic acid, calcium salt [(L-5-MTHF-Ca)]; (6S)-5-methyltetrahydrofolic acid, calcium salt [(6S)-5-MTHF-Ca]; (6S)-5-methyl-5,6,7,8-tetrahydropteroyl-L-glutamic acid, calcium salt, and L-5-methyl-tetrahydrofolic acid (L-5-MTHF) without the cation specified.

Structural formula:

Characteristics

Purity: > 95 % (Dry basis)

Water: ≤ 17,0 %

Calcium (on anhydrous and solvent free basis): 7,0 - 8,5 %

Calcium D-methylfolate (6R, α S isomer): $\leq 1,0 \%$

Other folates and related substances: ≤ 2,5 %

Ethanol: $\leq 0.5 \%$

Contaminants

CFU: colony forming units

02017R2470
EN01.05.2024

<u>M137</u>	

Calcidiol monohydrate

Authorised Novel Food

Description/Definition:

Infants and young children

Lead: $\leq 1 \text{ mg/kg}$

Boron: $\leq 10 \text{ mg/kg}$

Cadmium ≤ 0.5 mg/kg

Mercury ≤ 1.0 mg/kg

Arsenic ≤ 1.5 mg/kg

Platinum $\leq 2 \text{ mg/kg}$

Microbiological criteria:

Total viable aerobic counts: ≤ 1 000 CFU/g Total yeast and mould count: ≤ 100 CFU/g

The novel food is calcidiol monohydrate (25-hydroxycholecalciferol monohydrate). The novel food contains the monohydrate form of the major circulating metabolite of vitamin D_3 in the body and is a source of 1,25-dihydroxyvitamin D, the biologically active form of vitamin D.

Specifications

General population excluding infants and young

Conversion factor: 1 μ g calcidiol = 2,5 μ g vitamin D₃ for doses up to 10 μ g/day.

children

Lead: $\leq 1 \text{ mg/kg}$

Boron: $\leq 10 \text{ mg/kg}$

Cadmium ≤ 0.5 mg/kg

Mercury $\leq 1.5 \text{ mg/kg}$

Arsenic $\leq 1.5 \text{ mg/kg}$

Platinum $\leq 10 \text{ mg/kg}$

The production process of the novel food starts with a yeast fermentation which results in a mixture of sterols, with trienol being the major sterol obtained. After the fermentation, purification and several chemical steps follow. The chemical steps include saponification and extraction, where the trienol is isolated from the biomass. This is followed by a hydroxylation step to separate the trienol from the other sterols. Trienol is then epoxidised and subsequently reduced to give 25-hydroxydehydrocholesterol. A photoreaction follows, to obtain a mixture of 25-hydroxy-previtamin D₃, 25-hydroxy-tachysterol and 25-hydroxy-lumisterol. Thereafter, the 25-hydroxy-previtamin D₃ is thermally isomerised to "Calcidiol" and recrystallized to obtain the novel food of the required purity.

The novel food is intended to be placed on the market as a diluted form "0,25 % w/w", containing 0,250-0,275 % w/w of calcidiol (anhydrous). The novel food needs to be placed on the market in a preparation guaranteeing its stability.

Chemical name according to IUPAC:

(1S,3Z)-3-[(2E)-2- $[(1R,3\alpha S,7\alpha R)$ -1-[(2R)-6-hydroxy-6-methylheptan-2-yl]-7 α -methyl-2,3,3 α ,5,6,7-hexahydro-1H-inden-4-ylidene]ethylidene]-4-methylidene-cyclohexan-1-ol; hydrate

CAS Number: 63283-36-3 (Calcifediol monohydrate)

Empirical formula: C₂₇H₄₄O₂.H₂O Molecular weight: 418,7 g/mol

▼<u>M137</u>

	Authorised Novel Food	Specifications
		Characteristics/composition:
		25(OH)D ₃ .H ₂ O: 97,0-100 %
		Total related substances: ≤ 1.5 %, of which: Δ^{22} -25(OH)D ₃ : ≤ 0.5 %; Lumisterol (31): ≤ 0.5 %; pre-25(OH)D ₃ (32): ≤ 0.5 %; Tachysterol (33): ≤ 0.5 %; trans-Vitamin D ₃ (34): ≤ 0.5 %
		Other impurities: $\leq 0.10\%$
		Water content: 3,8-5,0 %
		Acetone: ≤ 1 000 mg/kg
		Isopropanol: ≤ 10 mg/kg
		Heavy metals:
		Arsenic: ≤ 1 mg/kg
T M106		
▼ <u>M106</u>	!	
	Dried nuts of Canarium ovatum	Description/Definition.
	Engl.	Description/Definition: The traditional food consists of the unreceted daied note of Canadium contains Engl. (femily, Powerscape) commonly, known or Bili note are produced.
	•	The traditional food consists of the unroasted dried nuts of <i>Canarium ovatum</i> Engl. (family: Burseraceae) commonly known as Pili nuts. Pili nuts are produced only by plants of <i>Canarium ovatum</i> Engl. varieties Laysa, Magnaye, M. Orolfo, Lanuza and Magayon and can be placed on the market with or without the shell. The edible part of the nut is the kernel.
		Typical composition range:
		Fat: 57-73 %
		Protein: 11-15 %
		Water: 1-5 %
		Carbohydrates: 8-16,5 %
		Ash: 2,8-3,4 %
		Microbiological criteria:
		Moulds and yeasts: ≤ 100 CFU/g
		Total colony count at 30 °C: ≤ 10 000 CFU/g
		Coliforms: ≤ 100 CFU/g
		Escherichia coli: ≤ 10 CFU/g
		Staphylococcus aureus: Absence in 25 g
		Salmonella spp.: Absence in 25 g
		Listeria monocytogenes: Absence in 25 g
		Sulphite reducing anaerobes: ≤ 10 CFU/g
		CFU: Colony Forming Units

Authorised Novel Food	Specifications
2	
Canarium indicum L. dried nuts	Description/Definition:
(Kenari) (Traditional food from a third country)	The traditional food is processed dried kenari nuts. The term 'Kenari Nuts' refers to the kernels of ripe Kenari Fruit, scientifically known as <i>Canarium indi</i> L. (or <i>Canarium amboinense</i> Hochr.; family: Burseraceae).
	Composition:
	Ash: $\leq 5 \ (g/100 \ g)$
	Moisture: $\leq 6 (g/100 g)$
	Protein: 12,8 – 14,4 g/100 g
	Carbohydrates: 11,0 – 16,4 g/100 g
	Fat: 59,3 - 66,3 g/100 g
	Dietary fibre: 4,4 – 9,8 g/100 g
	Microbiological criteria:
	Aerobic Plate Count: $\leq 5.0 \times 10^3$ CFU/g
	Coliforms: < 3 MPN/g
	E. coli: < 3 MPN/g
	Yeasts and moulds: < 10 CFU/g
	Salmonella: Absent in 25 g
	Staphylococcus aureus (absent/25 g)
	Listeria monocytogenes (absent/25 g)
	Aflatoxins
	Aflatoxins B1: ≤ 2 mcg/kg
	Aflatoxins (Sum of B1, B2, G1, G2): ≤ 4 mcg/kg
	Dioxins and dioxin like PCBs
	Sum of dioxins: $\leq 0.75 \text{ pg/g fat}$
	Sum of dioxins and dioxin-like PCBs: ≤ 1,5 pg/g fat
	Heavy metals
	Cadmium (Cd): ≤ 0.02 mg/kg
	Lead (Pb): ≤ 0.07 mg/kg
	CFU: Colony Forming Units

	Authorised Novel Food	Specifications
▼ M114		
•	Cellobiose	Description/Definition:
		Cellobiose is a disaccharide with two glucose monomers linked by a β -(1–4) glucosidic bond, that is produced from sucrose and glucose in a two-step enzymatic reaction, followed by a series of purification steps.
		Characteristics/composition:
		Cellobiose DM (%): ≥ 99
		Moisture (%): < 1
		Other identified sugars (%): ≤ 1
		Optical rotation $[\alpha]_D$ (c 10, water): +33–36
		Ash $(g/100 g)$: < 0,1
		Protein content $(g/100 \text{ g})$: < 0,01
		Heavy metals:
		Arsenic: < 0,1 mg/kg
		Microbiological criteria:
		Total aerobic count (cfu/g): ≤ 1 000
		Yeast and moulds (cfu/g): ≤ 100
		Salmonella (in 25 g): n.d.
		Coliforms (cfu/g): ≤ 10
		E. coli (in 10 g): n.d.
		cfu: colony forming units
		n.d.: not detected

▼ <u>M82</u>		
	4	
	Cetylated fatty acids	Description/Definition:
		The novel food concerns primarily a mixture of cetylated myristic acid and cetylated oleic acid synthesised from cetyl alcohol, myristic acid and oleic acid and to a lesser degree, other cetylated fatty acids and other compounds from olive oil.
		Characteristics/composition:
		Ester content: 70-80 %, of which: Cetyl oleate: 22-30 %, Cetyl myristate: 41-56 %
		Triglycerides: 22-25 %
		Acid value (mg KOH/g): ≤ 5
		Saponification value (mg KOH/g): 130-150
		Microbiological criteria:
		Total aerobic microbial count: ≤ 1 000 CFU/g
		Yeasts and moulds: ≤ 100 CFU/g
		KOH: potassium hydroxide
		CFU: colony forming units
V M0		
▼ <u>M9</u>		
	Chewing gum base (monome-thoxypolyethylene glycol)	Description/Definition: The novel food ingredient is a synthetic polymer (Patent number WO2006016179). It consists of branched polymers of monomethoxypolyethyleners.
		glycol (MPEG) grafted onto polyisoprene-graft-maleic anhydride (PIP-g-MA), and unreacted MPEG (less than 35 % by weight).
		White to off-white colour.
		CAS No.: 1246080-53-4

Authorised Novel Food	Specifications
	Characteristics:
	Moisture: < 5,0 %
	Aluminium: < 3,0 mg/kg
	Lithium: < 0,5 mg/kg
	Nickel: < 0,5 mg/kg
	Residual anhydride: < 15 μmol/g
	Polydispersity index: < 1,4
	Isoprene: < 0,05 mg/kg
	Ethylene oxide: < 0,2 mg/kg
	Free maleic anhydride: < 0,1 %
	Total oligomeres (less than 1 000 Dalton): ≤ 50 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/kg
	Formaldehyde: < 10 mg/kg
Chawing gum base (Mathyl vinyl	Description/Definition:
Chewing gum base (Methyl vinyl ether-maleic anhydride copolymer)	
cinci-maicic annyuriue coporymer)	Free-flowing, white to white-off powder
	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: $\leq 500 \text{ ppm}$
	Methanol: ≤ 500 ppm
	Dilauroyl peroxide: ≤ 15 ppm
	Total heavy metals: ≤ 10 ppm

Authorised Novel Food	Specifications
	Microbiological criteria:
	Total aerobic plate count: ≤ 500 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 <i>Salvia hispanica</i>	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: ≤ 2,0 %
	Peroxide value (PV): ≤ 10 meq/kg
	Insoluble impurities: ≤ 0,05 %
	Alpha linolenic acid: ≥ 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.
Chitin-glucan from Aspergillus	Description/Definition:
niger	Chitin-glucan is obtained from the mycelium of <i>Aspergillus niger</i> ; 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: ≤ 10 %
	Chitin-glucan: ≥ 90 %
	Ratio of chitin to glucan: 30:70 to 60:40
	Ash: $\leq 3.0 \%$
	Lipids: ≤ 1,0 %
	Proteins: $\leq 6,0 \%$
Chitin-glucan complex from Fomes	Description/Definition:
fomentarius	Chitin-glucan complex is obtained from the cell walls of the fruit bodies of the fungus <i>Fomes fomentarius</i> . 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: ≤ 15 %
	Ash: $\leq 3.0 \%$
	Chitin-glucan: ≥ 90 %
	Ratio of chitin to glucan: 70:20
	Total carbohydrates, excluding glucans: ≤ 0,1 %

Specifications

Authorised Novel Food	Specifications
	Proteins: ≤ 2,0 %
	Lipids: $\leq 1,0\%$
	Melanins: $\leq 8,3\%$
	Additives: None
	pH: 6,7-7,5
	Heavy metals:
	Lead (ppm): ≤ 1,00
	Cadmium (ppm): $\leq 1,00$
	Mercury (ppm): ≤ 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	The chitosan extract (containing mainly poly(D-glucosamine)) is obtained from stems of Agaricus bisporus or from the mycelium of Aspergillus niger
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: (C ₆ H ₁₁ NO ₄) _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

Authorised Novel Food	Specifications
	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): ≤ 3,0 Proteins (% w/w dry weight): ≤ 2,0 Particle size: > 100 nm Tapped density (g/cm³): 0,7-1,0 Fat binding capacity 800 × (w/w wet weight): pass Heavy metals: Mercury (ppm): ≤ 0,1 Lead (ppm): ≤ 1,0 Arsenic (ppm): ≤ 1,0 Cadmium (ppm): ≤ 0,5 Microbiological criteria: Aerobic count (CFU/g): ≤ 10³ Yeast and mould count (CFU/g): ≤ 10³ Escherichia coli (CFU/g): ≤ 10
	Yeast and mould count (CFU/g): $\leq 10^3$
	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 <i>Escherichia coli</i> 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 (ΔDi-68) (%): ≤ 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

▼<u>M56</u>

Authorised Novel Food	Specifications
hromium 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(C ₆ H ₄ NO ₂) ₃
	Chemical characteristics:
	Chromium Picolinate: ≥ 95 %
	Chromium (III): 12-13 %
	Chromium (VI): not detected
	Water: ≤ 4,0 %
Chromium-containing yeast (<i>Yarrowia lipolytica</i>) biomass	Description/Definition:
	The novel food is the dried and heat-killed chromium-containing biomass of the yeast Yarrowia lipolytica.
	The novel food is produced by fermentation in the presence of chromium chloride followed by a number of purification steps and a heat-killing step of the provential to ensure the absence of viable Varrania linguistica cells in the povel food

yeast to ensure the absence of viable Yarrowia lipolytica cells in the novel food.

Characteristics/Composition:

Total chromium: 18-23 µg/g

Chromium (VI): < 10 µg/kg (i.e. limit of detection)

Protein: 40-50 g/100 g Dietary fibre: 24-32 g/100 g

Sugars: < 2 g/100 g Fat: 6-12 g/100 g Total ash: ≤ 15 % Water: ≤ 5 % Dry matter: ≥ 95 % Heavy metals: Lead: $\leq 3.0 \text{ mg/kg}$ Cadmium: ≤ 1,0 mg/kg

Mercury: ≤ 0.1 mg/kg

▼<u>M56</u>

Authorised Novel Food	Specifications
	Microbiological criteria:
	Total aerobic microbial count: $\leq 5 \times 10^3$ CFU/g
	Total yeast and mould count: $\leq 10^2$ CFU/g
	Viable <i>Yarrowia lipolytica</i> cells (14): < 10 CFU/g (i.e. limit of detection)
	Coliforms: ≤ 10 CFU/g
	Salmonella spp.: Absence in 25 g
	CFU: colony forming units
-	er er evitetig fermining unite
<u>85</u>	
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.
	The novel food consists of the dried and cut aerial parts (young shoots with woody parts) of Cistus incanus L. Pandalis
<u>)</u>	
Citicoline	Description/Definition:
Citiconne	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: C ₁₄ H ₂₆ N ₄ O ₁₁ P ₂
	Molecular weight: 488,32 g/mol
	CAS No.: 987-78-0
	pH (sample solution of 1 %): 2,5-3,5
	Purity:
	Assay value: ≥ 98 % of dry matter
	Loss on drying (100 °C for 4 hours): ≤ 5.0 %
	Ammonium: $\leq 0.05 \%$
	Arsenic: Not more than 2 ppm
	Free phosphoric acids: ≤ 0,1 %
	5'-Cytidylic acid: ≤ 1,0 %
	Microbiological criteria:
	Total plate count: $\leq 10^3$ CFU/g
	Yeast and moulds: $\leq 10^2 \text{ CFU/g}$
	Escherichia coli: Absence in 1 g

02017R2470	
R2470 —	
Z	
05 2024	
— 046 001	
1 182	

Authorised Novel Food	Specifications
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: ≤ 10³ 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: ≤ 10² CFU/g

▼M79

Coffea arabica L. and/or Coffea canephora Pierre ex A.Froehner dried cherry pulp and its infusion (Traditional food from a third country)

Description/Definition:

The traditional food consists of the dried unroasted coffee cherry pulp of Coffea arabica L. and/or Coffea canephora Pierre ex A.Froehner (genus: Coffea family: Rubiaceae) and its infusion. The infusion can be used as such or concentrated or dried.

Ripe coffee cherries are collected, and then the coffee beans are mechanically removed, prior or after a drying process, leaving the dried coffee cherry pulp, which can be milled to a powder.

The separated coffee cherry pulp is also known as 'cascara', from the Spanish 'cáscara', meaning 'husk'.

Typically, the infusion is prepared by mixing up to 6 g of cascara pulp or husk in 100 ml of hot water (> 75 °C) for a few minutes and then pouring through a strainer, or using corresponding amounts in dried or instant infusions.

Composition of the dried coffee cherry pulp:

Water: < 18 %

Water activity (a_w) : ≤ 0.65 Ash: < 10.4 % DM

Protein: < 15 % DM Fat: < 5 % DM

Carbohydrates: < 85 % DM

Microbiological criteria:

Aerobic Plate Count: $< 10^4$ CFU/g Total yeasts and moulds: < 100 CFU/g

Enterobacteriaceae: < 50 CFU/g Salmonella: Absence in 25 g Bacillus cereus: < 100 CFU/g

▼<u>M79</u>

Authorised Novel Food	Specifications
	Mycotoxins:
	Ochratoxin A: < 5,0 µg/kg
	Aflatoxin B1: < 2,0 μg/kg
	Aflatoxin B1, B2, G1, G2 (as sum): < 4,0 μg/kg
	Heavy metals:
	Cadmium (Cd): < 0,05 mg/kg
	Lead (Pb): < 1,0 mg/kg
	Copper: ≤ 50 mg/kg
	Mercury: ≤ 0.02 mg/kg
	Arsenic: ≤ 0,2 mg/kg
	Impurities:
	Benzo(a)pyrene: < 10,0 µg/kg
	Sum of benzo(a)pyrene, benz(a)anthracene, benzo(b)fluoranthene and chrysene: < 50,0 µg/kg
	Pesticides:
	Pesticide levels in the traditional food shall comply with levels set by Regulation (EC) No 396/2005 for '0639000' for 'Herbal infusions from any other parts of the plant'.
	CFU: Colony Forming Units
	DM: Dry Matter
▼ <u>M30</u>	
D-ribose	Description
	D-ribose is an aldopentose monosaccharide which is produced by fermentation using a transketolase-deficient strain of Bacillus subtilis.
	Chemical formula: C ₅ H ₁₀ O ₅
	CAS No: 50-69-1
	Molecular mass: 150,13 Da

Authorised Novel Food	Specifications
	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 (8) 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: ≤ 0,1 mg/kg
	Cadmium: ≤ 0,1 mg/kg
	Mercury: ≤ 0.1 mg/kg
	Microbiological criteria
	Total plate count: ≤ 100 CFU (9)/g
	Yeast: $\leq 100 \text{ CFU/g}$
	Moulds: ≤ 100 CFU/g
	Coliforms: ≤ 10 CFU/g
	Salmonella sp: Negative/25 g

	Authorised Novel Food	Specifications
▼ <u>M54</u>		
	Dried Euglena gracilis	Description/Definition:
		The novel food is dried whole cell Euglena, which is the dried biomass of the microalga Euglena gracilis.
		The novel food is produced by fermentation followed by filtration and a heat-killing step of the microalga to ensure the absence of viable <i>Euglena gracilis</i> cells in the novel food.
		Characteristics/Composition:
		Total carbohydrates: ≤ 75 %
		β-glucan: > 50 %
		Protein: ≥ 15 %
		Fat: ≤ 15 %
		Ash: $\leq 10\%$
		Moisture: ≤ 6 %
		Heavy metals:
		Lead: $\leq 0.5 \text{ mg/kg}$
		Cadmium: $\leq 0.5 \text{ mg/kg}$
		Mercury: $\leq 0.05 \text{ mg/kg}$
		Arsenic: ≤ 0,02 mg/kg
		Microbiological criteria:
		Aerobic plate count: ≤ 10 000 CFU/g
		Coliforms: ≤ 100 MPN/g
		Yeast and mould: ≤ 500 CFU/g
		Escherichia coli: Absence in 10 g
		Staphylococcus aureus: Absence in 10 g
		Salmonella: Absence in 25 g
		Listeria monocytogenes: Absence in 25 g
		CFU: colony forming units.
		MPN: most probable number

Authorised Novel Food	Specifications
Extract of defatted cocoa powder	Cocoa (Theobroma cacao L.) Extract
P	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/cm ³
	pH: 5,0-6,5
	Residual solvent: Max 500 ppm
Low fat cocoa extract	Low fat Cocoa (Theobroma cacao 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 %
Coriander seed oil from Coriandrum	Description/Definition:
sativum	Coriander seed oil is an oil containing glycerides of fatty acids that is produced from the seeds of the coriander plant Coriandrum sativum L.
	Yellowish to brown colour, bland taste
	CAS No.: 8008-52-4
	Composition of fatty acids:
	Palmitic acid (C16:0): 2-5 %

▼<u>M70</u>

▼<u>M15</u>

<u>)</u>	
Authorised Novel Food	Specifications
	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: ≤ 4 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
5	
Cranberry 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 <i>Vaccinium macrocarpon</i> .
	Characteristics/Composition
	Moisture (% w/w): ≤ 4
	Proanthocyanidins — PACs (% w/w dry weight)
	— OSC-DMAC method (3) (5): 55.0-60.0 or
	— BL-DMAC method (4) (5): 15.0-18.0
	Total phenolics (GAE (6), % w/w dry weight) (5)

— Folin-Ciocalteau method: > 46.2

Solubility (water): 100 %, with no visible insoluble particles

▼<u>M15</u>

▼<u>M9</u>

Authorised Novel Food	Specifications
	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 (⁷)/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 g
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 re-precipitation, steam-stripping of the

02017R2
7R247
0 — EN
_
- 01.05.2024
024 -
- 046.00
001 —
189

Authorised Novel Food Specifications complexant, and crystallisation of α-cyclodextrin from the solution; or chromatography with ion-exchange or gel filtration followed by crystallisation of αcyclodextrin from the purified mother liquor; or membrane separation methods such as ultra-filtration and reverse osmosis: Description: Virtually odourless, white or almost white crystalline solid. Synonyms: α-cyclodextrin, α-dextrin, cyclohexaamylose, cyclomaltohexaose, α-cycloamylase Chemical name: Cyclohexaamylose CAS 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: $\left[\alpha\right]_{D}^{25}$: 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 a-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: ≤ 20 mg/kg (1-decanol) Reducing substances: $\leq 0.5 \%$ (as glucose) Sulphated ash: $\leq 0.1 \%$ Lead: $\leq 0.5 \text{ mg/kg}$

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

Method of assay:

Determine by liquid chromatography using the following conditions:

Authorised Novel Food	Specifications
	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-NH ₂ (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: $10 \mu l$ Procedure: 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: 17465-86-0
	Chemical formula: (C ₆ H ₁₀ O ₅) ₈
	Assay: ≥ 98 % (dry basis)

	Authorised Novel Food	Specifications
		Identification: Melting range: Decomposes above 285 °C Solubility: Freely soluble in water; very slightly soluble in ethanol Specific rotation: $[\alpha]_D^{2.5}$: between + 174° and + 180° (1 % solution) Purity: Water: ≤ 11 % Residual complexant (8-cyclohexadecen-1-one (CHDC)): ≤ 4 mg/kg Residual solvent (n-decane): ≤ 6 mg/kg Reducing substances: ≤ 0.5 % (as glucose) Sulphated ash: ≤ 0.1 %
▼ <u>M22</u>		
	Decorticated grains of <i>Digitaria</i> 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: ≤ 2,1 mg/g
▼ <u>M9</u>	Dextran preparation produced by Leuconostoc mesenteroides	1. Powdered form: Carbohydrates: 60 % with: (Dextran: 50 %, Mannitol: 0,5 %, Fructose: 0,3 %, Leucrose: 9,2 %) Protein: 6,5 %

Authorised Novel Food	Specifications
	Lipid: 0,5 %
	Lactic acid: 10 %
	Ethanol: traces
	Ash: 13 %
	Moisture: 10 %
	2. 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 %
	Moisture: 80 %
Diacylglycerol oil of plant origin	Description/Definition:
	Manufactured from glycerol and fatty acids derived from edible vegetable oils, in particular from soybean oil (Glycine max) or rapeseed oil (Brassica campestris, Brassica napus) using a specific enzyme.
	Acylglycerol Distribution:
	Diacylglycerols (DAG): ≥ 80 %
	1,3-Diacylglycerols (1,3-DAG): ≥ 50 %
	Triacylglycerols (TAG): ≤ 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): ≤ 15 %
	Saturated fatty acids: ≤ 10 %

Authorised N	Novel Food	Specifications
		Others:
		Acid value: ≤ 0,5 mg KOH/g
		Moisture and volatile: $\leq 0.1 \%$
		Peroxide value (PV): ≤ 1,0 meq/kg
		Unsaponifiables: ≤ 2,0 %
		Trans fatty acids≤ 1,0 %
		MAG = monoacylglycerols, DAG = diacylglycerols, TAG = triacylglycerols
Dihydrocapsiate (D	OHC)	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: C ₁₈ H ₂₈ O ₄
		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 %
<u>M13</u>		
Dried aerial parts	of <i>Hoodia</i>	Description/Definition:
parviflora		It is the whole dried aerial parts of <i>Hoodia parviflora</i> N.E.Br., (family <i>Apocynaceae</i>)
		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
		$oldsymbol{\cdot}$

Authorised Novel Food	Specifications
	pH: < 5,0
	Protein: < 4,5 g/100 g
	Fat: $< 3 \text{ g}/100 \text{ g}$
	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/kg
	Mercury: < 0,1 mg/kg
	Cadmium: < 0,1 mg/kg
	Lead: < 0,5 mg/kg
	Microbiological criteria:
	Aerobic plate count: $< 10^5$ CFU/g
	Escherichia coli: < 10 CFU/g
	Staphylococcus aureus: < 50 CFU/g
	Total coliforms: < 10 CFU/g
	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

	Authorised Novel Food	Specifications
	Dried extract of <i>Lippia citriodora</i> 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:		Description/Definition:
		Extract of the roots of <i>Echinacea angustifolia</i> obtained from plant tissue culture which is substantially equivalent to a root extract from <i>Echinacea angustifolia</i> obtained in ethanol-water titrated to 4 % echinacoside.
▼ <u>M32</u>		
	Echinacea purpurea extract from cell cultures	Description/Definition:
		Dried extract of <i>Echinacea purpurea</i> from cell cultures EchiPure-PC TM
▼ <u>M9</u> Echium plantagineum oil Description/Definition:		
		Description/Definition:
		Echium oil is the pale yellow product obtained by refining oil extracted from the seeds of <i>Echium plantagineum</i> L. Stearidonic acid: ≥ 10 % w/w of total fatty acids
		Trans fatty acids: ≤ 2,0 % (w/w of total fatty acids)
		Acid value: ≤ 0,6 mg KOH/g
		Peroxide value (PV): ≤ 5.0 meq O_2/kg
		Unsaponifiable content: ≤ 2,0 %
		Protein content (total nitrogen): ≤ 20 μg/ml
		Pyrrolizidine alkaloids: Not detectable with a detection limit 4,0 μg/kg

▼<u>M52</u>

Authorised Novel Food	Specifications
_	
Ecklonia 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

02017R2470
0 - EN -
01.05.2024
-046.001
-197

Authorised Novel Food	Specifications
-----------------------	----------------

▼M18

Egg 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

Total nitrogen-containing compounds (% w/w): ≥ 88

Collagen (% w/w): ≥ 15 Elastin (% w/w): ≥ 20

Total glycosaminoglycans (% w/w): ≥ 5

Calcium: ≤ 1 %

Physical parameters

pH: 6,5 - 7,6

Ash (\% w/w): ≤ 8

Moisture (% w/w): ≤ 9

Water activity: ≤ 0.3

Solubility (in water): soluble

Bulk density: ≥ 0,6 g/cc

Heavy metals

Arsenic ≤ 0.5 mg/kg

Microbiological criteria

Aerobic plate count: ≤ 2 500 CFU/g

Escherichia coli: ≤ 5 MPN/g

Salmonella: Negative (in 25 g)

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

 $\label{eq:Staphylococcus} \textit{Staphylococcus aureus} : \leq 10 \text{ CFU/g}$ $\mbox{Mesophilic spore count} : \leq 25 \text{ CFU/g}$

Thermophilic spore count: ≤ 10 CFU/10 g

Methods

Combustion according to AOAC 990.03 and AOAC 992.15

SircolTM Soluble Collagen Assay

FastinTM Elastin Assay

USP26 (chondroitin sulphate K0032 method)

▼M18

Authorised Novel Food		Specificat	ons	
	Yeast: ≤ 10 CFU/g			
	Mould: ≤ 200 CFU/g			
	_	MPN = Most Probable Number; USP: United State	Pharmaconeja.	
	, ,	,		
Epigallocatechin gallate as a	Description/Definition:			
purified extract from green tea leaves (Camellia sinensis)	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			
	Appearance: off-white to pale	e pink powder		
	Chemical name: polyphenol	(-) epigallocatechin-3-gallate		
	Synonyms: epigallocatechin g	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 m	naterial)		
	max. 0,1 % caffeine			
	Solubility: EGCG is fairly so	pluble in water, ethanol, methanol and acetone		
L-ergothioneine	Definition			
	Chemical name (IUPAC): (2S)-3-(2-thioxo-2,3-dihydro-1 <i>H</i> -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	$[\alpha]_D \ge (+) \ 122^{\circ} \ (c = 1, H_2O)^{a)}$	Polarimetry	

Authorised Novel Food		Specifications	
	Chemical purity	≥ 99,5 %	HPLC [Eur. Ph. 2,2.29]
		≥ 99,0 %	1H-NMR
	Identification	Compliant with the structure	1H-NMR
		C: $47,14 \pm 0,4 \%$	Elemental analysis
		H: $6,59 \pm 0,4 \%$	
		N: $18,32 \pm 0,4 \%$	
	Total residual solvents	[Eur. Ph. 01/2008:50400]	Gas chromatography
	(methanol, ethyl acetate, isopropanol, ethanol)	< 1 000 ppm	[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)	$\leq 1 \times 10^3 \text{ CFU/g}$	[Eur. Ph. 01/2011:50104]
	Total yeast and mould count (TYMC)	$\leq 1 \times 10^2 \text{ CFU/g}$	
	Escherichia coli	Absence in 1 g	

▼ <u>M9</u>		
	Authorised Novel Food	Specifications
		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. $[\alpha]_D = (+) 126,6^{\circ} (c = 1, H_2O)$
		b) Analyses conducted on each batch
		c) Maximum levels in accordance with Regulation (EC) No 1881/2006
▼ M108		
	Roasted and popped kernels from the seeds of <i>Euryale ferox</i> Salisb. (makhana) (Traditional food from a third country)	Description/Definition
		The traditional food consists of the roasted and popped kernels of the seeds of the fresh plants of <i>Euryale ferox</i> Salisb. (family: Nymphaeaceae, commonly referred to also as prickly water lily) to be consumed as a snack. The traditional food is produced via a series of steps involving the collection, washing, and drying of the seeds, a first roasting in oil, tempering at ambient temperatures, a subsequent second roasting in oil to pop the kernels, followed by hitting of the hot seeds to release the popped kernels. The traditional food is also known as makhana or fox nuts.
		Typical nutritional components:

Fat: 13,0 g/100 g

Carbohydrates: 75,0 g/100 g

Fibre: 2,5 g/100 g Protein: 7 g/100 g

Moisture (% w/w): < 5,0

Ash: < 0.5 g/100 g

Microbiological criteria:

Total plate count: $< 10^3$ CFU/g

Total yeast and mould count: < 100 CFU/g

Total Enterobacteriaceae: < 10 CFU/g Salmonella spp.: Absence in 25 g

Listeria monocytogenes: Absence in 25 g

Heavy metals:

Selenium: ≤ 0,8 mg/kg Copper: ≤ 30,0 mg/kg Lead: $\leq 0.1 \text{ mg/kg}$ Arsenic: ≤ 0,1 mg/kg

▼<u>M108</u>

	Authorised Novel Food	Specifications
		Cadmium: ≤ 0,1 mg/kg
		Tin: $\leq 3.5 \text{ mg/kg}$
		Mercury: ≤ 0.025 mg/kg
		Mycotoxins:
		Aflatoxin B1: ≤ 2,0 μg/kg
		Sum of aflatoxins B1, B2, G1, and G2: ≤ 4,0 µg/kg
		Ochratoxin A: ≤ 1,0 µg/kg
		Citrinin: $\leq 20,0 \mu g/kg$
		Cyanotoxins:
		Microcystins: ≤ 0.0015 mg/kg
		Pesticides:
		Pesticides: ≤ 0,01 mg/kg
		Process contaminants:
		Acrylamide: ≤ 40,0 µg/kg
		Sum of PAHs: $\leq 10.0 \mu \text{g/kg}$
		Sum of dioxin-like PCBs: $\leq 0.35 \text{ pg/g}$
		3 -MCPD: $\leq 20,0 \mu g/kg$
		Glycidyl fatty acid esters (expressed as glycidol): ≤ 500,0 µg/kg
		Sum of 3-MCPD and 3-MCPD fatty acid esters: ≤ 750,0 µg/kg
		CFU: Colony Forming Units; PAHs: Polycyclic Aromatic Hydrocarbons; PCBs: Polychlorinated Biphenyls; 3-MCPD: 3-MonoChloroPropane Diol.
▼ <u>M52</u>		
	Extract 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)

	Authorised Novel Food	Specifications
		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 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
▼ <u>M9</u>	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

Authorised Novel Food	Specifications
	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: ≤ 0,1 % Nitrilo-triacetic acid: ≤ 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.: $10101\text{-}60\text{-}7$ Chemical formula: FeNH ₄ PO ₄ Chemical characteristics: pH of 5 % suspension in water: 6,8-7,8 Iron (total): ≥ 28 % Iron (II): $22\text{-}30$ % (w/w) Iron (III): $\leq 7,0$ % (w/w) Ammonia: $5\text{-}9$ % (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 (<i>Sardinops sagax</i>) 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): ≥ 85 g/100 g Val-Tyr (dipeptide): 0,1-0,16 g/100 g Ash: ≤ 10 g/100 g Moisture: ≤ 8 g/100 g (¹) Kjeldahl method

	Authorised Novel Food	Specifications
	Flavonoids from Glycyrrhiza glabra	Description/Definition: Flavonoids derived from the roots or rootstock of <i>Glycyrrhiza glabra</i> 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: ≥ 99 % Protein: < 0,1 % Carbohydrates: not detectable
<u>M42</u>	Fruit pulp, pulp juice, concentrated pulp juice from <i>Theobroma cacao</i> L. (Traditional food from a third country)	Description/Definition The traditional food is the fruit pulp from the cocoa (<i>Theobroma cacao</i> 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 (y/100 g): 0,0 to 2,0 Total fat (g/100 g): 0,0 to 0,2 Total fat (g/100 g): 0,0 to 0,2 Total sugars (g/100 g): > 11,0 Brix level (° Brix): ≥ 14 pH: 3,3 to 4,0 Microbiological criteria Total Plate Count (aerobic): < 10 000 cfu (*9)/g Enterobacteriaceae: < 10 cfu/g Salmonella: Absence in 25 g

Specifications

Frozen, dried and powder forms of Locusta migratoria (migratory locust)

Description/Definition:

The novel food consists of the frozen, dried and powder forms of migratory locust. The term 'migratory locust' refers to the adult of *Locusta migratoria*, an insect species that belongs to the Acrididae family (subfamily Locustinae).

The novel food is intended to be marketed in three different forms, namely: (i) thermally processed and frozen *L. migratoria* (LM frozen); (ii) thermally processed and freeze-dried *L. migratoria* (LM dried), and (iii) thermally processed freeze-dried and ground whole *L. migratoria* (whole LM powder). The LM dried may be marketed as such or in powder.

For LM frozen and LM dried, legs and wings must be removed to reduce the risk of intestinal constipation that could be possibly caused by ingestion of the large spines on the insect tibia. The whole LM powder is obtained via mechanical grinding of the insect with legs and wings, and sieving to reduce particle size below 1 mm.

A minimum 24 hours fasting period is required before killing the insects by freezing, to allow the adults to discard their bowel content.

Parameters	LM frozen	LM dried	Whole LM powder		
Characteristics/Composition	Characteristics/Composition				
Ash (% w/w)	0,6-1,0	2,0-3,1	1,8-1,9		
Moisture (% w/w)	67-73	≤ 5	≤ 5		
Crude protein (N × 6,25) (% w/w)	11-21	43-53	50 - 60		
Fat (% w/w)	7-13	31-41	31-41		
Saturated fatty acids (% fat)	35-43	35-43	35-43		
Digestible carbohydrates (% w/w)	0,1-2,0	0,1-2,0	1,0-3,5		
(18) Dietary fibre (% w/w)	1,5-3,5	5,5-9,0	5,5-9,0		
Chitin (% w/w)	1,7-2,4	6,4-10,4	10,5-13,9		
Peroxide value (Meq O ₂ /kg fat)	≤ 5	≤ 5	≤ 5		

Authorised Novel Food	Specifications				
	Contaminants				
	Lead (mg/kg)	≤ 0,07	≤ 0,07	≤ 0,07	
	Cadmium (mg/kg)	≤ 0,05	≤ 0,05	≤ 0,05	
	Aflatoxins (Sum of B1, B2, G1, G2) (μg/kg)	≤ 4	≤ 4	≤ 4	
	Aflatoxin B1 (μg/kg)	≤ 2	≤ 2	≤ 2	
	Deoxynivalenol (μg/kg)	≤ 200	≤ 200	≤ 200	
	Ochratoxin A (µg/kg)	≤ 1	≤ 1	≤ 1	
	Sum of dioxins and dioxins-like PCBs UB ((19) WHO ₂₀₀₅ PCDD/ F-PCB-TEQ) (pg/g fat)	≤ 1,2	≤ 1,2	≤ 1,2	
	Microbiological criteria				
	Total aerobic colony count ((7) CFU/g)	≤ 10 ⁵	≤ 10 ⁵	≤ 10 ⁵	
	Enterobacteriaceae (presumptive) (CFU/g)	≤ 100	≤ 100	≤ 100	
	Escherichia coli (CFU/g)	≤ 50	≤ 50	≤ 50	
	Listeria monocytogenes	Not detected in 25g	Not detected in 25g	Not detected in 25g	
	Salmonella spp.	Not detected in 25g	Not detected in 25g	Not detected in 25g	
	Bacillus cereus (presumptive) (CFU/g)	≤ 100	≤ 100	≤ 100	
	Coagulase positive <i>Staphylococci</i> (CFU/g	≤ 100	≤ 100	≤ 100	
	Sulfite-reducing Anaerobes (CFU/g)	≤ 30	≤ 30	≤ 30	
	Yeasts and moulds (CFU/g)	≤ 100	≤ 100	≤ 100	

Authorised Novel Food	Specifications		
Fucoidan extract from the seaweed	Description/Definition:		
Fucus vesiculosus	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 %		

Authorised Novel Food	Specifications		
Fucoidan extract from the seaweed	Description/Definition:		
Undaria pinnatifida	Fucoidan from seaweed <i>Undaria pinnatifida</i> 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 %		

Authorised Novel Food	Specifications
	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 %
'-Fucosyllactose	Definition:
ynthetic)	Chemical name: α -L-Fucopyranosyl- $(1\rightarrow 2)$ - β -D-galactopyranosyl- $(1\rightarrow 4)$ - D-glucopyranose
,	Chemical formula: C ₁₈ H ₃₂ O ₁₅
	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: $\leq 1.0 \text{ w/w } \%$
	L-Fucose: ≤ 1,0 w/w %
	Difucosyl- D-lactose isomers: ≤ 1,0 w/w %
	2'-Fucosyl- D-lactulose: ≤ 0,6 w/w %
	pH (20 °C, 5 % solution): 3,2-7,0
	Water (%): $\leq 9.0 \%$
	Ash, sulphated: ≤ 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: ≤ 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: ≤ 10 CFU/g
	Residual endotoxins: ≤ 10 EU/mg

Aut	thorised Novel Food		Specifications		
▼ <u>M110</u>					
Specification	ns				Data protection
		Definition: Chemical name: α-L-Fucopyranosyl-(Chemical formula: C ₁₈ H ₃₂ O ₁₅ CAS No: 41263-94-9 Molecular weight: 488,44 g/mol	$1\rightarrow 2$)-β-D-galactopyranosyl-($1\rightarrow 4$)-D-glucopyranose		2'-Fucosyllactose produced with a genetically modified strain of Corynebacterium glutamicum ATCC 13032 authorised on 16 May 2023. This inclusion is based on proprietary scientifications.
2'-Fucosylla source)	Source: Genetically modified strain of Escherichia coli K-12 Source: Genetically modified strain coli BL-21	Source: Genetically modified strain of <i>Escherichia</i> coli BL-21	Source: Genetically modified strain of Corynebacterium glutamicum ATCC 13032	evidence and scientific data protected in accordance with Article 26 of Regulation (EU) 2015/2283. Applicant: 'Advanced Protein	
		Description: 2'-Fucosyllactose is a white to offwhite powder that is produced by a microbiological 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, Difusoryl D-lactose, L-Fucose, D-Lactose, D-Lactose, L-Fucose, D-Lactose, D	Description: 2'-Fucosyllactose is a white to off white powder and the liquid concentrate (45 % ± 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: ≤ 5,0 % Fucose: ≤ 3,0 % 3-Fucosyllactose: ≤ 5,0 % Fucosylgalactose: ≤ 3,0 %	Description: 2'-Fucosyllactose is a white to off white/ivory powder that is produced by a microbiological process. Purity: 2'-Fucosyllactose (w/w dry matter): ≥ 94,0 % D-Lactose (w/w dry matter): ≤ 3,0 % L-Fucose (w/w dry matter): ≤ 3,0 % 3-Fucosyllactose (w/w dry	Technologies Corporation', 7th Floor GyeongGi-BioCenter, 147, Gwanggyo-ro, Yeongtonggu, Suwon-si Gyeonggi-do, 16229 South Korea. During the period of data protection, 2'-Fucosyllactose produced with a genetically modified strain of Corynebacterium glutamicum ATCC 13032 is authorised for placing on the market within the Union only by 'Advanced Protein Technologies Corporation' unless a subsequent applicant obtains
		Difucosyl-D-lactose, 2'-Fucosyl-D-lactulose): ≥ 90 % pH (20 C, 5 % solution): 3,0-7,5 Water: ≤ 9,0 % Sulphated ash: ≤ 2,0 % Acetic acid: ≤ 1,0 % Residual proteins: ≤ 0,01 %	Difucosyllactose: ≤ 5,0 % Glucose: ≤ 3,0 % Galactose: ≤ 3,0 % Water: ≤ 9,0 % (powder) Ash, sulphated: ≤ 0,5 % (powder and liquid) Residual proteins: ≤ 0,01 % (powder and liquid)	matter): ≤ 3,0 % Difucosyllactose (w/w dry matter): ≤ 2,0 % D-Glucose (w/w dry matter): ≤ 3,0 % D-Galactose (w/w dry matter): ≤ 3,0 % Water: ≤ 9,0 % Ash: ≤ 0,5 % Residual proteins: ≤ 0,005 %	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 'Advanced Protein Technologies Corporation'.

▼<u>M110</u>

Authorised Novel Food	Specifications			
	Microbiological criteria: Aerobic mesophilic bacteria total count: ≤ 3 000 CFU/g Yeasts: ≤ 100 CFU/g Moulds: ≤ 100 CFU/g Endotoxins: ≤ 10 EU/mg CFU: Colony Forming Units; EU: Endotoxin Units	Heavy Metals: Lead: ≤ 0,02 mg/kg (powder and liquid) Arsenic: ≤ 0,2 mg/kg (powder and liquid) Cadmium: ≤ 0,1 mg/kg (powder and liquid) Mercury: ≤ 0,5 mg/kg (powder and liquid) Microbiological criteria: Total plate count: ≤ 104 CFU/g (powder), ≤ 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: ≤ 0,025 μg/kg (powder and liquid) CFU: Colony Forming Units; EU: Endotoxin Units	Contaminants: Arsenic: ≤ 0,03 mg/kg Aflatoxin M1: ≤ 0,025 µg/kg Ethanol: ≤ 1 000 mg/kg Microbiological criteria: Total plate count: ≤ 500 CFU/g Yeasts and Moulds: ≤ 100 CFU/g Enterobacteriaceae: absence in 10 g Salmonella: absence in 25 g Cronobacter spp.: absence in 10 g Endotoxins: ≤ 100 EU/g CFU: Colony Forming Units; EU: Endotoxin Units	End date of the data protection 16 May 2028.
2'-Fucosyllactose/Difucosyllactose mixture ('2'-FL/DFL') (microbial source)	Source: Genetically modified Escheric Characteristics/Composition: Appearance: White to off white power Sum of 2'-Fucosyllactose, Difucosyllactose, Difucosy			

▼<u>M58</u>

	Authorised Novel Food	Specifications
		2'-Fucosyllactose (% of dry matter): ≥ 75,0 % (w/w)
		Difucosyllactose (% of dry matter): ≥ 5,0 % (w/w)
		D-Lactose: $\leq 10.0 \% \text{ (w/w)}$
		L-Fucose: $\leq 1.0 \% \text{ (w/w)}$
		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: ≤ 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
▼ <u>M75</u>		
	3-Fucosyllactose ('3-FL')	Description:
	(microbial source)	3-Fucosyllactose (3-FL) is a purified, white to off-white powder that is produced by microbial fermentation and contains limited levels of D-Lactose, L-Fucose, D-Galactose, and D-Glucose.
		Source: Genetically modified strain of <i>Escherichia coli</i> K-12.
		Definition:
		Chemical formula: C ₁₈ H ₃₂ O ₁₅
		Chemical name: β -D-galactopyranosyl- $(1\rightarrow 4)[-\alpha$ -L-fucopyranosyl- $(1\rightarrow 3)]$ -D-glucopyranose
		Molecular mass: 488,44 Da
		CAS No 41312-47-4
		Characteristics/Composition:
		3-Fucosyllactose (% of dry matter): ≥ 90.0 % (w/w)
		D-Lactose (% of dry matter): ≤ 5.0 % (w/w)
		L-Fucose (% of dry matter): ≤ 3.0 % (w/w)

Authorised Novel Food	Specifications
	Sum of D-Galactose/D-Glucose (% of dry matter): ≤ 3,0 % (w/w)
	Sum of other carbohydrates ^a (% of dry matter): ≤ 3,0 % (w/w)
	Moisture: $\leq 5.0 \%$ (w/w)
	pH (20 °C, 5 % solution): 3,0-7,5
	Residual protein: $\leq 0.01 \% \text{ (w/w)}$
	Ash (%): ≤ 0.5
	Heavy metals/Contaminants:
	Arsenic: ≤ 0,2 mg/kg
	Cadmium: ≤ 0,05 mg/kg
	Lead: $\leq 0.05 \text{ mg/kg}$
	Mercury: $\leq 0.1 \text{ mg/kg}$
	Aflatoxin M1: $\leq 0.025 \mu g/kg$ Aflatoxin B1: $\leq 0.1 \mu g/kg$
	Residual endotoxins: ≤ 0.3 EU/mg
	Microbiological criteria:
	Total plate count: ≤ 1 000 CFU/g
	Enterobacteriaceae: Absence in 10 g
	Salmonella sp.: Absence in 25 g
	Cronobacter (Enterobacter) sakazakii: Absence in 10 g
	Listeria monocytogenes: Absence in 25 g
	Bacillus cereus: ≤ 10 CFU/g
	Yeast: \leq 100 CFU/g
	Mould: $\leq 100 \text{ CFU/g}$
	CFU: Colony Forming Units; EU: Endotoxin Units; ^a Sum of other carbohydrates: 3-Fucosyllactose isomer, difucosyllactose isomer, and oligomers.
M102	
	Description
3-Fucosyllactose ('3-FL') (produced by a derivative strain o	Description:
E. coli BL21(DE3))	^f 3-Fucosyllactose (3-FL) is a purified, white to off-white powder that is produced by microbial fermentation and contains limited levels of D-Lactose, L-Fucose, D-Galactose, and D-Glucose.
	Definition:
	Chemical name: β -D-Galactopyranosyl- $(1\rightarrow 4)$ - $[\alpha$ -L-fucopyranosyl- $(1\rightarrow 3)$]- D-glucopyranose
	Chemical formula: C ₁₈ H ₃₂ O ₁₅
	Molecular mass: 488,44 Da
	Molecular mass. 700,777 Da

▼<u>M102</u>

Authorised Novel Food	Specifications
	CAS No: 41312-47-4 Source: A genetically modified strain of <i>Escherichia coli</i> BL21(DE3) Characteristics/Composition: 3-Fucosyllactose (% of dry matter): ≥ 90.0 % (w/w) D-Lactose (% of dry matter): ≤ 5.0 % (w/w) D-glucose (% of dry matter): ≤ 3.0 % (w/w) D-galactose (% of dry matter): ≤ 3.0 % (w/w) L-Fucose (% of dry matter): ≤ 3.0 % (w/w) Sum of other carbohydrates (% of dry matter) (24): ≤ 5.0 % (w/w) Moisture: ≤ 9.0 % (w/w) Residual protein: ≤ 0.01 % (w/w)
	Heavy metals and contaminants: Arsenic: ≤ 0,2 mg/kg Aflatoxin M1: ≤ 0,025 μg/kg
	Microbiological criteria: Standard plate count: ≤ 1 000 CFU (25)/g Enterobacteriaceae: ≤ 10 CFU/g Salmonella spp.: Absence in 25 g Yeast and mould: ≤ 100 CFU/g Cronobacter (Enterobacter) sakazakii.: Absence in 10 g Residual endotoxins: ≤ 10 EU (26)/mg
▼ <u>M125</u>	
3-Fucosyllactose ('3-FL') (produced by derivative strain of E. coli K-12 DH1)	Description: 3-Fucosyllactose (3-FL) is a purified and concentrated white to off-white powder produced by microbial fermentation and contains limited levels of D-Lactose, 3-Fucosyllactulose, and L-Fucose. Definition: Chemical name: β-D-Galactopyranosyl-(1→4)- [α-L-fucopyranosyl-(1→3)]- D-glucopyranose Chemical formula: C ₁₈ H ₃₂ O ₁₅ Molecular mass: 488,44 Da CAS No: 41312-47-4 Source: Genetically modified strain of Escherichia coli K-12 DH1

▼<u>M125</u>

Authorised Novel Food	Specifications
	Characteristics/Composition:
	3-Fucosyllactose (% w/w of dry matter): ≥ 90,0
	D-Lactose (% w/w): ≤ 5.0
	3-Fucosyllactulose ($\%$ w/w): ≤ 1.5
	L-Fucose (% w/w): ≤ 1.0
	Sum of 3-Fucosyllactose, 3-Fucosyllactulose, D-Lactose and L-Fucose, (% w/w dry matter): ≥ 92,0
	Sum of other carbohydrates (% w/w): ≤ 5.0
	Moisture (% w/w): ≤ 6.0
	pH (20 °C, 5 % solution): 3,2 -7,0
	Ash (% w/w): ≤ 0.5
	Acetic acid (% w/w): $\leq 1,0$
	Residual protein (% w/w): ≤ 0.01
	Heavy metals and contaminants:
	Arsenic: ≤ 0.2 mg/kg
	Aflatoxin M1: $\leq 0.025 \mu \text{g/kg}$
	Microbiological criteria:
	Total plate count: ≤ 1 000 CFU/g
	Enterobacteriaceae: Absence in 10 g
	Salmonella spp.: Absence in 25 g
	Yeast and mould: ≤ 100 CFU/g
	Cronobacter spp.: Absence in 10 g
	Listeria monocytogenes: Absence in 25 g
	Presumptive Bacillus cereus: $\leq 50 \text{ CFU/g}$
	Endotoxins: ≤ 10 EU/mg
	CFU: Colony Forming Units; EU: Endotoxin Units
	Cro. Colony rollining Units, Ed. Endoloxin Units
<u>M132</u>	
Galacto-oligosaccharide	Description/Definition:
	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 and Papiliotrema terrestris.
	GOS: min. 46 % Dry Matter (DM)
	Lactose: max. 40 % DM
	Glucose: max. 22 % DM
	Ash: max. 4,0 % DM
	Protein: max. 4,5 % DM
	Nitrite: max. 2 mg/kg

Authorised Novel Food	Specifications
Glucosamine HCl from Aspergillus niger and genetically modified strain of E. coli K-12	White crystalline odourless powder Molecular formula: $C_6H_{13}NO_5 \cdot HCl$ Relative molecular mass: 215,63 g/mol D-Glucosamine HCl 98,0-102,0 % of reference standard (HPLC) Specific rotation + 70.0° - + 73.0°
Glucosamine sulphate KCl from Aspergillus niger and genetically modified strain of E. coli K-12	White crystalline odourless powder Molecular formula: (C ₆ H ₁₄ NO ₅) ₂ SO ₄ · 2KCl Relative molecular mass: 605,52 g/mol D-Glucosamine Sulphate 2KCl 98,0-102,0 % of reference standard (HPLC) Specific Rotation +50,0° to +52,0°
Glucosamine sulphate NaCl from Aspergillus niger and genetically modified strain of E. coli K-12	White crystalline odourless powder Molecular formula: (C ₆ H ₁₄ NO ₅) ₂ SO ₄ · 2NaCl Relative molecular mass: 573,31 g/mol D-Glucosamine HCl: 98-102 % of reference standard (HPLC) 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 (¹) & 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 (²).

Authorised Novel Food Specifications	
	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	Description/Definition:
fermented with Bacteroides xylani- solvens	Heat-treated fermented milk products are produced with <i>Bacteroides xylanisolvens</i> (DSM 23964) as starter culture.

Authorised Novel Food	Specifications		
	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 wi <i>Bacteroides xylanisolvens</i> (DSM 23964). The resulting fermented milk product is homogenised and then heat-treated to inactivate <i>Bacteroides xylanisolvens</i> (DS 23964). The final product does not contain viable cells of <i>Bacteroides xylanisolvens</i> (DSM 23964)(1).		
	() Modified DIN EN ISO 21328-2.		
ydroxytyrosol	Description/Definition:		
	Hydroxytyrosol is a pale yellow viscous liquid obtained by chemical synthesis		
	Molecular formula: C ₈ H ₁₀ O ₃		
	Molecular weight: 154,6 g/mol		
	CAS No: 10597-60-1		
	Moisture $\leq 0.4 \%$		
	Odour: CharacteristicTaste: Slightly bitter		
	Solubility (water): Miscible with water		
	pH: 3,5-4,5		
	Refractive Index: 1,571-1,575		
	Purity:		
	Hydroxytyrosol: ≥ 99 %		
	Acetic acid: ≤ 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: $\leq 0.01 \text{ mg/kg}$		
	Residual Solvents		
	Ethyl acetate: ≤ 25,0 mg/kg		
	Isopropanol: $\leq 2,50 \text{ mg/kg}$		
	Methanol: $\leq 2,00 \text{ mg/kg}$		
	Tetrahydrofuran: ≤ 0,01 mg/kg		

Authorised Novel Food

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 leaves of	Description/Definition:		
	Ilex guayusa	Dark brown liquid. Aqueous extracts of dried leaves of Ilex guayusa.		
		Composition:		
		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		
▼ <u>M49</u>				
	Infusion from coffee leaves of	Description/Definition:		
	Coffea arabica L. and/or Coffea	The traditional food consists of an infusion of leaves from Coffea arabica L. and/or Coffea canephora Pierre ex A.Froehner (family: Rubiaceae).		
	canephora Pierre ex A. Froehner	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		
	(Traditional food from a third country)	of hot water. Leaves are removed and the infusion is then subjected to pasteurization (at least 71 °C for 15 seconds).		

Specifications

Authorised Novel Food	Specifications	
	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	
94		
Iron hydroxide adipate tartrate	Description/Definition:	
	Iron hydroxide adipate tartrate (IHAT) is an odourless, engineered nanomaterial in powder form that is insoluble in water and is manufactured by a chemical synthesis involving a series of steps involving acid-base reaction, precipitation, filtration, and drying.	
	The food supplements containing the novel food are manufactured in capsular form. Excess adipate, tartrate and sodium chloride are used at levels resulting from the production process to help stabilise IHAT and ensure the authorised particle size distribution. If other forms of food supplements (e.g. tablets pastilles, sachets of powders, gummies, syrups, etc.) are used in combination with adipate, tartrate and sodium chloride or in combination with other substances, or if other substances are used in the capsular form food supplements containing the novel food, it must be ensured that the authorised IHAT particle size distribution is maintained.	

Authorised Novel Food

		•
		Į.

Specifications		
Common name	Iron oxo-hydroxide adipate tartrate	
Other names	Iron hydroxide adipate tartrate, Iron oxyhydroxide adipate tartrate	
Trade name	IHAT	
CAS number	2460638-28-0	
Molecular formula (calculated)	$FeO_m(OH)_n(H_2O)_x(C_4H_6O_6)_y(C_6H_{10}O_4)_z$ where: m and n are undefined as per accepted practice for ferric iron oxohydroxides (*) $x=0.28\text{-}0.88$ $y=0.78\text{-}1.50$ $z=0.04\text{-}0.19$ Tartaric (C ₄ H ₆ O ₆) and adipic (C ₆ H ₁₀ O ₄) acid are represented in their protonated form.	
Molecular weight	Average molecular weight: 35 803,4 Da (lower-upper bound: 27 670,5-45 319,4 Da)	

Specifications

Characteristics/Composition:

Physical/chemical

Iron (% dry matter): 24,0-36,0 Adipate: (% dry matter): 1,5-4,5 Tartrate: (% dry matter): 28,0-40,0 Water content (%): 10,0-21,0 Sodium (% dry matter): 9,0-11,0 Chloride (% dry matter): 2,6-4,2

▼<u>M94</u>

Authorised Novel Food	Specifications
	Phase distribution
	Soluble (%): 2,0-4,0
	Nano (%): 92,0-98,0
	Micro (%): 0,0-3,0
	Primary particle size
	Median diameter (20): 1,5-2,3 nm
	Mean diameter (20): 1,8-2,8 nm
	Dv(10) (²¹): 1,5-2,5 nm
	Dv(50) (21): 2,5-3,5 nm
	Dv(90) (21): 5,0-6,0 nm
	Heavy metals
	Arsenic: < 0,80 mg/kg
	Nickel: < 50,0 mg/kg
	Residual solvents
	Ethanol: < 500 mg/kg
	Microbiological criteria
	Total aerobic microbial count: < 10 CFU/g
	Total yeast and mould count: < 10 CFU/g
▼ <u>M116</u>	
Iron milk caseinate	Description:
	Iron milk caseinate is an iron-casein-phosphate complex in the form of a creamy or beige powder produced by the dissolution of ferric iron salts (ferric sulfate or ferric chloride) in a casein solution obtained from bovine milk in the presence of potassium orthophosphate following a series of steps involving pasteurisation, concentration, and drying.
	Characteristics/Composition:
	Protein (%): 50,0 - 65,0
	Ash (%): 20,0 – 40,0
	Moisture (%): < 8,0
	Fat (%): < 1,0
	Iron (%): 2,0 – 4,0

▼<u>M116</u>

	Authorised Novel Food	Specifications		
		Potassium (%): 5,0 – 15,0Phosphorus (%): 2,0 – 6,0		
		Sodium (%): < 4,0		
		Heavy metals:		
		Lead: < 0,5 mg/kg		
		Arsenic: ≤ 1,0 mg/kg		
		Cadmium: < 0,5 mg/kg		
		ercury: < 0.1 mg/kg		
		lycotoxins:		
		flatoxin M1: ≤ 0.02 mg/kg		
		Microbiological criteria:		
		Aerobic plate count: ≤ 1 000 CFU/g		
		Coliforms: ≤ 10 CFU/g		
		Salmonella spp.: Absence in 25 g		
		Yeast and mould: ≤ 10 CFU/g		
		Escherichia coli: ≤ 10 CFU/g		
		Staphylococcus aureus: Absence in 1 g		
		CFU: Colony Forming Units		
▼ <u>M9</u>				
	Isomalto-oligosaccharide	Powder:		
		Solubility (water) (%): > 99		
		Glucose (% dry basis): ≤ 5.0		
		Isomaltose + DP3 to DP9 (% dry basis): ≥ 90		
		Moisture (%): ≤ 4.0		
		Sulphated $ash(g/100 g)$: ≤ 0.3		
		Heavy metals:		
		Lead (mg/kg): ≤ 0.5		
		Arsenic (mg/kg): ≤ 0.5		

•	by	an	-

Syrup:
Dried solids (g/100 g): > 75
Glucose (% dry basis): ≤ 5.0
Isomaltose + DP3 to DP9 (% dry basis): ≥ 90
pH: 4 - 6
Sulphated $ash(g/100 g)$: ≤ 0.3
Heavy metals:
Lead (mg/kg): ≤ 0.5
 Arsenic (mg/kg): ≤ 0.5

Isomaltulose

Authorised Novel Food

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 taste

Specifications

Chemical name: $6\text{-O-}\alpha\text{-D-glucopyranosyl-D-fructofuranose}$, monohydrate

CAS No.: 13718-94-0

Chemical formula: C₁₂H₂₂O₁₁ · H₂O

Structural formula

Formula weight: 360,3 (monohydrate)

0201
7R247
0
三
01.05
05.2024
- 04
6.001
<u></u>

Authorised Novel Food	Specifications
	Purity: Assay: ≥ 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'
	(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, 322 pp., English, ISBN 92-5-102991-1.

▼M90

Jatropha curcas L. (edible variety) kernels

Description:

The kernels are obtained from the seeds of the ripe fruits of the edible variety of the *Jatropha curcas* L. plants that produce kernels with non-detectable levels of phorbol esters, following a series of steps involving the cleaning and de-husking of the fruits to obtain the seeds, the drying of the seeds, the cleaning of the seeds to remove debris and other residues, mechanical deshelling of the seeds to obtain the kernels, and the hydrothermal treatment (> 120 °C for 40 minutes) of the kernels to reduce anti-nutrients and the microbiological load.

As the edible variety of the *Jatropha curcas* L. plants, producing kernels that contain non-detectable levels of phorbol esters, are phenotypically undistinguishable from the non-edible variety, only the appropriate edible variety of *Jatropha curcas* L. plants should be used in the production of the novel food. The entire production process must ensure that the mixing of edible and non-edible kernels does not occur.

The absence of mixing of edible with non-edible kernels shall be confirmed by analytical controls for phorbol esters carried on each batch of the seeds after the seed-drying step and before the deshelling step according to the sampling procedure of Table A. Five laboratory samples extracted from each aggregate sample are de-shelled, ground, and analysed for phorbol esters using a validated UHPLC-UV-MS^(b) method. Only the batches in which phorbol esters are undetectable in all five samples are further processed to the seed deshelling and kernel hydrothermal treatment steps.

Table A

1400 12		
Batch weight (tons) Weight or number of sublots		Number of incremental samples
≥ 500 100 tons		100
> 100 and < 500	5 sub-lots	100
> 10 and ≤ 100	5 sub-lots	100
> 5,0 and \le 10	-	80
$> 1 \text{ and } \le 5,0$	-	60
$> 0.1 \text{ and} \le 1.0$	-	30
≤ 0,1	-	10

Each sub-lot shall be sampled separately. Aggregate samples are composed by a minimum of 10 incremental samples. The minimum amount of an aggregate sample shall be 3,5 kg. This amount may increase proportionally according to the number of incremental samples taken.

	Authorised Novel Food	Specifications
-		Characteristics/Composition: Moisture: ≤ 3,0 % Total fat: 54,0 – 61,0 % Total protein: 21,0 – 32,0 % Total fibre: 6,0 – 10,0 % Ash: 3,0 – 5,0 % Contaminants: Phorbol esters (μg TPA eq ^(a) /g kernel) ^(b) : ≤ 0,75 (LOD) ^(c) Lead: ≤ 0,20 mg/kg Cadmium: ≤ 0,20 mg/kg Sum of aflatoxins B1, B2, G1, G2: ≤ 4,0 μg/kg Microbiological criteria: Total aerobic microbial count: ≤ 1 000 CFU/g Total yeast/moulds count: ≤ 100 CFU/g Enterobacteriaceae: ≤ 10 CFU/g Salmonella sp.: Absent in 25 g Listeria monocytogenes: ≤ 100 CFU/g (a) TPAeq: 12-O-tetradecanoylphorbol-13-acetate equivalent; (b)Validated Ultra-High-Performance Liquid Chromatography coupled to Ultraviolet Spectrophotometry and Mass Spectrometry (UHPLC-UV-MS) method for detection of phorbol ester peaks; (c) Limit of Detection (Only batches with concentrations of PEs below the LOD can be fully processed.); CFU: Colony Forming Units
▼ <u>M9</u>	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: $4\text{-O-}\beta\text{-D-Galactopyranosyl-D-glucitol}$ Chemical formula: $C_{12}H_{24}O_{11}$ Molecular weight: $344,31$ g/mol CAS No: 585-86-4 Purity: Solubility (in water): Very soluble in water Specific rotation $[\alpha]_D^{20} = +13^\circ$ to $+16^\circ$ Assay: ≥ 95 % d.b (d.b — expressed on the dry weight basis) Water: $\leq 10,5$ % Other polyols: $\leq 2,5$ % d.b Reducing sugars: $\leq 0,2$ % d.b Chlorides: ≤ 100 mg/kg d.b Sulphates: ≤ 200 mg/kg d.b

Authorised Novel Food	Specifications
	Sulphated ash: ≤ 0,1 % d.b Nickel: ≤ 2,0 mg/kg d.b Arsenic: ≤ 3,0 mg/kg d.b Lead: ≤ 1,0 mg/kg d.b
Lacto-N-neotetraose (synthetic)	Definition: Chemical name: β-D-Galactopyranosyl-(1→4)-2-acetamido-2-deoxy-β-D-glucopyranosyl-(1→3)-β-D-galactopyranosyl-(1→4)- D-glucopyranose Chemical formula: C ₂₆ H ₄₅ NO ₂₁ 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): ≥ 96 % D-Lactose: ≤ 1,0 % Lacto-N-triose II: ≤ 0,3 % Lacto-N-neotetraose fructose isomer: ≤ 0,6 % pH (20 °C, 5 % solution): 5,0-7,0 Water: ≤ 9,0 % Ash, sulphated: ≤ 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: ≤ 10 CFU/g Moulds: ≤ 10 CFU/g Moulds: ≤ 10 CFU/g
123	Residual endotoxins: ≤ 10 EU/mg
Lacto-N-neotetraose (microbial source)	Definition Chemical name: β-D-Galactopyranosyl-(1→4)-2-acetamido-2-deoxy-β-D-glucopyranosyl-(1→3)-β-D-galactopyranosyl-(1→4)-D-glucopyranose Chemical formula: C ₂₆ H ₄₅ NO ₂₁ CAS No: 13007-32-4 Molecular weight: 707,63 g/mol

	Authorised Novel Food	Specifications
		Description/Source
		Lacto-N-neotetraose is a white to off-white powder that is produced by a microbiological process using genetically modified strain of <i>Escherichia coli</i> K-12, and/or of <i>Escherichia coli</i> BL21(DE3). An additional optional genetically modified degradation strain of <i>Escherichia coli</i> BL21(DE3) may be used in the production process to degrade intermediate carbohydrate by-products and remaining starting carbohydrate substrates.
		Purity
		Assay (water free): ≥ 80 %
		D-Lactose: $\leq 10.0 \%$
		Lacto-N-triose II: ≤ 3,0 %
		para-Lacto-N-neohexaose: ≤ 5,0 %
		Lacto- <i>N</i> -neotetraose fructose isomer: ≤ 1,0 %
		Sum of saccharides (Lacto- <i>N</i> -neotetraose, D-Lactose, Lacto- <i>N</i> -triose II, <i>para</i> -Lacto- <i>N</i> -neohexaose, Lacto- <i>N</i> -neotetraose fructose isomer): ≥ 92 % (% w/w dry matter)
		pH (20 °C, 5 % solution): 4,0-7,0
		Water: $\leq 9.0 \%$
		Ash, sulphated: $\leq 1.0 \%$
		Residual solvents (methanol): ≤ 100 mg/kg
		Residual proteins: ≤ 0,01 %
		Microbiological criteria
		Aerobic mesophilic bacteria total count: ≤ 500 CFU/g
		Yeasts and moulds: ≤ 50 CFU/g
		Residual endotoxins: ≤ 10 EU/mg
		CFU: Colony Forming Units; EU: Endotoxin Units
▼ M45		
▼ <u>M46</u>		
	T A NAME OF THE PARTY OF THE PA	
	Lacto-N-tetraose ('LNT') (microbial source)	Definition:
	(iniciobiai source)	Chemical formula: $C_{26}H_{45}NO_{21}$ 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 or agglomerates that is produced by a microbial process.
		Lacto-ty-tetraose is a purified, write to oil-write amorphous powder or aggiomerates that is produced by a microbial process.

Authorised Novel Food	Specifications
	Source:
	Genetically modified strain of Escherichia coli strain K-12 DH1
	Characteristics/Composition:
	Appearance: White to off white powder or agglomerates
	Sum of lacto-N-tetraose, D-Lactose and lacto-N-triose II (% of dry matter): ≥ 90.0 % (w/w)
	Lacto-N-tetraose (% of dry matter): ≥ 70.0 % (w/w)
	D-Lactose: $\leq 12.0 \%$ (w/w)
	Lacto-N-triose II: ≤ 10.0 % (w/w)
	$Para$ -lacto- N -hexaose-2: $\leq 3.5 \%$ (w/w)
	Lacto- <i>N</i> -tetraose fructose isomer: ≤ 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 spp.: Negative/25 g
	Yeast: ≤ 100 CFU/g
	Mould: $\leq 100 \text{ CFU/g}$
	Residual endotoxins: ≤ 10 EU/mg
	CFU: Colony Forming Units
▼ <u>M101</u>	
Lacto-N-tetraose ('LNT')	Description:
(produced by derivative strains of	Lacto- <i>N</i> -tetraose is a purified and concentrated white to off-white powder that is produced by a microbial fermentation process.
E. coli BL21(DE3))	Definition:
` "	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
	Chemical formula: $C_{26}H_{45}NO_{21}$
	CAS No: 14116-68-8
	Molecular mass: 707.63 Da
	Source: Two genetically modified strains (a production strain and an optional degradation strain) of <i>Escherichia coli</i> BL21(DE3)
	Source 1.10 generally meaning that he production stain and an optional degladation stain) of Estimated Stains (a production stain and an optional degladation stain) of Estimated Stains (a production stain and an optional degladation stain) of Estimated Stains (a production stain and an optional degladation stain) of Estimated Stains (a production stain and an optional degladation stain) of Estimated Stains (a production stain and an optional degladation stain) of Estimated Stains (a production stain and an optional degladation stain) of Estimated Stains (a production stain and an optional degladation stain and an option and an opt

▼<u>M101</u>

Authorised Novel Food	Specifications
	Characteristics/Composition:
	Lacto-N-tetraose (% of dry matter): ≥ 75.0 % (w/w)
	D-Lactose (% of dry matter): ≤ 5.0 % (w/w)
	Lacto-N-triose II (% of dry matter): ≤ 5.0 % (w/w)
	Para-lacto-N-hexaose (% of dry matter): ≤ 5.0 % (w/w)
	D-galactose and D-glucose (% of dry matter): ≤ 5.0 % (w/w)
	Sum of other carbohydrates ^a : $\leq 15,0 \% (w/w)$
	Moisture: $\leq 9.0 \% \text{ (w/w)}$
	Ash: $\leq 1,0 \% \text{ (w/w)}$
	Residual protein: $\leq 0.01 \% (w/w)$
	Heavy metals and contaminants:
	Arsenic: $\leq 0.2 \text{ mg/kg}$
	Aflatoxin M1: $\leq 0.025 \mu \text{g/kg}$
	Microbiological criteria:
	Standard plate count: ≤ 1 000 CFU/g
	Enterobacteriaceae: ≤ 10 CFU/g
	Salmonella spp.: Absence in 25 g
	Yeast and mould: ≤ 100 CFU/g
	Cronobacter (Enterobacter) sakazaki: Absence in 10 g
	Residual endotoxins: ≤ 10 EU/mg ^a Sum of the most should to a 100 (0/ (m/m) of dry most so) and should to a (0/ (m/m) of dry most so). Ash (0/ (m/m) of dry most so). CEU Color
	a Sum of other carbohydrates = 100 (% (w/w) of dry matter) – quantified carbohydrates (% (w/w) of dry matter) – Ash (% (w/w) of dry matter). CFU: Colon Forming Units; EU: Endotoxin Units
<u> </u>	
_	
Lonicera caerulea L. berries (haskap)	Description/Definition:
(Traditional food from a third	The traditional food are fresh and frozen berries from Lonicera caerulea var. edulis.
country)	Lonicera caerulea L. is a deciduous shrub belonging to the Caprifoliaceae family.
•	Typical nutritional components of haskap berries (given in fresh berries):
	Carbohydrates: 12,8 %
	Fibre: 2,1 %
	Lipids: 0,6 %
	Proteins: 0,7 %
	110tcms. 0,7 /0

▼<u>M21</u>

▼<u>M9</u>

Authorised Novel Food	Specifications
	Ash: 0,4 % Water: 85,5 %
Lucerne leaf extract from	Description/Definition:
Medicago sativa	The Lucerne (<i>Medicago sativa</i> 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

Authorised Novel Food	Specifications		
Lycopene from Blakeslea trispora	Description/Definition:		
	The purified lycopene from <i>Blakeslea trispora</i> 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: C ₄₀ H ₅₆		
	Formula weight: 536,85 Da		
Lycopene from tomatoes	Description/Definition:		
	The purified lycopene from tomatoes (<i>Lycopersicon esculantum</i> 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: C ₄₀ H ₅₆		
	Formula weight: 536,85 Da		
Lycopene oleoresin from tomatoes	Description/Definition:		
V	Lycopene oleoresin from tomatoes is obtained by solvent extraction of ripe tomatoes (<i>Lycopersicon esculentum Mill.</i>) 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 %		

	J		
-	_		

02017R2470 — EN — 01.05.2024 — 046.001 — 233

▾	M52

Hen	egg	white	lysozyme
hydr	olys	ate	

Authorised Novel Food

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.

Specifications

Specification

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

Tryptophan: 5-7 %

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

Degree of hydrolysis: 19-25 %

Moisture: < 5 %
Ash: < 10 %
Sodium: < 6 %

Heavy metals

Arsenic: < 1 ppm Lead: < 1 ppm

Cadmium: < 0,5 ppm Mercury: < 0,1 ppm

Microbiological criteriaTotal aerobic count: < 10³ CFU/g

Total combined yeasts/moulds count: < 10² 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

Authorised Novel Food	Specifications
Magnesium citrate malate	Description/Definition:
	Magnesium citrate malate is a white to yellowish-white, amorphous powder. Chemical formula: Mg ₅ (C ₆ H ₅ O ₇) ₂ (C ₄ H ₄ O ₅) 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): ≤ 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: ≤ 0,05 %
	Sulphate: ≤ 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 <i>Magnolia officinalis</i> 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: ≥ 85,2 %
	Honokiol: $\geq 0.5 \%$

Authorised Novel Food	Specifications
	Magnolol & Honokiol: ≥ 94 %
	Total Eudesmol: ≤ 2 %
	Moisture: 0,50 %
	Heavy metals:
	Arsenic (ppm): ≤ 0.5
	Lead (ppm): ≤ 0.5
	Methyl eugenol (ppm): ≤ 10
	Tubocurarine (ppm): $\leq 2,0$
	Total Alkaloid (ppm): ≤ 100
Maize-germ oil high in unsapo-	Description/Definition:
ifiable matter	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: ≥ 1,3 g/100 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: ≤ 6,0 mg KOH/g
	Peroxide value (PV): $\leq 10 \text{ mEq O}_2/\text{kg}$

Authorised Novel Food	Specifications		
	Heavy metals:		
	Iron (Fe): < 1 500 μg/kg		
	Copper (Cu): < 100 µg/kg		
	Impurities:		
	Polycyclic aromatic hydrocarbons (PAH) Benzo(a)pyrene: < 2 µg/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 unsaponifiable matter'		
Iethylcellulose	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:		
	- H		
	- CH ₃ or		
	- CH2CH3		
	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 Ol		
	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 acet acid.		
	Purity:		
	Loss on drying: ≤ 10 % (105 °C, 3 hours)		
	Sulphated Ash: ≤ 1.5 % determined at 800 ± 25 °C		
	pH: ≥ 5.0 and ≤ 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		

▼<u>M11</u>

Authorised Novel Food	Specifications
1-Methylnicotinamide chloride	Definition:
	Chemical name: 3-carbamoyl-1-methyl-pyridinium chloride
	Chemical formula: C ₇ H ₉ N ₂ OCl
	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: ≥ 98,5 %
	Trigonelline: ≤ 0,05 %
	Nicotinic Acid: ≤ 0,10 %
	Nicotinamide: ≤ 0,10 %
	Largest unknown impurity: ≤ 0,05 %
	Sum of unknown impurities: ≤ 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

Authorised Novel Food	Specifications	
(6S)-5-methyltetrahydrofolic acid, glucosamine salt	Description/Definition: Chemical name: N-[4-[[[(6S)-2-amino-1,4,5,6,7,8-hexahydro-5-methyl-4-oxo-6-pteridinyl]methyl]amino]benzoyl]-L-glutamic acid, glucosamine salt Chemical formula: C ₃₂ H ₅₁ N ₉ O ₁₆ 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: ≤ 8,0 % Heavy metals: Lead: ≤ 2,0 ppm Cadmium: ≤ 1,0 ppm Mercury: ≤ 0,1 ppm Mercury: ≤ 0,1 ppm Microbiological criteria: Total aerobic microbial count: ≤ 100 CFU/g Yeasts and moulds: ≤ 100 CFU/g Yeasts and moulds: ≤ 100 CFU/g Escherichia coli: Absence in 10g	
Monomethylsilanetriol (Organic Silicon)	Description/Definition: Chemical name: Silanetriol, 1-methyl- Chemical formula: CH ₆ O ₃ Si Molecular weight: 94,14 g/mol CAS No: 2445-53-6	

▼ M9

Authorised Novel Food	Specifications
	Purity:
	Organic Silicon (monomethylsilanetriol) preparation (aqueous solution):
	Acidity (pH): 6,4-6,8
	Silicon: 100-150 mg Si/l
	Heavy metals:
	Lead: $\leq 1.0 \mu \text{g/l}$
	Mercury: $\leq 1.0 \mu g/l$
	Cadmium: ≤ 1,0 μg/l
	Arsenic: $\leq 3.0 \mu \text{g/l}$
	Solvents:
	Methanol: ≤ 5,0 mg/kg (residual presence)

▼ M133

Monosodium salt of L-5-methyltetrahydrofolic acid

Description/Definition:

The novel food is produced by chemical synthesis and consists of L-5-methyltetrahydrofolic acid.

Molecular formula: C₂₀H₂₄N₇NaO₆

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

CAS number: 2246974-96-7 Molecular weight: 481,44 g/mol Characteristics/composition:

Appearance: White to yellow or beige powder

Assay & related compounds: Assay 5-MeTHFA-Na on dry basis: > 95 %; Sum of folate-related substances: ≤ 2,5

Sodium: 4 %–5 % w/w

Water: ≤ 1,0 %

Residual solvents: Ethanol: \leq 0,5 %; Isopropanol: \leq 0,5 % Diastereomeric purity: (6R)-Mefolinate: \leq 1,0 % area

Elemental impurities: Boron: ≤ 10 mg/kg

Platinum: ≤ 10 mg/kg (for foods intended for infants and young children and food supplements intended for pregnant women then ≤ 2 mg/kg)

Arsenic: $\leq 1,5$ mg/kg

▼ <u>M133</u>

	Authorised Novel Food	Specifications		
		Cadmium: ≤ 0,5 mg/kg		
		Lead: $\leq 1.0 \text{ mg/kg}$		
		Mercury: ≤ 1,5 mg/kg (for foods intended for infants and young children and food supplements intended for pregnant women then ≤ 1 mg/kg)		
	Microbiological criteria:			
		Total aerobic microbial count: ≤ 100 CFU/g		
		Total yeast and moulds count: ≤ 100 CFU/g		
		E. coli: Not detected in 10 g		
		Abbreviations: CFU: colony forming unit; IR: infra-red; MeTHFA: methyltetrahydrofolic acid.		
▼ <u>M87</u>				
-				
	Mung bean (Vigna radiata) protein			
		The novel food is mung bean protein powder extracted from seeds of the plant <i>Vigna radiata</i> by several processing steps followed by pasteurization and spray drying.		
		Characteristics/composition:		
		Moisture: ≤ 6 %		
		Protein $(w/w)^{(a)}$: $\geq 84\%$		
		Ash (w/w) : $\leq 6.0 \%$		
		Fat (w/w) : $\leq 5.5 \%$		
		Carbohydrate (w/w): ≤ 5.0 by calculation		
		Microbiological criteria:		
		Aerobic plate count: < 5 000 CFU/g (b)		
		Yeasts and moulds: < 100 CFU/g		
		Coliforms: < 100 CFU/g		
		Escherichia coli: < 10 CFU/g		
		Listeria monocytogenes: Not detected in 25 g		
		Salmonella spp.: Not detected in 25 g		
		(a) w/w: weight per weight.		
		(b) CFU: colony forming units.		

rm,	the	-

02017R2470 — EN

▼	M92
•	

Nicotino	mida	riboside	ablarida
Nicoting	miae	rinasiae	chiariae

Authorised Novel Food

Mycelial extract from Shiitake mushroom (Lentinula edodes)

Description/Definition:

Description/Definition:

tertiary structure.

Moisture: 98 % Dry matter: 2 %

brown, slightly turbid liquid.

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

Lentinan: 0.8 - 1.2 mg/ml

(1) Bradford method
 (2) Kjeldahl method

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

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.

Specifications

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

Lentinan is a β -(1-3) β -(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

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: C11H15N2O5Cl Molecular weight: 290,7 g/mol Characteristics/Composition: Colour: White to light brown

Form: Powder

Identification: Conforms by NMR (nuclear magnetic resonance)

Purity/Composition of the mycelial extract from Lentinula edodes:

Nicotinamide riboside chloride: ≥ 90 %

Water content: $\leq 2 \%$

Authorised Novel Food	Specifications
	Residual solvents:
	Acetone: ≤ 5 000 mg/kg
	Methanol: ≤ 1 000 mg/kg
	Acetonitrile: ≤ 50 mg/kg
	Methyl tert-butyl ether: ≤ 500 mg/kg
	Reaction by-products:
	Methyl acetate: ≤ 1 000 mg/kg
	Acetamide: ≤ 27 mg/kg
	Acetic acid: ≤ 5 000 mg/kg
	Heavy metals:
	Arsenic: ≤ 1 mg/kg
	Mercury*: $\leq 0.1 \text{ mg/kg}$
	Cadmium*: ≤ 1 mg/kg
	Lead*: ≤ 0,5 mg/kg
	Microbiological criteria:
	Total Plate Count: ≤ 1 000 CFU/g
	Yeast and Mould: ≤ 100 CFU/g
	Escherichia coli: Absence in 10 g
	CFU: colony forming units
	(*) only for foods for special medical purposes, total diet replacement for weight control and meal replacements
)	
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

Authorised Novel Food

Noni fruit juice powder (Morinda	Description/Definition:
citrifolia)	Seeds and skin of the sun-dried fruits of <i>Morinda citrifolia</i> 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).
New Contament	
Noni fruit puree and concentrate (Morinda citrifolia)	Description/Definition:
(Community of the Community of the Commu	The fruits of <i>Morinda citrifolia</i> 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.
	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: $\leq 0.4 \text{ g/}100 \text{ g}$
	Ash: $< 1.0 \text{ g}/100 \text{ g}$
	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 %

Specifications

Authorised Novel Food	Specifications	
	P 2.2.5 (100	
	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 <i>Morinda citrifolia</i> 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/kg	
	Rubiadin: non detectable, ≤ 10 µg/kg	
	Lucidin: non detectable, ≤ 10 μg/kg	
Noni fruit powder (<i>Morinda</i> citrifolia)	Description/Definition:	
··· •	Noni fruit powder is made from pulped noni (<i>Morinda citrifolia L.</i>) 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.	

Authorised Novel Food	Specifications
	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): $\leq 2,0 \mu g/ml$
	(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/	Description/Definition:
phytostanols	Oil enriched with phytosterols/phytostanols is composed of an oil fraction and a phytosterol fraction.
	Acylglycerol Distribution:
Free fatty acids (expressed as oleic acid): ≤ 2,0 %	
	M 11 1 000 00 1000
	Monoacylglycerols (MAG): ≤ 10 %
	Monoacylglycerols (MAG): ≤ 10 % Diacylglycerols (DAG): ≤ 25 %
	Diacylglycerols (DAG): ≤ 25 %
	Diacylglycerols (DAG): ≤ 25 % Triacylglycerols (TAG): Making up the balance
	Diacylglycerols (DAG): $\leq 25 \%$ Triacylglycerols (TAG): Making up the balance Phytosterol fraction: β-sitosterol: $\leq 80 \%$ β-sitostanol: $\leq 15 \%$
	Diacylglycerols (DAG): $\leq 25 \%$ Triacylglycerols (TAG): Making up the balance Phytosterol fraction: β -sitosterol: $\leq 80 \%$ β -sitostanol: $\leq 15 \%$ campesterol: $\leq 40 \%$
	Diacylglycerols (DAG): $\leq 25 \%$ Triacylglycerols (TAG): Making up the balance Phytosterol fraction: β -sitosterol: $\leq 80 \%$ β -sitostanol: $\leq 15 \%$ campesterol: $\leq 40 \%$ campestanol: $\leq 5,0 \%$
	Diacylglycerols (DAG): $\leq 25 \%$ Triacylglycerols (TAG): Making up the balance Phytosterol fraction: β -sitosterol: $\leq 80 \%$ β -sitostanol: $\leq 15 \%$ campesterol: $\leq 40 \%$ campestanol: $\leq 5,0 \%$ stigmasterol: $\leq 30 \%$
	Diacylglycerols (DAG): $\leq 25 \%$ Triacylglycerols (TAG): Making up the balance Phytosterol fraction: β -sitosterol: $\leq 80 \%$ β -sitostanol: $\leq 15 \%$ campesterol: $\leq 40 \%$ campestanol: $\leq 5,0 \%$

Authorised Novel Food	Specifications			
	Others: Moisture and volatile: ≤ 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 m than 99 %.			
Oil extracted from squids VM126 Partially defatted chia seed (Salvia hispanica) powders	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 % 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 % Powder with high protein content Powder with high fibre content			
	Particle size	≤ 130 µm	≤ 400 μm	
	Chemical composition:			
		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 %	

▼<u>M126</u>

	Authorised Novel Food	Specifications		
Microbiological criteria: Total plate count: ≤ 10 000 CFU/g Yeasts: ≤ 500 CFU/g Moulds: ≤ 500 CFU/g Staphylococcus aureus: ≤ 10 CFU/g Coliforms: < 100 MPN/g Enterobacteriaceae: ≤ 100 CFU/g Bacillus cereus: ≤ 50 CFU/g Escherichia coli: Not detected in 10 g Listeria monocytogenes: Not detected in 25 g Salmonella spp.: Absence in 25 g Contaminants: Arsenic: ≤ 0,1 ppm Cadmium: ≤ 0,1 ppm Lead: ≤ 0,1 ppm Mercury: ≤ 0,1 ppm Total aflatoxins: ≤ 4 ppb Ochratoxin A: ≤ 1 ppb		Total plate count: ≤ 10 000 CFU/g Yeasts: ≤ 500 CFU/g Moulds: ≤ 500 CFU/g Staphylococcus aureus: ≤ 10 CFU/g Coliforms: < 100 MPN/g Enterobacteriaceae: ≤ 100 CFU/g Bacillus cereus: ≤ 50 CFU/g Escherichia coli: Not detected in 10 g Listeria monocytogenes: Not detected in 25 g Salmonella spp.: Absence in 25 g Contaminants: Arsenic: ≤ 0,1 ppm Cadmium: ≤ 0,1 ppm Lead: ≤ 0,1 ppm Mercury: ≤ 0,1 ppm Total aflatoxins: ≤ 4 ppb		
▼ <u>M63</u>				
	Partially defatted rapeseed powder from <i>Brassica rapa</i> L. and <i>Brassica napus</i> L.	Definition: The powder is produced from the partially defatted seeds of non-genetically modified <i>Brassica rapa</i> L. and <i>Brassica napus</i> L. double low (00) cultivars through a series of processing steps to reduce glucosinolates and phytates. Source: Brassica rapa L. and Brassica napus L. seeds Characteristics/Composition: Protein (N × 6,25): 33,0-43,0 % Lipids: 14,0 - 22,0 % Total Carbohydrates(*): 33,0 - 40,0 % Total Fibre(**): 33,0 - 43,0 %		

Authorised Novel Foo	d Specifications
	Moisture: < 7,0 %
	Ash: 2,0–5,0 %
	Total Glucosinolates: < 0,3 mmol/kg (≤ 120 mg/kg)
	Phytate: < 1,5 %
	Peroxide value (in novel food weight): ≤ 3.0 mEq O_2/kg
	Heavy Metals:
	Lead: < 0,2 mg/kg
	Arsenic (inorganic): < 0,2 mg/kg
	Cadmium: < 0,2 mg/kg
	Mercury: < 0,1 mg/kg
	Aluminium: < 35,0 mg/kg
	Microbiological criteria:
	Total plate count (30 °C): < 5 000 CFU/g
	Enterobacteriaceae: < 10 CFU/g
	Salmonella sp.: Negative/25 g
	Yeast and mould: < 100 CFU/g
	Bacillus cereus: < 100 CFU/g
	(*) By difference: 100 % – [protein % + moisture % + fat % + ash %]
	(**) AOAC 2011.25 (Enzymatic gravimetry)
	CFU: Colony Forming Units, AOAC: Association of Official Agricultural Chemists
▼ <u>M55</u>	
Extract from Panax notog	inseng Description/Definition:
and Astragalus membrana	The novel food contains two extracts. One is an ethanol extract of the roots of <i>Astragalus membranaceus</i> (Fisch.) Bunge. The other is a hot water extract of the roots of <i>Panax notoginseng</i> (Burkill) F.H. Chen that is further concentrated using absorption on a resin and subsequent elution with 60 % ethanol. At the end of the manufacturing process both extracts are mixed (45–47,5 % of each extract) with maltodextrin (5–10 %).
	Characteristics/Composition:
	Total saponins: 1,5-5 %
	Ginsenoside Rb1: 0,1-0,5 %
	Astragaloside I: 0,01-0,1 %

▼<u>M55</u>

	Authorised Novel Food		Specifications			
Carbohydrates: ≥ 90 %						
Protein: ≤ 4,5 %						
Ash: ≤ 1 %						
		Moisture: ≤ 5 %				
		Fat: $\leq 1.5 \%$ Heavy metals:				
		Arsenic: ≤ 0,3 mg/kg				
		Microbiological criteria:				
		Total plate count: ≤ 5 000 CFU/g				
		Total yeast and mould count: ≤ 500	CFU/g			
	Enterobacteriaceae: < 10 CFU/g					
	Escherichia coli: Absence in 25 g					
Salmonella: Absence in 375 g						
	Staphylococcus aureus: Absence in 25 g					
		CFU: colony forming units				
▼ M9						
V <u>IVI</u>						
	Pasteurised fruit-based prep- arations produced using high-pressure treatment	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		
		pH	3,2 to 4,2			
		° Brix	7 to 42	Assured by added sugars		
		\mathbf{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		

Authorised Novel Food	Specifications
0	
Pea and rice protein fermented by	Description:
Lentinula edodes (Shiitake mushroom) mycelia	The novel food is produced from the fermentation of a mixture of 65 % pea and 35 % rice protein concentrates by the mycelia of the Shiitake mushro (<i>Lentinula edodes</i>) followed by heat treatment to terminate the fermentation and a series of drying steps to form a powder.
	Characteristics/Composition:
	Protein (% dry weight, N x 6,25): ≥ 75,0
	Moisture: ≤ 7.0
	Total fat (% dry weight): ≤ 10,0
	Ash (% dry weight): ≤ 10.0
	Carbohydrates (% by calculation): ≤ 15,0
	Mycotoxins:
	Aflatoxin B1 (μ g/kg): < 1,0
	Aflatoxin B2 (μ g/kg): < 1,0
	Aflatoxin G1 (μ g/kg): < 1,0
	Aflatoxin G2 (μ g/kg): < 1,0
	Aflatoxin total (B1+B2+G1+G2) (μ g/kg): < 3,0
	Heavy metals:
	Arsenic ($\mu g/g$): < 0,1
	Cadmium (μ g/g): < 0,1
	Lead ($\mu g/g$): < 0,3
	Mercury $(\mu g/g)$: < 0,1
	Microbiological criteria:
	Total aerobic microbial count: < 1 000 CFU/g
	Total yeast/moulds count: < 100 CFU/g
	Coliforms: $\leq 10 \text{ CFU/g}$
	Salmonella spp.: Absent in 25 g
	Escherichia coli: < 10 CFU/g
	Listeria monocytogenes: Absent in 25 g
	*CFU: Colony Forming Units

	Authorised Novel Food	Specifications
▼ <u>M37</u>		
]	Phenylcapsaicin	Description/Definition:
	, v. p. v.	Phenylcapsaicin (<i>N</i> -[(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): ≥ 98 %
		Moisture: ≤ 0,5 %
		Total synthesis related production by-products: ≤ 1,0 %
		N,N-dimethyl formamide: ≤ 880 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: ≤ 10 CFU/g
		Escherichia coli: Negative/10 g
		Salmonella sp.: Negative/10 g
		Yeast and mould: ≤ 10 CFU/g
		CFU: Colony Forming Units
		,

▼M112

02017R2470 -
-EN - 01.05.2024
-046.001 - 252

Phosphated	wheat	starch

Authorised Novel Food

Phosphated maize starch

Description:

pH: 4,5-7,5

Phosphated distarch phosphate produced from wheat starch (phosphated wheat starch) is a chemically modified resistant starch derived from wheat starch by combining chemical treatments to create phosphate cross-links within and between individual starch molecules.

Specifications

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 near white free flowing powder.

Residual bound phosphorus: ≤ 0.4 % (as phosphorus) 'high amylose maize' as source

Characteristic/Composition:

CAS No: 11120-02-8

Description/Definition:

CAS No: 11120-02-8

Loss on drying: 10-14 %

Dietary fibre: $\geq 70 \%$ Starch: 7-14 % Protein: $\leq 0.8 \%$ Lipids: $\leq 0.8 \%$

Chemical formula: $(C_6H_{10}O_5)_n [(C_6H_9O_5)_2PO_2H]_x [(C_6H_9O_5)PO_3H_2]_v$

n = number of glucose units; x, y = degrees of substitution

The novel food ingredient is a white or nearly white powder.

n = number of glucose units; x, y = degrees of substitutionThe chemical characteristics of phosphated distarch phosphate:

Chemical formula: $(C_6H_{10}O_5)_n [(C_6H_9O_5)_2PO_2H]x [(C_6H_9O_5)PO_3H_2]y$

Parameter	Powder form 1	Powder form 2
Phosphated Distarch Phosphate (Dry basis)	≥ 85 %	≥ 75 %
Unmodified Wheat Starch (Dry basis)	≤ 15 %	≤ 25 %
Moisture	9-12 %	

▼<u>M112</u>

Authorised Novel Food	Spe	ecifications	
	Total dietary fibre (dry matter basis)	≥ 76,0 %	≥ 66,0 %
	Ash	≤ 3	%
	Protein	≤ 0,	5 %
	Total fat	≤ 0,50 %	≤ 0,34 %
	Residual bound phosphorus	≤ 0,4 % (as	phosphorus)
	pH (25 % slurry)	4,5 -	- 6,5
	Heavy metals:	1	
	Arsenic: ≤ 1 mg/kg		
	Lead: ≤ 2 mg/kg		
	Mercury: ≤ 0,1 mg/kg		
	Microbiological criteria:		
	Total viable aerobic counts: ≤ 10 ⁴ CFU/g		
	Total yeast and mould count: ≤ 200 CFU/g		
	Escherichia coli: Negative to test		
	Salmonella spp.: Negative to test		
	CFU: Colony Forming Units		
19			
Phosphatidylserine from fish	Description/Definition:		
phospholipids	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: ≥ 75 %		
	Phosphatidylserine: ≥ 35 %		
	Glycerides: < 4,0 %		
	Free L-serine: < 1,0 %		
	Tocopherols: $< 0.5 \% (^1)$		
	Peroxide value (PV): < 5,0 meq O ₂ /kg		
	(1) Tocopherols may be added as antioxidants according to Commission Regulation	(EU) No 1129/2011	

Authorised Novel Food Specifications	
Phosphatidylserine from soya	Description/Definition:
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: ≥ 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 equal amounts of phosphati-	Description/Definition:
dylserine and phosphatidic acid	The product is manufactured through enzymatic conversion of soy lecithin. The phospholipid product is a highly concentrated, yellow-brown powder form of phosphatidylserine and phosphatidic acid at an equal level.
	Specification of the product:
	Moisture: ≤ 2,0 %

Authorised Novel Food	Specifications
	Total phospholipids: ≥ 70 %
	Phosphatidylserine: ≥ 20 %
	Phosphatidic acid: ≥ 20 %
	Glycerides: ≤ 1,0 %
	Free L-serine: ≤ 1,0 %
	Tocopherols: $\leq 0.3\%$
	Phytosterols: $\leq 2.0 \%$
	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	Description/Definition:
	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 %

Authorised Novel Food	Specifications
	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: ≥ 800 mg/g
hydrolysates thereof	Protein (N * 6,25): \geq 600 mg/g (dry substance)
	Ash: $\leq 400 \text{ mg/g (dry substance)}$
	Glycoalkaloid (total): ≤ 150 mg/kg
	Lysinoalanine (total): ≤ 500 mg/kg
	Lysinoalanine (free): ≤ 10 mg/kg
Prolyl oligopeptidase (enzyme	Specification of the enzyme:
preparation)	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

Authorised Novel Food	Specifications
	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: ≤ 1,0 mg/kg
	Arsenic: ≤ 1,0 mg/kg
	Cadmium: ≤ 0,5 mg/kg
	Mercury: $\leq 0.1 \text{ mg/kg}$
	Microbiological criteria:
	Total aerobic plate count: $\leq 10^3$ CFU/g
	Total yeasts and moulds: $\leq 10^2 \text{ CFU/g}$
	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: Absent Mycotoxins: Below limits of detection: Aflatoxin B1, B2, G1, G2 (< 0,25 µg/kg), total Aflatoxins (< 2,0 µg/kg), Ochratoxin A (< 0,20 µg/kg), T-2 Toxin (< 5 µg/kg), Zearalenone (< 2,5 µg/kg), Fumonisin B1 and B2 (< 2,5 µg/kg)
	(¹) PPI – Protease Picomole International
	(²) PPU – Prolyl Peptidase Units or Proline Protease Units

02017
7R247(
) — E
$\sqrt{-01}$
.05.202
4 - 0
46.001
-258

Authorised Novel Food	Specifications
-----------------------	----------------

▼ M136

Protein concentrate from Lemna gibba and Lemna minor

Description/Definition:The novel food is a protein concentrate produced from the *Lemna gibba* (70–100 %) and *Lemna minor* (0–30 %) plant species. The manufacturing process of the protein concentrate involves mechanical separation of the protein fraction from insoluble fibres, followed by precipitation under acidic conditions, pasteurisation and spray drying.

The cultivation is carried out in basins in greenhouses under controlled conditions. The water used for the cultivation is filtered and UV-treated. The cultivation conditions are monitored to control the growth of algae, yeast and fungi. The pH is maintained between 5,5 and 6,5.

Characteristics/composition:

Appearance: green powder

Moisture: 1,5-8 %

Protein (Nx6,25): 60-75 %

Ash: 4-12 %

Fat: 2-11 %

Fibre: 6-17 % Ash: 4-12 %

Vitamins:

β-Carotene: < 755 mg/kg

Vitamin K_1 (Phylloquinone): < 16 mg/100 g

Minerals:

Boron: < 10 mg/kg

Copper: < 12 mg/kg

Molybdenum: < 40 mg/kg

Iron: < 670 mg/kg

Zinc: < 50 mg/kg

Manganese: < 100 mg/kg

Antinutritional factors:

Oxalic acid: < 1 900 mg/kg

▼<u>M136</u>

Authorised Novel Food	Specifications
	Heavy metals:
	Lead (mg/kg): ≤ 0.3
	Cadmium (mg/kg): ≤ 0.2
	Mercury (mg/kg): ≤ 0.1
	Arsenic (mg/kg): ≤ 0.2
	Cyanotoxins:
	Microcystins-/Nodularin: < 0,19 mg/kg
	Other contaminants:
	Lysino-alanine (bound): < 500 mg/kg
	Lysino-alanine (free): < 10 mg/kg
	Nitrate: < 3 000 mg/kg
	Pesticides:
	Pesticide levels in accordance with Code number 0254000 ('Subgroup (d) watercresses' in the group of Leaf vegetables, herbs and edible flowers) set out in Regulation (EC) No 396/2005.
	Microbiological criteria:
	Total colony count: < 10 ⁴ CFU/g
	Bacillus cereus: < 100 CFU/g
	Clostridium perfringens: < 100 CFU/g
	Coagulase-positive Staphylococci: < 100 CFU/g
	Escherichia coli: < 10 CFU/g
	Enterobacteriaceae: < 10 CFU/g
	Listeria monocytogenes: Not detected in 25 g
	Salmonella spp.: Not detected in 25 g
	Yeasts and moulds: < 10 CFU/g
	CFU: colony forming units.

020
17R2
170 —
- EN
— 01.
1.05.20
- 01.05.2024 -
- 04 -
6.001
-260

Authorised Novel Food Specifications

▼<u>M11</u>8

Protein 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 extract 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 extract with natural content of DAO (E.C. 1.4.3.22) in an

enteric coated formulation:

Description/Definition:

The protein extract is obtained from homogenised pig kidneys through a series of steps involving a number of acetone washes to defat and dehydrate the homogenized pig kidneys, followed by draining, drying, milling, and sieving to produce a powder containing essentially proteins with a 7-9 % (on average) content of the enzyme diamine oxidase (enzyme nomenclature E.C. 1.4.3.22). The pig kidney extract powder is formulated either in enteric coated capsules or as encapsulated enteric coated pellets or enteric coated tablets to reach the active sites of digestion.

Basic Product:

Specification: pig kidney protein extract with natural content of Diamine oxidase (DAO):

Physical condition: powder

Colour: pale brown

Enzymatic activity: ≥ 0,10 mU/mg (UHPLC-FLD (Ultra High Performance Liquid Chromatography linked with Fluorescent Detection)).

Humidity: < 10 %

Residual Solvents:

Acetone: < 5 000 mg/kg Microbiological criteria:

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

▼<u>M118</u>

Authorised Novel Food	Specifications	
Authorised Novel Food	Physical condition: solid Colour: yellow grey Appearance: micropellets or tablets Enzymatic activity: 110-220 kHDU DAO/g pellet or g tablet (DAO REA (DAO Radio	Listeria monocytogenes: absence in 25 g Final product: Specification pig kidney protein extract with natural content of DAO (E.C. 1.4.3.22) in an enteric coated formulation:
	Extraction Assay)) 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 Radio Extraction Assay)) Humidity: < 10 % Microbiological criteria: Staphylococcus aureus: < 100 CFU/g	Physical condition: solid Colour: pale brown Appearance: micropellets, capsules, or tablets Enzymatic activity (micropellets, capsules or tablets): 2,29 – 4,6 mU/g pellet or g tablet or g capsule (UHPLC-FLD (Ultra High Performance Liquid Chromatography linked with Fluor-
	Escherichia coli: < 10 CFU/g Total aerobic microbiological count: < 10 ⁴ CFU/g Total combined yeasts/moulds count: < 10 ³ CFU/g Salmonella: Absence/10g	escent Detection)). Acid stability 15 min 0,1M HCl followed by 60 min Borat pH = 9,0: > 1,4 mU DAO/g pellet or g tablet or g capsule (UHPLC-FLD (Ultra High Performance Liquid Chromatography linked with Fluorescent Detection)) Humidity: < 10 %
	Bile salt resistant enterobacteriaceae: < 10 ² CFU/g PCR: Polymerase Chain Reaction; HDU (Histamine Degrading Units);	Microbiological criteria: Staphylococcus aureus: < 100 CFU/g Escherichia coli: < 10 CFU/g Total aerobic microbiological count: < 10 ⁴ CFU/g Total combined yeasts/moulds count: < 10 ³ CFU/g Salmonella: Absence/10g Bile salt resistant enterobacteriaceae: < 10 ² CFU/g
		Listeria monocytogenes: absence in 25 g mU: milliUnit (expressed in mU/mg) measures nanomols (nmol) of histamine degraded by the DAO per minute using Ultra High Performance Liquid Chromatography linked with Fluorescent Detection (UHPLC-FLD) (O. Comas-Basté et al. Analytical and Bioanalytical Chemistry 411:7595-7602 (2019)). 1 mU corresponds to 48 000 HDU of the DAO Radio Extraction Assay (REA) method.

Authorised Novel Food	Specifications
Pyrroloquinoline quinone disodium	Definition:
salt	Chemical name: disodium 9-carboxy-4,5-dioxo-1 <i>H</i> -pyrrolo[5,4-f]quinoline-2,7-dicarboxylate
	Chemical formula: C ₁₄ H ₄ N ₂ Na ₂ O ₈
	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 <i>Hyphomicrobium denitrificans</i> s 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: $\leq 0.05 \%$
	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

Authorised Novel Food	sed Novel Food Specifications	
Rapeseed oil high in unsapo-	Description/Definition:	
ifiable matter	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 triglyceride 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: ≤ 6,0 mg KOH/g	
	Peroxide value (PV): ≤ 10 mEq O ₂ /kg	
	Heavy metals:	
	Iron (Fe): $< 1\ 000\ \mu g/kg$	
	Copper (Cu): < 100 μg/kg	
	Impurities:	
	Polycyclic aromatic hydrocarbons (PAH) Benzo(a)pyrene: < 2 μg/kg	
	Treatment with active carbon is required to ensure that polycyclic aromatic hydrocarbons (PAH) are not enriched in the production of 'rapeseed oil high i unsaponifiable matter.	

Authorised Novel Food	Specifications
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 I
	Description:
	White to off-white, spray dried powder
	Total protein: ≥ 90 %
	Soluble protein: ≥ 85 %
	Moisture: $\leq 7.0 \%$
	Carbohydrates: ≤ 7,0 %
	Fat: $\leq 2.0 \%$
	Ash: ≤ 4,0 %
	Fibre: ≤ 0,5 %
	Total glucosinolates: ≤ 1 mmol/kg
	Purity:
	Total phytate: ≤ 1,5 %
	Lead: ≤ 0,5 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

▼<u>M17</u>

Authorised Novel Food	Specifications
1	
-	
Refined shrimp peptide	Description
concentrate	Refined shrimp peptide concentrate is a peptide mixture obtained from northern shrimp (<i>Pandalus borealis</i>) shells and heads via a series of purification steps following enzymatic proteolysis using a protease from <i>Bacillus licheniformis</i> and/or <i>Bacillus amyloliquefaciens</i> .
	Characteristics/Composition
	Total Dry matter (%): ≥ 95,0 %
	Peptides (w/weight dry matter): ≥ 87,0 % of which peptides with molecular weight < 2 kDa: ≥ 99,9 %
	Fat (w/w) : $\leq 1,0 \%$
	Carbohydrates (w/w): $\leq 1.0 \%$
	Ash (w/w): $\leq 15.0 \%$
	Calcium: ≤ 2,0 %
	Potassium: ≤ 0,15 %
	Sodium: ≤ 3,5 %
	Heavy Metals
	Arsenic (inorganic): ≤ 0,22 mg/kg
	Arsenic (organic): ≤ 51,0 mg/kg
	Cadmium: ≤ 0,09 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: ≤ 20 CFU/g
	CFU: Colony Forming Units
	ND: Not Detectable

Authorised Novel Food	Specifications
Trans-resveratrol	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: C ₁₄ H ₁₂ O ₃
	Molecular weight: 228,25 Da
	CAS No: 501-36-0
	Purity:
	Trans-resveratrol: $\geq 98\%-99\%$
	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
	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.

Authorised Novel Food	Specifications
	Hyaluronic acid: 60-80 %
	Chondroitin sulphate A: ≤ 5,0 %
	Dermatan sulphate (chondroitin sulphate B): ≤ 25 %
	pH: 5,0-8,5
	Purity:
	Chlorides: ≤ 1,0 %
	Nitrogen: $\leq 8.0 \%$
	Loss on drying: (105 °C for 6 hours): \leq 10 %
	Heavy metals:
	Mercury: $\leq 0.1 \text{ mg/kg}$
	Arsenic: ≤ 1,0 mg/kg
	Cadmium: ≤ 1,0 mg/kg
	Chromium: ≤ 10 mg/kg
	Lead: $\leq 0.5 \text{ mg/kg}$
	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	Description/Definition:
volubilis	Sacha inchi oil is a 100 % cold pressed vegetable oil obtained from the seeds of <i>Plukenetia volubiis</i> 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

Authorised Novel Food	Specifications
	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 meq O ₂ /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.
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 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 %

	Authorised Novel Food	Specifications
		MOLE % SCFA (short chain fatty acids): 30-67 % Saturated long chain fatty acids: < 70 % by weight Trans fatty acids: ≤ 1,0 % Free fatty acids as oleic acid: ≤ 0,5 % Triacylglycerol profile: Triesters (short/long of 0,5 to 2,0): ≥ 90 % Triesters (short/long = 0): ≤ 10 % Unsaponifiable material: ≤ 1,0 % Moisture: ≤ 0,3 % Ash: ≤ 0,1 % Colour: ≤ 3,5 Red (Lovibond) Peroxide value (PV): ≤ 2,0 Meq/Kg
	Schizochytrium sp. oil rich in DHA and EPA	Acid value: ≤ 0,5 mg KOH/g Peroxide value (PV): ≤ 5,0 meq/kg oil Oxidative stability: All food products containing <i>Schizochytrium sp.</i> oil rich in DHA and EPA should demonstrate oxidative stability by appropriate and recognised national/international test methodology (e.g. AOAC) Moisture and volatiles: ≤ 0,05 % Unsaponifiables: ≤ 4,5 % Trans-fatty acids: ≤ 1 % DHA content: ≥ 22,5 % EPA content: ≥ 10 %
M27	Schizochytrium 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 %

V 1V17		
	Authorised Novel Food	Specifications
▼ <u>M65</u>		
	Schizochytrium sp. (WZU477) oil	Description/Definition:
		The novel food is an oil produced from the strain WZU477 of the microalgae <i>Schizochytrium</i> sp. Composition:
		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 %
		Docosahexaenoic acid (DHA): ≥ 32,0 %
		p-anisidine value: ≤ 10
▼ <u>M23</u>		
	Syrup from Sorghum bicolor (L.)	Description/Definition
	Moench.	The traditional food is syrup from Sorghum bicolor (L.) Moench (genus, Sorghum; family, Poaceae (alt. Gramineae)).
	(Traditional food from a third country)	The syrup is obtained from stalks of <i>S. bicolor</i> , 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
▼ <u>M9</u>		
	Fermented soybean 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 K ₂ 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

	Authorised Novel Food	Specifications
		Condition: No offensive taste or smell Loss on drying: $\leq 10 \%$ Vitamin K_2 : $\leq 0,1 \text{ mg/kg}$ Heavy metals:
		Lead: ≤ 5,0 mg/kg
		Arsenic: ≤ 3,0 mg/kg
		Microbiological criteria:
		Total viable aerobic count: $\leq 10^3 \text{ CFU}(^3)/\text{g}$
		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).
▼ <u>M57</u>		
	Selenium-containing yeast (Yarrowia lipolytica) biomass	Description/Definition:
		The novel food is the dried and heat-killed selenium-containing biomass of the yeast <i>Yarrowia lipolytica</i> .
		The novel food is produced by fermentation in the presence of sodium selenite followed by a number of purification steps including a heat-killing step of the yeast to ensure the absence of viable <i>Yarrowia lipolytica</i> cells in the novel food.
		Characteristics/Composition:
		Total selenium: 165–200 μg/g
		Se-methionine (¹³): 100–140 μg/g
		Protein: 40–50 g/100 g
		Dietary fibre: 24–32 g/100 g
		Sugars: < 1 g/100 g
		Fat: 6–12 g/100 g
		Total ash: ≤ 15 %
		Water: ≤ 5 %
		Dry matter: ≥ 95 %

▼<u>M57</u>

	Authorised Novel Food	Specifications
		Heavy metals:
		Lead: $\leq 3.0 \text{ mg/kg}$
		Cadmium: ≤ 1,0 mg/kg
		Mercury: ≤ 0.1 mg/kg
		Microbiological criteria:
		Total aerobic microbial count: $\leq 5 \times 10^3$ CFU/g
		Total yeast and mould count: $\leq 10^2$ CFU/g
		Viable <i>Yarrowia lipolytica</i> cells (¹⁴): < 10 CFU/g (i.e. limit of detection)
		Coliforms: ≤ 10 CFU/g
		Salmonella spp.: Absence in 25 g
		CFU: colony forming units
▼ <u>M61</u>		
	3'-Sialyllactose (3'-SL) sodium salt	Description:
	(microbial source)	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: C ₂₃ H ₃₈ NO ₁₉ Na
		Chemical name: N -Acetyl- α -D-neuraminyl- $(2\rightarrow 3)$ - β -D-galactopyranosyl- $(1\rightarrow 4)$ -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): ≥ 90,0 % (w/w)
		3'-Sialyllactose sodium salt (% of dry matter): ≥ 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: ≤ 1,0 % (w/w)
		pH (20 °C, 5 % solution): 4,5 -6,0
		Residual protein: $\leq 0.01 \%$ (w/w)

▼<u>M61</u>

Authorised Novel Food	Specifications
	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: ≤ 100 CFU/g
	Residual endotoxins: ≤ 10 EU/mg
	CFU: Colony Forming Units; EU: Endotoxin Units.
<u>05</u>	
3'-Sialyllactose ('3'-SL') sodium	Description:
salt	3'-Sialyllactose (3'-SL) sodium salt is a purified, white to off-white powder or agglomerate produced by a microbial process and contains limited levels
(produced by derivative strains of	lactose, 3'-sialyl-lactulose, and sialic acid.
E. coli BL21(DE3))	Definition:
	Chemical name: N-Acetyl- α -D-neuraminyl- $(2\rightarrow 3)$ - β -D-galactopyranosyl- $(1\rightarrow 4)$ -D-glucose, sodium salt
	Chemical formula: C ₂₃ H ₃₈ NO ₁₉ Na
	Molecular mass: 655,53 Da
	CAS No: 128596-80-5
	Source: Two genetically modified strains (a production strain and an optional degradation strain) of Escherichia coli BL21(DE3)
	Characteristics/Composition:
	3'-Sialyllactose sodium salt (% of dry matter): ≥ 88.0 % (w/w)
	3'-Sialyl-lactulose (% of dry matter): ≤ 5.0 % (w/w)
	D-Lactose (% of dry matter): ≤ 5.0 % (w/w)
	Sialic acid (% of dry matter): ≤ 1.5 % (w/w)
	N-acetyl-D-glucosamine (% of dry matter): ≤ 1,0 % (w/w)
	Sum of other carbohydrates (% of dry matter) ^a : ≤ 5.0 % (w/w)
	Moisture: $\leq 9.0 \%$ (w/w)
	Ash: $\leq 8.5 \%$ (w/w)
	Residual protein: $\leq 0.01 \%$ (w/w)
	Sodium: $\leq 4.2 \% \text{ (w/w)}$
	Microbiological criteria:
	Standard plate count: ≤ 1 000 *CFU/g
	Enterobacteriaceae: ≤ 10 CFU/g
	Salmonella spp.: Absence in 25 g

▼<u>M105</u>

	Authorised Novel Food	Specifications
		Yeast and mould: ≤ 100 CFU/g
		Cronobacter (Enterobacter) sakazakii: Absence in 10 g
		Residual endotoxins: ≤ 10 **EU/mg
		Sum of other carbohydrates = 100 (% (w/w) of dry matter) – 3'-Sialyllactose sodium salt (% (w/w) of dry matter) – quantified carbohydrates (% (w/w) of dry matter) – Ash (% (w/w) of dry matter);
		* CFU: Colony Forming Units;
		** EU: Endotoxin Units
▼M135		
111100	•	
	3'-Sialyllactose (3'-SL) sodium salt	Description:
	(produced using a derivative strain	3'-Sialyllactose (3'-SL) sodium salt is a purified, concentrated, white to off-white powder that is produced by a microbial process. It contains limited levels of
	of E. coli W (ATCC 9637))	Sialic acid, D-Lactose, D-Glucose, and 3'-Sialyllactulose and 6'-Sialyllactose sodium salts.
		Source:
		Genetically modified strain of Escherichia coli W (ATCC 9637)
		Definition:
		Chemical formula: C ₂₃ H ₃₈ NO ₁₉ Na
		Chemical name: N-Acetyl- α -D-neuraminyl- $(2\rightarrow 3)$ - β -D-galactopyranosyl- $(1\rightarrow 4)$ -D-glucose, sodium salt
		Molecular mass: 655,53 Da
		CAS N°: 128596-80-5
		Characteristics/Composition:
		3'-Sialyllactose sodium salt (% w/w of dry matter): ≥ 82,0
		Sialic acid (% w/w of dry matter): ≤ 6,0
		D-Lactose (% w/w of dry matter): ≤ 3,0
		D-Glucose (% w/w of dry matter): ≤ 3,0
		Sum of 3'- Sialyllactulose and 6'-Sialyllactose sodium salts (% w/w of dry matter): ≤ 5,0
		Sum of other carbohydrates ^a (% w/w of dry matter): ≤ 12,0
		Moisture (% w/w): ≤ 10.5
		Sodium (% w/w): ≤ 5.0
		pH (25 °C, 5 % solution): 4,5 -7,5
		Residual protein (% w/w): ≤ 0.01
		Heavy metals and contaminants:
		Arsenic (mg/kg): ≤ 0.2
		Lead (mg/kg): ≤ 0.2
		Cadmium (mg/kg): ≤ 0.2

▼<u>M135</u>

•	Authorised Novel Food	Specifications
•		Mercury (mg/kg): ≤ 0.1
		Aflatoxin M1: $< 0.025 (\mu g/kg)$
		Microbiological criteria:
		Total plate count: ≤ 1 000 CFU/g
		Enterobacteriaceae: Absence in 10 g
		Cronobacter spp.: Absence in 10 g
		Salmonella spp.: Absence in 25 g
		Yeasts and moulds: ≤ 100 CFU/g
		Listeria monocytogenes: Absence in 25 g
		Presumptive Bacillus cereus: ≤ 50 CFU/g
		Residual endotoxins: ≤ 10 EU/mg
		^a Sum of other carbohydrates = 100 % w/w of dry matter – 3'-Sialyllactose (acid, % w/w of dry matter) –quantified carbohydrates ((% w/w of dry matter), Sialic acid + D-Lactose + D-Glucose + (3'- Sialyllactulose and 6'-Sialyllactose (acids)) – sodium (w/w of dry matter); CFU: Colony Forming Units; EU: Endotoxin Units
▼ <u>M60</u>		
	6'-Sialyllactose ('6'-SL') sodium	Description:
	salt	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
	(microbial source)	of lactose, 6'-sialyl-lactulose, and sialic acid.
		Source: Genetically modified strain of Escherichia coli K-12 DH1
		Definition:
		Chemical formula: C ₂₃ H ₃₈ NO ₁₉ Na
		Chemical name: N-Acetyl-α-D-neuraminyl-(2→6)-β-D-galactopyranosyl-(1→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): ≥ 94,0 % (w/w)
		6'-Sialyllactose sodium salt (% of dry matter): ≥ 90,0 % (w/w)
		D-Lactose: \(\le 5,0 \% \) (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: $\leq 0.01 \%$ (w/w)

▼<u>M60</u>

Authorised Novel Food	Specifications
	Microbiological criteria:
	Aerobic mesophilic bacteria total plate count: ≤ 1 000 CFU/g
	Enterobacteriaceae: ≤ 10 CFU/g
	Salmonella sp.: Absence in 25 g
	Yeast: ≤ 100 CFU/g
	Mould: ≤ 100 CFU/g
	Residual endotoxins: ≤ 10 EU/mg
	CFU: Colony Forming Units; EU: Endotoxin Units.
<u>5</u>	
6'-Sialyllactose ('6'-SL') sodium	Description:
salt (produced by derivative strains of	6'-Sialyllactose (6'-SL) sodium salt is a purified, white to off-white powder or agglomerate produced by a microbial process and contains limited levels of lactose, 6'-sialyl-lactulose, and sialic acid.
E. coli BL21(DE3))	Definition:
	Chemical name: N-Acetyl-α-D-neuraminyl-(2→6)-β-D-galactopyranosyl-(1→4)-D-glucose, sodium salt
	Chemical formula: C ₂₃ H ₃₈ NO ₁₉ Na
	Molecular mass: 655,53 Da
	CAS No: 157574-76-0
	Source: Two genetically modified strains (a production strain and an optional degradation strain) of Escherichia coli BL21(DE3)
	Characteristics/Composition:
	6'-Sialyllactose sodium salt (% of dry matter): ≥ 90,0 % (w/w)
	6'-Sialyl-lactulose (% of dry matter): ≤ 3,0 % (w/w)
	D-Lactose (% of dry matter): $\leq 5,0$ % (w/w)
	Sialic acid (% of dry matter): ≤ 2.0 % (w/w)
	N-acetyl-D-glucosamine (% of dry matter): ≤ 3,0 % (w/w)
	Sum of other carbohydrates (% of dry matter) (28): ≤ 5.0 % (w/w)
	Moisture: $\leq 9.0 \%$ (w/w)
	Ash: $\leq 8.5 \%$ (w/w)
	Residual protein: $\leq 0.01 \%$ (w/w)
	Sodium: $\leq 4.2 \%$ (w/w)
	Contaminants:
	Arsenic: $\leq 0.2 \text{ (mg/kg)}$
	Aflatoxin M1: $\leq 0.025 \; (\mu g/kg)$

▼<u>M115</u>

Authorised Novel Food	Specifications
	Microbiological criteria:
	Standard plate count: ≤ 1 000 CFU/g
	Enterobacteriaceae: ≤ 10 CFU/g
	Salmonella spp.: Absence in 25 g
	Yeast and mould: ≤ 100 CFU/g
	Cronobacter spp.: Absence in 10 g
	Residual endotoxins: ≤ 10 EU/mg
27	
6' -Sialyllactose (6'-SL) sodium salt	Description:
(produced by derivative strain of <i>E. coli</i> W (ATCC 9637))	6'-Sialyllactose (6'-SL) sodium salt is a purified, white to off-white powder that is produced by a microbial process, is further isolated, purified a concentrated. It contains limited levels of Sialic acid, D-Lactose, D-Glucose, 6'-Sialyllactulose, and 3'-Sialyllactose sodium salt.
	Source: Genetically modified strain of Escherichia coli W (ATCC 9637)
	Definition:
	Chemical formula: C ₂₃ H ₃₈ NO ₁₉ Na
	Chemical name: N-Acetyl-α-D-neuraminyl-(2→6)-β-D-galactopyranosyl-(1→4)-D-glucose, sodium salt
	Molecular mass: 655,53 Da
	CAS No 157574-76-0
	Characteristics/Composition:
	6'-Sialyllactose sodium salt (% w/w of dry matter): ≥ 82,0
	Sialic acid (% w/w of dry matter): ≤ 6.0
	D-Lactose (% w/w of dry matter): ≤ 3,0
	D-Glucose (% w/w of dry matter): ≤ 3.0
	Sum of 6'- Sialyllactulose and 3'-Sialyllactose sodium salt (% w/w of dry matter): ≤ 5,0
	Sum of other carbohydrates ^a (% w/w of dry matter): ≤ 13,0
	Moisture (% w/w): ≤ 10.5
	Sodium (% w/w): ≤ 5.0
	pH (25 °C, 5 % solution): 4,5–7,5
	Residual protein (% w/w): ≤ 0.01
	Heavy metals and contaminants:
	Arsenic (mg/kg): ≤ 0.2
	Aflatoxin M1: $< 0.025 (\mu g/kg)$

▼<u>M127</u>

	Authorised Novel Food	Specifications
		Microbiological criteria:
		Total plate count: ≤ 1 000 CFU/g
		Enterobacteriaceae: Absence in 10 g
		Cronobacter spp.: Absence in 10 g
		Salmonella spp.: Absence in 25 g
		Yeasts and moulds: ≤ 100 CFU/g
		Listeria monocytogenes: Absence in 25 g
		Presumptive Bacillus cereus: ≤ 50 CFU/g
		Residual endotoxins: ≤ 10 EU/mg
		^a Sum of other carbohydrates = 100 % w/w of dry matter – 6'-Sialyllactose (acid, % w/w of dry matter) –quantified carbohydrates ((% w/w of dry matter), Sialic acid + D-Lactose + D-Glucose + (6'- Sialyllactulose and 3'-Sialyllactose (acids)) – sodium (w/w of dry matter); CFU: Colony Forming Units; EU: Endotoxin Units
▼ M43		
	Spermidine-rich wheat germ extract (Triticum aestivum)	Description/Definition: Spermidine-rich wheat germ extract is obtained from non-fermented, non-sprouting wheat germs (<i>Triticum aestivum</i>) by the process of solid-liquid 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 μg/g Putrescine: < 0,3 mg/g Cadaverine: ≤ 16,0 μg/g Mycotoxins: Aflatoxins (total): < 0,4 μg/kg Microbiological criteria: Total aerobic bacteria: < 10 000 CFU/g Yeast and moulds: < 100 CFU/g Salmonella: Absence/25g Listeria monocytogenes: Absence/25g
▼ M9		
	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 <i>Leuconostoc citreum</i> or by means of a recombinant strain of the production organism <i>Bacillus licheniformis</i> . The resulting oligosaccharides are characterised by the presence of α-(1→6) and α-(1→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 %

Authorised Novel Food	Specifications		
	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 non-cellulosic components, bleaching of purified fibres, acid washing and neutralization. Moisture: ≤ 7,0 % Ash: ≤ 0,3 % Total Dietary Fibre (AOAC) dry basis (all insoluble): ≥ 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): ≤ 1,0 Arsenic (ppm): ≤ 1,0 Arsenic (ppm): ≤ 1,0 Microbiological criteria: Yeast and moulds (CFU/g): ≤ 1 000 Salmonella: Absence Listeria monocytogenes: Absence		

▼<u>M53</u>

Authorised Novel Food	Specifications
3	
=	
Sugars obtained from cocoa	Description/Definition:
(Theobroma cacao L.) pulp	Sugars are obtained from the concentrated cocoa pulp (<i>Theobroma cacao</i> 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): < 10 ⁴
	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 (<i>Theobroma cacao</i> L.) pulp:
	Total Plate Count (aerobic) (cfu/g): < 10 ⁴
	Salmonella spp.: Absence in 25 g

Authorised Novel Food	Specifications		
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 th sunflower, <i>Helianthus Annuus</i> L.		
	Composition:		
	Oleic acid (C18:1): 20 %		
	Linoleic acid (C18:2): 70 %		
	Unsaponifiable matter: 8,0 %		
	Phytosterols: 5,5 %		
	Tocopherols: 1,1 %		
73			
Synsepalum dulcificum dried fruits	Description/Definition:		
	The novel food is lyophilised pulp and skin of pitted fruits of <i>Synsepalum dulcificum</i> (Schumach. & Thonn.) Daniell that belongs to the Sapotaceae famil The resulting dried cake is milled into a powder.		
	Characteristics/Composition:		
	Moisture $(g/100 \text{ g})$: < 6		
	Ash (g/100 g): 3,5-8,5		
	Total carbohydrates (g/100 g): 70-87		
	Sugars (g/100 g): 50-75		
	Fibre (g/100 g): 1-6,5		
	Total protein (g/100 g): 3,5-6,0		
	Miraculin (16) (g/100 g): 1,5-2,5		
	Total fat (g/100 g): 0,50-3,50		
	Microbiological criteria:		
	Total aerobic colony count: < 10 ⁴ CFU (⁷)/g		
	Bacillus cereus (presumptive): < 100 CFU/g		
	Sulfite-reducing Clostridia: ≤ 30 CFU/g		
	Total Enterobacteriaceae: < 100 CFU/g		
	Yeasts and moulds: < 500 CFU/g		

▼M73

Authorised Novel Food	Specifications	
	Pesticides:	
	Pesticide levels in accordance with Code number 0820990 ('others' in the group of fruit spices) set out in Regulation (EC) No 396/2005 (17)	

▼<u>M66</u>

Dried *Tenebrio molito*r larva (yellow mealworm)

Description/Definition:

The novel food is the whole, thermally dried yellow mealworm, either whole (blanched, oven-dried larva) or in the form of a powder (blanched, oven-dried, ground larva). The term 'mealworm' refers to the larval form of *Tenebrio molitor*, an insect species that belongs to the family of *Tenebrionidae* (darkling beetles).

The entire mealworms are meant for human consumption and no parts are removed.

A minimum 24 hours fasting period is required before the thermal drying step, to allow the larvae to discard their bowel content.

Characteristics/Composition:

Ash (% w/w): 3,5 - 4,5

Moisture (% w/w): 1-8

Crude protein (N x 6,25) (% w/w): 56-61

Digestible Carbohydrates (15) (% w/w): 1-6

Fat (% w/w): 25-30

of which saturated (% w/w): 4-9

Peroxide value (Meq O_2/kg fat): ≤ 5

Dietary fibre (% w/w): 4-7

Chitin (% w/w): 4-7

Heavy metals:

Lead: $\leq 0.075 \text{ mg/kg}$

Cadmium: ≤ 0.1 mg/kg

Mycotoxins:

Aflatoxins (Sum of B1, B2, G1, G2): ≤ 4 µg/kg

Aflatoxin B1: $\leq 2 \mu g/kg$

Deoxynivalenol: ≤ 200 μg/kg

Ochratoxin A: $\leq 1 \mu g/kg$

Tenebrio Linnaeus.	

▼<u>M81</u>

Frozen, dried and powder forms of yellow mealworm (*Tenebrio molitor* larva)

Authorised Novel Food

Description/Definition:

Microbiological criteria:

Yeasts and moulds: ≤ 100 CFU/g Escherichia coli: ≤ 50 CFU/g

Salmonella spp.: Not detected in 25 g

Total aerobic colony count: $\leq 10^5$ CFU (7)/g

Listeria monocytogenes: Not detected in 25 g
Sulfite-reducing Anaerobes: ≤ 30 CFU/g
Bacillus cereus (presumptive): ≤ 100 CFU/g
Enterobacteriaceae (presumptive): < 10 CFU/g
Coagulase-positive staphylococci: ≤ 100 CFU/g

The novel food are frozen, dried and powder forms of yellow mealworm (*Tenebrio molitor* larva). The term 'mealworm' refers to the larval form of *Tenebrio molitor*, an insect species that belongs to the family of Tenebrionidae (darkling beetles). Another identified scientific synonym is *Tenebrio molitor* Linnaeus.

Specifications

The entire mealworms are meant for human consumption, no parts are removed.

A minimum 24 hours fasting period is required before killing the insects by freezing, to allow the larvae to discard their bowel content.

The novel food is intended to be placed on the market in three different forms, namely: whole, blanched and frozen *T. molitor* larva (frozen); whole, blanched and freeze-dried *T. molitor* larva (dried) which may be in powder form (powder).

Parameters	Frozen	Dried or powder			
Characteristics/Composition					
Ash	0,9-1,10	3,6-4,1			
Moisture (% w/w)	69-75	≤ 5			
Crude protein (N x 6,25) (% w/w)	14-19	54-60			

▼<u>M81</u>

Authorised Novel Food		Specifications	
	Fat (% w/w) — of which saturated fatty acids (% fat)	7-12,5 20-29	27-30 20-29
	Digestible carbohydrates (% w/w)	1-2	4-8
	Dietary fibre (% w/w)	1,2-3,5	4-6
	Chitin(*) (% w/w)	≤ 3	4-9
	Peroxide value (Meq O ₂ /kg fat)	≤ 5	≤ 5
	Contaminants		
	Heavy metals		
	Lead (mg/kg)	≤ 0,01	≤ 0,075
	Cadmium (mg/kg)	≤ 0,05	≤ 0,1
	Mycotoxins		
	Aflatoxins (Sum of B1, B2, G1, G2) (µg/kg)	≤ 4	≤ 4
	Aflatoxin B1 (μg/kg)	≤ 2	≤ 2
	Deoxynivalenol (μg/kg)	≤ 200	≤ 200
	Ochratoxin A (μg/kg)	≤ 1	≤ 1
	Dioxins and PCBs		
	Sum of dioxins and dl-PCBs (UB, WHO-TEQ2005)(**) (pg/g fat)	≤ 0,75	≤ 0,75

0201
7R247
0 — EN
_
01.05.202
)24 —
046.00
01 - 2
286

Authorised Novel Food	Specifications Microbiological criteria		
	Total aerobic colony count (CFU/g)	≤ 10 ⁵	≤ 10 ⁵
	Enterobacteriaceae (presumptive) (CFU/g)	≤ 100	≤ 100
	Escherichia coli (CFU/g)	≤ 50	≤ 50
	Listeria monocytogenes	Absence in 25g	Absence in 25g
	Salmonella spp.	Absence in 25g	Absence in 25g
	Bacillus cereus (presumptive) (CFU/g)	≤ 100	≤ 100
	Coagulase positive Staphylococci (CFU/g	≤ 100	≤ 100
	Sulfite-reducing Anaerobes (CFU/g)	≤ 30	≤ 30
	Yeasts and moulds (CFU/g)	≤ 100	≤ 100

^(*) Chitin calculated as the difference between the Acid Detergent Fibre fraction and the Acid Detergent Lignin fraction (ADF-ADL), as described by Hahn et al. (2018).

CFU: colony forming units.

▼M89

Tetrahydrocurcuminoids

Description:

The tetrahydrocurcuminoids are produced via a series of steps involving the extraction of curcuminoids from the dried, pulverised rhizomes of turmeric (Curcuma longa L.), hydrogenation (using palladium on carbon (Pd/C) as a catalyst), concentration, crystallisation, drying, and milling into a powder.

Characteristics/Composition:

Total tetrahydrocurcuminoids (dry basis) (% w/w): > 95,0

Moisture (% w/w): ≤ 1.0

Ash (\% w/w): ≤ 1.0

Palladium (mg/kg): < 5,0

^(**) Upper bound sum of polychlorinated dibenzo-para-dioxins (PCDDs)-polychlorinated dibenzofurans (PCDFs) and dioxin-like polychlorinated biphenyls (PCBs) expressed as World Health Organization toxic equivalent (using WHO-TEFs of 2005)).

▼<u>M89</u>

	Authorised Novel Food	Specifications
		Microbiological criteria:
		Total aerobic microbial count: ≤ 5 000 CFU/g
		Total yeast/moulds count: ≤ 100 CFU/g
		Escherichia coli: < 10 CFU/g
		Staphylococcus aureus: ≤ 10 CFU/g
		Enterobacteriaceae: ≤ 10 CFU/g
		Salmonella spp.: Absence in 25 g
		Coliforms: ≤ 10 CFU/g
		CFU: Colony Forming Units
▼ M9		
<u> </u>		
	Dried Tetraselmis chuii microalgae	Description/Definition:
		The dried product is obtained from the marine microalgae <i>Tetraselmis chuii</i> , belonging to the <i>Chlorodendraceae</i> 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

Authorised Novel Food	Specifications	
herapon barcoo/Scortum	Description/Definition:	
nerupon bureoo/scoreum	Scortum/ <i>Therapon barcoo</i> 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: <i>Therapon</i> or <i>Scortum barcoo</i>	
	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-3/n-6: 1,5-15,0	
	Total omega 3 acids: 1,6-40,0	
	Total omega 6 acids: 2,6-10,0	
-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	

Authorised Novel Food	Specifications
s	Synonym: D- <i>lyxo</i> -Hexulose
	CAS number: 87-81-0
C	Chemical formula: C ₆ H ₁₂ O ₆
F	Formula weight: 180,16 (g/mol)
P	Purity:
A	Assay: ≥ 98 % on a dry weight basis
L	Loss on drying: ≤ 0,5 % (102 °C, 2 hours)
s	Specific Rotation: $[\alpha]_D^{20}$: -4 to -5.6° (1 % aqueous solution)(1)
N	Melting range: 133– 137 °C
Н	Heavy metals:
L	Lead: $\leq 1.0 \text{ mg/kg(*)}$
(*	(*) 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. 'Instrumental methods' (1).
(¹	(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
►M52 Taxifolin-rich extract ◀ D	Description:
	Taxifolin-rich extract from the wood of Dahurian Larch (<i>Larix gmelinii</i> (Rupr.) Rupr) is a white to pale-yellow powder that crystallizes from hot aqueous solutions.
	► M52 Definition:
	Chemical name: $[(2R,3R)-2-(3,4 \text{ dihydroxyphenyl})-3,5,7-\text{trihydroxy-}2,3-\text{dihydrochromen-}4-one, also called (+) trans (2R,3R)- dihydroquercetin] and with no more than 2 % of the cis-form \blacktriangleleft$
s	Specifications:
P	Physical parameter
N	Moisture: ≤ 10 %Compound analysis
Т	Taxifolin (m/m): ≥ 90.0 % of the dry weight

Authorised Novel Food		Specifications
	Heavy Metals, Pesticide	
	Lead: ≤ 0,5 mg/kg	
	Arsenic: $\leq 0.02 \text{ mg/kg}$	
	Cadmium: $\leq 0.5 \text{ mg/kg}$	
	Mercury: ≤ 0,1 mg/kg	
	Dichlorodiphenyltrichloroethane (Γ	DDT): $\leq 0.05 \text{ mg/kg}$
	Residual solvents	
	Ethanol: < 5 000 mg/kg	
	Microbiological criteria	
	Total Plate Count (TPC): $\leq 10^4$ CF	$\mathrm{FU/g}$
	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	
		e 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

02017R2470
EN-0
01.05.2024 -
-046.001
-291

Description/Definition:

Authorised Novel Food

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

Specifications

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: \geq 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 mm
diameter: 10 mm
temperature: 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:

Authorised Novel Food	Specifications
	% trehalose = $100 \times (R_U/R_S) (W_S/W_U)$
	where
	$R_{\rm S}$ = peak area of trehalose in the standard preparation
	$R_{\rm U}$ = peak area of trehalose in the sample preparation
	W_S = weight in mg of trehalose in the standard preparation
	W_U = weight of dry sample in mg
	Characteristics:
	Identification:
	Solubility: Freely soluble in water, very slightly soluble in ethanol
	Specific rotation: $\left[\alpha\right]_{D}^{20} = +179^{\circ}$ (5 % aqueous solution, dihydrate), $+199^{\circ}$ (5 % aqueous solution, anhydrous substance)
	Melting point: 97 °C (dihydrate)
	Purity:
	Loss on drying: ≤ 1,5 % (60 °C, 5h)
	Total ash: ≤ 0,05 %
	Heavy metals:
	Lead: ≤ 1,0 mg/kg
52	
<u></u>	
UV-treated mushrooms (Agaricus	Description/Definition
bisporus)	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: (3β,5Z,7E,22E)-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 D ₂ in the final product: 5-20 μg/100 g fresh weight at the expiration of shelf life.

Authorised Novel Food	Specifications
<u> </u>	
UV-treated baker's yeast	Description/Definition
(Saccharomyces cerevisiae)	Baker's yeast (Saccharomyces cerevisiae) is treated with ultraviolet light to induce the conversion of ergosterol to vitamin D_2 (ergocalciferol). Vitamin content in the yeast concentrate varies between 800 000 – 3 500 000 IU vitamin D/100 g (200-875 μ g/g).
	The yeast shall be inactivated for use in infant formula and follow-on formula, processed cereal-based food and foods for special medical purposes as defin by Regulation (EU) No 609/2013, while for use in other foods the yeast may or may not be inactivated.
	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 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
	Microbiological criteria for the yeast concentrate
	Coliforms: $\leq 10^3/g$
	Escherichia coli: ≤ 10/g
	Salmonella: Absence in 25 g
<u> </u>	
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 convergosterol to vitamin D ₂ (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/ci
	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 D ₂ (ergocalciferol) in the final product: $0.75-3 \mu g/100 g(^1)$
	Yeast in dough: 1-5 g/100 g (²)
	(¹) EN 12821, 2009, European Standard.
	(²) Recipe calculation.

Authorised Novel Food

Description/Definition:

UV-treated milk

markets in , p. 671).	
mp. The	
ovel food	

		isation. 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 :
		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-methyl-idenecyclohexan-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
▼M51		
•	Vitamin D ₂ mushroom powder	Description/Definition
		Vitamin D ₂ 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_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 D ₂ content: 1 000–1 300 μg/g of mushroom powder (12)
		Moisture: ≤ 10,0 %
		Ash: $\leq 13.5 \%$

Specifications

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 pasteur-

▼<u>M51</u>

Authorised Novel Food	Specifications
	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 µg/kg
	Microbiological criteria:
	Total plate count: $\leq 5~000~\text{CFU}~(^7)/\text{g}$
	Yeast and mould: ≤ 100 CFU/g
	Salmonella sp.: Absent in 25 g
	Staphylococcus aureus: ≤ 10 CFU/g
	Escherichia coli: ≤ 10 CFU/g
	Coliforms: ≤ 10 CFU/g
	Enterobacteriaceae: ≤ 10 CFU/g
	Listeria monocytogenes: Absent in 25 g
▼ <u>M76</u>	
Vitamin D ₂ mushroom powder	Description/Definition:
	The novel food is mushroom powder produced from the dried whole <i>Agaricus bisporus</i> mushrooms. The process includes drying, milling and the controlled exposure of the mushroom powder to UV irradiation.
	UV radiation: A process of radiation in ultraviolet light within a range of wavelength similar to those UV-treated novel foods authorised under Regulation (EU) 2015/2283.
	Characteristics/composition:
	Vitamin D ₂ content: 580-595 μg/g of mushroom powder
	Ash: $\leq 13.5 \%$
	Water activity: < 0,5
	Moisture content: ≤ 7,5 %
	Carbohydrates: ≤ 35,0 %

	Authorised Novel Food	Specifications
		Total Dietary Fibre: ≥ 15 %
		Crude protein (N × 6,25): \geq 22 %
		Fat: ≤ 4,5 %
		Heavy metals:
		Lead: $\leq 0.5 \text{ mg/kg}$
		Cadmium: ≤ 0,5 mg/kg
		Mercury: $\leq 0.1 \text{ mg/kg}$
		Arsenic: ≤ 0,3 mg/kg
		Mycotoxins:
		Aflatoxin B1: ≤ 0,10 µg/kg
		Aflatoxins (sum of B1 + B2 + G1 + G2): $< 4 \mu g/kg$
		Microbiological criteria:
		Total plate count: ≤ 5 000 CFU (17) Total yeast and mould count: < 100 CFU/g E. 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
▼ <u>M98</u>		
	Vitamin D ₂ mushroom powder	Description/Definition:
		The novel food is produced by controlled exposure of sliced/diced <i>Agaricus bisporus</i> mushrooms to UV irradiation followed by dehydration and grinding into a powder.
		UV radiation: A process of radiation in ultraviolet light within a range of wavelength similar to those UV-treated novel foods authorised under Regulation (EU) 2015/2283.
		Characteristics/composition:
		Vitamin D ₂ content: 125–375 μg/g
		Moisture: ≤ 7 %

▼<u>M98</u>

▼<u>M9</u>

Authorised Novel Food	Specifications
	Ash: ≤ 13,5 %
	Water activity: < 0,5
	Fat: $\leq 4.5 \%$
	Total carbohydrates: ≤ 60 %
	Protein: ≤ 40 %
	Heavy metals:
	Lead: $\leq 0.5 \text{ mg/kg}$
	Cadmium: ≤ 0,5 mg/kg
	Mercury: $\leq 0.1 \text{ mg/kg}$
	Arsenic: ≤ 0.3 mg/kg
	Mycotoxins:
	Aflatoxin B1: $\leq 2 \mu g/kg$
	Aflatoxins (sum of B1 + B2 + G1 + G2): $< 4 \mu g/kg$
	Microbiological criteria:
	Total aerobic microbial count: ≤ 5 000 CFU/g
	Total yeast and mould count: < 100 CFU/g
	Coliforms: < 100 MPN/g
	Salmonella spp.: Absence in 25 g
	Staphylococcus aureus: Absence in 10 g
	Escherichia coli: Absence in 10 g
	Listeria monocytogenes: Absence in 25 g
	CFU: colony forming units. MPN: most probable number.
itamin K2 (menaquinone)	This novel food is produced by a synthetic or microbiological process.
ramm 142 (menaqumone)	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 homologue 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-(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-octacosaheptaenyl)-3-methyl-1,4-naphtalenedione
	CAS Number: 2124-57-4
	Molecular formula: C ₄₆ H ₆₄ O ₂

Authorised Novel Food	Specifications
	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 <i>Triticum aestivum</i> 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

▼ M9

Authorised Novel Food	Specifications
	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
8	

▼M78

Wolffia arrhiza and/or Wolffia globosa fresh plants (Traditional food from a third country)

Description/Definition:

The traditional food consists of fresh plants of Wolffia arrhiza (L.) Horkel ex Wimm. and/or of Wolffia globosa (Roxb.) Hartog & Plas (family: Araceae).

Microbiological criteria:

Total plate count: $< 10^3 \text{ CFU/g}$

Total yeast and mould count: < 100 CFU/g Total Enterobacteriaceae: < 100 CFU/g Escherichia coli: < 100 CFU/g

Salmonella: Absence in 25 g

Listeria monocytogenes: Absence in 25 g Staphylococcus aureus: Absence/10 g

Heavy metals:

Lead: < 0.3 mg/kg

Arsenic (inorganic): < 0,10 mg/kg

Cadmium: < 0,2 mg/kg

Chromium: < 1 mg/kg

Mercury: < 0,10 mg/kg

Trace elements:

Copper: < 0,8 mg/kg

Molybdenum: < 0,3 mg/kg

Zinc: < 5 mg/kg

▼<u>M19</u> ▼<u>M20</u>

Xylo-oligosaccharides

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	-
Dry material (%)	-	-	70-75
Protein (g/100 g)		< 0,2	•
Ash (%)		≤ 0,3	
pH		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

Authorised Novel Food	Specifications			
	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 ^c /25 g)	Negative		
	E. coli (MPN ^d /100 g)	Negative		
	Yeast (CFU/g)	< 10		
	Mould (CFU/g)	< 10		
	 a Other carbohydrates include monosaccharid b Maltodextrin content is calculated according DP: Degree of polymerization c CFU: Colony Forming Units d MPN: Most Probable Number 	·-		

Authorised Novel Food

		•	
▼ <u>M31</u>			
	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: ≤ 12 %	
		Water content: ≤ 5 %	
		Dry matter content: ≥ 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 (10): < 10 CFU/g (i.e. limit of detection)	
		Coliforms: $\leq 10 \text{ CFU/g}$	
		Salmonella spp.: Absence in 25 g	
▼ <u>M9</u>			
	Yeast beta-glucans	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)-\(\beta\)-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 <i>Saccharomyces cerevisiae</i> 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.	

Specifications

ı	7.0
ı	20
ı	
ı	7
ı	17R24
ı	\sim
ı	4
ı	7
ı	0
ı	-
ı	E
ı	_
ı	0
ı	01.
ı	0
ı)5.
ı	· .
ı	.05.20
ı	.2024
J	4
J	1
J	
ı	ے ا
ı	7
ı	6
ı	۔َ ا
J	001
J] (
J	
	3
ı	0

Authorised Novel Food Specifications 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) beta-glucans: 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

Authorised Novel Food	Specifications
	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:
	► <u>M32</u> Lead: < 0,2 mg/kg
	Arsenic: < 0,2 mg/kg
	Mercury: $< 0.1 \text{ mg/kg}$
	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: C ₄₀ H ₅₆ O ₂
	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

Authorised Novel Food	Specifications
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
	Molecular formula: (C ₅ H ₆ NO ₃) ₂ Zn
	Relative anhydrous molecular mass: 321,4
	Appearance: White to slightly white powder
	Purity:
	Zinc L-pidolate (purity): ≥ 98 %
	pH (10 % aqueous sol.): 5,0-6,0
	Specific rotation: 19,6°- 22,8°
	Water: ≤ 10,0 %
	Glutamic acid: < 2,0 %
	Heavy metals:
	Lead: ≤ 3.0 ppm
	Arsenic: ≤ 2,0 ppm
	Cadmium: ≤ 1,0 ppm
	Mercury: ≤ 0,1 ppm

02017R247
0 — EN —
- 01.05.2024
4 - 046.001
-306

Microbiological criteria:
Total viable mesophilic count: ≤ 1 000 CFU/g
Yeasts and moulds: ≤ 100 CFU/g
Pathogen: Absence

Specifications

- (*) Cornell RM and Schwertmann U, 2003. The Iron Oxides: Structure, Properties, Reactions, Occurrences and Uses. 2nd Edition. Wiley. https://doi.org/10.1002/3527602097
- (1) 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).
- (2) 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).
- ▶ M15 (3) OSC-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.
- (4) 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, Pavne MJ. Reed J. J Sci Food Agric. 2010 Jul;90(9):1473-8.
- (5) The different values for these three parameters are due to the different methods used.
- (6) GAE: Gallic Acid Equivalents.

Authorised Novel Food

- (7) CFU: Colony Forming Units. ◀
- M30 (8) HPLC/RI: High-performance liquid chromatography coupled with refractive index detection.
- (9) CFU: Colony-forming unit. ◀
- (10) To 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.
- (11) 2'-Fucosyl-galactose, Glucose, Galactose, Mannitol, Sorbitol, Galactitol, Trihexose, Allo-lactose and other structurally related carbohydrates.
- ► M51 (12) Converted from International Units (IU) using the conversion factor of 0.025 ug = 1 IU.
- (13) Expressed as selenium.
- (14) Applicable at all stages after the heat-treatment step to guarantee the absence of viable Yarrowia lipolytica cells and to be first 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 novel food.
- (15) Digestible carbohydrates = 100 (crude protein + fat + dietary fibre + ash + moisture).
- (16) Miraculin is part of the total protein content.
- (17) Regulation (EC) No 396/2005 of the European Parliament and of the Council of 23 February 2005 on maximum residue levels of pesticides in or on food and feed of plant and animal origin and amending Council Directive 91/414/EEC (OJ L 70, 16.3.2005, p. 1).
- (18) Dietary fibre may not include chitin due to different analytical methods.
- (19) Upper bound sum of polychlorinated dibenzo-para-dioxins (PCDDs)-polychlorinated dibenzofurans (PCDFs) and dioxin-like polychlorinated biphenyls (PCBs) expressed as World Health Organization toxic equivalent (using WHO-TEFs of 2005)).
- (20) Number-based (by Transmission Electron Microscopy (TEM)).
- (21) Volume-based (hydrodynamic diameter by Dynamic Light Scattering (DLS)); CFU: Colony Forming Units.
- (22) Chitin calculated as the difference between the Acid Detergent Fibre fraction and the Acid Detergent Lignin fraction (ADF-ADL), as described by Hahn et al. (2018).
- (23) Upper bound sum of polychlorinated dibenzo-para-dioxins (PCDDs)-polychlorinated dibenzofurans (PCDFs) and dioxin-like polychlorinated biphenyls (PCBs) expressed as World Health Organization Toxic Equivalent Factors (using WHO-TEFs of 2005)).
 - CFU: Colony Forming Units.
- (24) Sum of other carbohydrates = 100 (% (w/w) of dry matter) quantified carbohydrates (% (w/w) of dry matter) Ash (% (w/w) of dry matter).
- (25) CFU: Colony Forming Units.

- (26) EU: Endotoxin Units.
- (27) Chitin calculated as Acid Detergent Fibre.
- (28) Sum of other carbohydrates = 100 (% (w/w) of dry matter) 6'-Sialyllactose sodium salt (% (w/w) of dry matter) quantified carbohydrates (% (w/w) of dry matter) Ash (% (w/w) of dry matter); CFU: Colony Forming Units; EU: Endotoxin Units.
- (29) Other carbohydrates (g/100g) = 100 (Dry residue) Ash Protein (Nitrogen × 6,25) Total fat Succinic acid L-malic acid Dietary fibre
- (30) Expressed as total dietary fibre.
- (31) 9b,10a-Cholesta-5,7-diene-3b,25-diol (25(OH)).
- (32) Cholesta-5,7-diene-3b,25-diol.
- (33) (6E)-9,10-Secocholesta-5(10),6,8-triene-3b,25-diol (iso-25(OH)).
- (34) (5E,7E)-9,10-Secocholesta-5,7,10(19)-triene-3b,25-diol.