Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Volume 3

Schedule 1—Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Note: See sections 5 and 6.

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
2153	FABIANA IMBRICATA	A, H			
2154	FAGOPYRUM ESCULENTUM	A, H			
2155	FAGUS GRANDIFOLIA	A, H			
2156	FAGUS SYLVATICA	A, H			
2157	FALLOPIA MULTIFLORA	A, H	When for oral use, the medicine requires the following warning statement on the medicine label: - (FALLMUL) 'Warning: Fallopia multiflora may harm the liver in some people. Use under the supervision of a healthcare professional.'		
2158	FARNESOL	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance. If used in a flavour the total		
			flavour concentration in a medicine must be no more than 5%.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.		
2159	FARNESYL ACETATE	Е	Permitted for use only in combination with other permitted ingredients as part of a flavour or fragrance proprietary excipient		

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			formulation.
			When used in a flavour, the total flavour proprietary excipient formulation in a medicine must be no more than 5%.
			When used in a fragrance, the total fragrance proprietary excipient formulation in a medicine must be no more than 1%.
2160	FAST GREEN FCF	Е	Permitted for use only as a colour in medicines limited to topical and oral routes of administration.
2161	FENCHONE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2162	FENCHYL ACETATE	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.

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Permissible ingredients and requirements			
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2163	FENCHYL ALCOHOL	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2164	FENNEL BITTER SEED DRY	A, E, H	When used in oral medicines, the following warning statements are required on the label:
			- (CHILD3) 'Use in children under 12 years is not recommended'
			- (PREGNT2) 'Do not use if pregnant or likely to become pregnant (or words to that effect)'
			- (BREASF) 'Do not use while breastfeeding.'
2165	FENNEL LEAF	E	
2166	FENNEL OIL	A, E, H	Methyl chavicol is a mandatory component of fennel oil.
			When the concentration of methyl chavicol in the medicine is more than 5%, the nominal capacity of the container must be no more than 25mL, a restricted flow insert must be fitted on the container,

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
	mg.conc.n. mm.c	T dispose	statement is required on the label:
			- (CHILD) 'Keep out of reach of children (or words to that effect).'
			The maximum daily dose must provide no more than 150 mg of fennel oil.
			When used in oral medicines, the following warning statements are required on the label:
			- (CHILD3) 'Use in children under 12 years is not recommended.'
			- (PREGNT2) 'Do not use if pregnant or likely to become pregnant (or words to that effect).'
			- (BREASF) 'Do not use while breastfeeding.'
2167	FENNEL SWEET SEED DRY	A, E, H	When used in oral medicines, the following warning statements are required on the label:
			- (CHILD3) 'Use in children under 12 years is not recommended'
			- (PREGNT2) 'Do not use if pregnant or likely to become pregnant (or words to that effect)'
			- (BREASF) 'Do not use while breastfeeding.'
2168	FENUGREEK	E	Permitted for use only in

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Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2169	FENUGREEK OIL	E	Fenugreek oil is permitted for use only in combination with other permitted ingredients as a flavour. If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2170	FERRIC AMMONIUM CITRATE	A, E, H	When for internal use, iron is a mandatory component of ferric ammonium citrate.
			When for internal use, the medicine must contain a daily dose of no more than 24 mg of iron.
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide when used as an excipient), the primary pack must contain no more than 750 mg of iron.
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).
			Divided preparations with a dose of more than 5 mg of elemental iron per dosage unit and more than 250 mg of

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritiona support that do not make specific iron-deficiency related claims, the following warning statement is required on the label: - (IRONDEF) 'Not for the treatment of iron deficiency
			conditions' (or words to that effect).
2171	FERRIC CHLORIDE	A, E, H	When for internal use, iron is a mandatory component of ferric chloride.
			When for internal use, the medicine must contain a daily dose of no more than 24 mg of iron.
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide when used as an excipient), the primary pack must contain no more than 750 mg of iron.

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Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).
			Divided preparations with a dose of more than 5 mg of elemental iron per dosage unit and more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritions support that do not make specific iron-deficiency related claims, the medicine requires the following statement on the medicine label:
			- (IRONDEF) 'Not for the treatment of iron deficiency conditions' (or words to that effect).
2172	FERRIC CHLORIDE HEXAHYDRATE	A, E, H	When for internal use, iron is a mandatory component of ferrichloride hexahydrate.
			When for internal use, the medicine must contain a daily dose of no more than 24 mg o

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			iron. If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide wher used as an excipient), the primary pack must contain no more than 750 mg of iron.
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).
			Divided preparations with a dose of more than 5 mg of elemental iron per dosage unit and more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritional support that do not make specific iron-deficiency related claims, the medicine requires the following statement on the medicine label:
			- (IRONDEF) 'Not for the treatment of iron deficiency

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Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			conditions' (or words to that effect).
2173	FERRIC GLYCEROPHOSPHATE	A, E, H	When for internal use, iron is a mandatory component of ferrioglycerophosphate.
			When for internal use, the medicine must contain a daily dose of no more than 24 mg of iron.
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide whe used as an excipient), the primary pack must contain no more than 750 mg of iron.
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).
			Divided preparations with a dose of more than 5 mg of elemental iron per dosage unit and more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products

Permissible in	Permissible ingredients and requirements			
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
			indicated for general nutritional support that do not make specific iron-deficiency related claims, the medicine requires the following statement on the medicine label: - (IRONDEF) 'Not for the	
			treatment of iron deficiency conditions' (or words to that effect).	
2174	FERRIC OXIDE	E		
2175	FERRIC PHOSPHATE	Н	Only for use as an active homoeopathic ingredient.	
2176	FERRIC PYROPHOSPHATE	A, H	When for internal use, iron is a mandatory component of ferric pyrophosphate.	
			When for internal use, the medicine must contain a daily dose of no more than 24 mg of iron.	
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide when used as an excipient), the primary pack must contain no more than 750 mg of iron.	
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).	
			Divided preparations with a dose of more than 5 mg of	

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Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			elemental iron per dosage unit and more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritional support that do not make specific iron-deficiency related claims, the medicine requires the following statement on the medicine label:
			- (IRONDEF) 'Not for the treatment of iron deficiency conditions' (or words to that effect).
2177	FERROSOFERRIC OXIDE	E	When used in undivided preparations for internal use and the concentration of iron oxide in the medicine is more than 1%, it is considered part of the total iron content. When used in divided preparations for internal use, the concentration in the medicine must be no more than 10 mg per dosage unit.
2178	FERROSOFERRIC PHOSPHATE	Н	Only for use as an active homoeopathic ingredient.

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2179	FERROUS FUMARATE	A, H	When for internal use, iron is a mandatory component of ferrous fumarate.
			When used as an active ingredient, the medicine must contain a daily dose of no more than 24 mg of iron.
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide when used as an excipient), the primary pack must contain no more than 750 mg of iron.
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).
			Divided preparations with a dose of more than 5mg of elemental iron per dosage unit and more than 250 milligrams of elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 milligrams of elemental iron ir the total contents of the container are required to have child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritions

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Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
			support that do not make specific iron-deficiency related claims, the medicine requires the following statement on the medicine label:	
			- (IRONDEF) 'Not for the treatment of iron deficiency conditions' (or words to that effect).	
2180	FERROUS GLUCONATE	A, E, H	When for internal use, iron is a mandatory component of ferrous gluconate.	
			When for internal use, the medicine must contain a daily dose of no more than 24 mg or iron.	
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide whe used as an excipient), the primary pack must contain no more than 750 mg of iron.	
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).	
			Divided preparations with a dose of more than 5 mg of elemental iron per dosage unit and more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.	
			Undivided preparations containing more than 250 mg	

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
пеш	ingredient name	1 ur pose	of elemental iron in the total contents of the container are required to have a child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritional support that do not make specific iron-deficiency related claims, the medicine requires the following statement on the medicine label: - (IRONDEF) 'Not for the treatment of iron deficiency conditions' (or words to that effect).
2181	FERROUS GLUCONATE DIHYDRATE	А, Е, Н	When for internal use, iron is a mandatory component of ferrous gluconate dihydrate. When for internal use, the medicine must contain a daily dose of no more than 24 mg of iron.
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide when used as an excipient), the primary pack must contain no more than 750 mg of iron.
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%). Divided preparations with a dose of more than 5 mg of

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Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			elemental iron per dosage unit and more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritiona support that do not make specific iron-deficiency related claims, the medicine requires the following statement on the medicine label:
			- (IRONDEF) 'Not for the treatment of iron deficiency conditions' (or words to that effect).
2182	FERROUS IODIDE	Н	Only for use as an active homoeopathic ingredient.
2183	FERROUS LACTATE TRIHYDRATE	A, E, H	When for internal use, iron is a mandatory component of ferrous lactate trihydrate.
			When used as an active ingredient, the medicine must contain a daily dose of no more than 24 mg of iron. If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			used as an excipient), the primary pack must contain no more than 750 mg of iron.
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).
			Divided preparations with a dose of more than 5mg of elemental iron per dosage unit and more than 250 milligrams of elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 milligrams of elemental iron in the total contents of the container are required to have child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritions support that do not make specific iron-deficiency related claims, the medicine requires the following statement on the medicine label:
			- (IRONDEF) 'Not for the treatment of iron deficiency conditions' (or words to that effect).
2184	FERROUS PHOSPHATE	A, E, H	When for internal use, iron is mandatory component of

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Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
	OCTAHYDRATE		ferrous phosphate octahydrate. When used as an active ingredient, the medicine must contain a daily dose of no mor than 24 mg of iron.
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide when used as an excipient), the primary pack must contain no more than 750 mg of iron.
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).
			Divided preparations with a dose of more than 5mg of elemental iron per dosage unit and more than 250 milligrams of elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 milligrams of elemental iron in the total contents of the container are required to have child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritions support that do not make specific iron-deficiency related claims, the medicine requires the following statement on the medicine label:

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
пеш	ingi culcut name	1 ur pose	- (IRONDEF) 'Not for the treatment of iron deficiency conditions' (or words to that effect).
2185	FERROUS PICRATE	Н	Only for use as an active homoeopathic ingredient.
2186	FERROUS SULFATE	A, E, H	When used as an active ingredient, the medicine must contain a daily dose of no more than 24 mg of iron.
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide wher used as an excipient), the primary pack must contain no more than 750 mg of iron. In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).
			Divided preparations with a dose of more than 5mg of elemental iron per dosage unit and more than 250 milligrams of elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 milligrams of elemental iron ir the total contents of the container are required to have

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Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			child resistant closure. When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritional support that do not make specific iron-deficiency related claims, the medicine requires the following statement on the medicine label: - (IRONDEF) 'Not for the treatment of iron deficiency conditions' (or words to that effect).
2187	FERROUS SULFATE HEPTAHYDRATE	A, E, H	When for internal use, iron is a mandatory component of ferrous sulfate heptahydrate. When for internal use, the medicine must contain a daily dose of no more than 24 mg of iron. If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide when used as an excipient), the primary pack must contain no more than 750 mg of iron. In undivided preparations, the primary pack must contain no
			more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%). Divided preparations with a dose of more than 5 mg of elemental iron per dosage unit and more than 250 mg of elemental iron in the total

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritional support that do not make specific iron-deficiency related claims, the medicine requires the following statement on the medicine label: - (IRONDEF) 'Not for the treatment of iron deficiency conditions' (or words to that effect).
2188	FERULA ASSA-FOETIDA	A, E, H	
2189	FERULA FOETIDA	A, E, H	
2190	FERULA GALBANIFLUA	A, E, H	
2191	FERULA RUBRICAULIS	A, E, H	
2192	FERULA SUMBUL	A, H	
2193	FERULIC ACID	Е	Only for use in topical medicines for dermal application.
2194	FESTUCA ELATIOR	A, H	
2195	FEVERFEW HERB DRY	A, H	

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Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
2196	FEVERFEW HERB POWDER	A, H			
2197	FICUS CARICA	A, E, H			
2198	FICUS PUMILA	A, H			
2199	FIG	E			
2200	FIG DRY	A, H			
2201	FILIPENDULA ULMARIA	A, H	Methyl salicylate is a mandatory component of Filipendula ulmaria. Not to be included in medicines for use in the eye or on damaged skin.		
			When used internally, the concentration of methyl salicylate in the medicine must not be more than 0.001%.		
			When the concentration of methyl salicylate in a liquid preparation is more than 5% and the dosage form is other than spray, the medicine requires child resistant packaging.		
			When the concentration of methyl salicylate in a liquid preparation is more than 5% and the dosage form is spray, the medicine does not require child resistant packaging if:		
			 the delivery device is engage into the container in such a way that prevents it from being readily removed; 		
			 direct suction through the delivery device results in delivery of no more than one dosage unit; and 		
			- actuation of the spray device is ergonomically difficult for		

Permissible in	Permissible ingredients and requirements			
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
			young children to accomplish. The following warning statement is required on the medicine label:	
			- (METSAL) 'Contains methyl salicylate' (or words to that effect).	
			When for use in topical medicines for dermal application:	
			i) the concentration of methyl salicylate in the medicine must not be more than 25%;	
			ii) the following warning statements are required on the medicine label:	
			- (PREGNT2) 'Do not use if pregnant or likely to become pregnant' (or words to that effect);	
			- (CHILD4) 'Do not use [this product/insert name of product in children 6 years of age or less';	
			- (SENS) 'Application to skin may increase sensitivity to sunlight.' (or words to that effect);	
			 - (AVOID) 'Avoid prolonged exposure in the sun' (or words to that effect); 	
			iii) if the concentration of methyl salicylate in the medicine is greater than 1%, the following warning statement is required on the medicine label:	

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Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			develops, discontinue use'.		
2202	FIR BALSAM ABSOLUTE	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.		
2203	FIR NEEDLE OIL CANADIAN	A, E			
2204	FIR NEEDLE OIL SIBERIAN	A, E			
2205	FIRMIANA SIMPLEX	A, E, H			
2206	FISH OIL - RICH IN OMEGA-3 ACIDS	A	Only for use in oral medicines.		
2207	FLEMINGIA MACROPHYLLA	A, H			
2208	FLOUVE OIL	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.		
2209	FLUORESCEIN SODIUM	Е			
2210	FOENICULUM VULGARE	A, E, H	When used in oral medicines, the following warning statements are required on the label:		
			- (CHILD3) 'Use in children under 12 years is not		

Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
			recommended' - (PREGNT2) 'Do not use if pregnant or likely to become pregnant (or words to that effect)' - (BREASF) 'Do not use while	
			breastfeeding.' When the plant preparation is oil or distillate, methyl chavicol is a mandatory component and the maximum daily dose must provide no more than 150 mg of the plant preparation. When the plant preparation is oil or distillate and the concentration of methyl chavicol in the medicine is more than 5%, the nominal capacity of the container must be no more than 25mL, a restricted flow insert must be fitted on the container, and the following warning statement is required on the label: - (CHILD) 'Keep out of reach of children' (or words to that effect).	
2211	FOLIC ACID	A	When for internal use, the maximum recommended daily dose must be no more than 500 micrograms of folic acid. When folic acid, folinic acid, levomefolate salts and/or their derivatives are used in	

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Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			total of 500 micrograms of folic acid, folinic acid, levomefolate salts and/or their derivatives in total per daily dose.
			When used in preparations indicated for reducing the risk of having a child with spina bifida/neural tube defects:
			a) the maximum daily dose must provide 400 – 500 micrograms of folic acid; and
			b) the following statement must be included on the label: - (NEUR) 'Warning: Do not exceed the stated dose except on medical advice. If you have had a baby with a neural tube defect/spina bifida - seek specific medical advice (or
2212	FOOD ORANGE 6	E	Permitted for use only as a colour in medicines limited to topical and oral routes of administration.
2213	FOOD ORANGE 7	E	Permitted for use only as a colour in medicines limited to topical and oral routes of administration.
2214	FOOD RED 13	Е	Permitted for use only as a colour for topical use.
2215	FORMALDEHYDE/MELAMINE/T OSYLAMIDE COPOLYMER	Е	Only for use in topical medicines for dermal application.

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			The concentration in the medicine must be no more than 10%.
2216	FORMIC ACID	Н	Only for use as an active homoeopathic ingredient.
2217	FORSYTHIA SUSPENSA	A, H	
2218	FORTIFIED WINE	Е	Ethanol is a mandatory component of fortified wine.
			When the concentration of ethanol in the medicine is more than 3%, the medicine requires the following warning statement on the medicine label:
			- (ETHAN) 'Contains ethanol or contains alcohol'.
2219	FRACTIONATED COCONUT OIL	Е	
2220	FRACTIONATED PALM KERNEL OIL	A, E	When used as an active ingredient, can only be supplied as an uncompounded medicine substance packed for retail sale, and must comply with an uncompounded substance monograph of the British Pharmacopoeia, as in force or existing from time to time.
2221	FRAGARIA CHILOENSIS	A, E, H	
2222	FRAGARIA VESCA	A, E, H	
2223	FRAGARIA VIRGINIANA	A, E, H	
2224	FRAGARIA X ANANASSA	A, E, H	

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2225	FRANGULA BARK DRY	А, Н	Glucofrangulins calculated as glucofrangulin A is a mandatory component of Frangula bark dry.
			When used in oral medicines, if the maximum recommended daily dose contains more than 10 mg of hydroxyanthracene derivatives the medicine requires the following warning statements on the medicine label:
			- (CHILD3) 'Use in children under 12 years is not recommended';
			 (LAX2) 'Prolonged use may cause serious bowel problems' and
			- (LAX3) 'Do not use when abdominal pain, nausea or vomiting are present, or if you develop diarrhoea. If you are pregnant or breast feeding, seek the advice of a healthcare professional before taking this product' [or words to that effect].
			When promoted or marketed a a laxative, the medicine requires the following warning statement on the medicine label:
			- (LAX1) 'Drink plenty of water' [or words to that effect]
			When not promoted or marketed as laxative, the medicine requires the following warning statements on the medicine label:
			- (LAX5) 'This product

Permissible in Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
icii	ingreuent name	1 ur pose	contains [name of the herb(s) or the chemical component(s)]'; and - (LAX4) 'This product may have laxative effect'.
			When used in oral medicines, if the maximum recommended daily dose contains less than 10 mg of hydroxyanthracene derivatives and is promoted or marketed as laxative, the medicine requires the following warning statements on the medicine label: - (CHILD3) 'Use in children under 12 years is not recommended'; - (LAX1) 'Drink plenty of water' [or words to that effect]; and
			- (LAX2) 'Prolonged use may cause serious bowel problems'.
2226	FRANGULA BARK POWDER	А, Н	Glucofrangulins calculated as glucofrangulin A is a mandatory component of Frangula bark powder.
			When used in oral medicines, if the maximum recommended daily dose contains more than 10 mg of hydroxyanthracene derivatives the medicine requires the following warning statements on the medicine label: - (CHILD3) 'Use in children
			under 12 years is not recommended;

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			- (LAX2) 'Prolonged use may cause serious bowel problems' and
			- (LAX3) 'Do not use when abdominal pain, nausea or vomiting are present, or if you develop diarrhoea. If you are pregnant or breast feeding, seek the advice of a healthcare professional before taking this product [or words to that effect]'.
			When promoted or marketed a a laxative, the medicine requires the following warning statement on the medicine label:
			- (LAX1) 'Drink plenty of water [or words to that effect]'
			When not promoted or marketed as laxative, the medicine requires the following warning statements on the medicine label:
			- (LAX5) 'This product contains [name of the herb(s) or the chemical component(s)]'; and
			- (LAX4) 'This product may have laxative effect'.
			When used in oral medicines, if the maximum recommended daily dose contains less than 1 mg of hydroxyanthracene derivatives and is promoted or marketed as laxative, the medicine requires the following warning statements on the medicine label:
			on the medicine label: - (CHILD3) 'Use in children

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			under 12 years is not recommended'; - (LAX1) 'Drink plenty of		
			water [or words to that effect]'; and		
			- (LAX2) 'Prolonged use may cause serious bowel problems'.		
2227	FRANGULA PURSHIANA	А, Н	When for oral use, hydroxyanthracene derivatives calculated as cascaroside A is a mandatory component of Frangula purshiana.		
			When used in oral medicines, if the maximum recommended daily dose contains more than 10 mg of hydroxyanthracene derivatives the medicine requires the following warning statements on the medicine label:		
			- (CHILD3) 'Use in children under 12 years is not recommended';		
			- (LAX2) 'Prolonged use may cause serious bowel problems'; and		
			- (LAX3) 'Do not use when abdominal pain, nausea or vomiting are present, or if you develop diarrhoea. If you are pregnant or breast feeding, seek the advice of a healthcare professional before taking this product' [or words to that effect].		
			When promoted or marketed as a laxative, the medicine		

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			requires the following warning statement on the medicine label:
			- (LAX1) 'Drink plenty of water' [or words to that effect]
			When not promoted or marketed as laxative, the medicine requires the following warning statements on the medicine label:
			- (LAX5) 'This product contains [name of the herb(s) or the chemical component(s)]'; and
			 - (LAX4) 'This product may have laxative effect'.
			When used in oral medicines, if the maximum recommended daily dose contains less than 1 mg of hydroxyanthracene derivatives and is promoted or marketed as laxative, the medicine requires the following warning statements on the medicine label:
			- (CHILD3) 'Use in children under 12 years is not recommended';
			- (LAX1) 'Drink plenty of water' [or words to that effect] and
			- (LAX2) 'Prolonged use may cause serious bowel problems'
2228	FRAXINUS AMERICANA	A, H	
2229	FRAXINUS CHINENSIS SUBSP. RHYNCHOPHYLLA	A, H	
2230	FRAXINUS EXCELSIOR	A, H	The components Nuzhenide and secoiridoid glucoside GL3

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			are only available when the plant part is seed.
2231	FRAXINUS ORNUS	A, H	
2232	FRITILLARIA CIRRHOSA	A, H	
2233	FRITILLARIA THUNBERGII	A, H	
2234	FRITILLARIA VERTICILLATA	A, H	
2235	FRUCTOOLIGOSACCHARIDES	A, E	
2236	FRUCTOSE	A, E, H	
2237	FUCUS VESICULOSUS	A, E, H	Iodine is a mandatory component of Fucus vesiculosus.
			Only for external use when the concentration of available iodine in the medicine (excluding salts derivatives or iodophors) is 2.5% or less.
			Only for internal use when the medicine contains less than 300 micrograms of iodine per maximum recommended daily dose.
2238	FUMARIA OFFICINALIS	A, E, H	
2239	FUMARIC ACID	Е, Н	Only for use as an active homoeopathic or excipient ingredient.
2240	FUMITORY HERB DRY	A, H	
2241	FUMITORY HERB POWDER	A, H	
2242	FURAMINTON	Е	Permitted for use only in combination with other permitted ingredients as a

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			flavour. If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2243	FURFURAL	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2244	FURFURYL ACETATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2245	FURFURYL ALCOHOL	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2246	FURFURYL MERCAPTAN	Е	Permitted for use only in combination with other

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2247	FUSEL OIL	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2248	GALBANUM OIL	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2249	GALBANUM PHENOL	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2250	GALBANUM RESIN	E	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2251	GALBANUM RESINOID	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2252	GALEGA OFFICINALIS	A, H	
2253	GALEOPSIS SEGETUM	A, H	
2254	GALIUM APARINE	A, H	
2255	GALIUM ODORATUM	A, H	When used as an active ingredient coumarin is a mandatory component of Galium odoratum and the concentration of coumarin in the medicine must be no more than 0.001%.
2256	GALIUM PALUSTRE	A, H	
2257	GALIUM VERUM	A, H	
2258	GALL STONE	Н	Only for use as an active homoeopathic ingredient.

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2259	GALPHIMIA GLAUCA	А, Н	
2260	GAMMA-4-DIMETHYL-3- CYCLOHEXENE-1-PROPANOL	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2261	GAMMA-BUTYROLACTONE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2262	GAMMA-CYCLODEXTRIN	E	
2263	GAMMA-DECALACTONE	Е	Permitted for use only: (a) in topical medicines for dermal application; and
			(b) in oral medicines in combination with other permitted ingredients as part of a flavour proprietary excipient formulation.
			When used in a flavour, the total flavour proprietary excipient formulation in a medicine must be no more than 5%.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2264	GAMMA-DODECALACTONE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
2265	GAMMA-HEPTALACTONE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2266	GAMMA-HEXALACTONE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2267	GAMMA-IONONE	Е	Permitted for use only in combination with other

	agredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2268	GAMMA-LINOLEIC ACID	Е	Only for use in topical medicines for dermal application.
2269	GAMMA-LINOLENIC ACID	Е	
2270	GAMMA-N-METHYL IONONE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2271	GAMMA-NONALACTONE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			fragrance concentration in a medicine must be no more 1%.
2272	GAMMA-OCTALACTONE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2273	GAMMA-TERPINENE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2274	GAMMA-TOCOPHEROL	E	
2275	GAMMA-UNDECALACTONE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			medicine must be no more 1%.		
2276	GAMMA-VALEROLACTONE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
2277	GANODERMA LUCIDUM	A, E, H			
2278	GARCINIA GUMMI-GUTTA	A	Only for use in oral medicines.		
			Must be obtained from the rind of the fruit only.		
			Must not contain any directions for use for children or pregnant or lactating women.		
2279	GARCINIA QUAESITA	A, H			
2280	GARDEN BEAN	Е			
2281	GARDENIA JASMINOIDES	A, E			
2282	GARDENIA TAHITENSIS FLOWER EXTRACT	Е	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye or on damaged skin.		
			The concentration in the medicine must be no more than 0.002%		
2283	GARLIC BULB DRY	A, E, H			
2284	GARLIC BULB FRESH	A, H			

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2285	GARLIC BULB POWDER	A, E, H	
2286	GARLIC CLOVE POWDER	A, H	
2287	GARLIC OIL	A, E, H	
2288	GASTRODIA ELATA	A, H	
2289	GAULTHERIA PROCUMBENS	A, E, H	Methyl salicylate is a mandatory component of Gaultheria procumbens. Not to be included in medicines for use in the eye or on damaged skin.
			When used internally, the concentration of methyl salicylate in the medicine must not be more than 0.001%.
			When the concentration of methyl salicylate in a liquid preparation is more than 5% and the dosage form is other than spray, the medicine requires child resistant packaging.
			When the concentration of methyl salicylate in a liquid preparation is more than 5% and the dosage form is spray, the medicine does not require child resistant packaging if:
			 the delivery device is engage into the container in such a way that prevents it from being readily removed;
			 direct suction through the delivery device results in delivery of no more than one dosage unit; and
			 actuation of the spray device is ergonomically difficult for young children to accomplish.

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			The following warning statement is required on the medicine label:
			 - (METSAL) 'Contains methyl salicylate' (or words to that effect).
			When for use in topical medicines for dermal application
			i) the concentration of methyl salicylate in the medicine must not be more than 25%;
			ii) the following warning statements are required on the medicine label:
			- (PREGNT2) 'Do not use if pregnant or likely to become pregnant' (or words to that effect);
			- (CHILD4) 'Do not use [this product/insert name of product in children 6 years of age or less';
			- (SENS) 'Application to skin may increase sensitivity to sunlight.' (or words to that effect);
			- (AVOID) 'Avoid prolonged exposure in the sun' (or words to that effect);
			iii) if the concentration of methyl salicylate in the medicine is greater than 1%, the following warning statement is required on the medicine label:
			 - (IRRIT) 'If irritation develops, discontinue use'.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2290	GELATIN	A, E	
2291	GELIDIUM AMANSII	A, H	Iodine is a mandatory component of Gelidium amansii.
			Only for external use when the concentration of iodine in the medicine (excluding salts derivatives or iodophors) is 2.5% or less.
			Only for internal use when the medicine contains less than 300 micrograms of iodine per maximum recommended daily dose.
2292	GELLAN GUM	E	
2293	GELSEMIUM DRY	А, Н	The concentration of Gelsemium dry in the medicine must be no more than 1mg/Kg or 1mg/L or 0.0001%.
2294	GELSEMIUM POWDER	A, H	
2295	GELSEMIUM SEMPERVIRENS	А, Н	The concentration of equivalent dry Gelsemium sempervirens in the product must be no more than 1mg/Kg or 1mg/L or 0.0001%.
2296	GENET ABSOLUTE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance. If used in a flavour the total flavour concentration in a medicine must be no more than 5%.

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2297	GENTIAN DRY	A, H	
2298	GENTIAN POWDER	A, H	
2299	GENTIANA LUTEA	A, E, H	
2300	GENTIANA MACROPHYLLA	A, H	
2301	GENTIANA RHODANTHA	A, H	
2302	GENTIANA SCABRA	A, H	
2303	GENTIANELLA AMARELLA	A, H	
2304	GERANIAL	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2305	GERANIC ACID	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2306	GERANIOL	E	Permitted for use only:
			(a) in topical medicines for dermal application; and
			(b) in oral medicines in combination with other

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			permitted ingredients as part of a flavour proprietary excipient formulation.
			When used in a flavour, the total flavour proprietary excipient formulation in a medicine must be no more than 5%.
2307	GERANIUM	E	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2308	GERANIUM MACULATUM	A, E, H	
2309	GERANIUM OIL	A, E, H	
2310	GERANIUM OIL SAPONIFIED	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2311	GERANIUM OIL TERPENELESS	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.

	ngredients and requirements	Colonia 2	Colonia A
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2312	GERANIUM ROBERTIANUM GERANIUM ROSE OIL	A, E, H E	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2314	GERANIUM SIBIRICUM	A, E, H	
2315	GERANYL ACETATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2316	GERANYL ACETONE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2317	GERANYL BUTYRATE	Е	Permitted for use only in

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2318	GERANYL CROTONATE	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2319	GERANYL ETHYL ETHER	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2320	GERANYL FORMATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			medicine must be no more 1%.
2321	GERANYL ISOBUTYRATE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2322	GERANYL ISOVALERATE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2323	GERANYL NITRILE	E	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2324	GERANYL PROPIONATE	E	Permitted for use only in

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible ingredients and requirements Column 2 Column 2 Column 4				
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
			combination with other permitted ingredients as a flavour or a fragrance.	
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.	
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.	
2325	GERANYL TIGLATE	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.	
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.	
2326	GEUM RIVALE	A, H		
2327	GEUM URBANUM	A, H		
2328	GHATTI GUM	A, E, H		
2329	GIGARTINA MAMILLOSA	A, H	Iodine is a mandatory component of Gigartina mamillosa.	
			Only for external use when the concentration of iodine in the medicine (excluding salts derivatives or iodophors) is 2.5% or less.	
			Only for internal use when the medicine contains less than 300 micrograms of iodine per maximum recommended daily dose.	

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2330	GINGER DRY	A, E, H	
2331	GINGER OIL	A, E, H	
2332	GINGER OLEORESIN	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in the medicine must be no more than 5%.
2333	GINGER POWDER	A, E, H	
2334	GINKGO BILOBA	A, E, H	The Ginkgo biloba leaf extract used in the manufacture of this medicine must comply with the requirement of Identification Test B of the monograph Powdered Ginkgo Extract in the United States Pharmacopeia 32 - National Formulary 27 (USP32-NF27), as in force or existing from time to time. This condition does not apply to powdered or dried leaf.
2335	GLACIAL ACETIC ACID	Е, Н	The concentration in the medicine must be no more than 1.5%.
2336	GLECHOMA HEDERACEA	A, H	
2337	GLECHOMA LONGITUBA	A, H	
2338	GLEDITSIA AUSTRALIS	A, H	
2339	GLEDITSIA SINENSIS	A, H	

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2340	GLEHNIA LITTORALIS	A, H	
2341	GLORIOSA SUPERBA	А, Н	Colchicine is a mandatory component of Gloriosa superba and must be declared in the application.
			The concentration of colchicine in the product must be no more than 10 mg/kg or 10 mg/L or 0.001%.
2342	GLUCOMANNAN	Е	Only for use when the dosage form is other than tablet.
2343	GLUCONOLACTONE	E	
2344	GLUCOSAMINE HYDROCHLORIDE	A, E	When derived from seafood, the medicine requires the following warning statement on the medicine label:
			 (SFOOD) 'Derived from seafood'.
2345	GLUCOSAMINE SULFATE	A	When derived from seafood, the medicine requires the following warning statement on the medicine label: - (SFOOD) 'Derived from seafood'.
2346	GLUCOSAMINE SULFATE POTASSIUM CHLORIDE	A	Potassium chloride is a mandatory component of glucosamine sulfate potassium chloride.
			When derived from seafood, the medicine requires the following warning statement on the medicine label: - (SFOOD) 'Derived from seafood'.

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
Tecm	ingredient manie	Turpose	When for oral use, the medicine requires the following warning statement on the medicine label: - (POTAS) 'Contains [amount of potassium in milligrams] mg of potassium. If you have kidney disease or are taking heart or blood pressure medicines - consult your doctor or pharmacist before use. Keep out of reach of children.'
2347	GLUCOSAMINE SULFATE SODIUM CHLORIDE	A	When derived from seafood, the medicine requires the following warning statement on the medicine label: - (SFOOD) 'Derived from seafood'.
2348	GLUCOSE	A, E, H	When the medicine is for oral ingestion and the total amount of all sugars (monosaccharides and disaccharides such as glucose, honey, invert sugar, lactose, maltose, and sucrose) is more than 100mg in the maximum daily dose, then the medicine requires the following warning statement on the medicine label: - (SUGARS) 'Contains [insert name of sugar]' if medicine contains one sugar OR 'Contains sugars' (or words to that effect) if medicine contains two or more sugars. If one of the sugars is lactose

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			then the medicine also requires the following warning statement on the medicine label:
			- (LACT) 'Contains lactose' (o words to that effect).
2349	GLUCOSE GLUTAMATE	E	Only for use in topical medicines for dermal application.
2350	GLUCOSE MONOHYDRATE	A, E, H	When the medicine is for oral ingestion and the total amount of all sugars (monosaccharides and disaccharides such as glucose monohydrate, honey, invert sugar, lactose, maltose, and sucrose) is more than 100mg in the maximum daily dose, then the medicine requires the following warning statement on the medicine label: - (SUGARS) 'Contains [insert name of sugar]' if medicine contains one sugar OR 'Contains sugars' (or words to that effect) if medicine contains two or more sugars. If one of the sugars is lactose then the medicine also requires the following warning statement on the medicine label: - (LACT) 'Contains lactose' (or words to that effect).
2351	GLUCOSYLRUTIN	E	Only for use in topical medicines for dermal

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			application and not to be included in medicines intended for use in the eye. The concentration in the medicine must be no more than
			0.1%.
2352	GLUTAMIC ACID	A, E	Only for use in topical medicines for dermal application.
2353	GLUTAMIC ACID HYDROCHLORIDE	A, E, H	
2354	GLUTAMINE	A, E, H	
2355	GLUTARAL	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2356	GLUTATHIONE	A, E	When used as an active ingredient, glutathione can only be used in medicines with an oral route of administration and must be indicated for use in adults only and not in pregnant or lactating women.
			The medicine requires the following warning statement on the medicine label:
			 - (PREGNT) 'Not recommended for use by pregnant and lactating women'

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			(or words to that effect) - (ADULT) 'Adults only' (or words to that effect).
2357	GLUTEN-FREE WHEAT STARCH	E	
2358	GLYCERETH-26	Е	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
			The concentration in the medicine must be no more than 7%.
2359	GLYCEROL	A, E	When used as an active ingredient, it is only for use in topical medicines for dermal application.
2360	GLYCEROL ESTER OF PARTIALLY HYDROGENATED GUM ROSIN	E	Only for use when the dosage form is 'chewing gum'. Must comply with: a) the Glycerol Ester of Partially Hydrogenated Gum Rosin monograph in the Food Chemicals Codex published by the United States Pharmacopeial Convention, as in force or existing from time to time; and b) the requirements for residual solvents and catalysts in the British Pharmacopeia or the United States Pharmacopeia- National Formulary, as in force or existing from time to time.

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2361	GLYCERYL BEHENATE	E	Behenic acid is a mandatory component of glyceryl behenate.
			When for oral ingestion, the maximum recommended daily dose must not provide more than 383.5 milligrams of behenic acid.
			In medicines for topical use, the concentration of glyceryl behenate must be no more than 5%.
2362	GLYCERYL CAPRYLATE	E	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye. The concentration in the medicine must be no more than 1%.
2363	GLYCERYL DIISOSTEARATE	Е	For use in topical medicines for dermal application.
2364	GLYCERYL DILAURATE	Е	Only for use in topical medicines for dermal application.
2365	GLYCERYL DIOLEATE	Е	Only for use in topical medicines for dermal application.
2366	GLYCERYL DISTEARATE	Е	Only for use in topical medicines for dermal application.

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Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2367	GLYCERYL GLUCOSIDE	Е	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
			The concentration in the medicine must be no more than 5%.
2368	GLYCERYL ISOSTEARATE	E	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
			The concentration in the medicine must be no more than 5.5%.
2369	GLYCERYL LAURATE	E	Only for use in topical medicines for dermal application.
2370	GLYCERYL LINOLEATE	E	Only for use in topical medicines for dermal application.
2371	GLYCERYL LINOLENATE	Е	Only for use in topical medicines for dermal application.
2372	GLYCERYL MONOOLEATE	E	
2373	GLYCERYL MONOSTEARATE	Е	
2374	GLYCERYL MYRISTATE	Е	Only for use in topical medicines for dermal application.

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2375	GLYCERYL OLEATE CITRATE	E	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
			The concentration in the medicine must be no more than 4% of the formulation.
2376	GLYCERYL PALMITO- STEARATE	E	
2377	GLYCERYL POLYACRYLATE	Е	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye. The concentration in the
			medicine must be no more than 0.15%.
2378	GLYCERYL POLYMETHACRYLATE	E	Only for use in topical medicines for dermal application.
2379	GLYCERYL RICINOLEATE	E	Only for use in topical medicines for dermal application.
2380	GLYCERYL ROSINATE	E	Only for use when the dosage form is 'chewing gum'. Must comply with: a) the Glycerol Ester of Gum Rosin monograph in the Food Chemicals Codex published by the United States

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			Pharmacopeial Convention, as in force or existing from time to time; and
			b) the requirements for residual solvents and catalysts in the British Pharmacopoeia or the United States Pharmacopeia National Formulary, as in forcor existing from time to time.
2381	GLYCERYL SORBITAN OLEOSTEARATE	E	Only for use in topical medicines for dermal application.
2382	GLYCERYL STARCH	Е	Only for use in topical medicines for dermal application.
			The concentration in the medicine must be no more than 4%.
			The residual levels of epichlorohydrin are to be kept below the level of detection.
2383	GLYCERYL STEARATE CITRATE	Е	Only for use in topical medicines for dermal application.
			The concentration in the medicine must be no more than 5%.
2384	GLYCERYL TRIACETYL HYDROXYSTEARATE	Е	Only for use in topical medicines for dermal application.
			The concentration in the medicine must be no more than 6%.

	ngredients and requirements		6.1
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2385	GLYCERYL TRIACETYL RICINOLEATE	Е	Only for use in topical medicines for dermal application.
2386	GLYCERYL TRINITRATE	Н	Only for use as an active homoeopathic ingredient.
2387	GLYCERYL UNDECYLENATE	E	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye or on damaged skin. The concentration of glyceryl undecylenate in a medicine must be no more than 3%.
2388	GLYCINE	A, E	
2389	GLYCINE MAX	A, E, H	
2390	GLYCOGEN	Е	Only for use in topical medicines for dermal application.
2391	GLYCOL DISTEARATE	Е	Only for use in topical medicines for dermal application.
2392	GLYCOLIC ACID	Е	Only for use in topical medicines for dermal application.
			Sponsors should consider the impact of excipients on the sensitivity of the skin to sunlight and should ensure the finished product is safe for its

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			intended purpose.
			When present as an excipient in sunscreens, the concentration in the medicine must be no more than 5%.
			When used as an excipient ingredient in other medicines the concentration in the medicine must be no more than 20%.
			If the concentration is more than 5% but no more than 20% the pH of the medicine must be 3.5 or greater.
2393	GLYCYRRHIZA GLABRA	A, E, H	
2394	GLYCYRRHIZA SPECIES	E	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2395	GLYCYRRHIZA URALENSIS	A, E, H	
2396	GLYCYRRHIZINIC ACID	Е	
2397	GNAPHALIUM AFFINE	A, H	
2398	GNAPHALIUM POLYCEPHALUM	A, H	
2399	GNAPHALIUM ULIGINOSUM	A, H	
2400	GOAT	Н	Only for use as an active homoeopathic ingredient.
2401	GOAT MILK	Е	If the product is for oral ingestion and contains lactose, then the medicine requires the

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			following warning statement on the medicine label: - (LACT) 'Contains lactose' (o words to that effect).		
2402	COLD	ЕП			
2402	GOLD	Е, Н	Only for use as an active homoeopathic or excipient ingredient.		
2403	GOLD CHLORIDE	Н	Only for use as an active homoeopathic ingredient.		
2404	GOLDEN ROD HERB DRY	A, E, H			
2405	GOLDEN SEAL ROOT DRY	A, H			
2406	GOLDEN SEAL ROOT POWDER	A, H			
2407	GOLDEN SYRUP	E	Sucrose is a mandatory component of Golden syrup when the route of administration of the medicine is oral or sublingual.		
			When the medicine is for oral ingestion and the total amount of all sugars (monosaccharides and disaccharides such as glucose, honey, invert sugar, lactose, maltose, and sucrose) is more than 100mg in the maximum daily dose, then the medicine requires the following warning statement on the medicine label: - (SUGARS) 'Contains [insert name of sugar' if medicine		
			name of sugar]' if medicine contains one sugar OR 'Contains sugars' (or words to that effect) if medicine		

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
		-	contains two or more sugars. If one of the sugars is lactose then the medicine also requires the following warning statement on the medicine label: - (LACT) 'Contains lactose' (or medicine labels and the theory of the statement of the stateme
2408	GOMPHRENA GLOBOSA	A, H	words to that effect).
2409	GOOSEBERRY	Е	
2410	GOSSYPIUM HERBACEUM	A, E, H	
2411	GRAPE	Е	
2412	GRAPE SEED OIL	Е	
2413	GRAPE WINE RED	E	Ethanol is a mandatory component of Grape wine red. When the concentration of ethanol in the medicine is mor than 3%, the medicine requires the following warning statement on the medicine label: - (ETHAN) 'Contains ethanol' or 'contains alcohol'
2414	GRAPE WINE SHERRY	Е	Ethanol is a mandatory component of Grape wine sherry. When the concentration of ethanol in the medicine is more than 3%, the medicine requires the following warning statement on the medicine label: - (ETHAN) 'Contains ethanol' or 'contains alcohol'

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2415	GRAPE WINE WHITE	E	Ethanol is a mandatory component of Grape wine white.
			When the concentration of ethanol in the medicine is more than 3%, the medicine requires the following warning statement on the medicine label:
			- (ETHAN) 'Contains ethanol' or 'contains alcohol'
2416	GRAPEFRUIT	Е	
2417	GRAPEFRUIT OIL	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2418	GRAPEFRUIT OIL COLDPRESSED	A , E, H	
2419	GRAPEFRUIT OIL CONCENTRATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
	ingredient name	1 ui pose	specific requirements
2420	GRAPEFRUIT OIL TERPENELESS	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more tha 5%.
2421	GRAPEFRUIT OIL TERPENES AND TERPENOIDS	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
2422	GRAPHITE	Н	Only for use as an active homoeopathic ingredient.
2423	GRATIOLA LINIFOLIA	A, H	
2424	GREATER NETTLE HERB DRY	A, H	
2425	GREATER NETTLE HERB POWDER	A, H	
2426	GREATER NETTLE ROOT DRY	A, H	
2427	GREATER NETTLE ROOT POWDER	A, H	
2428	GREEN LIPPED MUSSEL	A	
2429	GREEN LIPPED MUSSEL DRIED	A	
2430	GREEN LIPPED MUSSEL OIL	A	

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2431	GREEN S	Е	Only for use as a colour in topical and oral medicines.
2432	GRIFOLA FRONDOSA	A	When the route of administration is oral or sublingual, the medicine requires the following warning statement on the medicine label: -(WARF) 'Do not take while on warfarin therapy without medical advice.'
2433	GRINDELIA CAMPORUM	А, Н	
2434	GRINDELIA ROBUSTA	A, H	
2435	GRISALVA	Е	Permitted for use only in combination with other permitted ingredients as a fragrance. If used in a fragrance the total
			fragrance concentration in a medicine must be no more than 1%.
2436	GROUND IVY HERB DRY	A, H	
2437	GROUND IVY HERB POWDER	A, H	
2438	GUAIAC WOOD OIL	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in Column 1	Column 2	Column 4	
Item	Ingredient name	Column 3 Purpose	Specific requirements
			fragrance concentration in a medicine must be no more 1%.
2439	GUAIACOL	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in the medicine must be no more than 5%.
2440	GUAIACUM OFFICINALE	A, E, H	
2441	GUAIACUM RESIN	A, E, H	
2442	GUAIACUM SANCTUM	A, H	
2443	GUAIACWOOD ACETATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance. If used in a flavour the total flavour concentration in a medicine must be no more than
			5%. If used in a fragrance the total
			fragrance concentration in a medicine must be no more than 1%.
2444	GUAIENE	Е	Permitted for use only in combination with other permitted ingredients as a
			fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2445	GUAIYL ACETATE	Е	Permitted for use only in combination with other permitted ingredients as a fragrance. If used in a fragrance the total fragrance concentration in a medicine must be no more than
2446	GUANINE	E	1%. Only for use as an excipient in
2440	GOANNE	L	topical medicines for dermal application.
2447	GUANOSINE	Е	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration must be no more than 0.01% in the medicine.
2448	GUAR GALACTOMANNAN	A	When for oral use:
			(a) the maximum daily dose must provide no more than 25 g of guar galactomannan;
			(b) the medicine requires the following dosage instructions:
			- (FIBRE) 'The dose of fibre should be increased gradually. Fluid intake should be increased with an increasing dose of fibre.' (or words to that effect)
			(c) when the dosage form is a powder preparation, the medicine requires the

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			following dosage instructions: - (DNTPOW) 'Do not take powder alone. Mix with food or fluid.' (or words to that effect).
2449	GUAR GUM	A, E, H	
2450	GUAR HYDROXYPROPYLTRIMONIUM CHLORIDE	Е	Only for use as an excipient in topical medicines for dermal application.
2451	GUAREA RUSBYI	A, H	
2452	GUAVA	Е	
2453	GURJUN BALSAM	E	Permitted for use only in combination with other permitted ingredients as a fragrance. If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2454	GYMNADENIA NIGRA	A	
2455	GYMNEMA SYLVESTRE	A, H	
2456	GYMNOCLADUS DIOICA	A, H	
2457	GYNOSTEMMA PENTAPHYLLUM	A	The herbal substance must be derived from the aerial parts of the vine only (stem, leaves, fruit).
2458	GYNURA JAPONICA	A, H	
2459	HAHNEMANN'S SOLUBLE MERCURY	Н	Only for use as an active homoeopathic ingredient.

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2460	HALIBUT-LIVER OIL	A, E	Colecalciferol and Vitamin A are mandatory components of Halibut-liver oil. When for internal use, the maximum recommended daily dose must be no more than 25 micrograms of Vitamin D.
			When for use in topical medicines, the concentration o Vitamin A in the medicine must be no more than 1%.
			When for internal use, the maximum daily dose must be no more than 3000 micrograms of Retinol Equivalents.
			When preparations for internal use in adults contain more than 33 micrograms of retinol equivalents per dosage unit in divided preparations or per gram of an undivided preparation, the medicine requires the following warning statements on the medicine label:
			- (VITA2) 'WARNING: If you are pregnant - or considering becoming pregnant - do not take Vitamin A supplements without consulting your doctor or pharmacist [or words to that effect].' NOTE: Position this warning at the beginning of the directions for use.
			- (VITA4) 'WARNING - When taken in excess of 3000 micrograms retinol equivalents - Vitamin A can cause birth defects.' NOTE: Position this

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			warning at the beginning of the directions for use.
			- (VITA3) 'The recommended daily amount of Vitamin A from all sources is 700 micrograms retinol equivalents for women and 900 micrograms retinol equivalents for men.'
2461	HAMAMELIS LEAF DRY	A, H	
2462	HAMAMELIS LEAF POWDER	A, H	
2463	HAMAMELIS VIRGINIANA	A, E, H	
2464	HAMAMELIS WATER	A, E, H	
2465	HANDROANTHUS HEPTAPHYLLUS	A, H	
2466	HANDROANTHUS IMPETIGINOSUS	A, E, H	
2467	HARD FAT	Е	
2468	HARD PARAFFIN	Е	
2469	HARICOT BEAN	Е	
2470	HARPAGOPHYTUM PROCUMBENS	A, E, H	
2471	HARUNGANA MADAGASCARIENSIS	A, H	
2472	HAZEL NUT	Е	
2473	HAZEL NUT OIL	Е	
2474	HEAVY KAOLIN	Е	
2475	HEAVY MAGNESIUM OXIDE	A, E, H	
2476	HECTORITE	Е	Only for use in topical medicines for dermal application.

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2477	HEDEOMA PULEGIOIDES	A	
2478	HEDERA HELIX	A, H	Emetine is a mandatory component of Hedera helix.
			The concentration of emetine in the medicine must be no more than 0.2%.
2479	HEDTA	Е	Only for use as an excipient in topical medicines for dermal application.
2480	HEKLA LAVA	Н	Only for use as an active homoeopathic ingredient.
2481	HELESTRALIS	E	Permitted for use only in combination with other permitted ingredients as a fragrance. If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2482	HELIANTHEMUM NUMMULARIUM	A, H	
2483	HELIANTHUS ANNUUS	A, E, H	
2484	HELIANTHUS TUBEROSUS	A, H	
2485	HELICHRYSUM ANGUSTIFOLIUM	A, E, H	
2486	HELICHRYSUM ARENARIUM	A, H	
2487	HELIOTROPYL ACETATE	E	Permitted for use only in combination with other permitted ingredients as a

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			fragrance. If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2488	HELLEBORUS NIGER	A, H	The maximum recommended daily dose must be no more than 1mg of the equivalent dry herbal material.
2489	HELLEBORUS VIRIDIS	А, Н	The maximum recommended daily dose must be no more than 1mg of the equivalent dry herbal material.
2490	HELONIAS RHIZOME DRY	A, H	
2491	HELONIAS RHIZOME POWDER	A, H	
2492	HEMIDESMUS INDICUS	A, E, H	
2493	HEPTANAL	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance. If used in a flavour the total flavour concentration in a medicine must be no more than 5%. If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2494	HEPTANAL DIMETHYL ACETAL	E	Permitted for use only in combination with other permitted ingredients as a flavour. If used in a flavour the total flavour concentration in a

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			medicine must be no more than 5%.
2495	HEPTANOIC ACID	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2496	HEPTENAL	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2497	HEPTYL ACETATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2498	HEPTYL BUTYRATE	Е	Permitted for use only in combination with other permitted ingredients as a

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

	agredients and requirements	Column 2	Column A
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			flavour. If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2499	HEPTYL UNDECYLENATE	Е	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye or on damaged skin.
			The concentration of the medicine must be no more than 25%.
2500	HERACLEUM HEMSLEYANUM	A, H	
2501	HERNIARIA GLABRA	A, H	
2502	HESPERIDIN	A, E	
2503	HEX-3-ENYL ACETATE	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2504	HEXAHYDRO-4,7- METHANOINDEN-6-YL PIVALATE	E	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.

Permissible ingredients and requirements			
Column 2	Column 3	Column 4	
Ingredient name	Purpose	Specific requirements	
HEXAMETHYLINDANOPYRAN	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance. If used in a flavour the total flavour concentration in a medicine must be no more than 5%.	
		If used in a fragrance the total fragrance concentration in a medicine must be no more 1%	
HEXAN-1-OL	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.	
		If used in a flavour the total flavour concentration in a medicine must be no more than 5%.	
		If used in a fragrance the total fragrance concentration in a medicine must be no more 1%	
HEXANE	Е	The concentration of the medicine must be no more than 0.029%.	
		When used for a route of administration other than topical, the residual solvent limit for Hexane is 2.9 mg per recommended daily dose.	
HEXANOATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance. If used in a flavour the total	
	Ingredient name HEXAMETHYLINDANOPYRAN HEXAN-1-OL HEXANE	HEXAMETHYLINDANOPYRAN E HEXAN-1-OL E HEXANE E	

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2509	HEXANOIC ACID	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2510	HEXASODIUM FYTATE	E	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye or on damaged skin.
			The concentration of Hexasodium fytate in the medicine must be no more than 1.0 %.
2511	HEXENAL	E	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2512	HEXYL 2-METHYLBUTYRATE	E	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2513	HEXYL ACETATE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2514	HEXYL BUTYRATE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2515	HEXYL CAPROATE	Е	Permitted for use only in combination with other permitted ingredients as a

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2516	HEXYL FORMATE	E	Permitted for use only in combination with other
			permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2517	HEXYL ISOBUTYRATE	 E	Permitted for use only in
-017	1121112100001111112	2	combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2518	HEXYL ISOVALERATE	E	Permitted for use only in
			combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2519	HEXYL LAURATE	E	Only for use as an excipient in topical medicines for dermal application.

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2520	HEXYL NICOTINATE	E	
2521	HEXYL PROPIONATE	Е	Only for use in medicines in combination with other permitted ingredients as a flavour proprietary excipient formulation.
			The total flavour proprietary excipient formulation in a medicine must not be more than 5%.
2522	HEXYL SALICYLATE	E	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2523	HEXYL TIGLATE	E	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2524	HEXYLDECANOL	Е	Only for use as an excipient in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			medicine must be no more than 3%.
2525	HEXYLENE GLYCOL	Е	Only for use as an excipient in topical medicines for dermal application.
2526	HEXYLRESORCINOL	A	Permitted for use only in medicated throat lozenges. The medicine of must not contain more than 2.5 mg of hexylresorcinol per lozenge. The maximum recommended daily dose of the medicine must not provide more than 30mg of hexylresorcinol. The medicine label must specify that the medicine is only to be used for 7 days (or less). The following warning statement must be included on the medicine label: - (PREGNT) 'Not recommended for use by pregnant and lactating women' (or words to that effect).
2527	HIBISCUS ESCULENTUS	A, H	
2528	HIBISCUS MUTABILIS	A, H	
2529	HIBISCUS ROSA-SINENSIS	Е	Permitted for use only in combination with other permitted ingredients as a flavour. If used in a flavour the total flavour concentration in a

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
		-	5%.
2530	HIBISCUS SABDARIFFA	A, E, H	
2531	HIERACIUM PILOSELLA	A, H	
2532	HIGH AMYLOSE MAIZE STARCH	A, E, H	
2533	HIGH CHROMIUM YEAST	A, E	Chromium is a mandatory component of high chromium yeast.
			The maximum recommended daily dose must not provide more than 50 micrograms of chromium from organic chromium sources.
			High chromium yeast is considered to be an organic form of chromium.
2534	HIGH FRUCTOSE MAIZE SYRUP	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2535	HIGH MOLYBDENUM YEAST	A, E	Molybdenum is a mandatory component of high molybdenum yeast.
			The maximum daily dose of molybdenum from high molybdenum yeast must be no more than 62.5 micrograms.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible ingredients and requirements Column 1				
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
2536	HIGH SELENIUM YEAST	A	When for oral or sublingual use, selenium is a mandatory component of high selenium yeast. Oral medicines must contain no more than 150 micrograms of selenium per maximum recommended daily dose. When for oral use, the medicine requires the following warning statement on the medicine label: - (SELE) 'This medicine contains selenium which is toxic in high doses. A daily dose of 150 micrograms for adults of selenium from dietary supplements should not be exceeded.'	
2537	HIMATANTHUS LANCIFOLIUS	A, E, H		
2538	HIPPOPHAE RHAMNOIDES	A, E, H		
2539	HIRSCHFELDIA INCANA	A, H	Allyl isothiocyanate is a mandatory component of Hirschfeldia incana when the plant part is seed. The concentration of allyl isothiocyanate from all ingredients in the product must be no more than 10 mg/kg or 10 mg/L or 0.001%.	
2540	HISTAMINE DIHYDROCHLORIDE	Н	Only for use as an active homoeopathic ingredient.	
2541	HISTIDINE	A		
2542	HISTIDINE HYDROCHLORIDE	A, E, H		

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2543	HO LEAF OIL	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2544	HO WOOD OIL	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2545	HOLCUS LANATUS	A, H	
2546	HOLY THISTLE HERB DRY	A, H	
2547	HOLY THISTLE HERB POWDER	A, H	
2548	HOMALOMENA OCCULTA	A, H	
2549	HOMOSALATE	A, E	For use as an active ingredient only in sunscreens for dermal application.
			For use as an excipient only in topical medicines for dermal application.
			Not to be included in

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			medicines intended for use in the eye.
			The concentration in the medicine must not be more than 15%.
			When used in primary sunscreen products, the following warning statements are required on the label:
			 - (AVOID) 'Avoid prolonged exposure in the sun' (or words to this effect); and
			 (SUNPRO) 'Wear protective clothing - hats and eyewear when exposed to the sun' (or words to this effect).
2550	HONEY	A, E	When the route of administration is oral, the medicine requires the following warning statement on the medicine label:
			- (BABY2) 'Not suitable for infants under the age of twelve months' (or words to that effect).
			When the medicine is for oral ingestion and the total amount of all sugars (monosaccharide and disaccharides such as glucose, honey, invert sugar, lactose, maltose, and sucrose) is more than 100mg in the maximum daily dose, then the medicine requires the following warning statement on the medicine label: (SLIGARS) 'Contains linser'
			- (SUGARS) 'Contains [insert name of sugar]' if medicine contains one sugar OR

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			'Contains sugars' (or words to that effect) if medicine contains two or more sugars.
			If one of the sugars is lactose then the medicine also requires the following warning statement on the medicine label:
			- (LACT) 'Contains lactose' (or words to that effect).
2551	HONEY BEE	Н	Only for use as an active homoeopathic ingredient.
2552	HONEY EXTRACT	Е	Not to be included in medicines intended for use in the eye.
			The concentration in the medicine must be no more than 1%.
			When the medicine is for oral ingestion and the total amount of all sugars (monosaccharides and disaccharides such as glucose, honey, invert sugar, lactose, maltose, and sucrose) is more than 100mg in the maximum daily dose, then the medicine requires the following warning statement on the medicine label:
			- (SUGARS) 'Contains [insert name of sugar]' if medicine contains one sugar OR 'Contains sugars' (or words to that effect) if medicine contains two or more sugars.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

	ngredients and requirements	Cala 2	Colonia 4
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2553	HONEY POWDER	E	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2554	HOP STROBILE DRY	A, H	
2555	HOP STROBILE POWDER	A, H	
2556	HOPS OIL	A, E, H	
2557	HORDEUM DISTICHON	A, E, H	Gluten is a mandatory component when the plant part is seed, and must be declared in the application when the route of administration is other than topical and mucosal.
2558	HORDEUM VULGARE	A, E, H	Gluten is a mandatory component when the plant part is seed, and must be declared in the application when the route of administration is other than topical and mucosal.
2559	HOREHOUND EXTRACT	Е	Permitted for use only in combination with other permitted ingredients as a flavour. If used in a flavour the total flavour concentration in a medicine must be no more than 5%.

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2560	HORSE RADISH	E, H	Volatile oil components (of Armoracia rusticana) is a mandatory component of Horse radish.
			The maximum recommended daily dose must be no more than 20 mg of volatile oil components (of Armoracia rusticana).
2561	HOTTONIA PALUSTRIS	A, H	
2562	HOUTTUYNIA CORDATA	A, H	
2563	HOVENIA DULCIS	A, H	
2564	HUMULUS LUPULUS	A, E, H	
2565	HYALURONIC ACID	Е	Only for use as an excipient in topical medicines for dermal application.
2566	HYDNOCARPUS ANTHELMINTICA	A, H	When the medicine is for other than topical use and the plant part is seed, the maximum recommended daily dose must be no more than 1 mg of the equivalent dry seed.
2567	HYDRANGEA ARBORESCENS	A, H	
2568	HYDRANGEA PANICULATA	A, H	
2569	HYDRASTIS CANADENSIS	A, E, H	
2570	HYDRATED SILICA	Е	Only for use when the route of administration is other than inhalation.
2571	HYDROCHLORIC ACID	Е	The concentration of the

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			medicine must be no more than 0.5%.
2572	HYDROCOTYLE UMBELLATA	A, H	
2573	HYDROFLUORIC ACID	Н	Only for use as an active homoeopathic ingredient.
2574	HYDROGEN CYANIDE	Н	Only for use as an active homoeopathic ingredient.
2575	HYDROGEN PEROXIDE	A, E	When used as the active ingredient, it is only for use in topical medicines for dermal application. The concentration of hydrogen peroxide in the medicine must be no more than 3%. When used as an active ingredient, can only be supplied as an uncompounded medicine substance packed for retail sale and must comply with an uncompounded substance monograph of the British Pharmacopoeia, as in force or existing from time to time.
2576	HYDROGENATED BUTYLENE/ETHYLENE/STYREN E COPOLYMER	E	Only for use in topical medicines for dermal application. The combined concentration of hydrogenated butylene/ethylene/stryene copolymer and hydrogenated ethylene/propylene/styrene copolymer in the medicine must be no more than 9%.

Item 2577	Ingredient name		
2577		Purpose	Specific requirements
	HYDROGENATED C6-14 OLEFIN POLYMERS	E	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
			The concentration in the medicine must be no more than 7%.
2578	HYDROGENATED CASTOR OIL	Е	
2579	HYDROGENATED COCO- GLYCERIDES	Е	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration in the medicine must be no more than 3%.
2580	HYDROGENATED COCONUT OIL	Е	
2581	HYDROGENATED COTTONSEED OIL	Е	
2582	HYDROGENATED DIMER DILINOLEYL/DIMETHYLCARBO NATE COPOLYMER	Е	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye. The concentration in the medicine must be no more than
2583	HYDROGENATED	E	4% in the product. The combined concentration of

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
	NE COPOLYMER		ethylene/propylene/styrene copolymer must be no more than 9%.
2584	HYDROGENATED LANOLIN	E	
2585	HYDROGENATED LECITHIN	Е	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
			The concentration in the medicine must be no more than 5%.
2586	HYDROGENATED PALM GLYCERIDES	Е	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
			The concentration in the medicine must be no more than 1.6%.
2587	HYDROGENATED PALM GLYCERIDES CITRATE	E	Only for use in topical medicines for dermal application and not to be included in medicines for use in the eye or on damaged skin.
			The concentration in the medicine must be no more than 0.01%.
2588	HYDROGENATED PALM KERNEL OIL	E	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
			The concentration in the medicine must be no more than

Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			1.2%.
2589	HYDROGENATED PALM OIL	Е	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
			The concentration in the medicine must be no more than 2%.
			Polycyclic aromatic hydrocarbons must be kept below the level of detection.
2590	HYDROGENATED POLYDECENE	Е	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
2591	HYDROGENATED POLYDEXTROSE	A	Only to be used in a medicine where Danisco Australia Pty Ltd (Client ID 54247), who applied to have the ingredient included in this Determination is the sponsor of the medicine or has given written authorisation to the sponsor of a medicine to include the ingredient in the medicine. This paragraph ceases to be a requirement for this ingredient after 2 March 2022.
			Only permitted for use in medicines: - limited to oral routes of administration; and

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			 when the maximum recommended daily dose does not provide more than 15g of hydrogenated polydextrose.
2592	HYDROGENATED POLYISOBUTENE	Е	Only for use in topical medicines for dermal application.
2593	HYDROGENATED SOYA OIL	Е	
2594	HYDROGENATED TALLOW GLYCERIDE	Е	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
			The concentration in the medicine must be no more than 3%.
2595	HYDROGENATED VEGETABLE OIL	Е	
2596	HYDROLIAC	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2597	HYDROLYSED ADANSONIA DIGITATA LEAF POLYSACCHARIDES	E	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration in the medicine must be no more than

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			0.01%
2598	HYDROLYSED ALGIN	Е	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration in the medicine must be no more than 0.02%
2599	HYDROLYSED CEREAL SOLIDS	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2600	HYDROLYSED COLLAGEN	A, E	
2601	HYDROLYSED ELASTIN	Е	Only for use in topical medicines for dermal application.
2602	HYDROLYSED GELATIN	A, E	
2603	HYDROLYSED GLYCOSAMINOGLYCANS	Е	Only for use in topical medicines for dermal application.
2604	HYDROLYSED JOJOBA ESTERS	E	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			The concentration in the medicine must be no more than 5%.
2605	HYDROLYSED KERATIN	Е	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
			The concentration in the medicine must be no more than 5%.
2606	HYDROLYSED MAIZE STARCH	E	
2607	HYDROLYSED MILK PROTEIN	Е	
2608	HYDROLYSED RICE	A, E, H	
2609	HYDROLYSED RICE PROTEIN	Е	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration in the medicine must be no more than 0.125%.
2610	HYDROLYSED SOY PROTEIN	Е	Only for use in topical medicines for dermal application not to be included in topical medicines intended for use in the eye.
			The concentration in the medicine must be no more than 0.5%.
2611	HYDROLYSED VEGETABLE PROTEIN	Е	

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2612	HYDROLYSED WHEAT PROTEIN	E	When the route of administration is other than topical or mucosal, the medicine requires the following warning statement on the medicine label: - (GLUTEN) 'Contains [insert name of ingredient]' or words to that effect.
2613	HYDROLYSED WHEAT PROTEIN/PVP CROSSPOLYMER	E	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration in the medicine must be no more than 1.2%.
2614	HYDROLYSED YEAST PROTEIN	E	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye or on damaged skin.
			The concentration in the medicine must be no more than 0.3%.
2615	HYDROQUINONE DIMETHYL ETHER	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2616	HYDROUS WOOL FAT	A, E	When used as an active ingredient, can only be supplied as an uncompounded medicine substance packed for retail sale, and must comply with an uncompounded substance monograph of the British Pharmacopoeia, as in force or existing from time to time.
2617	HYDROXOCOBALAMIN	A	
2618	HYDROXYACETOPHENONE	Е	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye or on damaged skin.
			The concentration in the medicine must be no more than 1%.
2619	HYDROXYAPATITE	A, E	
2620	HYDROXYCITRATE COMPLEX	A	Hydroxycitrate complex must contain one or more of the three salts (calcium, sodium or potassium hydroxycitrate) of hydroxycitric acid.
2621	HYDROXYCITRIC ACID	A	
2622	HYDROXYCITRONELLAL	Е	Permitted for use only in combination with other permitted ingredients as a

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2623	HYDROXYCITRONELLAL DIMETHYL ACETAL	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2624	HYDROXYCITRONELLAL- METHYLANTHRANILATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
2625	HYDROXYCITRONELLOL	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible ii	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2626	HYDROXYETHYL CETEARAMIDOPROPYLDIMONI UM CHLORIDE	E	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration in the medicine must be no more than 0.1%.
2627	HYDROXYETHYL UREA	E	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration must be no more than 1%.
2628	HYDROXYLATED LANOLIN	Е	
2629	HYDROXYLATED MILK GLYCERIDES	Е	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration must be no more than 0.1%.
2630	HYDROXYLYSINE	A, E	
2631	HYDROXYMETHYLCELLULOSE	E	
2632	HYDROXYOCTACOSANYL HYDROXYSTEARATE	Е	Only for use in topical medicines for dermal application.

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2633	HYDROXYPALMITOYL SPHINGANINE	E	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye or on damaged skin.
			The concentration must be no more than 0.1%.
2634	HYDROXYPROLINE	A, E	
2635	HYDROXYPROPYL DISTARCH PHOSPHATE	Е	Only permitted for:
			 use in topical medicines for dermal application; and
			- medicines for internal use.
			When for use in topical medicines for dermal application:
			- not to be included medicines intended for use in the eye or damaged skin; and
			- the concentration of hydroxypropyl distarch phosphate in the medicine mus be no more than 4%.
			When for internal use, the maximum recommended daily dose must not contain more than 240mg of hydroxypropyl distarch phosphate.
2636	HYDROXYPROPYL STARCH	E	
2637	HYDROXYPROPYLBETADEX	Е	Only for use in topical medicines for dermal

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			application.
2638	HYDROXYSTEARIC ACID	E	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration must be no more than 9%.
2639	HYETELLOSE	E	
2640	HYLOCEREUS LEMAIREI	Е	Permitted for use only as a colour for oral and topical use
2641	HYLOCEREUS UNDATUS	A, H	
2642	HYMETELLOSE	Е	
2643	HYOSCYAMUS LEAF DRY	A, H	Alkaloids calculated as hyoscyamine and hyoscine are mandatory components of Hyoscamus leaf dry. The concentration of alkaloids calculated as hyoscyamine in the medicine must be no more than 300 micrograms/Kg or 300 micrograms/L or 0.00003%.
			The concentration of hyoscine in the medicine must be no more than than 300 micrograms/kg or 300 micrograms/L or 0.00003%.
2644	HYOSCYAMUS LEAF POWDER	А, Н	Alkaloids calculated as hyoscyamine and hyoscine are mandatory components of Hyoscamus leaf powder. The concentration of alkaloids

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			calculated as hyoscyamine in the medicine must be no more than 300 micrograms/Kg or 300 micrograms/L or 0.00003%.
			The concentration of hyoscine in the medicine must be no more than 300 micrograms/kg or 300 micrograms/L or 0.00003%.
2645	HYOSCYAMUS NIGER	A, H	Alkaloids calculated as hyoscyamine and hyoscine are mandatory components of Hyoscyamus niger.
			The concentration of hyoscyamine in the medicine must be no more than 3 micrograms/kg or 3 micrograms/L or 0.3%.
			The concentration of hyoscine in the medicine must be no more than 300 micrograms/kg or 300 micrograms/L or 0.00003%.
2646	HYPERICUM ASCYRON	A, H	
2647	HYPERICUM JAPONICUM	A, H	
2648	HYPERICUM PERFORATUM	A, E, H	When used for oral ingestion, the medicine requires the following warning statement on the medicine label: - (STJOHN) 'St John's Wort affects the way many prescription medicines work including oral contraceptives.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible ingredients and requirements				
Column 2	Column 3	Column 4		
Ingredient name	Purpose	Specific requirements		
HADBUI USE	F			
	A, H			
ICHTHAMMOL	Н	Only for use as an active homoeopathic ingredient.		
ILEX AQUIFOLIUM	A, H			
ILEX CHINENSIS	A, H			
ILEX PARAGUARIENSIS	A, E, H	Caffeine is a mandatory component of Ilex paraguariensis. When the medicine is packaged for supply as an undivided preparation and is for internal use or oral application, the medicine mus not contain a concentration of total caffeine greater than 4%. When the medicine is packaged for supply as a divided preparation and is for internal use or oral application the medicine must not contain a concentration of total caffeine greater than 33%. The requirements specified in paragraphs (a) to (e) below apply in relation to a medicine that contains the ingredient that:		
	Column 2 Ingredient name HYPROLOSE HYPROMELLOSE HYPROMELLOSE PHTHALATE HYPTIS SUAVEOLENS HYSSOPUS OFFICINALIS IBERIS AMARA ICHTHAMMOL ILEX AQUIFOLIUM ILEX CHINENSIS	Column 2 Ingredient name Purpose HYPROLOSE HYPROMELLOSE HYPROMELLOSE PHTHALATE HYPTIS SUAVEOLENS A, H HYSSOPUS OFFICINALIS A, E, H IBERIS AMARA ICHTHAMMOL H ILEX AQUIFOLIUM A, H ILEX CHINENSIS A, H		

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			- is supplied after 2 March 2021.
			A medicine that contains the ingredient and that:
			- was listed in the Register before 2 September 2019; and
			- is supplied before 2 March 2021;
			may comply with the requirements in paragraphs (a) to (e) below.
			a) When for internal use or ora application, the maximum recommended daily dose of the medicine must provide no more than 400mg of total caffeine.
			b) When the medicine is packaged for supply as an undivided preparation and is for internal use or oral application, the medicine must not contain a concentration of total caffeine greater than 1%.
			c) When the medicine is for internal use or oral application, a maximum recommended dose of the medicine must not provide more than 100 mg of total caffeine within a 3 hour period.
			d) When the maximum recommended daily dose of the medicine provides greater than 10 mg of total caffeine and the medicine is for internal use or oral application, the following

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			required on the label:
			 - (ADULT) 'Adults only' (or words to that effect).
			- (CAFF) 'Contains [state quantity per dosage unit or per mL or per gram of product] total caffeine [per dosage unit or per mL or per gram]. A cup of instant coffee contains approximately 80mg of caffeine.'
			- (CAFFPREG) 'Caffeine intake more than 200 mg per day is not recommended during pregnancy or breastfeeding.'
			e) When the maximum recommended daily dose of the medicine provides greater than 80 mg of total caffeine and the medicines is for internal use or oral application, the following warning statements are required on the label:
			- (CAFFLMT) 'Limit the use o caffeine-containing products (including tea and coffee) when taking this product.'
			- (CAFFCYP) 'Caffeine interacts with enzyme CYP1A2 in the liver. Consult your health professional before taking with other medicines' (or words to that effect).
2659	ILEX ROTUNDA	A, H	
2660	ILEX VERTICILLATA	A, H	
2661	ILLICIUM VERUM	А, Н	When the plant preparation is oil or distillate, the nominal capacity of the container must

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			be no more than 50 millilitres. When the concentration of Illicium verum oil or distillate in the preparation is greater than 50%, a restricted flow insert must be fitted on the container and the medicine must include the following warning
			statement on the medicine label:
			- (CHILD) 'Keep out of reach of children' (or words to that effect).
2662	IMIDUREA	E	Only for use in topical medicines for dermal application.
2663	IMMORTELLE	E	Permitted for use only in combination with other permitted ingredients as a flavour. If used in a flavour the total flavour concentration in a medicine must be no more than
			5%.
2664	IMMORTELLE OIL	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1 Column 2 Column 3 Column 4				
Item	Ingredient name	Purpose	Specific requirements	
2665	IMPATIENS	E	Permitted for use only in combination with other permitted ingredients as a flavour.	
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.	
2666	IMPATIENS BALSAMINA	A, H		
2667	IMPATIENS GLANDULIFERA	A, H		
2668	IMPERATA CYLINDRICA	A, E, H		
2669	INDIGO CARMINE	Е	Permitted for use only as a colour in medicines limited to topical and oral routes of administration.	
2670	INDIGO CARMINE ALUMINIUM LAKE	Е	Permitted for use only as a colour in medicines limited to topical and oral routes of administration.	
2671	INDIGOFERA TINCTORIA	A, H		
2672	INDISAN	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.	
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.	
2673	INDOLE	Е, Н	Only for use as an active homoeopathic or excipient ingredient.	
			The maximum recommended	

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			daily dose must contain no more than 75 mg indole.
2674	INDOLENE	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2675	INDUSTRIAL METHYLATED SPIRIT	Е	
2676	INOSITOL	A, E	
2677	INULA BRITANNICA	A, H	
2678	INULA HELENIUM	A, E, H	
2679	INULA RACEMOSA	A, H	
2680	INULIN	A, E	
2681	INULIN LAURYL CARBAMATE	E	Only for use in topical medicines for dermal application and not to be included in medicines for use in the eye or on damaged skin.
			The concentration in the medicine must be no more than 1.2%.
2682	INVERT SUGAR	E	When the medicine is for oral ingestion and the total amount of all sugars (monosaccharides and disaccharides such as glucose, honey, invert sugar, lactose, maltose, and sucrose) is more than 100 mg in the maximum daily dose, then the

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			medicine requires the following warning statement on the medicine label:
			- (SUGARS) 'Contains [insert name of sugar]' if medicine contains one sugar OR 'Contains sugars' (or words to that effect) if medicine contains two or more sugars.
			If one of the sugars is lactose then the medicine also requires the following warning statement on the medicine label:
			- (LACT) 'Contains lactose' (or words to that effect).
2683	INVERT SYRUP	E	Glucose is a mandatory component of Invert syrup when the route of administration is oral or sublingual. When the medicine is for oral ingestion and the total amount of all sugars (monosaccharides and disaccharides such as glucose, honey, invert sugar, lactose, maltose, and sucrose) is more than 100 mg in the maximum daily dose, then the medicine requires the
			following warning statement on the medicine label: - (SUGARS) 'Contains [insert name of sugar]' if medicine contains one sugar OR 'Contains sugars' (or words to that effect) if medicine contains two or more sugars. If one of the sugars is lactose

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			then the medicine also requires the following warning statement on the medicine label:
			- (LACT) 'Contains lactose' (or words to that effect).
2684	IODINE	Н	Only for use as an active homoeopathic ingredient.
			Only for external use when the concentration of iodine in the medicine (excluding salts derivatives or iodophors) is 2.5% or less.
			Only for internal use when the medicine contains less than 300 micrograms of iodine per maximum recommended daily dose.
2685	IODOPROPYNYL BUTYLCARBAMATE	Е	For use as an excipient ingredient in topical medicines only.
			The concentration in aqueous medicines must be no more than 10%.
2686	IONONE	E	Permitted for use only:
			(a) in topical medicines for dermal application; and
			(b) in oral medicines in combination with other permitted ingredients as part of a flavour proprietary excipient formulation.
			When used in a flavour, the total flavour proprietary

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			excipient formulation in a medicine must be no more than 5%.
2687	IOPAMIDOL	E	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2688	IPECACUANHA DRY	А, Н	Emetine is a mandatory component of Ipecacuanha Dry.
			The concentration of emetine in the medicine must be no more than 0.2%.
2689	IPECACUANHA POWDER	A, H	Emetine is a mandatory component of Ipecacuanha Powder.
			The concentration of emetine in the medicine must be no more than 0.2%.
2690	IPECACUANHA PREPARED	А, Н	Emetine is a mandatory component of Ipecacuanha Prepared.
			The concentration of emetine in the medicine must be no more than 0.2%.
2691	IPECACUANHA ROOT LIQUID EXTRACT	A, H	Emetine is a mandatory component of Ipecacuanha roo liquid extract.

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			The concentration of emetine in the medicine must be no more than 0.2%.
2692	IPOMOEA BATATAS	A, H	
2693	IPOMOEA JALAPA	A, H	
2694	IRIDOPHYCUS FLACCIDUM	А, Н	Iodine is a mandatory component of Iridophycus flaccidum.
			Only for external use when the concentration of iodine in the medicine (excluding salts derivatives or iodophors) is more than 2.5%.
			Only for internal use when the medicine contains less than 300 micrograms of iodine per maximum recommended daily dose.
2695	IRIS DOMESTICA	А, Н	
2696	IRIS FLORENTINA	A, H	
2697	IRIS GERMANICA	A, H	
2698	IRIS PALLIDA	A, H	
2699	IRIS TENAX	Н	
2700	IRIS VERSICOLOR	A, H	
2701	IRON	A, H	Only for use in oral medicines.
			When used as an active ingredient, the medicine must contain a daily dose of no more than 24 mg of iron.
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			up to 10 mg of iron oxide wher used as an excipient), the primary pack must contain no more than 750 mg of iron.
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).
			Divided preparations with a dose of more than 5mg of elemental iron per dosage unit and more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritiona support that do not make specific iron-deficiency related claims, the following warning statement is required on the label:
			- (IRONDEF) 'Not for the treatment of iron deficiency conditions' (or words to that effect).
2702	IRON (II) BISGLYCINE SULFATE	A	Only for use in oral medicines.
	TRIHYDRATE		Iron is a mandatory component

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			of iron (II) bisglycine sulfate trihydrate. When for internal use, the		
			medicine must contain a daily dose of no more than 24 mg of iron.		
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide when used as an excipient), the primary pack must contain no more than 750 mg of iron. In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).		
			Divided preparations with a dose of more than 5 mg of elemental iron per dosage unit and more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.		
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritional support that do not make specific iron-deficiency related claims, the following warning statement is required on the label:		
			- (IRONDEF) 'Not for the treatment of iron deficiency conditions' (or words to that		

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			effect).
2703	IRON (II) GLYCINATE	A	Only for use in oral medicines. Iron is a mandatory component
			of iron (II) glycinate.
			When for internal use, the medicine must contain a daily dose of no more than 24 mg of iron.
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide when used as an excipient), the primary pack must contain no more than 750 mg of iron.
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).
			Divided preparations with a dose of more than 5 mg of elemental iron per dosage unit and more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritional support that do not make specific iron-deficiency related claims, the following warning statement is required on the label:

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			treatment of iron deficiency conditions' (or words to that effect).
2704	IRON (III) GLYCINATE	A	Only for use in oral medicines. Iron is a mandatory component
			of iron (III) glycinate. When for internal use, the medicine must contain a daily dose of no more than 24 mg of iron.
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide when used as an excipient), the primary pack must contain no more than 750 mg of iron.
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).
			Divided preparations with a dose of more than 5 mg of elemental iron per dosage unit and more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritiona support that do not make specific iron-deficiency related claims, the following warning statement is required on the label: - (IRONDEF) 'Not for the
			treatment of iron deficiency conditions' (or words to that effect).
2705	IRON AMINO ACID CHELATE	A, H	Only for use in oral medicines. When used internally, iron is a mandatory component of iron amino acid chelate.
			The concentration of iron in iron amino acid chelate must be no more than 25%.
			When for internal use, the medicine must contain a daily dose of no more than 24 mg of iron.
			If the divided dosage form contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide when used as an excipient), the primary pack must contain no more than 750 mg of iron.
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%). Divided preparations with a

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			elemental iron per dosage unit and more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritional support that do not make specific iron-deficiency related claims, the following warning statement is required on the label:
			- (IRONDEF) 'Not for the treatment of iron deficiency conditions' (or words to that effect).
2706	IRON OXIDE BLACK	E	Permitted for use only as a colour in medicines limited to topical and oral routes of administration.
			When used in undivided preparations for internal use and the concentration of iron oxide in the medicine is more than 1%, it is considered part of the total iron content. When used in divided
			preparations for internal use, the concentration in the

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			medicine must be no more than 10 mg per dosage unit.
2707	IRON OXIDE RED	Е	Permitted for use only as a colour in medicines limited to topical and oral routes of administration.
			When used in undivided preparations for internal use and the concentration of iron oxide in the medicine is more than 1%, it is considered part of the total iron content.
			When used in divided preparations for internal use, the concentration in the medicine must be no more than 10 mg per dosage unit.
2708	IRON OXIDE YELLOW	Е	Permitted for use only as a colour in medicines limited to topical and oral routes of administration.
			When used in undivided preparations for internal use and the concentration of iron oxide in the medicine is more than 1%, it is considered part of the total iron content.
			When used in divided preparations for internal use, the concentration in the medicine must be no more than 10 mg per dosage unit.
2709	IRON PHOSPHATE	A, E, H	When used internally, iron is a mandatory component of iron phosphate and must be declared.

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			When for internal use, the medicine must contain a daily dose of no more than 24 mg of iron. If the divided dosage form
			contains more than 5 mg of iron per dosage unit (excluding up to 10 mg of iron oxide when used as an excipient), the primary pack must contain no more than 750 mg of iron.
			In undivided preparations, the primary pack must contain no more than 750 mg of iron (excluding iron oxides when present as an excipient at a quantity of no more than 1%).
			Divided preparations with a dose of more than 5 mg of elemental iron per dosage unit and more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			Undivided preparations containing more than 250 mg of elemental iron in the total contents of the container are required to have a child resistant closure.
			When for internal use except for iron-containing multivitamin/mineral products indicated for general nutritiona support that do not make specific iron-deficiency related claims, the following warning statement is required on the

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			label:		
			 (IRONDEF) 'Not for the treatment of iron deficiency conditions' (or words to that effect). 		
2710	IRONE	E			
2711	IRVINGIA GABONENSIS SEED TRIGLYCERIDES	Е	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.		
			The concentration must be no more than 0.375%.		
2712	ISATIS TINCTORIA	A, H			
2713	ISOAMBRETTOLIDE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.		
2714	ISOAMYL 2-METHYLBUTYRATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2715	ISOAMYL ACETATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2716	ISOAMYL ALCOHOL	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2717	ISOAMYL BENZOATE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

	Column 2	Column 3	Column 4
Column 1 Item	Ingredient name		
2718	ISOAMYL BUTYRATE	Purpose E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
2719	ISOAMYL CAPRYLATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
2720	ISOAMYL CINNAMATE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
2721	ISOAMYL CITRONELLYL KETONE	E	Permitted for use only in combination with other

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2722	ISOAMYL FORMATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2723	ISOAMYL HEXANOATE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2724	ISOAMYL ISOBUTYRATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			medicine must be no more than 5%.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.		
2725	ISOAMYL ISOVALERATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.		
2726	ISOAMYL LAURATE	Е	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.		
			The concentration must be no more than 12%.		
2727	ISOAMYL METHOXYCINNAMATE	A	Only for use as an active ingredient in sunscreens for dermal application and not to be included in medicines intended for use in the eye.		
			The concentration in the medicine must not be more than 10%.		
			When used in primary sunscreen products, the following warning statements are required on the label:		

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
Item	ingredient name	1 ui pose	- (AVOID) 'Avoid prolonged exposure in the sun' (or words to this effect); and - (SUNPRO) 'Wear protective clothing - hats and eyewear when exposed to the sun' (or words to this effect).
2728	ISOAMYL PHENYLACETATE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance. If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2729	ISOAMYL PHENYLETHYL ETHER	E	Permitted for use only in combination with other permitted ingredients as part of a fragrance proprietary excipient formulation.
			The total fragrance proprietary excipient formulation in a medicine must be no more than 1%.
2730	ISOAMYL PROPIONATE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance. If used in a flavour the total flavour concentration in a medicine must be no more than

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
	5	•	5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2731	ISOAMYL SALICYLATE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2732	ISOBERGAMIATE	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2733	ISOBORNEOL	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2734	ISOBORNYL ACETATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2735	ISOBORNYL CYCLOHEXANOL	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2736	ISOBUTANE	E	Only for use in topical medicines for dermal application.
2737	ISOBUTYL ACETATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
2738	ISOBUTYL ALCOHOL	Е	The residual solvent limit for Isobutyl alcohol is 50mg per recommended daily dose.		
			The concentration of isobutyl alcohol must be no more than 0.5% of the formulation.		
2739	ISOBUTYL BENZOATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.		
			If used as a flavour the total flavour concentration in a medicine must be no more than 5%.		
2740	ISOBUTYL BENZYL CARBINOL	Е	Permitted for use only in combination with other permitted ingredients as a flavour.		
			If used as a flavour the total flavour concentration in a medicine must be no more tha 5%.		
2741	ISOBUTYL BUTYRATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.		
			If used in a flavour the total flavour concentration in a medicine must be no more tha 5%.		
2742	ISOBUTYL CAPROATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.		

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
2743	ISOBUTYL CINNAMATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
2744	ISOBUTYL FORMATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
2745	ISOBUTYL HYDROXYBENZOATE	E	Only for use in topical medicines for dermal application.		
			Medicines containing hydroxybenzoates require the following warning statement on the medicine label:		
			- (TOTBNZ) 'Contains hydroxybenzoates' (or words to this effect) if the medicine contains more than one hydroxybenzoate source OR 'Contains [insert the approved name of hydroxybenzoate		

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			used]' (or words to this effect) if product contains one hydroxybenzoate source.		
2746	ISOBUTYL ISOBUTYRATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%		
2747	ISOBUTYL ISOVALERATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
2748	ISOBUTYL PHENYLACETATE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%		
2749	ISOBUTYL PROPIONATE	Е	Permitted for use only in		

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			combination with other permitted ingredients as a flavour.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
2750	ISOBUTYL QUINOLINE	E	Permitted for use only in combination with other permitted ingredients as a fragrance.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.		
2751	ISOBUTYL SALICYLATE	E	Only for use in topical medicines for dermal application.		
2752	ISOBUTYLENE/ISOPRENE COPOLYMER	E	Only for oral use when the dosage form is chewing gum.		
			The concentration must be consistent with best practice for the production of gum delivery systems.		
2753	ISOBUTYRALDEHYDE	E	Permitted for use only in combination with other permitted ingredients as a flavour.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2754	ISOBUTYRIC ACID	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2755	ISOCETYL ALCOHOL	Е	Only for use in topical medicines for dermal application.
2756	ISOCETYL LINOLEOYL STEARATE	Е	Only for use in topical medicines for dermal application.
2757	ISOCETYL STEARATE	Е	Only for use in topical medicines for dermal application.
2758	ISOCETYL STEAROYL STEARATE	E	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration must be no more than 10%.
2759	ISOCYCLOCITRAL	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2760	ISODECYL ISONONANOATE	Е	Only for use in topical medicines for dermal application.
2761	ISODECYL NEOPENTANOATE	Е	Only for use in topical medicines for dermal application.
2762	ISODECYL OLEATE	Е	Only for use in topical medicines for dermal application.
2763	ISODECYL SALICYLATE	Е	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration must be no more than 2%.
2764	ISODODECANE	E	Only for use in topical medicines for dermal application.
2765	ISOEICOSANE	Е	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration must be no more than 2%.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
2766	ISOEUGENOL	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.		
2767	ISOEUGENYL ACETATE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.		
2768	ISOEUGENYL BENZYL ETHER	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.		
2769	ISOHEXADECANE	Е	Only for use in topical medicines for dermal application.		

Permissible in Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2770	ISOJASMONE	E	Permitted for use only in combination with other permitted ingredients as a flavour proprietary excipient formulation or fragrance proprietary excipient formulation. The total flavour proprietary excipient formulation in a medicine must not be more
			than 5%. The total fragrance proprietary excipient formulation in a medicine must not be more 1%.
2771	ISOLEUCINE	A, E	
2772	ISOMALT	Е	When the quantity of sugar alcohols per maximum recommended daily dose is more than 2g, the quantity of the sugar alcohols must be declared on the label and the medicine requires the following warning statement on the medicine label:
			- (SUGOLS) 'Products containing [insert name of sugar alcohol(s) may have a laxative effect or cause diarrhoea [or words to that effect]'.
2773	ISOMENTHONE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%		
2774	ISOMETHYLIONONE	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.		
2775	ISONONYL ACETATE	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.		
			If used as in a fragrance the total fragrance concentration in a medicine must be no more than 1%.		
2776	ISONONYL ISONONANOATE	E	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye or on damaged skin.		
			The concentration must be no more than 15%.		
2777	ISOPENTANE	E	For dental use only.		
			The concentration must be no more than 2%.		

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2778	ISOPENTANOIC ACID	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2779	ISOPHORONE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2780	ISOPHYTOL	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
2781	ISOPROPYL 2- METHYLBUTYRATE	Е	Permitted for use only in combination with other permitted ingredients as a

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

	ngredients and requirements		~ .
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			flavour or a fragrance. If used in a flavour the total flavour concentration in a medicine must be no more tha 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
2782	ISOPROPYL 4- HYDROXYBENZOATE	Е	Only for use in topical medicines for dermal application.
			Medicines containing hydroxybenzoates require the following warning statement on the medicine label:
			- (TOTBNZ) 'Contains hydroxybenzoates' (or words to this effect) if the medicine contains more than one hydroxybenzoate source OR 'Contains [insert the approved name of hydroxybenzoate used]' (or words to this effect) if product contains one hydroxybenzoate source.
2783	ISOPROPYL ACETATE	E	Only for use in topical medicines for dermal application.
2784	ISOPROPYL ALCOHOL	E	
2785	ISOPROPYL CAPROATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a

	ngredients and requirements	<u> </u>	C.1. 4
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	medicine must be no more than 5%.
2786	ISOPROPYL CINNAMATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
2787	ISOPROPYL ISOSTEARATE	E	Only for use in topical medicines for dermal application.
2788	ISOPROPYL LANOLATE	E	Only for use in topical medicines for dermal application.
2789	ISOPROPYL LAUROYL SARCOSINATE	Е	Only for use in topical medicines for dermal application and not to be included in topical medicines intended for use in the eye.
			The concentration must be no more than 5.6%.
2790	ISOPROPYL MYRISTATE	E	
2791	ISOPROPYL PALMITATE	Е	Only for use in topical medicines for dermal application.
2792	ISOPROPYL PPG-2 ISODECETH-7 CARBOXYLATE	Е	Only for use in topical medicines for dermal

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			application and not to be included in topical medicines intended for use in the eye.
			The concentration must be no more than 10%.
2793	ISOPROPYL STEARATE	Е	Only for use in topical medicines for dermal application.
2794	ISOPROPYL TITANIUM TRIISOSTEARATE	E	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
			The concentration must be no more than 0.2%.
2795	ISOPROPYL-3-METHYL- BUTANE THIOATE	E	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2796	ISOPULEGOL	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2797	ISORALDEINE 70	E	Permitted for use only in combination with other permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2798	ISOSTEARIC ACID	Е	Only for use in topical medicines for dermal application.
2799	ISOSTEAROYL HYDROLYSED COLLAGEN	E	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.
			The concentration must be no more than 0.3%.
2800	ISOSTEARYL ALCOHOL	E	Only for use in topical medicines for dermal application.
2801	ISOSTEARYL NEOPENTANOATE	Е	Only for use in topical medicines for dermal application.
2802	ISOSTEARYL PALMITATE	E	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye. The concentration must be no

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			more than 2%.
2803	ISOTRIDECYL ALCOHOL	Е	Permitted for use only in combination with other permitted ingredients as a fragrance. If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
2804	ISOVALERALDEHYDE	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance. If used in a flavour the total flavour concentration in a medicine must be no more than 5%. If used in a fragrance the total fragrance concentration in a
2805	ISOVALERIC ACID	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance. If used in a flavour the total flavour concentration in a
2806	ISPAGHULA HUSK DRY	А, Н	medicine must be no more than 5%. If used in a fragrance the total fragrance concentration in a medicine must be no more 1%. The requirement specified in paragraph (a) below applies to

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			ingredient that:
			- is listed in the Register on or after 2 March 2020; or
			- is supplied after 2 March 2021.
			(a) When a dose for children is stated, the following warning statement is required on the label:
			- (PSYLL1) 'Should only be used for children on medical advice' (or words to that effect).
			The requirement specified in paragraph (b) below applies in relation to a medicine that contains the ingredient that:
			- is listed in the Register before 2 March 2020; and
			- is supplied before 2 March 2021; and
			- does not have the warning statement (PSYLL1) on the label.
			(b) When a dose for children is stated, the following warning statement is required on the label:
			- (PSYLL) 'On medical advice' (or words to that effect).
2807	ISPAGHULA HUSK POWDER	А, Н	The requirement specified in paragraph (a) below applies to a medicine that contains the ingredient that:
			- is listed in the Register on or after 2 March 2020; or

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			- is supplied after 2 March 2021.
			(a) When a dose for children is stated, the following warning statement is required on the label:
			- (PSYLL1) 'Should only be used for children on medical advice' (or words to that effect).
			The requirement specified in paragraph (b) below applies in relation to a medicine that contains the ingredient that:
			- is listed in the Register before 2 March 2020; and
			- is supplied before 2 March 2021; and
			 does not have the warning statement (PSYLL1) on the label.
			(b) When a dose for children is stated, the following warning statement is required on the label:
			- (PSYLL) 'On medical advice' (or words to that effect).
2808	IVA AXILLARIS	A, H	
2809	JAMAICA DOGWOOD BARK DRY	А, Н	
2810	JAMAICA DOGWOOD BARK POWDER	A, H	
2811	JASMINE ABSOLUTE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavour the total

Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
			flavour concentration in a medicine must be no more than 5%.	
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.	
2812	JASMINE LACTONE	Е	Only for use in topical medicines for dermal application.	
2813	JASMINE OIL	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.	
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.	
2814	JASMINUM GRANDIFLORUM	E	Permitted for use only in combination with other permitted ingredients as a flavour.	
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.	
2815	JASMINUM OFFICINALE	A, E, H		
2816	JASSOLIA	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.	
			If used in a fragrance the total fragrance concentration in a	

Schedule 1 Specified permissible ingredients and requirements applying to these ingredients when contained in a medicine

Permissible ingredients and requirements					
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			medicine must be no more than 1%.		
2817	JATEORHIZA PALMATA	A, H			
2818	JATROPHA CURCAS	Н	Only for use as an active homoeopathic ingredient		
2819	JERUSALEM ARTICHOKE	Е			
2820	JOJOBA ESTERS	E	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye. The concentration in the medicine must be no more than 25%.		
2821	JUGLANS CINEREA	A, E, H			
2822	JUGLANS NIGRA	A, E, H			
2823	JUGLANS REGIA	A, H			
2824	JUNCUS EFFUSUS	A, H			
2825	JUNIPER BERRY OIL	A, E, H			
2826	JUNIPER BERRY OIL TERPENELESS	E	Permitted for use only in combination with other permitted ingredients as a fragrance. If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.		
2827	JUNIPERUS CALIFORNICA	A, H			
2828	JUNIPERUS COMMUNIS	A, E, H			
2829	JUNIPERUS MEXICANA	Е	Permitted for use only in		

Permissible ingredients and requirements					
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			combination with other permitted ingredients as a fragrance.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.		
2830	JUNIPERUS OXYCEDRUS	A, H			
2831	JUNIPERUS VIRGINIANA	A, E, H			
2832	JUSTICIA ADHATODA	A, H			