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

#### Volume 4

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

Note: See sections 5 and 6.

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2859	KADSURA COCCINEA	A, H	
2860	KAEMPFERIA GALANGA	A, H	
2861	KALMIA LATIFOLIA	A, H	Beta-arbutin is a mandatory component of Kalmia latifolia.
			When for oral use, the maximum recommended daily dose must not provide more than 500 mg of beta-arbutin.
			When for dermal application exclusively to the face:
			a) the concentration of beta- arbutin in the medicine must not be more than 7%;
			b) hydroquinone is a mandatory component; and
			c) the concentration of hydroquinone must not be more than 10 mg/kg or 10 mg/L or 0.001%.
			When for use other than oral or dermal application exclusively to the face, the concentration of beta-arbutin in the medicine must not be more than 10 mg/kg or 10 mg/L or 0.001%.
2862	KAOLIN	Е	
2863	KELP DRY	A, H	Iodine is a mandatory component of Kelp dry.
			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

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Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			maximum recommended daily dose.
2864	KELP POWDER	A, E, H	Iodine is a mandatory component of Kelp powder.
			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.
2865	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%.
2866	KEROSENE	E, H	Only for use as a homoeopathic ingredient.
			When used in liquid preparations, the concentration in the medicine must be no more than 25%.
2867	KHAYA SENEGALENSIS	A, E	The maximum daily dose of the medicine must not contain more than the equivalent of 1 g dry bark of Khaya senegalensis.
			The following warning statements are required on the medicine label:
			<ul> <li>- (PREGNT2) 'Do not use if pregnant or likely to become pregnant' (or words to that effect);</li> </ul>

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		Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4			
Item	Ingredient name	Purpose	Specific requirements			
			<ul> <li>(LONGUSE) 'Not for prolonged use. May harm liver';</li> </ul>			
			<ul> <li>- (GEN2) 'If symptoms persis' seek the advice of a healthcare professional';</li> </ul>			
			- (CHILD3) 'Use in children under 12 years is not recommended'; and			
			- (7DAYS) 'Do not use for more than 7 days'.			
2868	KIDNEY BEAN	Е				
2869	KIRSCH	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%.			
2870	KIWI FRUIT	Е				
2871	KNAUTIA ARVENSIS	A, H				
2872	KOREAN GINSENG ROOT DRY	A, H				
2873	KOREAN GINSENG ROOT POWDER	A, H				
2874	KRAMERIA IXIENA	A, H				
2875	KRAMERIA LAPPACEA	A, H				
2876	KUNZEA AMBIGUA	A	Only for use when the plant preparation is essential oil.			
			Only for use when the route of administration is topical or inhalation.			
			When the dosage form is essential oil, a restricted flow insert must be fitted on the container and the medicine requires the following warning statements on the medicine label:  - (CHILD) 'Keep out of reach			

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			of children'
			<ul> <li>- (EXTERN) 'For external use only'</li> </ul>
			<ul> <li>(UNDILU) 'Not to be applied undiluted to the skin except on the advice of a health care practitioner'.</li> </ul>
			When the dosage form is other than essential oil, the maximum concentration in topical medicines must be no more than 25% w/w and the medicine requires the following warning statements on the medicine label:
			- (CHILD) 'Keep out of reach of children'
			- (EXTERN) 'For external use only'.
2877	L-BORNEOL	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%
2878	L-BORNYL 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%.
2879	L-CARVONE	Е	Permitted for use only in combination with other permitted ingredients as a

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Permissible in	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%
2880	L-LIMONENE	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 tha 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
2881	L-LINALOOL	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 tha 5%.
2882	L-MENTHONE	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 tha 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
2883	L-MENTHYL ACETATE	Е	Permitted for use only in combination with other

Permissible in	Permissible ingredients 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%		
2884	L-ROSE OXIDE	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%		
2885	LABDANUM 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%.		
2886	LABDANUM GUM EXTRACT ETHYL ESTER	E	Permitted for use only in combination with other permitted ingredients as a fragrance.		
			If used in a fragrance and the total fragrance concentration in a medicine is no more than 1%		
2887	LABDANUM OIL	A, E, H			
2888	LABURNUM ANAGYROIDES	A, H	Sparteine is a mandatory component of Laburnum anagyroides.		

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Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
			The concentration of sparteine in the medicine must be no more than 0.001%.	
2889	LACTALBUMIN	Е		
2890	LACTIC ACID	A, E, H	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 form time to time.  Sponsors should consider the impact of excipients containing alpha hydroxy acids on the sensitivity of the skin to sunlight and should ensure the finished medicine is safe for its intended purpose.	
2891	LACTITOL	Е		
2892	LACTITOL MONOHYDRATE	Е		
2893	LACTOBACILLUS ACIDOPHILUS	A		
2894	LACTOBACILLUS AMYLOVORUS	A		
2895	LACTOBACILLUS BREVIS	A		
2896	LACTOBACILLUS CASEI	A		
2897	LACTOBACILLUS CASEI SUBSP. BIOVAR CASEI	A		
2898	LACTOBACILLUS CRISPATUS	A		
2899	LACTOBACILLUS DELBRUECKII SSP BULGARICUS	A		
2900	LACTOBACILLUS DELBRUECKII SSP LACTIS	A		
2901	LACTOBACILLUS FERMENTUM	A		
2902	LACTOBACILLUS GALLINARUM	A		
2903	LACTOBACILLUS GASSERI	A		
2904	LACTOBACILLUS HELVETICUS	A		

Permissible ingredients and requirements			
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2905	LACTOBACILLUS JOHNSONII	A	
2906	LACTOBACILLUS KEFIRANOFACIENS	A	
2907	LACTOBACILLUS KEFIRGRANUM	A	
2908	LACTOBACILLUS KEFIRI	A	
2909	LACTOBACILLUS PARACASEI	A	
2910	LACTOBACILLUS PARACASEI SUBSP. PARACASEI	A	
2911	LACTOBACILLUS PLANTARUM	A	
2912	LACTOBACILLUS REUTERI	A	
2913	LACTOBACILLUS RHAMNOSUS	A	
2914	LACTOBACILLUS SALIVARIUS SSP SALICINIUS	A	
2915	LACTOBACILLUS SALIVARIUS SSP SALIVARIUS	A	
2916	LACTOBIONIC ACID	Е	Only for use in topical medicines for dermal application.
2917	LACTOSCATONE	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%.
2918	LACTOSE	Е	
2919	LACTOSE MONOHYDRATE	Е	
2920	LACTUCA SATIVA	A, H	
2921	LACTUCA VIROSA	A, H	
2922	LACTULOSE	E	
2923	LACTULOSE SOLUTION	A	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

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Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			British Pharmacopoeia, as in force or existing form time to time.
2924	LAGENARIA VULGARIS	A, H	
2925	LAMINARIA CLOUSTONI	A, E, H	Iodine is a mandatory component of Laminaria cloustoni.
			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.
2926	LAMINARIA DIGITATA	<b>A</b> , E, H	Iodine is a mandatory component of Laminaria digitata.
			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.
2927	LAMINARIA JAPONICA	A, E, H	Iodine is a mandatory component of Laminaria japonica.
			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

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Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
			maximum recommended daily dose.		
2928	LAMIUM ALBUM	A, H			
2929	LANETH-5	Е	Only for use in topical medicines for dermal application.		
2930	LANOLIN ALCOHOL	Е	Only for use in topical medicines for dermal application.		
2931	LANOLIN OIL	Е	Only for use in topical medicines for dermal application.		
2932	LANOLIN WAX	Е	Only for use in topical medicines for dermal application.		
2933	LANTANA CAMARA	А, Н	The maximum recommended daily dose must contain no more than 1mg of the equivalent dry herbal material of Lantana camara.		
2934	LARIX ARABINOGALACTAN	A, E	The concentration of polysaccharides in the ingredient must be greater than or equal to 85%.		
			The ingredient must be derived from Larix occidentalis or Larix larcinia.		
			Only for use in oral medicines or topical medicines for derma application, and not to be included in topical products intended for use in the eye.		
			The maximum recommended daily dose of Larix arabinogalactan in oral medicines must not be more than 15 grams.		

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Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
			The concentration of Larix arabinogalactan in topical medicines for dermal application must not exceed 5.0%.	
2935	LARIX DECIDUA	А, Н		
2936	LARIX KAEMPFERI	A, H	The maximum recommended daily dose must be no more than 1 mg of the equivalent dry herbal material of Larix kaempferi.	
2937	LARREA TRIDENTATA	А, Н	The medicine requires the following warning statement on the medicine label:	
			- (CHAP) 'WARNING: Chaparral may harm the liver in some people - use only under supervision of a health care professional'.	
2938	LATHYRUS SATIVUS	А, Н	The maximum recommended daily dose must be no more than 1mg of the equivalent dry herbal material of Lathyrus sativus.	
			The medicine must not contain lathyrogenic amino acids.	
2939	LAURAMINE OXIDE	Е		
2940	LAUREL LEAF OIL	A, H		
2941	LAURETH-10	Е	Only for use in topical medicines for dermal application.	
2942	LAURETH-12	Е	Only for use in topical medicines for dermal application.	
2943	LAURETH-2	Е	Only for use in topical medicines for dermal application and not to be	

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Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			included in medicines intended for use in the eye.
			The concentration in the medicine must be no more than 0.4%.
			Residual levels of ethylene oxide (and related substances) must be kept below the level of detection.
2944	LAURETH-23	Е	Only for use in topical medicines for dermal application.
2945	LAURETH-3	Е	Only for use in topical medicines for dermal application.
2946	LAURETH-4	Е	Only for use in topical medicines for dermal application.
2947	LAURETH-7	Е	Only for use in topical medicines for dermal application.
2948	LAURETH-8	E	
2949	LAURIC ACID	A, E	When for use as an active ingredient is for use in oral medicines only and the maximum recommended daily dose must not exceed 1500 mg.
2950	LAURIL MACROGOL 400 DIMETICONE	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 5%.

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Permissible ii	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2951	LAUROMACROGOL 400	Е	Only for use in topical medicines for dermal application.
2952	LAUROYL LYSINE	Е	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 5.0%.
2953	LAURUS NOBILIS	A, E, H	When the plant preparation is oil or distillate, the nominal capacity of the container must be no more than 25 millilitres.
			When the concentration of Laurus nobilis oil or distillate in the preparation is greater than 25% and the nominal capacity of the container is less than or equal to 15 millilitres, a restricted flow insert must be fitted on the container.  When the concentration of
			Laurus nobilis oil or distillate in the preparation is greater than 25% and the nominal capacity of the container is greater than 15 millilitres, a child resistant closure must be fitted on the container.
			When the concentration of Laurus nobilis oil or distillate in the preparation is greater than 25%, the medicine must include the following warning statements on the medicine label:
			<ul> <li>(CHILD) 'Keep out of reach of children' (or words to that effect); and</li> </ul>

Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
2954	LAURYL ALDEHYDE	E	Permitted for use only in combination with other permitted ingredients as a coating solution, 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%.	
2955	LAURYL BETAINE	Е	Only for use in topical medicines for dermal application.	
2956	LAURYL GLUCOSIDE	E	Only for use as an excipient ingredient 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 12%.	
2957	LAURYL LACTATE	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 3%.	
			Sponsors should consider the impact of excipients containing alpha hydroxy acids on the sensitivity of the skin to sunlight and should ensure the finished medicine is safe for its intended purpose.	
2958	LAURYL PCA	E	Only for use in topical	

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Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements 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%.		
2959	LAURYL PEG-10 TRIS(TRIMETHYLSILOXY)SILYL ETHYL DIMETICONE	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 in the medicine must be no more that		
2960	LAURYL PEG-9 POLYDIMETHYLSILOXYETHYL DIMETICONE	E	Only for use in topical medicines for dermal application and not to be		
			included in medicines intender for use in the eye or damaged skin. The concentration in the medicine must be no more that 3.5%.		
2961	LAURYL PEG/PPG-18/18 METHICONE	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 tha 9%.		
			Residual levels of ethylene oxide (and related substances) must be kept below the level of detection.		
2962	LAURYL POLYGLUCOSE	Е	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.		

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			The concentration must not exceed 1% in leave-on medicines and 3% in wash-on/wash-off medicines.
2963	LAURYL PYRROLIDONE	Е	Only for use in topical medicines for dermal application.
2964	LAURYLDIMONIUM HYDROXYPROPYL HYDROLYSED COLLAGEN	Е	Only for use in topical medicines for dermal application.
2965	LAURYLDIMONIUM HYDROXYPROPYL HYDROLYSED SOY PROTEIN	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 0.007%.
2966	LAURYLMETICONE COPOLYOL	Е	Only for use in topical medicines for dermal application.
2967	LAVANDIN 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%.
2968	LAVANDIN OIL ABRIAL	A, E, H	
2969	LAVANDIN OIL GROSSO	E	Permitted for use only in combination with other permitted ingredients as a fragrance.

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Permissible ingredients and requirements  Column 1 Column 2 Column 2 Column 4				
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%.	
2970	LAVANDULA ANGUSTIFOLIA	A, E, H	Camphor is a mandatory component of Lavandula angustifolia.	
			In solid and semi solid preparations, the concentration of camphor must be no more than 12.5%.	
			In liquid preparations other than essential oils or distillates the concentration of camphor must be no more than 2.5%.	
2971	LAVANDULA ANGUSTIFOLIA SUBSP. ANGUSTIFOLIA	A, E, H	Camphor is a mandatory component of Lavandula angustifolia subsp. angustifolia.	
			In solid and semi solid preparations, the concentration of camphor must be no more than 12.5%.	
			In liquid preparations other than essential oils or distillates the concentration of camphor must be no more than 2.5%.	
2972	LAVANDULA X INTERMEDIA	A, E, H	Camphor is a mandatory component of Lavandula x intermedia.	
			In solid and semi solid preparations, the concentration of camphor must be no more than 12.5%.	
2973	LAVENDER OIL	A, E, H		
2974	LAWSONIA INERMIS	A, H		
2975	LEAD	Н	Only for use as an active homoeopathic ingredient.  The concentration in the	

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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 0.001%.
2976	LEAD ACETATE	Н	Only for use as an active homoeopathic ingredient.
2977	LEAF ACETAL	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%.
2978	LECITHIN	A, E	
2979	LEDEBOURIELLA SESELOIDES	A, H	
2980	LEDUM PALUSTRE	A, H	Beta-arbutin is a mandatory component of Ledum palustre.
			When for dermal application exclusively to the face:
			a) the concentration of beta- arbutin in the medicine must not be more than 7%;
			b) hydroquinone is a mandatory component; and
			c) the concentration of hydroquinone must not be more than 10 mg/kg or 10 mg/L or 0.001%.
			When for topical use other than dermal application exclusively to the face, the concentration of beta-arbutin in the medicine must not be more than 10 mg/kg or 10 mg/L or 0.001%.
			When the route of administration is other than topical, the maximum recommended daily dose must not contain more than 0.001 mg of the equivalent dry herba material of Ledum palustre.

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	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2001	I DI DI LI MULOD		
2981	LEMNA MINOR	A, H	
2982	LEMON	Е	When used internally, oxedrine is a mandatory component of lemon.
			The quantity of oxedrine in the maximum recommended daily dose must be no more than 30 milligrams.
2983	LEMON BALM LEAF DRY	A, H	
2984	LEMON BALM LEAF POWDER	A, E, H	
2985	LEMON OIL	A, E, H	When used internally, oxedrine is a mandatory component of lemon oil.
			The quantity of oxedrine in the maximum recommended daily dose must be no more than 30 milligrams.
			The warning statement (SENS) 'Application to skin may increase sensitivity to sunlight' (or words to that effect) must be included on the medicine label unless the medicine is:
			<ul><li>a) steam distilled or rectified;</li><li>or</li></ul>
			b) for internal use; or
			c) contains 0.05% or less of lemon oil; or
			d) for use in soaps or bath or shower gels that are washed of the skin.
2986	LEMON OIL DISTILLED	A, E, H	When used internally, oxedrine is a mandatory component of lemon oil distilled.
			The quantity of oxedrine in the maximum recommended daily dose must be no more than 30 milligrams.
2987	LEMON OIL TERPENELESS	A, E, H	When used internally, oxedrine

Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			is a mandatory component of lemon oil terpeneless.
			The quantity of oxedrine in the maximum recommended daily dose must be no more than 30 milligrams.
2988	LEMON 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%.
2989	LEMON PEEL DRIED	A, E, H	When used internally, oxedrine is a mandatory component of lemon peel dried.
			The quantity of oxedrine in the maximum recommended daily dose must be no more than 30 milligrams.
2990	LEMONGRASS OIL	A, E, H	
2991	LENS CULINARIS	A, H	
2992	LENTIL	E	
2993	LENTINULA EDODES	A, E, H	
2994	LEONTOPODIUM ALPINUM	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%.
2995	LEONURUS CARDIACA	A, E, H	
2996	LEONURUS SIBIRICUS	A, E, H	

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Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
2997	LEPIDIUM APETALUM	A, H	
2998	LEPIDIUM MEYENII	A	Only for use in oral medicines when the plant part is tuber and the plant preparation is dry.  The maximum recommended daily dose must be no more than 3.5g of Lepidium meyenii dried tuber (or its extract equivalent).
2999	LEPTOSPERMUM PETERSONII	E	Only for use in topical medicines for dermal application.
			The concentration in the medicine must be no more 5%.
3000	LEPTOSPERMUM SCOPARIUM OIL	A	Only for use as an active ingredient when the route of administration is topical or oral application in a mouthwash preparation.
			If the concentration is more than 25%, the nominal capacity of the container must be no more than 25mL.
			When the concentration is more than 25%, and the nominal capacity of the container less than 15mL, a restricted flow insert must be fitted on the container and requires the following warning statements on the medicine label:
			<ul> <li>(CHILD) 'Keep out of reach of children' (or word to that effect)</li> </ul>
			- (NTAKEN) 'Not to be taken'
			When the concentration is more than 25%, the nominal capacity of the container is more than 15 mL but no more than 25 mL, a child resistant closure and restricted flow

	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements  insert must be fitted on the container and requires the following warning statements on the medicine label:  - (CHILD) 'Keep out of reach of children' (or word to that effect)  - (NTAKEN) 'Not to be taken
3001	LESPEDEZA CAPITATA	A, H	
3002	LETTUCE	Е	
3003	LEUCINE	A, E	
3004	LEUZEA UNIFLORUM	A, H	
3005	LEVISTICUM OFFICINALE	A, H	
3006	LEVOCARNITINE	A	
3007	LEVOCARNITINE FUMARATE	A	
3008	LEVOCARNITINE HYDROCHLORIDE	A	
3009	LEVOCARNITINE MAGNESIUM CITRATE	A	
3010	LEVOCARNITINE TARTRATE	A	
3011	LEVOMEFOLATE CALCIUM	A	Available for medicines intended for internal use only.
			Levomefolic acid is a mandatory component of levomefolate calcium.
			The maximum recommended daily dose must not provide more than 500 micrograms of levomefolic acid from levomefolate calcium.
			When the medicine contains a combination of folic acid, folinic acid or levomefolic acid, the medicine must not provide more than a combined total of 500 micrograms of folic acid, folinic acid and levomefolic acid per maximum recommended daily dose.
3012	LEVOMEFOLATE	A	Available for medicines

**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
	GLUCOSAMINE		intended for internal use only.
			Levomefolic acid is a mandatory component of levomefolate glucosamine.
			The maximum recommended daily dose must not provide more than 500 micrograms of levomefolic acid from levomefolate glucosamine.
			When the medicine contains a combination of folic acid, folinic acid or levomefolic acid, the medicine must not provide more than a combined total of 500 micrograms of folic acid, folinic acid and levomefolic acid per maximum recommended daily dose.
3013	LEVOTHYROXINE SODIUM	Н	Only for use as an active homoeopathic ingredient.
3014	LEVULINIC ACID	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 tha 5%.
3015	LIGHT KAOLIN	E	
3016	LIGHT LIQUID PARAFFIN	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.
3017	LIGHT MAGNESIUM OXIDE	<b>A</b> , E, H	The requirements specified in

Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			paragraphs (a) to (c) below apply to a medicine that contains the ingredient that is:
			- listed in the Register on or after 1 March 2021; or
			<ul> <li>released for supply after 1</li> <li>March 2022. (a) Magnesium i</li> <li>a mandatory component of light magnesium oxide.</li> </ul>
			(b) When used in a medicine:
			(i) with an oral route of administration;
			(ii) not indicated for laxative (or related) use; and
			(iii) where the maximum recommended daily dose for:
			(A) children aged between 1 and 3 years (inclusive) provides 65 mg or more total magnesium from inorganic magnesium salts;
			(B) children aged between 4 and 8 years (inclusive) provides 110 mg or more total magnesium from inorganic magnesium salts; or
			(C) individuals aged 9 years of older provides 350 mg or more total magnesium from inorganic magnesium salts;
			the following warning statement is required on the medicine label:
			- (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).
			(c) When the route of administration is oral, the medicine must not be directed for use in infants younger than 12 months of age.

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

	ngredients and requirements	C-1 2	Column 4
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
3018	LIGUSTICUM SINENSE	A, H	
3019	LIGUSTICUM STRIATUM	A, E, H	
3020	LIGUSTRUM LUCIDUM	A, H	
3021	LILIUM BROWNII	A, H	
3022	LILIUM CANDIDUM	A, E, H	
3023	LILIUM LANCIFOLIUM	A, H	
3024	LILIUM LONGIFLORUM	A, H	
3025	LIME FRUIT	Е	
3026	LIME 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%.
3027	LIME OIL COLDPRESSED	A, E, H	The warning statement (SENS) 'Application to skin may increase sensitivity to sunlight' (or words to that effect) must be included on the medicine label unless the medicine is:
			a) for internal use; or
			b) contains 0.5% or less of lime oil coldpressed; or
			<ul> <li>c) for use in soaps or bath or shower gels that are washed of the skin.</li> </ul>
3028	LIME OIL DISTILLED	A, E, H	The warning statement (SENS) 'Application to skin may increase sensitivity to sunlight' (or words to that effect) must be included on the medicine label unless the medicine is: a) for internal use; or
			b) contains 0.5% or less of lim oil distilled; or

Column 1	ngredients and requirements  Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
Item	ingredient name	T air pose	c) for use in soaps or bath or shower gels that are washed of the skin.
3029	LIME OIL TERPENELESS	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%.
3030	LIME OIL TERPENES AND TERPENOIDS	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%.
3031	LIME TREE FLOWER DRY	A, H	
3032	LIME TREE FLOWER POWDER	A, H	
3033	LIME, ESSENCE	Е	
3034	LIMES TERPENES	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%.
3035	LIMONENE	Е	When for oral use, the quantity must be no more than 10 mg per maximum recommended daily dose.

**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
3036	LINALOOL	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%.
3037	LINALOOL OXIDE	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%.
3038	LINALYL 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%.
3039	LINALYL ACETATE	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.

**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
			When used in a flavour, the total flavour proprietary excipient formulation in a medicine must be no more tha 5%.
3040	LINALYL BENZOATE	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%.
3041	LINALYL BUTYRATE	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 tha 1%.
3042	LINALYL CINNAMATE	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 tha 1%.
3043	LINALYL 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 tha 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	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
3044	LINALYL 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%.
3045	LINALYL 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 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
3046	LINDERA STRYCHNIFOLIA	A, H	
3047	LINOLEAMIDOPROPYL PG- DIMONIUM CHLORIDE PHOSPHATE	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 0.5%.
3048	LINOLEIC ACID	E	
3049	LINOLENIC ACID	Е	
3050	LINSEED DRY	A, E, H	
3051	LINSEED OIL	A, E, H	
3052	LINSEED POWDER	A, E, H	
3053	LINUM USITATISSIMUM	A, E, H	
3054	LIPASE	A	Lipase must only be derived

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements from Rhizopus oryzae and must comply with the relevant compositional guideline.
3055	LIPPIA DULCIS	A, H	
3056	LIQUID GLUCOSE	Е	
3057	LIQUID PARAFFIN	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.
3058	LIQUIDAMBAR FORMOSANA	A, H	
3059	LIQUIDAMBAR ORIENTALIS	A, H	
3060	LIQUIDAMBAR STYRACIFLUA	A, E, H	
3061	LIQUIDAMBAR STYRACIFLUA RESIN	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%.
3062	LIQUIDAMBAR TAIWANIANA	A, H	
3063	LIQUORICE	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%.
3064	LIQUORICE DRY	A, E, H	
3065	LIQUORICE LIQUID EXTRACT	A, E, H	
3066	LIQUORICE POWDER	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
3067	LITCHI CHINENSIS	A, H	
3068	LITHIUM CARBONATE	Н	Only for use as an active homoeopathic ingredient.
3069	LITHOSPERMUM OFFICINALE	A, H	The maximum recommended daily dose must be no more than 1 mg of the equivalent dry herbal material of Lithospermum officinale.
3070	LITSEA CUBEBA	A, E, H	
3071	LITSEA CUBEBA 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%.
3072	LOBARIA PULMONARIA	A, H	
3073	LOBELIA DRY	A, H	The concentration in the medicine must be no more than 0.001% or 10mg/kg or 10ml/L or 10 ppm unless the medicine is administered by inhalation.
3074	LOBELIA INFLATA	A, H	The concentration in the medicine must be no more than 0.001% or 10mg/kg or 10ml/L or 10 ppm unless the medicine is administered by inhalation.
3075	LOBELIA POWDER	А, Н	The concentration in the medicine must be no more than 0.001% or 10mg/kg or 10ml/L or 10 ppm unless the medicine is administered by inhalation.
3076	LOLIUM PERENNE	A, H	

Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
3077	LOLIUM TEMULENTUM	A, H		
3078	LONGIFOLENE	E	Permitted for use only in combination with other permitted ingredients as a fragrance.  If used in a fragrance the total longifolene concentration in a medicine must be no more than 1%.	
3079	LONICERA CAPRIFOLIUM	A, E, H		
3080	LONICERA JAPONICA	A, E, H		
3081	LONICERA PERICLYMENUM	A, H		
3082	LOPHATHERUM GRACILE	A, H		
3083	LOQUAT	Е		
3084	LORANTHUS PARASITICUS	A, H		
3085	LOROPETALUM CHINENSIS	A, H		
3086	LOTUS CORNICULATUS	A, H		
3087	LOVAGE OIL	A, E, H		
3088	LOVAGE ROOT DRY	A, H		
3089	LOVAGE ROOT POWDER	A, H		
3090	LUDWIGIA PROSTRATA	A, H		
3091	LUFFA CYLINDRICA	A, H		
3092	LUFFA PURGANS	A, H		
3093	LUTEIN	A, E, H	When used as an excipient, permitted for use only as a colour in medicines limited to topical and oral routes of administration.	
3094	LYCHEE	E		
3095	LYCIUM BARBARUM	A, H		
3096	LYCIUM CHINENSE	A, E, H		
3097	LYCOPENE	A, E		
3098	LYCOPERSICON ESCULENTUM	A, E, H	Steroidal alkaloids calculated as solanine is a mandatory component of Lycopersicon esculentum.  The maximum daily dose must	

**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
			not provide more than 10 mg of steroidal alkaloids calculated as solanine.
3099	LYCOPODIUM ANNOTINUM	A, H	
3100	LYCOPODIUM CLAVATUM	A, H	
3101	LYCOPODIUM COMPLANATUM	A, H	
3102	LYCOPUS EUROPAEUS	A, H	
3103	LYCOPUS LUCIDUS	A, H	
3104	LYCOPUS VIRGINICUS	A, H	Pulegone is a mandatory component of Lycopus virginicus.  The concentration of pulegone in the medicine must be no more than 4%.
3105	LYGODIUM JAPONICUM	A, H	
3106	LYSIMACHIA CHRISTINAE	A, H	
3107	LYSIMACHIA VULGARIS	A, H	
3108	LYSINE	A, E	
3109	LYSINE HYDROCHLORIDE	A, E	
3110	LYTHRUM HYSSOPIFOLIA	A, H	
3111	LYTHRUM SALICARIA	A, H	
3112	LYTHRUM VERTICILLATUM	A, H	
3113	MACADAMIA INTEGRIFOLIA	A, E	
3114	MACADAMIA NUT	Е	
3115	MACADAMIA NUT OIL	Е	
3116	MACADAMIA TERNIFOLIA	A, E, H	
3117	MACE	E	Safrole is a mandatory component of Mace.  When used internally, the concentration of safrole in the medicine must be no more than 0.1%.  When used topically, the concentration of safrole in the medicine must be no more than 1.0%.
3118	MACE OIL	A, H	Safrole is a mandatory

Permissible in	igredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			component of Mace oil.  When used internally, the concentration of safrole in the medicine must be no more than 0.1%.
			When used topically, the concentration of safrole in the medicine must be no more than 1.0%.
			When the concentration of mace oil in the preparation is more than 50% and the nominal capacity of the container is 25 mL or less, a restricted flow insert must be fitted on the container.
3119	MACROCYSTIS PYRIFERA	A, E, H	Iodine is a mandatory component of Macrocystis pyrifera.
			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.
3120	MACROGOL 1000	E	
3121	MACROGOL 1450	Е	Only for use in topical medicines for dermal application.
3122	MACROGOL 1500	E	
3123	MACROGOL 1500 CASTOR OIL	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 in the

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

	ngredients and requirements	G.1. 2	
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements medicine must be no more than 2%.
3124	MACROGOL 200	Е	Only for use in topical medicines for dermal application.
3125	MACROGOL 20000	E	
3126	MACROGOL 300	Е	
3127	MACROGOL 3000	Е	
3128	MACROGOL 3350	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 form time to time.
3129	MACROGOL 40	E	Only for use in topical medicines for dermal application.
3130	MACROGOL 400	E	
3131	MACROGOL 4000	Е	
3132	MACROGOL 45000	Е	Only for use in topical medicines for dermal application.
3133	MACROGOL 600	Е	
3134	MACROGOL 6000	E	
3135	MACROGOL 600000	E	
3136	MACROGOL 800	Е	
3137	MACROGOL 8000	Е	
3138	MACROGOL 900	Е	Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.

Permissible in	gredients 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 0.95%.
3139	MACROGOL POLY(VINYL ALCOHOL) GRAFTED POLYMER	Е	Only for use in oral medicines.  The concentration in the
			medicine must be no more than 5%.
3140	MACROPIPER EXCELSUM VAR EXCELSUM	A, H	
3141	MAGNESIUM AMINO ACID	A, E, H	Only for use in oral medicines.
	CHELATE		The concentration of magnesium must be no more than 25% of the magnesium amino acid chelate.
			The requirements specified in paragraphs (a) to (c) below apply to a medicine that contains the ingredient that is:
			- listed in the Register on or after 1 March 2021; or
			- released for supply after 1 March 2022.
			<ul><li>(a) Magnesium is a mandatory component of magnesium amino acid chelate.</li></ul>
			(b) When used in a medicine:
			(i) not indicated for laxative (o related) use; and
			(ii) where the maximum recommended daily dose for:
			(A) children aged between 1 and 3 years (inclusive) provides 65 mg or more total magnesium from inorganic magnesium salts;
			(B) children aged between 4 and 8 years (inclusive) provides 110 mg or more total magnesium from inorganic magnesium salts; or
			(C) individuals aged 9 years or

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

	ngredients and requirements	C.1. 2	C.1. 4
Column 1	Column 2	Column 3	Column 4
<u>Item</u>	Ingredient name	Purpose	Specific requirements older provides 350 mg or more total magnesium from inorganic magnesium salts;
			the following warning statement is required on the medicine label:
			- (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).
			(c) the medicine must not be directed for use in infants younger than 12 months of age
3142	MAGNESIUM ASCORBATE	A, E, H	
3143	MAGNESIUM ASCORBATE MONOHYDRATE	A, E, H	
3144	MAGNESIUM ASCORBYL PHOSPHATE	Е	Only for use in topical medicines for dermal application.
3145	MAGNESIUM ASPARTATE	A, E, H	
3146	MAGNESIUM ASPARTATE DIHYDRATE	A, E, H	
3147	MAGNESIUM ASPARTATE TETRAHYDRATE	A, E, H	
3148	MAGNESIUM CARBONATE HYDRATE	A, E, H	
3149	MAGNESIUM CHLORIDE 4.5- HYDRATE	A	The requirements specified in paragraphs (a) to (c) below apply to a medicine that contains the ingredient that is:
			- listed in the Register on or after 1 March 2021; or
			- released for supply after 1 March 2022.
			(a) Magnesium is a mandatory component of magnesium chloride 4.5-hydrate.
			<ul><li>(b) When used in a medicine:</li><li>(i) with an oral route of administration;</li></ul>

Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
			(ii) not indicated for laxative (or related) use; and	
			(iii) where the maximum recommended daily dose for:	
			(A) children aged between 1 and 3 years (inclusive) provides 65 mg or more total magnesium from inorganic magnesium salts;	
			(B) children aged between 4 and 8 years (inclusive) provides 110 mg or more total magnesium from inorganic magnesium salts; or	
			(C) individuals aged 9 years of older provides 350 mg or more total magnesium from inorganic magnesium salts;	
			the following warning statement is required on the medicine label:	
			- (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).	
			(c) When the route of administration is oral, the medicine must not be directed for use in infants younger than 12 months of age.	
3150	MAGNESIUM CHLORIDE HEXAHYDRATE	A, E, H	The requirements specified in paragraphs (a) to (c) below apply to a medicine that contains the ingredient that is:	
			- listed in the Register on or after 1 March 2021; or	
			- released for supply after 1 March 2022.	
			(a) Magnesium is a mandatory component of magnesium chloride hexahydrate.	
			(b) When used in a medicine:	

**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	
			(i) with an oral route of administration;	
			(ii) not indicated for laxative (or related) use; and	
			(iii) where the maximum recommended daily dose for:	
			(A) children aged between 1 and 3 years (inclusive) provides 65 mg or more total magnesium from inorganic magnesium salts;	
			(B) children aged between 4 and 8 years (inclusive) provides 110 mg or more total magnesium from inorganic magnesium salts; or	
			(C) individuals aged 9 years of older provides 350 mg or more total magnesium from inorganic magnesium salts;	
			the following warning statement is required on the medicine label:	
			- (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).	
			(c) When the route of administration is oral, the medicine must not be directed for use in infants younger than 12 months of age.	
3151	MAGNESIUM CITRATE	A, E, H		
3152	MAGNESIUM CITRATE NONAHYDRATE	A, E, H		
3153	MAGNESIUM CITRATE TETRADECAHYDRATE	A, E, H		
3154	MAGNESIUM DIGLUTAMATE	A, E, H		
3155	MAGNESIUM GLUCONATE	A, E, H		
3156	MAGNESIUM GLYCEROPHOSPHATE	A, E, H		

Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
3157	MAGNESIUM GLYCINATE	A	Only for use in oral medicines
3158	MAGNESIUM GLYCINATE DIHYDRATE	A	Only for use in oral medicines
	DIHYDKATE		Magnesium is a mandatory component of Magnesium glycinate dihydrate.
			The percentage of Magnesium from Magnesium glycinate dihydrate should be calculated based on the molecular weight of Magnesium glycinate dihydrate.
3159	MAGNESIUM HYDROGEN PHOSPHATE	Н	The requirements specified in paragraphs (a) to (c) below apply to a medicine that contains the ingredient that is:
			- listed in the Register on or after 1 March 2021; or
			- released for supply after 1 March 2022.
			<ul><li>(a) Magnesium is a mandatory component of magnesium hydrogen phosphate.</li></ul>
			(b) When used in a medicine:
			(i) with an oral route of administration;
			(ii) not indicated for laxative (or related) use; and
			(iii) where the maximum recommended daily dose for:
			(A) children aged between 1 and 3 years (inclusive) provides 65 mg or more total magnesium from inorganic magnesium salts;
			(B) children aged between 4 and 8 years (inclusive) provides 110 mg or more total magnesium from inorganic magnesium salts; or
			(C) individuals aged 9 years o older provides 350 mg or mor

**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 total magnesium from inorganic magnesium salts;
			the following warning statement is required on the medicine label:
			- (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).
			(c) When the route of administration is oral, the medicine must not be directed for use in infants younger than 12 months of age.
3160	MAGNESIUM HYDROXIDE	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.
			The requirements specified in paragraph (a) below apply to a medicine that contains the ingredient that is:
			- listed in the Register before March 2021;
			<ul> <li>released for supply before or on 1 March 2022; and</li> </ul>
			- the following warning statement is not specified on the label:
			- (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).
			(a) When the medicine is not promoted or marketed as laxative, contains more than 2

Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			g magnesium hydroxide per maximum recommended daily dose, the following warning statements are required on the label:
			- (LAX5) 'This product contains [name of the herb(s) or the chemical component(s)
			- (LAX4) 'This product may have laxative effect'.
			The requirements specified in paragraphs (b) to (d) below apply to a medicine that contains the ingredient that is:
			- listed in the Register on or after 1 March 2021; or
			- released for supply after 1 March 2022.
			<ul><li>(b) Magnesium is a mandator component of magnesium hydroxide.</li></ul>
			(c) When used in a medicine:
			(i) with an oral route of administration;
			(ii) not indicated for laxative (or related) use; and
			(iii) where the maximum recommended daily dose for:
			(A) children aged between 1 and 3 years (inclusive) provides 65 mg or more total magnesium from inorganic magnesium salts;
			(B) children aged between 4 and 8 years (inclusive) provides 110 mg or more total magnesium from inorganic magnesium salts; or
			(C) individuals aged 9 years of older provides 350 mg or more total magnesium from inorganic magnesium salts;
			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		
			medicine label: - (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).		
			(d) When the route of administration is oral, the medicine must not be directed for use in infants younger than 12 months of age.		
3161	MAGNESIUM LYSINATE	A	Only for use in oral medicines		
3162	MAGNESIUM METHIONINATE	A	Only for use in oral medicines		
3163	MAGNESIUM NITRATE	E	Only for use in topical medicines for dermal application.		
3164	MAGNESIUM OROTATE	A, E, H			
3165	MAGNESIUM OROTATE DIHYDRATE	A, E, H			
3166	MAGNESIUM OXIDE	A, E, H	The requirements specified in paragraphs (a) to (c) below apply to a medicine that contains the ingredient that is: - listed in the Register on or		
			after 1 March 2021; or - released for supply after 1		
			March 2022.  (a) Magnesium is a mandatory component of magnesium oxide.		
			<ul><li>(b) When used in a medicine:</li><li>(i) with an oral route of</li></ul>		
			administration;		
			(ii) not indicated for laxative (or related) use; and		
			(iii) where the maximum recommended daily dose for:		
			(A) children aged between 1		

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			and 3 years (inclusive) provides 65 mg or more total magnesium from inorganic magnesium salts;
			(B) children aged between 4 and 8 years (inclusive) provides 110 mg or more total magnesium from inorganic magnesium salts; or
			(C) individuals aged 9 years o older provides 350 mg or mor- total magnesium from inorganic magnesium salts;
			the following warning statement is required on the medicine label:
			- (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).
			(c) When the route of administration is oral, the medicine must not be directed for use in infants younger than 12 months of age.
3167	MAGNESIUM PHOSPHATE PENTAHYDRATE	A, E, H	The requirements specified in paragraphs (a) to (c) below apply to a medicine that contains the ingredient that is:
			- listed in the Register on or after 1 March 2021; or
			- released for supply after 1 March 2022.
			<ul><li>(a) Magnesium is a mandatory component of magnesium phosphate pentahydrate.</li></ul>
			(b) When used in a medicine:
			(i) with an oral route of administration;
			(ii) not indicated for laxative (or related) use; and
			(iii) where the maximum

**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
			recommended daily dose for:  (A) children aged between 1 and 3 years (inclusive) provides 65 mg or more total magnesium from inorganic magnesium salts;
			(B) children aged between 4 and 8 years (inclusive) provides 110 mg or more total magnesium from inorganic magnesium salts; or
			(C) individuals aged 9 years of older provides 350 mg or more total magnesium from inorganic magnesium salts;
			the following warning statement is required on the medicine label:
			- (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).
			(c) When the route of administration is oral, the medicine must not be directed for use in infants younger than 12 months of age.
3168	MAGNESIUM PHOSPHATE TRIBASIC	<b>A</b> , E, H	Magnesium is a mandatory component of magnesium phosphate tribasic.
			The percentage of magnesium from magnesium phosphate tribasic should be calculated based on the molecular weight of magnesium phosphate tribasic.
			The requirements specified in paragraphs (a) to (b) below apply to a medicine that contains the ingredient that is:
			- listed in the Register on or after 1 March 2021; or
			- released for supply after 1

Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements March 2022.	
			(a) When used in a medicine:	
			(i) with an oral route of administration;	
			(ii) not indicated for laxative (or related) use; and	
			(iii) where the maximum recommended daily dose for:	
			(A) children aged between 1 and 3 years (inclusive) provides 65 mg or more total magnesium from inorganic magnesium salts;	
			(B) children aged between 4 and 8 years (inclusive) provides 110 mg or more total magnesium from inorganic magnesium salts; or	
			(C) individuals aged 9 years of older provides 350 mg or more total magnesium from inorganic magnesium salts;	
			the following warning statement is required on the medicine label:	
			- (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).	
			(b) When the route of administration is oral, the medicine must not be directed for use in infants younger than 12 months of age.	
3169	MAGNESIUM PYRUVATE	A	Only for use in oral medicines	
			The maximum recommended daily dose must be no more than 7 grams.	
3170	MAGNESIUM STEARATE	E		
3171	MAGNESIUM SULFATE	A, E, H	When used internally, the	

**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	
	DIHYDRATE		maximum recommended daily dose must not be more than 1.5g.	
			The requirements specified in paragraphs (a) to (c) below apply to a medicine that contains the ingredient that is:	
			- listed in the Register on or after 1 March 2021; or	
			- released for supply after 1 March 2022.	
			(a) Magnesium is a mandatory component of magnesium sulfate dihydrate.	
			(b) When used in a medicine:	
			(i) with an oral route of administration;	
			(ii) not indicated for laxative (or related) use; and	
			(iii) where the maximum recommended daily dose for:	
			(A) children aged between 1 and 3 years (inclusive) provides 65 mg or more total magnesium from inorganic magnesium salts;	
			(B) children aged between 4 and 8 years (inclusive) provides 110 mg or more tota magnesium from inorganic magnesium salts; or	
			(C) individuals aged 9 years of older provides 350 mg or more total magnesium from inorganic magnesium salts;	
			the following warning statement is required on the medicine label:	
			- (LAX6) 'Contains magnesium, which may have laxative effect or cause diarrhoea' (or words to that effect).	
			(c) When the route of	

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			administration is oral, the medicine must not be directed for use in infants younger than 12 months of age.
3172	MAGNESIUM SULFATE HEPTAHYDRATE	A, E, H	When used internally, the maximum recommended daily dose must not be more than 1.5 g.  The requirements specified in paragraphs (a) to (c) below apply to a medicine that contains the ingredient that is:  - listed in the Register on or after 1 March 2021; or  - released for supply after 1 March 2022.  (a) Magnesium is a mandatory component of magnesium sulfate heptahydrate.  (b) When used in a medicine:  (i) with an oral route of administration;  (ii) not indicated for laxative (or related) use; and  (iii) where the maximum recommended daily dose for:  (A) children aged between 1 and 3 years (inclusive) provides 65 mg or more total magnesium from inorganic magnesium salts;  (B) children aged between 4 and 8 years (inclusive) provides 110 mg or more total magnesium from inorganic magnesium salts; or  (C) individuals aged 9 years or older provides 350 mg or more total magnesium from inorganic magnesium salts; 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	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			- (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).
			(c) When the route of administration is oral, the medicine must not be directed for use in infants younger than 12 months of age.
3173	MAGNESIUM SULFATE MONOHYDRATE	A, E, H	When used internally, the maximum recommended daily dose must not be more than 1.5 g.
			The requirements specified in paragraphs (a) to (c) below apply to a medicine that contains the ingredient that is:
			- listed in the Register on or after 1 March 2021; or
			- released for supply after 1 March 2022.
			<ul><li>(a) Magnesium is a mandatory component of magnesium sulfate monohydrate.</li></ul>
			(b) When used in a medicine:
			(i) with an oral route of administration;
			(ii) not indicated for laxative (or related) use; and
			(iii) where the maximum recommended daily dose for:
			(A) children aged between 1 and 3 years (inclusive) provides 65 mg or more total magnesium from inorganic magnesium salts;
			(B) children aged between 4 and 8 years (inclusive) provides 110 mg or more total magnesium from inorganic magnesium salts; or
			(C) individuals aged 9 years or

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			older provides 350 mg or more total magnesium from inorganic magnesium salts;
			the following warning statement is required on the medicine label:
			- (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).
			(c) When the route of administration is oral, the medicine must not be directed for use in infants younger than 12 months of age.
3174	MAGNESIUM SULFATE TRIHYDRATE	A, E, H	When used internally, the maximum recommended daily dose must not be more than 1.5 g.
			The requirements specified in paragraphs (a) to (c) below apply to a medicine that contains the ingredient that is:
			- listed in the Register on or after 1 March 2021; or
			- released for supply after 1 March 2022.
			<ul><li>(a) Magnesium is a mandatory component of magnesium sulfate trihydrate.</li></ul>
			(b) When used in a medicine:
			(i) with an oral route of administration;
			(ii) not indicated for laxative (or related) use; and
			(iii) where the maximum recommended daily dose for:
			(A) children aged between 1 and 3 years (inclusive) provides 65 mg or more total magnesium from inorganic magnesium salts;

**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
			(B) children aged between 4 and 8 years (inclusive) provides 110 mg or more total magnesium from inorganic magnesium salts; or
			(C) individuals aged 9 years of older provides 350 mg or more total magnesium from inorganic magnesium salts;
			the following warning statement is required on the medicine label:
			- (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).
			(c) When the route of administration is oral, the medicine must not be directed for use in infants younger than 12 months of age.
3175	MAGNESIUM TRISILICATE	Е	The requirements specified in paragraphs (a) to (c) below apply to a medicine that contains the ingredient that is:
			- listed in the Register on or after 1 March 2021; or
			- released for supply after 1 March 2022.
			<ul><li>(a) Magnesium is a mandatory component of magnesium trisilicate.</li></ul>
			(b) When used in a medicine:
			(i) with an oral route of administration;
			(ii) not indicated for laxative (or related) use; and
			(iii) where the maximum recommended daily dose for:
			(A) children aged between 1 and 3 years (inclusive) provides 65 mg or more total

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			magnesium from inorganic magnesium salts;
			(B) children aged between 4 and 8 years (inclusive) provides 110 mg or more total magnesium from inorganic magnesium salts; or
			(C) individuals aged 9 years o older provides 350 mg or mor- total magnesium from inorganic magnesium salts;
			the following warning statement is required on the medicine label:
			- (LAX6) 'Contains magnesium, which may have a laxative effect or cause diarrhoea' (or words to that effect).
			(c) When the route of administration is oral, the medicine must not be directed for use in infants younger than 12 months of age.
3176	MAGNOLIA GLAUCA	A, H	
3177	MAGNOLIA LILIFLORA	A, H	
3178	MAGNOLIA OBOVATA	A, H	
3179	MAGNOLIA OFFICINALIS	A, E, H	
3180	MAGNOLIA SALICIFOLIA	A, H	
3181	MAIZE	Е	
3182	MAIZE BRAN	Е	
3183	MAIZE OIL	A, E, H	
3184	MAIZE STARCH	A, E, H	
3185	MALACHITE GREEN	Е	Permitted for use only as a colour for topical use.
3186	MALIC ACID	Е	Sponsors should consider the impact of excipients on the sensitivity of the skin to sunlight and should ensure the finished medicine is safe for it

**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.
3187	MALPIGHIA GLABRA	A, E, H	
3188	MALT EXTRACT	E	
3189	MALTITOL	E	
3190	MALTITOL SOLUTION	E	
3191	MALTODEXTRIN	E	Gluten is a mandatory component of Maltodextrin where the ingredient is derived from gluten containing grains such as wheat, barley, rye and oats.
3192	MALTOL	E	
3193	MALTONE	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 tha 5%.
3194	MALTOSE	Е	
3195	MALUS DOMESTICA	A, E, H	The concentration of amygdalin in the medicine must be no more than 0%.
3196	MALUS PUMILA	A, E, H	A medicine that contains the ingredient must not be listed in the Register on or after 2 March 2020 or be released for supply after 2 March 2021.
3197	MALUS SYLVESTRIS	A, H	
3198	MALVA MOSCHATA	A, H	
3199	MALVA SYLVESTRIS	A, E, H	
3200	MALVA VERTICILLATA	A, H	
3201	MANDARIN	Е	
3202	MANDARIN OIL	E	Permitted for use only in combination with other

ngredients and requirements		
Column 2	Column 3	Column 4
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%
MANDARIN OIL COLDPRESSED	A, E, H	When used internally, oxedring is a mandatory component of mandarin oil coldpressed.  The quantity of oxedrine in the maximum recommended daily dose must be no more than 30 milligrams.
MANDARIN OIL TERPENES	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%
MANDARIN RESIDUE	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%.
MANDARINAL 32048	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.  If used in a fragrance the total
	MANDARIN OIL COLDPRESSED  MANDARIN OIL TERPENES  MANDARIN RESIDUE	MANDARIN OIL COLDPRESSED A, E, H  MANDARIN OIL TERPENES E  MANDARIN RESIDUE 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
			fragrance concentration in a medicine must be no more than 1%.
3207	MANDRAGORA OFFICINARUM	А, Н	Atropine, hyoscine and hyoscyamine are mandatory components of Mandragora officinarum.
			The concentration in the medicine must be no more than 10 mg/kg or 10 mL/L or 0.001%.
			The concentration of atropine in the medicine must be no more than 100 micrograms/kg or 100 micrograms/L or 0.00001%.
			The concentration of hyoscine in the medicine must be no more than 300 micrograms/kg or 300 micrograms/L or 0.00003%.
			The concentration of hyoscyamine in the medicine must be no more than 300 micrograms/kg or 300 micrograms/L or 0.00003%.
3208	MANGANESE	Н	Only for use as an active homoeopathic ingredient.
3209	MANGANESE (II) DIASPARTATE	А, Н	Only for use in oral medicines.
3210	MANGANESE (II) GLYCINATE	A, H	Only for use in oral medicines.
3211	MANGANESE ACETATE TETRAHYDRATE	Н	Only for use as an active homoeopathic ingredient.
3212	MANGANESE AMINO ACID CHELATE	A, E, H	Only for use in oral medicines.  The concentration of  Manganese must be no more than 25% of the manganese

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			amino acid chelate.
3213	MANGANESE CHLORIDE TETRAHYDRATE	A, E, H	
3214	MANGANESE DIASPARTATE	A, E, H	Only for use in oral medicines.
3215	MANGANESE GLUCONATE	A, E, H	
3216	MANGANESE GLYCEROPHOSPHATE	A, E, H	
3217	MANGANESE OXIDE	A, E, H	
3218	MANGANESE SULFATE MONOHYDRATE	A, E, H	
3219	MANGANESE SULFATE TETRAHYDRATE	A, E, H	
3220	MANGIFERA INDICA	A, E, H	
3221	MANGO	E, H	
3222	MANIHOT ESCULENTA	A, H	
3223	MANNITOL	E	
3224	MARANTA ARUNDINACEA	A, H	
3225	MARINE SPONGE	Н	Only for use as an active homoeopathic ingredient.
3226	MARJORAM OIL SPANISH	A, E, H	When the concentration in the preparation is more than 50%, the nominal capacity of the container must be no more than 50 mL, the medicine must have a restricted flow insert fitted to the container and requires the following warning statement on the medicine label:  - (CHILD) 'Keep out of reach of children' (or words to that effect).
3227	MARJORAM OIL SWEET	A, E, H	When the concentration in the preparation is more than 50%, the nominal capacity of the container must be no more than 50 mL, the medicine must have a restricted flow insert fitted to

**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
	<b></b>		the container and requires the following warning statement on the medicine label:  - (CHILD) 'Keep out of reach of children' (or words to that effect).
3228	MARRUBIUM VULGARE	A, E, H	
3229	MARSDENIA CUNDURANGO	A, H	
3230	MARSHMALLOW ROOT DRY	A, H	
3231	MARSHMALLOW ROOT POWDER	A, H	
3232	MASSOIA LACTONE	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%.
3233	MASTIC	A, H	
3234	MATE ABSOLUTE	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%
3235	MATRICARIA CHAMOMILLA	A, E, H	
3236	MATRICARIA FLOWER DRY	A, E, H	
3237	MEADOWSWEET HERB DRY	А, Н	Methyl salicylate is a mandatory component of meadowsweet herb dry.
			Not to be included in medicines for use in the eye or on damaged skin.

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			When used internally, the concentration of methyl salicylate in the medicine mus 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:
			<ul> <li>the delivery device is engage into the container in such a way that prevents it from bein readily removed;</li> </ul>
			<ul> <li>direct suction through the delivery device results in delivery of no more than one dosage unit; and</li> </ul>
			<ul> <li>actuation of the spray device is ergonomically difficult for young children to accomplish.</li> </ul>
			The following warning statement is required on the medicine label:
			<ul> <li>(METSAL) 'Contains methy salicylate' (or words to that effect).</li> </ul>
			When for use in topical medicines for dermal application
			i) the concentration of methyl salicylate in the medicine mus not be more than 25%
			<ul><li>ii) the following warning statements are required on the medicine label:</li></ul>
			- (PREGNT2) 'Do not use if pregnant or likely to become

**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
			pregnant' (or words to that effect);
			<ul> <li>- (CHILD4) 'Do not use [this product/insert name of product in children 6 years of age or less';</li> </ul>
			<ul> <li>- (SENS) 'Application to skin may increase sensitivity to sunlight.' (or words to that effect);</li> </ul>
			<ul> <li>- (AVOID) 'Avoid prolonged exposure in the sun' (or words to that effect);</li> </ul>
			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'.
3238	MECOBALAMIN (CO- METHYLCOBALAMIN)	A	Only for use in oral medicines.
3239	MEDICAGO SATIVA	A, E, H	The level of l-canavanine must be no more than that of the dried leaf.
			When fresh leaf extract is used and the extraction ratio is between 34:1 and 46:1, the quantity of l-canavanine in the extract must not be more than that in the fresh leaf.
3240	MEDIUM CHAIN TRIGLYCERIDES	Е	
3241	MELALEUCA ALTERNIFOLIA	A, E, H	Cineole is a mandatory component of Melaleuca alternifolia.
			In liquid preparations when the concentration of cineole OR the concentration of oil or distillate in the preparation is more than 25%:

**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
			<ul><li>a) the nominal capacity of the container must be no more than</li><li>25 millilitres;</li></ul>
			b) a restricted flow insert must be fitted on the container; and
			c) the container must include the following warning statements on the medicine label:
			<ul> <li>- (CHILD) 'Keep out of reach of children' (or words to that effect); and</li> </ul>
			- (NTAKEN) 'Not to be taken'.
			In liquid preparations, when the concentration of cineole OR the concentration of oil or distillate in the preparation is more than 25% and the nominal capacity of the container is more than 15 millilitres but less than or equa to 25 millilitres the medicine must also have a child resistant closure.
3242	MELALEUCA CAJUPUTI	A, E, H	Cineole is a mandatory component of Melaleuca cajuputi.
			In liquid preparations, when the concentration of cineole OR the concentration of oil or distillate in the preparation is more than 25%:
			<ul><li>a) the nominal capacity of the container must be no more than 25 millilitres;</li></ul>
			b) a restricted flow insert must be fitted on the container; and
			c) the container must include the following warning statements on the medicine label:
			- (CHILD) 'Keep out of reach

**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
			of children' (or words to that effect); and
			- (NTAKEN) 'Not to be taken'. In liquid preparations, when the concentration of cineole OR the concentration of oil or distillate in the preparation is more than 25% and the nominal capacity of the container is more than 15 millilitres but less than or equa to 25 millilitres the medicine must also have a child resistant closure.
3243	MELALEUCA CITRINA	A, H	
3244	MELALEUCA DISSITIFLORA	А, Н	Cineole is a mandatory component of Melaleuca dissitiflora.
			In liquid preparations, when the concentration of cineole OR the concentration of oil or distillate in the preparation is more than 25%:
			<ul><li>a) the nominal capacity of the container must be no more than 25 millilitres;</li></ul>
			b) a restricted flow insert must be fitted on the container; and
			c) the container must include the following warning statements on the medicine label:
			- (CHILD) 'Keep out of reach of children' (or words to that effect); and
			- (NTAKEN) 'Not to be taken'.
			In liquid preparations, when the concentration of cineole OR the concentration of oil or distillate in the preparation is more than 25% and the nominal capacity of the container is more than 15 millilitres but less than or equa

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			to 25 millilitres the medicine must also have a child resistan closure.
3245	MELALEUCA ERICIFOLIA	A, E, H	Cineole is a mandatory component of Melaleuca ericifolia.
			In liquid preparations, when the concentration of cineole OR the concentration of oil or distillate in the preparation is more than 25%:
			<ul> <li>a) the nominal capacity of the container must be no more than 25 millilitres;</li> </ul>
			b) a restricted flow insert must be fitted on the container; and
			c) the container must include the following warning statements on the medicine label:
			- (CHILD) 'Keep out of reach of children' (or words to that effect); and
			- (NTAKEN) 'Not to be taken'
			In liquid preparations, when the concentration of cineole OR the concentration of oil or distillate in the preparation is more than 25% and the nominal capacity of the container is more than 15 millilitres but less than or equa to 25 millilitres the medicine must also have a child resistan closure.
3246	MELALEUCA LINARIIFOLIA	А, Н	Cineole is a mandatory component of Melaleuca linariifolia.
			In liquid preparations, when the concentration of cineole OR the concentration of oil or distillate in the preparation is

**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
			more than 25%:  a) the nominal capacity of the container must be no more than 25 millilitres;
			b) a restricted flow insert must be fitted on the container; and
			c) the container must include the following warning statements on the medicine label:
			<ul> <li>- (CHILD) 'Keep out of reach of children' (or words to that effect); and</li> </ul>
			- (NTAKEN) 'Not to be taken'.
			In liquid preparations, when the concentration of cineole OR the concentration of oil or distillate in the preparation is more than 25% and the nominal capacity of the container is more than 15 millilitres but less than or equa to 25 millilitres the medicine must also have a child resistant closure.
3247	MELALEUCA OIL	<b>A</b> , E, H	Cineole and cajuput oil are a mandatory components of Melaleuca Oil.
			When the plant preparation is oil and the concentration in the medicine is more than 25%, the nominal capacity of the container must be no more than 25 mL and the medicine requires the following warning statements on the medicine label:
			<ul> <li>- (CHILD) 'Keep out of reach of children' (or word to that effect)</li> </ul>
			- (NTAKEN) 'Not to be taken'
			When the nominal capacity of the container is 15 mL or less, then a restricted flow insert

Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
		•	must be fitted on the container.
			Where the nominal capacity of the container is more than 15 mL but less than or equal to 25 mL, then a child resistant closure and restricted flow insert must be fitted on the container.
3248	MELALEUCA QUINQUENERVIA	A, E, H	Cineole is a mandatory component of Melaleuca quinquenervia.
			In liquid preparations, when the concentration of cineole OR the concentration of oil or distillate in the preparation is more than 25%:
			<ul><li>a) the nominal capacity of the container must be no more than 25 millilitres;</li></ul>
			b) a restricted flow insert must be fitted on the container; and
			c) the container must include the following warning statements on the medicine label:
			- (CHILD) 'Keep out of reach of children' (or words to that effect); and
			- (NTAKEN) 'Not to be taken'
			In liquid preparations, when the concentration of cineole OR the concentration of oil or distillate in the preparation is more than 25% and the nominal capacity of the container is more than 15 millilitres but less than or equato 25 millilitres the medicine must also have a child resistan closure.
3249	MELICOPE PTELEIFOLIA	А, Н	
3250	MELILOTUS OFFICINALIS	A, E, H	Coumarin is a mandatory

**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
			component of Melilotus officinalis.
			The concentration of coumarin in the medicine must be no more than 0.001%.
3251	MELISSA OFFICINALIS	A, E, H	
3252	MELON	Е	
3253	MENADIONE SODIUM BISULFITE	Е	
3254	MENAQUINONE 7	A	For oral use only.
			The medicine must not provide more than 180 micrograms per maximum daily dose in adults, 90 micrograms per maximum daily dose in children between 10-18 years, and 45 micrograms per maximum daily dose in children less than 10 years of age.
3255	MENISPERMUM CANADENSE	A, H	
3256	MENTHA AQUATICA	A, H	Menthol is a mandatory component of Mentha aquatica
			When the medicine is for topical use for dermal application:
			(i) the medicine must not be intended for use in the eye or on damaged skin;
			(ii) the medicine must not deliver more than 25% total menthol when administered according to the directions for use;
			(iii) the following warning statement is required on the medicine label:
			- (EYE) Avoid contact with eyes (or words to that effect).
			(iv) if the medicine delivers more than 1% total menthol when administered according

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			to the directions for use, the following warning statements are required on the medicine label:
			<ul> <li>- (SKTEST) If you have sensitive skin, test this product on a small area of skin before applying it to a large area;</li> <li>- (IRRIT) If irritation developed discontinue use.</li> </ul>
			(v) if the medicine delivers more than 5% total menthol when administered according to the directions for use, the following warning statement i required on the medicine label
			<ul> <li>(MENTH) Contains a high concentration of menthol, which can cause severe skin irritation.</li> </ul>
			When the medicine is for internal use, the maximum recommended daily dose must not contain more than 1 gram of menthol.
3257	MENTHA ARVENSIS	A, E, H	Menthol is a mandatory component of Mentha arvensis
			When the medicine is for topical use for dermal application:
			(i) the medicine must not be intended for use in the eye or on damaged skin;
			(ii) the medicine must not deliver more than 25% total menthol when administered according to the directions for use;
			(iii) the following warning statement is required on the medicine label:
			- (EYE) Avoid contact with eyes (or words to that effect).

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Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			(iv) if the medicine delivers more than 1% total menthol when administered according to the directions for use, the following warning statements are required on the medicine label:
			<ul> <li>- (SKTEST) If you have sensitive skin, test this produc on a small area of skin before applying it to a large area;</li> <li>- (IRRIT) If irritation developed discontinue use.</li> </ul>
			(v) if the medicine delivers more than 5% total menthol when administered according to the directions for use, the following warning statement i required on the medicine label
			<ul> <li>(MENTH) Contains a high concentration of menthol, which can cause severe skin irritation.</li> </ul>
			When the medicine is for internal use, the maximum recommended daily dose must not contain more than 1 gram of menthol.
3258	MENTHA ARVENSIS LEAF OIL	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 be no more tha 5%.
			The total fragrance proprietary excipient formulation in a medicine must be no more 1% Menthol is a mandatory
			component of Mentha arvensis

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			leaf oil.  When the medicine is for topical use for dermal application:
			<ul><li>(i) the medicine must not be intended for use in the eye or on damaged skin;</li></ul>
			(ii) the medicine must not deliver more than 25% total menthol when administered according to the directions fo use;
			(iii) the following warning statement is required on the medicine label:
			- (EYE) Avoid contact with eyes (or words to that effect).
			(iv) if the medicine delivers more than 1% total menthol when administered according to the directions for use, the following warning statements are required on the medicine label:
			<ul> <li>(SKTEST) If you have sensitive skin, test this produce on a small area of skin before applying it to a large area;</li> </ul>
			- (IRRIT) If irritation develop
			(v) if the medicine delivers more than 5% total menthol when administered according to the directions for use, the following warning statement required on the medicine laber – (MENTH) Contains a high concentration of menthol,
			which can cause severe skin irritation.
			When the medicine is for internal use, the maximum recommended daily dose must not contain more than 1 gram of menthol.

**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
3259	MENTHA ARVENSIS OIL	E	Permitted for use only 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%.
			Menthol is a mandatory component of Mentha arvensis oil.
			When the medicine is for topical use for dermal application:
			(i) the medicine must not be intended for use in the eye or on damaged skin;
			(ii) the medicine must not deliver more than 25% total menthol when administered according to the directions for use;
			(iii) the following warning statements is required on the medicine label:
			- (EYE) Avoid contact with eyes (or words to that effect).
			(iv) if the medicine delivers more than 1% total menthol when administered according to the directions for use, the following warning statements are required on the medicine label:
			- (SKTEST) If you have sensitive skin, test this product on a small area of skin before applying it to a large area;
			- (IRRIT) If irritation develops, discontinue use.
			(v) if the medicine delivers more than 5% total menthol when administered according

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			to the directions for use, the following warning statement is required on the medicine label:  – (MENTH) Contains a high concentration of menthol, which can cause severe skin irritation.  When the medicine is for internal use, the maximum recommended daily dose must
			not contain more than 1 gram of menthol.
3260	MENTHA HAPLOCALYX	A, E, H	Menthol is a mandatory component of Mentha haplocalyx.
			When the medicine is for topical use for dermal application:
			(i) the medicine must not be intended for use in the eye or on damaged skin;
			(ii) the medicine must not deliver more than 25% total menthol when administered according to the directions for use;
			(iii) the following warning statement is required on the medicine label:
			- (EYE) Avoid contact with eyes (or words to that effect).
			(iv) if the medicine delivers more than 1% total menthol when administered according to the directions for use, the following warning statements are required on the medicine label:
			- (SKTEST) If you have sensitive skin, test this product on a small area of skin before applying it to a large area;
			- (IRRIT) If irritation develops, discontinue use.

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

1 CI IIII SSIDIC II	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	(v) if the medicine delivers more than 5% total menthol when administered according to the directions for use, the following warning statement is required on the medicine label – (MENTH) Contains a high concentration of menthol, which can cause severe skin irritation.  When the medicine is for internal use, the maximum recommended daily dose must
3261	MENTHA PULEGIUM	A, H	not contain more than 1 gram of menthol.  D-pulegone, menthol and volatile oil components (of Mentha pulegium) are mandatory components of
			Mentha pulegium.  When the nominal capacity of the container is more than 15 millilitres, the concentration o d-pulegone in the medicine must be no more than 4%.
			When the concentration of d- pulegone in the preparation is more than 4% and the nomina capacity of the container is 15 millilitres or less, the medicine must have a child resistant closure and restricted flow insert fitted on the container.
			The medicine requires the following warning statements on the medicine label: - (NTAKEN) 'Not to be taken'
			- (CHILD) 'Keep out of reach of children' (or words to that effect).
			When the medicine is for topical use for dermal application:
			a) the maximum recommende

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			daily dose must not contain more than 150 mg of Mentha pulegium oil or distillate;
			b) the medicine must not be intended for use in the eye or on damaged skin;
			c) the medicine must not deliver more than 25% total menthol when administered according to the directions for use;
			d) the following warning statement is required on the medicine label:
			- (EYE) Avoid contact with eyes (or words to that effect).
			e) if the medicine delivers more than 1% total menthol when administered according to the directions for use, the following warning statements are required on the medicine label:
			<ul> <li>- (SKTEST) If you have sensitive skin, test this produc on a small area of skin before applying it to a large area;</li> </ul>
			<ul> <li>(IRRIT) If irritation develops discontinue use.</li> </ul>
			f) if the medicine delivers more than 5% total menthol when administered according to the directions for use, the following warning statement is required on the medicine label
			<ul> <li>(MENTH) Contains a high concentration of menthol, which can cause severe skin irritation.</li> </ul>
			When the medicine is for internal use:
			<ul> <li>a) the maximum recommended daily dose must not contain more than 50 mg of Mentha pulegium oil or distillate;</li> </ul>

**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
			b) the maximum recommended daily dose must not contain more than 1 gram of menthol.
3262	MENTHA SPICATA	A, E, H	Menthol is a mandatory component of Mentha spicata.
			When the medicine is for topical use for dermal application:
			(i) the medicine must not be intended for use in the eye or on damaged skin;
			(ii) the medicine must not deliver more than 25% total menthol when administered according to the directions for use;
			(iii) the following warning statement is required on the medicine label:
			- (EYE) Avoid contact with eyes (or words to that effect).
			(iv) if the medicine delivers more than 1% total menthol when administered according to the directions for use, the following warning statements are required on the medicine label:
			<ul> <li>(SKTEST) If you have sensitive skin, test this product on a small area of skin before applying it to a large area;</li> </ul>
			<ul> <li>(IRRIT) If irritation develops discontinue use.</li> </ul>
			(v) if the medicine delivers more than 5% total menthol when administered according to the directions for use, the following warning statement is required on the medicine label
			<ul> <li>(MENTH) Contains a high concentration of menthol, which can cause severe skin irritation.</li> </ul>

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			When the medicine is for internal use, the maximum recommended daily dose must not contain more than 1 gram of menthol.
3263	MENTHA X CARDIACA	A, E, H	Menthol is a mandatory component of Mentha x cardiaca.
			When the medicine is for topical use for dermal application:
			(i) the medicine must not be intended for use in the eye or on damaged skin;
			(ii) the medicine must not deliver more than 25% total menthol when administered according to the directions for use;
			(iii) the following warning statement is required on the medicine label:
			- (EYE) Avoid contact with eyes (or words to that effect).
			(iv) if the medicine delivers more than 1% total menthol when administered according to the directions for use, the following warning statements are required on the medicine label:
			<ul> <li>(SKTEST) If you have sensitive skin, test this product on a small area of skin before applying it to a large area;</li> </ul>
			<ul> <li>(IRRIT) If irritation develops discontinue use.</li> </ul>
			(v) if the medicine delivers more than 5% total menthol when administered according to the directions for use, the following warning statement is required on the medicine label:

**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	
			<ul> <li>(MENTH) Contains a high concentration of menthol, which can cause severe skin irritation.</li> </ul>	
			When the medicine is for internal use, the maximum recommended daily dose must not contain more than 1 gram of menthol.	
3264	MENTHA X PIPERITA	A, E, H	Menthol is a mandatory component of Mentha x piperita.	
			When the medicine is for topical use for dermal application:	
			(i) the medicine must not be intended for use in the eye or on damaged skin;	
			(ii) the medicine must not deliver more than 25% total menthol when administered according to the directions for use;	
			(iii) the following warning statement is required on the medicine label:	
			- (EYE) Avoid contact with eyes (or words to that effect).	
			(iv) if the medicine delivers more than 1% total menthol when administered according to the directions for use, the following warning statements are required on the medicine label:	
			<ul> <li>(SKTEST) If you have sensitive skin, test this produc on a small area of skin before applying it to a large area;</li> </ul>	
			<ul> <li>(IRRIT) If irritation develops discontinue use.</li> </ul>	
			(v) if the medicine delivers more than 5% total menthol when administered according	

Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			to the directions for use, the following warning statement is required on the medicine label:
			<ul> <li>(MENTH) Contains a high concentration of menthol, which can cause severe skin irritation.</li> </ul>
			When the medicine is for internal use, the maximum recommended daily dose must not contain more than 1 gram of menthol.
3265	MENTHADIENYL ACETATE	E	Menthadienyl acetate must only be included in medicines when in combination with other permitted ingredients as a flavour proprietary excipient formulation.
			The total concentration of the flavour proprietary excipient formulation containing menthadienyl acetate must not be more than 5% of the total medicine.
3266	MENTHANYL 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%.
3267	MENTHOFURAN	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

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
3268	MENTHOL	A, E	When the medicine is for topical use for dermal application:
			<ul><li>(i) the medicine must not be intended for use in the eye or on damaged skin;</li></ul>
			(ii) the medicine must not deliver more than 25% total menthol when administered according to the directions for use;
			(iii) the following warning statement is required on the medicine label:
			- (EYE) Avoid contact with eyes (or words to that effect)
			(iv) if the medicine delivers more than 1% total menthol when administered according to the directions for use, the following warning statement are required on the medicine label:
			<ul> <li>(SKTEST) If you have sensitive skin, test this produ on a small area of skin before applying it to a large area;</li> </ul>
			<ul> <li>(IRRIT) If irritation develo discontinue use.</li> </ul>
			(v) if the medicine delivers more than 5% total menthol when administered according to the directions for use, the following warning statement required on the medicine lab
			<ul> <li>(MENTH) Contains a high concentration of menthol, which can cause severe skin irritation.</li> </ul>
			When the medicine is for internal use, the maximum recommended daily dose mu not contain more than 1 gran of menthol.

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
3269	MENTHONE	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%.
3270	MENTHONE GLYCERINE 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 medicine must be no more than 5%.
3271	MENTHONE THIOL FRACTION	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%.
3272	MENTHOXYPROPANEDIOL	Е	For oral use only.
			The concentration in the medicine must be no more than 0.04%.
3273	MENTHYL 2-HYDROXYETHYL CARBONATE	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

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
3274	MENTHYL 2-HYDROXYPROPYL CARBONATE	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%.
3275	MENTHYL ANTHRANILATE	A	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 not be more than 5%.
			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).
3276	MENTHYL 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%.
3277	MENTHYL LACTATE	Е	
3278	MENYANTHES TRIFOLIATA	A, H	
3279	MERCURIC CHLORIDE	Н	Only for use as an active homoeopathic ingredient.

1 CI IIIISSIDIC II	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
3280	MERCURY	Н	Only for use as an active homoeopathic ingredient.
3281	METACRESOL	E	Only for use in topical medicines for dermal application.
3282	METHACRYLIC ACID COPOLYMER	Е	Only for use in oral medicines.
3283	METHANOL	Е	The residual solvent limit is 30 mg per recommended daily dose.
			The concentration in the medicine must be no more than 0.3%.
3284	METHICONE	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%.
3285	METHIONINE	A, E	
3286	METHYL 2,6,6- TRIMETHYLCYCLOHEX-2-ENE- 1-CARBOXYLATE	E	Only for use in medicines in combination with other permitted ingredients as a fragrance proprietary excipient formulation.
			The total fragrance proprietary excipient formulation in a medicine must not be more than 1%.
3287	METHYL 2-METHYLBUTYRATE	Е	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.  If used in a flavour the total

**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	
			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%.	
3288	METHYL 2-OCTYNOATE	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%.	
3289	METHYL 3,6- DIMETHYLRESORCYLATE	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%.	
3290	METHYL ACETATE	Е	The residual solvent limit is 50 mg per recommended daily dose.	
			The concentration in the medicine must be no more than 0.5%.	
3291	METHYL ACETOPHENONE	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	

	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%.
3292	METHYL ACETYL RICINOLEATE	Е	Only for use in topical medicines for dermal application.
3293	METHYL ANISATE	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%.
3294	METHYL ANTHRANILATE	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%.
3295	METHYL BENZOATE	Е	Only for use in topical medicines for dermal application.
3296	METHYL 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 than 5%.
3297	METHYL CAPROATE	Е	Permitted for use only in combination with other

**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		
			permitted ingredients as a flavour.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
3298	METHYL CAPRYLATE	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%		
3299	METHYL CARBITOL	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%.		
3300	METHYL CEDRYL KETONE	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%.		
3301	METHYL CHAVICOL	E	Permitted for use only in combination with other permitted ingredients as part o a fragrance proprietary excipient formulation.		
			The ingredient is not to be included in medicines intended		

Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
			for oral use.	
			The quantity of methyl chavicol in a medicine must be no more than 0.01%.	
			The total fragrance proprietary excipient formulation in a medicine must be no more than 1%.	
3302	METHYL CINNAMATE	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%.	
3303	METHYL CIS-5-OCTENOATE	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%.	
3304	METHYL CYCLOPENTENOLONE	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%.	
3305	METHYL CYCLOPENTYLIDENEACETATE	Е	Permitted for use only in combination with other	

**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
			permitted ingredients as a fragrance.
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.
3306	METHYL DI-TERT-BUTYL-4- HYDROXYHYDROCINNAMATE	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%.
3307	METHYL DIHYDROABIETATE	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%.
3308	METHYL DIISOPROPYL PROPIONAMIDE	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%.
3309	METHYL ETHER	Е	Only for use in topical medicines for dermal application.
3310	METHYL ETHYL KETONE	Е	The residual solvent limit is 50 mg per maximum recommended daily dose.
			The concentration in the medicine must be no more than 0.5%.

**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		
3311	METHYL EUGENOL	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%.		
3312	METHYL FUROATE	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%.		
3313	METHYL GLUCETH-10	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%.		
			Residue levels of ethylene oxide are to be kept below the level of detection.		
3314	METHYL GLUCETH-20	Е	Only for use in topical medicines for dermal application.		
3315	METHYL GLUCETH-20 BENZOATE	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.		
			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	Permissible ingredients 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 than 1%.		
3316	METHYL GLUCETH-20 SESQUIHYDRATE	Е	Only for use in topical medicines for dermal application.		
3317	METHYL GLUCOSE DIOLEATE	E	Only for use in topical medicines for dermal application.		
3318	METHYL GLUCOSE SESQUIOLEATE	E	Only for use in topical medicines for dermal application.		
3319	METHYL GLUCOSE SESQUISTEARATE	E	Only for use in topical medicines for dermal application.		
3320	METHYL HEPTANOATE	Е	Permitted for use only 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%.		
3321	METHYL HEPTENONE	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%.		
3322	METHYL HEPTYL KETONE	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 or a fragrance.		
			If used in a flavour the total flavour concentration in a medicine must be no more th 5%.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more 19		
3323	METHYL HEXYL CARBINOL	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 the 5%.		
3324	METHYL HEXYL KETONE	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 th 5%.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more 19		
3325	METHYL HYDROGENATED ROSINATE	Е	Only for use in topical medicines for dermal application.		
3326	METHYL HYDROJASMONATE	E	Only for use in topical medicines for dermal application.		
3327	METHYL HYDROXYBENZOATE	Е			
3328	METHYL IONONE	Е	Permitted for use only in combination with other		

**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
			permitted ingredients as a 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%
3329	METHYL ISOBUTYL KETONE	Е	The residual solvent limit is 50 mg per maximum daily dose.
			The concentration in the medicine must be no more that 0.5%.
3330	METHYL ISOEUGENOL	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 tha 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
3331	METHYL 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 tha 5%.
3332	METHYL JASMONATE	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 that

Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
			5%.	
3333	METHYL LAURATE	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%.	
3334	METHYL LINOLEATE	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 tha 1%.	
3335	METHYL LINOLENATE	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 tha 1%.	
3336	METHYL MAGNESIUM CHLORIDE	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%.	
3337	METHYL METHACRYLATE	E		
3338	METHYL METHACRYLATE CROSSPOLYMER	Е	Only for use in topical medicines for dermal application and not to be included in medicines intended	

**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
			for use in the eye.  When the concentration of methyl methacrylate crosspolymer is greater than 1%, the medicine must not be intended for use on damaged skin.
			The concentration in the medicine must not be more than 4.85%.
3339	METHYL METHOXY PYRAZINE	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%.
3340	METHYL MYRISTATE	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%.
3341	METHYL NAPHTHYL KETONE	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%.
3342	METHYL NONYL KETONE	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

**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%.		
3343	METHYL NONYLENATE	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%.		
3344	METHYL OCTIN CARBONATE	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%.		
3345	METHYL PALMITATE	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%.		
3346	METHYL PHENYL CARBINOL	Е	Permitted for use only in combination with other permitted ingredients as a flavour.		
			If used in a flavour 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
			flavour concentration in a medicine must be no more than 5%.
3347	METHYL PHENYL CARBINYL- ISO-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 than 5%.
3348	METHYL PHENYL GLYCIDATE	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%.
3349	METHYL PHENYLACETATE	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%.
3350	METHYL PHENYLCARBINYL 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

**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 1%
3351	METHYL ROSINATE	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%.
3352	METHYL SALICYLATE	A, E	Not to be included in medicines for use in the eye or on damaged skin.
			When used internally, the concentration 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:
			<ul> <li>the delivery device is engage into the container in such a way that prevents it from being readily removed;</li> </ul>
			<ul> <li>direct suction through the delivery device results in delivery of no more than one dosage unit; and</li> </ul>
			<ul> <li>actuation of the spray device is ergonomically difficult for young children to accomplish.</li> <li>The following warning statement is required on the medicine label:</li> </ul>

**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
			<ul> <li>- (METSAL) 'Contains methyl salicylate' (or words to that effect).</li> </ul>
			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:
			<ul> <li>- (PREGNT2) 'Do not use if pregnant or likely to become pregnant' (or words to that effect);</li> </ul>
			<ul> <li>(CHILD4) 'Do not use [this product/insert name of product in children 6 years of age or less';</li> </ul>
			<ul> <li>(SENS) 'Application to skin may increase sensitivity to sunlight' (or words to that effect);</li> </ul>
			<ul> <li>- (AVOID) 'Avoid prolonged exposure in the sun' (or words to that effect);</li> </ul>
			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'.
3353	METHYL STEARATE	E	
3354	METHYL THIOBUTYRATE	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 that

**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	
			5%.	
3355	METHYL TRIMETICONE	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%.	
3356	METHYL-3- METHYLTHIOPROPIONATE	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%.	
3357	METHYL-BETA-METHYL THIOLPROPIONATE	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%.	
3358	METHYL-PARA-TERT-BUTYL PHENYLACETATE	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%.	
3359	METHYLBENZYL ACETATE	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

	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 than 1%.
3360	METHYLCELLULOSE	A, E	
3361	METHYLCHLOROISOTHIAZOLI NONE	Е	Only for use in topical medicines for dermal application that are rinsed off the skin.
			The total concentration of methylchloroisothiazolinone and methylisothiazolinone in the medicine must be no more than 0.0015%.
3362	METHYLCYCLOHEXADIENE	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%.
3363	METHYLENE BIS- BENZOTRIAZOLYL TETRAMETHYLBUTYLPHENOL	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:
			<ul> <li>- (AVOID) 'Avoid prolonged exposure in the sun' (or words to this effect); and</li> </ul>
			- (SUNPRO) 'Wear protective clothing - hats and eyewear when exposed to the sun' (or words to this effect).

the skin.  The total concentration of methylchloroisothiazolinon and methylisothiazolinon and methylisothiazolinon in the medicine must be no methan 0.0015%.  METHYLMERCAPTAN  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 to 5%.  METHYLPROPANEDIOL  E  Only for use in topical medicines for dermal application and not to be included in medicines inten for use in the eye.  The concentration in the medicine must be no more to 10%.  METHYLSILANOL/SILICATE  CROSSPOLYMER  Donly for use in topical medicines inten for use in the eye or on damaged skin.  The concentration in the	Permissible in	gredients and requirements		
METHYLISOTHIAZOLINONE  Body length of the skin.  The total concentration of methylehloroisothiazolinon and methylehloroisothiazolinon in the medicine must be no methan 0.0015%.  METHYLMERCAPTAN  Body length of the skin.  The total concentration of methylesothiazolinon in the medicine must be no methan 0.0015%.  METHYLMERCAPTAN  Body length of the skin.  If used in a flavour the total flavour concentration in a medicine must be no more to 5%.  METHYLPROPANEDIOL  Body length of the skin.  The concentration in the medicine must be no more to 10%.  METHYLSILANOL/SILICATE  CROSSPOLYMER  METHYLSILANOL/SILICATE  CROSSPOLYMER  Only for use in topical medicines for dermal application and not to be included in medicines inten for use in the eye or on damaged skin.  The concentration in the medicine must be no more to 10%.	Column 1	Column 2	Column 3	Column 4
medicines for dermal application that are rinsed of the skin.  The total concentration of methylchloroisothiazolinon and methylisothiazolinon and methylisothiazolinon and methylisothiazolinon and methylisothiazolinon in the medicine must be no methan 0.0015%.  3365  METHYLMERCAPTAN  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 to 5%.  3366  METHYLPROPANEDIOL  E  Only for use in topical medicines inten for use in the eye.  The concentration in the medicine must be no more to 10%.  3367  METHYLSILANOL/SILICATE  CROSSPOLYMER  E  Only for use in topical medicines for dermal application and not to be included in medicines inten for use in the eye.  The concentration in the medicines of the eye or on damaged skin.  The concentration in the medicine must be no more to 0.1%.	Item	Ingredient name	Purpose	Specific requirements
methylchloroisothiazolinona and methylisothiazolinone in the medicine must be no methan 0.0015%.  METHYLMERCAPTAN  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 to 5%.  METHYLPROPANEDIOL  E  Only for use in topical medicines for dermal application and not to be included in medicines inten for use in the eye.  The concentration in the medicine must be no more to 10%.  METHYLSILANOL/SILICATE  CROSSPOLYMER  METHYLSILANOL/SILICATE  E  Only for use in topical medicines for dermal application and not to be included in medicines for dermal application and not to be included in medicines inten for use in the eye or on damaged skin.  The concentration in the medicine must be no more to 0.1%.	3364	METHYLISOTHIAZOLINONE	E	medicines for dermal application that are rinsed off the skin.
combination with other permitted ingredients as a flavour.  If used in a flavour the total flavour concentration in a medicine must be no more to 5%.  3366  METHYLPROPANEDIOL  E  Only for use in topical medicines for dermal application and not to be included in medicines inten for use in the eye.  The concentration in the medicine must be no more to 10%.  3367  METHYLSILANOL/SILICATE  CROSSPOLYMER  METHYLSILANOL/SILICATE  E  Only for use in topical medicines for dermal application and not to be included in medicines inten for use in the eye or on damaged skin.  The concentration in the medicine must be no more to 0.1%.				methylchloroisothiazolinone and methylisothiazolinone in the medicine must be no more
flavour concentration in a medicine must be no more to 5%.  METHYLPROPANEDIOL  E  Only for use in topical medicines for dermal application and not to be included in medicines intent for use in the eye.  The concentration in the medicine must be no more to 10%.  METHYLSILANOL/SILICATE  CROSSPOLYMER  METHYLSILANOL/SILICATE  E  Only for use in topical medicines for dermal application and not to be included in medicines intent for use in the eye or on damaged skin.  The concentration in the medicine must be no more to 0.1%.	3365	METHYLMERCAPTAN	E	combination with other permitted ingredients as a
medicines for dermal application and not to be included in medicines intent for use in the eye.  The concentration in the medicine must be no more to 10%.  METHYLSILANOL/SILICATE E Only for use in topical medicines for dermal application and not to be included in medicines intent for use in the eye or on damaged skin.  The concentration in the medicine must be no more to 0.1%.				medicine must be no more tha
medicine must be no more to 10%.  METHYLSILANOL/SILICATE E Only for use in topical medicines for dermal application and not to be included in medicines inten for use in the eye or on damaged skin.  The concentration in the medicine must be no more to 0.1%.	3366	METHYLPROPANEDIOL	E	medicines for dermal application and not to be included in medicines intended
CROSSPOLYMER  medicines for dermal application and not to be included in medicines inten for use in the eye or on damaged skin.  The concentration in the medicine must be no more to 0.1%.				medicine must be no more tha
medicine must be no more to 0.1%.	3367		E	medicines for dermal application and not to be included in medicines intended for use in the eye or on damaged skin.
3368 METHYLSTYRENE/VINYLTOLU E Only for use in topical				medicine must be no more tha
ENE COPOLYMER medicines for dermal application.	3368		E	medicines for dermal
3369 MICA E Only for use when the route				

**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
			administration is oral, dental or topical.
			The concentration in oral medicines must be no more than 2.5%.
			The concentration in dental toothpastes must be no more than 0.5%.
3370	MICROCALICIUM ARENARIUM	A, H	
3371	MICROCOCCUS LUTEUS LYSATE	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 0.005%.
3372	MICROCOS PANICULATA	A, H	
3373	MICROCRYSTALLINE CELLULOSE	Е	
3374	MICROCRYSTALLINE WAX	E	Only for use as an excipient in medicines for topical, oral or oral application routes of administration.
			When microcrystalline wax is used as an excipient ingredient the route of administration 'oral' is only permitted when the dosage form is 'chewing gum'.
3375	MILK FAT	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%.
3376	MILK THISTLE FRUIT DRY	A, H	

Column 1	ngredients and requirements  Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
3377	MILK THISTLE FRUIT POWDER	A, H	Specific requirements
3378	MILLET	E	
3379	MILLETTIA DIELSIANA	A, H	
3380	MIMOSA ABSOLUTE	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%.
3381	MIMULUS GUTTATUS	A, H	
3382	MINT OIL DEMENTHOLISED	A, E, H	Menthol is a mandatory component of mint oil dementholised.
			When the medicine is for topical use for dermal application:
			(i) the medicine must not be intended for use in the eye or on damaged skin;
			(ii) the medicine must not deliver more than 25% total menthol when administered according to the directions for use;
			(iii) the following warning statement is required on the medicine label:
			- (EYE) Avoid contact with eyes (or words to that effect).
			(iv) if the medicine delivers more than 1% total menthol when administered according to the directions for use, the following warning statements are required on the medicine label:
			<ul> <li>(SKTEST) If you have sensitive skin, test this product on a small area of skin before applying it to a large area;</li> </ul>

**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
			- (IRRIT) If irritation develops, discontinue use.
			(v) if the medicine delivers more than 5% total menthol when administered according to the directions for use, the following warning statement is required on the medicine label:  – (MENTH) Contains a high concentration of menthol, which can cause severe skin irritation.
			When the medicine is for internal use, the maximum recommended daily dose must not contain more than 1 gram of menthol.
3383	MINTLACTONE	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%.
3384	MITCHELLA REPENS	A, H	
3385	MIXED (HIGH-ALPHA TYPE) TOCOPHEROLS CONCENTRATE	A, E	
3386	MIXED (LOW-ALPHA TYPE) TOCOPHEROLS CONCENTRATE	A, E	
3387	MIXED TERPENES	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%.
3388	MODIFIED FOOD STARCH	E	
3389	MOLASSES	Е	Permitted for use only in

**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  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%.
3390	MOLYBDENUM	Н	Only for use as an active homoeopathic ingredient.
			When Molybdenum is sourced from Molybdenum trioxide then the maximum daily dose must be no more than 125 micrograms.
			When Molybdenum is sourced from yeast - high molybdenum then the maximum recommended daily dose must be no more than 62.5 micrograms.
3391	MOLYBDENUM TRIOXIDE	A	Molybdenum is a mandatory component of Molybdenum trioxide.
			The maximum daily dose of molybdenum from Molybdenum trioxide must be no more than 125 micrograms
			The percentage of molybdenum from molybdenum trioxide should be calculated based on the molecular weight of molybdenum trioxide.
3392	MOMORDICA BALSAMINA	A, H	
3393	MOMORDICA CHARANTIA	A, H	
3394	MOMORDICA COCHINCHINENSIS	A, H	When Lycopene, Lutein or Betocarotene are declared as components, the plant part is restricted to fruit flesh, fruit peel or seed aril.

**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
3395	MONARDA DIDYMA	A, H	
3396	MONO- AND DI- GLYCERIDES	E	
3397	MONOBASIC AMMONIUM PHOSPHATE	E	Only for use in topical medicines for dermal application.
3398	MONOBASIC CALCIUM PHOSPHATE	A, E, H	
3399	MONOBASIC POTASSIUM PHOSPHATE	А, Е, Н	When used in a solid medicine containing this ingredient, the pH of a 10 g/L aqueous solution must not be more than 11.5.  When used in a liquid or a semi-solid medicine containing this ingredient, the pH of the medicine must be no more than 11.5.
3400	MONOBASIC SODIUM PHOSPHATE	A, E, H	When used in a solid preparation, the pH of a 10 g/L aqueous solution must not be more than 11.5.
			When used in a liquid or a semi-solid preparation, the pH of the preparation must not exceed 11.5.
3401	MONOBASIC SODIUM PHOSPHATE DIHYDRATE	Е	When used in a solid preparation, the pH of a 10 g/L aqueous solution must not be more than 11.5.
			When used in a liquid or a semi-solid preparation, the pH of the preparation must not exceed 11.5.
3402	MONOETHANOLAMINE	Е	Only for use in topical medicines for dermal application.
			The concentration in the medicine must be no more than 5%.

Permissible in	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4		
Item	Ingredient name	Purpose	Specific requirements		
3403	MONOMENTHYL SUCCINATE	E	Monomenthyl succinate must only be included in medicines when in combination with other permitted ingredients as a flavour proprietary excipient formulation.		
			The total concentration of the flavour proprietary excipient formulation containing monomenthyl succinate must not be more than 5% of the total medicine.		
3404	MONOPHOSPHOTHIAMINE	A			
3405	MONOPHOSPHOTHIAMINE DIHYDRATE	A			
3406	MONOPOTASSIUM GLUTAMATE	A, E			
3407	MONOSODIUM DIHYDROGEN CITRATE	Е			
3408	MONOSODIUM GLUTAMATE MONOHYDRATE	A, E			
3409	MONSTERA DELICIOSA	A, H			
3410	MONTAN WAX	Е			
3411	MORDANT RED 11	Е	Permitted for use only as a colour for topical use.  The concentration in the medicine must be no more than 0.05%		
3412	MORINDA CITRIFOLIA	A, H	Only for use when the plant part is fruit and the plant preparation is fruit juice or fruit powder.		
			Fruit powder must be produced by freeze drying the whole fruit (excluding the seeds).		
3413	MORINDA OFFICINALIS	A, H			
3414	MORINGA OLEIFERA	A, H			
3415	MORUS ALBA	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
3416	MORUS BOMBYCIS	A, H	
3417	MORUS NIGRA	A, E, H	
3418	MOSKENE	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%.
3419	MOTHERWORT HERB DRY	A, H	
3420	MOTHERWORT HERB POWDER	A, H	
3421	MUCUNA PRURIENS	A	Levodopa is a mandatory component of Mucuna pruriens.
			The concentration of levodopa in the medicine must not be more than 10 mg/kg or 10 mg/L or 0.001%.
3422	MULBERRY	E	
3423	MUNG BEAN	Е	
3424	MURRAYA KOENIGII	A, H	
3425	MURRAYA PANICULATA	A, H	
3426	MUSA X PARADISIACA	A, H	
3427	MUSK KETONE	E	Only for use in topical medicines for dermal application.
3428	MUSK TIBETENE	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%.
3429	MUSK XYLOL	E	Only for use in topical medicines for dermal application.

**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
3430	MUSKS	Н	Only for use as an active homoeopathic ingredient.
3431	MUSTARD	Е	Allyl isothiocyanate is a mandatory component of mustard 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%.
3432	MUSTARD OIL	Е	Allyl isothiocyanate is a mandatory component of mustard oil 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%.
3433	MUSTARD SEED OIL	Е	Allyl isothiocyanate is a mandatory component of mustard seed oil when the plan 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%.
3434	MYOSOTIS ARVENSIS	A, H	
3435	MYRCENE	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%.

**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%
3436	MYRCENYL 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%.
3437	MYRICA CERIFERA	A, E, H	
3438	MYRISTIC ACID	E	
3439	MYRISTIC ALDEHYDE	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%
3440	MYRISTICA FRAGRANS	A, E, H	Safrole is a mandatory component of Myristica fragrans.
			When for internal use then the concentration of safrole in the medicine must be no more than 0.1%.
			When for topical use then the concentration of safrole in the medicine must be no more than 1%.
			When the plant preparation is oil or distillate and the concentration in the medicine is more than 50%, the nominal capacity of the container must be no more than 25 millilitres,

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			the medicine must have a restricted flow insert fitted on the container and requires the following warning statement on the medicine label:
			- (CHILD) 'Keep out of reach of children' (or word to that effect).
3441	MYRISTYL ALCOHOL	Е	Only for use in topical medicines for dermal application.
3442	MYRISTYL LACTATE	E	Only for use in topical medicines for dermal application.
3443	MYRISTYL MYRISTATE	Е	Only for use in topical medicines for dermal application.
3444	MYROXYLON BALSAMUM	A, E, H	
3445	MYROXYLON BALSAMUM VAR. PEREIRAE	А, Н	
3446	MYRRH	A, H	
3447	MYRRH OIL	A, E, H	
3448	MYRRH RESIN	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%.
3449	MYRRHIS ODORATA	A, H	
3450	MYRSINE AFRICANA	A, H	
3451	MYRTENAL	Е	Permitted for use only in combination with other

**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		
			permitted ingredients as a flavour.		
			If used in a flavour the total flavour concentration in a medicine must be no more than 5%.		
3452	MYRTENYL ACETATE	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%.		
3453	MYRTLE ESSENCE MAX	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%.		
3454	MYRTLE OIL	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.		
			If used as a fragrance the total fragrance concentration in a medicine must be no more than 1%.		
3455	MYRTUS COMMUNIS	A, E, H			
3456	N,N'- BIS(SALICYLIDENE)PROPYLEN EDIAMINE	E	N,N'- Bis(salicylidene)propylenedia mine must only be included in medicines when in combination with other permitted ingredients as a proprietary excipient formulation in medicines with		

**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		
			administration for topical application.		
3457	N-BUTYL SULFIDE	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%		
3458	N-GLUCONYL ETHANOLAMINE	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%.		
3459	N-HEXYL 2-BUTENOATE	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%.		
3460	N-NONYL 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%		

**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
3461	NAPHTHALENE	Н	Only for use as an active homoeopathic ingredient.
3462	NARDOSTACHYS CHINENSIS	A, H	
3463	NARINGIN	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%.
3464	NASTURTIUM OFFICINALE	A, E, H	
3465	NATURAL FISH OIL	A, E	When therapeutic indications for this product are made against Vitamin A or colecalciferol (Vitamin D), they are mandatory components of natural fish oil. When for use in topical medicines, the concentration of 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

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			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 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.'
			When for internal use, the maximum recommended daily dose must be no more than 25 micrograms of Vitamin D.
3466	NAUCLEA OFFICINALIS	A, H	
3467	NELUMBO NUCIFERA	A, H	
3468	NELUMBO NUCIFERA FLOWER WAX	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 0.1%.
3469	NEOHESPERIDIN- DIHYDROCHALCONE	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 in the medicine must be no more than 0.1%

**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	
3470	NEOMENTHOL	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%.	
3471	NEOPENTYL GLYCOL DIHEPTANOATE	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 tha 25%.	
3472	NEOPENTYL GLYCOL DIISOSTEARATE	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%.	
3473	NEOPENTYL GLYCOL DIOCTANOATE	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 not be more than 8.1%.	
			When the concentration of neopentyl glycol dioctanoate i greater than 5%, the medicine must not be intended for use o damaged skin.	
3474	NEOPENTYL GLYCOL DIOCTANOATE/DIDECANOATE	E	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.
3475	NEOPICRORHIZA SCROPHULARIIFLORA	А, Н	
3476	NEPETA CATARIA	A, H	Pulegone is a mandatory component of Nepeta cataria and must be declared in the application.
			The concentration of pulegone in the medicine must be no more than 4%.
3477	NERAL	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%.
3478	NERIUM OLEANDER	А, Н	The concentration of equivalent dry Nerium oleander in the product must b no more than 1mg/Kg or 1mg/L or 0.0001%.
3479	NEROL	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 tha 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
3480	NEROL OXIDE	Е	Permitted for use only in combination with other permitted ingredients as part of a flavour or fragrance

**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	
			proprietary excipient formulation.	
			When used in a flavour, the total flavour proprietary excipient formulation in a medicine must be no more that 5%.	
			When used in a fragrance, the total fragrance proprietary excipient formulation in a medicine must be no more that 1%.	
3481	NEROLIDOL	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 that 5%.	
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%	
3482	NERONE	Е	Permitted for use only in combination with other permitted ingredients as part of a fragrance proprietary excipient formulation.	
			The total fragrance proprietar excipient formulation in a medicine must be no more that 1%.	
3483	NERYL 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 that 5%.	
			If used in a fragrance the total	

**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
		•	fragrance concentration in a medicine must be no more 1%.
3484	NERYL-ISO-BUTYRATE	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%.
3485	NICKEL	Н	Only for use as an active homoeopathic ingredient.
3486	NICOTIANA TABACUM	Н	Only for use as an active homoeopathic ingredient.
3487	NICOTINAMIDE	A, E, H	
3488	NICOTINAMIDE ASCORBATE	A, E	
3489	NICOTINAMIDE RIBOSIDE CHLORIDE	A	Only to be used in a medicine where Chromadex Inc (Client ID 68566), 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 02 December 2021.  Ribose is a mandatory
			component of nicotinamide riboside chloride.  Only permitted for use in medicines limited to oral routes of administration.
			The maximum recommended daily dose of the medicine must not provide more than 300 mg of nicotinamide riboside chloride.

**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
			The following warning statement (or words to the same effect) is required on the medicine label:
			<ul> <li>- (NTAKEN12) 'Not to be taken by children under 12 years old.'</li> </ul>
			When the maximum recommended daily dose of the medicine provides greater than 230 mg of nicotinamide riboside chloride, the following warning statement is required on the medicine label:  - (PREG) 'Not recommended for use during pregnancy or lactation'.
3490	NICOTINIC ACID	A, E	The medicine must contain no more than 100 mg of nicotinic acid per dosage unit.
3491	NIGELLA DAMASCENA	A, H	
3492	NIGELLA SATIVA	A, E, H	
3493	NITRIC ACID	E, H	The concentration of nitric acid in the medicine must be no more than 0.5%.
3494	NONADIENOL	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%
3495	NONANAL	E	Permitted for use only in combination with other permitted ingredients as a

Permissible in	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 than 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
3496	NONANOIC 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%
3497	NONFAT DRY MILK	Е, Н	
3498	NONIVAMIDE	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%.
3499	NONOXINOL 10	E	Only for use in topical medicines for dermal application.
3500	NONOXINOL 12	Е	For use in hand scrub formulations for healthcare professionals only.
			Only for use in topical medicines for dermal application and not to be included in medicines intended for use in the eye.

**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
			The concentration in the medicine must be no more tha 5%.
3501	NONOXINOL 5	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%.
3502	NONOXINOL 9	Е	Only for use in topical medicines for dermal application.
			The concentration in the medicine must be no more than 25%.
3503	NONYL 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%.
3504	NOOTKATONE	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%.
3505	NOPYL ACETATE	Е	Permitted for use only in combination with other permitted ingredients as a fragrance.  If used in a fragrance the total

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 than 1%.
3506	NORDIHYDROGUAIARETIC ACID	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 0.3%.
3507	NOTOPTERYGIUM FORBESII	A, H	
3508	NOTOPTERYGIUM INCISIUM	A, H	
3509	NUPHAR JAPONICA	A, H	
3510	NUPHAR LUTEA	A, H	
3511	NUTMEG DRY	A, E, H	Safrole is a mandatory component of Nutmeg Dry.
			When for internal use then the concentration of safrole from all ingredients in the medicine must be no more than 0.1%.
			When for topical use then the concentration of safrole from all ingredients in the medicine must be no more than 1%.
3512	NUTMEG OIL	A, E, H	Safrole is a mandatory component of Nutmeg oil.
			When for internal use then the concentration of safrole in the medicine must be no more than 0.1%.
			When for topical use then the concentration of safrole in the medicine must be no more than 1%.
			When the concentration of Nutmeg oil in the medicine is more than 50%, the nominal capacity of the container must be no more than 25 mL, 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 have a restricted flow insert fitted on the container and requires the following warning statement on the medicine label:  - (CHILD) 'Keep out of reach of children' (or words to that effect).
3513	NUTMEG POWDER	А, Е, Н	Safrole is a mandatory component of Nutmeg powder. When for internal use then the concentration of safrole in the medicine must be no more than 0.1%.  When for topical use then the concentration of safrole in the medicine must be no more than 1%.
3514	NUX VOMICA DRY	A, H	Strychnine (of Strychnos spp.) is a mandatory component of Nux Vomica Dry.  The concentration of in the medicine must be no more than 1mg/Kg or 1mg/L or 0.0001%.
3515	NUX VOMICA POWDER	Н	Only for use as an active homoeopathic ingredient.  Strychnine (of Strychnos spp.) is a mandatory component of Nux vomica powder.  The concentration in the medicine must be no more than 1mg/Kg or 1mg/L or 0.0001%.
3516	NYCTANTHES ARBOR-TRISTIS	A, H	When the plant part is leaf:  a) methyl salicylate is a mandatory component of Nyctanthes arbor-tristis;  b) not to be included in medicines for use in the eye or on damaged skin;

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			c) when used internally, the concentration of methyl salicylate in the medicine mus not be more than 0.001%;
			d) 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;
			e) 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:
			<ul> <li>the delivery device is engaged into the container in such a way that prevents it from being readily removed;</li> </ul>
			<ul> <li>direct suction through the delivery device results in delivery of no more than one dosage unit; and</li> </ul>
			<ul> <li>actuation of the spray device is ergonomically difficult for young children to accomplish</li> </ul>
			f) the following warning statement is required on the medicine label:
			<ul> <li>(METSAL) 'Contains methy salicylate' (or words to that effect); and</li> </ul>
			g) when for use in topical medicines for dermal application:
			i) the concentration of methyl salicylate in the medicine must not be more than 25%
			<ul><li>ii) the following warning statements are required on the medicine label:</li></ul>
			- (PREGNT2) 'Do not use if pregnant or likely to become

**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
			pregnant' (or words to that effect);
			<ul> <li>- (CHILD4) 'Do not use [this product/insert name of product in children 6 years of age or less';</li> </ul>
			<ul> <li>- (SENS) 'Application to skin may increase sensitivity to sunlight' (or words to that effect);</li> </ul>
			<ul> <li>- (AVOID) 'Avoid prolonged exposure in the sun' (or words to that effect);</li> </ul>
			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'.
3517	NYLON	E	Only for use in topical medicines for dermal application.
3518	NYLON 6/12	Е	Only for use in topical medicines for dermal application.
3519	NYLON-12	Е	Only for use in topical medicines for dermal application.
3520	NYMPHAEA ALBA	A, E, H	
3521	NYMPHAEA CAERULEA	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 to be no more than 0.3%.

**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
			Only for use in liquid extracts where the plant part is the flower and the solvent in 100% water.
3522	NYMPHAEA ODORATA	А, Н	
3523	OAK CHIPS EXTRACT	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%.
3524	OAKMOSS 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%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%.
3525	OAT	E, H	Only for use as an active homoeopathic or excipient ingredient.
			Gluten is a mandatory component of Oat when the route of administration is other than topical and mucosal.
3526	OAT BRAN	E	Gluten is a mandatory component of Oat bran when the route of administration is other than topical and mucosal.
3527	OATMEAL COLLOIDAL	А, Е	Gluten is a mandatory component of Oatmeal colloidal when the route of

**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
			administration is other than topical and mucosal.
3528	OCIMENE	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 tha 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
3529	OCIMENYL 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 tha 5%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
3530	OCIMUM BASILICUM	A, E, H	When the plant preparation is oil or distillate, Methyl chavicol, eugenol, methyleugenol and cineole are mandatory components of Ocimum basilicum.
			The concentration of methyleugenol in the medicine must not exceed 1%.
			When the concentration of Methyl chavicol in the medicine is more than 5%, the nominal capacity of the container must be no more that 25 millilitres.
			When the concentration of Methyl chavicol in the

Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements medicine is more than 5% and
			the nominal capacity of the container is 25 millilitres or less, a restricted flow insert must be fitted on the container and requires the following warning statement on the medicine label:
			- (CHILD) 'Keep out of reach of children' (or words to that effect). When the concentration of cineole OR eugenol in the preparation is more than 25%, the nominal capacity of the container must not be more than 25 millilitres and the following warning statements must be included on the medicine label:
			<ul> <li>(CHILD) 'Keep out of reach of children' (or words to that effect); and</li> </ul>
			- (NTAKEN) 'Not to be taken
			When the concentration of cineole OR eugenol in the preparation is more than 25% and the nominal capacity of the container is more than 15 millilitres but less than or equ to 25 millilitres, the medicine must have a child resistant closure and restricted flow insert fitted on the container.
			When the concentration of cineole OR eugenol in the preparation is more than 25% and the nominal capacity of the container is no more than 15 millilitres, the medicine must have a restricted flow insert fitted on the container.
			When the preparation is for topical use in the mouth, the preparation may not contain more than 5 millilitres of eugenol and the concentration

**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
			of eugenol in the product must not be greater than 25%.
3531	OCIMUM KILIMANDSCHARICUM	А, Н	Camphor is a mandatory component of Ocimum kilimandscharicum.
			In solid and semi solid preparations, the concentration of camphor must be no more than 12.5%.
			In liquid preparations, the nominal capacity of the container must be no more than 25 millilitres.
			In liquid preparations other than essential oils or distillates the concentration of camphor must be no more than 2.5%.
			In essential oil or distillate preparations when the concentration of camphor is more than 2.5%, the medicine must have a restricted flow insert fitted on the container and include the following warning statements on the medicine label:
			- (CHILD) 'Keep out of reach of children' (or words to that effect); and
			- (NTAKEN) 'Not to be taken'
			In essential oil or distillate preparations, if the concentration of camphor is more than 10%, and the nominal capacity of the container is more than 15 millilitres but less than or equa to 25 millilitres, the medicine must also have a child resistan closure fitted on the container.
3532	OCIMUM MINIMUM	A, H	
3533	OCIMUM TENUIFLORUM	A, H	When the plant part is oil or distillate, eugenol is a

Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			mandatory component of Ocimum tenuiflorum.
			When the concentration of eugenol in the preparation is more than 25%, the nominal capacity of the container must not be more than 25 millilitres and the following warning statements must be included of the medicine label:
			<ul> <li>(CHILD) 'Keep out of reach of children' (or words to that effect); and</li> </ul>
			- (NTAKEN) 'Not to be taken'
			When the concentration of eugenol in the preparation is more than 25% and the nominal capacity of the container is more than 15 millilitres but less than or equato 25 millilitres, the medicine must have a child resistant closure and restricted flow insert fitted on the container.
			When the concentration of eugenol in the preparation is more than 25% and the nominal capacity of the container is no more than 15 millilitres, the medicine must have a restricted flow insert fitted on the container.
			When the preparation is for topical use in the mouth, the preparation may not contain more than 5 mL of eugenol an the concentration of eugenol in the product must not be greate than 25%.
3534	OCOTEA ODORIFERA	A, H	Safrole is a mandatory component of Ocotea odorifera.
			When for internal use then the concentration of safrole 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 0.1%.
			When for topical use then the concentration of safrole in the medicine must be no more than 1%.
3535	OCTACOSANOL	E	
3536	OCTADECANAL	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%.
3537	OCTADECENE/MA COPOLYMER	Е	Only for use in topical medicines for dermal application.
3538	OCTAHYDRO-4,7-METHANO- 3AH-INDENE-3A-CARBOXYLIC ACID, ETHYL ESTER	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%.
3539	OCTAHYDROCOUMARIN	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%.
3540	OCTAN-1-OL	E	Permitted for use only in

Ingredient name	Column 3 Purpose	Column 4  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 tha 5%.  If used in a fragrance the total
Ingredient name	Purpose	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 that 5%.  If used in a fragrance the total
		If used in a flavour the total flavour concentration in a medicine must be no more that 5%.  If used in a fragrance the total
		fragrance concentration in a medicine must be no more 1%
OCTANAL DIMETHYL ACETAL	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 that 5%.
		If used in a fragrance the tota fragrance concentration in a medicine must be no more 19
OCTANOHYDROXAMIC ACID	Е	Only for use in topical medicines for dermal application and not to be included in medicines intende for use in the eye.
		The concentration in the medicine must be no more than 0.5%.
OCTANOIC ACID	A, E	When for topical use, the concentration in the medicine must be no more than 2% (w/w).
		When for excipient use, permitted for use only in combination with other permitted ingredients as part a flavour or fragrance proprietary excipient formulation.
	OCTANOHYDROXAMIC ACID	OCTANOHYDROXAMIC ACID E

**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 total flavour proprietary excipient formulation in a medicine must be no more tha 5%.
			When used in a fragrance, the total fragrance proprietary excipient formulation in a medicine must be no more than 1%.
3544	OCTENE-1	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%.
3545	OCTHILINONE	Е	Only for use in topical medicines for dermal application.
3546	OCTOCRYLENE	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: - (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).

Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
3547	OCTOXINOL 10	Е	Only for use in topical medicines for dermal application.
3548	OCTYL 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%.
3549	OCTYL CROTONATE	Е	Octyl crotonate must only be included in medicines when in combination with other permitted ingredients as a fragrance proprietary excipient formulation.
			The total concentration of the fragrance proprietary excipient formulation containing octyl crotonate must not be more than 1% of the total medicine.
3550	OCTYL HYDROXYSTEARATE	Е	Only for use in topical medicines for dermal application.
3551	OCTYL ISOBUTYRATE	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%.
3552	OCTYL ISONONANOATE	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.
3553	OCTYL 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:
			<ul> <li>- (AVOID) 'Avoid prolonged exposure in the sun' (or words to this effect); and</li> </ul>
			- (SUNPRO) 'Wear protective clothing - hats and eyewear when exposed to the sun' (or words to this effect).
3554	OCTYL PALMITATE	Е	Only for use in topical medicines for dermal application.
3555	OCTYL SALICYLATE	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 5%.
			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
			<ul> <li>(SUNPRO) 'Wear protective clothing - hats and eyewear when exposed to the sun' (or</li> </ul>

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

	rigredients and requirements	Colone 2	Column 4
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements words to this effect).
3556	OCTYL STEARATE	E	Only for use in topical medicines for dermal application.
3557	OCTYLBICYCLOHEPTENEDICA RBOXIMIDE	E	Only for use in topical medicines for dermal application.  The medicine requires the following warning statement on the medicine label:  - (OBCARB) 'Contains octylbicycloheptenedicarboxin ide' (or words to that effect).
3558	OCTYLDODECANOL	Е	Only for use in topical medicines for dermal application.
3559	OCTYLDODECETH-25	E	Only for use in topical medicines for dermal application and not to be included in medicines intender for use in the eye.  The concentration in the medicine must be no more that 5%.  Residual levels of 1,4-dioxane and ethylene oxide (and relate substances) are to be kept below the level of detection.
3560	OCTYLDODECYL CITRATE CROSSPOLYMER	E	Only for use in topical medicines for dermal application and not to be included in medicines intender for use in the eye or on damaged skin.  The concentration in the medicine must be no more that 12%.

**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
3561	OCTYLDODECYL NEOPENTANOATE	Е	Only for use in topical medicines for dermal application.
3562	OCTYLDODECYL STEARATE	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%.
3563	OCTYLDODECYL XYLOSIDE	Е	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 1.5%.
3564	OENANTHATE	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%.
3565	OENANTHE AQUATICA	Н	Only for use as an active homoeopathic ingredient.
			The maximum recommended daily dose must be no more than 1 mg of the equivalent dry herbal material.
3566	OENANTHE CROCATA	А, Н	The maximum recommended daily dose must be no more than 1 mg of the equivalent dry herbal material.

Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4	
Item	Ingredient name	Purpose	Specific requirements	
3567	OENOTHERA BIENNIS	A, E, H		
3568	OENOTHERA STRICTA	A, H		
3569	OKOUBAKA AUBREVILLEI	A, H		
3570	OLDENLANDIA DIFFUSA	A, E, H		
3571	OLEA EUROPAEA	A, E, H		
3572	OLEIC ACID	Е		
3573	OLETH-10	Е	Only for use in topical medicines for dermal application.	
3574	OLETH-2	Е	Only for use in topical medicines for dermal application.	
			Dioxane and Ethylene oxide are mandatory components of Oleth-2.	
			The concentration of Dioxane in the medicine must be no more than 10 mg/kg or 10 mg/L or 0.001%.	
			The concentration of Ethylene oxide in the medicine must be no more than 1 mg/kg or 1 mg/L or 0.0001%.	
3575	OLETH-20	Е	Only for use in topical medicines for dermal application.	
3576	OLETH-3	Е	Only for use in topical medicines for dermal application.	
3577	OLETH-3 PHOSPHATE	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 0.12%.	

**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
3578	OLETH-5	E	Only for use in topical medicines for dermal application.
3579	OLEYL ALCOHOL	Е	Only for use in topical medicines for dermal application.
3580	OLIBANUM OIL	A, E, H	
3581	OLIGOFRUCTOSE	A, E	
3582	OLIVE	Е	
3583	OLIVE OIL	A, E, H	
3584	OMEGA-3 FISH OIL PHYTOSTEROL ESTERS	A	The medicine requires the following warning statement on the medicine label: - (PREGNT) 'Not recommended for use by pregnant and lactating women (or words to that effect).'
3585	OMEGA-3-ACID ETHYL ESTERS 60	A	Docosahexaenoic acid, docosapentaenoic acid and eicosapentaenoic acid are mandatory components of omega-3-acid ethyl esters 60.  Only to be used in a medicine where DSM Nutritional Products Pty Ltd (Client ID 31685), 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 30 June 2021.  Only permitted for use in medicines that are for oral routes of administration.  The maximum recommended

Permissible in	gredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			must not provide more than 3750 milligrams of docosahexaenoic acid, docosapentaenoic acid and eicosapentaenoic acid combined.
			The following warning statements are required on the medicine label:  - (PREGNT) 'Not recommended for use by pregnant and lactating women (or words to that effect);  - (ACOAG) 'Individuals taking anticoagulants should seek medical advice before taking this product' (or words to that effect);  - (CHILD3) 'Use in children under 12 years is not recommended';  - (FOOD) 'To be taken with
2506	OMECA 2 ACID ETHNI ECTERC		food' (or words to that effect).
3586	OMEGA-3-ACID ETHYL ESTERS 90	A	Only for use in oral medicines The maximum recommended daily dose of the medicine must not provide more than: a) 4000 mg of omega-3-acid ethyl esters 90; and
			b) 3750 mg EPA, DHA and DPA combined, when used alone or in combination with other sources of omega-3 fatty acids.
			The following warning statements (or words to the same effect) are required on the medicine label:
			<ul> <li>- (ACOAG) 'Individuals takin anticoagulants should seek medical advice before taking this product.'</li> </ul>
			- (FOOD) 'To be taken with

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

Permissible in	gredients and requirements	Permissible ingredients and requirements				
Column 1	Column 2	Column 3	Column 4			
Item	Ingredient name	Purpose	Specific requirements			
			food.' - (PREG) 'Not recommended for use during pregnancy or lactation.' - (CHILD3) 'Use in children			
			under 12 years is not recommended.'			
3587	ONION	E				
3588	ONION OIL	A, H				
3589	ONONIS SPINOSA	A, E, H				
3590	ONOPORDUM ACANTHIUM	A, H				
3591	ONOSMODIUM VIRGINIANUM	A, H				
3592	OPHIOPOGON JAPONICUS	A, H				
3593	OPOPANAX CHIRONIUM	A, E, H	When used as an excipient, permitted for use only in combination with other permitted ingredients as part of a flavour or a fragrance proprietary excipient formulation.			
			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%			
3594	OPOPANAX 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 that 5%.			
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%			
3595	OPUNTIA FICUS-INDICA	A, H				

**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
3596	ORANGE	Е	
3597	ORANGE FLOWER 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%.
			If used in a fragrance the total fragrance concentration in a medicine must be no more 1%
3598	ORANGE FLOWER OIL	A, E, H	When used internally, oxedring is a mandatory component of orange flower oil.
			The quantity of oxedrine in the maximum recommended daily dose must be no more than 30 milligrams.
3599	ORANGE JUICE	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%.
3600	ORANGE JUICE 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%
3601	ORANGE OIL	A, E, H	When used internally, oxedring

**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
			is a mandatory component of orange oil.
			The quantity of oxedrine in the maximum recommended daily dose must be no more than 30 milligrams.
3602	ORANGE OIL BITTER	E	Permitted for use only in combination with other permitted ingredients as a flavour or a fragrance.
			If used in a flavor, 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%.
			The warning statement (SENS) 'Application to skin may increase sensitivity to sunlight' or words to that effect must be include on the medicine label unless the medicine is:
			a) for internal use;
			b) in preparations containing 1.4% or less of orange oil bitter;
			<ul> <li>c) for use in soaps or bath or shower gels that are washed of the skin.</li> </ul>
3603	ORANGE OIL BITTER COLDPRESSED	A, E, H	When used internally, oxedrine is a mandatory component of orange oil bitter coldpressed.
			The quantity of oxedrine in the maximum recommended daily dose must be no more than 30 milligrams.
			The warning statement (SENS 'Application to skin may increase sensitivity to sunlight' (or words to that effect) must be included on the medicine

Permissible in	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
Item	Ingredient name	Purpose	Specific requirements
			label unless the medicine is:
			a) for internal use; or
			<ul><li>b) in preparations containing</li><li>1.4% or less of orange oil bitte</li><li>coldpressed; or</li></ul>
			<ul> <li>c) for use in soaps or bath or shower gels that are washed of the skin.</li> </ul>
3604	ORANGE OIL COLD PRESSED	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%.
3605	ORANGE OIL DISTILLED	A, E, H	When used internally, oxedring is a mandatory component of orange oil distilled.
			The quantity of oxedrine in the maximum recommended daily dose must be no more than 30 milligrams.
3606	ORANGE OIL SWEET	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%.
3607	ORANGE OIL TERPENELESS	A, E, H	When used internally, oxedring is a mandatory component of

**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	
			orange oil terpeneless.  The quantity of oxedrine in the maximum recommended daily dose must be no more than 30 milligrams.	
3608	ORANGE PEEL	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 that 5%.	
3609	ORANGE PEEL DRIED BITTER	A, E, H	When used internally, oxedring is a mandatory component of orange peel dried bitter.	
			The quantity of oxedrine in the maximum recommended daily dose must be no more than 30 milligrams.	
3610	ORANGE PEEL OIL SWEET TERPENELESS	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%	
3611	ORANGE ROUGHY OIL	E	Only for use in topical medicines for dermal application.	
3612	ORIGANUM MAJORANA	A, H	Beta-arbutin is a mandatory component of Origanum majorana.  When for oral use, the	

	ngredients and requirements		
Column 1	Column 2	Column 3	Column 4
<u>Item</u>	Ingredient name	Purpose	Specific requirements
			maximum recommended daily dose must not provide more than 500 mg of beta-arbutin.
			When for dermal application exclusively to the face:
			a) the concentration of beta- arbutin in the medicine must not be more than 7%;
			b) hydroquinone is a mandatory component; and
			c) the concentration of hydroquinone must not be more than 10 mg/kg or 10 mg/L or 0.001%.
			When for use other than oral or dermal application exclusively to the face, the concentration of beta-arbutin in the medicine must not be more than 10 mg/kg or 10 mg/L or 0.001%.
			When the plant preparation is oil or distillate, and the concentration of Origanum majorana oil or distillate withit the medicine is more than 50%
			a) the nominal capacity of the container must not be more than 50 mL;
			b) a restricted flow insert must be fitted on the container; and
			c) the following warning statement is required on the label:
			- (CHILD) 'Keep out of reach of children' (or words to that effect).
3613	ORIGANUM OIL	Е	Permitted for use only in combination with other ingredients as a fragrance.
			If used as a fragrance the total concentration in the medicine must be no more than 1%.

**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		
3614	ORIGANUM OIL SPANISH	A, E, H			
3615	ORIGANUM VULGARE	A, E, H			
3616	ORNITHINE	A, E			
3617	ORNITHINE ASPARTATE	A, E			
3618	ORNITHINE MONOHYDROCHLORIDE	A, E			
3619	ORNITHOGALUM UMBELLATUM	A, H			
3620	OROSTACHYS FIMBRIATA	A, H			
3621	OROXYLUM INDICUM	A, H			
3622	ORRIS	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%.		
3623	ORRIS CONCRETE	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%.		
3624	ORRIS ROOT 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 tha 5%.		
3625	ORRIS ROOT OIL	A, E, H			
3626	ORRIS ROOT RESIN	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%.		
3627	ORTHO-TERT- BUTYLCYCLOHEXYL 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%.		
3628	ORTHOSIPHON ARISTATUS	A, H			
3629	ORYZA SATIVA	A, E, H			
3630	ORYZANOL	E			
3631	OSBECKIA CHINENSIS	A, H			
3632	OSMANTHUS 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%.		
3633	OSMANTHUS FRAGRANS	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%.		
3634	OTTELIA ALISMOIDES	A, H			
3635	OXACYCLOHEPTADEC-11-EN-2- ONE	E	Permitted for use only in combination with other		

**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		
			permitted ingredients as a fragrance.		
			If used in a fragrance the total fragrance concentration in a medicine must be no more than 1%.		
3636	OXACYCLOHEXADECAN-2-ONE	Е	Only for use in topical medicines for dermal application.		
3637	OXACYCLOHEXADECEN-2-ONE	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%.		
3638	OXALIC ACID	Н	Only for use as an active homoeopathic ingredient.		
3639	OXALIS ACETOSELLA	A, H			
3640	OXIDISED MAIZE STARCH	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%.		
3641	OXIDISED TAPIOCA STARCH	E			
3642	OXYBENZONE	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		

**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
			sunscreen products, the following warning statements are required on the label:
			<ul> <li>- (AVOID) 'Avoid prolonged exposure in the sun' (or words to this effect); and</li> </ul>
			<ul> <li>(SUNPRO) 'Wear protective clothing - hats and eyewear when exposed to the sun' (or words to this effect).</li> </ul>
3643	OYSTER	E	
3644	OYSTER SHELL	A, E, H	