

IFRA Standards





Acetic acid, anhydride, reaction products with 1,5,10-trimethyl-1,5,9cyclododecatriene

CAS N°:	144020-22-4 28371-99-5*	Empirical formula: Structure:	C17H26O
Synonyms:	Trimofix O (commerci Fixamber (commercia	ial name) al name)	

* This substance was previously erroneously identified as CAS 28371-99-5, however this CAS number is still used on certain commercial qualities today and as such this Standard is also applicable to that CAS number, which is an isomer of CAS 144020-22-4.

History:	Initial reviews:	New Standard	
	Current revision date:	2015	
	Implementation date:	For new submissions**:	August 10, 2015
		For existing fragrance compounds**:	August 10, 2016
	Next review date	2020	

** This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:							
Category 1 See Note box (1)	0.16 %	Category 7	0.42 %				
Category 2	0.20 %	Category 8	2.00 %				
Category 3	0.83 %	Category 9	5.00 %				
Category 4	2.49 %	Category 10	2.50 %				
Category 5	1.31 %	Category 11	See Note box (2)				
Category 6 3.99 %							
Note box:							
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOEI (International Organisation of the Elavor Industry -							

ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry www.iofi.org)

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

Acetic acid, anhydride, reaction products with 1,5,10-trimethyl-1,5,9cyclododecatriene

CRITICAL EFFECT:

DERMAL SENSITIZATION

RIFM SUMMARIES:

			Human Data		
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
5177 [1]	Weak	5510	NA	NA	5500

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No Observed Effect Level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = Lowest Observed Effect Level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003. ²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Acetic acid, anhydride, reaction products with 1,5,10trimethyl-1,5,9-cyclododecatriene and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 5500 μ g/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Acetic acid, anhydride, reaction products with 1,5,10-trimethyl-1,5,9-cyclododecatriene in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api, A. M., Basketter, D. A., Cadby, P. A., Cano, M-F., Ellis, G., Gerberick, G. F. et al, 2008. Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients. Regulatory Toxicology and Pharmacology 52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 2007. Acetic acid, anhydride, reaction products with 1,5,10-trimethyl-1,5,9-cyclododecatriene (Trimofix O): Assessment of skin sensitization potential using the local lymph node assays in the mouse. Unpublished study from I.F.F., Report number 56699 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2011. Repeated insult patch test with acid, anhydride, reaction products with 1,5,10-trimethyl-1,5,9-cyclododecatriene. Unpublished study from I.F.F., Report number 63990 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2011. Repeated insult patch test with acid, anhydride, reaction products with 1,5,10-trimethyl-1,5,9-cyclododecatriene. Unpublished study from I.F.F., Report number 63992 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. Repeated insult patch test with acid, anhydride, reaction products with 1,5,10-trimethyl-1,5,9-cyclododecatriene (Trimofix O). Unpublished study from I.F.F., Report number 64145 (RIFM, Woodcliff Lake, NJ, USA).

Acetyl ethyl tetramethyl tetralin



History:	Initial reviews:	November 1977, February 1980	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

NEUROTOXICITY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke, D.L.J. (1979), Food and Cosmetics Toxicology 17, 357-360. Spencer, P.S., Sterman, A.B et al. (1979), Neurotoxicology 1(1)



5-Acetyl-1,1,2,3,3,6-hexameth	yl indan ((AHMI)
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CAS N°:	15323-35-0	Empirical formula: Structure:	C ₁₇ H ₂₄ O H ₃ C H ₃ C H ₃ C H ₃ C CH ₃ CH ₃ CH ₃ CH ₃
Synonyms:	Acetyl hexamethyl indan 6-Acetyl-1,1,2,3,3,5-hexamethyl 1-(2,3-Dihydro-1,1,2,3,3,6-hexa Ethanone, 1-(2,3-dihydro-1,1,2,3 1,1,2,3,3,6-Hexamethylindan-5- Phantolid (commercial name)	lindane methyl-1h-inden-5-yl)e 3,3,6-hexamethyl-1H-iı yl methylketone	thanone nden-5-yl)-

History:	Initial reviews:	October 1978, October 1987, September 2001	
	Current revision date:	2015	
	Implementation date:	For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave-on products:	2%	Rinse- Includi	off products: ng household cleaning products	No restriction		
Non skin-contact products:	No restriction					
Note box:						
The Standard is set due to the phototoxic effects of the material. The limit only applies to applications on skin exposed to sunshine, excluding rinse- off products (please refer to Table 4 of the QRA booklet for more detailed information).						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

5-Acetyl-1,1,2,3,3,6-hexamethyl indan (AHMI)

CRITICAL EFFECT:

PHOTOTOXICITY

RIFM SUMMARIES:

Human studies - phototoxicity

The IFRA Standard is based upon two photoirritation studies in humans. In the first study, 10 volunteers were treated with 10% solution of 5-Acetyl-1,1,2,3,3,6-hexamethyl indan (AHMI) in 75% ethanol plus 25% diethyl phthalate on each forearm. Twenty-four hours later, one arm was irradiated (UVA) and the other served as a control. Observations immediately after radiation, at 24 hrs, and at 48 hours showed no phototoxic effects (RIFM, 1986). In the second study, 10 volunteers were treated with a 10% solution in 75% ethanol plus 25% diethyl phthalate on the back. After 30 minutes, the site was irradiated (UVA and UVB). Observations at 5 minutes after irradiation, and at 3, 24, 48, and 72 hours showed no phototoxic effects (RIFM, 1987).

Animal studies - phototoxicity

- 5, 20, 50 % in guinea pigs, photoirritation observed 20 and 50% (RIFM, 1978a).
- 5, 20% in rabbits, photoirritation observed at 5 and 20% (RIFM, 1978a).
- 1, 5, 10, 20% in guinea pigs and rabbits, photoirritation observed in guinea pigs and rabbits at 5, 10, and 20% (Ogoshi *et al.*, 1980; Ohkoshi *et al.*, 1981).
- 10% in guinea pigs, no photoirritation observed (Guillot et al., 1985).
- 1% in rabbits, photoirritation observed (RIFM, 1978).
- 1, 2, 4 % in rabbits, photoirritation observed (RIFM, 1985a; 1985b).
- 0.01, 1, 10, 25, 50% in hairless mice, photoirritation observed at 10, 25, 50% (RIFM, 1978c).

<u>Animal studies – photoallergy</u>

2% in guinea pigs, no photoallergy observed, 1/10 showed sensitization (RIFM, 1985c).

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 5-Acetyl-1,1,2,3,3,6-hexamethyl indan (AHMI) and recommended no change to the Standard (September 2001).

REFERENCES:

Guillot, J.P., Gonnet, J.F., Loquerie, J.F., Martini, M.C., Convert, P., and Cotte, J. (1985). A new method for the assessment of phototoxic and photoallergic potentials by topical applications in the albino guinea pig. J. Toxicol.-Cut. Ocu. Toxicol., 4(2), 117-133.

Ogoshi, K., Tanaka, N., and Sekine, A. (1980). A study on the phototoxicity of musk type fragrances. Unpublished. Presented at Society of Cosmetic Chemists, Japan. Report number 7465, 17 November.

Ohkoshi, K., Watanabe, A., and Tanaka, N. (1981). Phototoxicity of musks in perfumery. J. Society Cosmetic Chemists, Japan, 15(3), 207-213.

Research Institute for Fragrance Materials, Inc. (1978a). Phototoxicity of synthetic musks. Unpublished report from Shiseido laboratories. Report number 4415, 26 August.

Research Institute for Fragrance Materials, Inc. (1978b). Phototoxicity tests with 5-acetyl-1,1,2,3,3,6-hexamethylindan in albino rabbits. Unpublished report from Quest International. Report number 8055, 1 January.

Research Institute for Fragrance Materials, Inc. (1978c). Phototoxicity studies. RIFM report number 2042, 12 May.

Research Institute for Fragrance Materials, Inc. (1985a). Photosensitization test with 2% and 4% 5-acetyl-1,1,2,3,3,6-hexamethylindan in albino rabbits. Unpublished report from PFW Aroma Chemicals. Report number 29705, 1 November.

Research Institute for Fragrance Materials, Inc. (1985b). Photosensitization test with 1% 5-acetyl-1,1,2,3,3,6-hexamethylindan in albino rabbits. Unpublished report from PFW Aroma Chemicals. Report number 29706, 1 November.

Research Institute for Fragrance Materials, Inc. (1985c). Photosensitization test with 5-acetyl-1,1,2,3,3,6-hexamethylindan (17179) in guinea pigs. Unpublished report from PFW Aroma Chemicals. Report number 29704, 1 November.

Research Institute for Fragrance Materials, Inc. (1986). Phototoxicity testing in human subjects. RIFM report number 5748, 19 December.

Research Institute for Fragrance Materials, Inc. (1987). Phototoxicity testing in human subjects. RIFM report number 5743, 23 January.



Acetyl isovaleryl (5-methyl-2,3-hexanedione)

CAS N°:	13706-86-0	Empirical formula: Structure:	
Synonyms:	2,3-Hexanedione, 5-methyl- Acetyl isopentanoyl		

History:	Initial reviews:	February 1980, May 1983	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002

REFERENCES:

Opdyke, D.L.J., Letizia, C. (1982), Food and Chemical Toxicology 20, 637.

Acetylated Vetiver oil

CAS N°:	117-98-6 62563-80-8 68917-34-0 73246-97-6 84082-84-8	Empirical formula: Structure:	N/A N/A
Synonyms:	6-Azulenol, 1,2,3,3a,4,5,6,8a-octah 6-Azulenol, 1,2,3,3a,4,5,6,8a-octah 2-Isopropylidene-4,8-dimethyl-1,2,3 Vetivert acetate, Vetivert acetate (H Vetiverol, acetate	ydro-4,8-dimethyl-2-(methylethy ydro-4,8-dimethyl-2-(1-methylet ,3a,4,5,6,8a-octahydroazulen-6 aiti), Vetyvenyl acetate	/lidene)-, acetate hylidene)-, acetate -yl acetate

History:	Initial reviews:	July 2009		
	Current revision date:	2015		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	2020		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.07 %	Category 7	0.17 %			
Category 2	0.08 %	Category 8	2.00 %			
Category 3	0.35 %	Category 9	5.00 %			
Category 4	1.04 %	Category 10	2.50 %			
Category 5	0.55 %	Category 11	See Note Box (2)			
Category 6	1.67 %					
Note box:						
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organization of the Flavor Industry - www.iofi.org)						
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to negligible skin contact the concentration of a fragrance ingredient should not exceed the usual concentration of the fragrance compound in the finished product.						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

Acetylated Vetiver oil

CRITICAL EFFECT:

DERMAL SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (µg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
2910 [4]	Weak	2362	NA	NA	2300

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No Observed Effect Level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test;

LOEL = Lowest Observed Effect Level; NA = Not Available.

¹ Data derived from HRIPT or HMT.

²Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

³ WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Acetylated vetiver oil and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 2300 mg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Acetylated vetiver oil in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Jones, L., Foxenberg, R., Lalko, J., Letizia, C., Api, A, 2008. Sensitization potential of Vetiveryl acetate evaluated using the Local Lymph Node Assay. The Toxicologist, 102, 298.

RIFM (Research Institute for Fragrance Materials, Inc.), 2008. Local Lymph Node Assay. RIFM report number 55336, July 29. (RIFM, Woodcliff Lake, NJ, USA).

Alantroot oil

CAS N°:	97676-35-2	Empirical formula:	N/A
Synonyms:	Alantroot oil (<i>Inula helenium)</i> Elecampane oil Inula helenium oil		

History:	Initial reviews:	June 1975	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke, D.L.J. (1976), Food and Chemical Toxicology 14, 307.

Allyl esters

CAS N°:	N/A	Empirical formula:	N/A
Synonyms:	N/A		

History:	Initial reviews:	February 1977	
	Current revision date:	2009	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	2014	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

SPECIFICATION

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products:	N/A				
Note box:					
Fragrance material specificatio	ns:	Allyl esters should o of free allylalcohol i 0.1%. This recomm delayed irritant pote	only be used when the level n the ester is less than rendation is based on the ential of allylalcohol.		

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SEE FRAGRANCE MATERIAL SPECIFICATION



REXPAN RATIONALE / CONCLUSION:

IFRA specification noted.

REFERENCES:

Fd. Cosmet, Toxicol, 15,611-21 (1977)



History:	Initial reviews:	April 1989, April 1999, April 2005, May 2007		
	Current revision date:	e: Not applicable		
	Implementation date:	For new submissions*:	Not applicable	
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000

Allyl isothiocyanate

CAS N°:	57-06-7	Empirical formula: Structure:	C ₄ H ₅ N ₅
Synonyms:	AITC Allyl isosulfocyanate Allyl thiocarbonimide 1-Propenal, 3-isothiocyanato- 2-Propenyl isothiocyanate		

History:	Initial reviews:	Not applicable		
	Current revision date:	Not applicable		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:							
Skin contact products:							
Leave on products:	N/A	Rinse-off products:	N/A				
	Including household cleaning products						
Non skin contact products:	N/A						
Note box:	-						
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.							
Fragrance material specification	ons:	N/A					

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000



Allyl phenoxyacetate

CAS N°:	7493-74-5	Empirical formula: Structure:	$C_{11}H_{12}O_3$ H_2C
Synonyms:	Acetate PA Acetic acid, phenoxy-, 2-propeny 2-Propenyl phenoxyacetate	l ester	

History:	Initial reviews:	New Standard	
	Current revision date:	2009	
	Implementation date:	For new submissions*:	August 7, 2009
		For existing fragrance compounds*:	August 7, 2011
	Next review date	2014	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED / SPECIFICATION

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.02 %	Category 7	0.05 %			
Category 2	0.03 %	Category 8	0.70 %			
Category 3	0.11 %	Category 9	3.50 %			
Category 4	0.32 %	Category 10	2.50 %			
Category 5	0.17 %	Category 11	See Note box (2)			
Category 6 0.51 %						
Note box:						
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to negligible skin contact the concentration of a fragrance ingredient should not exceed the usual concentration of the fragrance compound in the finished product. 						
Fragrance material specifications: Please also refer to the IFRA Standard ALLYL ESTERS. Purity requirement: Allyl esters should only be used when the level of free allylalcohol in the ester is less than 0.1%. This recommendation is based on the delayed irritant potential of allylalcohol.						

Allyl phenoxyacetate

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (μg/cm ²)
775 [1] ⁴	Moderate	709	690	NA	700

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Data derived from HRIPT or HMT.

²Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

³WoE NESIL limited to two significant figures.

⁴EC3 value from one LLNA, not the mean.

⁵LOEL from human maximization test, not a human repeated insult patch test.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Allyl phenoxyacetate and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 700 mg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Allyl phenoxyacetate in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Food and Cosmetic Toxicology 15, 611-21 (1977)

RIFM (Research Institute for Fragrance Materials, Inc.), 1974a. Maximization study with allyl phenoxyacetate. RIFM report number 1801, April 16. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1974b. Maximization study with allyl phenoxyacetate. RIFM report number 1779, November 19. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007. Local Lymph Node Assay. RIFM report number 52909, May 21. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2008. Human repeated insult patch test. RIFM report number 54680, April 16a. (RIFM, Woodcliff Lake, NJ, USA).

alpha-Amyl cinnamic alcohol

CAS N°:	101-85-9	Empirical formula: Structure:	C ₁₄ H ₂₀ O HO
Synonyms:	Amylcinnamy α-Amylcinnar 2-Amyl-3-phe 2-Benzyliden 1-Heptanol, 2 α-Pentylcinna	1 alcohol nyl alcohol enyl-2-propen-1-ol eheptanol 2-(phenylmethylene)- amyl alcohol	

History:	Initial reviews:	New Standard	
	Current revision date:	2007	
	Implementation date:	For new submissions*:	June 16, 2007
		For existing fragrance compounds*:	June 16, 2009
	Next review date	2012	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.1 %	Category 7	0.3 %			
Category 2	0.1 %	Category 8	2.0 %			
Category 3	0.5 %	Category 9	5.0%			
Category 4	1.6 %	Category 10	2.5 %			
Category 5	0.8 %	Category 11	See Note box (2)			
Category 6 2.5 %						
Note box:						
(1) IFRA would recommend that any material us	sed to impart perfume or	flavour in products intended for human ir	ngestion should consist of			

ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry www.iofiorg.org)

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

alpha-Amyl cinnamic alcohol

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
>6250 [1]	Weak	3543	NA	NA	3500

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Data derived from HRIPT or HMT.

²Gerberick et al., 2001

³WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for alpha-amyl cinnamic alcohol and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 3500 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of alpha-amyl cinnamic alcohol in the various product categories. These were derived from the application of the exposurebased quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161)

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 2004a. Local Lymph Node Assay on alpha-Amylcinnamyl Alcohol. RIFM report number 45128, April 16. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004b. Repeated Insult Patch Test on alpha-Amylcinnamyl Alcohol. RIFM report number 46097, July 7A. (RIFM, Woodcliff Lake, NJ, USA).

alpha-Amyl cinnamic aldehyde

CAS N°:	122-40-7	Empirical formula: Structure:	C ₁₄ H ₁₈ O
Synonyms:	Amyl cinnamal Amyl cinnamic aldehyde α-Amylcinnamaldehyde α-Amyl β-phenylacrolein Heptanal, 2-phenylmethy Heptanal, 2-(phenylmethy α-Pentylcinnamaldehyde α-Pentyl-β-phenylacroleir 2-(Phenylmethylene)hept Flomine (commercial nam	lene)- ylene) n canal ne)	

History:	Initial reviews:	October 2009	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.7 %	Category 7	1.8 %			
Category 2	0.9 %	Category 8	2.0 %			
Category 3	3.6 %	Category 9	5.0 %			
Category 4	10.7 %	Category 10	2.5 %			
Category 5	5.6 %	Category 11	See Note box (2)			
Category 6	17.1 %					
Note box:						
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. 						
Fragrance material specifications: N/A						

alpha-Amyl cinnamic aldehyde

CONTRIBUTION FROM OTHER SOURCES:

See Annex II

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

LLNA weighted h (μg/α [no. st	iean EC3 values cm²) udies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
2942	2 [3]	Extremely weak	23622	NA	NA	23600
NOTI Na shaamiad a				an Massimal attac	TestulOFL	laura at

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹Data derived from HRIPT or Human Max tests ²Gerberick *et al.*, 2001 ³WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for alpha-amyl cinnamic aldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 23600 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of alpha-amyl cinnamic aldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Basketter, D.A. 2004. Unpublished data. Private communication to RIFM.

Elahi, E.N., Wright, Z., Hinselwood, D., Hotchkiss, S.A.M., Basketter D.A., Smith Pease, C.K., 2004. Protein binding and metabolism influence the relative skin sensitization potential of cinnamic compounds. Chemical Research in Toxicology, 17(3), 301-310.

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161

RIFM (Research Institute for Fragrance Materials, Inc.), 1994. Repeated Insult Patch Test on alpha-Amylcinnamaldehyde. RIFM report number 26498, July 29. (RIFM, Woodcliff Lake, NJ, USA).

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Repeated Insult Patch Test on alpha-Amylcinnamaldehyde. RIFM report number 47874, February 23. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2006. Local Lymph Node Assay on alpha-Amylcinnamaldehyde. DRAFT (RIFM, Woodcliff Lake, NJ, USA).

Smith, C. K. and Hotchkiss, S. A. M., 2001. Allergic Contact Dermatitis: Chemical and Metabolic Mechanisms. Taylor & Francis Ltd, London.

 $r \alpha$

Amylcyclopentenone

CAS N°:	25564-22-1	Empirical formula: Structure:	C10H16O
Synonyms:	2-Cyclopenten-1-one, 2-pentyl- 2-Pentyl-2-cyclopentenone 2-Pentylcyclopent-2-en-1-one		

History:	Initial reviews:	Nov. 1987, July 1994, May 2007	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.						
Fragrance material specificatio	ons:	N/A				

CONTRIBUTION FROM OTHER SOURCES:

N/A

Amylcyclopentenone

CRITICAL EFFECT:

SEE NOTE BOX

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000



CAS N°:	8015-64-3	Empirical formula: Structure :	N/A N/A
Synonyms:	Angelica archangelica oil Angelica archangelica root oil Angelica root oil (Angelica archangelica	L.)	

History:	Initial reviews:	June 1975, October 1978, September 2001			
	Current revision date:	2015			
	Implementation date:	For new submissions*: Not applicable			
		For existing fragrance compounds*:	Not applicable		
	Next review date	2020			

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave-on products:	0.8%	Rinse-off products:	No restriction			
		Including household	cleaning products			
Non skin contact products:	No restriction					
Note box:						
The Standard is set due to the phototoxic effect off products (please refer to Table 4 of the QRA	s of the material. The lim booklet for more detaile	nit only applies to applications or ad information).	n skin exposed to sunshine, excluding rinse-			
If combinations of phototoxic fragrance ingredients are used, the use levels have to be reduced accordingly. The sum of the concentrations of all phototoxic ingredients, expressed in % of their recommended maximum level in the consumer product shall not exceed 100.						
Note: See remark on phototoxic ingredients in the Introduction to the IFRA Standards (Appendix 8 to the IFRA Code of Practice) and the Standard on Citrus oil and other furocoumarins-containing essential oils.						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

PHOTOTOXICITY



RIFM SUMMARIES:

Two human phototoxicity studies were conducted. In one study, the test material at concentrations of 1% and 5% was applied to the backs of 30 male volunteers for 48 hours, under occlusion. Twenty-three hours after patch removal the sites were irradiated. Observations were made at 72 and 96 hours after application. No phototoxic reactions were observed in any subjects with either 1 or 5% concentrations of the test material (RIFM, 1975a). In a second study, the test material was applied neat to 13 male and female volunteers. Six hours later, the test sites were exposed to UVA radiation. Positive reactions were observed in 5/13 subjects (Kaidbey and Kligman, 1978, 1980).

- 4% on guinea pigs, UVA, photoirritation observed in all animals, 20/20 (Guillot, et al, 1985).
- 100% on hairless mice, UV, photoirritation observed (RIFM, 1974. Forbes, et al, 1977). 0.78, 1.56, 3.125, 6.25, 12.5, 25,
- 50% on hairless mice. UV. Photoirritation observed at concentrations of 1.56% and higher (RIFM, 1975b).
- 0.375, 0.75, and 1.5% on hairless mice. Photoirritation observed at all concentrations (RIFM, 1987).

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Angelica root oil and has no concerns with the current limit of 0.8% (September 2001).

REFERENCES:

Forbes P.D., Urbach F., and Davies R.E. (1977). Phototoxicity testing of fragrance raw materials. Food and Cosmetics Toxicology, 15, 55-60.

Guillot, J.P., Gonnet, J.F., Loquerie, J.F., Martini, M.C., Convert, P., and Cotte, J. (1985). A new method for the assessment of phototoxic and photoallergic potentials by topical applications in the albino guinea pig. Journal of Toxicology: Cutaneous and Ocular Toxicology, 4(2), 117-133.

Kaidbey, K.H. and Kligman, A.M. (1978). Identification of topical photosensitizing agents in humans. JID 70(3), 149-151.

Kaidbey, K.H. and Kligman, A.M. (1980). Identification of contact photosensitizers by human assay. Current Concepts in Cutaneous Toxicity, 55-68. Academic Press, NY.

Research Institute for Fragrance Materials, Inc. (1974). Phototoxicity and irritation test of fragrance materials in the mouse and miniature swine. RIFM report number 2037, 17 July.

Research Institute for Fragrance Materials, Inc. (1975a). Phototoxicity and irritation test of fragrance materials in the mouse and miniature swine. RIFM report number 2038, 4 February.

Research Institute for Fragrance materials, Inc. (1975b). Primary skin irritation and phototoxicity evaluation in human subjects with fragrance materials. RIFM report number 15092, December.

Research Institute for Fragrance Materials, Inc. (1987). Phototoxicity dilution assay of angelica root oil in hairless mice. RIFM report number 5147, 26 May.

Anisyl alcohol

CAS N°:	105-13-5 1331-81-3	Empirical formula: Structure:	C ₈ H ₁₀ O ₂
			OCH ₃
Synonyms:	Anisalcohol Anise alcohol Anisic alcohol Benzyl alcohol, p-methoxy p-Methoxybenzyl alcohol		

History:	Initial reviews:	May 2007	
	Current revision date:	2015	
	Implementation date:	For new submissions*:	August 10, 2015
		For existing fragrance compounds*:	August 10, 2016
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1	0.04 %	Category 7	0.11 %		
Category 2	0.06 %	Category 8	1.52 %		
Category 3	0.23 %	Category 9	5.00 %		
Category 4	0.68 %	Category 10	2.50 %		
Category 5	0.36 %	Category 11	See Note box (2)		
Category 6 See Note box (1)	1.09 %				
Note hox:					

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofi.org).

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

Anisyl alcohol

CRITICAL EFFECT:

DERMAL SENSITIZATION

RIFM SUMMARIES:

Anisyl alcohol - Sensitization Potency Estimation Based on Weight of Evidence

LLNA weighted mean EC3 values (µg/cm ²) [no. studies] Potency Classification ²		i			
	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – MAX (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)	
1475 [1]	Weak	NA	3448	NA	1500

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No Observed Effect Level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = Lowest Observed Effect Level; NA = Not Available

¹Data derived from HRIPT or Human Max tests ²Gerberick *et al.*, 2001 ³WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Anisyl alcohol and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1500 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Anisyl alcohol in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161)

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 1971. Maximization Test on Anisyl Alcohol. RIFM report number 1805, May 24. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Local Lymph Node Assay on Anisyl Alcohol. RIFM report number 48750, January 28. (RIFM, Woodcliff Lake, NJ, USA).

CAS N°:	943-88-4	Empirical formula: Structure:	
Synonyms:	3-Butene-2-one, 4-(4-methoxyph Methyl p-methoxycinnamyl ketor	nenyl) ester ne	

Anisylidene acetone (4-(p-methoxyphenil)-3-butene-2-one)

History:	Initial reviews:	November 1974	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke, D.L.J. (1975), Food and Chemical Toxicology 13, 456.



History:	Initial reviews:	December 1991		
	Current revision date:	Not applicable		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	December 1991	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED / RESTRICTED

cis- and trans-Asarone

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products:	N/A				
Note box:					
cis- and trans-Asarone as such should not be used as fragrance ingredients ; essential oils containing cis- or trans-asarone (e.g. calamus oils) should not be used at a level such that the total concentration of cis- and trans-asarone exceeds 0.01% in consumer products. This recommendation is based on similar biological effects to those of safrole (R.W. Wiseman, E.C. Miller et al. (1987), Cancer Res. 47,2275-2283).					
Fragrance material specificatio	ns:	N/A			

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX

REXPAN RATIONALE / CONCLUSION:

N/A

REFERENCES:

N/A

Benzaldehyde

CAS N°:	100-52-7	Empirical formula: Structure:	C7H6O
Synonyms:	Benzenecarbonal Benzene carboxaldehyde Benzenecarboxaldehyde Benzenemethylal Benzoic aldehyde Bitter almond oil, synthetic Phenylformaldehyde Phenylmethanol aldehyde		

History:	Initial reviews:	June 2009	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.02 %	Category 7	0.05 %			
Category 2	0.02 %	Category 8	0.60 %			
Category 3	0.09 %	Category 9	3.00 %			
Category 4	0.27 %	Category 10	2.50 %			
Category 5	0.14 %	Category 11	See Note box (2)			
Category 6	0.43 %					
Note box:						
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to negligible skin contact the concentration of a fragrance ingredient should not exceed the usual concentration of the fragrance compound in the finished product. 						
Fragrance material specification	ns:	N/A				

Benzaldehyde

CONTRIBUTION FROM OTHER SOURCES:

See Annex I and Annex II

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
> 6250 [1]4	Weak	590	NA	2760 ⁵	590

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Data derived from HRIPT or HMT.

²Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

³WoE NESIL limited to two significant figures.

⁴EC3 value from one LLNA, not the mean.

⁵LOEL from human maximization test, not a human repeated insult patch test.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for benzaldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 590 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of benzaldehyde in the various product categories. These were derived from the application of the exposurebased quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Basketter, D.A., Wright, Z., Gilmour, N.J., Ryan, C.A., Gerberick, G.F., Robinson, M.K., Dearman, R.J., Kimber, I., 2002. Prediction of human sensitization potency using local lymph node assay EC3 values. The Toxicologist, 66(1-S), 240.

RIFM (Research Institute for Fragrance Materials, Inc.), 1973. Maximization study with benzaldehyde. RIFM report number 1802, October 11a. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2009. Human repeated insult patch test. RIFM report number 57360. (RIFM, Woodcliff Lake, NJ, USA).



CAS N°:	71-43-2	Empirical formula: Structure:	C ₆ H ₆
Synonyms:	Benzol		

History:	Initial reviews:	1988	
	Current revision date:	January 2004	
	Implementation date:	For new submissions*:	May 6, 2004
		For existing fragrance compounds*:	May 6, 2005
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
The material should not be use	ed as fragrance i	ngredient for any app	olication.			
Fragrance material specificatio	ons:	The level of benzene practicable and shou fragrance compound. Since the introduction the use of benzene b been significant chan that permit the reduct level of this substanc technological improve of this solvent for the materials and in elimit impurity in alternative	has to be kept as low as Id never exceed 1 ppm in the of the original restriction on y IFRA in 1988, there have ges in manufacturing practices tion of the maximum permitted e. These include use of ements allowing replacement extraction of fragrance inating its presence as an e extraction solvents.			

Benzene

CONTRIBUTION FROM OTHER SOURCES:

None to consider

CRITICAL EFFECT:

CARCINOGENICITY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted - REXPAN, January 2004.

REFERENCES:

1) IARC (International Agency for Research on Cancer) Monographs Vol 7, p. 203 (1974); Vol 29, p. 93 and 391 (1982); Suppl. 7, p. 120 (1987)

2) CSTEE (Scientific Committee on Toxicity, Ecotoxicity and the Environment), Opinion on the results of the Risk Assessment of Benzene carried out in the framework of Council Regulation (EEC) 793/93 as adopted on Feb., 6, 2003



Benzyl alcohol C7H8O CAS N°: Empirical formula: 100-51-6 Structure: ΌН Benzenemethanol Synonyms: Benzylic alcohol alpha-Hydroxytoluene Phenylcarbinol Phenyl carbinol Phenylmethanol Phenylmethyl alcohol alpha-Toluenol

History:	Initial reviews:	New Standard	
	Current revision date:	2007	
	Implementation date:	For new submissions*:	June 16, 2007
		For existing fragrance compounds*:	June 16, 2009
	Next review date	2012	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:			
Category 1 See Note box (1)	0.2 %	Category 7	0.4 %
Category 2	0.2 %	Category 8	2.0 %
Category 3	0.9 %	Category 9	5.0 %
Category 4	2.7 %	Category 10	2.5 %
Category 5	1.4 %	Category 11	See Note box (2)
Category 6	4.3 %		
Note box:			
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. 			
Fragrance material specifications:		N/A	
CONTRIBUTION FROM OTHER SOURCES:			
See Annex I			
Benzyl alcohol

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

			Human Data		
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
>12500 [1]	Weak	5906	6897	8858	5900

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Data derived from HRIPT or HMT. ²Gerberick *et al.*, 2001 ³WeE NESU limited to two significant figures

³WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for benzyl alcohol and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 5900 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of benzyl alcohol in the various product categories. These were derived from the application of the exposurebased quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 1970. Maximization test on Benzyl Alcohol. RIFM report number 1760, October 7. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Repeated Insult Patch Test on Benzyl Alcohol. RIFM report number 45131, April 30. (RIFM, Woodcliff Lake, NJ, USA)

ÙSA).



Benzyl benzoate



History:	Initial reviews:	New Standard	
	Current revision date:	2007	
	Implementation date:	For new submissions*:	June 16, 2007
		For existing fragrance compounds*:	June 16, 2009
	Next review date	2012	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Category 1 See Note box (1)	1.7 %	Category 7	4.5 %	
Category 2	2.2 %	Category 8	2.0 %	
Category 3	8.9 %	Category 9	5.0 %	
Category 4	26.7 %	Category 10	2.5 %	
Category 5	14 %	Category 11	See Note box (2)	
Category 6	42.8 %			
Note box:				
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. 				
Fragrance material specifications: N/A				
CONTRIBUTION FROM OTHER SOURCES:				
See Annex I				

Benzyl benzoate

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

			Human Data		
LLNA weighted mean EC3 values (µg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
>12500 [1]	Extremely weak	59050	20690	NA	59000

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Data derived from HRIPT or HMT. ²Gerberick *et al.*, 2001 ³WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for benzyl benzoate and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 59000 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of benzyl benzoate in the various product categories. These were derived from the application of the exposure based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 1970. Maximization test on Benzyl Benzoate. RIFM report number 1760, June 1. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Repeated Insult Patch Test on Benzyl Benzoate. RIFM report number 47159, July 7A. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Local Lymph Node Assay on Benzyl Benzoate. RIFM report number 47377, January 20. (RIFM, Woodcliff Lake, NJ, USA).



Benzyl cinnamate



History:	Initial reviews:	New Standard	
	Current revision date:	2007	
	Implementation date:	For new submissions*:	June 16, 2007
		For existing fragrance compounds*:	June 16, 2009
	Next review date	2012	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.1 %	Category 7	0.4 %		
Category 2	0.2 %	Category 8	2.0 %		
Category 3	0.7 %	Category 9	5.0 %		
Category 4	2.1 %	Category 10	2.5 %		
Category 5	1.1 %	Category 11	See Note box (2)		
Category 6	3.4 %				
Note box:					

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org)

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

Benzyl cinnamate

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

			Human Data		
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
4600 [1]	Weak	4720	5517	NA	4700

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Data derived from HRIPT or HMT. ²Gerberick *et al.*, 2001

³WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for benzyl cinnamate and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 4700 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of benzyl cinnamate in the various product categories. These were derived from the application of the exposure based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 1972. Maximization test on Benzyl Cinnamate. RIFM report number 1804, June 1. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1975. Maximization test on Benzyl Cinnamate. RIFM report number 1799, March 27A. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005a. Local Lymph Node Assay on Benzyl Cinnamate. RIFM report number 48751, January 26. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005b. Repeated Insult Patch Test on Benzyl Cinnamate. RIFM report number 49109, June 23. RIFM, Woodcliff Lake, NJ, USA).

Benzyl cyanide



History:	Initial reviews:	New Standard**	
	Current revision date:	October 2003	
	Implementation date:	For new submissions*:	Not applicable**
		For existing fragrance compounds*:	Not applicable**
	Next review date	Not applicable**	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	N/A	Rinse-off products:	N/A	
		Including household	cleaning products	
Non skin contact products:	N/A			
Note box:				
The material as such should not be used as fragrance ingredient for any application.				
**This material has previously been included in the list of 'Other Materials'; the material therefore has already been prohibited before.				
Fragrance material specifications: N/A				



Benzyl cyanide

CONTRIBUTION FROM OTHER SOURCES:

On the basis of established maximum levels of this substance in commercially available natural sources (like essential oils and extracts), exposure to this substance from the use of these oils and extracts is not significant and the use of these oils is authorized as long as the level of benzyl cyanide in the finished product does not exceed 100 ppm. Furthermore, these natural extracts should not be used as substitutes for this substance.

Examples for potential natural sources (with maximum levels) of benzyl cyanide are provided below:

Fleur d'oranger absolute 1%

Frangipani (plumeria acutifolia) absolute 0.2%

Jasmine grandiflorum (Egypt) absolute traces

Jasmine multiflorum (India) absolute 0.3%

Karo karoundé absolute (ĆAS 68916-95-0) 4.75%

Tuberose (India) absolute 0.8%

CRITICAL EFFECT:

RELEASE OF CYANIDE

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted - REXPAN, October 2003

REFERENCES:

Potter et al., 2001, Food and Chemical Toxicology 39 (2), page 141-146.

Potter et al., 2001, Food and Chemical Toxicology 39 (2), page 147-151.



Benzylidene acetone (4-Phenyl-3-buten-2-one)

CAS N°:	122-57-6	Empirical formula: Structure:	
Synonyms:	3-Buten-2-one, 4-phenyl- Benzilideneacetone Methyl styryl ketone		

History:	Initial reviews:	lune 1974	
rinstory.	Current revision data:	Contembor 2002	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke, D.L.J. (1973), Food and Chemical Toxicology 11, 1021.

Private communication to IFRA.





History:	Initial reviews:	New Standard	
	Current revision date:	2007	
	Implementation date:	For new submissions*:	June 16, 2007
		For existing fragrance compounds*:	June 16, 2009
	Next review date	2012	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.5 %	Category 7	1.3 %			
Category 2	0.7 %	Category 8	2.0 %			
Category 3	2.7 %	Category 9	5.0 %			
Category 4	8.0 %	Category 10	2.5 %			
Category 5	4.2 %	Category 11	See Note box (2)			
Category 6	12.8 %					
Note box:						
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragmene ingredient in the finished product. 						

Fragrance material specifications:

N/A

Benzyl salicylate

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data				
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)	
725 [1]	Weak	17717	20690	NA	17700	

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Data derived from HRIPT or HMT. ²Gerberick *et al.*, 2001

³WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for benzyl salicylate and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 17700 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of benzyl salicylate in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161

RIFM (Research Institute for Fragrance Materials, Inc.), 1970. Maximization test on Benzyl Salicylate. RIFM report number 1760, October 7. (RIFM, Woodcliff Lake, NJ, USA).

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 1975a. Maximization test on Benzyl Salicylate. RIFM report number 1798, March 28. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1975b. Maximization test on Benzyl Salicylate. RIFM report number 1799, March 27. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Repeated Insult Patch Test on Benzyl Salicylate. RIFM report number 45129, May 3. RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Local Lymph Node Assay on Benzyl Salicylate. RIFM report number 47378, January 20. (RIFM, Woodcliff Lake, NJ, USA)

Bergamot oil expressed

CAS N°:	908007-75-8	Empirical formula: Structure :	N/A N/A
Synonyms:	N/A		

History:	Initial reviews:	October 1974, June 1992	
	Current revision date:	2015	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave-on products:	0.4%	Rinse-off products:	No Restriction		
		Including household	cleaning products		
Non skin contact products:	No Restriction				
Note box:					
The Standard is set due to the phototoxic effect off products (please refer to Table 4 of the QRA	s of the material. The lin booklet for more detaile	nit only applies to applications or ad information).	n skin exposed to sunshine, excluding rinse-		
If combinations of phototoxic fragrance ingredie phototoxic ingredients, expressed in % of their	ents are used, the use lev recommended maximum	vels have to be reduced accordir level in the consumer product s	ngly. The sum of the concentrations of all hall not exceed 100.		
Note: See remark on phototoxic ingredients in the Introduction to the IFRA Standards (Appendix 8 to the IFRA Code of Practice) and the Standard on Citrus oil and other furocoumarins-containing essential oils.					
For qualities of the expressed oil in which the less volatile components have been concentrated by partial or total removal of the terpene fraction, this limit should be reduced in proportion to the degree of concentration.					
Fragrance material specification	ons:	N/A			

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

PHOTOTOXICITY



Bergamot oil expressed

REXPAN RATIONALE / CONCLUSION:

For applications on areas of skin exposed to sunshine, excluding bath preparations, soaps and other products which are washed off the skin, bergamot oil expressed should not be used such that the level in the consumer products exceeds 0.4%.

REFERENCES:

These recommendations are based on the published literature on the phototoxicity of this material, summarized by D.L. Opdyke, Fd. Cosm. Toxicol. 11,1031 (1973) and other investigations published in Contact Dermatitis 3,225 (1977).



Birch wood pyrolysate

CAS N°:	8001-88-5 84012-15-7 85940-29-0 68917-50-0	Empirical formula:	N/A
Synonyms:	For the crude material banned: Birch tar oil, crude For the distillates specified: Birch tar oil dephenolated Birch tar oil rectified Essence bouleau dephenolisée Essence bouleau (Goudron) rect.		

History:	Initial reviews:	December 1996, October 2003	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

SPECIFICATION / PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
	Including household cleaning products				
Non skin contact products:	N/A				
Note box:					
Crude birch wood (bark) pyrolysates (oils) deriv Betula Lenta or Betula Alba should not be use	ed by pyrolysis (destruct d as a fragrance ingree	tive distillation) of the wood or ba dient.	ark of Betula Pubescens, Betula Pendula,		
Only rectified (purified) birch tar oils being in co	mpliance with the below	limitations for polynuclear aroma	atic hydrocarbons should be used.		
Fragrance material specifications: Limit content of polynuclear aromatic hydrocarbons (PAH) resulting from the use of rectified oils according to Good Manufacturing Practice.					
Benzopyrene and 1,2-Benzanthracene are to be used as markers for PAH. If used alone or in combination with rectified Cade oil, rectified					
Styrax oil or rectified Opoponax oil, the total concentration of both o the markers should not exceed 1 ppb in the final product.					

CONTRIBUTION FROM OTHER SOURCES:

N/A

Birch wood pyrolysate

CRITICAL EFFECT:

CARGINOGENICITY, GENOTOXICITY*

*Some of the polynuclear aromatic hydrocarbons are known to be carcinogen or genotoxic materials.

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, October 15, 2003

Bitter orange peel oil expressed

CAS N°:	68916-04-1 72968-50-4	Empirical formula: Structure :	N/A N/A
Synonyms:	Orange Peel Oil, Bitter (<i>Citrus aural</i> Bitter orange oil (<i>Citrus aurantium</i> L Citrus aurantium peel oil Curacao peel oil (<i>Citrus aurantium</i> L Daidai peel oil (<i>Citrus aurantium</i> L.)	<i>ntium</i> L. subsp amara L.) . subsp. amara L.))	

History:	Initial reviews:	October 1975, June 1992, July 2002	
	Current revision date:	2015	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	1.25%	Rinse-off products:	No Restriction		
		Including household	cleaning products		
Non skin contact products:	No Restriction				
Note box:					
The Standard is set due to the phototoxic effect off products (please refer to Table 4 of the QRA	s of the material. The lin booklet for more detaile	nit only applies to applications or ad information).	n skin exposed to sunshine, excluding rinse-		
If combinations of phototoxic fragrance ingredients are used, the use levels have to be reduced accordingly. The sum of the concentrations of all phototoxic fragrance ingredients, expressed in % of their recommended maximum level in the consumer product, shall not exceed 100.					
Note: See remark on phototoxic ingredients in the Introduction to the IFRA Standards (Appendix 8 to the IFRA Code of Practice) and the Standard on Citrus oil and other furocoumarins-containing essential oils.					
Fragrance material specificatio	ns:	N/A			
Note: See remark on phototoxic ingredients in the Introduction to the IFRA Standards (Appendix 8 to the IFRA Code of Practice) and the Standard on Citrus oil and other furocoumarins-containing essential oils. Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on the contributions from other sources in the Introduction to the IFRA Standards).

Bitter orange peel oil expressed

CRITICAL EFFECT:

PHOTOTOXICITY

RIFM SUMMARIES:

Human Studies: The material was tested for phototoxic potential in human volunteers (Kaidbey and Kligman, 1980). Five µL/cm² of 100% bitter orange oil was applied to 2 cm² under occlusive tape. One cm circular sites were exposed to visible light or 20 J/ cm² UVA.

Reactions were read at 24 and 48 hours. All 8 subjects reacted.

Animal studies: The NOEL was based on studies conducted with pooled samples of bitter orange oil in one miniature swine and hairless mice, which showed NOEL of 6.25%.

		Human Data			
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
N/A	N/A	N/A	N/A	N/A	N/A

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No Observed Effect Level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = Lowest Observed Effect Level; NA = Not Available.

¹Data derived from HRIPT or HMT.

²Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

³WoE NESIL limited to two significant figures.

⁴EC3 value from one LLNA, not the mean.

⁵LOEL from human maximization test, not a human repeated insult patch test.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for orange peel oil, bitter, and recommended that the skin contact level should change to 1.25%, incorporating a 5 fold uncertainty factor.

REFERENCES:

P.D. Forbes, F. Urbach and R.E. Davies (1977). Phototoxicity testing of fragrance raw materials. Food and Cosmetics Toxicology, 15, 55-60. Report number 1422.

Kaidbey, K.H. and Kligman, A.M. (1980). Identification of contact photosensitizers by human assay. Current Concepts in Cutaneous Toxicity, 55-68. Academic Press, NY. Report number 1995.

Research Institute for Fragrance Materials, Inc. (1972). Phototoxicity and irritation studies of fragrance materials in hairless mice and miniature swine. RIFM report number 2034, May 26.

Research Institute for Fragrance Materials, Inc. (1978). Phototoxicity and irritation studies of mice and pigs with fragrance materials. RIFM report number 2042, April 14.

Boldo oil

CAS N°:	8022-81-9	Empirical formula:	N/A
Synonyms:	Boldo leaf oil (Peumus boldus Mol.) Oil, boldo leaf Peumus boldus oil		

History:	Initial reviews:	New Standard	
	Current revision date:	2009	
	Implementation date:	For new submissions*:	August 7, 2009
		For existing fragrance compounds*:	August 7, 2010
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

INSUFFICIENT DATA

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use.

This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Regulatory Toxicology and Pharmacology, 31, 166-181, 2000



3-Bromo-1,7,7-trimethylbicyclo[2.2.1]heptane-2-one

CAS N°:	76-29-9	Empirical formula: Structure:	C10H15BrO
Synonyms:	Bicyclo[2.2.1]heptan-2-one, 3-b 2-Bornanone, 3-bromo- 3-Bromobornan-2-one 3-Bromo-2-bornanone 3-Bromocamphor Camphor bromide Camphor, 3-bromo-	promo-1,7,7-trimethyl-	

History:	Initial reviews:	Not applicable		
	Current revision date:	Not applicable		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

N/A



3-Bromo-1,7,7-trimethylbicyclo[2.2.1]heptane-2-one

CRITICAL EFFECT:

SEE NOTE BOX

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or

2) adverse data on the material itself and/or

3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000



CAS N°:	103-64-0	Empirical formula: Structure:	C ₈ H ₇ Br
Synonyms:	Benzene, (2-bromoethenyl)- α-Bromo-β-phenylethylene β-Bromostyrene β-Bromovinylbenzene Omega-Bromstyrene Bromstyrol Bromstyrolene		

History:	Initial reviews:	Not applicable		
	Current revision date:	Not applicable		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



Bromostyrene

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000

alpha-Butylcinnamaldehyde

CAS N°:	7492-44-6	Empirical formula: Strucutre:	C ₁₃ H ₁₆ O
Synonyms:	2-Benzylidenehexanal Butyl cinnamic aldehyde α-Butyl-β-phenylacrolein Hexanal, 2-(phenylmethyler alpha-butylcinnamaldehyde	ne)-	

History:	Initial reviews:	New Standard		
	Current revision date:	June 20, 2011		
	Implementation date:	For new submissions*: August 20, 2011		
		For existing fragrance compounds*:	August 20, 2012	
	Next review date	2016		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
0.03%	Category 7	0.08%			
0.04%	Category 8	1.01%			
0.15%	Category 9	5.00%			
0.45%	Category 10	2.50%			
0.24%	Category 11	See Note Box (2)			
0.72%					
Note box:					
 See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products with the potential of ingestion. Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. 					
	0.03% 0.04% 0.15% 0.45% 0.24% 0.72%	0.03% Category 7 0.04% Category 8 0.15% Category 9 0.45% Category 10 0.24% Category 11 0.72% Category 11			

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None known at the time of the publication of the Standard

CRITICAL EFFECT:

SENSITIZATION

alpha-Butylcinnamaldehyde

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
2775 [2]	Weak	NA	5520	NA	1000

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003. ²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures. A default value based on the LLNA data was employed because the material is used a very low volume and there are no HRIPT data.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for α -butylcinnamaldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1000 µg/cm², which is a default value based on the LLNA data. They recommend the limits for the 11 different product categories, which are the acceptable use levels of α -butylcinnamaldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick G, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.



3-(m-tert-Butylphenyl)-2-methylpropionaldehyde (m-BMHCA)

CAS N°:	62518-65-4	Empirical formula:	C ₁₄ H ₂₀ O
		Structure:	H ₃ C CH ₃
Synonyms:	Benzenepropanal, 3-(3-(3-tert-Butylphenyl)- m-BMHCA	1,1-dimethylethyl)-α-methyl- 2-methylpropanal	

History:	Initial reviews:	New Standard	
	Current revision date:	2015	
	Implementation date:	For new submissions*:	August 10, 2015
		For existing fragrance compounds*:	August 10, 2016
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.12 %	Category 7	0.31 %		
Category 2	0.15 %	Category 8	2.00 %		
Category 3	0.62 %	Category 9	5.00 %		
Category 4	1.86 %	Category 10	2.50 %		
Category 5	0.98 %	Category 11	See Note box (2)		
Category 6	2.97 %				
Note box:					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organization of the Flavor Industry - www.iofi.org).					

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

3-(m-tert-Butylphenyl)-2-methylpropionaldehyde (m-BMHCA)

CRITICAL EFFECT:

DERMAL SENSITIZATION

RIFM SUMMARIES:

LLNA weighted mean EC3 values			Human Data		
(μg/cm²) [no. studies] for p-BMHCA	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
2376 [6]4	Weak	4125	NA	29528	4100

All data in this table are for p-BMHCA and and are listed in the RIFM Database and available from RIFM.

NOEL = No Observed Effect Level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test;

LOEL = Lowest Observed Effect Level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003. ²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴EC3 value from the mean of the individual LLNAs

REXPAN RATIONALE / CONCLUSION:

The Standard for 3-(m-tert-Butylphenyl)-2-methylpropionaldehyde (m-BMHCA) is based on a concern for sensitization potential due to the presence of structural alerts, the fact that structurally similar materials are sensitizers, and the lack of sensitization test data on this material.

The RIFM Expert Panel recommends the limits for the 11 different product categories, which are the acceptable use levels of 3-(m-tert-Butylphenyl)-2-methylpropionaldehyde (m-BMHCA) in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008, based on a No Expected Sensitization Induction Level (NESIL) of 4100 µg/cm² derived from the data on p-BMHCA as provided above.

REFERENCES:

Api, A. M., Basketter, D. A., Cadby, P. A., Cano, M-F., Ellis, G., Gerberick, G. F. et al, 2008. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. Regulatory Toxicology and Pharmacology 52(1): 3-23.

Gerberick, G.F. et al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161

Kimber, I., Ryan, C. A., Gerberick, G. F., White, I. R., 2001. Human potency predictions for aldehydes using the local lymph node assay. Contact Dermatitis, 45(2), 89-94.

RIFM (Research Institute for Fragrance Materials, Inc.), 1980. Repeated Insult Patch Test on BMHCA. Unpublished report from IFF, Inc., 14 February. Report number 15029 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1999. Repeated Insult Patch Test on BMHCA. RIFM report number 34405, May 4. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001a. Local Lymph Node Assay on BMHCA. RIFM report number 37065, March 12. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001b. Local Lymph Node Assay on BMHCA. RIFM report number 37066, May 9. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001c. Local Lymph Node Assay on BMHCA. RIFM report number 37068, May 9. (RIFM, Woodcliff Lake, NJ, USA).



3-(m-tert-Butylphenyl)-2-methylpropionaldehyde (m-BMHCA)

RIFM (Research Institute for Fragrance Materials, Inc.), 2001d. Local Lymph Node Assay on BMHCA. RIFM report number 37067, May 9. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001e. Local Lymph Node Assay on BMHCA. RIFM report number 41235, September 27. (RIFM, Woodcliff Lake, NJ, USA).



p-tert-Butyldihydrocinnamaldehyde (Bourgeonal)



History:	Initial reviews:	April 1991, July 1994, May 2007 (42nd Amendment)		
	Current revision date:	2008		
	Implementation date:	For new submissions*:	August 16, 2008	
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.03 %	Category 7	0.1 %		
Category 2	0.04 %	Category 8	0.6 %		
Category 3	0.2 %	Category 9	0.6 %		
Category 4	0.5 %	Category 10	0.6 %		
Category 5	0.3 %	Category 11	Not Restricted (2)		
Category 6	0.8 %				
Note box:					
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again. (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.					
Fragrance material specificatio	ns:	N/A			

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

p-tert-Butyldihydrocinnamaldehyde (Bourgeonal)

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

p-tert-Butyl-dihydrocinnamaldehyde - Sensitization Potency Estimation Based on Weight of Evidence

			Human Data		
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
1075 [1]	Weak	1181	4138	7087	1100

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Data derived from HRIPT or HMT.

²Gerberick et al., 2001

³WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for p-tert-butyldihydrocinnamaldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1100 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of p-tertbutyldihydrocinnamaldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161

RIFM (Research Institute for Fragrance Materials, Inc.), 1980. Maximization test on p-tert- Butyldihydrocinnamaldehyde. RIFM report number 1790, June 25a. (RIFM, Woodcliff Lake, NJ, USA).

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 2002. Repeated Insult Patch Test on p-tert-Butyldihydrocinnamaldehyde. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003. Repeated Insult Patch Test on p-tert-Butyldihydrocinnamaldehyde. RIFM report number 47269, March 21. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003. Repeated Insult Patch Test on p-tert-Butyldihydrocinnamaldehyde. RIFM report number 47272, February 10. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Repeated Insult Patch Test on p-tert-Butyldihydrocinnamaldehyde. RIFM report number 45134, March 11. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2006. Local Lymph Node Assay on p-tert-Butyldihydrocinnamaldehyde. DRAFT. (RIFM, Woodcliff Lake, NJ, USA).



CAS N°:	80-54-6	Empirical formula: Structure:	C ₁₄ H ₂₀ O
Synonyms:	Benzenepropanal, 4-(1,1-dir p-t-Bucinal 2-(4-tert-Butylbenzyl)propion p-t-Butyl-alpha-methylhydrod Butylphenyl methylpropional alpha-Methyl-ß-(p-t-butylphe Lilestralis, Lilial, Lysmeral (c	nethylethyl)-alpha-methyl- naldehyde cinnamaldehyde enyl)propionaldehyde ommercial names)	

p-tert-Butyl-alpha-methylhydrocinnamic aldehyde (p-BMHCA)

History:	Initial reviews:	April 2003, May 2007, October 2008, June 2013		
	Current revision date:	2015		
	Implementation date:	For new submissions*:	August 10, 2015	
	-	For existing fragrance compounds*:	August 10, 2016	
	Next review date	2020		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.12 %	Category 7	0.31 %		
Category 2	0.15 %	Category 8	2.00 %		
Category 3	0.62 %	Category 9	5.00 %		
Category 4	1.86 %	Category 10	2.50 %		
Category 5	0.98 %	Category 11	See Note box (2)		
Category 6	2.97 %				
Note box:					
(1) IFRA would recommend that any material uningredients that are in compliance with appropriates are lacking, with the recommendations la www.iofi.org).	sed to impart perfume or iate regulations for foods id down in the Code of F	flavour in products intended for human ir and food flavourings in the countries of p ractice of IOFI (International Organization	ngestion should consist of Janned distribution and, where n of the Flavor Industry -		
(2) Category 11 includes all non-skin contact or there is no justification for a restriction of the co	incidental skin contact p ncentration of this fragra	products. Due to the negligible skin contact not not not not not not not not not no	t from these types of products		
Fragrance material specifications: N/A					
CONTRIBUTION FROM OTHER SOURCES:					
See Annex II.					

p-tert-Butyl-alpha-methylhydrocinnamic aldehyde (p-BMHCA)

CRITICAL EFFECT:

DERMAL SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
2372 [6]	Weak	4125	NA	29528	4100

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No Observed Effect Level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = Lowest Observed Effect Level; NA = Not Available

¹Data derived from HRIPT or Human Max tests ²Gerberick et al., 2001

³WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for p-tert-Butyl-alphamethylhydrocinnamic aldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 4100 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of p-tert-Butylalpha-methylhydrocinnamic aldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api et al., 2008.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161

Kimber, I., Ryan, C. A., Gerberick, G. F., White, I. R., 2001. Human potency predictions for aldehydes using the local lymph node assay. Contact Dermatitis, 45(2), 89-94.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 1980. Repeated Insult Patch Test on BMHCA. Unpublished report from IFF, Inc., 14 February. Report number 15029 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1999. Repeated Insult Patch Test on BMHCA. RIFM report number 34405, May 4. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001a. Local Lymph Node Assay on BMHCA. RIFM report number 37065, March 12. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001b. Local Lymph Node Assay on BMHCA. RIFM report number 37066, May 9. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001c. Local Lymph Node Assay on BMHCA. RIFM report number 37068, May 9. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001d. Local Lymph Node Assay on BMHCA. RIFM report number 37067, May 9. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001e. Local Lymph Node Assay on BMHCA. RIFM report number 41235, September 27. (RIFM, Woodcliff Lake, NJ, USA).

p-tert-Butylphenol

CAS N°:	98-54-4	Empirical formula: Structure:	C ₁₀ H ₁₄ O
Synonyms:	4-tert-Butylphenol 4-(1,1-Dimethylethyl) phenol 1-Hydroxy-4-tert-butylbenzene Phenol, 4-(1,1-dimethylethyl)- Phenol, p-tert-butyl		

History:	Initial reviews:	June 1975	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION, DEPIGMENTATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke, D.L.J. (1975), Food and Chemical Toxicology 12, 835.

Cade oil

CAS N°:	8013-10-3 90046-02-9	Empirical formula:	N/A
Synonyms:	For the crude material banned: Juniper tar (CAS) For the distillates specified: Juniper tar oil Juniperus oxycedrus oil		

History:	Initial reviews:	July 1990, October 2003	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

SPECIFICATION / PROHIBITED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	N/A	Rinse-off products:	N/A	
		Including household	cleaning products	
Non skin contact products:	N/A			
Note box:				
Crude cade oil derived by pyrolysis of the wood	and twigs of Juniperus	oxycedrus L. should not be used	as a fragrance ingredient.	
Only rectified (purified) cade oils being in comp	liance with the maximum	limit for polynuclear aromatic h	ydrocarbons should be used.	
Fragrance material specifications: Limit content of polynuclear aromatic hydrocarbons (PAH) resulting from the use of rectified oils according to Good Manufacturing Practice.				
Benzopyrene and 1,2-Benzanthracene are to be used as markers fo <u>PAH. If used alone or in combination with rectified Birch tar oils,</u> <u>rectified Opoponax oil or rectified Styrax oil, the total concentration</u> <u>of both of the markers should not exceed 1 ppb in the final product</u>				

CONTRIBUTION FROM OTHER SOURCES:

N/A

Cade oil

CRITICAL EFFECT:

CARGINOGENICITY, GENOTOXICITY*

*Some of the polynuclear aromatic hydrocarbons are known to be carcinogen or genotoxic materials.

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, October 15, 2003

CAS N°:	Carvone: 99-49-0 d-Carvone: 2244-16-8 I-Carvone: 6485-40-1	Empirical formula: Structure:	C ₁₀ H ₁₄ O Carvone:
Synonyms:	2-Cyclohexen-1-one, 2-meth 6,8(9)-p-Menthadien-2-one p-Mentha-6,8-dien-2-one 1-Methyl-4-isopropenyl-6-cy	nyl-5-(1-methylethenyl)- clohexen-2-one	

Carvone

History:	Initial reviews:	New Standard	
	Current revision date:	2008	
	Implementation date:	For new submissions*: August 16, 2008	
		For existing fragrance compounds*:	August 16, 2010
	Next review date	2013	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Category 1 See Note box (1)	0.08 %	Category 7	0.2 %	
Category 2	0.1 %	Category 8	2.0 %	
Category 3	0.4 %	Category 9	5.0 %	
Category 4	1.2 %	Category 10	2.5 %	
Category 5	0.6 %	Category 11	Not Restricted (2)	
Category 6	1.9 %			
Note box:				
 IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. 				
Fragrance material specification	ons:	N/A		

Carvone

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (µg/cm ²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
2675 [1] ⁵	Weak	2657 ⁴	1379 ⁴	NA	2650

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for carvone aldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 2650 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of carvone aldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1976 . Human Maximization Test RIFM report number 1797, April 9. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007a. Local Lymph Node Assay. RIFM report number 52902, May 10. (RIFM, Woodcliff Lake, NJ. USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007b. Human Repeated Insult Patch Test. RIFM report number 52896, May 15. (RIFM, Woodcliff Lake, NJ, USA).

Carvone oxide

CAS N°:	33204-74-9	Empirical formula: Strucutre:	
Synonyms:	Carvone epoxide 1,6-Epoxy-p-menth-8-en-2-one 1-Methyl-4-(1-methylvinyl)-7-oxabicyclo[4. 7-Oxabicyclo[4.1.0]heptan-2-one, 1-methy	1.0]heptan-2-one /l-4-(1-methylethenyl)	-

History:	Initial reviews:	New Standard**	
	Current revision date:	October 2003	
	Implementation date:	For new submissions*: Not applicable**	
		For existing fragrance compounds*:	Not applicable**
	Next review date	Not applicable**	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	N/A	Rinse-off products:	N/A	
		Including household	cleaning products	
Non skin contact products:	N/A			
Note box:				
This material should not be used as fragrance ingredient for any application.				
**This material has previously been included in the list of 'Other Materials', the material therefore has already been prohibited before.				

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None to consider.

CRITICAL EFFECT:

SENSITIZATION


Carvone oxide

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, October 2003

REFERENCES:

Letizia et al., 2000, Food and Chemical Toxicology, Volume 38, Supplement 3, Special Issue IX, pages S25-26.



CAS N°:	8006-99-3	Empirical formula:	N/A
Synonyms:	American wormseed oil Chenopodium ambrosioides L. var anthe	elminticum	

History:	Initial reviews:	Not applicable		
	Current revision date:	Not applicable		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
	Including household cleaning products				
Non skin contact products:	N/A				
Note box:					
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.					
Fragrance material specificatio	ns:	N/A			

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



Chenopodium oil

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000

Cinnamic alcohol



History:	Initial reviews:	October 1987, June 1992, September 2002, May 2007 (4: Amendment)		
	Current revision date:	2008		
	Implementation date: For new submissions*:		August 16, 2008	
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.09 %	Category 7	0.2 %			
Category 2	0.1 %	Category 8	0.4 %			
Category 3	0.4 %	Category 9	0.4 %			
Category 4	0.4 %	Category 10	0.4 %			
Category 5	0.4 %	Category 11	Not Restricted (2)			
Category 6	2.2 %					
Note box:						
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again. (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. There is also a separate Standard for Styrax that should be taken into account.						
Fragrance material specificatio	Fragrance material specifications: N/A					

Cinnamic alcohol

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

Cinnamic alcohol - Sensitization Potency Estimation Based on Weight of Evidence

		Human Data			
LLNA weighted mean EC3 values (µg/cm ²) [no. studies] Based on Animal Data ²		NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
5250 [1]	Weak	3000	2759	4724	3000

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Data derived from HRIPT or HMT.

²Gerberick et al., 2001

³WoE NESIL limited to three significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for cinnamic alcohol and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 3000 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of cinnamic alcohol in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Basketter, D.A., Wright, Z., Gilmour, N.J., Ryan, C.A., Gerberick, G.F., Robinson, M.K., Dearman, R.J., Kimber, I., 2002. Prediction of human sensitization potency using local lymph node assay EC3 values. *The Toxicologist*, 66(1-S), 240.

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. *American Journal of Contact Dermatitis*, 12(3), 156-161

Greif, N., 1967. Cutaneous safety of fragrance materials as measured by the maximization test. American Perfumer and Cosmetics, 82, 54.

RIFM (Research Institute for Fragrance Materials, Inc.), 1979. Human Maximization Test on Cinnamyl Alcohol. RIFM report number 1697, July 6a. (RIFM, Woodcliff Lake, NJ, USA).

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 2001a. Repeated Insult Patch Test on Cinnamyl Alcohol. RIFM report number 40696, August 20. (RIFM, WoodcliffLake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001b. Repeated Insult Patch Test on Cinnamyl Alcohol. RIFM report number 40695, August 20. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2002. Repeated Insult Patch Test on Cinnamyl Alcohol. RIFM report number 40697, March 5. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Repeated Insult Patch Test on Cinnamyl Alcohol. RIFM report number 47241, December 16. (RIFM, Woodcliff Lake, NJ, USA).



Cinnamic aldehyde

CAS N°:	104-55-2	Empirical formula: Structure:	C₀H₀O
Synonyms:	Cinnamal Cinnamaldehyde Phenylacrolein 3-Phenyl-2-propenal 3-Phenyl-2-propen-1-al 'Cassia aldehyde'		

History:	Initial reviews:	March 1978, April 2004, May 2006, May 2007, June 2008		
	Current revision date:	June 2013		
	Implementation date:	For new submissions*:	August 10, 2013	
		For existing fragrance compounds*:	August 10, 2014	
	Next review date	2018		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.02 %	Category 7	0.04 %		
Category 2	0.02 %	Category 8	0.05 %		
Category 3	0.05 %	Category 9	0.05 %		
Category 4	0.05 %	Category 10	0.05 %		
Category 5	0.05 %	Category 11	Not Restricted (2)		
Category 6	0.4 %		'		
Note hov:					

For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again.

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org)

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

The Standard on cinnamic aldehyde covers and replaces the former existing Standards on cassia oil, cinnamon bark oil as well as cinnamic aldehyde – methyl anthranilate schiff base. The existing Standards for cinnamic aldehyde, cassia oil, cinnamic aldehyde – methyl anthranilate schiff base and cinnamon bark oil are no longer valid.

47th Amendment



Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

See Annex I and Annex II

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data				
LLNA weighted mean EC3 values (µg/cm ²) [no. studies] Based on Animal Data ²		NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)	
262 [23]	Moderate	591 ⁴	NA	775	590	

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested,

not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for cinnamic aldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 590 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of cinnamic aldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Basketter, D.A., Wright, Z.M., Warbrick, E.V. Dearman, R.J., Kimber, I., Ryan, C.A., Gerberick, G.F., White, I.R., 2001. Human potency predictions for aldehydes using the local lymph node assay. *Contact Dermatitis*, 45(2), 89-94.

Basketter, D.A., Wright, Z., Gilmour, N.J., Ryan, C.A., Gerberick, G.F., Robinson, M.K., Dearman, R.J., Kimber, I., 2002. Prediction of human sensitization potency using local lymph node assay EC3 values. *The Toxicologist*, 66(1-S), 240.

Elahi, E.N., Wright, Z., Hotchkiss, S.A.M., Basketter, D.A., Smith Pease, C.K., 2002. Protein binding and metabolic inhibition reveals clues on the mechanisms surrounding relative potency of sensitizing cinnamic compounds. Toxicology, 178, 52.

Elahi, E.N., Wright, Z., Hinselwood, D., Hotchkiss, S.A.M., Basketter D.A., Smith Pease , C.K., 2004. Protein binding and metabolism influence the relative skin sensitization potential of cinnamic compounds. *Chemical Research in Toxicology*, 17(3), 301-310.

RIFM (Research Institute for Fragrance Materials, Inc.), 1973. Repeated Insult Patch Test on Cinnamaldehyde. RIFM report number 12509, January 23. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2002a. Repeated Insult Patch Test on Cinnamaldehyde. RIFM report number 41692, August 27. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2002b. Repeated Insult Patch Test on Cinnamaldehyde. RIFM report number 41693, August 27. (RIFM, Woodcliff Lake, NJ, USA).

Cinnamic aldehyde

RIFM (Research Institute for Fragrance Materials, Inc.), 2003a. Local Lymph Node Assay on Cinnamaldehyde. RIFM report number 42032, February 18. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003b. Local Lymph Node Assay on Cinnamaldehyde. RIFM report number 42033, February 18. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003c. Local Lymph Node Assay on Cinnamaldehyde. RIFM report number 42040, February 18. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003d. Local Lymph Node Assay on Cinnamaldehyde. RIFM report number 42034, February 18. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003e. Local Lymph Node Assay on Cinnamaldehyde. RIFM report number 42036, February 18. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003f. Local Lymph Node Assay on Cinnamaldehyde. RIFM report number 42035, February 18. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003g. Local Lymph Node Assay on Cinnamaldehyde. RIFM report number 42037, February 18. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003h. Local Lymph Node Assay on Cinnamaldehyde. RIFM report number 42038, February 18. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003i. Local Lymph Node Assay on Cinnamaldehyde. RIFM report number 42039, February 18. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003j. Local Lymph Node Assay on Cinnamaldehyde. RIFM report number 42041, February 18. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Repeated Insult Patch Test on Cinnamaldehyde. RIFM report number 47158, April 22a. (RIFM, Woodcliff Lake, NJ, USA).

Smith, C.K., Hotchkiss, S.A.M., 2001. Allergic Contact Dermatitis. Chemical and Metabolic Mechanisms. Taylor and Francis, London.

Wright, Z.M., Basketter, D.A., Blaikie, L., Cooper, K.J., Warbrick, E.V., Dearman, R.J., Kimber, I., 2001. Vehicle effects on skin sensitization potency of four chemicals

assessment using the local lymph node assay. *Journal International Journal of Cosmetic Science*, 23(2), 75-83.

Cinnamic Aldehyde Dimethyl Acetal

CAS N°: 4364-06-1 Empirical formula: Structure: C11H14O2 Synonyms: Benzene, (3,3-dimethoxy-1-propenyl)-Cinnamic aldehyde dimethyl acetal (3,3-Dimethoxypropen-1-yl)benzene (3,3-Dimethoxyprop-1-en-1-yl)benzene 3-Phenyl-2-propenal dimethyl acetal Synonyms

History:	Initial reviews:	New Standard		
	Current revision date:	2009		
	Implementation date:	For new submissions*: August 7, 2009		
		For existing fragrance compounds*:	August 7, 2011	
	Next review date	2014		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.02 %	Category 7	0.06 %			
Category 2	0.03 %	Category 8	0.80 %			
Category 3	0.12 %	Category 9	4.10 %			
Category 4	0.37 %	Category 10	2.50 %			
Category 5	0.20 %	Category 11	See Note Box (2)			
Category 6	0.59 %					
Note box:						
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. 						
Fragrance material specifications: N/A						

Cinnamic Aldehyde Dimethyl Acetal

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		l l			
LLNA weighted mean EC3 values (µg/cm ²) [no. studies] Based on	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
NA	NA	827	NA	1938	820

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or HMT

² Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

³WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for cinnamic aldehyde dimethyl acetal and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 820 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of cinnamic aldehyde dimethyl acetal in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1965. Human repeated insult patch test. Unpublished study from IFF Inc., 7 September. Report number 48396. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2008. Human repeated insult patch test. RIFM report number 55346, July 30. (RIFM, Woodcliff Lake, NJ, USA).

Cinnamylidene acetone



History:	Initial reviews:	Not applicable	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
	Including household cleaning products				
Non skin contact products:	N/A				
Note box:					
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

N/A

IFRA STANDARD

Cinnamylidene acetone

CRITICAL EFFECT:

SEE NOTE BOX

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000



Cinnamyl nitrile



History:	Initial reviews:	March 2002 (36th Amendment)	
	Current revision date:	2008	
	Implementation date:	For new submissions*:	August 16, 2008
		For existing fragrance compounds*:	August 16, 2010
	Next review date	2013	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.03 %	Category 7	0.08 %		
Category 2	0.04 %	Category 8	0.125 %		
Category 3	0.125 %	Category 9	0.125 %		
Category 4	0.125 %	Category 10	0.125 %		
Category 5	0.125 %	Category 11	See Note Box (2)		
Category 6	0.80 %				
Note box:					
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the					

position will be reevaluated again.

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) (http://www.iofiorg.org/).

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

Cinnamyl nitrile

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		1	Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)	
>2500	Weak	1063 ⁴	3448 ⁴	1938	1060	

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

²Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Cinnamyl nitrile and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1060 μ g/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Cinnamyl nitrile in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1965. Human Repeated Insult Patch Test. Unpublished report from IFF International, 1 April. Report number 1981 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1981. Human Maximization Test. RIFM report number 1792, October 27 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Local Lymph Node Assay. RIFM report number 51626, January 28 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007. Human Repeated Insult Patch Test. RIFM report number 54076, December 13 (RIFM, Woodcliff Lake, NJ, USA).

Citral



History:	Initial reviews:	March 2002 (36 th Amendment), June 2008		
	Current revision date:	June 2013		
	Implementation date:	For new submissions*: August 10, 2013		
		For existing fragrance compounds*:	August 10, 2014	
	Next review date	2018		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Category 1	0.04 %	Category 7	0.1 %	
Category 2	0.05 %	Category 8	1.4 %	
Category 3	0.2 %	Category 9	5.0 %	
Category 4	0.6 %	Category 10	2.5 %	
Category 5	0.3 %	Category 11	See Note Box	
Category 6	1.0 %			
Note box:				
Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.				
This Standard cancels and replaces the existing one on citral, which was based on the no longer supported 'quenching' phenomenon.				
Fragrance material specifications: N/A				

CONTRIBUTION FROM OTHER SOURCES:

See Annex I and Annex II

Citral

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

Citral - Sensitization Potency Estimation Based on Weight of Evidence

			Human Data		
LLNA weighted mean EC3 values (µg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
1414 [11]	Weak	1400	NA	3876	1400 μg/cm ²

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max Tests

² Gerberick et al., 2001

³ WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for citral and based on the weight of evidence established the No Expected Sensitization Induction Level (NESIL) as 1400 μ g/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of citral in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of March 15, 2006.

REFERENCES:

Basketter, D. A., Wright, Z., Gilmour, N. J., Ryan, C. A., Gerberick, G. F., Robinson, M. K., Dearman, R. J., Kimber, I., 2002a. Prediction of human sensitization potency using local lymph node assay EC3 values. The Toxicologist, 66(1-S), 240.

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

Research Institute for Fragrance Materials, Inc (1964). Repeated insult patch test of citral in human subjects. Unpublished report from International Flavors and Fragrances Inc., Report number 14576 (RIFM, Woodcliff Lake, NJ USA). Research Institute for Fragrance Materials, Inc (2004). Repeated insult patch test in human subjects with citral. RIFM report number 47157 (RIFM, Woodcliff Lake, NJ USA).

Research Institute for Fragrance Materials, Inc (2004). Local Lymph Node Assay on Citral. RIFM report number 45126 (RIFM, Woodcliff Lake, NJ USA).

		Citronelloi		
CAS N°:	106-22-9 1117-61-9 26489-01-0 6812-78-8 141-25-3 68916-43-8 7540-51-4	Empirical formula: Structure:	C ₁₀ H ₂₀ O (+)-Citronellol :	
Synonyms:	106-22-9 Citronellol dl-Citronellol Dihydrogeraniol 3,7-Dimethyl-6-octen-1-ol 6-Octen-1-ol, $3,7$ -dimethyl- 1117-61-9 (+)- β -Citronellol (+)- (R) -Citronellol (R)- $3,7$ -Dimethyloct-6-en-1-ol 6-Octen-1-ol, $3,7$ -dimethyl-, (I 26489-01-0 dl-Citronellol 6812-78-8 α -Citronellol 141-25-3 3,7-Dimethyl-(6-or 7-)octen-1- 3,7-Dimethyl-7-octen-1-ol 7-Octen-1-ol, $3,7$ -dimethyl-, (S 7-Octen-1-ol, $3,7$ -dimethyl-, (S Rhodinol 68916-43-8 Geranium oil, saponified Rhodinol 7540-51-4 I-Citronellol (-)- $3,7$ -Dimethyl-6-octen-1-ol (S)- $3,7$ -Dimethyl-6-octen-1-ol 6-Octen-1-ol, $3,7$ -dimethyl-, (S	R)- -Ol S)- somer unspecified) S)-		

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History:	Initial reviews:	New Standard	
	Current revision date:	2007	
	Implementation date:	For new submissions*: June 16, 2007	
		For existing fragrance compounds*:	June 16, 2009
	Next review date	2012	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.8 %	Category 7	2.2 %		
Category 2	1.1 %	Category 8	2.0 %		
Category 3	4.4 %	Category 9	5.0 %		
Category 4	13.3 %	Category 10	2.5 %		
Category 5	7.0 %	Category 11	See Note Box (2)		
Category 6	21.4 %				

Note box:

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry). Further information about IOFI can be found on its website (www.iofiorg.org).

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

Citronellol

RIFM SUMMARIES:

dl-Citronellol - Sensitization Potency Estimation Based on Weight of Evidence

		l l	Human Data		
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
10875 [1]	Extremely weak	29528	4138	NA	29500
				-	

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max Tests

² Gerberick et al., 2001

³ WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for dl-citronellol and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 29500 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of dl-citronellol in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

Greif, N., 1967. Cutaneous safety of fragrance materials as measured by the maximization test. American Perfumer and Cosmetics, 82, 54.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 2005a. Repeated Insult Patch Test on dl-Citronellol. RIFM report number 47277, January 28. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005b. Local Lymph Node Assay on dl-Citronellol. RIFM report number 48752, January 6. (RIFM, Woodcliff Lake, NJ, USA).

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Citrus oils and other furocoumarins containing essential oils

History:	Initial reviews:	December 1996	
	Current revision date:	2015	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:			
Skin contact products:			
Leave-on products:	15 ppm 5-MOP	Rinse-off products:	No Restriction
		Including household	cleaning products
Non skin contact products:	No Restriction		
Note box:			
The Standard is set due to the phototoxic effect off products (please refer to Table 4 of the QRA	s of the material. The lin booklet for more detaile	nit only applies to applications or ad information).	n skin exposed to sunshine, excluding rinse-
If combinations of phototoxic fragrance ingreduphototoxic ingredients, expressed in % of their in	ents are used, the use I recommended maximum	evels have to be reduced accor I level in the consumer product s	dingly. The sum of the concentrations of all shall not exceed 100.
Note: See remark on phototoxic ingredients in t	he Introduction to the I	FRA Standards (Appendix 8 to	o the IFRA Code of Practice).
Fragrance material specificatio	ns:	Where the bergapten (5-methopresent in a compound has be applications on areas of skin e preparations, soaps and other total level of bergapten in the 0.0015% (15 ppm). This is equ compound used at 20% in the Where the level of bergapten I methods, the limits specified in apply. In those cases, where such oil phototoxic ingredients, the add consideration and the use leve sum of the concentrations of a expressed in % of their recom product, shall not exceed 100. Restrictions for furocoumaring Bitter orange oil expres Bitter orange oil expressed Cumin oil, Grapefruit oil expressed Lemon oil cold pre Lime oil expressed Rue oil.	oxypsoralen) content of all relevant oils seen determined, it is recommended that for exposed to sunshine, excluding bath products which are washed off the skin, the consumer products should not exceed uivalent to 0.0075% (75 ppm) in a fragrance consumer product. has not been determined by appropriate in the guidelines on individual oils should ls are used in combination with other ditive effect has to be taken into els have to be reduced accordingly. The ull phototoxic fragrance ingredients, mended maximum level in the consumer containing essential oils have been essed, spressed, d, ntain small amounts of phototoxic re not bich enough to require special

Citrus oils and other furocoumarins containing essential oils

restrictions if used alone, but if used in combination with one or the other phototoxic essential oil, attention should be paid that the total level of bergapten (5-MOP) in the consumer product does not exceed 15 ppm. It is the responsibility of fragrance manufacturers to ensure that the level is observed.

Typical levels of 5-MOP are the following: Petitgrain Mandarin oil - 50 ppm, Tangerine oil cold pressed - 50 ppm, Mandarin oil cold pressed - 250 ppm, Parsley leaf oil - 20 ppm.

CRITICAL EFFECT:

PHOTOTOXICITY

REFERENCES:

These recommendations are based on the published phototoxic effects of bergapten and the established dose-effect relationships (Young at al., J. Photochem. Photobiol. B,7, 231 (1990); Dubertret et al.ibid 7, 251 (1990), idem, ibid, 7, 362 (1990).

Colophony

CAS N°:	8050-09-7	Empirical formula:	N/A
Synonyms:	Colophonium Rosin		

History:	Initial reviews:	December 1992	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Hausen. B.M. (1989), Contact Dermatitis (20), 41-50; 133-143; 295-301.

Costus root oil, absolute and concrete

CAS N°:	8023-88-9	Empirical formula:	N/A
Synonyms:	Costus root essential oil, absolute and c Oils, costus Saussurea lappa root oil Spiral flag oil	oncrete (<i>Saussurea lappa</i> (Clarke)

History:	Initial reviews:	October 1974, May 1998	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke D.L. (1974), Food and Cosmetics Toxicology 12, 867.

Mitchell J.C. and Epstein W.L (1974), Archives of Dermatology, 110, 871-872.

Foussereau, J., Muller J.C. and Benezra C. (1975), Contact Dermatitis, 1, 223-230.

Epstein, W.L., Reynolds G.W. and Rodriguez, E. (1980), Archives of Dermatology, 116, 59-60.

Cheminat, A., Benezra, C., Farral M.J. and Frechet, J.M.J. (1981), Canadian Journal of Chemistry, 59, 1405-1414.



Coumarin

CAS N°:	91-64-5	Empirical formula: Structure:	C9H6O2
Synonyms:	2H-1-Benzopyran-2-one 1,2-Benzopyrone cis-o-Coumaric acid lacto Coumarin Coumarinic anhydride 2-Oxo-1,2-benzopyran Tonka bean camphor	one	

History:	Initial reviews:	New Standard		
	Current revision date:	2008		
	Implementation date:	For new submissions*: August 16, 2008		
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.1 %	Category 7	0.3 %		
Category 2	0.13 %	Category 8	2.0 %		
Category 3	0.5 %	Category 9	5.0 %		
Category 4	1.6 %	Category 10	2.5 %		
Category 5	0.8 %	Category 11	See Note Box (2)		
Category 6	2.5 %				
Note box:					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/.					
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.					
Fragrance material specifications: N/A					

Coumarin

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
>12 500 [2]	Weak	3 543 ⁴	5517 ⁴	8 858	3 500

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Coumarin and, based on the

weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 3 500 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Coumarin in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Greif, N., 1967. Cutaneous safety of fragrance material as measured by the maximization test. American Perfumer and Cosmetics, 82, 54-57.

RIFM (Research Institute for Fragrance Materials, Inc.), 2003. Local Lymph Node Assay with coumarin. Unpublished report from Rhodia Services, 26a May. Report number 47072. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004a. Human Repeated Insult Patch Test with coumarin. RIFM report number 47161, March 10. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004b. Local Lymph Node Assay with coumarin. Unpublished report from Rhodia Services, 3a December. Report number 47073. (RIFM, Woodcliff Lake, NJ, USA). RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Human Repeated Insult Patch Tests with coumarin. RIFM report number 47274, January 6 and 6b. (RIFM, Woodcliff Lake, NJ, USA). $r \cap$

Cumin oil

CAS N°:	8014-13-9	Empirical formula: Structure :	N/A N/A
Synonyms:	Cumin seed oil Cuminum cyminum (Cumin) seed oil Cuminum cyminum L. Cuminum cyminum oil Oils, cumin (<i>Cuminum cyminum</i>)		

History:	Initial reviews:	October 1975, June 1986, September 2001		
	Current revision date:	2015		
	Implementation date:	For new submissions*:	Not applicable	
		For existing fragrance compounds*:	Not applicable	
	Next review date	2020		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	0.4%	Rinse-off products:	No Restriction	
		Including household	cleaning products	
Non skin contact products:	No Restriction			
Note box:				
The Standard is set due to the phototoxic effect off products (please refer to Table 4 of the QRA	s of the material. The lin booklet for more detaile	nit only applies to applications or ad information).	a skin exposed to sunshine, excluding rinse-	
If combinations of phototoxic fragrance ingredients are used, the use levels have to be reduced accordingly. The sum of the concentrations of all phototoxic fragrance ingredients, expressed in % of their recommended maximum level in the consumer product, shall not exceed 100.				
Note: See remark on phototoxic ingredients in the Introduction to the IFRA Standards (Appendix 8 to the IFRA Code of Practice) and the Standard on Citrus oil and other furocoumarins-containing essential oils.				
Fragrance material specifications: N/A				

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

Cumin oil

CRITICAL EFFECT:

PHOTOTOXICITY

RIFM SUMMARIES:

The NOEL for phototoxicity is 50% based on a study in 23 volunteers patched under occlusion on the back for 24 hours. Patches were removed after 10 minutes followed by irradiation with 16-20 J/cm² of UVA. Readings were made at 1, 24, 48 & 72 hours after irradiation. No photoirritation was observed (RIFM, 1986).

- 100% in miniature swine, UV, distinct photoirritant effects were observed (RIFM 1972; Forbes et al., 1977)
- 100% in hairless mice, UV, distinct photoirritant effects were observed (RIFM 1972; Forbes et al., 1977).
- 100% and 25% in hairless mice, UV, no reactions at 25% 0/12, 6/12 reactions at 100% (RIFM, 1983).
- 100%, 75%, 50%, and 25% in hairless mice, UV, no reactions 0/6 at 25%, 5/6 reactions at 50%, 6/6 reactions at 75% and 100% (RIFM, 1983).
- 30% in guinea pigs, UV, no reactions 0/10 (RIFM, 1984)
- 3% and 10% in guinea pigs, UV, no reactions 0/10 at 3%, and 4/10 reactions at 10% (RIFM, 1984).

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Cumin oil and has no concerns with the current limit of 0.4% (September 2001).

REFERENCES:

Research Institute for Fragrance Materials, Inc. (1986). Human phototoxicity study of cumin oil, tagetes minuta absolute, thyme concrete and pentyl acetate. RIFM report number 4348, 21 August.

Research Institute for Fragrance Materials, Inc. (1985). Cumin oil: A photoirritation test in humans. Unpublished report from the Givaudan-Roure Corp. Report number 3877, 7 January.

Research Institute for Fragrance Materials, Inc. (1972). Phototoxicity and irritation tests of fragrance materials in the hairless mice and miniature swine. Report number 2035, 26 July.

P.D.Forbes, F.Urbach and R.E.Davies. (1977). Phototoxicity testing of fragrance raw materials. Food and Cosmetics Toxicology, 15, 55-60. Report number 1422.

K.H.Kaidbey and A.M.Kligman (1978). Identification of topical photosensitizing agents in humans. Journal of Investigative Dermatology, 70(3), 149-151. Report number 3090.

Research Institute for Fragrance Materials, Inc. (1983). Phototoxicity study of fragrance materials in hairless mice. RIFM report number 2043, 31 January.

Research Institute for Fragrance Materials, Inc. (1984). Determination of phototoxicity of cumin oil in guinea pigs. Unpublished report from the Givaudan-Roure Corp. Report number 3875, 23 February.

Research Institute for Fragrance Materials, Inc. (1984). Determination of phototoxicity of cumin oil in guinea pigs. Unpublished report from the Givaudan-Roure Corp. Report number 3876, 17 July.



History:	Initial reviews:	New Standard	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.03 %	Category 7	0.08 %		
Category 2	0.04 %	Category 8	1.11 %		
Category 3	0.17 %	Category 9	5.00 %		
Category 4	0.50 %	Category 10	2.50 %		
Category 5	0.26 %	Category 11	See Note box (2)		
Category 6	0.80 %				
Note box:					
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. 					

Fragrance material specifications:

N/A

Cuminaldehyde

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
> 2500 [1] ⁴	Weak	1181	2760	NA	1100

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴EC3 value from one LLNA, not the mean.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Cuminaldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1100 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Cuminaldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick G, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. Regulatory Toxicology and Pharmacology 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 1972. Maximization test. RIFM report number 1804, November 22. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1975. Maximization test. RIFM report number 1804, March 27a. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1973. Maximization test. RIFM report number 1804, November 11. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. Repeat Insult Patch Test. Draft RIFM Report number 63810. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.) 2012. Local Lymph Node Assay. Draft RIFM Report number 63814. (RIFM, Woodcliff Lake, NJ, USA).



Cyclamen alcohol (3-(4-lsopropylphenyl)-2-methylpropanol)

CAS N°:	4756-19-8	Empirical formula: Structure:	C13H18O
Synonyms:	Benzenepropanol, .βmethyl-4-(1-met Cyclamen alcohol 3-(<i>p</i> -lsopropyl)phenyl-2-methyl-1-prop 3-(4-lsopropylphenyl)-2-methylpropan-	hylethyl)- anol ·1-ol	

History:	Initial reviews:	November 1977, October 1978	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED / SPECIFICATION

RESTRICTIONS:

Cyclamen alcohol should not be used as a fragrance ingredient as such, but a level of up to 1.5% in cyclamen aldehyde is accepted.

CRITICAL EFFECT:

SENSITIZATION

REFERENCES:

This recommendation is based on the sensitizing potential of cyclamen alcohol and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction in a sample of cyclamen alcohol, and the absence of sensitization reaction is a sample of cyclamen alcohol, and the absence of sensitization reaction is a sample of cyclamen alcohol, and the absence of sensitization reaction is a sample of cyclamen alcohol, and the absence of sensitization reaction is a sample of cyclamen alcohol, and the absence of sensitization reaction is a sample of cyclamen alcohol, and the absence of sensitization reaction alcohol, and the abse

Cyclamen aldehyde

CAS N°:	103-95-7	Empirical formula: Structure:	
Synonyms:	Benzenepropanal, α-methyl-4 Benzenepropanol, .αmethyl- 3-p-Cumenyl-2-methylpropion Cyclamal Cyclaviol Cyclosal p-Isopropyl-α-methylhydrocinr 3-(4-Isopropylphenyl)-2-methy 2-Methyl-3-(p-isopropylphenylpro α-Methyl-p-isopropylphenylpro α-Methyl-4-(1-methylethyl)ber 2-Methyl-3-(pisopropylphenyl)	-(1-methylethyl)- 4-(1-methylethyl)- aldehyde /lpropanal)propionaldehyde pylaldehyde izenepropanal propionaldehyde	

History:	Initial reviews:	June 2013	
	Current revision date:	2015	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.17 %	Category 7	0.45 %		
Category 2	0.22 %	Category 8	2.00 %		
Category 3	0.89 %	Category 9	5.00 %		
Category 4	2.67 %	Category 10	2.50 %		
Category 5	1.40 %	Category 11	See Note box (2)		
Category 6	4.28 %				
Note box:	-				
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofi.org)					
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.					
Fragrance material specification	ons:	Cyclamen aldehyde should 1.5% of Cyclamen alcohol.	not contain more than		

Cyclamen aldehyde

CONTRIBUTION FROM OTHER SOURCES:

See Annex I.

CRITICAL EFFECT:

DERMAL SENSITIZATION

RIFM SUMMARIES:

			Human Data		
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
5413 [1] ⁴	Weak	5905	2069	NA	5900

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No Observed Effect Level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test;

LOEL = Lowest Observed Effect Level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴EC3 value from one LLNA, not the mean.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Cyclamen aldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 5900 μ g/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Cyclamen aldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick GF, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 1971. Maximization study. RIFM report number 1805, March 25. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. Repeat Insult Patch Test. Draft RIFM Report number 63811. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.) 2012. Local Lymph Node Assay. Draft RIFM Report number 63815. (RIFM, Woodcliff Lake, NJ, USA).

	C S	yciopentadecanolide	
CAS N°:	106-02-5	Empirical formula: Structure:	C ₁₅ H ₂₈ O ₂
Synonyms:	Angelica lactone Cyclopentadecanolide 15-Hydroxypentadecanoic a Oxacyclohexadecan-2-one Pentadecalactone w-Pentadecalactone Pentadecanolide Cyclopentadecanolid Supra Exaltex (commercial name) Exaltolide (commercial name Muskalactone (commercial name Thibetolide (commercial name	icid, ω-lactone (commercial name) e) e) name) e) ne)	

History:	Initial reviews:	New Standard	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Category 1 See Note box (1)	0.16 %	Category 7	0.42 %	
Category 2	0.20 %	Category 8	2.00 %	
Category 3	0.83 %	Category 9	5.00 %	
Category 4	2.50 %	Category 10	2.50 %	
Category 5	1.31 %	Category 11	See Note box (2)	
Category 6	3.93 %			



Cyclopentadecanolide

Note box:

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org)

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
>12,500 [1] ⁴	Weak	5500⁵	6900⁵	NA	5500

All data in this table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴EC3 value from one LLNA conducted on a high purity material, not the mean. LLNA data from three commercial samples, with varying degrees of impurities, resulted in a range of EC3 values (<2500 – 6375 μg/cm²). ⁵HRIPT and HMT conducted on commercial materials.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Cyclopentadecanolide and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 5500 µg/cm². The material may contain unidentified impurities that have potential to induce sensitization. While a purified material shows no potential for dermal sensitization in an LLNA, the impurities remain unidentified. As such the NESIL is based on the commercial material. The Panel recommends the limits for the 11 different product categories, which are the acceptable use levels of Cyclopentadecanolide in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick GF, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 1974. Report on human maximization studies. RIFM report number 1779 06/05A. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2006. Repeated insult patch test with ω -pentadecalactone. Unpublished study from Symrise GmbH & Co. KG, 21 October. Report number 60740. (RIFM, Woodcliff Lake, NJ, USA).

Cyclopentadecanolide

RIFM (Research Institute for Fragrance Materials, Inc.), 2009. Local Lymph Node Assay. Unpublished study from Symrise GmbH & Co. KG, 21 October. Report number 60740. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2009. Local Lymph Node Assay. Unpublished study from Symrise GmbH & Co. KG, 21 October. Report number 60741. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2010. Local Lymph Node Assay. Unpublished study from Symrise GmbH & Co. KG, 2 December. Report number 60742. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2010. Local Lymph Node Assay. Unpublished study from Symrise GmbH & Co. KG, 21 January. Report number 60743. (RIFM, Woodcliff Lake, NJ, USA).



Dibenzyl ether

CAS N°:	103-50-4	Empirical formula: Structure:	C14H14O
Synonyms:	Benzene, 1,1'-[oxybis(methyler Benzyl ether Benzyl oxide Dibenzyl oxide 1,1'-[Oxybis(methylene)]dibenz	ne)]bis- zene	

History:	Initial reviews:	New Standard		
	Current revision date:	2009		
	Implementation date:	For new submissions*:	August 7, 2009	
		For existing fragrance compounds*:	August 7, 2011	
	Next review date	2014		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Category 1 See Note box (1)	0.07 %	Category 7	0.17 %	
Category 2	0.08 %	Category 8	2.00 %	
Category 3	0.35 %	Category 9	5.00 %	
Category 4	1.04 %	Category 10	2.50 %	
Category 5	0.55 %	Category 11	See Note Box (2)	
Category 6	1.67 %			
Note box:				
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry http://www.iofiorg.org/) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no untification for a restriction of the concentration of this fragmence intendion in the finished product. 				

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)
Dibenzyl ether

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
1575 [1]⁴	Weak	2362	2760	NA	2300

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or HMT

²Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

³WoE NESIL limited to three significant figures

⁴EC3 value from one LLNA, not the mean.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Dibenzyl ether and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 2300 □g/cm2. They recommend the limits for the 11 different product categories, which are the acceptable use levels of Dibenzyl ether in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1974. Maximization study with dibenzyl ether. RIFM report number 1779, June 4a. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007. Local Lymph Node Assay. RIFM report number 52907, May 21. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2008. Human repeated insult patch test. RIFM report number 54679, April 16a. (RIFM, Woodcliff Lake, NJ, USA). 3rd Amendment

1,3-Dibromo-2-methoxy-4-nitro-5-(1,1-dimethylethyl)-6-methyl-benzene (Musk alpha)

CAS N°:	63697-53-0	Empirical formula:	$C_{12}H_{15}Br_2NO_3$
Synonyms:	Musk alpha		
	Benzene,1,3-dibromo-5-(1,1-dimethy	lethyl)-2- methoxy-4-met	thyl-6-nitro-

History:	Initial reviews:	Not applicable	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products:	N/A				
Note box:					
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.					
Fragrance material specifications:		N/A			

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX

1,3-Dibromo-2-methoxy-4-nitro-5-(1,1-dimethylethyl)-6-methyl-benzene (Musk alpha)

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or

3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000

CAS N°:	62265-99-0	Empirical formula: Structure:	C ₈ H ₇ Br ₂ NO ₃ Br Br
Synonyms:	Benzene, 1,3-dibromo-2-methoxy-4-met Bromorose 1,3-Dibromo-2-methoxy-5-nitro-6-methy 2,4-Dibromo-3-methoxy-6-nitrotoluene 2,6-Dibromo-3-methyl-4-nitroanisole 6-Nitro-2,4-dibromo-3-methoxytoluene Musk KS	hyl-5-nitro- Ibenzene	

1,3-Dibromo-2-methoxy-4-methyl-5-nitrobenzene (Musk KS)

History:	Initial reviews:	Not applicable	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
	Including household cleaning products				
Non skin contact products:	N/A				
Note box:					
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.					
Fragrance material specificatio	ns:	N/A			

CONTRIBUTION FROM OTHER SOURCES:

N/A



CRITICAL EFFECT:

SEE NOTE BOX

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000

2,2-Dichloro-1-methylcyclopropylbenzene



History:	Initial reviews:	Not applicable	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



2,2-Dichloro-1-methylcyclopropylbenzene

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or

2) adverse data on the material itself and/or

3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000



	2,4-Dier	als	
CAS N°:	Including but not limited to: 764-40-9 142-83-6 80466-34-8 5910-85-0 30361-28-5 6750-03-4 2363-88-4 13162-46-4 21662-16-8 25152-84-5 30361-29-6 4313-03-5 (including all geometric isomers)	Empirical formula: Structure:	C _{5+n} H _{6+2n} O o
Synonyms:	Including but not limited to: 2,4-Pentadienal 2,4-Hexadienal 2,4-Heptadienal 2,4-Octadienal 2,4-Octadienal 2,4-Decadienal 2,4-Undecadienal 2,4-Dodecadienal trans,trans-2,4-Decadienal trans,trans-2,4-Undecadienal 2,4-Heptadien-1-al		

History:	Initial reviews:	New Group Standard	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*: See Note Box	
		For existing fragrance compounds*:	See Note Box
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products:	N/A				



Note box:

This IFRA Standard represents the group of 2-4-Dienals and replaces the existing individual IFRA Standards for the materials listed above. This new group also includes any other 2,4-Dienals. Note that 2,4-Hexadienal, 2,4-Heptadienal, 2,4-Octadienal, 2,4-Nonadienal, 2,4-Decadienal and 2,4-Undecadienal have already been banned via individual IFRA Standards therefore no implementation time is necessary. For all other 2,4-dienals (such as 2,4-Pentadienal and 2,4-Dodecadienal), the implementation date is August 10, 2013 for new submissions and August 10, 2014 for existing fragrance compounds.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None to consider

CRITICAL EFFECT:

INSUFFICIENT DATA

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000



History:	Initial reviews:	June 1975	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.						
Fragrance material specificatio	ns:	N/A				

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION



Diethyl maleate

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002

REFERENCES:

Opdyke, D.L.J. (1976), Food and Cosmetics Toxicology 14, 443.



Dihydrocoumarin

CAS N°:	119-84-6	Empirical formula: Structure:	C ₉ H ₈ O ₂
Synonyms:	1,2-Benzodihydropyrone 2H-1-Benzopyran-2-one, 3,4- Chroman-2-one 2-Chromanone 3,4-Dihydro-2H-1-benzopyrar o-Hydroxydihydrocinnamic ac Melilotic acid lactone	dihydro- n-2-one cid lactone	

History:	Initial reviews:	October 1974	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.029%	Category 7	0.08 %			
Category 2	0.037 %	Category 8	1.01 %			
Category 3	0.15 %	Category 9	5 %			
Category 4	0.45 %	Category 10	2.5 %			
Category 5	0.24 %	Category 11	See Note box (2)			
Category 6	0.72 %					
Note box:						
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. 						
Fragrance material specifications: N/A						
CONTRIBUTION FROM OTHER SOURCES:						

See Annex I

Dihydrocoumarin

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm ²) [no. studies] Based on Animal Data		NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
1070 [3]4	Moderate	NA⁵	NA	2000	1000

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴EC3 value from the mean of the individual LLNAs.

⁵ An HRIPT conducted on 49 subjects produced 2 questionable reactions which were not confirmed.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Dihydrocoumarin and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1000 µg/cm², which is a default value based on the LLNA data. They recommend the limits for the 11 different product categories, which are the acceptable use levels of Dihydrocoumarin in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick GF, et al, 2008. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 52(1): 3-23.

Kimber, I., Hilton, J., Weisenberger, C., 1989. The murine local lymph node assay for identification of contact allergens: A preliminary evaluation of in situ measurement of lymphocyte proliferation. *Contact Dermatitis*, 21, 215-220.

Kimber, I., Weisenberger, C., 1991. Anamnestic responses to contact allergens: Application in the murine local lymph node assay. *Journal of Applied Toxicology*, 11(2), 129-133.

Opdyke, D.L.J. (1974), Food and Cosmetics Toxicology 12, 521.

RIFM (Research Institute for Fragrance Materials, Inc.), 2003. Evaluation of skin sensitization potential of coumarin in mice using the Local Lymph Node Assay (LLNA). Unpublished study from Rhodia, Inc. Report number 47072. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. Local Lymph Node Assay on Dihydrocoumarin. Draft RIFM report number 63816. (RIFM, Woodcliff Lake, NJ, USA).



6,7-Dihydro-1,1,2,3,3-pentamethyl-4(5H)-indanone (DPMI)

CAS N°:	33704-61-9	Empirical formula: Structure:	C14H22O H ₃ C H ₃ C H ₃ C H ₃ C
Synonyms:	DPMI 1,2,3,5,6,7-Hexahydr 4H-Inden-4-one, 1,2, 1,1,2,3,3-Pentamethy Cashmeran (commen	o-1,1,2,3,3-pentamethyl-4H-ind 3,5,6,7-hexahydro-1,1,2,3,3-per /l-1,2,3,5,6,7-hexahydro-4H-ind rcial name)	len-4-one ntamethyl- len-4-one

History:	Initial reviews:	New Standard	
	Current revision date:	2015	
	Implementation date:	For new submissions*:	August 10, 2015
		For existing fragrance compounds*:	August 10, 2016
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.34 %	Category 7	0.91 %		
Category 2	0.44 %	Category 8	2.00 %		
Category 3	1.81 %	Category 9	5.00 %		
Category 4	5.43 %	Category 10	2.50 %		
Category 5	2.86 %	Category 11	See Note box (2)		
Category 6	8.70 %				
Note box:					

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofi.org)

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

6,7-Dihydro-1,1,2,3,3-pentamethyl-4(5H)-indanone (DPMI)

CRITICAL EFFECT:

DERMAL SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (µg/cm ²) [no. studies] Based on Animal Date		NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
8250 [1]	Weak	12,121	NA	NA	12,000

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No Observed Effect Level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = Lowest Observed Effect Level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003. ²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 6,7-Dihydro-1,1,2,3,3-pentamethyl-4(5H)-indanone (DPMI) and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 12,000 μ g/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of 6,7-Dihydro-1,1,2,3,3-pentamethyl-4(5H)-indanone (DPMI) in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api, A. M., Basketter, D. A., Cadby, P. A., Cano, M-F., Ellis, G., Gerberick, G. F. et al., 2008. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. Regulatory Toxicology and Pharmacology 52(1): 3-23.

Roberts, D. W., Patlewicz, G., Kern, P. S., Gerberick, F., Kimber, I., Daerman, R. J., Ryan, C. A., Basketter, D. A. and Aptula, A. O., 2007. Mechanistic applicability domain classification of a local lymph node assay dataset for skin sensitization. Chem Res Toxicol. Jul;20(7): 1019-30.

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. 6,7-Dihydro-1,1,2,3,3-pentamethyl-4(5H)-indanone (cashmeran): Local Lymph Node Assay in the mouse. Unpublished study from I.F.F., Report number 65177 (RIFM, Woodcliff Lake, NJ, USA).

Deziel, N. C., Wei, W. Q. and Abnet C. C., 2013. A multi-day environmental study of polycyclic aromatic hydrocarbon exposure in a high-risk region for esophageal cancer in China. J Expo Sci Environ Epidemiol. Jan-Feb; 23(1): 52-9.

(ifra

2,4-Dihydroxy-3-methylbenzaldehyde



History:	Initial reviews:	February 1980, April 1989		
	Current revision date:	September 2002		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		
* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.				

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products:	N/A				
Note box:					
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.					
Fragrance material specification	ons:	N/A			

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002

REFERENCES:

Ford, R.A. (1988), Food and Chemical Toxicology 26, 303.



Dimethylcyclohex-3-ene-1-carbaldehyde (mixed isomers)

CAS N°:	68737-61-1 (mixed isomers) 68039-49-6 68039-48-5 27939-60-2 67801-65-4 36635-35-5 68084-52-6 35145-02-9	Empirical formula:	C ₉ H ₁₄ O
Synonyms:	Dimethylcyclohex-3-ene-1-carbalde 2,4-Dimethyl-3-cyclohexen-1-carbo 3,5-Dimethylcyclohex-3-ene-1-carba Dimethylcyclohex-3-ene-1-carbalde 3,6-Dimethyl-3-cyclohexene-1-carbo 3-Cyclohexene-1-carboxaldehyde, o 2,4-Dimethyltetrahydrobenzaldehyde (is Triplal, Vertocitral, Vertoliff, Tricycla Aldehyde AA (commercial names)	hyde (isomer mixture) (68737 xaldehyde (68039-49-6) aldehyde (68039-48-5) hyde (isomer unspecified) (21 oxaldehyde (67801-65-4) dimethyl- (isomer mixture) e somer mixture) I, Hivertal, Agrumen Aldehydd	7-61-1) 7939-60-2) e, Cyclovertal, Ligustral,

History:	Initial reviews:	June 11, 2010	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.17 %	Category 7	0.45 %			
Category 2	0.22 %	Category 8	2.0 %			
Category 3	0.89 %	Category 9	5.0 %			
Category 4	2.7 %	Category 10	2.5 %			
Category 5	1.4 %	Category 11	See Note box (2)			
Category 6 4.3 %						
Note box:						
The above limits apply to Dimethylcyclohexen-3-ene-1-carbaldehyde (mixed isomers) used individually or in combination. (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of						

ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry www.iofiorg.org)

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.



Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

See Annex II

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data				
LLNA weighted mean EC3 values (µg/cm ²) [no. studies] Based on Animal Data ¹		NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (μg/cm ²)	
2500-5875 [7] ⁴	Weak	5905⁵	1380-6900 ⁶	N/A	5900	

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴A range of values and not the weighted mean was provided because the seven studies were performed on 7 different materials – all are isomers but the isomeric mixtures were different. There are currently 4 isomers of this material in the RIFM Database. Each material is typically a mixture of two or more isomers. In order to confirm the potency and characterization of these materials, a set of LLNAs was conducted on different commercial samples provided by individual suppliers with isomer ratios specified. The LLNA data showed all materials with similar dermal sensitization potency.

⁵MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL. The HRIPT MT-NOEL reported is for the fragrance material with the highest reported use in perfumery (IFRA Survey, 2004); the LLNA data showed all materials with similar dermal sensitization potency.

⁶A range of values was provided; three human maximization studies were conducted. No dermal sensitization was observed in any of the tests. The 3 studies were performed on 3 different isomer mixtures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for dimethylcyclohex-3-ene-1-carbaldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 5900 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of dimethylcyclohex-3-ene-1-carbaldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick GF, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 2010a. Local Lymph Node Assay. RIFM report number 58108, Draft Report. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2010b. Local Lymph Node Assay. RIFM report number 58112, Draft Report. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2010c. Local Lymph Node Assay. RIFM report number 58113, Draft Report. (RIFM, Woodcliff Lake, NJ, USA).



Dimethylcyclohex-3-ene-1-carbaldehyde (mixed isomers)

RIFM (Research Institute for Fragrance Materials, Inc.), 2009. Human repeated insult patch test. RIFM report number 58150, December 14. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1982. Maximization study. RIFM report number 1643, October 5. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1978. Maximization study. RIFM report number 1698, February 27b. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1977. Maximization study. RIFM report number 1702, May 4c. (RIFM, Woodcliff Lake, NJ, USA).



1-(5,5-Dimethyl-1-cyclohexen-1-yl)pent-4-en-1-one



History:	Initial reviews:	New Standard		
	Current revision date:	2009		
	Implementation date:	For new submissions*: August 7, 2009		
		For existing fragrance compounds*:	August 7, 2011	
	Next review date	2014		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.07 %	Category 7	0.19 %		
Category 2	0.09 %	Category 8	2.00 %		
Category 3	0.38 %	Category 9	5.00 %		
Category 4	1.13 %	Category 10	2.50 %		
Category 5	0.60 %	Category 11	See Note Box (2)		
Category 6	1.81 %				
Note box:					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry http://www.iofiorg.org/)					
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)

CRITICAL EFFECT:

SENSITIZATION

1-(5,5-Dimethyl-1-cyclohexen-1-yl)pent-4-en-1-one

RIFM SUMMARIES:

		Human Data				
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)	
745 [1] ⁴⁻⁵	Moderate	2500	NA	NA	2500	

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹ Data derived from HRIPT or HMT.

²Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

³ WoE NESIL limited to two significant figures.

⁴ EC3 value from one LLNA, not the mean.

⁵ LLNA used a very wide spacing of doses. Based on the response at each dose the EC3 value would be slightly on the low side. The response in the LLNA was:

0.1% SI of 0.9 1.0% SI of 0.9 10% SI of 10.4 100% SI of 17.5

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 1-(5,5-Dimethyl-1-cyclohexen-1- yl)pent-4-en-1-one and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 2500 mg/cm2. They recommend the limits for the 11 different product categories, which are the acceptable use levels of 1-(5,5-Dimethyl-1-cyclohexen-1-yl)pent-4-en-1-one in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1999. Human repeated insult patch test. Unpublished study from Firmenich Inc., 29a June. Report number 42138. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001a. Local Lymph Node Assay. Unpublished study from Givaudan, 16 May. Report number 42073. (RIFM, Woodcliff Lake,NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001b. Human repeated insult patch test. Unpublished study from IFF Inc., 29 October. Report number 51118. (RIFM, Woodcliff Lake, NJ, USA).



4,6-Dimethyl-8-tert-butylcoumarin

CAS N°:	17874-34-9	Empirical formula:	$C_{15}H_{18}O_2$
CAS N°:	17874-34-9	Empirical formula: Structure:	
			0
Synonyms:	2H-1-Benzopyran-2-one, 8-(1,1-dimethy Butolia	lethyl)-4,6-dimethyl-	

History:	Initial reviews:	February 1979, June 1981		
	Current revision date:	September 2002		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		
* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.				

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
	Including household cleaning products				
Non skin contact products:	N/A				
Note box:					
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.					
Fragrance material specification	Fragrance material specifications: N/A				

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

PHOTOALLERGY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002



4,6-Dimethyl-8-tert-butylcoumarin

REFERENCES:

Opdyke, D.L.J. (1980), Food and Cosmetics Toxicology 18, 671.



3,7-Dimethyl-2-octen-1-ol



History:	Initial reviews:	New Standard**	
	Current revision date:	October 2003	
	Implementation date:	For new submissions*: Not applicable**	
		For existing fragrance compounds*:	Not applicable**
	Next review date	Not applicable**	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products:	N/A				
Note box:					
This material should not be use	d as fragrance i	ngredient for any app	blication.		
**The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.					
Fragrance material specificatio	Fragrance material specifications: N/A				

CONTRIBUTION FROM OTHER SOURCES:

None Known.

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, October 2003

REFERENCES:

Ford et al., 1992, Food and Chemical Toxicology, Volume 30, Supplement, Special Issue VIII, page 19S.

2,2-Dimethyl-3-(3-tolyl)propan-1-ol

CAS N°:	103694-68-4	Empirical formula: Strucutre:	C ₁₂ H ₁₈ O OH
Synonyms:	Benzenepropanol,.ß.,. ß.,3-trim 2,2-Dimethyl-3-(3-methylpheny Majantol (commercial name) Linlan alcohol (commercial nam	iethyl I)propanol ne)	

History:	Initial reviews:	July 2008 (43rd Amendment), March 2010		
	Current revision date:	June 11, 2010		
	Implementation date:	For new submissions*: January 11, 2011		
		For existing fragrance compounds*:	January 11, 2012	
	Next review date	March 2015		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED / SPECIFICATION

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.28%	Category 7	0.8%		
Category 2	0.36%	Category 8	2.0%		
Category 3	1.5%	Category 9	5.0%		
Category 4	4.5%	Category 10	2.5%		
Category 5	2.4%	Category 11	See Note Box (2)		
Category 6	7.2%				
Note box:					
 (1) See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products with the potential of ingestion. (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. This Standard replaces the existing one on 2,2-Dimethyl-3-(3-tolyl)propan-1-ol distributed with the 43rd Amendment, which only contained the restrictions based on the QRA. 					
Fragrance material specifications:2,2-Dimethyl-3-(3-tolyl)propan-1-ol should only be used as a fragrance ingredient if traces of organochlorine compounds are restricted. Total chlorine, which can be measured by Atomic Absorption Spectroscopy, must not exceed 25 pp in the raw material.					

CONTRIBUTION FROM OTHER SOURCES:

None known at the time of the publication of the Standard

2,2-Dimethyl-3-(3-tolyl)propan-1-ol

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (µg/cm ²) [no. studies] Potency Classification Based on Animal Data ¹		NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
>7500	Weak	9900 ⁴	N/A	N/A	9900

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Dose reported reflect the highest concentration tested, not necessarily the highest achievable NOEL.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 2,2-Dimethyl-3-(3-tolyl)propan-1-ol and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 9900 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of 2,2-Dimethyl-3-(3-tolyl)propan-1-ol in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 2002. Local Lymph Node Assay. Unpublished report from Symrise GmbH & Co. KG, 9 December. Report number 49523 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Repeated Insult Patch Test. Unpublished report from Symrise GmbH & Co. KG,. Report number 49526 (RIFM, Woodcliff Lake, NJ, USA)

RIFM (Research Institute for Fragrance Materials, Inc.), 2007. Repeated Insult Patch Test. Unpublished report from Symrise GmbH & Co. KG, 26 September. Report number 53799 (RIFM, Woodcliff Lake, NJ, USA).



History:	Initial reviews:	October 1976	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke, D.L.J. (1976), Food and Cosmetics Toxicology 14, 749

Diphenylamine

CAS N°:	122-39-4	Empirical formula:	$C_{12}H_{11}N$
		Structure:	
			\land
Synonyms:	Benzeneamine, N-phenyl-		

History:	Initial reviews:	New Standard**	
	Current revision date:	October 2003	
	Implementation date:	For new submissions*:	Not applicable**
		For existing fragrance compounds*:	Not applicable**
	Next review date	Not applicable**	
* This date applies t	o the supply of fragrance com	pounds (formulas) only, not to the finished product	s in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products: N/A					
Note box:					

This material should not be used as fragrance ingredient for any application.

**The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None Known.

CRITICAL EFFECT:

TOXICITY, TERATOGENICITY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, October 2003

REFERENCES:

Opdyke, 1978, Food and Cosmetics Toxicology, Volume 16, Supplement 1, Special Issue IV, page 723-727.

2,4-Dodecadien-1-ol, (2E, 4E)

CAS N°:	18485-38-6	Empirical formula: Structure:	C12H22O
Synonyms:	2,4-Dodecadien-1-ol		

History:	Initial reviews:	New Standard		
	Current revision date:	2015		
	Implementation date:	For new submissions*:	August 10, 2015	
		For existing fragrance compounds*:	August 10, 2016	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave-on products:	ucts: N/A Rinse-off products: N/A				
	Including household cleaning products				
Non-skin contact products:	<u>:</u> N/A				
Fragrance material specificatio	ns:	N/A			

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

INSUFFICIENT DATA

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

1) presence of structural alerts as defined in Api et al. (2014) and/or

2) adverse data on the material itself and/or

3) adverse data for a structurally related material.



2,4-Dodecadien-1-ol, (2E, 4E)

REFERENCES:

Api A.M., Belsito D., Bruze M., Cadby P., Calow P., Dagli M. L., Dekant W., Dent M., Ellis G., Fryer A. D., Fukayama M., Griem P., Hickey C., Kromidas L., Lalko J., Liebler D.C., Miyachi Y., Politano V.T., Renskers K., Ritacco G., Salvito D., Schultz T.W., Sipes I. G., Smith B., Vitale D., Wilcox D.K. (2014). Food and Chemical Toxicology, <u>http://dx.doi.org/10.1016/j.fct.2014.11.014</u>.

Esters of 2-nonynoic acid (Except methyl octtine carbonate)

CAS N°:	10031-92-2	Empirical formula:	N/A
Synonyms:	Ethyl 2-nonynoate Ethyl octine carbonate Ethyl octyne carbonate 2-Nonynoic acid, ethyl ester		

History:	Initial reviews:	Not applicable	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	N/A	Rinse-off products:	N/A	
		Including household	cleaning products	
Non skin contact products:	N/A			
Note box:				
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.				
Methyl octine carbonate is not Prohibited because it is IFRA Restricted.				
Fragrance material specifications: N/A				

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



Esters of 2-nonynoic acid (Except methyl octtine carbonate)

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000



CAS N°:	10484-32-9 10519-20-7	Empirical formula:	C13H22O2 & C10H16O2
Synonyms:	 10519-20-7: Amyl heptine carbonate 2-Octynoic acid, pentyl ester Pentyl 2-octynoic acid Principal Vert de violette 10519-20-7: Ethyl heptine carbonate Ethyl 2-octynoate Principal 2-Octynoic acid, ethyl ester 		

History:	Initial reviews:	Not applicable	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	N/A	Rinse-off products:	N/A	
		Including household	cleaning products	
Non skin contact products:	N/A			
Note box:				
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.				
Methy hopfine carbonate is not i foliblica because it is if the Restituted.				
Fragrance material specificatio	ns:	N/A		

CONTRIBUTION FROM OTHER SOURCES:

N/A



CRITICAL EFFECT:

SEE NOTE BOX

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or

2) adverse data on the material itself and/or

3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000

Estragole

CAS N°:	140-67-0	Empirical formula: Structure:	C ₁₀ H ₁₂ O
Synonyms:	<i>p</i> -Allylanisole 1-Allyl-4-methoxybenzene Benzene, 1-methoxy-4-(2-propenyl)- Chavicyl methyl ether Isoanethole p-Methoxyallylbenzene 1-Methoxy-4-(2-propen-1-yl)benzene Methyl chavicol		

History:	Initial reviews:	October 2009	
	Current revision date:	2015	
	Implementation date:	For new submissions*: Not Applicable	
		For existing fragrance compounds*:	Not Applicable
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave-on products:	See note box	Rinse-off products:	See note box	
Non-skin contact products:	See note box Including house	hold cleaning products	5	
Note box:				
The total concentration of Estragole should not exceed the following limitations in the finished product: Fine fragrance and Eau de Toilette: 0.2% Other leave-on and rinse-off cosmetic products: 0.01% Non-skin, incidental skin contact products: 0.2%				
Fragrance material specifications: N/A				

CONTRIBUTION FROM OTHER SOURCES:

See Annex I.

CRITICAL EFFECT:

CARCINOGENICITY

Estragole

RFIM SUMMARIES:

Although Estragole has been shown to cause tumors in laboratory animals (NTP, 2008) the studies have a number of limitations that have an impact on its direct application to human risk and particularly risk from dermal contact:

- 1. The current studies are confounded by high dose toxicity including significant hepatotoxicity, gastric damage and malnutrition in both mice and rats. This makes the distinction between primary and secondary mechanisms of tumour formation difficult.
- 2. Current scientific evidence supports a non-linear relationship between dose and the potential for carcinogenicity of Estragole and related substances (Smith *et al.*, 2002). This is due to differences in the way that Estragole is metabolized at high versus low doses. Studies indicate that all these events are likely to be minimal in the dose range of 1-10 mg/kg body weight (Zangouras *et al.*, 1981; Anthony *et al.*, 1987). The lowest dose in the NTP 90-day study used in this assessment was 37.5 mg/kg body weight/day, thus care needs to be taken in extrapolating to lower doses relevant to fragrance exposure. This non-linear response has also been used to support the risk assessment for exposure to Estragole in Flavours.

Consideration also needs to given to differences between dermal and oral exposure. Introduction of a bolus dose of test material into the stomach leads to higher peak blood plasma levels and increased metabolic demand compared with the slower, more steady absorption of the substance from the skin. Furthermore, although no data exist on the skin metabolism of Estragole or related compounds there is evidence that many enzymatic processes, particularly oxidative ones, are much lower in the skin than in the liver (Bronaugh *et al.*, 1995). Thus the relevance of reported tumours resulting from skin painting studies or subcutaneous injection (Miller *et al.*, 1983) with putative genotoxic metabolites of Estragole needs to be put into perspective. Although data indicate that the most potent metabolite for inducing skin tumours in rodents is the1'-hydroxy epoxide metabolite, characterization of dermal metabolism has not been established to show that the epoxide metabolites used in the skin painting and subcutaneous injection studies would be the metabolite of concern in either rat or mouse, nor has it been established that the level of exposure is relevant as it is unlikely that significant local tissue concentrations for metabolites would result from a realistic oral ingestion or dermal application of Estragole.

REXPAN RATIONALE / CONCLUSION:

The total dermal exposure resulting from the limited use of Estragole as described in this Standard is 0.04 mg/kg body weight/day. Making the following conservative assumptions:

- 100% dermal absorption
- Metabolism human = rodent
- Metabolism skin = liver
- Oral LOEL = 37.5 mg/kg/day based on rat oral low dose observed hyperplasia (oval, bile cell) and it assumes that the hyperplasia will progress to tumor formation
- UF = 1,000 (10 for LOEL to NOAEL, 10 for species, 10 for inter-individual variability)

Worst Case Risk Assessment:

• Systemic RfD = 37.5/1,000 = 0.04 mg/kg/day

The IFRA Standard reflects the potential for the higher presence of Estragole in hydroalcoholic and air freshener products whilst ensuring that the RfD for cumulative exposure through all product types will not be exceeded.

REFERENCES:

Anthony, A., Caldwell, J., Hutt, A.J., Smith R.L., 1987. Metabolism of Estragole in rat and mouse and influence of dose size on excretion of the proximate carcinogen 1'-hydroxyestragole. Food and Chemical Toxicology 25, 799-806.

Bronaugh, R.L., 1995. Methods for in Vitro Skin Metabolism Studies. Toxicology Mechanisms and Methods, Volume 5, Issue 4, pages 275 – 281.

Miller, E.C., Swanson, A.B., Phillips, D.H., Fletcher, T.L., Liem, A., Miller, J.A., 1983. Structure-activity studies of the carcinogenicities in the mouse and rat of some naturally occurring and synthetic alkenylbenzene derivatives related to Safrole and Estragole. Cancer Research 43, 1124-1134.
Estragole

National Toxicology Program, 2008. NTP Technical Report on the 3-month toxicity studies of Estragole administered by gavage to rats and mice. National Toxicology Program Toxicity Report Series Number 82. NIH Publication No. 08-5966.

Phillips, D.H., Miller, J.A., Miller, E.C., Adams, B., 1981. Structures of the DNA adducts formed in mouse liver after administration of the proximate hepatocarcinogen 1'-hydroxyestragole. Cancer Research 41, 176-186.

Smith R.L., Adams T.B., Doullc J., Ferond V.J., Goodmane J.I., Marnettf L.J., Portogheseg P.S., Waddellh W.J., Wagneri B.M., Rogers A.E., Caldwellk J., and Sipes I.G. 2002. Safety assessment of Allylalkoxybenzene derivatives used as flavoring substances — Methyl eugenol and Estragole. Food and Chemical Toxicology 40: 851–870.

Swanson, A.B., Miller, E.C., Miller, J.A., 1981. The side-chain epoxidation and hydroxylation of the hepatocarcinogens Safrole and Estragole and some related compounds by rat and mouse liver microsomes. Biochemica et Biophysica Acta 673, 504-516.

Wiseman, R.W., Fennell, T.R., Miller, J.A., Miller, E.C., 1985. Further characterization of the DNA adducts formed by electrophilic esters of the hepatocarcinogens 1'-Hydroxysafrole and 1'-Hydroxyestragole *in-vitro* and in mouse liver *in-vivo*, including new adducts at C-8 and N-7 of guanine residues. Cancer Research 45, 3096-3105.

Zangouras, A., Caldwell, J., Hutt, A.J., Smith, R.L., 1981. Dose dependent conversion of Estragole in the rat and mouse to the carcinogenic metabolite, 1'-hydroxyestragole. Biochemical Pharmacology, 30, 1383-1386.

2-Ethoxy-4-methylphenol



History:	Initial reviews:	New Standard	
	Current revision date:	2008	
	Implementation date:	For new submissions*: August 16, 2008	
		For existing fragrance compounds*:	August 16, 2010
	Next review date	2013	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.01 %	Category 7	0.02 %		
Category 2	0.01 %	Category 8	0.2 %		
Category 3	0.03 %	Category 9	1.2 %		
Category 4	0.1 %	Category 10	1.9 %		
Category 5	0.1 %	Category 11	See Note Box (2)		
Category 6	0.2 %				
Note box:					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry http://www.iofiorg.org/)					
there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.					
Fragrance material specifications: N/A					
CONTRIBUTION FROM OTHER SOURCES:					

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)

2-Ethoxy-4-methylphenol

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (µg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
	Moderate	236 ⁴	2362 ⁴	NA	230

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed

effect level; NA = Not Available.

² Data derived from HRIPT or HMT.

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

³ WoE NESIL limited to three significant figures.

⁴ MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 2-Ethoxy-4-methylphenol and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 230 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of 2-Ethoxy-4-methylphenol in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 2003a. Human Repeated Insult Patch Test. RIFM report number 44237, December 10 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003b. Human Repeated Insult Patch Test. RIFM report number 44238, December 10 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Human Repeated Insult Patch Test. RIFM report number 45135, March 30 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007. Human Repeated Insult Patch Test. RIFM report number 52894, May 15 (RIFM, Woodcliff Lake, NJ, USA).



Ethyl acrylate

CAS N°:	140-88-5	Empirical formula: Structure:	C5H8O2
Synonyms:	Ethyl propenoate 2-Propenoic acid, ethyl ester		

History:	Initial reviews:	November 1974	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke, D.L.J. (1975), Food and Cosmetics Toxicology 13, 801.



p-Ethylbenzaldehyde 4748-78-1 CAS N°: $C_9H_{10}O$ **Empirical formula:** Structure: 0 4-Ethylbenzaldehyde Synonyms: Benzaldehyde, 4-ethyl (CAS)

History:	Initial reviews:	New Standard		
	Current revision date:	June 2013		
	Implementation date:	For new submissions*: August 10, 2013		
		For existing fragrance compounds*:	August 10, 2014	
	Next review date	2018		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Category 1 See Note box (1)	0.03 %	Category 7	0.08 %	
Category 2	0.04 %	Category 8	1.11 %	
Category 3	0.17 %	Category 9	5.00 %	
Category 4	0.50 %	Category 10	2.50 %	
Category 5	0.26 %	Category 11	See Note box (2)	
Category 6	0.80 %			
Note hov:				

(1) See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products

there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None to consider.

p-Ethylbenzaldehyde

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

Cuminaldehyde has been designated as a read across material for p-Ethylbenzaldehyde for the sensitization endpoint. As such, the data for Cuminaldehyde are below:

		Human Data			
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
> 2500 [1] ⁴	Weak	1181	2760	NA	1100⁵

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴EC3 value from one LLNA, not the mean.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Cuminaldehyde and p-Ethylbenzaldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as $1100 \ \mu g/cm^2$ based on the critical sensitization data on Cuminaldehyde by read-across. They recommend the limits for the 11 different product categories, which are the acceptable use levels of p-Ethylbenzaldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick G, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. Regulatory Toxicology and Pharmacology 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 1972. Maximization test. RIFM report number 1804, November 22. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1975. Maximization test. RIFM report number 1804, March 27a. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1973. Maximization test. RIFM report number 1804, November 11. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. Repeat Insult Patch Test. Draft RIFM Report number 63810. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.) 2012. Local Lymph Node Assay. Draft RIFM Report number 63814. (RIFM, Woodcliff Lake, NJ, USA).



Ethylene glycol monoethyl ether and its acetate

CAS N°:	110-80-5 111-15-9 (acetate)	Empirical formula: Structure:	C ₄ H ₁₀ O ₂ ; C ₆ H ₁₂ O ₃ 0 Acetate: 0 0 0 0 0 0 0 0 0 0 0 0 0
Synonyms:	Ethylene glycol ethyl ether 2-Ethoxyethanol Ethanol, 2-ethoxy- Cellosolve Oxitol Ethylene glycol ethyl ether acetate 2-Ethoxyethyl acetate Ethyl cellosolve acetate Ethanol, 2-ethoxy-, acetate 1-Acetoxy-2-ethoxyethane		

History:	Initial reviews:	New Standard**		
	Current revision date:	October 2003		
	Implementation date:	For new submissions*: Not applicable**		
		For existing fragrance compounds*:	Not applicable**	
	Next review date	Not applicable**		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:	Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
This material should not be used as fragrance ingredient for any application.						
**The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.						
Fragrance material specificatio	ns:	N/A				

Ethylene glycol monoethyl ether and its acetate

CONTRIBUTION FROM OTHER SOURCES:

None to consider

CRITICAL EFFECT:

REPRODUCTIVE TOXICITY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, October 2003

REFERENCES:

NIOSH, 1983, Current Intelligence bulletin, No. 39, page 1-20.

EPA, 1984b, EPA/540/1-86/052; PB86-134632.

ECETOC, 1985, ECETOC Technical Report, 17.

CAS N°:	109-86-4 110-49-6 (acetate)	Empirical formula: Structure:	$C_3H_8O_2$; $C_5H_{10}O_3$ HO Acetate:
Synonyms:	Ethylene glycol met 2-Methoxyethanol Ethanol, 2-methoxye Methyl cellosolve Ethylene glycol met 2-Methoxyethanol a 2-Methoxyethyl ace Methyl cellosolve ac Ethanol, 2-methoxye	hyl ether - hyl ether acetate cetate tate cetate cetate -, acetate	

Ethylene glycol monomethyl ether and its acetate

History:	Initial reviews:	New Standard**	
	Current revision date:	October 2003	
	Implementation date:	For new submissions*:	Not applicable**
		For existing fragrance compounds*:	Not applicable**
	Next review date	Not applicable**	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	N/A	Rinse-off products:	N/A	
		Including household	cleaning products	
Non skin contact products:	N/A			
Note box:				
This material should not be use	d as fragrance i	ngredient for any app	lication.	
**The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.				
Fragrance material specifications: N/A				

Ethylene glycol monomethyl ether and its acetate

CONTRIBUTION FROM OTHER SOURCES:

None Known.

CRITICAL EFFECT:

REPRODUCTIVE TOXICITY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, October 2003

REFERENCES:

NIOSH, 1983, Current Intelligence bulletin, No. 39, page 1-20.

EPA, 1984b, EPA/540/1-86/052; PB86-134632.

ECETOC, 1985, ECETOC Technical Report, 17.



Eugenol

CAS N°:	97-53-0	Empirical formula: Structure:	C10H12O2
Synonyms:	4-Allylcatechol-2-methyl ether 1-Allyl-4-hydroxy-3-methoxybenzene 4-Allyl-2-methoxyphenol Caryophyllic acid 2-Hydroxy-5-allylanisole 1-Hydroxy-2-methoxy-4-allylbenzene 4-Hydroxy-3-methoxy-1-allylbenzene 1-Hydroxy-2-methoxy-4-propenylben 2-Methoxy-4-allylphenol 2-Methoxy-4-(2-propenyl)phenol Phenol, 2-methoxy-4-(2-propenyl)- Eugenic acid Allylguaiacol, 4-Allylguaiacol	zene	

History:	Initial reviews:	April 2004, May 2006, May 2007		
	Current revision date:	2008		
	Implementation date:	For new submissions*:	August 16, 2008	
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.2 %	Category 7	0.4 %		
Category 2	0.2 %	Category 8	0.5 %		
Category 3	0.5 %	Category 9	0.5 %		
Category 4	0.5 %	Category 10	0.5 %		
Category 5	0.5 %	Category 11	See Note Box (2)		
Category 6	4.3 %				
Note box:					
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again.					



(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

Eugenol - Sensitization Potency Estimation Based on Weight of Evidence

		Human Data			
LINA weighted mean EC3 values (μg/cm²) [no. studies]	an EC3 values ²⁾ lies] Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
2703 [6]	Weak	5906	NA	NA	5900

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max tests

² Gerberick et al., 2001

³WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for eugenol and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 5900 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of eugenol in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Basketter, D.A., Lea, L.J., Dickens, A., Briggs, D., Pate, I., Dearman, R.J., Kimber, I., 1999. A comparison of statistical approaches to the derivation of EC3 values from local lymph node assay dose responses. Journal of Applied Toxicology, 19(4), 261-266.

Basketter, D.A., Gilmour, N., Dearman, R.J., Kimber, I., Ryan, C.A., Gerberick, F., 2003. Classification of skin sensitisation potency using the Local Lymph Node Assay. The Toxicologist, 72(S-1), 101.

Isola, D., Lalko, J., 2001a. Vehicle effects in the murine local lymph node assay (LLNA). American College of Toxicology Meeting, November 4-7. Washington DC.

Isola, D., Lalko, J., 2001b. Vehicle effects in the murine local lymph node assay (LLNA). International Journal of Toxicology, 20(6), 401.

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 2001. Repeated Insult Patch Test on dl-Citronellol. RIFM report number 39081, May 15. (RIFM, Woodcliff Lake, NJ, USA).





History:	Initial reviews:	October 1979, February 1980, 2002	
	Current revision date:	2006	
	Implementation date:	For new submissions*:	June 11, 2007
		For existing fragrance compounds*:	June 11, 2008
	Next review date	2011	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED / SPECIFICATION

RESTRICTIONS:

Limits in the finished product:					
Category 1	0.08 %	Category 7	0.2 %		
Category 2	0.11 %	Category 8	2.0 %		
Category 3	0.4 %	Category 9	5.0 %		
Category 4	1.2 %	Category 10	2.5 %		
Category 5	0.6 %	Category 11	See Note Box		
Category 6	2.0 %				
Note box:					
Category 11 includes all non-skin contact or inc is no justification for a restriction of the concent	idental skin contact proc ration of this fragrance ir	lucts. Due to the negligible skin contact front and the finished product.	om these types of products there		
This Standard replaces the existing one on I	Farnesol, which only co	ntained the purity criterion as outlined belo	ow.		
Fragrance material specifications:Farnesol should only be used as a fragrance ingredient if it contains a minimum of 96% of farnesol isomers as determined by GLC.					

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

Farnesol

RIFM SUMMARIES:

Farnesol - Sensitization Potency Estimation Based on Weight of Evidence

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
1200 [2]	Weak	2755	NA	68974	2700

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max tests

² Gerberick et al., 2001

³WoE NESIL limited to two significant figures

LOEL from human maximization test, not a human repeated insult patch test.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Farnesol and based on the weight of evidence established the No Expected Sensitization Induction Level (NESIL) as 2700 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Farnesol in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group, Technical Dossier of March 15, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

Research Institute for Fragrance Materials, Inc (1976). Human maximization test with Farnesol. RIFM report number 1797, 11b November (RIFM, Woodcliff Lake, NJ USA).

Research Institute for Fragrance Materials, Inc (1977). Human maximization test with Farnesol. RIFM report number 1702, 7 February (RIFM, Woodcliff Lake, NJ USA).

Research Institute for Fragrance Materials, Inc (2004). Local Lymph Node Assay on Farnesol. Unpublished report from Symrise GmbH & Co., Report number 47136 (RIFM, Woodcliff Lake, NJ USA).

Research Institute for Fragrance Materials, Inc (2004a). Local Lymph Node Assay on Farnesol. Unpublished report from Symrise GmbH & Co., Report number 47137 (RIFM, Woodcliff Lake, NJ USA).

Research Institute for Fragrance Materials, Inc (2004b). Repeated insult patch test of Farnesol in human subjects. Unpublished report from Symrise GmbH & Co., Report number 47190 (RIFM, Woodcliff Lake, NJ USA).

Fig leaf absolute

CAS N°:	68916-52-9	Empirical formula:	N/A
Synonyms:	Ficus carica absolute Fig leaf absolute (<i>Ficus carica</i>)		

History:	Initial reviews:	October 1980, May 1983	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION, PHOTOTOXICITY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke, D.L.J., Letizia, C. (1982), Food and Chemical Toxicology 20, 691

Furfural

CAS N°:	98-01-1	Empirical formula: Structure:	C5H4O2
Synonyms:	2-Formylfuran Fural Furaldehyde 2-Furaldehyde 2-Furancarbonal 2-Furancarboxaldehyde Furfural Furfuraldehyde α-Furfuraldehyde 2-Furylcarboxaldehyde Pyromucic aldehyde		

History:	Initial reviews:	New Standard	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Skin contact	0.001 %				
Non skin contact (1)	0.05 %	0.05 %			
Note box:					
These restrictions are not based on dermal sen	sitization QRA.				
(1) The non skin contact level should apply to the same product types as contained in QRA category 11, which includes all non-skin contact or incidental skin contact products.					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

See Annex I



CRITICAL EFFECT:

CARCINOGENICITY

REFERENCES:

This recommendation is based on fragrance industry data which were reviewed by the EU Scientific Committee on Consumer Safety at its 14th plenary meeting of March 27th, 2012 in the form of SCCS Opinion 1461/12.

Furfuryl alcohol

CAS N°:	98-00-0	Empirical formula: Structure:	C5H6O2
Synonyms:	2-Furancarbinol 2-Furanmethanol Furfuralcohol Furfuryl alcohol α-Furylcarbinol 2-Furylcarbinol 2-Furylmethanol 2-Hydroxymethylfuran		

History:	Initial reviews:	2009	
	Current revision date:	2015	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

Contributions from other sources like Coffee extracts or certain types of Cade oil have been evaluated. On the basis of the established maximum level of Furfuryl alcohol in these commercially available natural sources, exposure to this substance from the use of these oils and extracts is not significant and not regarded of concern from a consumer safety point of view.

See also the note on contributions from other sources in the Introduction to the IFRA Standards.

CRITICAL EFFECT:

INSUFFICIENT DATA

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

1) presence of structural alerts as defined in Api et al. (2014) and/or

2) adverse data on the material itself and/or

3) adverse data for a structurally related material.



REFERENCES:

Api A.M., Belsito D., Bruze M., Cadby P., Calow P., Dagli M. L., Dekant W., Dent M., Ellis G., Fryer A. D., Fukayama M., Griem P., Hickey C., Kromidas L., Lalko J., Liebler D.C., Miyachi Y., Politano V.T., Renskers K., Ritacco G., Salvito D., Schultz T.W., Sipes I. G., Smith B., Vitale D., Wilcox D.K. (2014). Food and Chemical Toxicology, http://dx.doi.org/10.1016/j.fct.2014.11.014.

Furfurylidene acetone

CAS N°:	623-15-4	Empirical formula: Structure:	C8H8O2
Synonyms:	3-Buten-2-one, 4-(2-furanyl)- Furfuralacetone 4-(2-Furyl)-3-buten-2-one		

History:	Initial reviews:	Not applicable		
	Current revision date:	Not applicable		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



Furfurylidene acetone

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or

2) adverse data on the material itself and/or

3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000



Geraniol



History:	Initial reviews:	New Standard		
	Current revision date:	2007		
	Implementation date:	For new submissions*: June 16, 2007		
		For existing fragrance compounds*:	June 16, 2009	
	Next review date	2012		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.3 %	Category 7	0.9 %		
Category 2	0.4 %	Category 8	2.0 %		
Category 3	1.8 %	Category 9	5.0 %		
Category 4	5.3 %	Category 10	2.5 %		
Category 5	2.8 %	Category 11	See Note Box (2)		
Category 6	8.6 %				

Note box:

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry). Further information about IOFI can be found on its website (http://www.iofiorg.org)

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.



Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
3525 [5]	Weak	11811	NA	NA	11800

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max tests

² Gerberick et al., 2001

³WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for geraniol and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 11800 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of geraniol in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Isola, D., Lalko, J., 2001a. Vehicle effects in the murine local lymph node assay (LLNA). American College of Toxicology Meeting, November 4-7. Washington DC.

Isola, D., Lalko, J., 2001b. Vehicle effects in the murine local lymph node assay (LLNA). International Journal of Toxicology, 20(6), 401.

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 2004a. Repeated Insult Patch Test on Geraniol. RIFM report number 46888, August 9a. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004b. Local Lymph Node Assay on Geraniol. RIFM report number 43812, December 17.

(RIFM, Woodcliff Lake, NJ, USA).



History:	Initial reviews:	November 2006 (41st Amendment)		
	Current revision date:	Not applicable		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products:	N/A				
Note box:					
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



Geranyl nitrile

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or

2) adverse data on the material itself and/or

3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000

Grapefruit oil expressed

CAS N°:	8016-20-4	Empirical formula: Structure :	N/A N/A
Synonyms:	N/A		

History:	Initial reviews:	June 1992	
	Current revision date:	2015	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	4%	Rinse-off products: No Restriction			
		Including household	cleaning products		
Non skin contact products:	No Restriction				
Note box:					
The Standard is set due to the phototoxic effect off products (please refer to Table 4 of the QRA	s of the material. The lin booklet for more detaile	nit only applies to applications or ad information).	n skin exposed to sunshine, excluding rinse-		
If combinations of phototoxic fragrance ingredie phototoxic fragrance ingredients, expressed in	ents are used, the use lev % of their recommended	vels have to be reduced accordin maximum level in the consume	ngly. The sum of the concentrations of all r product, shall not exceed 100.		
Note: See remark on phototoxic ingredients in the Introduction to the IFRA Standards (Appendix 8 to the IFRA Code of Practice) and the Standard on Citrus oil and other furocoumarins-containing essential oils.					
For qualities of the expressed oil in which the less volatile components have been concentrated by partial or total removal of the terpene fraction, this limit should be reduced in proportion to the degree of concentration.					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

PHOTOTOXICITY

REFERENCES:

These recommendations are made in order to promote good manufacturing practice (GMP) considering the large variations in the Bergapten content of commercial samples of grapefruit oil expressed (private communication to IFRA).



trans-2-Heptenal

CAS N°:	18829-55-5	Empirical formula: Structure:	C7H12O
Synonyms:	beta-Butylacrolein 3-Butylacrolein (E)-2-Hepten-1-al 2-Heptenal, (E)-		

History:	Initial reviews:	February 1985, April 1989	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Ford, R.A. (1988), Food and Chemical Toxicology 26, 331.



2-Heptylidene cyclopentan-1-one



History:	Initial reviews:	New Standard	
	Current revision date:	June 20, 2011	
	Implementation date:	: For new submissions*: August 20, 201	
		For existing fragrance compounds*:	August 20, 2012
	Next review date	2016	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Category 1 See Note box (1)	0.03%	Category 7	0.08%	
Category 2	0.04%	Category 8	1.01%	
Category 3	0.15%	Category 9	5.00%	
Category 4	0.45%	Category 10	2.50%	
Category 5	0.24%	Category 11	See Note Box (2)	
Category 6	0.72%			
Note box:				
 See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products with the potential of ingestion. Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. 				
Fragrance material specifications: N/A				

CONTRIBUTION FROM OTHER SOURCES:

None known at the time of the publication of the Standard

CRITICAL EFFECT:

SENSITIZATION

2-Heptylidene cyclopentan-1-one

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
4100 [1]	Weak	NA	NA	NA	1000

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003. ²Data derived from HRIPT or HMT. ³WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 2-Heptylidene cyclopentan-1-one and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1000 μ g/cm², which is a default value based on the LLNA data. They recommend the limits for the 11 different product categories, which are the acceptable use levels of 2-Heptylidene cyclopentan-1-one in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick G, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.



2,4-Hexadien-1-ol CAS N°: 111-28-4 17102-64-6 Empirical formula: Structure: C₆H₁₀O Synonyms: 1-Hydroxy-2,4-hexadiene Hexa-2,4-dien-1-ol Sorbic alcohol Sorbic alcohol

History:	Initial reviews:	New Standard		
	Current revision date:	2015		
	Implementation date:	e: For new submissions*: August 10, 2015		
		For existing fragrance compounds*:	August 10, 2016	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

Sorbyl alcohol

Hexadienol (commercial name)

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave-on products:	on products: N/A Rinse-off products: N/A				
	cleaning products				
Non-skin contact products: N/A					
Fragrance material specification	ons:	N/A			

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

INSUFFICIENT DATA



2,4-Hexadien-1-ol

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in Api et al. (2014) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material.

REFERENCES:

Api A.M., Belsito D., Bruze M., Cadby P., Calow P., Dagli M. L., Dekant W., Dent M., Ellis G., Fryer A. D., Fukayama M., Griem P., Hickey C., Kromidas L., Lalko J., Liebler D.C., Miyachi Y., Politano V.T., Renskers K., Ritacco G., Salvito D., Schultz T.W., Sipes I. G., Smith B., Vitale D., Wilcox D.K. (2014). Food and Chemical Toxicology, <u>http://dx.doi.org/10.1016/i.fct.2014.11.014</u>.

Hexahydrocoumarin

CAS N°:	700-82-3	Empirical formula: Structure:	C9H12O2
Synonyms:	2H-1-Benzopyran-2-one, 3,4,5,6,7,8-hexah Coumarin, hexahydro- Coumarin, 3,4,5,6,7,8-hexahydro- 1-Cyclohexene-1-propanoic acid, 2-hydrox 3,4,5,6,7,8-Hexahydro-2H-1-benzopyran-2	nydro- y-, d-lactone -one	

History:	Initial reviews:	February 1980	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Private communication to IFRA.

trans-2-Hexenal

CAS N°:	6728-26-3	Empirical formula: Structure:	C ₆ H ₁₀ O
Synonyms:	2-Hexenal, (E)- Hexen-2-al Leaf aldehyde beta-Propyl acrolein		

History:	Initial reviews:	April 1989, June 1992, May 2006, May 2007		
	Current revision date:	2008		
	Implementation date:	For new submissions*:	August 16, 2008	
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.001 %	Category 7	0.002 %		
Category 2	0.001 %	Category 8	0.002 %		
Category 3	0.002 %	Category 9	0.002 %		
Category 4	0.002 %	Category 10	0.002 %		
Category 5	0.002 %	Category 11	See Note Box (2)		
Category 6	0.02 %				
Note box:					
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again.					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry). Further information about IOFI can be found on its website (http://www.iofiorg.org)					
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

trans-2-Hexenal

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

trans-2-Hexenal - Sensitization Potency Estimation Based on Weight of Evidence

		Human Data			
LENA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	ication nimal Data ² NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
1012 [2]	Strong	24	NA	236	24

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max tests

² Gerberick et al., 2001

³WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for trans-2-hexenal and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as $24 \mu g/cm^2$. They recommend the limits for the 11 different product categories, which are the acceptable use levels of trans-2-hexenal in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Estrada, E., Patlewicz, G., Chamberlain, M., Basketter, D. and Larbey, S. (2003) Computer aided Knowledge Generation for Understanding Skin Sensitization Mechanisms: The TOPS-MODE Approach. Chem. Res. Toxicol., 16, 1226-1235

RIFM (Research Institute for Fragrance Materials, Inc.), 1989. Repeated Insult Patch Test on trans-2-Hexenal. RIFM report number 27821, May 22. (RIFM, Woodcliff Lake, NJ, USA).

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161. QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 1990. Repeated Insult Patch Test on trans-2-Hexenal. RIFM report number 27822, January 9b. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005a. Repeated Insult Patch Test on trans-2-Hexenal. RIFM report number 49111, July 14. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005b. Local Lymph Node Assay on trans-2-Hexenal. RIFM report number 48756, January 26. (RIFM, Woodcliff Lake, NJ, USA).

trans-2-Hexenal diethyl acetal

CAS N°:	67746-30-9	Empirical formula: Structure:	
Synonyms:	1,1-Diethoxy- (E) 2-Hexena 2-Hexene, 1,7	trans-2-hexene I diethyl acetal 1-diethoxy-, (2E)-	

History:	Initial reviews:	February 1985, April 1989	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Ford, R.A. (1988), Food and Chemical Toxicology 26, 345.



History:	Initial reviews:	February 1985, April 1989	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Ford, R.A. (1988), Food and Chemical Toxicology 26, 347.
alpha-Hexyl cinnamic aldehyde

CAS N°:	101-86-0	Empirical formula: Structure:	C ₁₅ H ₂₀ O
Synonyms:	2-Benzylideneoctanal Hexyl cinnamal α-Hexyl cinnamaldehyde Hexyl cinnamic aldehyde α-n-Hexylcinnamic aldehyde Hexyl cinnamyl α-n-Hexyl-β-phenylacrolein Octanal, 2-(phenylmethylene)- Jasmonal H (commercial name)		

History:	Initial reviews:	April 2007	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Category 1 See Note box (1)	0.7 %	Category 7	1.8 %	
Category 2	0.9 %	Category 8	2.0 %	
Category 3	3.6 %	Category 9	5.0 %	
Category 4	10.7 %	Category 10	2.5 %	
Category 5	5.6 %	Category 11	See Note box (2)	
Category 6 17.1 %				
Note box:				
(1) IFRA would recommend that any material us	(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of			

ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry www.iofiorg.org)

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

alpha-Hexyl cinnamic aldehyde

CONTRIBUTION FROM OTHER SOURCES:

See Annex II

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data				
LLNA weighted mean EC3 values (µg/cm ²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)	
2372 [>5]	Weak	23622	NA	NA	23600	

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹Data derived from HRIPT or Human Max tests ²Gerberick *et al.*, 2001 ³WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for alpha-hexyl cinnamic aldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 23600 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of alpha-hexyl cinnamic aldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api et al., 2008.

REFERENCES:

Basketter, D.A., Lea, L.J., Dickens, A., Briggs, D., Pate, I, Dearman, R.J., Kimber, I., 1999. A comparison of statistical approaches to the derivation of EC3 values from local lymph node assay dose responses. Journal of Applied Toxicology, 19(4), 261-266. Basketter, D.A., Wright, Z.M., Warbrick, E.V., Dearman, R.J., Kimber, I., Ryan, C.A., Gerberick, G.F., White, I.R., 2001. Human potency predictions for aldehydes using the local lymph node assay. Contact Dermatitis, 45(2), 89-94.

Basketter, D.A., Gilmour, N., Dearman, R.J., Kimber, I., Ryan, C.A., and Gerberick, F., 2003. Classification of skin sensitisation potency using the Local Lymph Node Assay. The Toxicologist, 72(S-1), 101.

Dearman, R.J., Hilton, J., Evans, P., Harvey, P., Basketter, D.A., Kimber, I., 1998. Temporal stability of local lymph node assay responses to hexyl cinnamic aldehyde. Journal of Applied Toxicology, 18(4), 281-284.

Dearman, R.J., Wright, Z.M., Basketter, D.A., Ryan, C.A., Gerberick, G.F., Kimber, I, 2001. The suitability of hexyl cinnamic aldehyde as a calibrant for the murine local lymph node assay. Contact Dermatitis, 44(6), 357-361.

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Repeated Insult Patch Test on alpha-hexylcinnamaldehyde. RIFM report number 51047, November 11. (RIFM, Woodcliff Lake, NJ, USA).



alpha-Hexylidene cyclopentanone



History:	Initial reviews:	May 1983, July 1994	
	Current revision date:	2008	
	Implementation date:	For new submissions*:	August 16, 2008
		For existing fragrance compounds*:	August 16, 2010
	Next review date	2013	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Category 1 See Note box (1)	0.01 %	Category 7	0.02 %	
Category 2	0.01 %	Category 8	0.06 %	
Category 3	0.05 %	Category 9	0.06 %	
Category 4	0.06 %	Category 10	0.06 %	
Category 5	0.06 %	Category 11	See Note Box (2)	
Category 6	0.2 %			
Note box:				
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again.				
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry). Further information about IOFI can be found on its website (http://www.iofiorg.org)				
(2) Category 11 includes all non-skin contact or there is no justification for a restriction of the co	incidental skin contact p ncentration of this fragra	roducts. Due to the negligible skin contact normalized ingredient in the finished product.	t from these types of products	

N/A

Fragrance material specifications:

alpha-Hexylidene cyclopentanone

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

			Human Data		
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
600	Weak	300 4	NA	500	300

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for alpha-Hexylidene cyclopentanone and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 300 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of alpha-Hexylidene cyclopentanone in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1982. Human Repeated Insult Patch Test. Unpublished report from IFF Incorporated, 8 January and 27 August. Report number 15002 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Human Repeated Insult Patch Test. Unpublished report from Firmenich Incorporated, 17 March. Report number 48712 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2008. Local Lymph Node Assay. RIFM report (RIFM, Woodcliff Lake, NJ, USA).



History:	Initial reviews:	New Standard	
	Current revision date:	2007	
	Implementation date:	For new submissions*:	June 16, 2007
		For existing fragrance compounds*:	June 16, 2009
	Next review date	2012	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	1.0 %	Category 7	2.7 %		
Category 2	1.3 %	Category 8	2.0 %		
Category 3	5.3 %	Category 9	5.0 %		
Category 4	16.0 %	Category 10	2.5 %		
Category 5	8.4 %	Category 11	See Note Box (2)		
Category 6	25.7 %				
Note box:					
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry). Further information about IOFI can be found on its website (http://www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification of the concentration of this fragrance ingradient in the finished product. 					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

Hexyl salicylate

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

Hexyl salicylate - Sensitization Potency Estimation Based on Weight of Evidence

		i	Human Data		
LLNA weighted mean EC3 values (μg/cm ²) [no. studies] Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)	
45 [1]	Weak	35433	2069	NA	35400

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max tests

² Gerberick et al., 2001

³WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for hexyl salicylate and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 35400 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of hexyl salicylate in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 1975. Human Maximization Test on Hexyl Salicylate. RIFM report number 1798, January 31. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Repeated Insult Patch Test on Hexyl Salicylate. RIFM report number 45130, May 3. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2006. Local Lymph Node Assay on Hexyl Salicylate. RIFM report number 51636, March 29a. (RIFM, Woodcliff Lake, NJ, USA).

Hydroabietyl alcohol, Dihydroabietyl alcohol

CAS N°:	13393-93-6 26266-77-3 1333-89-7	Empirical formula:	C ₂₀ H ₃₆ O
Synonyms:	Abitol (mixture of different hydroabietyl	alcohols)	

History:	Initial reviews:	October 1974, May 1976, December 19	95
	Current revision date:	May 2003	
	Implementation date:	For new submissions*:	May 6, 2004
		For existing fragrance compounds*:	May 6, 2005
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products:	N/A				
Note box:					
So far only the use in cosmetic products was banned. This ban is now extended to all kinds of fragrance application (use).					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

None Known.

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted - REXPAN, May 2003

REFERENCES:

RIFM Monograph 323, Fd. Cosmet. Toxicol. 12, 919-921 (1974)

Hydroquinone monoethyl ether

CAS N°:	622-62-8	Empirical formula: Structure:	C ₈ H ₁₀ O ₂
Synonyms:	1-Ethoxy-4-hydroxybenzene p-Ethoxyphenol Phenol, 4-ethoxy-		

History:	Initial reviews:	June 1982, October 1983	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	
* This date applies	to the supply of fragrance com	pounds (formulas) only, not to the finished product	s in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:	Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A			
	Including household cleaning products					
Non skin contact products:	N/A					
Note box:						
So far only the use in cosmetic products was banned. This ban is now extended to all kinds of fragrance application (use).						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

DEPIGMENTATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted - REXPAN, September 2002

REFERENCES:

E. Frenk, (1969), Arch. Klin. Exp. Derm. 235, 16.

E. Frenk (1970), Ann. Derm. Syph (Paris) 97, 287.

E. Frenk & F. Ott (1971), Journal of Investigative Dermatology 56, 287.



Hydroquinone monoethyl ether

W. Wohlrab and R.P. Zaumseil (1976), Derm. Monatsschr. 162, 908.

Private communication to IFRA.

Hydroquinone monomethyl ether

CAS N°:	150-76-5	Empirical formula: Structure:	C7H8O2
Synonyms:	4-Hydroxyanisole p-Hydroxyanisole 4-Methoxyphenol p-Methoxyphenol Phenol, p-methoxy-		

History:	Initial reviews:	October 1983	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

DEPIGMENTATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Private communication to IFRA.



Hydroxycitronellal

CAS N°:	107-75-5	Empirical formula: Structure:	C ₁₀ H ₂₀ O ₂ OH
Synonyms:	Citronellalhydrate 3,7-Dimethyl-7-hydroxyoct Octanal, 7-hydroxy-3,7-dir Oxydihydrocitronellal Laurinal, Laurine	anal nethyl-	

History:	Initial reviews:	March 1987, September 2000, April 2005, May 2007, June 2008		
	Current revision date:	June 2013		
	Implementation date:	For new submissions*:	August 10, 2013	
		For existing fragrance compounds*:	August 10, 2014	
	Next review date	2018		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.1 %	Category 7	0.4 %		
Category 2	0.2 %	Category 8	1.0 %		
Category 3	0.8 %	Category 9	1.0 %		
Category 4	1.0 %	Category 10	1.0 %		
Category 5	1.0 %	Category 11	See Note box (2)		
Category 6	3.6 %				

Note box:

For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again.

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry). (http://www.iofiorg.org)

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

See Annex II

Hydroxycitronellal

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
5612 [9]	Weak	5000 ⁴	NA	5906	5000

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴ MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Hydroxycitronellal and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 5000 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Hydroxycitronellal in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Ashby, J., Basketter, D. A., Paton, D., Kimber, I., 1995. Structure activity relationships in skin sensitization using the murine local lymph node assay. *Toxicology*, 103(3), 177-194. Basketter, D. A., Wright, Z., Warbrick, E. V., Dearman, R. J., Kimber, I., Ryan, C. A.,

Gerberick, G. F., White, I. R., 2001. Comparison of the local lymph node assay with the guinea-pig maximization test for the detection of a range of contact allergens. *Food and Chemical Toxicology*, 30(1), 65-69.

Basketter, D. A., Wright, Z., Gilmour, N. J., Ryan, C. A., Gerberick, G. F., Robinson, M. K., Dearman, R. J., Kimber, I., 2002. Prediction of human sensitization potency using local lymph node assay EC3 values. *Contact Dermatitis*, 45(2), 89-94.

Estrada, E., Patlewicz, G., Chamberlain, M., Basketter, D., Larbey, S., 2003. Computeraided knowledge generation for understanding skin sensitization mechanisms: The TOPSMODE approach. *Chemical Research in Toxicology*, 16(10), 1226-1235.

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. *American Journal of Contact Dermatitis*, 12(3), 156-161.

Isola, D., Lalko, J., 2001. Vehicle effects in the murine local lymph node assay (LLNA). *International Journal of Toxicology*, 20(6), 401

RIFM (Research Institute for Fragrance Materials, Inc.), 1990. Repeated insult patch testwith hydroxycitronellal in human subjects. RIFM report number 28267, October, 24. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1991. Repeated insult patch test with hydroxycitronellal in human subjects. RIFM report number 28273, April, 17. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Material, Inc.), 2006. Repeated insult patch test with hydroxycitronellal. RIFM report number 49736, January 5. (RIFM, Woodcliff Lake, NJ, USA).

Hydroxycitronellal

Smith, C. K., Hotchkiss, S. A. M., 2001. Allergic Contact Dermatitis: Chemical and Metabolic Mechanisms. Taylor and Francis Ltd, London.

3 and 4-(4-Hydroxy-4-methylpentyl)-3-cyclohexene-1-carboxaldehyde (HMPCC)

CAS N°:	31906-04-4 51414-25-6	Empirical formula:	C ₁₃ H ₂₂ O ₂
Synonyms:	3-Cyclohexen-1-carboxaldehyde, 4 3-Cyclohexen-1-carboxaldehyde, 5 Hydroxyisohexyl 3-cyclohexene ca 4-(4-Hydroxy-4-methylpenyl) cyclo 3-(4-Hydroxy-4-methylpentyl) cyclo Lyral, Kovanol, Mugonal, Landola	4-(4-hydroxy-4-methylpentyl)- 3-(4-hydroxy-4-methylpentyl)- arboxaldehyde (INCI) hex-3-enecarbaldehyde bhex-3-ene-1-carbaldehyde , Cyclohexal (trade names)	

History:	Initial reviews:	April 2003, July 2008, October 2009	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.02 %	Category 7	0.02 %		
Category 2	0.02 %	Category 8	0.2 %		
Category 3	0.2 %	Category 9	0.2 %		
Category 4	0.2 %	Category 10	0.2 %		
Category 5	0.2 %	Category 11	See Note box (2)		
Category 6	0.2 %				
Note box:					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org). (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the peoligible skin contact from these types of products.					

there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. The restrictions as given for the individual categories are NOT based on the QRA but solely represent a pragmatic approach to address the specific

situation for HMPCC. These pragmatic measures will be reviewed over time in view of the development of sensitization rates (clinical data) to the material.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

See Annex II

3 and 4-(4-Hydroxy-4-methylpentyl)-3-cyclohexene-1-carboxaldehyde (HMPCC)

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

See Note Box

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel approved the pragmatic approach proposed by IFRA.



p-lsobutyl-alpha-methyl hydrocinnamaldehyde

CAS N°:	6658-48-6	Empirical formula: Structure:	C14H20O
Synonyms:	p-IsobutyI-α-methyl hydro cinna Benzenepropanal, α-methyl-4-(3-(4-IsobutyI-phenyI)-2-methyl-p Rhodial Suzaral	mic aldehyde 2-methylpropyl)- propionaldehyde	

History:	Initial reviews:	New Standard	
	Current revision date:	2009	
	Implementation date:	For new submissions*:	August 7, 2009
		For existing fragrance compounds*:	August 7, 2011
	Next review date	2014	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.07 %	Category 7	0.17 %		
Category 2	0.08 %	Category 8	2.00 %		
Category 3	0.35 %	Category 9	5.00 %		
Category 4	1.04 %	Category 10	2.50 %		
Category 5	0.55 %	Category 11	See Note Box (2)		
Category 6	1.67 %				
Note box:					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/					
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.					
Fragrance material specificatio	ns:	N/A			

p-lsobutyl-alpha-methyl hydrocinnamaldehyde

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

			Human Data		
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
<2500 [1]4	Weak	2362	5520	NA	2300

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed

effect level; NA = Not Available.

¹ Data derived from HRIPT or HMT.

² Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

³WoE NESIL limited to two significant figures.

⁴EC3 value from one LLNA, not the mean.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for *p*-lsobutyl- α -methyl hydrocinnamaldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 2300 mg/cm2. They recommend the limits for the 11 different product categories, which are the acceptable use levels of *p*-lsobutyl- α methyl hydrocinnamaldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1977. Maximization study with *p*-Isobutyl-α-methyl hydrocinnamaldehyde. RIFM report number 1702, May 9c. (RIFM,

Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001. Local Lymph Node Assay. Unpublished study from Givaudan, 18 May. Report number 41055. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2008. Human repeated insult patch test. RIFM report number 55563, August 25a. (RIFM, Woodcliff Lake, NJ, USA).

Isobutyl N-methylanthranylate



History:	Initial reviews:	New Standard	
	Current revision date:	2009	
	Implementation date:	For new submissions*:	August 7, 2009
		For existing fragrance compounds*:	August 7, 2010
	Next review date	2014	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
	Including household cleaning products					
Non skin contact products:	N/A					
Note box:						
The material has been identified for having the potential of forming nitrosamines in nitrosating systems. Downstream users therefore have to be notified of the presence of the material and its potential to be able to consider adequate protective measures.						
Fragrance material specification	ons:	N/A				

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)

CRITICAL EFFECT:

POTENTIAL OF NITROSAMINE FORMATION

REXPAN RATIONALE / CONCLUSION:

IFRA measures regarding potential nitrosamine formation noted - REXPAN April 2009.

REFERENCES:

Nitrosamine policy as contained in the EU Cosmetics Directive 76/768/EEC and its Amendments.

		Isocyclocitra	d	
CAS N°:	1335-66-6 1423-46-7 67634-07-5	Empirical formula: Structure:	C ₁₀ H ₁₆ O 1335-66-6:	. 0
Synonyms:	1335-66-6 1-Formyl-[2,4,6-]&[3,5,6-]tr Isocyclocitral [2,4,6-]&[3,5,6-]Trimethyl-3 1423-46-7 3-Cyclohexene-1-carboxal Neocyclocitral 2 4 6-Trimethylcyclohex-3-	imethyl-3-cyclohexer 3-cyclohexene-1-carb dehyde, 2,4,6-trimeth enecarbaldehyde	ne oxaldehyde nyl-	
	2,4,6-Trimethyl-3-cyclohex 2,4,6-Trimethyl-3-cyclohex 67634-07-5 3-Cyclohexene-1-carboxal 3,5,6-Trimethylcyclohex-3-	enylcarboxaldehyde ene-1-carbaldehyde dehyde, 3,5,6-trimeth ene-1-carbaldehyde	ıyl-	

History:	Initial reviews:	New Standard		
	Current revision date:	2007		
	Implementation date:	For new submissions*:	June 16, 2007	
		For existing fragrance compounds*:	June 16, 2009	
	Next review date	2012		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

Isocyclocitral

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.2 %	Category 7	0.5 %		
Category 2	0.3 %	Category 8	2.0 %		
Category 3	1.1 %	Category 9	5.0 %		
Category 4	3.2 %	Category 10	2.5 %		
Category 5	1.7 %	Category 11	See Note Box (2)		
Category 6	5.1 %				

Note box:

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

Isocyclocitral - Sensitization Potency Estimation Based on Weight of Evidence

			Human Data			
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)	
1825 [1]	Weak	7087	2759	NA	7000	

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max tests

² Gerberick *et al.*, 2001

³WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for isocyclocitral and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 7000 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of isocyclocitral in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.



Isocyclocitral

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 1972. The contact-sensitization potential of fragrance materials by maximization testing in humans. RIFM report number 1804, November 01 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Repeated insult patch test with isocyclocitral. RIFM report number 47260, April 16 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Repeated insult patch test with isocyclocitral. RIFM report number 47590, January 24 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007. Murine local lymph node assay on isocyclocitral. RIFM DRAFT (RIFM, Woodcliff Lake, NJ, USA).



Isocyclogeraniol CAS N°: 68527-77-5 Empirical formula: Structure: $C_{10}H_{18}O$ Synonyms: 3-Cyclohexene-1-methanol, 2,4,6-trimethyl-2,4,6-Trimethyl-3-cyclohexene-1-methanol

History:	Initial reviews:	December 1995, April 2005		
	Current revision date:	2008		
	Implementation date:	For new submissions*: August 16, 2008		
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.11 %	Category 7	0.3 %		
Category 2	0.14 %	Category 8	0.5 %		
Category 3	0.5 %	Category 9	0.5 %		
Category 4	0.5 %	Category 10	0.5 %		
Category 5	0.5 %	Category 11	See Note Box (2)		
Category 6	2.8 %				
Note box:					
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again.					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/					
(2) Category 11 includes all non-skin contact or there is no justification for a restriction of the co	ncentration of this fragra	products. Due to the negligible skin contac nce ingredient in the finished product.	ct from these types of products		

Fragrance material specifications:

N/A

Isocyclogeraniol

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

LINA weighted mean EC2 values		Human Data				
LINA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)	
>6250	Weak	3898 ⁴	NA	7752	3800	

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Isocyclogeraniol and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 3800 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Isocyclogeraniol in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1982. Evaluation of potential irritation and sensitization hazards by dermal contact of isocyclogeraniol in humans. Unpublished report from International Flavors and Fragrances, 22 July. Report number 21790 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1983. Evaluation of potential hazards by dermal contact of isocyclogeraniol in humans. Unpublished report from International Flavors and Fragrances, 06 January. Report number 21792 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. 2,4,6-Trimethyl-3cyclohexene-1-methanol diluted with vehicle 1:3 EtOH:DEP: Local Lymph Node Assay. RIFM report number 48755, January 28 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005a. Repeated insult patch test with 2,4,6-trimethyl-3-cyclohexene-1-methanol. RIFM report number 49110, September 07 (RIFM, Woodcliff Lake, NJ, USA).



CAS N°:	97-54-1	Empirical formula: Structure:	C10H12O2
Synonyms:	1-Hydroxy-2-methoxy-4-propen-1-ylb 4-Hydroxy-3-methoxy-1-propen-1-ylb 4-Hydroxy-3-methoxy-1-propenylben iso-Eugenol 3-Methoxy-4-hydroxy-1-propen-1-ylbo 2-Methoxy-4-propenylphenol 2-Methoxy-4-(1-propenyl)phenol Phenol, 2-methoxy-4-(1-propenyl)- 4-Propenylguaiacol	enzene enzene zene enzene	

History:	Initial reviews:	May 1980, May 1998, 2001, April 2004, May 2007		
	Current revision date:	2008		
	Implementation date:	For new submissions*:	August 16, 2008	
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.01 %	Category 7	0.02 %			
Category 2	0.01 %	Category 8	0.02 %			
Category 3	0.02 %	Category 9	0.02 %			
Category 4	0.02 %	Category 10	0.02 %			
Category 5	0.02 %	Category 11	See Note Box (2)			
Category 6	0.2 %					
Note box:						

For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again.

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/

Isoeugenol

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
498 [18]	Moderate	250 ⁴	NA	775	250

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴ MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Isoeugenol and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 250 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Isoeugenol in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Basketter, D. A., Lea, L. J., Dickens, A., Briggs, D., Pate, I., Dearman, R. J.,Kimber, I., 1999. A comparison of statistical approaches to the derivation of EC3 values from local lymph node assay dose responses. *Journal of Applied Toxicology*, 19(4), 261-266. Basketter, D. A., Wright, Z., Gilmour, N. J., Ryan, C. A., Gerberick, G. F., Robinson, M. K.,Dearman, R. J.,Kimber, I., 2002. Prediction of human sensitization potency using local lymph node assay EC3 values. *The Toxicologist*, 66(1-S), 240

Loveless, S. E., Ladics, G. S., Gerberick, G. F., Ryan, C. A., Basketter, D. A., Scholes, E. W., House, R. V., Hilton, J., Dearman, R. J., Kimber, I., 1996. Further evaluation of the local lymph node assay in the final phase of an international collaborative trial. *Toxicology*, 108 (1-2), 141-152

RIFM (Research Institute for Fragrance Materials, Inc.), 1991. Murine local lymph node assay on isoeugenol. Unpublished report from Firmenich Incorporated, 24 August. Report number 40676 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001. Murine local lymph node assay with methoxy dicyclopentadiene carboxaldehyde. Unpublished report from Firmenich Incorporated, 08 October. Report number 42120 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001a. Murine local lymph node assay with 3-cyclopentene-1-butanal, alpha,2,2,3-tetramethyl-.gamma.-methylene. Unpublished report from Firmenich Incorporated, 30 November. Report number 42122 (RIFM, Woodcliff Lake, NJ, USA).



RIFM (Research Institute for Fragrance Materials, Inc.), 2001b. 4-Penten-2-ol, 3,3dimethyl-5-(2,2,3-trimethyl-3-cyclopenten-1-yl)-: Murine local lymph node assay. Unpublished report from Firmenich Incorporated, 11 November. Report Number 42130 (RIFM, Woodcliff Lake, NJ, USA)

RIFM (Research Institute for Fragrance Materials, Inc.), 2002. Murine local lymph node assay with methyl hexadecanoate. Unpublished report from Firmenich Incorporated, 22 January. Report number 42123 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2002a. Murine local lymph node assay of delta-1-(2,6,6-trimethyl-3-cyclohexen-1-yl)-2-buten-1-one. Unpublished report from Firmenich Incorporated, 02 April. Report number 42139 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2002b. 2(3H)-Naphthalenone, 4,4a,5,6,7,8-hexahydro-4,4a-dimethyl-6-(1-methylethylidene)-, (4R,4aS)-: Murine Local Lymph Node Assay. Unpublished report from Firmenich Incorporated, 31 January. Report number 42145 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2002c. Methyl octanoate: Murine lymph node assay. Unpublished report from Firmenich Incorporated, 12 September. Report number 42131 (RIFM, Woodcliff Lake, NJ, USA).

Thompson, G. R., Booman, K. A., Dorsky, J., Kohrman, K. A., Rothenstein, A. S., Schwoeppe, E. A., Sedlak, R. I., Steltenkamp, R. J., 1983. Isoeugenol: A survey of consumer patch-test sensitization. Food and Chemical Toxicology, 21(6), 735-740.

Wright, Z. M., Basketter, D. A., Blaikie, L., Cooper, K. J., Warbrick, E. V., Dearman, R. J., Kimber, I., 2001. Vehicle effects on skin sensitization potency of four chemicals assessment using the local lymph node assay. *International Journal of Cosmetic Science*, 23(2), 75-83.



Isophorone

CAS N°:	78-59-1	Empirical formula: Structure:	C9H14O
Synonyms:	2-Cyclohexen-1-one, 3,5,5-trime Isoacetophorone 3,5,5-Trimethyl-2-cyclohexen-1-	ethyl- one	

History:	Initial reviews:	Not applicable		
	Current revision date:	Not applicable		
	Implementation date:	For new submissions*:	Not applicable	
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		
* This date applies	to the supply of fragrance com	pounds (formulas) only, not to the finished product	s in the marketplace.	

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
	Including household cleaning products				
Non skin contact products:	N/A				
Note box:					
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



Isophorone

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or

3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000

6-Isopropyl-2-decalol

CAS N°:	34131-99-2	Empirical formula: Structure:	C ₁₃ H ₂₄ O OH
Synonyms:	Decahydro-6-is Decahydro-6-(6-Isopropyl-2-c 6-Isopropyldec 2-Naphthaleno Decatol	sopropyl-2-naphthol 1-methylethyl)-2-naph lecahydronaphthalen alol I, decahydro-6-(1-me	nthalenol ol thylethyl)-

History:	Initial reviews:	June 1979, April 1989	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Ford, R.A., (1988), Food and Chemical Toxicology 26, 367.

Jasmine absolute (grandiflorum)

CAS N°:	8022-96-6 8024-43-9 90045-94-6 84776-64-7	Empirical formula:	N/A
Synonyms:	Jasmine absolute (<i>Jasminum grand</i> Jasminum grandiflorum absolute Jasmin officinale var. grandiflorum	iflorum L.)	

History:	Initial reviews:	New Standard		
	Current revision date:	2008		
	Implementation date:	For new submissions*: August 16, 2008		
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.04 %	Category 7	0.1 %		
Category 2	0.05 %	Category 8	1.5 %		
Category 3	0.22 %	Category 9	5.0 %		
Category 4	0.7 %	Category 10	2.5 %		
Category 5	0.4 %	Category 11	See Note Box (2)		
Category 6	1.1 %				
Note box:					
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/ (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is an intificiant of the contact from these types of products. 					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SENSITIZATION

Jasmine absolute (grandiflorum)

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
1475	Weak	1475 ⁴	NA	2069	1470

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴ MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Jasmine Absolute Grandiflorum and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1470 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Jasmine Absolute Grandiflorum in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1972. Human Maximization Test. RIFM report number 1804, November 1c (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1977. Human Maximization Test. RIFM report number 1702, May 4a (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1980. Human Maximization Test. RIFM report number 1791, November 6 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2006. Local Lymph Node Assay. RIFM report number 53024, May 19 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007. Human Repeated Insult Patch Test. RIFM report number 52893, May 18 (RIFM, Woodcliff Lake, NJ, USA).

Jasmine absolute (sambac)

CAS N°:	91770-14-8	Empirical formula:	N/A
Synonyms:	Jasmin sambac extract		

History:	Initial reviews:	New Standard		
	Current revision date:	2008		
	Implementation date:	For new submissions*: August 16, 2008		
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.25 %	Category 7	0.7 %		
Category 2	0.32 %	Category 8	2.0 %		
Category 3	1.33 %	Category 9	5.0 %		
Category 4	4.0 %	Category 10	2.5 %		
Category 5	2.1 %	Category 11	See Note Box (2)		
Category 6	6.4 %				
Note box:					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/					
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.					
Fragrance material specifications: N/A					
CONTRIBUTION FROM OTHER SOURCES:					

N/A

CRITICAL EFFECT:

SENSITIZATION

Jasmine absolute (sambac)

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
9100	Weak	8858 ⁴	NA	NA	8850

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴ MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Jasmine absolute Sambac and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 8850 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Jasmine absolute Sambac in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 2006. Local Lymph Node Assay. RIFM report number 52885, November 16 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007. Human Repeated Insult Patch Test. RIFM report number 52897, May 15 (RIFM, Woodcliff Lake, NJ, USA).

Lemon oil cold pressed

CAS N°:	8008-56-8	Empirical formula: Structure :	N/A N/A
Synonyms:	N/A		

History:	Initial reviews:	October 1975, June 1992	
	Current revision date:	2015	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	2%	Rinse-off products:	No Restriction			
		Including household	cleaning products			
Non skin contact products:	No Restriction					
Note box:						
The Standard is set due to the phototoxic effects of the material. The limit only applies to applications on skin exposed to sunshine, excluding rinse- off products (please refer to Table 4 of the QRA booklet for more detailed information).						
If combinations of phototoxic fragrance ingredients are used, the use levels have to be reduced accordingly. The sum of the concentrations of all phototoxic fragrance ingredients, expressed in % of their recommended maximum level in the consumer product, shall not exceed 100.						
Note: See remark on phototoxic ingredients in the Introduction to the IFRA Standards (Appendix 8 to the IFRA Code of Practice) and the Standard on Citrus oil and other furocoumarins-containing essential oils.						
For qualities of the expressed oil in which the less volatile components have been concentrated by partial or total removal of the terpene fraction, this limit should be reduced in proportion to the degree of concentration.						
Fragrance material specificatio	Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

PHOTOTOXICITY

Lemon oil cold pressed

REFERENCES:

These recommendations are based on results of RIFM on the phototoxicity of lemon oil cold pressed (Fd. Cosm. Toxicol. 12,725(1974), its low bergapten content (C.K. Shu et al. VI Int. Congress of Essential oils 1974) and the observed no-effect level of pooled samples in tests using the animal model (private communication to IFRA).

8th Amendment

Lime oil expressed

CAS N°:	8008-26-2	Empirical formula: Structure :	N/A N/A
Synonyms:	N/A		

History:	Initial reviews:	October 1975, June 1992	
	Current revision date:	2015	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	0.7%	Rinse-off products:	No Restriction	
		Including household cleaning products		
Non skin contact products:	No Restriction	n		
Note box:				
The Standard is set due to the phototoxic effects of the material. The limit only applies to applications on skin exposed to sunshine, excluding rinse- off products (please refer to Table 4 of the QRA booklet for more detailed information).				
If combinations of phototoxic fragrance ingredients are used, the use levels have to be reduced accordingly. The sum of the concentrations of all phototoxic fragrance ingredients, expressed in % of their recommended maximum level in the consumer product, shall not exceed 100.				
Note: See remark on phototoxic ingredients in the Introduction to the IFRA Standards (Appendix 8 to the IFRA Code of Practice) and the Standard on Citrus oil and other furocoumarins-containing essential oils.				
For qualities of the expressed oil in which the less volatile components have been concentrated by partial or total removal of the terpene fraction, this limit should be reduced in proportion to the degree of concentration.				
Fragrance material specifications:		N/A		
CONTRIBUTION FROM OTHER SOURCES:				

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

ΡΗΟΤΟΤΟΧΙCITY
\bigcirc

Lime oil expressed

REFERENCES:

These recommendations are based on results of RIFM on the phototoxicity of lime oil expressed (Fd. Cosm. Toxicol. 12,731(1974), its bergapten content reported in J.A.O.A.C.52,(4),727(1969) and the observed no-effect level of pooled samples in tests using the animal model (private communication to IFRA).



CAS N°: 138-86-3 Synonyms:

History:	Initial reviews:	December 1995	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

SPECIFICATION

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	N/A	Rinse-off products:	N/A	
		Including household	cleaning products	
Non skin contact products:	N/A			
Note box:				
Fragrance material specifications:		<i>d</i> -, <i>I</i> -and <i>dI</i> -Limonene containing substantia used when the level of lowest practical level, antioxidants at the tim products should have 20 millimoles peroxid according to the FMA downloaded from the <u>Methods</u>).	and natural products amounts of it, should only be of peroxides is kept to the , for instance by adding ne of production. Such a peroxide value of less than les per liter, determined a method, which can be IFRA website (see <u>Analytical</u>	



Limonene

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SENSITIZATION

REFERENCES:

This recommendation is based on the published literature, mentioning sensitizing properties when containing peroxides and oxidation products. (D.L.J. Opdyke(1975), Fd. Cosmet. Toxicol. 13,825).

Linalool

CAS N°:	78-70-6 126-90-9 (d-linalool) 126-91-0 (l-linalool)	Empirical formula: Structure:	
Synonyms:	Coriandrol 2,6-Dimethyl-2,7-octadien-6-ol 3,7-Dimethyl-1,6-octadien-3-ol Licareol Linalol Linalyl alcohol 1,6-Octadien-3-ol, 3,7-dimethyl (CA	S)	

History:	Initial reviews:	New Standard	
	Current revision date:	November 2003	
	Implementation date:	For new submissions*:	May 6, 2004
		For existing fragrance compounds*:	May 6, 2005
	Next review date	November 2008	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

SPECIFICATION

RESTRICTIONS:

Limits in the finished product:			
Skin contact products:			
Leave on products:	N/A	Rinse-off products:	N/A
		Including household	cleaning products
Non skin contact products:	N/A		
Note box:			
Fragrance material specifications:		Limit peroxide level to 20 mmol/l.	
		Linalool and natural produ as bois de rose, coriander when the level of peroxide is recommended to add ar the raw material. The addi for example has shown gru level for products in use sh The (hydro) peroxide conte FMA method.	cts known to be rich in linalool, such or ho wood oil, should only be used s is kept to the lowest practical level. It tioxidants at the time of production of tion of 0.1% BHT or alpha-tocopherol eat efficiency. The maximum peroxide nould be 20 mmol/l. ent can be determined by using the

Linalool

CONTRIBUTION FROM OTHER SOURCES:

See fragrance material specification

CRITICAL EFFECT:

SENSITIZATION*

*Pure linalool is not a sensitizer while hydroperoxides and other oxidation products have shown sensitizing properties.

One of the major oxidation products of linalool was isolated and identified as 7-hydroperoxy-3,7-dimethyl-octa-1,5-diene-3-ol. In sensitization studies in guinea pigs, linalool of high purity gave no reactions, while linalool that had been oxidized for 10 weeks sensitized the animals. It was concluded that autoxidation of linalool is essential for its sensitizing potential (Skold et al., 2002).

REXPAN RATIONALE / CONCLUSION:

The GMP recommended by IFRA have been noted and approved by REXPAN, (November 17, 2003).

REFERENCES:

M.Skold, A.Borje, M.Matura and A.-T.Karlberg., 2002. Studies on the autoxidation and sensitizing capacity of the fragrance chemical linalool, identifying a linalool hyperperoxide.

Contact Dermatitis, 46(5), 267-272.

M.Skold, A.Borje, M.Matura and A.-T.Karlberg., 2002. Sensitization studies on the fragrance chemical linalool, with respect to auto-oxidation. Contact Dermatitis, 46 (Suppl. 4), 20.

Massoia bark oil

CAS N°:	85085-26-3	Empirical formula:	N/A
Synonyms:	Cryptocarya massoio oil Cryptocarya massoy bark extract Cryptocarya massoy, ext. Massoia bark oil (<i>Cryptocarya massoio</i>)		

History:	Initial reviews:	Not applicable	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	N/A	Rinse-off products:	N/A	
	Including household cleaning products			
Non skin contact products:	N/A			
Note box:				
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.				
Fragrance material specifications:		N/A		

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



Massoia bark oil

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000



History:	Initial reviews:	2008	
	Current revision date:	2015	
	Implementation date:	For new submissions*: August 10, 2015	
		For existing fragrance compounds*:	August 10, 2016
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave-on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non-skin contact products:	N/A				
Fragrance material specifications: N/A					
CONTRIBUTION FROM OTHER SOURCES:					

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

INSUFFICIENT DATA



Massoia lactone

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in Api et al. (2014) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material.

REFERENCES:

Api A.M., Belsito D., Bruze M., Cadby P., Calow P., Dagli M. L., Dekant W., Dent M., Ellis G., Fryer A. D., Fukayama M., Griem P., Hickey C., Kromidas L., Lalko J., Liebler D.C., Miyachi Y., Politano V.T., Renskers K., Ritacco G., Salvito D., Schultz T.W., Sipes I. G., Smith B., Vitale D., Wilcox D.K. (2014). Food and Chemical Toxicology, http://dx.doi.org/10.1016/j.fct.2014.11.014.

Melissa oil (genuine Melissa officinalis L.)

CAS N°:	8014-71-9 84082-61-1	Empirical formula:	N/A
Synonyms:	Balm oil (Melissa officinalis L.) Lemon balm oil Melissa officinalis leaf oil Melissa oil (Melissa officinalis L.) Oil of balm		

History:	Initial reviews:	July 2008 (43th Amendment)	
	Current revision date:	2009	
	Implementation date:	For new submissions*:	August 7, 2009
		For existing fragrance compounds*:	August 7, 2011
	Next review date	2014	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.04 %	Category 7	0.11 %			
Category 2	0.05 %	Category 8	1.40 %			
Category 3	0.21 %	Category 9	5.00 %			
Category 4	0.63 %	Category 10	2.50 %			
Category 5	0.33 %	Category 11	See Note Box (2)			
Category 6	1.01 %					
Note box:						
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/						
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SENSITIZATION

Melissa oil (genuine Melissa officinalis L.)

RIFM SUMMARIES:

		Human Data				
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)	
4500 [1] ⁵	Weak	1470 ⁴	NA	NA	1400	

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or HMT

²Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

³WoE NESIL limited to two significant figures

⁴MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL ⁵EC3 value from one LLNA, not the mean.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Melissa oil and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1400 mg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Melissa oil in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 2001. Human repeated insult patch test. Unpublished study from Robertet, 21 February. Report number 36641. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2008a. Local Lymph Node Assay. Unpublished study from Robertet. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2008b. Human repeated insult patch test. Unpublished study from Robertet. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2009 Repeated insult patch test on melissa oil in humans. Unpublished summary report from Robertet, June 2009 (RIFM, Woodcliff Lake, NJ, USA).

CAS N°:	2111-75-3	Empirical formula: Structure:	C ₁₀ H ₁₄ O
Synonyms:	1-Cyclohexene-1-carboxaldeh Dihydrocuminic aldehyde 4-Isopropenylcyclohex-1-ene-7 4-Isopropenyl-1-cyclohexene-7 p-Mentha-1,8-dien-7-al Perilla aldehyde Perillaldehyde	yde, 4-(1-methylethenyl)- 1-carbaldehyde 1-carboxaldehyde	

p-Mentha-1,8-dien-7-al (Perilla aldehyde)

History:	Initial reviews:	Oct. 1979, July 1994	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.02 %	Category 7	0.05 %			
Category 2	0.03 %	Category 8	0.1 %			
Category 3	0.1 %	Category 9	0.1 %			
Category 4	0.1 %	Category 10	0.1 %			
Category 5	0.1 %	Category 11	See Note Box (2)			
Category 6	0.5 %					
Note box [.]						

For this material, for pragmatic reasons, the restricted levels allowed by the QRA for certain categories that are actually higher than those that were in place before applying the QRA, will not be implemented for now. This position will be reevaluated as appropriate in the future.

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) (http://www.iofiorg.org/).

p-Mentha-1,8-dien-7-al (Perilla aldehyde)

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data				
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (μg/cm ²)	
2175 [2]	Moderate	7094	690⁴	2760	700	

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴ MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for p-Mentha-1,8-dien-7-al and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 700 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of p-Mentha-1,8-dien-7-al in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1978. Human Maximization Tests. RIFM report number 1698, August 25 and November 21a (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1979. Human Maximization Test. RIFM report number 1697, August 31 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007. Human Repeated Insult Patch Test. RIFM report number 53802, October 3 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2008. Local Lymph Node Assay. RIFM report 54428 (RIFM, Woodcliff Lake, NJ, USA).

Roberts, D.W., Patlewicz, G., Kern, P.S., Gerberick, F., Kimber, I., Dearman, R.J., Ryan, C.A., Basketter, D.A., Aptula, A.O., 2007. Mechanistic applicability domain classification of a local lymph node assay dataset for skin sensitization. *Chemical Research in Toxicology* 20, 1019-1030.

Menthadiene-7-methyl formate



History:	Initial reviews:	February 1986, July 1994			
	Current revision date:	2008			
	Implementation date:	For new submissions*:	August 16, 2008		
		For existing fragrance compounds*:	August 16, 2010		
	Next review date	2013			

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.03 %	Category 7	0.08 %		
Category 2	0.04 %	Category 8	0.1 %		
Category 3	0.1 %	Category 9	0.1 %		
Category 4	0.1 %	Category 10	0.1 %		
Category 5	0.1 %	Category 11	See Note Box (2)		
Category 6	0.8 %				
Note box:					
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again.					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/					
(2) Category 11 includes all non-skin contact or there is no justification for a restriction of the co	incidental skin contact p ncentration of this fragra	roducts. Due to the negligible skin contac nce ingredient in the finished product.	t from these types of products		

Fragrance material specifications:

N/A

Menthadiene-7-methyl formate

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

LINA weighted mean EC2 values		l	Human Data		
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
>2025	Weak	1063⁴	690⁴	6900	1060

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴ MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Menthadiene-7-methyl formate and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1060 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Menthadiene-7-methyl formate in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22,2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1977. Human Maximization Test. RIFM report number 1691, July 1b (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1978a. Human Maximization Test. RIFM report number 1698, April 28 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1978b. Human Maximization Test. RIFM report number 1787, October 26 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1979. Human Maximization Tests. RIFM report number 1775, September 11 and December 7 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007. Human Repeated Insult Patch Test. RIFM report number 53723, August 22a (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2008. Local Lymph Node Assay. DRAFT RIFM data, report 54429 (RIFM, Woodcliff Lake, NJ, USA).

p-Methoxybenzaldehyde

CAS N°:	123-11-5	Empirical formula: Structure:	C ₈ H ₈ O ₂
Synonyms:	Anisaldehyde <i>p</i> -Anisaldehyde Anisic aldehyde Benzaldehyde, 4-methoxy 4-Methoxybenzaldehyde p-Methoxybenzaldehyde Aubepine P Cresol (commer Aubepine liquid (commercial	cial name) name)	

History:	Initial reviews:	New Standard		
	Current revision date:	June 2013		
	Implementation date:	For new submissions*:	August 10, 2013	
		For existing fragrance compounds*:	August 10, 2014	
	Next review date	2018		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.10 %	Category 7	0.27 %		
Category 2	0.13 %	Category 8	2.00 %		
Category 3	0.54 %	Category 9	5.00 %		
Category 4	1.61 %	Category 10	2.50 %		
Category 5	0.84 %	Category 11	See Note box (2)		
Category 6	2.53 %				

Note box:

(1) See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products with the potential of ingestion.

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

p-Methoxybenzaldehyde

See Annex I

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

			Human Data		
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	NA weighted mean EC3 values (μg/cm ²) [no. studies] Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
>6250 [1]4	Weak	3543	6900	4724	3500

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴EC3 value from one LLNA, not the mean.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for p-Methoxybenzaldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as $3500 \ \mu g/cm^2$. They recommend the limits for the 11 different product categories, which are the acceptable use levels of p-Methoxybenzaldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick GF, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 2009a. Human repeated insult patch test. RIFM report number 58028, December 11. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2009b. Human repeated insult patch test. RIFM report number 58029, December 11. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. Human repeated insult patch test. Draft RIFM report number 63812. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007. Local Lymph Node Assay. RIFM report number 52910, May 21. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1975. Maximization study. RIFM report number 1799, March 27. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1973. Maximization study. RIFM report number 1802, October 31b. (RIFM, Woodcliff Lake, NJ, USA).

o-Methoxycinnamaldehyde

CAS N°:	1504-74-1	Empirical formula: Strucutre:	C ₁₀ H ₁₀ O ₂
Synonyms:	2'-Methoxycinnamaldehyde ortho-Methoxycinnamic aldeh β-(o-Methoxyphenyl)acrolein 3-(2-Methoxyphenyl)acrylalde 3-(o-Methoxyphenyl)-2-prope 2-Propenal, 3-(2-methoxyphe	yde ehyde nal nyl)-	

History:	Initial reviews:	New Standard	
	Current revision date:	June 20, 2011	
	Implementation date:	For new submissions*:	August 20, 2011
		For existing fragrance compounds*:	August 20, 2012
	Next review date	2016	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.03%	Category 7	0.08%		
Category 2	0.04%	Category 8	1.01%		
Category 3	0.15%	Category 9	5.00%		
Category 4	0.45%	Category 10	2.50%		
Category 5	0.24%	Category 11	See Note Box (2)		
Category 6	0.72%				
Note box:					

(1) See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products with the potential of ingestion.

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

o-Methoxycinnamaldehyde

RIFM SUMMARIES:

			Human Data		
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
NA	Weak	NA	2760	NA	1000

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003. ²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures. A default value based on the LLNA data for α -butylcinnamaldehyde (CAS 7492-44-6) was employed because the material is used a very low volume and there are no HRIPT data.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for *o*-methoxycinnamaldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1000 μ g/cm², which is a default value based on the LLNA data for a structurally related material, α -butylcinnamaldehyde (CAS 7492-44-6). They recommend the limits for the 11 different product categories, which are the acceptable use levels of *o*-methoxycinnamaldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick G, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.

7-Methoxycoumarin

CAS N°:	531-59-9	Empirical formula: Structure:	C10H8O3
Synonyms:	2H-1-Benzopyran-2-one, 7-methoxy- Herniarin		

History:	Initial reviews:	June 1979, April 1989	
	Current revision date:	2008	
	Implementation date:	For new submissions*:	August 16, 2008
		For existing fragrance compounds*:	August 16, 2009
	Next review date	2013	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products:	N/A				
Note box:					
The material as such should not be used as a f	agrance ingredient for a	ny application.			
On the basis of established maximum levels of exposure to this substance from the use of thes finished product does not exceed 100 ppm . Furthermore, these natural extracts should not	this substance in comme e oils and extracts is reg be used as substitutes fo	ercially available natural sources arded acceptable as long as the or this substance.	(like essential oils, extracts and absolutes), e level of 7-Methoxy-coumarin in the		
Examples for potential natural sources (with inc	licative maximum levels)	of 7-Methoxycoumarin are:			
 Camomilla matricaria EO 0,1 % Camomilla matricaria absolute (volatile part) : 5 % Lavandin absolute: 5 % (on the total absolute). Lavender and lavandin essential oils : <0,02 % Lavender absolute: 5 % (on the total absolute). Lime cold pressed oil: 0,1 % Tarragon absolute: volatile part: 5 % 					
Fragrance material specification	ns:	N/A			

7-Methoxycoumarin

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SENSITIZATION, PHOTOSENSITIZATION

REXPAN RATIONALE / CONCLUSION:

Based on the findings of RIFM on the potential for this material to induce allergic and photoallergic reactions (R.A. Ford et al. (1988), Fd. Chem. Toxic 26,375) the material as such is prohibited for use in fragrance compounds.

REFERENCES:

R.A. Ford et al. (1988), Fd. Chem. Toxic. 26, 375



Methoxy dicyclopentadiene carboxaldehyde (Scentenal)



History:	Initial reviews:	May 1998, May 2007 (42nd Amendment)		
	Current revision date:	2008		
	Implementation date:	For new submissions*: August 16, 2008		
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.1 %	Category 7	0.4 %		
Category 2	0.2 %	Category 8	0.5 %		
Category 3	0.5 %	Category 9	0.5 %		
Category 4	0.5 %	Category 10	0.5 %		
Category 5	0.5 %	Category 11	See Note Box (2)		
Category 6	3.6 %				
Note box:					
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again.					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/					
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.					

Fragrance material specifications:

Methoxy dicyclopentadiene carboxaldehyde (Scentenal)

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

Methoxy dicyclopentadiene carboxaldehyde - Sensitization Potency Estimation Based on Weight of Evidence

LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
>2500	Weak	5000 (DEP)	NA	NA	5000

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max Test

² Gerberick et al., 2001

³ WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for methoxy dicyclopentadiene carboxaldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 5000 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of methoxy dicyclopentadiene carboxaldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 1997. Repeated insult patch test of methoxy dicyclopentadiene carboxaldehyde in human subjects. Unpublished report from Firmenich, Inc., 23 January. RIFM report number 30026. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001. Murine local lymph node assay with methoxy dicyclopentadiene carboxaldehyde. Unpublished report from Firmenich, Inc., 8 October. RIFM report number 42120. (RIFM, Woodcliff Lake, NJ, USA).

V		V	
			7

CAS N°:	5462-06-6	Empirical formula: Structure:	C ₁₁ H ₁₄ O ₂
Synonyms:	2-Anisylpropional Benzenepropanal, 4-methy Hydrocinnamaldehyde, <i>p</i> -r <i>p</i> -Methoxyhydratropaldehy 4-Methoxy-α-methylbenze <i>p</i> -Methoxy-α-methylbydroo 3-(4-Methoxyphenyl)-2-methyl- 3-(<i>p</i> -Methoxyphenyl)-2-methyl- 2-Methyl-3-(<i>p</i> -methoxyphethyl- 2-Methyl-3-(4-methoxyphethyl- Canthoxal, Fennaldehyde,	oxy-α-methyl- methoxy-a-methyl /de nepropanal cinnamaldehyde ethylpropionaldehyde enyl)propanal enyl)propionaldehyde Foliaver (commercial nar	mes)

4-Methoxy-alpha-methylbenzenepropanal

History:	Initial reviews:	June 2009	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.17 %	Category 7	0.45 %			
Category 2	0.22 %	Category 8	2.00 %			
Category 3	0.89 %	Category 9	5.00 %			
Category 4	2.67 %	Category 10	2.50 %			
Category 5	1.40 %	Category 11	See Note box (2)			
Category 6	4.28 %					
Note box:						
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org) (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. 						
En		N1/A				

4-Methoxy-alpha-methylbenzenepropanal

CONTRIBUTION FROM OTHER SOURCES:

See Annex II

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
5900 [1] ⁴	Weak	5905	1380	NA	5900

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed

effect level; NA = Not Available.

¹ Data derived from HRIPT or HMT

² Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

³ WoE NESIL limited to two significant figures

⁴ EC3 value from one LLNA, not the mean

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 4-Methoxy- α -methylbenzenepropanal and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 5900 mg/cm2. They recommend the limits for the 11 different product categories, which are the acceptable use levels of 4-Methoxy- α -methylbenzenepropanal in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api et al., 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick GF, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 1980. Maximization study with 4-Methoxy-α-methylbenzenepropanal. RIFM report number 1790, August 26. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Local Lymph Node Assay. Unpublished study from IFF Inc., 22 November. Report number 47809. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2008. Human repeated insult patch test. RIFM report number 55562, July 30a. (RIFM, Woodcliff Lake, NJ, USA).

2-Methoxy-4-methylphenol				
CAS N°:	93-51-6	Empirical formula: Structure:		
Synonyms:	Creosol p-Creosol p-Cresol, 2-methoxy- Homoguaiacol 1-Hydroxy-2-methoxy-4-n 4-Hydroxy-3-methoxytolu 2-Methoxy-p-cresol 3-Methoxy-4-hydroxytolue 4-Methylguaiacol p-Methylguaiacol 4-Methyl-2-methoxyphene Phenol, 2-methoxy-4-met Valspice	nethylbenzene ene ene ol hyl-		

History:	Initial reviews:	January 1999, April 2005, May 2007	
	Current revision date:	2008	
	Implementation date:	For new submissions*: August 16, 2	
		For existing fragrance compounds*:	August 16, 2010
	Next review date	2013	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.003 %	Category 7	0.009 %			
Category 2	0.004 %	Category 8	0.01 %			
Category 3	0.01 %	Category 9	0.01 %			
Category 4	0.01 %	Category 10	0.01 %			
Category 5	0.01 %	Category 11	See Note Box (2)			
Category 6	0.09 %					
Note box:						
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again						



(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

2-Methoxy-4-methylphenol - Sensitization Potency Estimation Based on Weight of Evidence

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
1450	Weak	118	NA	NA	118

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max Test

² Gerberick et al., 2001

³ WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 2-methoxy-4-methylphenol and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 118 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of 2-methoxy-4-methylphenol in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Basketter, D. A., Gilmour, N., Dearman, R. J., Kimber, I., Ryan, C. A., Gerberick, E., 2003. Classification of skin sensitisation potency using the Local Lymph Node Assay. The Toxicologist, 72(S-1), 101.

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 1998. Repeated insult patch test of 2-methoxy-4-methylphenol in human subjects. RIFM report number 33100, June 15. (RIFM, Woodcliff Lake, NJ, USA).

alpha-Methyl anisylidene acetone

CAS N°:	104-27-8	Empirical formula: Structure:	C ₁₂ H ₁₄ O ₂
Synonyms:	1-(p-Methoxy p-Methoxyst alpha-Methy 1-Penten-3-0 Ethone	yphenyl)-1-penten-3-one yryl ethyl ketone lanisalacetone one, 1-(4-(methoxyphen	e yl)-

History:	Initial reviews:	November 1977, May 1980	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke, D.L.J. (1979), Food and Chemical Toxicology 17, 863.



alpha-Methyl-1,3-benzodioxole-5-propionaldehyde (MMDHA)

CAS N°:	1205-17-0	Empirical formula: Structure:	C ₁₁ H ₁₂ O ₃
Synonyms:	Heliofolal, Heliogan, Helional 1,3-Benzodioxole-5-propanal 3-(1,3-Benzodioxol-5-yl)-2-m 2-Methyl-3-(3,4-methylenedio α-Methyl-3,4-(methylenediox α-Methyl-1,3-benzodioxole-5 α-Methyl-1,3-benzodioxole-5 3-(3,4-Methylenedioxyphenyl α-Methyl-3,4-methylene-diox	, Tropional (commercial na , α-methyl- ethylpropanal oxyphenyl)- propionaldehyd oxyphenyl)propanal y)-hydrocinnamaldehyde -propanal -propionaldehyde)-2-methylpropanal yhydrocinnamic aldehyde	mes) le

History:	Initial reviews:	June 11, 2012		
	Current revision date:	June 2013		
	Implementation date:	For new submissions*: August 10, 2013		
		For existing fragrance compounds*:	August 10, 2014	
	Next review date	2018		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.34%	Category 7	0.89%		
Category 2	0.43%	Category 8	2.0%		
Category 3	1.78%	Category 9	5.0%		
Category 4	5.3%	Category 10	2.5%		
Category 5	2.8%	Category 11	See Note box (2)		
Category 6	8.6%				
Note box:					
 (1) See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products with the potential of ingestion. (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. 					

Fragrance material specifications:

N/A

alpha-Methyl-1,3-benzodioxole-5-propionaldehyde (MMDHA)

CONTRIBUTION FROM OTHER SOURCES:

See Annex II

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
4100 [1] ⁴	Weak	11,811	13,800	15,000	11,800

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴EC3 value from one LLNA, not the mean.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for α -methyl-1,3-benzodioxole-5-propionaldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 11800 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of α -methyl-1,3-benzodioxole-5-propionaldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick GF, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 2009. Human repeated insult patch test. RIFM report number 57514, July 16. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Local Lymph Node Assay. RIFM report number 50886, November 7. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2002. Human repeated insult patch test. Unpublished study from IFF, Inc., November 19. Report number 46969. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1985. Maximization study in humans. RIFM report number 1919, January 7c. (RIFM, Woodcliff Lake, NJ, USA).

alpha-Methyl cinnamic aldehyde

CAS N°:	101-39-3	Empirical formula: Structure:	C10H10O
Synonyms:	 α-Methylcinnamaldehyde α-Methylcinnamyl aldehyde α-Methylcinnamic aldehyde 2-Methyl-3-phenyl-2-propenal 3-Phenyl-2-methylacrolein 2-Propenyl, 2-methyl-3-phenyl- 		

History:	Initial reviews:	New Standard		
	Current revision date:	2007		
	Implementation date:	For new submissions*:	June 16, 2007	
		For existing fragrance compounds*:	June 16, 2009	
	Next review date	2012		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.1 %	Category 7	0.3 %		
Category 2	0.1 %	Category 8	2.0 %		
Category 3	0.5 %	Category 9	5.0 %		
Category 4	1.6 %	Category 10	2.5 %		
Category 5	0.8 %	Category 11	See Note Box (2)		
Category 6	2.5 %				
Note box:					
 (1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/ (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products 					
there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

alpha-Methyl cinnamic aldehyde

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

alpha-Methyl cinnamic aldehyde - Sensitization Potency Estimation Based on Weight of Evidence

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
1125 [1]	Extremely weak	3543	5517	NA	3500

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max Test

² Gerberick et al., 2001

³ WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for alpha-methyl cinnamic aldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 3500 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of alpha-methyl cinnamic aldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Elahi, E. N., Wright, Z., Hinselwood, D., Hotchkiss, S. A. M., Basketter, D. A., Pease, C. K. S., 2004. Protein binding and metabolism influence the relative skin sensitization potential of cinnamic compounds. Chemical Research in Toxicology, 17(3), 301-310.

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 1973. Report on human maximization studies. RIFM report number 1802, July 10 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003. Repeated insult patch test with alpha-methylcinnamaldehyde. RIFM report number 47264, February 28 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2003a. Repeated insult patch test with alpha-methylcinnamaldehyde. RIFM report number 47265, February 28 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Repeated insult patch test with alpha-methylcinnamaldehyde. RIFM report number 45133, March 11 (RIFM, Woodcliff Lake, NJ, USA).

6-Methylcoumarin

CAS N°:	92-48-8	Empirical formula: Structure:	C ₁₀ H ₈ O ₂
Synonyms:	2H-1-Benzopyran-2-one, 6-methyl 6-Methyl-2h-1-benzopyran-2-one 6-Methylbenzopyrone 6-Methyl coumarin 6-Methyl-cis-o-coumarinic lactone 5-Methyl-2-hydroxyphenylpropenoic acid Toncarine	lactone	

History:	Initial reviews:	March 1978, October 1978, February 1980	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

PHOTOALLERGY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Kaidbay, K.H. &. Kligman, A.M. (1978), Contact Dermatitis 4, No 5, 277. Opdyke, D.L.J. (1979), Food and Cosmetics Toxicology 17, 275.



11	1 10 1 1		
History:	Initial reviews:	February 1979, May 1983	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION, PHOTOALLERGY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted - REXPAN, September 2002.

REFERENCES:

Opdyke, D.L.J., Letizia, C.S. (1982), Food and Chemical Toxicology 20, 747.

Synonyms:



2-Butenoic acid, methyl ester, (E)-

Methyl trans-2-butenoate



History:	Initial reviews:	March 1978, May 1980	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke, D.L.J. (1979), Food and Cosmetics Toxicology 17, 865.

CAS N°:87-05-8Empirical formula:
Structure: $C_{12}H_{12}O_3$ Synonyms:2H-1-Benzopyran-2-one, 7-ethoxy-4-methyl-
Coumarin, 7-ethoxy-4-methyl-
7-Ethoxy-4-methyl-
7-ethoxybenzopyrone
MaraniolC12H12O3

4-Methyl-7-ethoxycoumarin

History:	Initial reviews:	June 1979	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

PHOTOALLERGY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Private communication to IFRA.
Methyl eugenol

CAS N°:	93-15-2	Empirical formula: Structure:	C11H14O2
Synonyms:	Eugenyl methyl ether Methyl eugenol ether Allylveratrole Veratrole methyl ether 4-Allyl-1,2-dimethoxybenzene Benzene, 1,2-dimethoxy-4-(2-propenyl)- 1,2-Dimethoxy-4-allylbenzene 1,2-dimethoxy-4-(2-propenyl)- benzene		

History:	Initial reviews:	December 2002		
	Current revision date:	2015		
	Implementation date:	For new submissions*: August 10, 2015		
		For existing fragrance compounds*:	August 10, 2016	
	Next review date	2020		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave-on products:	See note box	Rinse-off products:	See note box		
Non-skin contact products:	See note box Including household cleaning products				
Note box:					
The Standard is based on long-term systemic effects and will therefore have a wider range of product type limitations as follows: Fine fragrance: 0.02% Eau de toilette: 0.008% Fragrancing cream: 0.004% Other leave-on: 0.0004% Rinse-off: 0.001% Non-skin, incidental skin contact: 0.01%					
The limitations apply to Methyl eugenol (ME) originating from all sources. Contributions from essential oils can be significant (see Contribution from other sources).					
Fragrance material specifications: N/A					

Methyl eugenol

CONTRIBUTION FROM OTHER SOURCES:

See Annex I.

CRITICAL EFFECT:

CARCINOGENICITY

RFIM SUMMARIES:

The currently available metabolic, biochemical and toxicological data found for Methyl eugenol in laboratory species provide clear evidence of non-linearity in the dose-response relationship for Methyl eugenol with respect to metabolic activation and mechanisms associated with carcinogenic effects. Consideration of these data indicates a No Observed Effect Level for Methyl eugenol in the rat in the dose-range of 1-10 mg/kg body weight/day.

REXPAN RATIONALE / CONCLUSION:

Based on the lower end of the NOEL and applying a 1000 times safety factor for systemic effects a daily dose for Methyl eugenol of 60 μ g/day is supported. Taking into account a dermal penetration factor of 40% leads to an acceptable dose of 150 μ g/day.

This daily dose, applying the calculation table for dermal exposure as attached (contained in Api *et al.* (2014)), results in the maximum concentrations for Methyl eugenol in certain product categories as outlined in the section 'limits' of this Standard.

REFERENCES:

Api A.M., Belsito D., Bruze M., Cadby P., Calow P., Dagli M. L., Dekant W., Dent M., Ellis G., Fryer A. D., Fukayama M., Griem P., Hickey C., Kromidas L., Lalko J., Liebler D.C., Miyachi Y., Politano V.T., Renskers K., Ritacco G., Salvito D., Schultz T.W., Sipes I. G., Smith B., Vitale D., Wilcox D.K. (2014). Food and Chemical Toxicology, <u>http://dx.doi.org/10.1016/j.fct.2014.11.014</u>.

6-Methyl-3,5-heptadien-2-one

CAS N°:	1604-28-0	Empirical formula: Structure:	C ₈ H ₁₂ O
Synonyms:	3,5-Heptadien-2-one, 6-methyl- Methylheptadienone 2-Methylhepta-2,4-dien-6-one 6-Methylhepta-3,5-dien-2-one		

History:	Initial reviews:	April 1989, April 1999		
	Current revision date:	2009		
	Implementation date:	For new submissions*: August 7, 2009		
		For existing fragrance compounds*:	August 7, 2011	
	Next review date	2014		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.002 %	Category 7	0.002 %		
Category 2	0.002 %	Category 8	0.002 %		
Category 3	0.002 %	Category 9	0.002 %		
Category 4	0.002 %	Category 10	0.002 %		
Category 5	0.002 %	Category 11	See Note Box (2)		
Category 6	0.100 %				
Note box:					
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be re-evaluated again.					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) (http://www.iofiorg.org/).					
(2) Category 11 includes all non-skin contact or there is no justification for a restriction of the co	incidental skin contact p ncentration of this fragra	roducts. Due to the negligible skin contac nce ingredient in the finished product.	t from these types of products		

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

6-Methyl-3,5-heptadien-2-one

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
> 1250 [1] ⁴	Weak	118	NA	1299	110

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed

effect level; NA = Not Available.

¹ Data derived from HRIPT or HMT

- ² Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003
- ³ WoE NESIL limited to two significant figures
- ⁴ EC3 value from one LLNA, not the mean

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 6-Methyl-3,5-heptadiene-2-one and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 110 mg/cm2. They recommend the limits for the 11 different product categories, which are the acceptable use levels of 6-Methyl-3,5-heptadiene-2-one in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 2008a. Local Lymph Node Assay. RIFM report number 55564, July 30. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2008b. Human repeated insult patch test. RIFM report number 55345, August 5. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2008c. Human repeated insult patch test. RIFM report number 55661, November 24a. (RIFM, Woodcliff Lake, NJ, USA).

Methyl heptine carbonate



History:	Initial reviews:	October 1976, April 2000, April 2005		
	Current revision date:	2008		
	Implementation date:	For new submissions*:	August 16, 2008	
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.003 %	Category 7	0.008 %		
Category 2	0.004	Category 8	0.01 %		
Category 3	0.01 %	Category 9	0.01 %		
Category 4	0.01 %	Category 10	0.01 %		
Category 5	0.01 %	Category 11	See Note Box (2)		
Category 6	0.08 %				
Note boy					

For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be re-evaluated again.

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) (http://www.iofiorg.org/).

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

When used in the same fragrance compound within a specific QRA category, the sum total of methyl heptine carbonate (MHC) and methyl octine carbonate (MOC) contributions must not exceed the maximum permitted level for MHC. At the same time, the contribution from methyl octine carbonate should always respect the maximum levels permitted in the respective categories as listed in the Standard for MOC. If the same compound is intended for more than one IFRA QRA category, then the most restrictive limitations (based on foreseen use concentrations and maximum permitted level) will apply.

Methyl heptine carbonate

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
< 125	Strong	118 ⁴	NA	194	110

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for methyl heptine carbonate and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 110 mg/cm2. They recommend the limits for the 11 different product categories, which are the acceptable use levels of methyl heptine carbonate in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1964. Human repeated patch test on methyl-2-nonynoate and isoeugenol. Unpublished report from IFF, Inc., 30 April. Report number 1808. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1989. Human repeated insult patch test of methyl 2-octynoate. RIFM report number 12368, November 16. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1990. Repeated insult patch test of methyl 2-octynoate in human subjects. RIFM report number 12452, April 27. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Methyl 2-octynoate diluted with vehicle 1:3 EtOH:DEP: Local Lymph Node Assay. RIFM report number 48753, January 28. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2006. Methyl 2-octynoate: Local Lymph Node Assay. RIFM report number 51627, April 6. (RIFM, Woodcliff Lake, NJ, USA).

p-Methylhydrocinnamic aldehyde



History:	Initial reviews:	Nov. 1987, July 1994, January 2002, May 2007	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



p-Methylhydrocinnamic aldehyde

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or

2) adverse data on the material itself and/or

3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000



Methyl ionone, mixed isomers

CAS N°:	1335-46-2 127-42-4 127-43-5 127-51-5 7779-30-8 79-89-0	Empirical formula: Structure:	C14H22O N/A		
Synonyms:	CAS 1335-46-2: Methy	ionone, mixture of isomers			
	CAS 127-42-4: Methyl-a alpha-Cetone alpha-Cyclocitrylideneb alpha-Cyclocitrylidenen Methyl-a-ionone alpha-Methylionone 1-Penten-3-one, 1-(2,6, (R-(E))-1-(2,6,6-Trimeth	alpha-ionone utanone nethyl ethyl ketone 6-trimethyl-2-cyclohexen-1-yl)-, [R-(nyl-2-cyclohexen-1-yl)pent-1-en-3-or	(E)]- ne		
	CAS 127-43-5: Methyl-beta-ionone Methyl-ß-ionone beta-Methylionone beta-Cetone beta-Cyclocitrylidenebutanone beta-Iraldeine 1-Penten-3-one, 1-(2,6,6-trimethyl-1-cyclohexen-1-yl)- 5-(2,6,6-Trimethyl-1-cyclohexen-1-yl)-4-penten-3-one 1-(2,6,6-Trimethyl-1-cyclohexen-1-yl)pent -1-en-3-one				
	CAS 127-51-5: alpha-is 3-Buten-2-one, 3-methy 3-Methyl-4-(2,6,6-trimer alpha-Isomethyl ionone gamma-Methylionone Iraldeine gamma Isoraldeine 95	o methylionone /I-4-(2,6,6-trimethyl-2-cyclohexen-1- thyl-2-cyclohexen-1-yl)-3-buten-2-or	yl)- ie		
	CAS 7779-30-8: 1-(2,6,6-Trimethyl-2-cyclohexen-1-yl)pent-1-en-3-one 1-Penten-3-one, 1-(2,6,6-trimethyl-2-cyclohexen-1-yl)-				
	CAS 79-89-0: iso-Meth 3-Buten-2-one, 3-meth 3-Methyl-4-(2,6,6-trime δ-Iraldeine	yl-beta-ionone /l-4-(2,6,6-trimethyl-1-cyclohexen-1- thylcyclohex-1-en-1-yl)but-3-en-2-or	yl)- ie		

History:	Initial reviews:	2007	
	Current revision date:	2015	
	Implementation date:	For new submissions*:	August 10, 2015
		For existing fragrance compounds*:	August 10, 2016
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

Methyl ionone, mixed isomers

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:							
Category 1 See Note box (1)	2.00 %	Category 7	5.30 %				
Category 2	2.59 %	Category 8	2.00 %				
Category 3	10.56 %	Category 9	5.00 %				
Category 4	31.67 %	Category 10	2.50 %				
Category 5	16.67 %	Category 11	See Note box (2)				
Category 6	50.72 %						
Also La							

Note box:

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofi.org)

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

The above limits apply to Methyl ionone isomers used individually or in combination.

Fragrance material specifications:

Pseudo methyl ionones (CAS numbers 26651-96-7, 72968-25-3, 1117-41-5) should not be used as fragrance ingredient as such, but a level of up to 2% as an impurity in methyl ionones is accepted.

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

DERMAL SENSITIZATION

RIFM SUMMARIES:

LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
5450	Weak	708864	NA	NA	70000

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No Observed Effect Level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test;

LOEL = Lowest Observed Effect Level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL.

Methyl ionone, mixed isomers

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Methyl ionone, mixed isomers and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 70000 μ g/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Methyl ionone, mixed isomers in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api, A. M., Basketter, D. A., Cadby, P. A., Cano, M-F., Ellis, G., Gerberick, G. F. *et al.*, 2008. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. alpha-iso-Methylionone diluted with vehicle 1:3 EtOH:DEP: Local Lymph Node Assay. RIFM report number 48749, January 26 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004a. Repeated insult patch test with alpha-iso-methylionone. RIFM report number 47278, March 10 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004b. Repeated insult patch test with alpha-iso-methylionone. RIFM report number 47279, March 10 (RIFM, Woodcliff Lake, NJ, USA).



Methyl methacrylate

CAS N°:	80-62-6	Empirical formula: Structure:	C5H8O2
Synonyms:	Methyl 2-methacrylate, 2-(metho Methyl 2-methyl-2-propenoate 2-Propenoic acid, 2-methyl-, met MMA	xycarbonyl)-1-propene hyl ester	

History:	Initial reviews:	Not applicable		
	Current revision date:	Not applicable		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:							
Skin contact products:							
Leave on products:	N/A	Rinse-off products:	N/A				
		Including household	cleaning products				
Non skin contact products:	N/A						
Note box:							
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.							
Fragrance material specifications: N/A							

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX

Methyl methacrylate

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000

Methyl N-methylanthranilate

CAS N°:	85-91-6	Empirical formula: Structure:	C9H11NO2
Synonyms:	Benzoic acid, 2-(methylamino)-, m Dimethyl anthranilate 2-Methylamino methyl benzoate N-Methylanthranilic acid, methyl e Methyl 2-(methylamino)benzoate Methyl 2-methylaminobenzoate Methyl o-methylaminobenzoate	ethyl ester ster	

History:	Initial reviews:	1978, 2001, 2002, 2006, 2009	
	Current revision date:	2015	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:							
Skin contact products:							
Leave-on products :	0.1 %	Rinse-off products: Including household cleaning products	No restriction				
Non-skin contact products:	No rest	riction					
Note box:							
The Standard is set due to the phototoxic effects of the material. The limit only applies to applications on skin exposed to sunshine, excluding rinse- off products (please refer to Table 4 of the QRA booklet for more detailed information).							
The material has been identified for having the potential of forming nitrosamines in nitrosating systems. Downstream users therefore have to be notified of the presence of the material and its potential to be able to consider adequate protective measures.							
Fragrance material specifications: N/A							

CONTRIBUTION FROM OTHER SOURCES:

See Annex I.

Methyl N-methylanthranilate

CRITICAL EFFECT:

PHOTOTOXICITY, POTENTIAL FOR NITROSAMINE FORMATION

RIFM SUMMARIES:

A human phototoxicity study at 0.5% in 75% ethanol/25% diethyl phthalate (DEP) resulted in 0/26 reactions (RIFM, 2001). Another human phototoxicity study with concentrations of 0.1, 0.3, and 0.5% resulted in 0/29 reactions (RIFM, 1998). Several other phototoxicity studies showed phototoxic reactions at 1% and 5% (Kaidbey and Kligman, 1980; Letizia and Api, 2003; RIFM, 1999).

A human photosensitization study at 0.5% in 75% ethanol/25% DEP resulted in 0/26 reactions (RIFM, 2001). Another human photosensitization study at 5.0% resulted in no photoallergic reactions. However, 14/18 phototoxic reactions were observed (RIFM, 1978a).

A phototoxicity study at 50% in methanol and 100% on hairless mice produced reactions at both dose levels (RIFM, 1978b).

An in vitro phototoxicity assay using a human skin model (Skin2[®]) with concentrations of methyl N-methylanthranilate ranging from 0.05% to 25% in corn oil showed that the material was phototoxic at dose levels above 5% (Api, 1997).

REXPAN RATIONALE / CONCLUSION:

IFRA measures regarding potential nitrosamine formation noted - REXPAN April 2009.

REFERENCES:

Api A.M. (1997). In vitro assessment of phototoxicity. In Vitro Toxicology: Journal of Molec. Cell. Toxicol., 10(3), 339-350.

Kaidbey K.H. and Kligman A.M. (1980). Identification of contact photosensitizers by human assay. In Current Concepts In Cutaneous Toxicity, Academic Press, New York, pages 55-68.

Letizia C.S. and Api A.M. (2003). Evaluation of the phototoxic and photoallergenic potential of methyl N-methyl anthranilate. The Toxicologist, 72(S1), 378-379.

Research Institute for Fragrance Materials, Inc. (1978a). Phototoxicity and contact photoallergy testing in human subjects. RIFM report number 1788, 18 January.

Research Institute for Fragrance Materials, Inc. (1978b). Phototoxicity and irritation studies of mice and pigs with fragrance materials. RIFM report number 2042, 13 April.

Research Institute for Fragrance Materials, Inc. (1998). Evaluation of phototoxicity of dimethyl anthranilate in humans. RIFM report number 34768, 8 December.

Research Institute for Fragrance Materials, Inc. (1999). Evaluation of phototoxicity of dimethyl anthranilate in humans. RIFM report number 34769, 20 July.

Research Institute for Fragrance Materials, Inc. (2001) Evaluation of human photoallergy by repeated insult patch test. RIFM report number 36789, 1 March.

Nitrosamine policy as contained in the EU Cosmetics Directive 76/768/EEC and its Amendments.

Methyl β -naphthyl ketone CAS N°: 93-08-3 Empirical formula: Structure: C12H10O Synonyms: 2-Acetonaphthone β -Acetylnaphthalene Cetone d Ethanone, 1-(2-naphthalenyl) (CAS) β -Methyl naphthyl ketone Oranger crystals C12H10O

History:	Initial reviews:	October 2004		
	Current revision date:	2015		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	2020		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:							
Skin contact products:							
Leave-on products :	0.2%	Rinse- Includi	off products: ng household cleaning products	No restriction			
Non-skin contact products:	No restriction						
Note box:							
The Standard is set due to the phototoxic effects of the material. The limit only applies to applications on skin exposed to sunshine, excluding rinse- off products (please refer to Table 4 of the QRA booklet for more detailed information).							
Fragrance material specifications: N/A							

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

PHOTOTOXICITY

Methyl β-naphthyl ketone

RIFM SUMMARIES:

Human Studies

A human phototoxicity study with Methyl β -naphthyl ketone (concentrations of 0.1, 1 and 10% in 3:1 DEP:EOH) was conducted. No reactions indicative of primary irritation were observed in this study. However, under irradiated conditions, Methyl β -naphthyl ketone at 10% in 3:1 DEP:EtOH produced moderate erythema in 5 subjects. These responses were stronger than those seen for the irradiated blank patch, which only produced slight to mild erythema. Under the conditions of the study, Methyl β -naphthyl ketone at 10% in 3:1 DEP:EtOH showed evidence of phototoxicity. Erythema scores for Methyl β -naphthyl ketone at 0.1% and 1.0% in 3:1 DEP:EtOH were similar to those seen for the blank patch under irradiated conditions. These reactions were not indicative of phototoxic responses (RIFM, 2004).

Other Studies

Methyl β -naphthyl ketone has been observed to absorb in the UV range of 290-400 nm and is positive in the Neutral Red Uptake Phototoxicity Assay (RIFM, 2002). However, it has been shown to be non-phototoxic in guinea pigs at concentrations up to 60% in 3:1 EtOH:DEP (RIFM, 2003).

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Methyl β -naphthyl ketone and recommended a limit of 0.2%, based on a no-effect level for phototoxicity in humans of 1% (May 18, 2004).

REFERENCES:

Research Institute for Fragrance Materials, Inc. (2002). Methyl β-naphthyl ketone: Neutral red uptake phototoxicity assay in BALB/C 3T3 mouse fibroblasts. RIFM report number 40279, May 30 (RIFM, Woodcliff Lake, NJ, USA).

Research Institute for Fragrance Materials, Inc. (2003). Topical photoallergy screening test of â-methyl naphthyl ketone in male albino hairless guinea pigs including primary irritation, phototoxicity and contact hypersensitivity evaluations. RIFM report number 44882, June 9 (RIFM, Woodcliff Lake, NJ, USA).

Research Institute for Fragrance Materials, Inc. (2004). Evaluation of phototoxicity of methyl â-naphthyl ketone in humans. RIFM report number 45136, March 16 (RIFM, Woodcliff Lake, NJ, USA).



3-Methyl-2(3)-nonenenitrile

CAS N°:	53153-66-5	Empirical formula: Structure:	C10H17N
Synonyms:	2-Nonenenit Citgrenile	rile, 3-methyl-	

History:	Initial reviews:	February 1980, May 1983, May 2007		
	Current revision date:	Not applicable		
	Implementation date:	For new submissions*:	Not applicable	
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:							
Skin contact products:							
Leave on products:	N/A	Rinse-off products:	N/A				
	Including household cleaning products						
Non skin contact products:	N/A						
Note box:							
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.							
Fragrance material specification	ons:	N/A					

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



3-Methyl-2(3)-nonenenitrile

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000

Methyl octine carbonate



History:	Initial reviews:	March 1988, April 2000		
	Current revision date:	2008		
	Implementation date:	For new submissions*: August 16, 2008		
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.001 %	Category 7	0.002 %		
Category 2	0.001 %	Category 8	0.002 %		
Category 3	0.002 %	Category 9	0.002 %		
Category 4	0.002 %	Category 10	0.002 %		
Category 5	0.002 %	Category 11	See Note Box (2)		
Category 6	0.02 %				
Note box:					
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be re-evaluated again.					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) (http://www.iofiorg.org/).					
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.					
When used in the same fragrance compound within a specific ORA category, the sum total of methyl octine carbonate (MOC) and methyl boating					

When used in the same fragrance compound within a specific QRA category, the sum total of methyl octine carbonate (MOC) and methyl heptine carbonate (MHC) contributions must not exceed the maximum permitted level for MHC. At the same time, the contribution from methyl octine carbonate should always respect the maximum levels permitted as listed in the table above. If the same compound is intended for more than one IFRA QRA category, then the most restrictive limitations (based on foreseen use concentrations and maximum permitted level) will apply.

Fragrance material specifications:

N/A

Methyl octine carbonate

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
< 1250 Estimated 625	Strong	244	NA	118	24

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³ WoE NESIL limited to three significant figures

⁴ MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Methyl octine carbonate and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as $24 \ \mu g/cm^2$. They recommend the limits for the 11 different product categories, which are the acceptable use levels of Methyl octine carbonate in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1989a. Repeated insult patch test of methyl octine carbonate in human subjects. RIFM report number 27280, May 22. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1989b. Human repeated insult patch test of methyl 2-octynoate and methyl 2-nonynoate. RIFM report number 12367, November 16. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1990a. Repeated insult patch test of methyl octine carbonate and t-2-hexenal in human subjects. RIFM report number 27822, January 9. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1990b. Repeat insult patch test of methyl 2-nonynoate in human subjects. RIFM report number 12454, April 27. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1990c. Repeat insult patch test of methyl 2-octynoate and methyl 2-nonynoate in human subjects. RIFM report number 12456, April 27. (RIFM, Woodcliff Lake, NJ, USA).

Ryan, C. A., Gerberick, G. F., Cruse, L. W., Basketter, D. A., Lea, L., Blaikie, L., Dearman, R. J., Warbrick, E. V., Kimber, I., 2000. Activity of human contact allergens in the murine local lymph node assay. *Contact Dermatitis*, 43(2), 95-102.



3-Methyl-2-(pentyloxy)cyclopent-2-en-1-one

CAS N°:	68922-13-4	Empirical formula: Strucutre:	
Synonyms:	2-Cyclopenten-1-one, 2-(pentyl	oxy)-3-methyl-	
	Pentyloxy Cyclopentenone (trac	de name)	

History:	Initial reviews:	New Standard		
	Current revision date:	June 20, 2011		
	Implementation date:	For new submissions*: August 20, 201		
		For existing fragrance compounds*:	August 20, 2012	
	Next review date	2016		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.03 %	Category 7	0.08 %			
Category 2	0.04 %	Category 8	1.11 %			
Category 3	0.17 %	Category 9	5.00 %			
Category 4	0.50 %	Category 10	2.50 %			
Category 5	0.26 %	Category 11	See Note Box (2)			
Category 6 0.80 %						
Note box:						
(1) See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products						

with the potential of ingestion. (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None known at the time of the publication of the Standard

CRITICAL EFFECT:

SENSITIZATION

3-Methyl-2-(pentyloxy)cyclopent-2-en-1-one

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
NA	NA	1181	NA	10% ³	1100

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003. ²Data derived from HRIPT or HMT. ³Conversion to μg/cm² was not possible due to absence of details

⁴WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 3-Methyl-2-(pentyloxy)cyclopent-2-en-1-one and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1100 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of 3-Methyl-2-(pentyloxy)cyclopent-2-en-1-one in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick G, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.

p-Methyltetrahydroquinoline



History:	Initial reviews:	New Standard			
	Current revision date:	2009			
	Implementation date:	For new submissions*:	August 7, 2009		
		For existing fragrance compounds*:	August 7, 2010		
	Next review date	2014			
* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.					

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:							
Skin contact products:							
Leave on products:	N/A	Rinse-off products:	N/A				
		Including household	cleaning products				
Non skin contact products:	N/A						
Note box:							
The material has been identified for having the potential of forming nitrosamines in nitrosating systems. Downstream users therefore have to be notified of the presence of the material and its potential to be able to consider adequate protective measures.							
Fragrance material specifications: N/A							

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)

CRITICAL EFFECT:

POTENTIAL FOR NITROSAMINE FORMATION

REXPAN RATIONALE / CONCLUSION:

IFRA measures regarding potential nitrosamine formation noted - REXPAN April 2009.

REFERENCES:

Nitrosamine policy as contained in the EU Cosmetics Directive 76/768/EEC and its Amendments.

CAS N°:	116-66-5	Empirical formula: Structure:	C14H18N2O4
			0 ^{N+} 0
Synonyms:	1H-Indene, 2,3-dihyd 1,1,3,3,5-Pentameth	dro-1,1,3,3,5-pentamet yl-4,6-dinitroindane	hyl-4,6,-dinitro-

Moskene (1,1,3,3,5-Pentamethyl-4,6-dinitroindane)

History:	Initial reviews:	39 th Amendment		
	Current revision date:	Not applicable		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	N/A	Rinse-off products:	N/A	
	Including household cleaning products			
Non skin contact products:	N/A			
Note box:				
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.				
Fragrance material specifications: N/A				

CONTRIBUTION FROM OTHER SOURCES:

N/A



Moskene (1,1,3,3,5-Pentamethyl-4,6-dinitroindane)

CRITICAL EFFECT:

SEE NOTE BOX

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or

2) adverse data on the material itself and/or

3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000

Musk ambrette



History:	Initial reviews:	June 1981, July 1994, December 1995	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products:	N/A	N/A			
Note box:					
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

N/A



CRITICAL EFFECT:

PHOTOSENSITIZATION, NEUROTOXICITY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Spencer, P.S., Bischoff-Fenton, M.C., Moreno, O.M., Opdyke D.L. and Ford, R.A. (1984), Toxicology and Applied Pharmacology 75, 571.

Musk tibetene (1-tert-Butyl-2,6-dinitro-3,4,5-trimethylbenzene)



History:	Initial reviews:	39 th Amendement	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	N/A	Rinse-off products:	N/A	
	Including household cleaning products			
Non skin contact products:	N/A			
Note box:				
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.				
Fragrance material specifications: N/A				

CONTRIBUTION FROM OTHER SOURCES:

N/A



CRITICAL EFFECT:

SEE NOTE BOX

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or

2) adverse data on the material itself and/or

3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000

Musk xylene



History:	Initial reviews:	New Standard	
	Current revision date:	2009	
	Implementation date:	For new submissions*:	August 7, 2009
		For existing fragrance compounds*:	August 7, 2010
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

vPvB

REXPAN RATIONALE / CONCLUSION:

Ongoing research on the fragrance ingredient musk xylene has provided evidence over time that it fulfills the criteria for being classified vPvB (Environmental half-life >180 days; BCF>5000).

Musk Xylene, as of October 8, 2008, has been identified by the European Chemicals Bureau as a material requiring authorization under REACH due to its properties as a vPvB

Based on its potential detrimental environmental impact the REXPAN has concluded that the material should not be used as a fragrance ingredient.



Musk xylene

REFERENCES:

PBT draft Addendum to the final report (2005) of the Risk Assessment (PBT assessment), January 2008 (the Netherlands National Institute for Public health and Environment, RIVM).

ECHA (European Chemicals Agency, , Member State Committee, Substances of Very High Concern support document for identification of 5-tert-butyl-2,4,6-trinitro-m-xylene, Adopted on October 8, 2008 (http://echa.europa.eu/doc/candidate_list/svhc_supdoc_muskxylene_publication.pdf.)



History:	Initial reviews:	June 1974		
	Current revision date:	September 2002		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	N/A	Rinse-off products:	N/A	
	Including household cleaning products			
Non skin contact products:	N/A			
Note box:				
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.				
Fragrance material specifications:		N/A		

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

ACUTE TOXICITY, SKIN TOXICITY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Christensen, H.E., Toxic Substances List, National Institute for Occupational Safety and Health (1972), p. 369.

2-Nonyn-1-al dimethyl acetal

CAS N°:	13257-44-8	Empirical formula: Strucutre:	C ₁₁ H ₂₀ O ₂
Synonyms:	1,1-Dimethoxynon-2-yne 2-Nonyn-1-al-Dimeth-Acetyl 2-Nonyne, 1,1-dimethoxy- Parmavert		

History:	Initial reviews:	New Standard	
	Current revision date:	June 20, 2011	
	Implementation date:	For new submissions*: August 20, 2011	
		For existing fragrance compounds*:	August 20, 2012
	Next review date	2016	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Category 1 See Note box (1)	0.66 %	Category 7	1.74 %	
Category 2	0.84 %	Category 8	2.00 %	
Category 3	3.47 %	Category 9	5.00 %	
Category 4	10.41 %	Category 10	2.50 %	
Category 5	5.48 %	Category 11	See Note Box (2)	
Category 6	16.67 %			
Note box:				
 See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products with the potential of ingestion. Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product. 				

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None known at the time of the publication of the Standard

CRITICAL EFFECT:

SENSITIZATION

2-Nonyn-1-al dimethyl acetal

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
>5,000 [1]	Weak	23,622	NA	NA	23,000

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003. ²Data derived from HRIPT or HMT. ³WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 2-Nonyn-1-al dimethyl acetal and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 23,000 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of 2-Nonyn-1-al dimethyl acetal in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick G, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.

Nootkatone

CAS N°:	4674-50-4	Empirical formula: Structure:	C ₁₅ H ₂₂ O O
Synonyms:	5,6-Dimethyl- 4a,5-Dimethyl 4betaH,5alph 4betaH,5alph (4R-(4alpha,4 methylvinyl)na 4,4a,5,6,7,8-H 2(3H)-Naphth (4R,4aS,6R)-	8-isopropenylbicyclo(I-1,2,3,4,4a,5,6,7-octa a-Eremophila-1(10),1 a-Eremorphila-1(10), a alpha,6beta))-4,4a, aphthalen-2(3H)-one lexahydro-6-isoprope alenone, 4,4a,5,6,7,8	4.4.0)dec-1-en-3-one ahydro-7-keto-3-isopropenylnaphthalene 1-dien-2-one 5,6,7,8-Hexahydro-4,4a-dimethyl-6-(1- enyl-4,4a-dimethyl-2(3H)-naphthalenone i-hexahydro-4,4a-dimethyl-6-(1-methylethenyl)-,

History:	Initial reviews:	October 1980	
	Current revision date:	August 2005	
	Implementation date:	For new submissions*:	December 11, 2006
		For existing fragrance compounds*:	December 11, 2007
	Next review date	2009	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

SPECIFICATION

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	N/A	Rinse-off products:	N/A	
		Including household cleaning products		
Non skin contact products:	N/A			
Note box:				
Nootkatone used as a fragrance ingredient should be at least 98% pure, with a melting point of at least 32°C. Lower purity grades may not be used as a fragrance ingredient.				
Fragrance material specifications:		N/A		

CONTRIBUTION FROM OTHER SOURCES:

None to consider.
Nootkatone

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the Standard on Nootkatone. After reviewing the critical effect data, the Panel concluded that only samples of Nootkatone that are at least 98% pure should be used. Lower purity grades may not be used as a fragrance ingredient. Sensitization was observed with the lower purity grades, but not with high purity material (>98% pure) samples of Nootkatone.

REFERENCES:

Research Institute for Fragrance Materials, Inc., 1971. Sensitization and irritation study of nootkatone. Unpublished report from Givaudan, May 24, Report number 41820.

Research Institute for Fragrance Materials, Inc., 1977. Report on human maximization studies. RIFM report number 1702, June 6c.

Research Institute for Fragrance Materials, Inc., 1978. Report on human maximization studies. RIFM report number 1698, January 13a.

Research Institute for Fragrance Materials, Inc., 1979. Report on human maximization studies. RIFM report number 1775, September 11.

Research Institute for Fragrance Materials, Inc., 2005. Repeated insult patch test with nootkatone. Unpublished report from Bedoukian Research, Inc., May 11. Report number 46155.

Oakmoss extracts

CAS N°:	90028-68-5 = Evernia prunastri extract 9000-50-4 = oils, Oakmoss resinoid 68917-10-2 =oils, Oakmoss	Empirical formula:	N/A
Synonyms:	Oakmoss absolute Evernia absolute <i>Evernia prunastri</i> , ext. Mousse de Chêne absolute Oakmoss absolute (<i>Evernia prunastri</i>) Evernia prunastri (Oakmoss) extract		

History:	Initial reviews:	April 1991, July 2001		
	Current revision date:	2008		
	Implementation date:	For new submissions*: February 16, 2009		
		For existing fragrance compounds*:	February 16, 2011	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.02 %	Category 7	0.1 %			
Category 2	0.03 %	Category 8	0.1 %			
Category 3	0.1 %	Category 9	0.1 %			
Category 4	0.1 %	Category 10	0.1 %			
Category 5	0.1 %	Category 11	See Note Box (2)			
Category 6	0.5 %					
Note box:						

For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again.

In the presence of tree moss extracts, the level of oak moss in the respective category has to be reduced accordingly such that the total amount of both extracts does not exceed the maximum permitted level in each category as listed in the table above. If the same compound is intended for more than one IFRA QRA category, then the most restrictive limitation (based on foreseen use concentrations and maximum permitted level) will apply.

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) (http://www.iofiorg.org/).

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Oakmoss extracts

Fragrance material specifications:	Oak moss extracts used in fragrance compounds must not contain added tree moss, which is a source of resin acids.
	Traces of resin acids may be carried over to commercial qualities of oak moss in the manufacturing process. These traces must not exceed 0.1% (1000 ppm) dehydroabietic acid (DHA) in the extract.
	The concentration of resin acids in oak moss can be measured with an HPLC Reverse Phase – spectrofluorometry method.
	Further, levels of atranol and chloroatranol should each be below 100 ppm in oak moss extracts.

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
970	Moderate	7004	1724⁴	1417	700

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³ WoE NESIL limited to three significant figures

⁴ MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Oakmoss extracts and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 700 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Oakmoss extracts in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1973. Human Maximization Test. RIFM report number 1802, October 9a (RIFM, Woodcliff Lake, NJ, USA)

Research Institute for Fragrance materials, Inc. (1989). Human repeated insult patch test of oakmoss absolute. RIFM report number 12360, 31 October.

Research Institute for Fragrance Materials, Inc. (1989). Human repeated insult patch test of oakmoss absolute. RIFM report number 12361, 31 October.

Research Institute for Fragrance Materials, Inc. (1990). Human repeated insult patch test of oakmoss absolute. RIFM report number 12380, 1 March.

Research Institute for Fragrance Materials, Inc. (1990). Human repeated insult patch test on oakmoss absolute. RIFM report number 14118, 26 November.

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Local Lymph Node Assay. RIFM report number 50881, June 30 (RIFM, Woodcliff Lake, NJ, USA).

OTNE (1-(1,2,3,4,5,6,7,8 Octahydro-2,3,8,8-tetramethyl-2-naphthalenyl)ethanone

CAS N°:	54464-57-2	Empirical formula: Structure:	
Synonyms:	1-(1,2,3,4,5,6,7,8-octahydro-2,3,8,8-te 1-(1,2,3,4,5,6,7,8-Octahydro-2,3,8,8-te 1-(1,2,3,4,5,6,7,8-Octahydro-2,3,8,8-te Amberonne Boisvelone Isocyclemone E Iso E super	tramethyl-2-naphthaleny etramethyl-2-aphthalenyl etramethyl-2-naphthyl)etl	I))ethanone nan-1-one

History:	Initial reviews:	New Standard		
	Current revision date:	2008		
	Implementation date:	For new submissions*: August 16, 2008		
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Limits in the infished product.						
Category 1 See Note box (1)	1.34 %	Category 7	3.6 %			
Category 2	1.73 %	Category 8	2.0 %			
Category 3	7.1 %	Category 9	5.0 %			
Category 4	21.4 %	Category 10	2.5 %			
Category 5	11.2 %	Category 11	See Note Box (2)			
Category 6	34.2 %					
Note hox:						

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry)

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

(http://www.iofiorg.org/).

OTNE (1-(1,2,3,4,5,6,7,8 Octahydro-2,3,8,8-tetramethyl-2-naphthalenyl)ethanone

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

LINA weighted mean EC2 values		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
6825	Weak	47 244 ⁴	NA	NA	47 200

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³ WoE NESIL limited to three significant figures

⁴ MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for OTNE and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 47200 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of OTNE in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Human Repeated Insult Patch Test with OTNE. RIFM report number 45124, June 28. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Local Lymph Node Assay with OTNE. RIFM report number 51630, November 9. (RIFM, Woodcliff Lake, NJ, USA).



CAS N°:	2442-10-6	Empirical formula: Structure:	C10H18O2
Synonyms:	3-Acetoxyoctene Amyl crotonyl acetate Amyl vinyl carbinyl acetate 1-Octen-3-ol, acetate Octenyl acetate beta-Octenyl acetate n-Pentyl vinyl carbinol acetate		

History:	Initial reviews:	July 1989, July 1994, May 2007		
	Current revision date:	2008		
	Implementation date:	For new submissions*: August 16, 2008		
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.1 %	Category 7	0.3 %			
Category 2	0.1 %	Category 8	0.3 %			
Category 3	0.3 %	Category 9	0.3 %			
Category 4	0.3 %	Category 10	0.3 %			
Category 5	0.3 %	Category 11	See Note Box (2)			
Category 6	2.5 %					
Note box:						

For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again.

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) (http://www.iofiorg.org/).

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

1-Octen-3-yl acetate

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
>7500 [1]	Extremely weak	3543	NA	6900	3500

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max tests

² Gerberick et al., 2001

³ WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 1-octen-3-yl acetate and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 3500 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of 1-octen-3-yl acetate in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

RIFM (Research Institute for Fragrance Materials, Inc.), 1974a. Report on human maximization studies. Report to RIFM. RIFM report number 1779, June 06 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1974b. Report on human maximization studies. Report to RIFM. RIFM report number 1779, August 20 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1985. Report on human maximization studies. Report to RIFM. RIFM report number 1779, January 7a (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1988. Repeat insult patch test of 1-octen-3-yl acetate in human subjects. RIFM report number 8516, December 07 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. 1-Octen-3-yl acetate: Local Lymph Node Assay. Unpublished report from International Flavors and Fragrances, 13 December. Report number 47816 (RIFM, Woodcliff Lake, NJ, USA).

Opoponax

CAS N°:	8021-36-1 9000-78-6 93384-32-8	Empirical formula:	N/A
Synonyms:	Opoponax (absolute, resinoid, oil, gu Bisabol-myrrh Sweet myrrh Opoponax chironium (L.) W.D.J. Koo Commiphora erythraea Engler var. g	um, tincture) ch glabrescens (Burseraceae)	

History:	Initial reviews:	March 1978, July 1994		
	Current revision date:	June 2013		
	Implementation date:	For new submissions*:	January 10, 2014	
		For existing fragrance compounds*:	January 10, 2015	
	Next review date	2018		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.03%	Category 7	0.08%			
Category 2	0.04%	Category 8	0.60%			
Category 3	0.15%	Category 9	0.60%			
Category 4	0.45%	Category 10	0.60%			
Category 5	0.24%	Category 11	See Note box (2)			
Category 6	0.60%					

Note box:

(1) See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products with the potential of ingestion.

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:	Opoponax oil can be obtained from solvent extraction or pyrolysis. Opoponax oil obtained through pyrolysis shall be rectified according to Good Manufacturing Practices and the content of polynuclear aromatic hydrocarbons (PAH) resulting from their use shall respect the following
	Benzopyrene and 1,2-Benzanthracene are to be used as markers for PAH. If used alone or in combination with rectified Cade oil, rectified Birch tar oils or rectified Styrax oil, the total concentration of both of the markers should not exceed 1 ppb in the final product.

Opoponax

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
4450 – 5625 [2] ⁴	Weak	NA		NA	1000

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures. A default value based on the LLNA data was employed because the material is used a very low volume and there are no HRIPT data. ⁴ A range of values and not the weighted mean was provided because three studies were performed on three materials having

⁴ A range of values and not the weighted mean was provided because three studies were performed on three materials having different compositions – opoponax essential oil, opoponax extract and opoponax pyrogenated. Of these LLNAs, opoponax essential oil and opoponax pyrogenated resulted in a positive response.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Opoponax (all forms) and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1000 μ g/cm², which is a default value based on the LLNA data. They recommend the limits for the 11 different product categories, which are the acceptable use levels of Opoponax (all forms) in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick G, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. Regulatory Toxicology and Pharmacology 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. Local Lymph Node Assay. Draft RIFM Report number 63817. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. Local Lymph Node Assay. Draft RIFM Report number 63818. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. Local Lymph Node Assay. Draft RIFM Report number 63819. (RIFM, Woodcliff Lake, NJ, USA).

1-(2,4,4,5,5-Pentamethyl-1-cyclopenten-1-yl)ethan-1-one

CAS N°:	13144-88-2	Empirical formula: Strucutre:	C ₁₂ H ₂₀ O
Synonyms:	2-Acetyl-1,3,3,4,4-pentamethyl-1 Ethanone, 1-(2,4,4,5,5-pentamet	-cyclopentene thyl-1-cyclopenten-1-yl)-	
	1-(2,4,4,5,5-Pentamethylcyclope Alpinone (trade name)	nt-1-en-1-yl)ethanone	

History:	Initial reviews:	New Standard	
	Current revision date:	June 20, 2011	
	Implementation date:	For new submissions*:	August 20, 2011
		For existing fragrance compounds*:	August 20, 2012
	Next review date	2016	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.03 %	Category 7	0.08 %		
Category 2	0.04 %	Category 8	1.01 %		
Category 3	0.15 %	Category 9	5.00 %		
Category 4	0.45 %	Category 10	2.50 %		
Category 5	0.24 %	Category 11	See Note Box (2)		
Category 6	0.72 %				
Note box:					
			0 1 1 1 1		

(1) See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products with the potential of ingestion.

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None known at the time of the publication of the Standard

CRITICAL EFFECT:

SENSITIZATION

1-(2,4,4,5,5-Pentamethyl-1-cyclopenten-1-yl)ethan-1-one

RIFM SUMMARIES:

		Human Data				
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)	
3600 [1]	Weak	NA	NA	NA	1000	

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003. ²Data derived from HRIPT or HMT. ³WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 1-(2,4,4,5,5-Pentamethyl-1-cyclopenten-1-yl)ethan-1-one and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1000 μ g/cm², which is a default value based on the LLNA data. They recommend the limits for the 11 different product categories, which are the acceptable use levels of 1-(2,4,4,5,5-Pentamethyl-1-cyclopenten-1-yl)ethan-1-one in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick G, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.



2-Pentylidene cyclohexanone

CAS N°:	25677-40-1	Empirical formula: Structure:	C11H18O
Synonyms:	Cyclohexanon	ie, 2-pentylidene-	

History:	Initial reviews:	February 1979, May 1983		
	Current revision date:	September 2002		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products:	N/A				
Note box:					
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.					
Fragrance material specifications:		N/A			

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke, D.L.J. and Letizia, C. (1982), Food and Chemical Toxicology, 20, 797.

Peru balsam crude

CAS N°:	8007-00-9	Empirical formula:	N/A
Synonyms:	Exudation of Myroxylon pereirae Klotsch	1 -	

History:	Initial reviews:	October 1974, December 1991	
	Current revision date:	2007	
	Implementation date:	For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:							
Skin contact products:	Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A				
		Including household	cleaning products				
Non skin contact products:	N/A						
Note box:							
Restrictions on the use of Peru balsam oil, absolute and anhydrol are contained in a separate Standard (Peru balsam extracts and distillates).							
Fragrance material specifications:		N/A					

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, January 2007.

REFERENCES:

D.L. Opdyke (1974), Fd. Cosmet. Toxicol. 12, 951 and 953

Peru balsam extracts and distillates

CAS N°:	8007-00-9	Empirical formula:	N/A
Synonyms:	Balsam oil, Peru (<i>Myroxylon pereirae</i> Ki Balsams, Peru <i>Myroxylon pereirae</i> (Balsam Peru) oil <i>Myroxylon pereirae</i> (Balsam Peru) resin <i>Myroxylon pereirae</i> oil Peru balsam absolute Peru balsam anhydrol	otzsch)	

History:	Initial reviews:	October 1974, December 1991, May 2007		
	Current revision date:	2008		
	Implementation date:	For new submissions*: August 16, 2008		
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.03 %	Category 7	0.07 %		
Category 2	0.04 %	Category 8	0.4 %		
Category 3	0.1 %	Category 9	0.4 %		
Category 4	0.4 %	Category 10	0.4 %		
Category 5	0.2 %	Category 11	See Note Box (2)		
Category 6	0.7 %				
Note box:					

For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again.

The use of <u>Peru balsam crude</u> is **PROHIBITED** in a separate Standard.

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) (http://www.iofiorg.org/).

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

Peru balsam extracts and distillates

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

LLNA weighted mean EC3 values (μg/cm²) [no. studies]		Human Data			
	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
Balsam oil, Peru (Myroxylon pereirae Klotzsch)					
987	Moderate	950	NA	NA	950
Peru balsam absolute					
625	Moderate	NA	NA	NA	950
Peru balsam anhydrol					
NA	Moderate	NA	NA	NA	950

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max tests

² Gerberick et al., 2001

³ WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Peru balsam extracts and distillates and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 950 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Peru balsam extracts and distillates in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

Peru balsam (myroxylon pereirae klotzch)

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Local Lymph Node Assay on Peru balsam (Myroxylon pereirae Klotzsch). RIFM report number 44372, February 16(RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2005. Repeated insult patch test with balsam, Peru (Myroxylon pereirae Klotzsch). RIFM report number 47380, January 20 (RIFM, Woodcliff Lake, NJ, USA).

Peru balsam absolute

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Local Lymph Node Assay on Peru balsam absolute. RIFM report number 44371, February 16 (RIFM, Woodcliff Lake, NJ, USA).

Phenylacetaldehyde

CAS N°:	122-78-1	Empirical formula: Structure:	C ₈ H ₈ O
Synonyms:	Benzeneacetaldehyde Benzylcarboxaldehyde Hyacinthin 1-Oxo-2-phenylethane Phenylacetic aldehyde Phenyl Acetic Aldehyde (pure) alpha-Tolualdehyde alpha-Toluic aldehyde		

History:	Initial reviews:	October 1975, February 1980	
	Current revision date:	2006	
	Implementation date:	For new submissions*: June 11, 2007	
		For existing fragrance compounds*:	June 11, 2008
	Next review date	2011	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.02 %	Category 7	0.04 %			
Category 2	0.02 %	Category 8	0.6 %			
Category 3	0.09 %	Category 9	3.0 %			
Category 4	0.3 %	Category 10	2.5 %			
Category 5	0.1 %	Category 11	See Note Box			
Category 6	0.4 %					
Note box:						
Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.						
This Standard cancels and replaces the existing one on phenylacetaldehyde , which was based on the no longer supported 'quenching' phenomenon.						

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the introduction to the IFRA Standards)

Phenylacetaldehyde

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

Phenylacetaldehyde - Sensitization Potency Estimation Based on Weight of Evidence

LLNA weighted mean EC3 values (µg/cm²) [no. studies] Based on Animal Data ²		Human Data			
	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)	
962 [2]	Moderate	591	NA	1181	590

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; MAX = Human Maximization Test;

LOEL = lowest observed effect level; NA = Not Available

¹ Data derived from HRIPT or Human Max tests

² Gerberick et al., 2001

³ WoE NESIL limited to two significant figures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for phenylacetaldehyde and based on the weight of evidence established the No Expected Sensitization Induction Level (NESIL) as 590 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of phenylacetaldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of March 15, 2006.

REFERENCES:

D.A.Basketter, Z.M.Wright, E.V.Warbrick, R.J.Dearman, I.Kimber, C.A.Ryan, G.F.Gerberick and I.R.White (2001). Human potency predictions for aldehydes using the local lymph node assay. Contact Dermatitis, 45(2), 89-94.

D.A.Basketter, N.Gilmour, R.J.Dearman, I.Kimber, C.A.Ryan and G.F. Gerberick (2003). Classification of skin sensitisation potency using the Local Lymph Node Assay. The Toxicologist, 72(S-1), 101.

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161

QRA Expert Group (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

Research Institute for Fragrance Materials, Inc (2003). Repeated insult patch test in human subjects with phenylacetaldehyde. RIFM report number 44245 (RIFM, Woodcliff Lake, NJ USA).

Research Institute for Fragrance Materials, Inc (2004). Repeated insult patch test in human subjects with phenylacetaldehyde. RIFM report number 45132 (RIFM, Woodcliff Lake, NJ USA).



CAS N°:	103-79-7	Empirical formula: Structure:	C9H10O
Synonyms:	Benzyl methyl ketone Methyl benzyl ketone 2-Propanone, 1-phenyl		

History:	Initial reviews:	Not applicable		
	Current revision date:	Not applicable		
	Implementation date:	For new submissions*: Not applicable		
		For existing fragrance compounds*:	Not applicable	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



Phenyl acetone (Methyl benzyl ketone)

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000



History:	Initial reviews:	Not applicable	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX



Phenyl benzoate

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

- 1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or
- 2) adverse data on the material itself and/or
- 3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000



3-Phenylbutanal

CAS N°:	16251-77-7	Empirical formula: Strucutre:	C ₁₀ H ₁₂ O H
Synonyms:	Benzenepropanal, β-methyl- 3-Phenylbutanal 3-Phenylbutyraldehyde 3-Phenyl-3-methylpropanal Trifernal (commercial name)		

History:	Initial reviews:	New Standard		
	Current revision date:	June 11, 2010		
	Implementation date:	For new submissions*:	January 11, 2011	
		For existing fragrance compounds*:	January 11, 2012	
	Next review date	March 2015		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.17%	Category 7	0.45%		
Category 2	0.22%	Category 8	2.0%		
Category 3	0.89%	Category 9	5.0%		
Category 4	2.7%	Category 10	2.5%		
Category 5	1.4%	Category 11	See Note Box (2)		
Category 6	4.3%				
Note box:					
(1) See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products with the potential of ingestion.					

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None known at the time of the publication of the Standard

CRITICAL EFFECT:

SENSITIZATION

3-Phenylbutanal

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
N/A	N/A	5906	N/A	12,500	5900

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003. ²Data derived from HRIPT or HMT. ³WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 3-phenylbutanal and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 5900 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of 3-phenylbutanal in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick G, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 2009. Human repeated insult patch test. RIFM report number 57513, July 16. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1983. Human repeated insult patch test. Unpublished study from Firmenich, Inc., August 24. Report number 40514. (RIFM, Woodcliff Lake, NJ, USA).

2-Phenylpropionaldehyde

CAS N°:	93-53-8	Empirical formula: Structure:	C ₉ H ₁₀ O O
Synonyms:	Benzeneacetaldehyde, α-meth Hydratropaldehyde Hydratropic aldehyde α-Methylphenylacetaldehyde α-Methyltolualdehyde 2-Phenylpropanal α-Phenylpropionaldehyde	nyl-	

History:	Initial reviews:	New Standard		
	Current revision date:	2009		
	Implementation date:	For new submissions*: August 7, 2009		
		For existing fragrance compounds*:	August 7, 2011	
	Next review date	2014		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.01 %	Category 7	0.03 %			
Category 2	0.01 %	Category 8	0.40 %			
Category 3	0.06 %	Category 9	1.90 %			
Category 4	0.17 %	Category 10	2.50 %			
Category 5	0.09 %	Category 11	See Note Box (2)			
Category 6	0.28 %					
Note box:						
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofiorg.org)						
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to negligible skin contact the concentration of a fragrance ingredient should not exceed the usual concentration of the fragrance compound in the finished product.						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the introduction to the IFRA Standards)

2-Phenylpropionaldehyde

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ²	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
1575 [1] ⁴	Weak	3885	1380	1938	380 ⁶

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed

effect level; NA = Not Available.

¹ Data derived from HRIPT or HMT.

²Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

³ WoE NESIL limited to two significant figures.

⁴ EC3 value from one LLNA, not the mean.

⁵ HRIPT with 38 subjects only; study with 100 subjects not done because the material is used at a low volume (1-5 metric tons) ⁶ WoE NESIL based on limited subject HRIPT which was lower than the default LLNA (1000 ug/cm2)

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 2-Phenylpropionaldehyde and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 380 mg/cm2. They recommend the limits for the 11 different product categories, which are the acceptable use levels of 2-Phenylpropionaldehyde in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Patlewicz, G., Roberts, D.W., Walker, J.D., 2003. QSARs for the skin sensitization potential of aldehydes and related compounds. QSAR & Combinatorial Science, 22, 196-203.

RIFM (Research Institute for Fragrance Materials, Inc.), 1964. Human repeated insult patch test. Unpublished study from IFF Inc., 3 April. Report number 51926. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1971a. Human repeated insult patch test. Unpublished study from IFF Inc., 7 July. Report number 51925. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1971b. Maximization study with 2-phenylpropionaldehyde. RIFM report number 1805, April 2a. (RIFM, Woodcliff Lake, NJ, USA).



Pinacea derivatives

CAS N°:	N/A	Empirical formula:	N/A
Synonyms:	N/A		

History:	Initial reviews:	May 1976, July 1994	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

SPECIFICATION

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

Essential oils (e.g. turpentine oil) and isolates (e.g. delta-3-carene) derived from the *Pinacea* family, including *Pinus* and *Abies* genera, should only be used when the level of peroxides is kept to the lowest practicable level, for instance by adding antioxidants at the time of production. Such products should have a peroxide value of less than 10 millimoles peroxide per liter, determined according to the FMA method, which can be downloaded from the IFRA website (see Analytical Methods).

REFERENCES:

This recommendation is based on the published literature, mentioning sensitizing properties when containing peroxides (Fd. Cosmet. Toxicol. 11, 1053 (1973); 16, 843 (1978);16, 853(1978).

This Guideline does not permit the use of colophony.

Synonyms: 1/369-59-4 Empirical formula: Structure: C11H10O2 Synonyms: 1(3H)-Isobenzofuranone, 3-propylidene-Propylidene phthalide 3-Propylidenephthalide 3-propylidene-Propylidenephthalide

History:	Initial reviews:	May 1977, July 1994 (28th Amendment)		
	Current revision date:	2008		
	Implementation date:	For new submissions*:	August 16, 2008	
		For existing fragrance compounds*:	August 16, 2010	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.01 %	Category 7	0.01 %		
Category 2	0.01 %	Category 8	0.01 %		
Category 3	0.01 %	Category 9	0.01 %		
Category 4	0.01 %	Category 10	0.01 %		
Category 5	0.01 %	Category 11	See Note Box (2)		
Category 6	0.7 %				
Note box:					
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be re-evaluated again.					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) (http://www.iofiorg.org/).					
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.					
Fragrance material specifications: N/A					

3-Propylidenephthalide

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

			Human Data		
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
350	Moderate	945 ⁴	3454	2760	920

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³ WoE NESIL limited to three significant figures

⁴ MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 3-Propylidenephthalide and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 920 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of 3-Propylidenephthalide in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Gerberick, G.F., Ryan, C.A., Kern, P.S., Dearman, R.J., Kimber, I., Patlewicz, G.Y.,

Basketter, D.A., 2004. A chemical dataset for evaluation of alternative approaches to skinsensitization testing. Contact Dermatitis 50, 274-288.

RIFM (Research Institute for Fragrance Materials, Inc.), 1975. Human Maximization Test. RIFM report number 1799, June 16a (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1978. Human Maximization Test. RIFM report number 1787, May 1 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007. Human Repeated Insult Patch Test. RIFM report number 53724, September 21a (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2008. Local Lymph Node Assay. DRAFT RIFM data, report 54427 (RIFM, Woodcliff Lake, NJ, USA).

Pseudoionone (2,6-Dimethylundeca-2,6,8-trien-10-one)

CAS N°:	141-10-6	Empirical formula: Structure:	C13H20O
Synonyms:	Citrylideneacetone 6,10-Dimethyl-3,5,9-unde 3,5,9-Undecatrien-2-one,	ecatrien-2-one 6,10-dimethyl-	

History:	Initial reviews:	February 1979, July 1987, April 1989	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED / SPECIFICATION

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
Pseudoionone should not be used as fragrance ingredient as such, but a level of up to 2% as an impurity in ionones is accepted.						
Fragrance material specification	ons:	N/A				

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)

CRITICAL EFFECT:

SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke D.L.J. (1975), Food and Cosmetics Toxicology 13, 549.

Ford R.A. et al. (1988), Food and Chemical Toxicology 26, 311.

Pseudo methylionones

CAS N°:	26651-96-7 72968-25-3 1117-41-5	Empirical formula:	C ₁₄ H ₂₂ O
Synonyms:	2,6-Dimethyldodeca-2,6,8-trien-10-one 7,11-Dimethyl-4,6,10-dodecatrien-3-on 7,11-Dimethyldodeca-4,6,10-trien-3-on 4,6,10-Dodecatrien-3-one, 7,11-dimeth 3,6,10-Trimethylundeca-3,5,9-trien-2-o	e e yl- ne	

History:	Initial reviews:	February 1979, October 1979, April 1989, September 2002, May 2006		
	Current revision date:	2009		
	Implementation date:	For new submissions*:	August 7, 2009	
		For existing fragrance compounds*:	August 7, 2010	
	Next review date	Not applicable		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED / SPECIFICATION

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
Pseudo methylionones should not be used as fragrance ingredients as such, but a level of up to 2% as an impurity in methylionones is accepted.						
Fragrance material specification	ons:	N/A				

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)

CRITICAL EFFECT:	SENSITIZATION

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Opdyke, D.L.J. (1975), Food and Cosmetics Toxicology 13, 863.

Ford R.A. et al. (1988), Food and Chemical Toxicology 26, 305 and 413.

Quinoline

CAS N°:	91-22-5	Empirical formula: Structure:	C9H7N
Synonyms:	1-Benzazine 2,3-Benzopyridine Benzo(b)pyridine Chinoleine Leucoline Quinoleine		

History:	Initial reviews:	New Standard	
	Current revision date:	June 11, 2010	
	Implementation date:	For new submissions*: August 11, 2010	
		For existing fragrance compounds*:	August 11, 2011
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:			
Skin contact products:			
Leave on products:	N/A	Rinse-off products:	N/A
		Including household	cleaning products
Non skin contact products:	N/A		
Note box:			
The material is considered as a carcinogenic / mutagenic agent and had only minor reported use in fragrances. REXPAN agreed to the IFRA policy to ban the material for the given effects and not further elaborate on a risk management policy.			
Fragrance material specifications: N/A			

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

CARCINOGENICITY, MUTAGENICITY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, June 2010.

REFERENCES:

Commission Directive 2009/2/EC (31st ATP to Directive 67/548/EEC)

	Ro	se ketones	
CAS N°:	23696-85-7	Empirical formula:	$C_{13}H_{20}O$
	23726-93-4		
	43052-87-5		
	24720-09-0		
	23726-94-5		
	23726-92-3		
	23726-91-2		
	57378-68-4		
	71048-82-3		
	39872-57-6		
	70266-48-7		
	33073-71-1		
	35047-49-1		
	33044-08-9		
Synonyme	23696-85-7 (C13H18O)		1
Cynonyms.	1-(2.6.6-Trimethylcyclohexa-1.3-di	envl)-2-buten-1-one	
	2-Buten-1-one, 1-(2,6,6-trimethyl-1	1,3-cyclohexadien-1-yl)	
	Damascenone, Floriffone		
	23726-93-4 (C ₁₃ H ₁₈ O)		
	(E)-1-(2,6,6-Trimethyl-1,3-cyclohez	xadien-1-yl)-2-buten-1-one	
	trans-1-(2,6,6-Trimethyl-1,3-cycloh	nexadien-1-yl)-2-buten-1-one	
	β-Damascenone		
	42052.97.5		
	43032-67-5	-1-v/l)-2-buten-1-one	
	2-Buten-1-one 1-(2.6.6-trimethyl-2	2-cvclobexen-1-vl)-	
	α -Damascone Dihydrofloriffone α		
	24720-09-0		
	(E)-1-(2,6,6-Trimethyl-2-cyclohexe	n-1-yl)-2-buten-1-one	
	trans-1-(2,6,6-Trimethyl-2-cyclohe	xen-1-yl)but-2-en-1-one	
	2-Buten-1-one, 1-(2,6,6-trimethyl-2	2-cyclohexen-1-yl)-, (2E)-	
	trans-α-Damascone		
	00700 04 5		
	23/26-94-5		
	(Z)-1-(Z,6,6-1 nmethyl-2-cyclonexe	n-1-yi)-2-buten-1-one	
	2-Buten-1-one 1-(2.6.6-trimethyl-2	$r_1 - r_2 r_2$	
	cis-a-Damascone		
	23726-92-3		
	1-(2,6,6-Trimethylcyclohex-1-en-1-	-yl)but-2-en-1-one	
	(Z)-β-1-(2,6,6-Trimethyl-1-cyclohe	xen-1-yl)-2-buten-1-one	
	(Z)-1-(2,6,6-Trimethyl-1-cyclohexe	n-1-yl)-2-buten-1-one	
	2-Buten-1-one, 1-(2,6,6-trimethyl-1	I-cyclohexen-1-yl)-, (2Z)-	
	cis-β-Damascone, Damasione		
	23/26-91-2		
	(2E)-1- $(2,6,6-I$ rimethyl-1-cyclohex		
	(\Box) -1-(2,0,0-1)IIIIet(I)/-1-Cyclonexe	$\frac{1}{1}$ $\frac{1}{2}$ $\frac{1}$	
	r-(∠,0,0-11) trans-β-Damascone Dibudrofloriff	nne B	
		nie h	



Rose ketones

57378-68-4

δ-1-(2,6,6-Trimethyl-3-cyclohexen-1-yl)-2-buten-1-one 2-Buten-1-one, 1-(2,6,6-trimethyl-3-cyclohexen-1-yl)-1-(2,6,6-Trimethyl-3-cyclohexen-1-yl)-2-buten-1-one

δ-Damascone, Dihydrofloriffone TD

71048-82-3

 $[1\alpha(E),2\beta]$ -1-(2,6,6-Trimethyl-3-cyclohexen-1-yl)-2-buten-1-one $[1\alpha(E),2\beta]$ -1-(2,6,6-Trimethylcyclohex-3-en-1-yl)but-2-en-1-one trans,trans- δ -Damascone

39872-57-6

1-(2,4,4-Trimethyl-2-cyclohexen-1-yl)-2-buten-1-one (E)-1-(2,4,4-Trimethyl-2-cyclohexen-1-yl)-2-buten-1-one 2-Buten-1-one, 1-(2,4,4-trimethyl-2-cyclohexen-1-yl)-, (2E)-2-Buten-1-one, 1-(2,4,4-trimethyl-2-cyclohexen-1-yl)-, (E)-Isodamascone (high α)

70266-48-7

1-(2,4,4-Trimethyl-1-cyclohexen-1-yl)-2-buten-1-one 2-Buten-1-one, 1-(2,4,4-trimethyl-1-cyclohexene-1-yl) Isodamascone (standard quality)

33673-71-1

1-(2,4,4-Trimethylcyclohex-2-en-1-yl)but-2-en-1-one 1-(2,4,4-Trimethyl-2-cyclohexen-1-yl)-2-buten-1-one 2-Buten-1-one, 1-(2,4,4-trimethyl-2-cyclohexen-1-yl)-Isodamascone (isomer unspecified)

35087-49-1

1-(2,2-Dimethyl-6-methylenecyclohexyl)but-2-en-1-one 2-Buten-1-one, 1-(2,2-dimethyl-6-methylenecyclohexyl)-Damascone g-, γ-Damascone

35044-68-9

2-Buten-1-one, 1-(2,6,6-trimethyl-1-cyclohexen-1-yl)-2,6,6-Trimethyl-1-(2-butenoyl)-1-cyclohexene 2,6,6-Trimethyl-1-crotonoyl-1-cyclohexene 1-(2,6,6-Trimethylcyclohexenyl)-2-buten-1-one 1-(2,6,6-Trimethyl-1-cyclohexen-1-yl)-2-buten-1-one Damascone b-, β -Damascone

History:	Initial reviews:	December 1991, December 1995, July 2007 (42th Amendment), July 2008 (43th Amendment)	
	Current revision date:	2009	
	Implementation date:	For new submissions*:	August 7, 2009
		For existing fragrance compounds*:	August 7, 2011
	Next review date	2014	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

Rose ketones

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Category 1 See Note box (1)	0.003 %	Category 7	0.008 %	
Category 2	0.004 %	Category 8	0.02 %	
Category 3	0.02 %	Category 9	0.02 %	
Category 4	0.02 %	Category 10	0.02 %	
Category 5	0.02 %	Category 11	See Note Box (2)	
Category 6	0.07 %			
Noto hov:				

For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be re-evaluated again.

The above limits apply to Rose Ketones used individually or in combination.

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) (http://www.iofiorg.org/).

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

See Enclosures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Rose Ketones and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 100 µg/cm2. They recommend the limits for the 11 different product categories, which are the acceptable use levels of Rose Ketones in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

Damascenone, 1-(2,6,6-Trimethylcyclohexa-1,3-dienyl)-2-buten-1-one (23696-85-7)

RIFM (Research Institute for Fragrance Materials, Inc.), 1978. Repeated insult patch test of 1-(2,6,6-trimethylcyclohexa-1,3-dienyl)-2-buten-1-one in human subjects. RIFM report number 15395, July 12 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001. Murine local lymph node assay with 1-(2,6,6-trimethylcyclohexa-1,3-dienyl)-2-buten-1one in mice. Unpublished report from Firmenich Incorporated, Report number 38813 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2002. 1-(2,6,6-Trimethylcyclohexa- 1,3-dienyl)-2-buten-1-one: local lymph node assay in mice. Unpublished report from Firmenich Incorporated, 22 October. Report number 41991 (RIFM, Woodcliff Lake, NJ, USA).

N/A

Rose ketones

alpha-Damascone, alpha-1-(2,6,6-Trimethyl-2-cyclohexen-1-yl)-2-buten-1-one (43052-87-5)

RIFM (Research Institute for Fragrance Materials, Inc.), 1979a. Evaluation of potential hazards of alpha-1-(2,6,6-trimethyl-2-cyclohexen-1-yl)-2-buten-1-one by dermal contact in human subjects. Unpublished report from IFF Inc., 17 September. RIFM report number 15397. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1979b. Repeated insult patch test of alpha-1-(2,6,6-trimethyl-2-cyclohexen-1-yl)-2-buten-1-one in humans. Unpublished report from Firmenich Inc., 19 December. RIFM report number 153414. (RIFM, Woodcliff Lake, NJ, USA). RIFM (Research Institute for Fragrance Materials, Inc.), 1985. Maximization study of alpha-1-(2,6,6-trimethyl-2-cyclohexen-1-yl)-2-buten-1-one in

humans. Unpublished report from Firmenich Inc., 29 July. RIFM report number 15416. (RIFM, Woodcliff Lake, NJ, USA).

trans-alpha-Damascone, (E)-1-(2,6,6-Trimethyl-2-cyclohexen-1-yl)-2-buten-1-one (24720-09-0)

RIFM (Research Institute for Fragrance Materials, Inc.), 2001. Murine local lymph node assay with 2-buten-1-one, 1-(2,6,6-trimethyl-2-cyclohexen-1-yl)-, (2E)- in mice.

Unpublished report from Firmenich Incorporated, Report number 38814 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001a. Repeated insult patch study of 2-buten-1-one, 1-(2,6,6-trimethyl-2-cyclohexen-1-yl)-, (2E)- at 0.5% in diethyl phthalate (DEP). Unpublished report from Firmenich Incorporated, Report number 38815 (RIFM, Woodcliff Lake, NJ, USA).

cis-beta-Damascone, (Z)-B-1-(2,6,6-Trimethyl-1-cyclohexen-1-yl)-2-buten-1-one (23726-92-3)

RIFM (Research Institute for Fragrance Materials, Inc.), 1979. Repeated insult patch test of beta-damascone in human subjects. Unpublished report from IFF, Inc., 19 February. RIFM report number 15394. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1980. Repeat insult patch test with (Z)-b-1-(2,6,6-trimethyl-1-cyclohexen-1-yl)-2-buten-1-one. Unpublished report from IFF, Inc., 17 June. Report number 47353. (RIFM, Woodcliff Lake, NJ, USA).

trans-beta-Damascone, (2E)-1-(2,6,6-Trimethyl-1-cyclohexen-1-yl)-2-buten-1-one (23726-91-2)

RIFM (Research Institute for Fragrance Materials, Inc.), 1979. Repeated insult patch test of (2E)-1-(2,6,6-trimethyl-1-cyclohexen-1-yl)-2-buten-1-one in humans. Unpublished report from Firmenich, Inc., 19 December. Report number 15407. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2001. Murine local lymph node assay with (2E)-1-(2,6,6-trimethyl-1-cyclohexen-1-yl)-2-buten-1-one.

Unpublished report from Firmenich, Inc. Report number 38811. (RIFM, Woodcliff Lake, NJ, USA).

delta-Damascone, delta-1-(2,6,6-Trimethyl-3-cyclohexen-1-yl)-2-buten-1-one (57378-68-4)

RIFM (Research Institute for Fragrance Materials, Inc.), 1982. Evaluation of potential irritation and sensitization hazards of delta-damascone by dermal contact in humans.

Unpublished report from International Flavors and Fragrances, 17 November. Report number 15399 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2002a. Murine local lymph node assay of delta-1-(2,6,6-trimethyl-3-cyclohexen-1-yl)-2-buten-1one. Unpublished report from Firmenich Incorporated, 9 July. Report number 41992 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2002b. Murine local lymph node assay of delta-1-(2,6,6-trimethyl-3-cyclohexen-1-yl)-2-buten-1one. Unpublished report from Firmenich Incorporated, 2 April. Report number 42139 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. delta-1-(2,6,6-Trimethyl-3- cyclohexen-1-yl)-2-buten-1-one: Local Lymph Node Assay. Unpublished report from International Flavors and Fragrances, 13 December. Report number 47819 (RIFM, Woodcliff Lake, NJ, USA).

trans,trans-delta-Damascone, [1alpha(E),2beta]-1-(2,6,6-Trimethyl-3-cyclohexen-1- yl)-2-buten-1-one (71048-82-3)

RIFM (Research Institute for Fragrance Materials, Inc.), 1978. Repeated insult patch test with [1.alpha.(E),2.beta.]-1-(2,6,6-trimethyl-3-cyclohexen-1yl)-2-buten-1-one. Unpublished report from International Flavors and Fragrance, 11 August. Report number 50614 (RIFM, Woodcliff Lake, NJ, USA).

Isodamascone (high alpha), (E)-1-(2,4,4-Trimethyl-2-cyclohexen-1-yl)-2-buten-1-one (39872-57-6)

RIFM (Research Institute for Fragrance Materials, Inc.), 1994. Repeated insult patch test on isodamascone on human subjects. RIFM report number 25751, July 25 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1995. Repeated insult patch test of isodamascone in human subjects. RIFM report number 25753, June 20 (RIFM, Woodcliff Lake, NJ, USA).

Isodamascone (standard quality), 1-(2,4,4-Trimethyl-1-cyclohexen-1-yl)-2-buten-1-One (70266-48-7)

RIFM (Research Institute for Fragrance Materials, Inc.), 1985. Repeated insult patch test with isodamascone. RIFM report number 41231, December 22 (RIFM, Woodcliff Lake, NJ, USA).

gamma-Damascone, 1-(2,2-Dimethyl-6-methylenecyclohexyl)but-2-en-1-one (35087-49-1)

RIFM (Research Institute for Fragrance Materials, Inc.), 2001. Murine local lymph node assay with 2-buten-1-one, 1-(2,2-dimethyl-6methylenecyclohexyl)- in mice. Unpublished report from Firmenich Inc. RIFM report number 38812. (RIFM, Woodcliff Lake, NJ, USA). ra

Rue oil

CAS N°:	8014-29-7	Empirical formula: Structure :	N/A N/A
Synonyms:	N/A		

History:	Initial reviews:	November 1974, October 1978, April 2001 2015 For new submissions*: Not applicable	
	Current revision date:		
	Implementation date:		
		For existing fragrance compounds*:	Not applicable
Next review date 2020		2020	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	0.15%	Rinse-off products:	No Restriction	
	Including household cleaning products			
Non skin contact products:	No Restriction			
Note box:				
The Standard is set due to the phototoxic effects of the material. The limit only applies to applications on skin exposed to sunshine, excluding rinse- off products (please refer to Table 4 of the QRA booklet for more detailed information).				
If combinations of phototoxic fragrance ingredients are used, the use levels have to be reduced accordingly. The sum of the concentrations of all phototoxic fragrance ingredients, expressed in % of their recommended maximum level in the consumer product, shall not exceed 100.				
Note: See remark on phototoxic ingredients in the Introduction to the IFRA Standards (Appendix 8 to the IFRA Code of Practice) and the Standard on Citrus oil and other furocoumarins-containing essential oils.				
Fragrance material specifications: N/A				

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

PHOTOTOXICITY

REFERENCES:

This recommendation is based on the fact that Rue oil is known to contain psoralens and on the no-effect level of 0.8% found in hairless mice (P.D. Forbes, F. Urbach, R.E. Davis (1977), Fd. Cosmet. Toxicol. 15, 55-60 and communication from RIFM).


Safrole, Isosafrole, Dihydrosafrole

CAS N°:	94-59-7 120-58-1 94-58-6	Empirical formula:	N/A
Synonyms:	N/A		

History:	Initial reviews:	October 1976, July 1987	
	Current revision date:	Not applicable	
	Implementation date:	For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED / RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	0.01%	Rinse-off products:	0.01%	
		Including household	cleaning products	
Non skin contact products:	0.01%			
Note box:				
Fragrance material specifications:		Safrole as such should not be oils containing safrole should concentration of safrole excee of essential oils with a high sa officinale Nees& Eberm.), Occ and certain qualities of Camph The total concentration of safr exceed 0.01% in consumer pr	used as a fragrance ingredient; essential not be used at a level such that the total ds 0.01% in consumer products. Examples frole content are Sassafras oil (<i>Sassafras</i> <i>otea Cymbarum</i> oil (<i>Ocotea pretiosa</i> Metz) nor oils. ole, isosafrole and dihydrosafrole should not oducts.	

CONTRIBUTION FROM OTHER SOURCES:

N/A

REFERENCES:

These recommendations are based on the conclusions of the Scientific Committee on Cosmetology of the EEC on safrole and on the similarity of the biological activity of these substances (Scientific Committee of Cosmetology of the EEC, opinion reached on September 2, 1980; Communication to the EEC Commission ENV/521/79 and IARC Monograph Vol. 10, 1976, 231-244).

Santolina oil

CAS N°:	84961-58-0	Empirical formula:	N/A
Synonyms:	N/A		

History:	Initial reviews:	May 2006 (40th Amendement)	
	Current revision date:	Not applicable	
	Implementation date:	e: For new submissions*: Not applicable	
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products:	N/A				
Note box:					
The material had been on the list of 'other materials' before, which for reasons of a unified Standard format for prohibited materials was disbanded.					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SEE NOTE BOX

REXPAN RATIONALE / CONCLUSION:

The material has been reviewed by the RIFM Expert Panel with the conclusion that it should not be used as or in fragrance ingredients until additional data is available and considered sufficient to support its safe use. This conclusion is based on:

1) presence of structural alerts as defined in the Human Health Criteria Document (Ford et al., 2000) and/or

2) adverse data on the material itself and/or

3) adverse data for a structurally related material

REFERENCES:

Human Health Criteria Document, Reg. Tox & Pharm., 31, 166-181, 2000



Savin oil

CAS N°:	8024-00-8	Empirical formula:	N/A
Synonyms:	N/A		

History:	Initial reviews:	May 1980, June 1982	
	Current revision date:	e: Not applicable	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED / SPECIFICATION

RESTRICTIONS:

Limits in the finished product:			
Skin contact products:			
Leave on products:	N/A	Rinse-off products:	N/A
		Including household	cleaning products
Non skin contact products:	N/A		
Note box:			
Fragrance material specifications:		Savin oil should not be used as a fragrance ingredient if prepared from <i>Juniperus Sabina</i> L. Only oils obtained from <i>Juniperus phoenicea</i> L. should be used.	
		In the absence of an international standard, the following specificiations for oils of <i>Juniperus phoenicea</i> L. are proposed:	
		Density d 20/20 0,864 - 0,873 Refraction n 20 D 1,4700 - 1,4720 Rotation alpha 20 D -1° - +4° Acid value 0,4 - 1 Ester value 2,5 - 7 Ester value after acetylation 10 - 23 Solubility 0.5-6 vol. in alcohol 96%, beyond that opalescence on dilution	

CONTRIBUTION FROM OTHER SOURCES:

N/A

REFERENCES:

This recommendation is based on the high acute toxicity of oils from J. sabina L. (R.E. Gosselin, H.C. Hodge, R.P. Smith & M.N. Gleason (1976), Clinical Toxicology of Commercial Products, 4th ed., Section II, p. 153, Williams & Wilkins Co., Baltimore), and the low acute toxicity of oils from J. phoenicea L. (private communication to IFRA).



CAS N°:	515-03-7	Empirical formula: Structure:	C ₂₀ H ₃₆ O ₂ OH OH OH
Synonyms:	Labd-14-ene-8,13-diol 1-Naphthalenepropanol,decahydro-alpha-6 (1R-(1-alpha(R*),2-beta,4a-beta,8a-alpha)	ethenyl-2-hydroxy- alı)-	bha,2,5,5,8apentamethyl-,

History:	Initial reviews:	February 1986	
	Current revision date:	e: September 2004	
	Implementation date:	For new submissions*: November 12, 200	
		For existing fragrance compounds*:	November 12, 2006
	Next review date	2009	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

SPECIFICATION

RESTRICTIONS:

Limits in the finished product:					
Skin contact products:					
Leave on products:	N/A	Rinse-off products:	N/A		
		Including household	cleaning products		
Non skin contact products:	N/A				
Note box:					
Fragrance material specifications:Sclareol used as a fragrance ingredient should have a minimum purity of 98%.			rance ingredient should have a		

CONTRIBUTION FROM OTHER SOURCES:

Contributions from other sources are known but not of relevance for this type of Standard.

Sclareol

REXPAN RATIONALE / CONCLUSION:

This recommendation is based on test results of RIFM with different qualities showing a sensitizing potential of samples with a lower purity and no sensitizing reactions for samples with a minimum purity of 98%.

REFERENCES:

Research Institute for Fragrance Materials, Inc. (1975a). Repeated Insult Patch Test with Sclareol. RIFM report number 45024, June 17. (RIFM, Woodcliff Lake, NJ, USA).

Research Institute for Fragrance Materials, Inc. (1975b). Repeated Insult Patch Test with Sclareol. RIFM report number 45025, June 18. (RIFM, Woodcliff Lake, NJ, USA).

Research Institute for Fragrance Materials, Inc. (1979a). Report on Human Maximization Studies. RIFM report number 1697, April 20. (RIFM, Woodcliff Lake, NJ, USA).

Research Institute for Fragrance Materials, Inc. (1979b). Report on Human Maximization Studies. RIFM report number 1697, November 6. (RIFM, Woodcliff Lake, NJ, USA).

Research Institute for Fragrance Materials, Inc. (1981). Report on Human Maximization Studies. RIFM report number 1792, March 18. (RIFM, Woodcliff Lake, NJ, USA).

Research Institute for Fragrance Materials, Inc. (1986). Report on Human Maximization Studies. RIFM report number 3100, January 15. (RIFM, Woodcliff Lake, NJ, USA).

Styrax

CAS N°:	8046-19-3 8024-01-9 94891-27-7 94891-28-8	Empirical formula:	N/A
Synonyms:	For the crude materials banned: Styrax crude gums For the distillates specified: Stryax resin Styrax oil Styrax oil, rectified		

History:	Initial reviews:	November 1977, July 1994		
	Current revision date:	June 2013		
	Implementation date:	For new submissions*: January 10, 2014		
		For existing fragrance compounds*:	January 10, 2015	
	Next review date	2018		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED / RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.04 %	Category 7	0.11 %			
Category 2	0.05 %	Category 8	0.60 %			
Category 3	0.23 %	Category 9	0.60 %			
Category 4	0.60 %	Category 10	0.60 %			
Category 5	0.36 %	Category 11	See Note box (2)			
Category 6	0.60 %					

Note box:

(1) See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products

(1) doe in a local of ingestion.
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:	Crude gums of Liquidambar styraficula L. var. macrophylla or Liquidambar orientalis Mill. should not be used as fragrance ingredients: Only extracts or distillates (resinoids, absolutes and oils), prepared from exudations of Liquidambar styraciflua L. var. macrophylla or Liquidambar orientalis Mill., can be used. This recommendation is made in order to promote good manufacturing practice (GMP) for the use of styrax derivatives as fragrance ingredients. It is based on a wide variety of RIFM test data with gums, resinoids, absolutes and oils of American and Asian styrax (private communication to IFRA). In addition, Styrax oil can be obtained from solvent extraction or pyrolysis. Styrax oil obtained through pyrolysis shall be rectified according to Good Manufacturing Practices and the content of polynuclear aromatic
	Manufacturing Practices and the content of polynuclear aromatic hydrocarbons (PAH) resulting from their use shall respect the following requirement:

Styrax

Benzopyrene and 1,2-Benzanthracene are to be used as markers for PAH. If used alone or in combination with rectified Cade oil, rectified Birch tar oils or rectified Opoponax oil, the total concentration of both of the markers should not exceed 1 ppb in the final product.

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
> 1525 [1] ^{4,5}	Moderate	1500 ⁵	NA	NA	1500

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴EC3 value from one LLNA , not the mean.

⁵Study conducted on a well-characterized sample of Styrax oil – pyrogenated. The same sample was evaluated in both the LLNA and

HRIPT reported within the Standard.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Styrax (all forms) and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as $1500 \ \mu g/cm^2$. They recommend the limits for the 11 different product categories, which are the acceptable use levels of Styrax (all forms) in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick GF, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. Local Lymph Node Assay on Styrax Oil - Pyrogenated. Draft RIFM Report number 64109. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. Human Repeated Insult Patch Test on Styrax Oil - Pyrogenated. Draft RIFM Report number 64110. (RIFM, Woodcliff Lake, NJ, USA



Tagetes oil and absolute

CAS N°:	91722-29-1	Empirical formula:	N/A
	8016-84-0	Structure:	N/A
Synonyms:	Tagetes absolute (Tagetes patula L.) Tagetes patula absolute Tagetes patula, ext. Tagetes minuta absolute Tagetes oil		

History:	Initial reviews:	October 1986, April 2001, January 2001			
	Current revision date:	2015			
	Implementation date:	For new submissions*:	Not applicable		
		For existing fragrance compounds*:	Not applicable		
	Next review date	2020			

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:							
Skin contact products:							
Leave-on products :	0.01%	Rinse- Includi	off products: ng household cleaning products	No restriction			
Non-skin contact products:	No restriction						
Note box:							
The Standard is set due to the phototoxic effects of the material. The limit only applies to applications on skin exposed to sunshine, excluding rinse- off products (please refer to Table 4 of the QRA booklet for more detailed information).							
Fragrance material specifications: N/A							

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

PHOTOTOXICITY

Tagetes oil and absolute

RIFM SUMMARIES:

Tagetes oils and absolutes obtained from Tagetes minuta L. (syn. Tagetes glandulifera Schrank and Tagetes patula L.) were evaluated by RIFM (Letizia and Api, 2000). A no effect level for phototoxicity of 0.05% was determined on humans using Egyptian Tagete minuta (RIFM, 1986a).

- At 0.003% in guinea pigs, no observable effects, 0/10 (RIFM, 1985a).
- At 0.01% in guinea pigs, phototoxicity observed, 8/10 (RIFM, 1985b).
- At 100% in mice, phototoxicity was observed, 6/6 (RIFM,1986b).
- At 1% in mice, phototoxicity was observed, 6/6 (RIFM, 1986c).
- At 0.1% in mice, phototoxicity was observed, 6/6 (RIFM, 1986c).
- At 0.01% in mice, phototoxicity was observed, 2/6 (RIFM, 1986c).

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Tagetes oil and absolute and recommended no change to the Standard (September 2001).

REFERENCES:

Letizia C.S. and Api A.M (2000). A dermal safety evaluation of extracts from Tagetes plants used in fragrances. The Toxicologist, 54(1), 397.

Research Institute for Fragrance Materials, Inc. (1985a). Guinea Pig Phototoxicity Test. Unpublished report from Givaudan. Report number 3361, 17 December.

Research Institute for Fragrance Materials, Inc. (1985b). Guinea Pig Phototoxicity Test. Unpublished report from Givaudan. Report number 3362, 17 December.

Research Institute for Fragrance Materials, Inc. (1986a). Human Photosensitization Test. RIFM report number 1690, 21 November.

Research Institute for Fragrance Materials, Inc. (1986b). Mouse Phototoxicity Test. RIFM report number 3828, 25 June.

Research Institute for Fragrance Materials, Inc. (1986c). Mouse Phototoxicity Test. RIFM report number 4343, 31 July.

Tea leaf absolute

CAS N°:	84650-60-2	Empirical formula:	N/A
Synonyms:	Camellia sinensis leaf extract Tea, ext. Tea sinensis absolute Thea chinensis ext. Thea sinensis ext.		

History:	Initial reviews:		
	Current revision date:	2006	
	Implementation date:	For new submissions*:	June 11, 2007
		For existing fragrance compounds*:	June 11, 2008
	Next review date	2011	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:						
Category 1 See Note box (1)	0.01 %	Category 7	0.04 %			
Category 2	0.02 %	Category 8	0.5 %			
Category 3	0.07 %	Category 9	2.4 %			
Category 4	0.2 %	Category 10	2.5 %			
Category 5	0.1 %	Category 11	See Note Box (2)			
Category 6	0.3 %					
Note box:						
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) (http://www.iofiorg.org/).						
(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.						
Fragrance material specifications: N/A						

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SENSITIZATION

IFRA Standard – Tea leaf absolute

Tea leaf absolute

RIFM SUMMARIES:

Tea Leaf Absolute - Sensitization Potency Estimation Based on Weight of Evidence

		Human Data			
LLNA weighted mean EC3 values (µg/cm ²) [no. studies] Potency Classification Based on Animal Data ²		NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ¹ (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
>1250[1]4	Moderate	480	NA	NA	480

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed

effect level; NA = Not Available.

¹ Data derived from HRIPT or HMT.

²Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

³WoE NESIL limited to two significant figures.

⁴ Irritation was observed at higher concentrations; EC3 value not calculable

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for tea leaf absolute and based on the weight of evidence established the No Expected Sensitization Induction Level (NESIL) as 480 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of tea leaf absolute in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of March 15, 2006.

REFERENCES:

Gerberick, GF. et. al. (2001) Contact allergenic potency: Correlation of human and local lymph node assay data. American Journal of Contact Dermatitis, 12(3), 156-161.

QRA Expert Group* (AM Api, DA Basketter, PA Cadby, M-F Cano, G Ellis, GF Gerberick, P Griem, PM McNamee, CA Ryan and R Safford), Dermal Sensitization Quantitative Risk Assessment (QRA) for Fragrance Ingredients, Technical Dossier, March 15, 2006, http://www.rifm.org/pub/publications.asp.

Research Institute for Fragrance Materials, Inc (1990). Delayed contact hypersensitivity study of tea leaf absolute in guinea pigs. RIFM report number 12409 (RIFM, Woodcliff Lake, NJ USA).

Research Institute for Fragrance Materials, Inc (2004). Repeated insult patch test of tea leaf absolute in human subjects. Unpublished report from Robertet Incorporated, Report number 44878 (RIFM, Woodcliff Lake, NJ USA).

Research Institute for Fragrance Materials, Inc (2005). Local Lymph Node Assay on tea leaf absolute. Unpublished report from Robertet Incorporated, Report number 47597 (RIFM, Woodcliff Lake, NJ USA).

1,2,3,4-Tetrahydro-4-methylquinoline

CAS N°:	19343-78-3	Empirical formula: Structure:	C10H13N
Synonyms:	4-Methyl-1,2,3,4-tetrahydroquinolir Quinoline, 1,2,3,4-tetrahydro-4-me 1,2,3,4-Tetrahydrolepidine 1,2,3,4-Tetrahydro-4-methylquinole	ne thyl- eine	

History:	Initial reviews:	New Standard		
	Current revision date:	2009		
	Implementation date:	For new submissions*:	August 7, 2009	
		For existing fragrance compounds*:	August 7, 2010	
	Next review date	2014		
* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.				

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
The material has been identified for having the potential of forming nitrosamines in nitrosating systems. Downstream users therefore have to be notified of the presence of the material and its potential to be able to consider adequate protective measures.						
Fragrance material specificatio	ns:	N/A				

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)

CRITICAL EFFECT:

POTENTIAL OF NITROSAMINE FORMATION

REXPAN RATIONALE / CONCLUSION:

IFRA measures regarding potential nitrosamine formation noted - REXPAN April 2009.

REFERENCES:

Nitrosamine policy as contained in the EU Cosmetics Directive 76/768/EEC and its Amendments.

o,m,p-Tolualdehydes and their mixtures

CAS N°:	529-20-4 620-23-5 104-87-0 1334-78-7	Empirical formula: Structure:	C_8H_8O 0 R1 R2 R3 With $R_1=Me, R_2=H, R_3=H$ or $R_1=H, R_2=Me, R_3=H$ or $R_1=H, R_2=H, R_3=Me$
Synonyms:	ortho-Tolualdehyde 2-Methyl-benzaldehyde		
	meta-Tolualdehyde 3-Methyl-benzaldehyde		
	para-Tolualdehyde 4-Methyl-benzaldehyde		
	Tolualdehydes (mixed o,m,p) o,m,p-Methyl-benzaldehydes		

History:	Initial reviews:	New Standard		
	Current revision date:	June 2013		
	Implementation date:	For new submissions*: August 10, 2013		
		For existing fragrance compounds*:	August 10, 2014	
	Next review date	2018		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

o,m,p-Tolualdehydes and their mixtures

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.03 %	Category 7	0.08 %		
Category 2	0.04 %	Category 8	1.11 %		
Category 3	0.17 %	Category 9	5.00 %		
Category 4	0.50 %	Category 10	2.50 %		
Category 5	0.26 %	Category 11	See Note box (2)		
Category 6	0.80 %				
NL-1-L-					

Note box:

(1) See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products with the potential of ingestion.

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

None to consider.

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

Cuminaldehyde has been designated as a read across material for the Tolualdehydes for the sensitization endpoint. As such, the data for Cuminaldehyde are below:

		Human Data			
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (μg/cm ²)
> 2500 [1] ⁴	Weak	1181	2760	NA	1100 ⁵

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴EC3 value from one LLNA, not the mean.



o,m,p-Tolualdehydes and their mixtures

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Cuminaldehyde and the Tolualdehydes and, based on the weight of evidence and read-across, established the No Expected Sensitization Induction Level (NESIL) as 1100 μ g/cm² based on the critical sensitization data on Cuminaldehyde. They recommend the limits for the 11 different product categories, which are the acceptable use levels of Tolualdehydes in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick G, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. Regulatory Toxicology and Pharmacology 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 1972. Maximization test. RIFM report number 1804, November 22. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1975. Maximization test. RIFM report number 1804, March 27a. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1973. Maximization test. RIFM report number 1804, November 11. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. Repeat Insult Patch Test. Draft RIFM Report number 63810. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.) 2012. Local Lymph Node Assay. Draft RIFM Report number 63814. (RIFM, Woodcliff Lake, NJ, USA).



CAS N°:	108-88-3	Empirical formula: Structure:	С7Н8
Synonyms:	Toluol Methylbenzol Methylbenzene		

History:	Initial reviews:	Not applicable	
	Current revision date:	October 2003	
	Implementation date:	For new submissions*:	May 6, 2004
		For existing fragrance compounds*:	May 6, 2005
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

DECOM	
REGUNN	

PROHIBITED

RESTRICTIONS:

Limits in the finished product:						
Skin contact products:						
Leave on products:	N/A	Rinse-off products:	N/A			
		Including household	cleaning products			
Non skin contact products:	N/A					
Note box:						
The material should not be used as f	The material should not be used as fragrance ingredient for any application.					
Fragrance material specificatio	ns:	The level of toluene has and should never excee compound.	s to be kept as low as practicable ed 100 ppm in the fragrance			

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

LIVER TOXICITY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted - REXPAN, October 15, 2003

REFERENCES:

1) Cosmetic Ingredient Review, Journal of the American College of Toxicology JACT 6 (1) 1987.

2) IARC (International Agency for Research on Cancer) Monographs Vol 47, p .79 (1989); Vol 71 p. 829 (1999)

Treemoss extracts

CAS N°:	90028-67-4 = Evernia furfuracea 68648-41-9 = oils, Treemoss 68917-40-8 = oils, Treemoss resinoid	Empirical formula:	N/A
Synonyms:	Treemoss absolute (Pseudevernia fur Treemoss (Usnea furfuracea) Treemoss Colourless Pseudevernia furfuracea extract Cedar moss	furacea)	

History:	Initial reviews:	1991, 2001		
	Current revision date:	2008		
	Implementation date:	For new submissions*:	February 16, 2009	
		For existing fragrance compounds*:	February 16, 2011	
	Next review date	2013		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED / SPECIFICATION

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.02 %	Category 7	0.1 %		
Category 2	0.03 %	Category 8	0.1 %		
Category 3	0.1 %	Category 9	0.1 %		
Category 4	0.1 %	Category 10	0.1 %		
Category 5	0.1 %	Category 11	See Note Box (2)		
Category 6	0.5 %				

Note box:

For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again.

In the presence of oak moss extracts, the level of tree moss in the respective category has to be reduced accordingly such that the total amount of both extracts does not exceed the maximum permitted level in each category as listed in the table above.

If the same compound is intended for more than one IFRA QRA category, then the most restrictive limitation (based on foreseen use concentrations and maximum permitted level) will apply.

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/.

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Treemoss extracts

Fragrance material specifications:

Tree moss extracts shall not contain more than 0.8% of dehydroabietic acid (DHA) as a marker of 2% of total resin acids. The concentration of DHA (about 40% of the total resin acids) in tree moss can be measured with an HPLC reverse Phase - spectrofluorometry method.

Further, levels of atranol and chloroatranol should each be below $100\ ppm$ in tree moss extracts.

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards)

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
> 5000	Moderate	700 ⁴	6896 ⁴	1417	700

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

¹ Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

² Data derived from HRIPT or HMT

³WoE NESIL limited to three significant figures

⁴MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Treemoss extracts and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 700 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Treemoss extracts in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1974. Human Maximization Test. RIFM report number 1779, September 12 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1990a. Human Repeated Insult Patch Test. RIFM report number 12382, March 1 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1990b. Human Repeated Insult Patch Test. RIFM report number 14120, November 26 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1990c. Human Repeated Insult Patch Test. RIFM report number 14118, November 26 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2004. Local Lymph Node Assay. RIFM report number 44368, March 25 (RIFM, Woodcliff Lake, NJ, USA).

ITra/...

2,6,6-Trimethylcyclohex-1,3-dienyl methanal

CAS N°:	116-26-7	Empirical formula: Structure:	
Synonyms:	2,6,6-Trimethylcyclohexa-1,3-diene-1-c 2,6,6-Trimethyl-1,3-cyclohexadienal 2,6,6-Trimethyl-1,3-cyclohexadien-1-ca 1,1,3-Trimethyl-2-formylcyclohexa-2,4- Dehydro-β-cyclocitral Safranal	carbaldehyde arboxaldehyde diene	

History:	Initial reviews:	December 1998	
	Current revision date:	June 2013	
	Implementation date:	For new submissions*:	August 10, 2013
		For existing fragrance compounds*:	August 10, 2014
	Next review date	2018	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.001 %	Category 7	0.002 %		
Category 2	0.001 %	Category 8	0.005 %		
Category 3	0.004 %	Category 9	0.005 %		
Category 4	0.005 %	Category 10	0.005 %		
Category 5	0.005 %	Category 11	See Note box (2)		
Category 6	0.02 %				
Note box:					
(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry - www.iofforg.org)					

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

CONTRIBUTION FROM OTHER SOURCES:

See Annex I

2,6,6-Trimethylcyclohex-1,3-dienyl methanal

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

		Human Data			
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
<250 [1]	Strong	29.5	NA	39	29

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for 2,6,6-Trimethylcyclohexa-1,3-dienyl methanal and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 29 μ g/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of 2,6,6-Trimethylcyclohexa-1,3-dienyl methanal in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick G, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. Regulatory Toxicology and Pharmacology 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 2012. Repeat Insult Patch Test. Draft RIFM Report number 63809. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.) 2012. Local Lymph Node Assay. Draft RIFM Report number 63813. (RIFM, Woodcliff Lake, NJ, USA).

Verbena absolute (Lippia citriodora Kunth.)

CAS N°:	8024-12-2 85116-63-8	Empirical formula: Strucutre:	N/A
Synonyms:	Lippia citriodora absolute Verbena absolute Aloysia triphylla absolute Lippia triphylla absolute Verbena triphylla absolute Zappania citrodora absolute		

History:	Initial reviews:	November 1987		
	Current revision date:	June 11, 2010		
	Implementation date:	For new submissions*:	August 11, 2010	
		For existing fragrance compounds*:	August 11, 2011	
	Next review date	March 2015		

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:					
Category 1 See Note box (1)	0.05%	Category 7	0.12%		
Category 2	0.06%	Category 8	0.2%		
Category 3	0.2%	Category 9	0.2%		
Category 4	0.2%	Category 10	0.2%		
Category 5	0.2%	Category 11	See Note Box (2)		
Category 6	1.2%				
Note box:					
For this material, for pragmatic reasons, restrictive levels allowed by the QRA for certain categories but actually being higher than those already in place before applying the QRA, will temporarily not be implemented until the end of a 5 year monitoring phase. At the end of the 5 years the position will be reevaluated again. (1) See the IFRA Code of Practice (Appendix 8, Introduction to the IFRA Standards) regarding the Note on Oral Care Products and other products with the potential of ingestion. (2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.					
Fragrance material specifications: N/A					

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SENSITIZATION

Verbena absolute (Lippia citriodora Kunth.)

RIFM SUMMARIES:

		Human Data				
LLNA weighted mean EC3 values (μg/cm ²) [no. studies]	Potency Classification Based on Animal Data ¹	NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)	
4500 [1] ⁴	Weak	1600	1380	8280 ⁵	1600	

All data in this Table are available from RIFM and are listed in the RIFM Database.

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available.

¹Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003.

²Data derived from HRIPT or HMT.

³WoE NESIL limited to two significant figures.

⁴EC3 value from one LLNA, not the mean.

⁵LOEL from human maximization test, not a human repeated insult patch test.

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for verbena absolute and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 1600 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of verbena absolute in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the publication by Api *et al.*, 2008.

REFERENCES:

Api AM, Basketter DA, Cadby PA, Cano M-F, Ellis G, Gerberick GF, et al. Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients. *Regulatory Toxicology and Pharmacology* 2008;52(1): 3-23.

RIFM (Research Institute for Fragrance Materials, Inc.), 1979. Maximization study with verbena absolute. RIFM report number 1697, October 24a. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2009. Cutaneous tolerance and sensitizing potential of verbena. Unpublished study from Robertet, September 23. Report number 58178. (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2009. Local Lymph Node Assay. Unpublished study from Robertet, January 23. Report number 54268. (RIFM, Woodcliff Lake, NJ, USA).

Verbena oil

CAS N°:	8024-12-2	Empirical formula:	N/A
Synonyms:	Lippia citriodora oils		

History:	Initial reviews:	December 1981	
	Current revision date:	September 2002	
	Implementation date:	For new submissions*:	Not applicable
		For existing fragrance compounds*:	Not applicable
	Next review date	Not applicable	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

PROHIBITED

RESTRICTIONS:

Limits in the finished product:				
Skin contact products:				
Leave on products:	N/A	Rinse-off products:	N/A	
		Including household cleaning products		
Non skin contact products:	N/A			
Note box:				
Verbena oil from <i>Lippia citriodora</i> Kunth should not be used as a fragrance ingredient, based on its sensitizing and phototoxic potential. Commercial compositions of the verbena type should meet the requirements of the IFRA Standards.				
Fragrance material specifications:		N/A		

CONTRIBUTION FROM OTHER SOURCES:

None to consider (see also the note on contributions from other sources in the Introduction to the IFRA Standards).

CRITICAL EFFECT:

SENSITIZATION, PHOTOXICITY

REXPAN RATIONALE / CONCLUSION:

IFRA ban noted – REXPAN, September 2002.

REFERENCES:

Private communication to IFRA.

Ylang Ylang extracts

CAS N°:	8006-81-3 68606-83-7 83863-30-3	Empirical formula:	N/A
Synonyms:	Cananga odorata (Lamark) (Hooker Cananga odorata extract Cananga odorata flower oil Cananga odorata oil Cananga oil Ylang ylang oil (Cananga odorata H Ylang ylang oil extra Ylang ylang oil extra Ylang ylang oil II Ylang ylang oil II Ylang ylang oil III Ylang ylang, Cananga odorata, ext.	et Thompson) (Anonaceae) ook. f. and Thomas)	

History:	Initial reviews:	New Standard	
	Current revision date:	2008	
	Implementation date:	For new submissions*:	August 16, 2008
	-	For existing fragrance compounds*:	August 16, 2010
	Next review date	2013	

* This date applies to the supply of fragrance compounds (formulas) only, not to the finished products in the marketplace.

RECOMMENDATION:

RESTRICTED

RESTRICTIONS:

Limits in the finished product:				
Category 1 See Note box (1)	0.05 %	Category 7	0.1 %	
Category 2	0.06 %	Category 8	1.8 %	
Category 3	0.27 %	Category 9	5.0 %	
Category 4	0.8 %	Category 10	2.5 %	
Category 5	0.4 %	Category 11	See Note Box (2)	
Category 6	1.3 %			
Note box:				

(1) IFRA would recommend that any material used to impart perfume or flavour in products intended for human ingestion should consist of ingredients that are in compliance with appropriate regulations for foods and food flavourings in the countries of planned distribution and, where these are lacking, with the recommendations laid down in the Code of Practice of IOFI (International Organisation of the Flavor Industry) http://www.iofiorg.org/.

(2) Category 11 includes all non-skin contact or incidental skin contact products. Due to the negligible skin contact from these types of products there is no justification for a restriction of the concentration of this fragrance ingredient in the finished product.

Fragrance material specifications:

N/A

Ylang Ylang extracts

CONTRIBUTION FROM OTHER SOURCES:

N/A

CRITICAL EFFECT:

SENSITIZATION

RIFM SUMMARIES:

	Potency Classification Based on Animal Data ¹	Human Data			
LLNA weighted mean EC3 values (μg/cm²) [no. studies]		NOEL – HRIPT (induction) (µg/cm ²)	NOEL – HMT (induction) (µg/cm ²)	LOEL ² (induction) (µg/cm ²)	WoE NESIL ³ (µg/cm ²)
1700	Moderate	1772 ⁴	6897 ⁴	7752	1770

NOEL = No observed effect level; HRIPT = Human Repeat Insult Patch Test; HMT = Human Maximization Test; LOEL = lowest observed effect level; NA = Not Available

1 Based on animal data using classification defined in ECETOC, Technical Report No. 87, 2003

2 Data derived from HRIPT or HMT

3 WoE NESIL limited to three significant figures

4 MT-NOEL = Maximum Tested No Effect Level. No sensitization was observed in human predictive studies. Doses reported reflect the highest concentration tested, not necessarily the highest achievable NOEL

REXPAN RATIONALE / CONCLUSION:

The RIFM Expert Panel reviewed the critical effect data for Ylang Ylang Extracts and, based on the weight of evidence, established the No Expected Sensitization Induction Level (NESIL) as 230 µg/cm². They recommend the limits for the 11 different product categories, which are the acceptable use levels of Ylang Ylang Extracts in the various product categories. These were derived from the application of the exposure-based quantitative risk assessment approach for fragrance ingredients, which is detailed in the QRA Expert Group Technical Dossier of June 22, 2006.

REFERENCES:

RIFM (Research Institute for Fragrance Materials, Inc.), 1971a. Human Maximization Test. RIFM report number 1805, April 20 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1971b. Human Repeated Insult Patch Test. RIFM report number 7906, April 13 and August 23 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1972a. Human Maximization Test. RIFM report number 1804, February 18 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1972b. Human Repeated Insult Patch Test. RIFM report number 14033, March 21 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1975. Human Maximization Test. RIFM report number 1798, March 28 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 1979. Human Maximization Test. RIFM report number 1697, July 6a (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007a. Human Repeated Insult

Patch Test. RIFM report number 52898, May 15 (RIFM, Woodcliff Lake, NJ, USA).

RIFM (Research Institute for Fragrance Materials, Inc.), 2007b. Local Lymph Node Assay. RIFM report number 52903, April 24 (RIFM, Woodcliff Lake, NJ, USA).



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