



Scientific Committee on Consumer Safety

SCCS

OPINION
on
Fragrance allergens in cosmetic products

The SCCS adopted this opinion at its 15th plenary meeting

of 26-27 June 2012

About the Scientific Committees

Three independent non-food Scientific Committees provide the Commission with the scientific advice it needs when preparing policy and proposals relating to consumer safety, public health and the environment. The Committees also draw the Commission's attention to the new or emerging problems which may pose an actual or potential threat.

They are: the Scientific Committee on Consumer Safety (SCCS), the Scientific Committee on Health and Environmental Risks (SCHER) and the Scientific Committee on Emerging and Newly Identified Health Risks (SCENIHR) and are made up of external experts.

In addition, the Commission relies upon the work of the European Food Safety Authority (EFSA), the European Medicines Agency (EMA), the European Centre for Disease prevention and Control (ECDC) and the European Chemicals Agency (ECHA).

SCCS

The Committee shall provide opinions on questions concerning all types of health and safety risks (notably chemical, biological, mechanical and other physical risks) of non-food consumer products (for example: cosmetic products and their ingredients, toys, textiles, clothing, personal care and household products such as detergents, etc.) and services (for example: tattooing, artificial sun tanning, etc.).

Scientific Committee members

Jürgen Angerer, Ulrike Bernauer, Claire Chambers, Qasim Chaudhry, Gisela Degen, Elsa Nielsen, Thomas Platzek, Suresh Chandra Rastogi, Vera Rogiers, Christophe Rousselle, Tore Sanner, Jan van Benthem, Jacqueline van Engelen, Maria Pilar Vinardell, Rosemary Waring, Ian R. White

Contact

European Commission
Health & Consumers
Directorate D: Health Systems and Products
Unit D5 - Risk Assessment
Office: B232 B-1049 Brussels
Sanco-SCCS-Secretariat@ec.europa.eu

© European Union, 2011

ISSN 1831-4767

Doi:10.2772/77628

ISBN 978-92-79-30752-2

ND-AQ-12-002-EN-N

The opinions of the Scientific Committees present the views of the independent scientists who are members of the committees. They do not necessarily reflect the views of the European Commission. The opinions are published by the European Commission in their original language only.

http://ec.europa.eu/health/scientific_committees/consumer_safety/index_en.htm

Acknowledgements

Dr. C. Chambers
Dr. Q. Chaudry
Dr. S.C. Rastogi
Dr. I.R. White (chairman)

External experts

| | |
|----------------------|--|
| Prof.. A. Börje | University of Gothenburg, Sweden |
| Prof. J. D. Johansen | Gentofte Hospital, University of Copenhagen, Denmark |
| Prof. A-T. Karlberg | University of Gothenburg, Sweden |
| Prof. C. Lidén | Karolinska Institutet, Sweden |
| Dr. D.W. Roberts | Liverpool John Moores University, UK |
| Prof. W. Uter | (rapporteur) Friedrich-Alexander University (FAU), Erlangen, Germany |

Keywords: SCCS, scientific opinion, labelling, fragrance allergens, directive 76/768/ECC

Opinion to be cited as: SCCS (Scientific Committee on Consumer Safety), opinion on fragrance allergens in cosmetic products, 26-27 June 2012

Table of contents

| | |
|--|----|
| Acknowledgements..... | 3 |
| Table of contents | 4 |
| Summary..... | 7 |
| 1. Background | 9 |
| 2. Terms of reference..... | 10 |
| 3. Introduction..... | 11 |
| 4. Clinical aspects of contact allergy to fragrance ingredients..... | 12 |
| 4.1. Spectrum of reactions..... | 12 |
| 4.1.1. Allergic contact dermatitis | 12 |
| 4.1.2. Irritant reactions (including contact urticaria) | 14 |
| 4.1.3. Pigmentary anomalies..... | 14 |
| 4.1.4. Photo-reactions..... | 14 |
| 4.1.5. General/respiratory | 14 |
| 4.2. Patch testing | 15 |
| 4.3. Epidemiology of fragrance allergy | 15 |
| 4.3.1. Substances used for screening of contact allergy to fragrance ingredients | 15 |
| 4.3.2. Clinical epidemiology | 16 |
| 4.3.3. Population-based epidemiology | 23 |
| 4.4. Consumer products as a cause of fragrance contact sensitisation and allergic contact dermatitis..... | 25 |
| 4.4.1. Clinical relevance | 25 |
| 4.4.2. Elicitation with clinical symptoms/signs, current and past..... | 26 |
| 4.4.3. Elicitation in diagnostic patch tests without clinical history..... | 28 |
| 4.5. Socio-economic impact of contact allergy..... | 29 |
| 4.5.1. Health related quality of life..... | 29 |
| 4.5.2. Occupational restrictions | 29 |
| 4.5.3. Costs to health care/health economics | 29 |
| 4.6. Allergen avoidance | 30 |
| 4.6.1. Primary prevention: limiting or eliminating exposure to allergens in the population | 30 |
| 4.6.2. Secondary prevention: avoiding re-exposure to (a) specific sensitiser(s) in clinically diagnosed individuals..... | 30 |
| 4.7. Conclusions..... | 32 |
| 5. Activation of weak or non-sensitising substances into sensitisers - prohaptens and prohaptens..... | 33 |
| 5.1. Prehaptens..... | 33 |
| 5.2. Prohaptens..... | 37 |
| 5.3. Conclusions | 39 |
| 6. Retrieval of evidence and classification of fragrance substances..... | 40 |
| 6.1. Retrieval of evidence | 40 |

| | | |
|---------|--|-----|
| 6.1.1. | Search strategy for clinical data | 40 |
| 6.1.2. | Collection of experimental (LLNA) data | 41 |
| 6.2. | Grading of evidence | 41 |
| 6.2.1. | Quality of a clinical study | 41 |
| 6.2.2. | Quality of an experimental study | 42 |
| 6.2.3. | Quality of "other" evidence | 42 |
| 6.3. | Aggregating evidence for a final conclusion | 42 |
| 6.3.1. | Established contact allergen in humans | 42 |
| 6.3.2. | Established contact allergen in animals | 43 |
| 6.3.3. | Likely contact allergen, if human, animal and other evidence is considered ... | 43 |
| 6.3.4. | Possible contact allergen, if human, animal and other evidence is considered | 43 |
| 6.4. | Conclusions | 44 |
| 7. | Reported fragrance allergens from the clinical perspective | 45 |
| 7.1. | Tabular summary of evaluated individual fragrance chemicals | 45 |
| 7.2. | Tabular summary of evaluated natural extracts/essential oils | 53 |
| 7.3. | Conclusions | 57 |
| 8. | Animal data | 58 |
| 8.1. | Predictive tests and sensitising potency categories | 58 |
| 8.1.1. | LLNA data | 59 |
| 8.1.2. | LLNA data on oxidised fragrance substances | 61 |
| 8.2. | Methodological considerations | 62 |
| 8.3. | Summary of animal data by LLNA | 63 |
| 8.4. | Conclusions | 64 |
| 9. | Structure activity relationships (SAR): grouping of substances based on expert judgement | 66 |
| 9.1. | General results | 71 |
| 9.2. | Conclusions | 71 |
| 10. | Exposure | 72 |
| 10.1. | Concentrations and quantities used | 72 |
| 10.2. | Global exposure (household and occupational exposures) | 81 |
| 10.3. | Exposures related to particular anatomical sites | 84 |
| 10.4. | Conclusion | 86 |
| 11. | Dose-response relationships and thresholds | 87 |
| 11.1. | Induction | 87 |
| 11.2. | Elicitation | 88 |
| 11.2.1. | General considerations | 88 |
| 11.2.2. | Studies on specific fragrance ingredients | 90 |
| 11.3. | Hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC) | 98 |
| 11.4. | Conclusion | 101 |
| 12. | Data gaps and research needed | 103 |
| 12.1. | Clinical and epidemiological research | 103 |

| | | |
|---|---|-----|
| 12.2. | Non-human studies | 104 |
| 13. | Opinion..... | 105 |
| 13.1. | Question 1 | 106 |
| | Conclusions - Question 1 | 114 |
| 13.2. | Question 2 | 115 |
| | Conclusions - Question 2 | 116 |
| 13.3. | Question 3 | 117 |
| | Conclusions - Question 3 | 119 |
| 14. | List of abbreviations | 121 |
| 15. | References | 123 |
| Annex I - Catalogue of fragrance allergens..... | | 141 |
| | Single chemicals | 142 |
| | Catalogue of single chemicals evaluated..... | 146 |
| | Natural extracts / essential oils | 237 |
| | Catalogue of natural extracts / essential oils evaluated..... | 238 |
| | References..... | 277 |
| Annex II - Animal Data | | 293 |
| | References..... | 309 |
| Annex III - Tabular summary of dose-elicitation studies in sensitised patients..... | | 315 |
| | Chloroatranol | 316 |
| | Cinnamal | 318 |
| | Hydroxycitronellal | 321 |
| | Hydroxyisohexyl 3-cyclohexenecarboxaldehyde (HICC) | 323 |
| | Isoeugenol | 329 |
| | References..... | 333 |

Summary

Contact allergy to fragrance ingredients may develop following skin contact with a sufficient amount of these substances, often through the use of cosmetic products. Contact allergy is an altered specific reactivity in the immune system, which entails recognition of the fragrance allergen(s) in question by immune cells. Contact allergy, which *per se* is a latent condition, i.e. without visible signs or symptoms, persists lifelong. Upon each re-exposure to sufficient amounts of the allergen(s) eczema develops (allergic contact dermatitis), which typically will involve the face, the armpits and/or the hand(s). The disease can be severe and generalised, with a significant impairment of quality of life and potential consequences for fitness for work.

Around 16% of eczema patients in the European population are sensitised to fragrance ingredients. From studies performed on sectors of the population it can be estimated that the frequency of contact allergy to fragrance ingredients in the general population in Europe is 1-3%. The overall trend of fragrance allergy has been stable during the last 10 years, as some causes of fragrance allergy have decreased and others increased.

Most individuals with contact allergy to fragrance ingredients are aware that they cannot tolerate scented products on their skin and are often able to specifically name product categories that initiated their disease. In this context colognes, eau de toilette, deodorants and lotions are named significantly more often by fragrance allergic eczema patients than by patients without fragrance contact allergy.

Commercially available fragrances and other scented cosmetic products can provoke allergic contact dermatitis under patch test as well as simulated use conditions.

Appropriate diagnostic procedures and patient information are cornerstones in secondary prevention of contact allergy. The SCCNFP identified in 1999 a set of 26 fragrance allergens with a well-recognised potential to cause allergy, for which information should be provided to consumers about their presence in cosmetic products.

This listing has shown to be important in the clinical management of patients who are allergic to one or more of these 26 fragrance chemicals. Listing of the 26 fragrances has also been shown to be beneficial for patients with contact allergy to one or more of the fragrance chemicals, because these are identified on the ingredient listings of cosmetic products, and can thus be avoided.

The present opinion updates the SCCNFP opinion with a systematic and critical review of the scientific literature to identify fragrance allergens, including natural extracts, relevant to consumers. Clinical, epidemiological and experimental studies were evaluated, as well as modelling studies performed, to establish lists of (i) established fragrance allergens, (ii) likely fragrance allergens and (iii) possible fragrance allergens.

The studies since the SCCNFP Opinion on fragrance allergy in consumers confirm that the fragrance allergens identified by SCCNFP in 1999 are still relevant fragrance allergens for consumers from their exposure to cosmetic products. The review of the clinical and experimental data published since then shows that many more fragrance substances have been shown to be sensitisers in humans. Based on the clinical experience alone, 82 substances can be classified as established contact allergens in humans, 54 single chemicals and 28 natural extracts. Of these, 12 chemicals and 8 natural extracts were found to pose a high risk of sensitisation to the consumer, considering the high number of reported cases. In particular one ingredient stood out, hydroxyisohexyl 3-cyclohexene carboxaldehyde, having been the cause of more than 1500 reported cases since the 1999 opinion.

Moreover, animal experiments indicate that additional fragrance substances can be expected to be contact allergens in humans, although human evidence is currently lacking. Additionally, limited *in vivo* evidence together with Structure-Activity Relationship analysis suggests that other fragrance ingredients may also be a cause of concern with regard to their potential of causing contact allergy in humans.

The review also lists fragrance substances that can act as prehapten or prohaptens, forming new or more potent allergens by air oxidation and/or metabolic activation. Such

activation processes are of concern as they increase the risk of sensitisation and also the risk for cross reactivity between fragrance substances. In addition to known prehapten fragrance substances, the SCCS performed SAR analyses to identify fragrance substances with structural alerts that indicate that they are possible prehapten. While in the case of prohapten the possibility of becoming activated is inherent to the molecule and cannot be avoided, the activation of prehapten can be prevented by appropriate measures.

The SCCS examined available elicitation dose-response data to decide whether safe thresholds can be established for the fragrance allergens of concern, i.e. those found to pose a high risk of sensitisation to consumers. The SCCS considers that thresholds based on elicitation levels in sensitised individuals will be sufficiently low to protect both the majority of sensitised individuals as well as most of the non-sensitised consumers from developing contact allergy. As data from human dose elicitation experiments are very limited in several respects, no levels that could be considered safe for the majority of contact allergic consumers could be established for individual substances. The studies available, however, indicate that a general level of exposure of up to 0.8 µg/cm² (0.01% in cosmetic products) may be tolerated by most consumers, including those with contact allergy to fragrance allergens. The SCCS is of the opinion that this level of exposure (up to 0.01%) would suffice to prevent elicitation for the majority of allergic individuals, unless there is experimental or clinical substance-specific data allowing the derivation of individual thresholds.

It was not possible to provide a safe threshold for natural extracts of concern, as no specific investigations exist and the model providing the general threshold (0.01%) has been based on individual chemicals only. However the SCCS considers that the maximum use concentration applies to the identified chemicals both if added as chemicals or as an identified constituent of a natural ingredient. This will also reduce the risk of sensitisation and elicitation from natural extracts.

The suggested general threshold, although limiting the problem of fragrance allergy in the consumer significantly, would not preclude that the most sensitive segment of the population may react upon exposure to these levels and does not remove the necessity for providing information to the consumer concerning the presence of the listed fragrance substance in cosmetics.

In the case of hydroxyisohexyl 3-cyclohexene carboxaldehyde, the SCCP had recommended limiting the concentration in cosmetics to 200 ppm. Recent voluntary restrictions (recommendations to lower use concentrations, at least for some product types, to the level recommended by the SCCS in 2003) are not reflected in available evidence and are considered insufficient. The SCCS considers that the number of cases of HICC allergy documented over the last decade is exceptionally high and that continued exposure to HICC by the consumer is not considered safe, even at concentrations as low as 200 ppm. Therefore, HICC should not be used in consumer products in order to prevent further cases of contact allergy to HICC and to limit the consequences to those who already have become sensitized.

The SCCP concluded in 2004 that chloroatranol and atranol, the main allergenic constituents of *Evernia prunastri* and *Evernia furfuracea*, should not be present in products for the consumer. The persistently high frequency of contact allergy to *Evernia prunastri* and *Evernia furfuracea* noted in eczema patients does point to a persisting problem with exposure to the allergenic constituents. The SCCS is of the opinion that the presence of the two constituents, chloroatranol and atranol, in cosmetic products are not safe.

1. Background

As a result of the public consultation on perfumery materials, which ended on 27 January 2007, there were further requests and information on important and/or frequently used allergens other than those proposed for regulation, such as farnesol, citral, linalool and hydroxyisohexyl-3-cyclohexenecarboxaldehyde. These substances were not part of the consultation, but they all belong to the 26 fragrance substances which should be labelled when present in cosmetic products under certain conditions.

The 26 fragrance substances were introduced into annex III of the Cosmetics Directive by the 7th amendment (2003/15/EC) on the basis of the SCCNFP draft opinion (SCCNFP/0017/98) published on 30 September 1999 for public consultation and the final opinion adopted by the SCCNFP during the plenary session of 8 December 1999.

Thirteen of the allergenic fragrance substances listed in this opinion have been frequently reported as well-recognised contact allergens in consumers and are thus of most concern; 11 others are less well documented. See the lists below from the opinion.

List A: *Fragrance chemicals, which according to existing knowledge, are most frequently reported and well-recognised consumer allergens.*

| Common name | CAS number |
|---|-------------------|
| Amyl cinnamal | 122-40-7 |
| Amylcinnamyl alcohol | 101-85-9 |
| Benzyl alcohol | 100-51-6 |
| Benzyl salicylate | 118-58-1 |
| Cinnamyl alcohol | 104-54-1 |
| Cinnamal | 104-55-2 |
| Citral | 5392-40-5 |
| Coumarin | 91-64-5 |
| Eugenol | 97-53-0 |
| Geraniol | 106-24-1 |
| Hydroxycitronellal | 107-75-5 |
| Hydroxymethylpentyl-cyclohexenecarboxaldehyde | 31906-04-4 |
| Isoeugenol | 97-54-1 |

List B: *Fragrance chemicals, which are less frequently reported and thus less documented as consumer allergens.*

| Common name | CAS number |
|--|-------------------|
| Anisyl alcohol | 105-13-5 |
| Benzyl benzoate | 120-51-4 |
| Benzyl cinnamate | 103-41-3 |
| Citronellol | 106-22-9 |
| Farnesol | 4602-84-0 |
| Hexyl cinnamaldehyde | 101-86-0 |
| Lilial | 80-54-6 |
| d-Limonene | 5989-27-5 |
| Linalool | 78-70-6 |
| Methyl heptine carbonate | 111-12-6 |
| 3-Methyl-4-(2,6,6-trimethyl-2-cyclohexen-1-yl)-3-buten-2-one | 127-51-5 |

Furthermore, two fragrances (natural mixtures) were added

| Common name | CAS number |
|--------------------|-------------------|
| Oak moss | 90028-68-5 |
| Tree moss | 90028-67-4 |

At the time there were insufficient scientific data to allow for the determination of dose-response relationships and/or thresholds for these allergens. Nevertheless, in a pragmatic administrative decision the limits of 0.01 and 0.001% were set, for rinse-off and leave-on products respectively.

Scientific information of both a general and a specific nature has been submitted to DG ENTR in order to ask the SCCS for a revision of the 26 fragrances with respect to further restrictions and possible even delisting.

2. Terms of reference

- 1. Does the SCCS still consider that the fragrance allergens currently listed in Annex III, entries 67-92, for labelling purposes represent those fragrance ingredients that the consumer needs to be made aware of when present in cosmetic products?*
- 2. Can the SCCS establish any threshold for their safe use based on the available scientific data?*
- 3. Can the SCCS identify substances where processes (e.g. metabolism, oxidation and hydrolysis) may lead to cross-reactivity and new allergens which are relevant for the protection of the consumer?*

3. Introduction

Fragrance ingredients

Fragrance and flavour substances are organic compounds with characteristic, usually pleasant, odours. They are ubiquitously used in perfumes and other perfumed cosmetic products, but also in detergents, fabric softeners, and other household products where fragrance may be used to mask unpleasant odours from raw materials. Flavourings are used in foods, beverages, and dental products. Fragrance substances are also used in aromatherapy and may be present in herbal products, and used as topical medicaments for their antiseptic properties.

Contact allergy to fragrance ingredients occurs when an individual has been exposed, on the skin, to a sufficient degree of fragrance contact allergens. Contact allergy is a life-long, specifically altered reactivity in the immune system. This means that once contact allergy is developed, cells in the immune system will be present which can recognise and react towards the allergen. As a consequence, symptoms, i.e. allergic contact dermatitis, may occur upon re-exposure to the fragrance allergen(s) in question. Allergic contact dermatitis is an inflammatory skin disease characterised by erythema, swelling and vesicles in the acute phase. If exposure continues it may develop into a chronic condition with scaling and painful fissures of the skin. Allergic contact dermatitis to fragrance ingredients is most often caused by cosmetic products and usually involves the face and/or hands. It may affect fitness for work and the quality of life of the individual.

Fragrance contact allergy has long been recognised as a frequent and potentially disabling problem. Prevention is possible as it is an environmental disease and if the environment is modified (e.g. by reduced use concentrations of allergens), the disease frequency and severity will decrease. Ingredient information is a cornerstone in the prevention of allergic contact dermatitis, as knowledge about the allergens which a patient has been exposed to is crucial for including the right substances in the allergy test, and for subsequent information on avoidance of re-exposure. However, the labelling rules in the Cosmetics Directive 76/768/EEC stipulated that perfume and aromatic compositions and their raw materials shall be referred to by the word "perfume" or "aroma", rather than being labelled individually. This is the reason why the SCCNFP in their opinion SCCNFP/0017/98 (1) identified 26 fragrance allergens for which information should be provided to consumers concerning their presence in cosmetic products. This was implemented in the Cosmetics Directive as individual ingredient labelling of the 26 fragrance allergens (Annex III, entries 67-92). However, safe use concentrations of these fragrances in cosmetic products had not yet been determined and much new evidence concerning fragrance allergy has been published since the 1999 opinion. The present request to review the list of recognised fragrance allergens which the consumer needs to be made aware of, to indicate thresholds for their safe use and to consider possible modification of allergens by metabolism and autoxidation, required a thorough review of all relevant scientific data. This includes both published scientific literature as well as unpublished scientific information on fragrances from the industry. The International Fragrance Association (IFRA), as representative of the fragrance industry, was contacted to provide relevant unpublished scientific data on fragrance ingredients. This information, together with the up-to-date published scientific literature, has been critically reviewed for the present SCCS opinion. The relevant data gaps are identified and recommendations for research addressing these gaps are made.

4. Clinical aspects of contact allergy to fragrance ingredients

4.1. Spectrum of reactions

Adverse reactions to fragrances in perfumes and in fragranced cosmetic products include allergic contact dermatitis, irritant contact dermatitis, photosensitivity, immediate contact reactions (contact urticaria), and pigmented contact dermatitis. Airborne and connubial contact dermatitis occur.

4.1.1. Allergic contact dermatitis

Mechanism

Allergic contact dermatitis (ACD) depends primarily on the activation of allergen-specific T-cells. In allergic contact dermatitis, a distinction is made between induction (sensitisation) and elicitation phases. A useful review is available (2).

The induction phase includes the events following initial contact with the allergen and is complete when the individual is sensitised and capable of giving a positive allergic contact dermatitis reaction.

The elicitation phase begins upon re-exposure to the allergen (challenge) and results in clinical manifestation of allergic contact dermatitis.

The entire process of the induction phase requires ca. 10 days to several weeks, whereas an elicitation phase reaction develops within 1–2 days.

Most contact allergens are small, chemically reactive compounds. As these compounds are too small to be directly immunogenic, they act as haptens; i.e. they react with higher molecular weight epidermal and/or dermal biomolecules to form immunogenic adducts. It is usually considered that the biomolecules involved are free or membrane bound proteins, which react via nucleophilic thiol, amino, and hydroxyl groups.

Dendritic cells (DCs) and the local tissue microenvironment are crucial factors in the development of ACD. Langerhans cells (LCs), as epidermal DCs, and dermal DCs are pivotal for the sensitisation and the elicitation phases of ACD. During sensitisation, DCs react with the immunogenic complexes by interaction with neighbouring keratinocytes, migration to the local draining lymph nodes and the priming of naïve T-cells. These reactions are mediated by inflammatory cytokines, chemokines and adhesion molecules. Antigen specific effector T-cells are then recruited into the skin upon contact with the same hapten (elicitation). Following their recruitment these T-cells are activated by antigen-presenting skin cells, including LCs, dermal DCs and keratinocytes, and macrophages.

Although most allergens can form hapten–carrier complexes directly, some need activation, e.g. by enzyme-induced metabolic conversion or abiotic oxidation. Such compounds are termed prohaptens and prehaptens, respectively, and are discussed in more detail in chapter 5. Well known examples of prehaptens and prohaptens are limonene and eugenol. Reduced enzyme activity in certain individuals, related to genetic enzyme polymorphisms, may give an increased or reduced risk of sensitisation to prohaptens (that need enzymatic activation) in certain individuals or populations.

Once sensitised, individuals can develop allergic contact dermatitis upon re-exposure to the contact allergen. Positive patch test reactions mimic this process of allergen-specific skin hyper-sensitivity. Skin contact induces an inflammatory reaction that is maximal within 2–3 days and, without further allergen supply, then declines.

Overview of clinical features

Perfumes and deodorants are the most frequent sources of sensitisation to fragrance ingredients in women, while aftershave products and deodorants are most often responsible in men (3). Thereafter, eczema may appear or be worsened by contact with other

fragranced products such as cosmetics, toiletries, household products, industrial contacts and flavourings.

Contact allergy to a particular product or chemical is established by means of diagnostic patch testing. When patients with suspected allergic cosmetic dermatitis are investigated, fragrances are identified as the most frequent allergens, not only in perfumes, after-shaves and deodorants, but also in other cosmetic products. Evaluation of perfume allergy may be difficult; a perfume compound may consist of ten to > 300 basic components selected from about 2500 materials.

Between 6 and 14% of patients routinely tested for suspected allergic contact dermatitis react to a standard indicator of fragrance allergy, the Fragrance Mix I (4), see also chapter 4.3.2. When tested with ten popular perfumes, 6.9% of female eczema patients proved to be allergic to them (5) and 3.2–4.2% were allergic to fragrances from perfumes present in various cosmetic products (6). The finding of a positive reaction to the Fragrance Mix I should be followed by a search for its relevance, i.e. is fragrance allergy the cause of the patient's current or previous complaints, or does it at least contribute to it? Between 50 and 65% of all positive patch test reactions to the mix are relevant. Sometimes, correlation with the clinical picture is lacking and many patients appear to tolerate perfumes and fragranced products without problems (7). This may be explained by: a) irritant (false-positive) patch test reactions to the mix; b) the absence of relevant allergens in those products; and c) the concentration being too low to elicit clinically visible allergic contact reactions. Contact allergy to fragrances often causes dermatitis of the hands (and aggravation of), face and neck, axillae and patches in areas where perfumes are dabbed on such as behind the ears, upper chest, elbow flexures and wrists. Depending on the degree of sensitivity and exposure, the severity of dermatitis may range from mild to severe with dissemination (8) [pp 158–170].

Clinical studies have shown a highly significant association between reporting a history of visible skin symptoms from using scented products and a positive patch test to the Fragrance Mix I (9). Provocation studies with perfumes and deodorants have also shown that fragrance-mix-positive eczema patients often react to use-tests with the products. Subsequent chemical analysis of such products has detected significant amounts of one or more Fragrance Mix I ingredients, confirming the relevance of positive patch tests to the Fragrance Mix I in these patients (5, 10).

Hands

Contact sensitisation may be the primary cause of hand eczema, or may be a complication of irritant or atopic hand eczema. The number of positive patch tests has been reported to correlate with the duration of hand eczema, indicating that long-standing hand eczema may often be complicated by sensitisation (11). The most common contact allergies in patients with hand eczema are metals, the Fragrance Mix, *Myroxylon pereirae*, and colophonium (12).

Fragrance allergy may be a relevant problem in patients with hand eczema; perfumes are present in consumer products to which their hands are exposed (13). A significant relationship between hand eczema and fragrance contact allergy has been found in some studies based on patients investigated for contact allergy (14). However, hand eczema is a multi-factorial disease and the clinical significance of fragrance contact allergy in (severe) chronic hand eczema may not be clear. A review on the subject has been published (15).

Axillae

Bilateral axillary dermatitis may be caused by perfume in deodorants and, if the reaction is severe, it may spread down the arms and to other areas of the body (8) [pp 158–170]. In individuals who consulted a dermatologist, a history of such first-time symptoms was significantly related to the later diagnosis of perfume allergy (9).

Face

Facial eczema is an important manifestation of fragrance allergy from the use of cosmetic products (16). In men, aftershave products can cause an eczematous eruption of the beard area and the adjacent part of the neck (8) [pp 158–170], and men using wet shaving as opposed to dry have been shown to have an increased risk of 2.9 of being fragrance allergic (17).

4.1.2. Irritant reactions (including contact urticaria)

Irritant effects of some individual fragrance ingredients, e.g. citral (18, 19), are known. Irritant contact dermatitis from perfumes is believed to be common, but there are no existing investigations to substantiate this (7). Many more people complain about intolerance or rashes to perfumes/perfumed products than are shown to be allergic by testing (9). This may be due to irritant effects or inadequate diagnostic procedures.

Fragrances may cause a dose-related contact urticaria of the non-immunological type (irritant contact urticaria). Cinnamal, cinnamic alcohol, and *Myroxylon pereirae* are well recognised causes of contact urticaria, but others, including menthol, vanillin and benzaldehyde have also been reported (20). The reactions to *Myroxylon pereirae* may be due to cinnamates (21).

A relationship to delayed contact hypersensitivity was suggested (22), but no significant difference was found between a fragrance-allergic group and a control group in the frequency of immediate reactions to fragrance ingredients (20), in keeping with a non-immunological basis for the reactions seen.

4.1.3. Pigmentary anomalies

The term “pigmented cosmetic dermatitis” was introduced in 1973 for what had previously been known as melanosis faciei feminae when the mechanism (type IV allergy) and causative allergens were clarified (23). It refers to increased pigmentation, usually on the face/neck, often following sub-clinical contact dermatitis. Many cosmetic ingredients were patch tested at non-irritant concentrations and statistical evaluation showed that a number of fragrance ingredients were associated: jasmine absolute, ylang-ylang oil, cananga oil, benzyl salicylate, hydroxycitronellal, sandalwood oil, geraniol, geranium oil (24).

4.1.4. Photo-reactions

Musk ambrette produced a considerable number of allergic photocontact reactions (in which UV-light is required) in the 1970s (25) and was later banned from use in the EU. Nowadays, photoallergic contact dermatitis is uncommon (26). Furocoumarins (psoralens) in some plant-derived fragrance ingredients caused phototoxic reactions with erythema followed by hyperpigmentation resulting in Berloque dermatitis (8) [pp 417–432]. There are now limits for the amount of furocoumarins in fragrance products. Phototoxic reactions still occur but are rare (27).

4.1.5. General/respiratory

Fragrances are volatile and therefore, in addition to skin exposure, a perfume also exposes the eyes and naso-respiratory tract. It is estimated that 2–4% of the adult population is affected by respiratory or eye symptoms by such an exposure (28). It is known that exposure to fragrances may exacerbate pre-existing asthma (29). Asthma-like symptoms can be provoked by sensory mechanisms (30). In an epidemiological investigation, a significant association was found between respiratory complaints related to fragrances and contact allergy to fragrance ingredients, in addition to hand eczema, which were independent risk factors in a multivariate analysis (31).

4.2. Patch testing

The diagnosis of contact sensitisation (or contact allergy – regarded here as synonymous) as the immunological alteration underlying allergic contact dermatitis is made by patch testing. This diagnostic tool involves the standardised application of small doses of a set of potential or individually suspected allergens for a period of 1 day or, mostly, 2 days. In the following days, exposed skin sites are checked for the occurrence of allergic reactions, which morphologically mimic allergic contact dermatitis occurring elsewhere, after exposure to culprit products. International guidelines for the application, reading and interpretation of the patch test exist (32). The present brief section does not intend to reiterate all technical and scientific aspects, but to outline some aspects of diagnostic patch testing which are often misunderstood (for a recent comment see also (33)).

- The patch test identifies whether the patient has contact allergy to a substance, but cannot contribute information on the clinical relevance of that contact allergy for the eczema that led to consultation and to patch testing (see 4.4.1).
- Exposure conditions of the patch test (one-time, prolonged occlusive application, usually in petrolatum or water, of a single substance) have been optimised to achieve above diagnostic aim, and thereby have nothing in common with exposures which lead to sensitisation and elicitation of allergic contact dermatitis. These are normally repetitive, often over weeks, months or years, non-occlusive, and to much lower concentrations and doses/area, respectively, but possibly on damaged or inflamed skin. In fact, the repeated open application test (ROAT), which is sometimes used after a positive patch test of uncertain validity to verify that contact allergy indeed exists mimicks these day-to-day exposure conditions, and typically involves single dosings which are a small fraction of the one-time patch test dose (see 11).
- It is self-evident that such (repeated, low-level) exposures must have occurred and have culminated in an adaptive immune response – therefore it is axiomatic that the substance involved is a skin sensitizer in humans (33).
- Repeated patch testing, which is a relatively rare event, does not contribute significantly to contact allergy (to fragrance allergens).
- Most allergen test preparations, and certainly those that are included in international baseline series, have evolved from studies critically (re-) appraising their diagnostic validity, i.e., sensitivity and specificity. Notwithstanding this, false-positive and false-negative reactions do occur (as with any diagnostic tool). While in the individual case such diagnostic misclassification may have unfortunate consequences, it will hardly impair epidemiological estimates of contact allergy frequency – at least as long as a reasonable balance between false-positive and false-negative reactions is achieved.

4.3. Epidemiology of fragrance allergy

4.3.1. Substances used for screening of contact allergy to fragrance ingredients

A fragrance formula may consist of ten to 300 or more different ingredients. The CosIng database lists 2587 ingredients used for perfuming¹, as well as several other materials classified as odour “masking” agents, which is equivalent with regard to allergy. A mixture of seven fragrance chemicals and one natural extract, which have been identified as major fragrance allergens in the past (34), are used for diagnosing contact allergy to fragrance

¹ <http://ec.europa.eu/enterprise/cosmetics/cosing/index.cfm?fuseaction=search.results&function=66&search>, last accessed 2009-10-14.

ingredients (Table 4-1). This mixture is called the Fragrance Mix (FM I) and is included in the standard patch test tray containing the most common allergens in Europe.

Table 4-1: Ingredients of Fragrance Mix I (FM I; 8% allergens in petrolatum).

| Single constituent: INCI name (common name) | Conc. (%) |
|--|-----------|
| Amyl cinnamal (alpha-amyl cinnamal) | 1 |
| Cinnamyl alcohol (cinnamic alcohol) | 1 |
| Cinnamal (cinnamic aldehyde) | 1 |
| Eugenol | 1 |
| Geraniol | 1 |
| Hydroxycitronellal | 1 |
| Isoeugenol | 1 |
| Oak moss absolute (a natural extract; INCI: <i>Evernia prunastri</i>) | 1 |
| Sorbitan sesquioleate (added as an emulsifier) | 5 |

Note: All single allergens of the above, when used for breakdown testing, are also in petrolatum.

However, due to the introduction of new fragrance ingredients (with allergenic potential), the above Fragrance Mix I was deemed not to be sufficient for the diagnosis of fragrance allergy. Thus, Fragrance Mix II was devised to supplement Fragrance Mix I in a European multicentre study (35, 36). Since then, FM II has been included in the European baseline series. Table 4-2 lists the ingredients of FM II. In addition to being tested in FM II, hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC) is also tested separately at 5% test concentration in the baseline series (37).

Table 4-2: Ingredients of Fragrance Mix II (FM II; 14% allergens in petrolatum).

| Single constituent: INCI name (common name) | Conc. (%) |
|---|-----------|
| Citronellol | 0.5 |
| Citral | 1 |
| Coumarin | 2.5 |
| Hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC) | 2.5 |
| Farnesol | 2.5 |
| Alpha-hexyl-cinnamal | 5 |

Note: All single allergens of the above, when used for breakdown testing, are also in petrolatum.

Patch test results in patients and in population samples with these two screening mixes, and single allergens, will be presented and discussed in the following two sections.

4.3.2. Clinical epidemiology

For a number of reasons the bulk of the evidence regarding the frequency of contact allergy to fragrance ingredients relies on clinical data, i.e. the history, clinical presentation and test results of patients patch tested for suspected allergic contact dermatitis – in general, and not specifically due to fragrance ingredients. The frequency of contact allergy to fragrance ingredients (or other contact allergies, for that matter) cannot be related to the population

directly, as it is derived from a subgroup (of patients) selected for specific morbidity. Nevertheless, these data can be examined epidemiologically assuming a largely similar selection process: (i) across time in a given department; and (ii) between departments at any point of time. If the notion of similarity, and thus direct comparability, does not appear valid, adjustment or standardisation techniques can be employed to account for differences, e.g. the average age of patients in a time series on a (fragrance) allergen with age-associated risk of sensitisation. In this situation, changes in the age composition of the patients tested may confound a time trend. A distinction must be made between patch testing "consecutive" patients, i.e. all patients who are patch tested for suspected contact sensitisation, and "aimed" patch testing, i.e. application of allergens only in the subset of patients in whom exposure to the particular allergens of the applied "special series" is suspected. For any given allergen, the latter "aimed" approach will usually yield higher sensitisation prevalences than the testing of not-further-selected "consecutive" patients. Thus, information on the inclusion of an allergen either in a baseline series (tested in virtually all patients) or in a special series (applied in an aimed fashion) must be considered and is given in the following tables, where available in the cited references.

Notwithstanding the potential pitfalls of clinical data, they have proven useful in identifying emerging trends or persisting problems, and also in evaluating the effect of preventive action – either regarding the entire population, or subgroups thereof, such as certain occupations. Regarding the fragrance mixes (FM I and FM II) mentioned above, evidence regarding sensitisation frequencies published since 1999 will be outlined below, thus supplementing the data presented in the SCCNFP opinion on Fragrance Allergy in 1999 (1).

Fragrance Mix I ("Larsen Mix")**Table 4-3:** Results with screening agents for contact allergy to fragrance ingredients reported since 1999 in patients patch tested for suspected allergic contact dermatitis in Europe: Fragrance Mix "I" (see Table 4-1). If not given in the publication, the confidence interval (CI) was calculated from the absolute numbers by the SCCS (§).

| Country (Ref.) | Population | Year(s) | No. tested | Crude % positive (95% CI) |
|-----------------------------|--|-----------------|------------|----------------------------------|
| Sweden (38) | Consecutive patients | 2000 | 3790 | 6.9 |
| Hungary (39) | | 1998-1999 | 3604 | 8.2 (7.3–9.1) [§] |
| Czech Republic (40) | | 1997-2001 | 12058 | 5.8 (5.4–6.2) [§] |
| Ljubljana, Slovenia (41) | Consecutive patients | 1989-1998 | 6129 | 5.9 (5.3–6.5) [§] |
| Germany (42) | Consecutive IVDK patients | 1996-2002 | 59298 | 11.3 (11.0–11.5) [§] |
| Germany (43) | Consecutive IVDK patients | 2005-2008 | 36961 | 7.3 (7.0–7.6) [§] |
| Vienna, Austria (16) | Consecutive patients of one clinic | 1997-2000 | 2660 | 9.1 (8.1–10.3) [§] |
| Groningen, Netherlands (44) | Patients (fragrance allergy suspected) | 04/2005-06/2007 | 295 | 5.8 (3.4–9.1) [§] |
| The Netherlands (45) | Consecutive patients | 09/1998-04/1999 | 1825 | 10.6 (9.2–12.1) |
| The Netherlands (46) | Patients (cosmetic allergy suspected) | 1994-1998 | 757 | 14.8 (12.3–17.5) [§] |
| Leuven, Belgium (47) | Consecutive patients | 1990-2005 | 10128 | 9.1 (8.6–9.7) [§] |
| Coimbra, Portugal (48) | Consecutive patients | 07/1989-06/1999 | 2600 | 10.9 (9.7–12.2) [§] |
| Spain (49) | Consecutive patients | 10/2005-06/2008 | 1253 | 4.5 (3.4–5.8) [§] |
| Sheffield, UK (50) | Consecutive patients | 1994-1995 | 744 | 11.4 (9.2–13.9) [§] |
| St. John's, London, UK (51) | Consecutive patients | 1980-2004 | 34072 | 7.7 (7.4–8.0) [§] |
| Copenhagen, Denmark (52) | Consecutive patients | 1985-2007 | 16173 | 7.2 (6.8–7.6) [§] |
| ESSCA (53) | Consecutive patients | 2002-2003 | 9663 | 7.1 (6.6–7.6) [§] |
| ESSCA (54) | Consecutive patients | 2004 | 9941 | 7.6 (7.1–8.2) [§] |
| ESSCA (55) | Consecutive patients | 2005-2006 | 18542 | 7.0 (6.6–7.4) [§] |

Table 4-4: Results with screening agents for contact allergy to fragrance ingredients reported since 1999 in patients patch tested for suspected allergic contact dermatitis in non-European countries: Fragrance Mix "I" (see Table 4-1). If not given in the publication, the confidence interval (CI) was calculated from the absolute numbers by the SCCS (§).

| Country (Ref.) | Population | Year(s) | No. tested | Crude % positive (95% CI) |
|------------------------------------|-------------------------------|-----------------------|------------|-------------------------------|
| South Korea (56) | Consecutive patients | 04/2002–06/2003 | 422 | 9.7 (7.1–13.0) [§] |
| Lahore, Pakistan (57) | Dermatitis patients | 2 years prior to 2002 | 350 | 7.7 (5.2–11.0) [§] |
| Manipal, India (58) | Dermatitis patients | 1989-1998 | 1780 | 3.1 (2.3–4.0) [§] |
| Tel Aviv, Israel [§] (59) | Consecutive patients | 1999-2000 | 943 | 8.5 (6.8–10.5) [§] |
| Tel Aviv, Israel (60) | Consecutive patients | 1998-2004 | 2156 | 7.1 (6.1–8.3) [§] |
| Tehran, Iran (61) | Consecutive patients | 2002-2004 | 250 | 4.0 (1.9–7.2) [§] |
| Ankara, Turkey (62) | Consecutive patients | 1992-2004 | 1038 | 2.1 (1.3–3.2) [§] |
| Beijing, China (63) | Consecutive patients | 2000-2003 | 378 | 15.9 (12.3–20.0) [§] |
| USA (Canada) (64) | Probably consecutive patients | 2003 | 1603 | 5.9 |
| NACDG 2009 (US and Canada) (65) | Consecutive patients | 2005-2006 | 4439 | 11.5 |

Note: § Possibly included in (60).

Beyond the studies discussed above, regarding a time trend of sensitisation to FM I, a significant increase of positive results to FM I until 1998, and a significant drop thereafter has been noted in the IVDK study covering 1996 to 2002 (42). A similar drop from 1999 to 2007 has been observed in female, but not male patients from Copenhagen (52). In accordance with these findings, the prevalence of positive reactions to FM I doubled, or thereabouts, from 1989-1993 to 1994-1998 in Ljubljana, Slovenia (41).

Within Europe, a comparison between different countries and clinical departments is possible. An EECDRG study covering 1996-2000 found 9.7% positives to FM I (range: 5.0–12.6% in ten departments from seven European countries (66). A different European study, covering 10/1997-10/1998, found 11.3% (95% CI: 9.9–12.9%) positive reactions to FM 1 in 1,855 patients; the variation between centres was marked: Gentofte 8.2% vs. Leuven 23.0% as extremes (67). In the first study of the European Surveillance System on Contact Allergies (ESSCA), covering 2002 and 2003, 9663 patients were patch tested with FM I, overall yielding 7.1% positive reactions with marked variation between participating departments. In Dortmund, Germany, the minimum frequency of 3.7% was noted, while in Lahti, Finland, the highest prevalence, namely 10.4%, was found (53). Subsequently, in the year 2004, the overall prevalence was 7.6%, i.e. largely unchanged (54). In the most recent study by ESSCA, based on 2005/2006 PT data across Europe, significant differences were again noted, this time on the aggregated level of European regions, with FM I sensitisation being the least frequent in the Southern countries (4.8% [95% CI: 3.9–5.5%] age- and sex-standardised prevalence) vs. 7.7% (95% CI: 7.0–8.4%) in the central European departments, with the Finnish, Polish and Lithuanian departments (5.7% [95%

CI: 4.6 – 6.8%]) and the UK network (6.8% [95% CI: 6.3 – 7.3%]) in an intermediate position (55).

Fragrance Mix II

Table 4-5: Results with screening agents for contact allergy to fragrance ingredients reported since 1999 in patients patch tested for suspected allergic contact dermatitis: Fragrance Mix "II" (see Table 4-2). The FM II was only conceived in 2005, so results are still sparse). If not given in the publication, the confidence interval (CI) was calculated from the absolute numbers by the SCCS ([§]).

| Country (Ref.) | Population | Year(s) | No. tested | Crude % positive (95% CI) |
|--|--|-----------------|-------------------|----------------------------------|
| EU (35) | Six clinical depts. | 10/2002-06/2003 | 1701 | 2.9 (2.2–3.9) [§] |
| IVDK, Germany (68) | Consecutive patients | 01/2005-12/2008 | 35633 | 4.9 (4.7–5.1) [§] |
| Groningen, Netherlands (44) | Patients (fragrance allergy suspected) | 04/2005-06/2007 | 227 | 9.3 (5.8–13.8) [§] |
| Leuven, Belgium (47) | Consecutive patients | 2005 only | 335 | 2.1 (0.8–4.3) [§] |
| Spain (49) | Consecutive patients | 10/2005-06/2008 | 1253 | 0.6 (0.2–1.1) [§] |
| Denmark (69) on behalf of the DCDG, 2010 | Consecutive patients | 2005-2008 | 12302 | 4.5 (4.1–4.9) [§] |

Hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC)

Hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC) has been the most frequently reported chemical causing fragrance allergy since the 1999 opinion on fragrance allergy. In total, reports of about 1500 cases have been published in the scientific literature (see section 7.1).

HICC was recognised as an allergen in 1995 (70) and later included in the new perfume mixture, Fragrance Mix II (71), which is routinely used for the diagnosis of perfume allergy, see above. Furthermore, it is recommended to test separately with HICC, because it is a very frequent allergen (37) and detects relevant fragrance sensitisation which would otherwise have been missed (49). In the studies performed in European dermatology clinics, 0.5-2.7% of eczema patients have been found to be allergic to HICC with the highest frequency in central Europe (55). For further details see Table 4-6.

Table 4-6: Results with fragrance contact allergy screening agents reported since 1999 in patients patch tested for suspected allergic contact dermatitis: **HICC** (5% pet. if not stated otherwise). If not given in the publication, the confidence interval (CI) was calculated from the absolute numbers by the SCCS ([§]).

| Country (Ref.) | Population | Year(s) | No. tested | Crude % positive (95% CI) |
|-----------------------|----------------------|-----------------|-------------------|----------------------------------|
| Lithuania (72) | Consecutive patients | 04/2006-10/2008 | 816 | 0.9 (0.3–1.8) [§] |
| Spain (49) | Consecutive patients | 10/2005-06/2008 | 852 | 0.8 (0.3–1.7) [§] |
| Germany (CH, AT) (73) | Consecutive patients | 03/2000-02/2001 | 3245 | 1.9 (1.5–2.4) [§] |

| Country (Ref.) | Population | Year(s) | No. tested | Crude % positive (95% CI) |
|-----------------------|-------------------------------|-----------------|------------|----------------------------|
| Germany (CH, AT) (74) | Consecutive patients | 01/2003-12/2004 | 21325 | 2.4 (2.2-2.6) [§] |
| Germany (CH, AT) (68) | Consecutive patients | 01/2005-12/2008 | 35582 | 2.3 (2.2-2.5) [§] |
| Belgium (47) | Consecutive patients | 2002-2005 | 2901 | 2.1 (1.6-2.7) [§] |
| Denmark (69) | Consecutive patients | 2005-2008 | 12302 | 2.4 (2.1-2.7) [§] |
| South Korea (56) | Consecutive patients | 04/2002-06/2003 | 422 | 1.7 (0.6-3.4) [§] |
| USA, Canada (64) | Probably consecutive patients | 2003 | 1603 | 0.4 (0.2-0.9) [§] |

Myroxylon pereirae (Balsam of Peru)

Myroxylon pereirae is a balm obtained from a Central American tree. It is used as a screening substance for fragrance allergy in Europe and other geographical areas. Although the crude balm is not used in Europe in cosmetics, extracts and distillates are used (75). This natural mixture has been employed as screening agent in the baseline series for many decades. Hence, a wealth of data is available; Table 4-7 summarises results of the past 10 years.

Table 4-7: Results with fragrance contact allergy screening agents reported since 1999 in patients patch tested for suspected allergic contact dermatitis: ***Myroxylon pereirae resin*** (Balsam of Peru) (25% pet.). If not given in the publication, the confidence interval (CI) was calculated from the absolute numbers by the SCCS ([§]).

| Country (Ref.) | Population | Year(s) | No. tested | Crude % positive (95% CI) [§] |
|-------------------------|------------------------------------|-------------------|------------|--|
| Tel Aviv, Israel (59) # | Consecutive patients | 1999-2000 | 943 | 6.6 (5.1-8.4) [§] |
| South Korea (56) | Consecutive patients | 04/2002 – 06/2003 | 422 | 7.3 (5.1-10.3) [§] |
| Tel Aviv, Israel (60) | Consecutive patients | 1998-2004 | 2156 | 3.6 (2.9-4.5) [§] |
| Manipal, India (58) | Dermatitis patients | 1989-1998 | 1780 | 1.0 (0.5 – 1.5) [§] |
| Tehran, Iran (61) | Consecutive patients | 2002-2004 | 250 | 2.4 (0.9-5.2) [§] |
| Sevilla, Spain (76) | Consecutive patients | 2002-2004 | 863 | 5.8 (4.3-7.6) [§] |
| Ankara, Turkey (62) | Consecutive patients | 1992-2004 | 1038 | 2.1 (1.3-3.2) [§] |
| Vienna, Austria (16) | Consecutive patients of one clinic | 1997-2000 | 2660 | 5.4 (4.6-6.3) [§] |
| Czech Republic (40) | Consecutive patients | 1997-2001 | 12058 | 7.3 (6.8-7.8) [§] |
| Spain (49) | Consecutive patients | 10/2005-06/2008 | 1253 | 6.4 (5.1-7.9) [§] |

| Country (Ref.) | Population | Year(s) | No. tested | Crude % positive (95% CI) [§] |
|--|-------------------------------|-----------|------------|--|
| Copenhagen, Denmark (52) | Consecutive patients | 1985-2007 | 16173 | 3.9 (3.6-4.2) [§] |
| Sweden (38) | Consecutive patients | 2000 | 3790 | 6.5 |
| Nine European countries (53) | Consecutive patients | 2002-2003 | 9672 | 6.1 |
| Germany, three Swiss and one Austrian Dept. (43) | Consecutive patients | 2005-2008 | 36919 | 8.0 (7.7-8.3) |
| Ten depts. From seven EU countries (66) | Consecutive patients | 1996-2000 | 26210 | 6.0 |
| USA (Canada) (64) | Probably consecutive patients | 2003 | 1603 | 6.6 |
| NACDG 2009 (65) | Consecutive patients | 2005-2006 | 4449 | 11.9 |

Oil of turpentine

This natural extract is not tested in all baseline series. It is considered as a minor screening allergen for fragrance contact allergy. Moreover, oil of turpentine is used as a raw material in perfumery (see Annex I). Table 4-8 summarises results of the past 10 years with patch testing of consecutive patients.

Table 4-8: Results with fragrance contact allergy screening agents reported since 1999 in patients patch tested for suspected allergic contact dermatitis: **Oil of turpentine** (10% pet.) patients patch tested for suspected allergic contact dermatitis. If not given in the publication, the confidence interval (CI) was calculated from the absolute numbers by the SCCS ([§]).

| Country | Population | Year(s) | No. tested | Crude % positive (95% CI) [§] |
|--|-------------------------------------|-------------------------|------------|--|
| Lisbon, Portugal (77); virtually no .delta.-3-carene | Consecutive patients | 1979-1983 | 4316 | 2.3 (1.9-2.8) [§] |
| Birmingham, UK (78) | Potters with occup. hand dermatitis | 6 months; prior to 1996 | 24 | 14/4 pos. to "Indonesian turpentine" |
| Austria/Germany (IVDK) (79) | Consecutive patients | 1992-1995 | 27658 | 0.47 (0.39-0.55) [§] |
| Austria/Germany (IVDK) (42) | Consecutive patients | 1996-2002 | 59478 | Annual prevalence 1.6 to 4.4% |
| Augsburg, Germany (80) | Population sample | 1998 | 1141 | 1.2% (on population level!) |
| Europe (ESSCA) (53) | Consecutive patients | 2002/03 | 3767 | 1.6% |
| Austria/Germany/Switzerland (IVDK) (43) | Consecutive patients | 2005-2008 | 37163 | 1.8% |

An "overall burden" of fragrance contact allergy, in terms of the prevalence of contact allergy to at least one of the up-to-five screening allergens present in the baseline series (FM I, FM II, HICC, *Myroxylon pereirae*, oil of turpentine) has not been given in the published studies. A re-analysis of data from the two published studies of the IVDK (43, 68), covering central Europe from 2005 to 2008 (Germany, Austria and Switzerland), yielded an estimate of such overall prevalence of 16.2% (95% CI: 15.8-16.6%) (IVDK technical report, 2011-11-18).

4.3.3. Population-based epidemiology

In principle, the examination of a representative sample of the population is the most valid approach for estimating disease frequency, as there is no systematic selection process. However, in practice, participation of much less than 70% of those approached introduces the possibility of self-selection and thus of biased morbidity (or risk) estimates. Moreover, the resources needed prohibit regular, e.g. yearly, patch test studies in a sample of several thousand persons. For these reasons few studies exist (see Table 4-9).

A Swedish study of hand eczema in an industrial city showed that among 1,087 individuals recruited from the general population with symptoms of present or previous hand eczema, 5.8% were positive to the Fragrance Mix (81). In Denmark, Fragrance Mix sensitivity was found in 1.1% (0.3-2.1%) of 567 persons drawn as a sample from the general Danish population; only nickel sensitivity was more prevalent (82). In Italy, female patients with hand eczema caused by contact with detergents were patch tested. Of 1100 women, 3.1% reacted to Fragrance Mix I (83). A control group of 619 female patients with no eczema disease were also patch tested; 1.3% were positive to the Fragrance Mix (83). On the other hand, in a sample of 593 healthy Italian recruits, only three positive reactions (0.50%) to FM I were observed (84). Among Danish school children, 14-15 years of age, fragrance contact allergy was detected in 1.8% by patch testing with Fragrance Mix I (85). A study of 85 American student nurses showed that 15 (17.6%) had a positive reaction to Fragrance Mix I; 12 of the individuals also had a positive history of contact dermatitis (86). In this study the concentration of Fragrance Mix I was 16% as opposed to the currently recommended concentration of 8% and the study included only young females. Both of these factors may have contributed to the high prevalence of fragrance sensitivity found.

In 1990, 1998 and 2006, samples of the Danish adult population living in the Copenhagen area were patch tested with the European baseline series. In total 4299 individuals aged 18-69 years (18-41 years only in 1998) completed a pre-mailed questionnaire and were patch tested with FM I and *Myroxylon pereirae* (82, 87, 88). In 1990, 1.1% were found positive to FM I and in 2006, 1.6% were positive, which means no general change. However, when the age group of 18-41 years was analysed, the prevalence of FM I sensitisation followed an inverted V-pattern among women, i.e. an increase from 0.7% in 1990 to 3.9% in 1998, followed by a decrease to 2.3% in 2006. The participation rate varied in the three samples from 71.5% in 1990 to 52.4% in 1998, and to 43.7% in 2006 (82, 87, 88).

Contact sensitisation to FM I is strongly age related, with the relative risk more than doubling in the older age groups, compared to younger PT patients. This has been found in both bivariate (89) and adjusted multifactorial analyses (90). Hence, in older samples of the population, the prevalence of contact allergy to fragrance ingredients in general, and to FM I in particular, can be expected to be higher than in younger samples. From this background, the strikingly high prevalence observed in the MONICA/KORA allergy study in Augsburg, Germany (see Table 4-9) (80), may be explained, together with some residual confounding from the rather complex sampling process.

Opinion on fragrance allergens in cosmetic products

Table 4-9: Results from patch testing with Fragrance Mix I in different population based groups.

| Country (Ref.) | Population | Year(s) | No. tested | % positive (95% CI) |
|------------------|---|--------------------|------------|---------------------|
| Italy (83) | Females without eczema | Not given | 619 | 1.3 |
| Italy (84) | Male recruits | Not given | 593 | 0.50 |
| Denmark (82) | Population sample adults, 15-69 years | 1990-91 | 567 | 1.1 |
| Denmark (85) | School children 12-16 years old | 1995/96 | 717 | 1.8 |
| Denmark (82, 87) | Population sample adults, 18-41 years | Jan-Nov 1998 | 414 | 2.7 |
| Denmark (88) | Population sample adults, 18-69 years | June 2006–May 2008 | 3460 | 1.6 |
| Norway (91) | Population sample adults, 18-69 years. (Results reported in 2007) | 1994 (92) | 1236 | 1.8 (1.1–2.7) |
| Germany (80) | Subgroup of MONICA sample, age 25-74 | 1994/95 | 1141 | 11.4 |
| USA (86) | Student nurses, females | 1980 | 85 | 17.6* |
| Sweden (81) | Population sample adults, age 20-65 years reporting hand eczema | 1983-84 | 1087 | 5.8* |

Note: * Testing performed with Fragrance Mix I, containing 16% allergens; the currently used Fragrance Mix I contains 8% allergens (see above).

Table 4-10: Results from patch testing with other fragrance allergens in different population based groups. If not given in the publication, the confidence interval (CI) was calculated from the absolute numbers by the SCCS (5).

| Country (Ref.) | Population | Year(s) | Fragrance allergen | No. tested | % positive (95% CI) [§] |
|----------------|---|--------------|--|-------------|---|
| Thailand (93) | Convenience sample (via advertisement), age 18-55 | Not given | Isoeugenol, <i>Evernia prunastri</i> , <i>Myroxylon pereirae</i> * | 2545 | Positive to at least one of three allergens: 2.5 (1.9–3.2) [§] |
| Germany (80) | Subgroup of MONICA sample, age 25-74 | 1994/95 | <i>Myroxylon pereirae</i> | 1141 | 2.4 |
| Denmark (88) | Population sample, age 18-69 | 1990 2006 | <i>Myroxylon pereirae</i> | 567 3460 | 1.1 0.1 |

Note: * *Myroxylon pereirae* is a balm obtained from a Central American tree. It is used as a screening substance for fragrance allergy in Europe and other geographical areas. Although the crude balm is not used in Europe in cosmetics, extracts and distillates are used (75).

4.4. Consumer products as a cause of fragrance contact sensitisation and allergic contact dermatitis

4.4.1. Clinical relevance

Clinical relevance is a concept used to describe the significance of a positive (allergic) patch test reaction for an individual patient: a reaction is deemed relevant if contact allergy to the substance is associated with previous or current episodes of allergic contact dermatitis. Thereby, the evaluation of clinical relevance links past exposure to morbidity. For the evaluation of relevance, past or recent exposure(s) to the allergen need to be identified in the patient's history. The success of this process generally depends on:

- The patient's understanding and awareness;
- The dermatologist's knowledge concerning exposures;
- Ingredient labelling; and
- Information about the actual chemical composition of the implicated product.

As these requirements may be met to a varying extent, the validity of relevance information as reported in clinical studies may also be variable. However, information on clinical relevance is important, in principle, because the proportion of currently relevant sensitisations reflects the amount of current exposure and resulting disease state, which may increase or decrease with time. In this way, current relevance also reflects the direct burden of a fragrance contact allergy to the individual and indirectly to society. Further important aspects of the evaluation of clinical relevance as a final step of patch testing have been discussed (32, 94-96).

Generally, clinical relevance is categorised as "current", "previous" or "unknown". Further differentiation has been introduced by adding information on:

- Occupational versus non-occupational causation; and
- The level of certainty of the relevance statement, e.g. as "certain", "probable", "possible".

In some cases, clinical relevance may not be established due to:

- Immunological cross-reactivity with an individual allergen, diagnosed or not;
- Active sensitisation by the patch testing;
- Contact sensitisation not caused by the substance, but by a contaminating constituent; or
- Failure to test with a true hapten (e.g. haptens formed from prehapten on exposure to air, see chapter 5).

It should be noted that this statement on clinical relevance refers to the past history of a patient. This implies that a lack of, or unknown, clinical relevance does not make future allergen avoidance unnecessary.

In the context of contact allergy to fragrance ingredients, a number of alternative concepts of relevance have been used, for example:

- A history of intolerance to perfume or to perfumed products;
- A history of intolerance to perfume actually containing the allergen diagnosed;
- Detection of the culprit allergen in a perfume previously used.

4.4.2. Elicitation with clinical symptoms/signs, current and past

In case reports or small series, the clinical relevance of positive patch test reactions is usually well established and presented in detail. Moreover, a few large-scale clinical studies on contact allergy to fragrance ingredients have reported results on clinical relevance, which will be presented and discussed in this section. The studies can be subdivided into those which focus on medical history, patch testing with consumer products or detection of specific allergens in consumer products used by patients.

Medical history

A series of studies conducted in the 1990s showed that most individuals with contact allergy to fragrance ingredients were aware that they could not tolerate fragranced products on their skin and were able to specifically name product categories that initiated their disease (9). In this context, colognes, deodorants and lotions were named significantly more often by fragrance allergic dermatitis patients than by patients without fragrance contact allergy (3). These studies are described in the SCCNFP opinion on fragrance allergy of 1999 (1). Newer studies are outlined below.

NACDG 2009 study (65)

The definition of "present" clinical relevance in this North American network study was strict, requiring:

- A positive use or patch test with the suspected item(s) for "definite" relevance; and
- Verification of the presence of the allergen in known skin contactants, and consistent clinical presentation for "probable".

If these conditions were not met, but skin contact to items generally containing the item was likely, "possible" was used.

Regarding fragrance allergens, the proportions were as described in Table 4-11.

Table 4-11: Extract from ((65) Table 3) regarding the proportion of patients with "present clinical relevance" (see text) and "past clinical relevance" (criteria not given).

| Fragrance allergen | n (tested) | % (pos.) | Current relevance (%) | | | Past relevance (%) |
|---------------------------|------------|----------|-----------------------|----------|----------|--------------------|
| | | | Definite | Probable | Possible | |
| <i>Myroxylon pereirae</i> | 4449 | 11.9 | 1.3 | 33 | 53 | 2.7 |
| FM I | 4439 | 11.5 | 2.0 | 29.4 | 54.3 | 4.3 |
| Cinnamal | 4435 | 3.1 | 1.5 | 33.8 | 50 | 2.9 |
| Ylang-Ylang oil | 4434 | 1.5 | 4.6 | 10.8 | 73.8 | 1.5 |
| Jasmine absolute | 4447 | 1.1 | 0 | 24.5 | 67.3 | 6.1 |

Frosch 2002 (a) study (67)

In this study, 1,855 consecutive patients were patch tested with FM I and a series of a further 14 fragrance chemicals. Prior to the test, the history of adverse reactions to fragrances was classified as "certain" (6.6%), "probable" (8.0%), "questionable" (9.2%) or "none" (76.1%) (see (71)).

Frosch 2002 (b) study (97)

A series of 18 essential oils or components thereof, together with FM I, was assessed in 1,606 consecutive patients. Similar to the above study, the proportions of patients with a "certain" or "probable" history (or otherwise) and positive reactions to either FM I or the

special series, or both, were cross-tabulated. Of note, 53.7% of patients with positive reactions to FM I only, had no history. Similarly 54.2% of patients with positive reactions only to one of the essential oils had no history. However, in cases of reactivity to both FM I and one of the essential oils, the proportion of patients with no history was only 36.5%.

Frosch 2005 study (35)

The diagnostic properties of FM I and the new FM II were evaluated in 1,701 consecutive patients patch tested in six European centres. Contrasting a "certain" (found in 8.7% of patients) with "no history" (75.3% of patients), the sensitivity of FM I was 25.2%, and the positive predictive value (PPV) 45.1%. In comparison, the sensitivity of FM II at 14% concentration was 13.5% and the PPV was 55.6%. The combination of the two mixes was important, as more patients with a "certain" history, but also independently from history, reacted to just one of the mixes rather than to both.

Danish Contact Dermatitis Group 2005-2008 (69)

In 12302 consecutive patients patch tested in seven dermatology clinics and three university hospitals, 10.6% were positive to one or more of the fragrance allergy markers (FM I, FM II, *Myroxylon pereirae* or hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC)). Clinical relevance covered current and/or past relevance based on: 1) medical history; 2) results of patch and/or use tests; 3) ingredient labelling; or 4) chemical analysis. Clinical relevance was found in 71.0% of cases positive to FM I, 72.2% of those positive to FM II and 76.7% of those positive to HICC. These proportions were higher than the average for other cosmetic allergens such as preservatives and hair dyes, which gave relevant reactions in about 50% of those positive, as did *Myroxylon pereirae*. *Myroxylon pereirae* itself is not used in cosmetics as it is banned, but sensitisation may be caused by exposures to related substances and thus relevance may be difficult to determine.

Cosmetic products

Fragrance formulae from cosmetic products

Popular fine fragrances (5), as well as toilet soaps, shampoos, lotions, deodorants, and aftershaves have been shown to provoke allergic contact dermatitis in patients when used for patch testing (5, 6, 98, 99). Moreover, commercially available fragrance formulae and dilutions of individual fragrance allergens were potent elicitors of allergic contact dermatitis under simulated use conditions (10, 100, 101).

More recently, deodorants spiked with the fragrance allergens cinnamal, hydroxycitronellal and HICC, respectively, in realistic in-use concentrations were shown to elicit allergic contact dermatitis in 89-100% of the fragrance allergic individuals tested (102-104). In 87.5% of HICC sensitised individuals the use of a cream (and in 82.8% the use of an ethanol solution) spiked with HICC provoked dermatitis (105). These studies are discussed in more detail in chapter 11 on quantitative aspects. Other new studies are mentioned below:

IVDK "own perfumes" study (106)

A different perspective on clinical relevance is provided by assessing the proportion of positive reactions to the FM I or single fragrance allergens in patients who had not tolerated certain perfumed products, such as deodorants and aftershaves and who were patch test positive to these cosmetics. The following two tables are taken from this publication.

Table 4-12: Extract from ((106) Table 2) on the frequency of positive reactions to fragrance allergens in patients with vs. without positive patch test reaction to their own deodorant.

| Fragrance allergen | Conc. (%) | Deodorant positive (n=66) | | Deodorant negative (n=855) | |
|---------------------------|-----------|---------------------------|------------------|----------------------------|------------------|
| | | n (test) | % pos. (95% CI) | n (test) | % pos. (95% CI) |
| Fragrance Mix I | 8 | 61 | 38.0 (24.1-51.9) | 805 | 15.0 (12.5-17.5) |
| <i>Myroxylon pereirae</i> | 25 | 60 | 22.9 (12.7-33.1) | 806 | 9.1 (7.2-11.0) |
| Hydroxycitronellal | 1 | 33 | 6.5 (0.7-12.3) | 204 | 4.3 (1.5-7.1) |
| Isoeugenol | 1 | 33 | 6.5 (0.7-12.3) | 204 | 7.2 (3.6-10.8) |
| Cinnamal | 1 | 29 | 11.3 (0-24.1) | 133 | 1.1 (0-2.7) |
| Geraniol | 1 | 29 | 8.3 (0-20.4) | 141 | 0 (0-2.1) |

Of the 66 patients with a positive patch test reaction to their own deodorant, most had positive reactions to one or more fragrance allergens. This was much more prevalent than those patients in whom no positive reaction to their deodorant was observed. This observation supports the notion that the respective fragrance allergens are important in contact allergy to fragrance ingredients caused by deodorants, supporting data regarding exposure (chapter 10.1).

Table 4-13: Extract from ((106) Table 2) on the frequency of positive reactions to fragrance allergens in patients with vs. without positive patch test reaction to their own aftershave, eau de toilette or perfume.

| Fragrance allergen | Conc. (%) | Product positive (n=63) | | Product negative (n=819) | |
|--|-----------|-------------------------|------------------|--------------------------|------------------|
| | | n (test) | % pos. (95% CI) | n (test) | % pos. (95% CI) |
| Fragrance Mix I | 8 | 56 | 57.1 (46.2-68.1) | 764 | 13.9 (11.4-16.4) |
| <i>Myroxylon pereirae</i> | 25 | 56 | 13.9 (7.3-20.4) | 766 | 8.8 (6.8-10.7) |
| HICC | 5 | 20 | 58.3 (37.5-79.0) | 310 | 1.3 (0-2.7) |
| <i>Evernia prunastri</i> | 1 | 28 | 22.1 (7.0-37.2) | 153 | 8.8 (4.2-13.4) |
| Hydroxycitronellal | 1 | 33 | 6.5 (0.7-12.3) | 204 | 4.3 (1.5-7.1) |
| <i>Cananga odorata</i> (ylang-ylang oil) | 10 | 7 | 16.3 (2.0-30.5) | 43 | 5.0 (0-11.3) |

Similar results were obtained from the subgroup of patients with a positive reaction to their eau de toilette, aftershave (hydroalcohol solutions) or perfumes (Table 4-13). However, notable differences were: (i) the greater relative importance of *Evernia prunastri* (Oak moss absolute); and (ii) generally an extremely high proportion of positive reactions to various other fragrance ingredients.

4.4.3. Elicitation in diagnostic patch tests without clinical history

In a variable proportion of patients, a positive patch test reaction does not correlate with recent or past episodes of presumptive allergic contact dermatitis. Apart from particular circumstances, such as cross-reactivity or reactivity to contaminants outlined above, there are several possible explanations for this:

- The patch test reaction was a false-positive (irritant).
- There was erroneous recall/interpretation of the patient's history (false-negative).
- Lack of knowledge concerning exposures.

- If the patient is weakly sensitised (e.g. by a low induction dose), the occlusive exposure during patch testing may have been the only exposure above the individual elicitation threshold capable of eliciting an unequivocal allergic contact reaction. In this situation, clinical relevance would be classified as “unknown”. Nevertheless, there is an alteration of the immune status of the individual.

Sometimes, a repeated open application or provocative use test is employed to mimic “normal” exposure to the allergen. A positive reaction to such a use-related test confirms actual sensitisation. Moreover, the positive result supports the necessity of future allergen avoidance. Apart from the risk of developing allergic contact dermatitis in the future, sensitisation means an alteration of the immune status of the individual.

4.5. Socio-economic impact of contact allergy

4.5.1. Health related quality of life

Skin diseases in general are known to affect quality of life significantly (107); this also applies to eczema, where most studies concern atopic dermatitis and hand eczema patients (108, 109). Hand eczema has a poor prognosis and may affect the self-image, limit social activities and lead to occupational restrictions (109, 110). The quality of life in hand eczema patients with fragrance contact allergy is affected in a similar degree as patients with other contact allergies (111).

In a questionnaire study of 117 patients recently diagnosed with contact allergy to fragrance ingredients, most presented with hand or facial eczema. In response to the question if and how fragrance allergy had affected their life situation, 67.5% replied that they often had to take special precautions, 47.0% replied that they were often bothered by eczema and itch, 17.1% said that they had had to take sick leave due to their fragrance contact allergy and 45.3% felt that fragrance contact allergy had significantly influenced their daily living (112).

4.5.2. Occupational restrictions

Contact allergy is known to influence severity and prognosis of hand eczema (113, 114) including risk of sick leave (111). Fragrance contact allergy is mostly of a non-occupational origin (90) related to the personal use of scented cosmetics, but may have secondary occupational consequences. This may be due to exposure to fragrance ingredients also in the work place or because hand eczema has developed. Hand eczema itself may make it impossible to remain in the trade even if protective equipment is used. In young people, fragrance allergy may limit the choice of occupations, as it will be difficult to work as a hairdresser, cosmetologist or in other occupations with a significant skin exposure to fragranced products.

4.5.3. Costs to health care/health economics

In a population based study of 3,460 individuals, contact allergy to FM I was found in 1.6%; logistic regression analyses showed that medical consultation due to cosmetic dermatitis (OR 3.37, 95% CI 1.83-6.20) and cosmetic dermatitis within the past 12 months (OR 3.53, CI 2.02-6.17) were significantly associated with sensitisation to FM I (88). Further, as mentioned above, fragrance allergy may lead to sick leave (112). No specific cost estimates for fragrance allergy exist, but the yearly total costs of contact dermatitis in Western Europe was estimated to be 5.2 billion Euro in 1997. Prices were based on the Allergy White Paper (1997) and on results of investigations and extrapolations of known data for Western Europe (115). Fragrance allergy is the second most frequent cause of contact allergy after nickel allergy and is seen in every 10th patient investigated for contact allergy. Even a modest reduction in nickel allergy has been estimated to have the value of 12 million Euro/year/million people in Denmark (Environmental Project Nr. 929, 2004; <http://www2.mst.dk/Udgiv/publications/2004/87-7614-295-7/pdf/87-7614-296-5.pdf>, last

accessed 2011-11-13). The costs are likely to differ in other countries, some with higher expenses and some with lower costs. These estimates show that the cost of contact allergy in the population may be considerable.

4.6. Allergen avoidance

Generally, “allergen avoidance” can be regarded as having two aspects: (i) primary prevention of the acquisition of contact allergy achieved by avoiding or limiting exposure of the general population, or certain parts of it, to allergens; and (ii) secondary prevention in terms of avoiding (re-)elicitation of allergic contact dermatitis in sensitised individuals.

4.6.1. Primary prevention: limiting or eliminating exposure to allergens in the population

The main aim of public health is the primary prevention of disease in populations. Allergic contact dermatitis (to fragrances) has the potential to have a significant impact on quality of life, including effects on fitness for work (chapter 4.5). Moreover, it is a common phenomenon and therefore a reduction of exposure to (fragrance) allergens must be an objective of effective Public Health measures.

Means of limiting or eliminating exposure to fragrance allergens include the following:

- *Prohibition* by regulatory measures or other means.
- *Restriction* by regulatory measures or other means of the maximum permissible concentration of a substance, or a critical component of natural mixtures, possibly according to different uses and product types, respectively.
- *Substitution* with suitable, but less or non-allergenic compounds. Substitution by a component which is chemically different, but effectively not different in terms of allergenicity or cross-reactivity, is not adequate (e.g. an ester) (chapter 5).
- *Formulating the fragrance* with the aim of limiting or eliminating those substances for which a sensitising potential has been shown. One difficulty with this approach is that sometimes no sensitisation data exist for those components of a fragrance formula which are used to replace a “known sensitiser”.
- *Deliberate avoidance* of the use of fragrances where they are not essential to the function of a finished product, but used merely to add to its appeal. Examples could include most cosmetics, topical medicaments, detergents etc., but obviously not perfumes, eau de toilette and other products used for their scent.
- *Information, e.g. labelling* so that the consumer may make an informed choice to avoid exposure to a particular ingredient.

4.6.2. Secondary prevention: avoiding re-exposure to (a) specific sensitiser(s) in clinically diagnosed individuals

In clinical dermatology, avoidance of re-exposure to an allergen is central to the care of sensitised patients. Contact sensitisation, as a latent condition, persists life-long, and therefore allergen avoidance is the only means of avoiding potentially severe and/or handicapping disease, which affects quality of life and may affect fitness for work, i.e. allergic contact dermatitis.

In this context, the valid diagnosis of sensitisation, by patch testing (32) with standardised materials, is a prerequisite of successful allergen avoidance.

In the case of fragrances, a history clearly indicative of “fragrance dermatitis” but in which patch testing with commercially available test preparations is negative, most probably reflects a shortcoming of the patch test procedure, namely, a false-negative investigation. An important cause is inadequate information on the presence of fragrance substances

present in cosmetic products (and consumer products in general). This means that patients cannot be tested for relevant substances.

A false-negative investigation can also be due to a number of other reasons: (i) non-adherence to scientific recommendations (32) or guidelines (e.g. (116)); (ii) sub-optimal patch test concentration; or (iii) use of non-oxidised material if oxidised material is the true allergen.

In an "ideal" case, from the point of view of successful patient management, the test procedure identifies all the allergen(s) to which the patient has developed contact allergy, according to the information on the culprit product(s) brought in by the patient. Such contact sensitisation is termed "clinically relevant" (65), and the need for allergen avoidance in the future is unequivocally evident in these cases. However, not infrequently, clinical relevance of an allergic patch test reaction cannot be ascertained for various reasons, which may be beyond control by the clinician (see chapter 4.4). Nevertheless, future elicitation of allergic contact dermatitis by sufficient contact with the identified "non-relevant" allergen may be expected. Hence, the patient will need to avoid the respective substance(s).

In a less "ideal" case, only part of the fragrance allergens having caused allergic contact dermatitis are identified (and can subsequently be avoided), while another part remains unidentified, for instance because it is: (i) not labelled on the product; and/or (ii) not available for routine diagnostic patch testing (special investigations such as chemical analysis of the culprit product, and break-down patch testing of its individual components, are performed rarely). Such "residual" undetermined sensitisation will hamper the success of secondary prevention of allergic contact dermatitis due to fragrances.

The above consideration raises the question for the patient of how to identify fragrance chemicals in cosmetics and other products coming into contact with the skin, such as detergents and household products, topical medicaments, products used professionally (e.g. by hairdressers, beauticians, masseurs, aromatherapists), and in other industrially used categories of products (7) (see also chapter 9). In this regard, the labelling with "perfume" or "contains fragrances" does not provide sufficient information. Moreover, such general labelling has two main disadvantages:

- It does not aid the identification of past exposure to specific agents when planning a patch test and later, when interpreting possible positive patch test results regarding clinical relevance.
- The diagnosis of allergic contact sensitisation to unidentified fragrance allergens will lead to unnecessary avoidance of other fragrance substances to which the patient is not sensitised, which are, however, included under the label "perfume".

Furthermore, the attribute "fragrance-free" may be misleading, as it merely states that no substance was added to the product to give it a scent, assuming it is used correctly at all. Nevertheless, fragrance substances used for other purposes, e.g. as preservatives, may expose the "fragrance allergic" patient to the allergen even in a "fragrance free" product (117). However, in terms of cosmetic ingredient labelling, such other uses are less problematic, as each ingredient not used as a fragrance component must be labelled. Also the use of natural products (essential oils) as preservatives must be considered in this context.

Ingredient labelling of 26 individual fragrance ingredients, identified as allergens in humans, was introduced for cosmetics in 2005. The intention was to provide a tool for clinicians for optimizing the investigation of patients with suspected fragrance allergy, as well as for fragrance allergic patients for avoiding products containing substances they have been shown to be allergic to. Both these aims are objectives of secondary prevention and seem to have been well accepted. In a study of fragrance allergic patients and their utilisation of ingredient labelling (112), most responded that they used the ingredient labelling (86.3%) and of those who used it, the majority (65.3%) found it helpful (112). Most allergic patients used the ingredient labelling (83.2%) to find out if the product was scented, while 35.6%

also looked for specific ingredients. Many (84.9%) found that a clearer labelling, e.g. easier names and a larger font size, would increase their benefit.

4.7. Conclusions

Contact allergy to fragrances is relatively common, affecting 1 to 3% of the general population, based on limited testing with eight common fragrance allergens and about 16 % of patients patch tested for suspected allergic contact dermatitis. Fragrance contact allergy is mostly non-occupational and related to the personal use of cosmetic products.

Allergic contact dermatitis can be severe and widespread, with a significant impairment of quality of life and potential consequences for fitness for work. Thus, prevention of contact sensitisation to fragrances, both in terms of primary prevention (avoiding sensitisation) and secondary prevention (avoiding relapses of allergic contact dermatitis in those already sensitised), is an important objective of public health risk management measures.

5. Activation of weak or non-sensitising substances into sensitisers - prehaptens and prohaptens

Fragrance allergens act as haptens, i.e. low molecular weight chemicals that are immunogenic only when attached to a carrier protein. However, not all sensitising fragrance chemicals are directly reactive, but require previous activation.

A prehapten is a chemical that itself is non- or low-sensitising, but that is transformed into a hapten outside the skin by simple chemical transformation (air oxidation, photoactivation) and without the requirement of specific enzymatic systems.

A prohaptent is a chemical that itself is non- or low-sensitising but that is transformed into a hapten in the skin (bioactivation) usually via enzyme catalysis.

It is not always possible to know whether a particular allergen that is not directly reactive acts as a prehapten or as a prohaptent, or both, because air oxidation and bioactivation can often give the same product (geraniol is an example).

Some chemicals might act by all three pathways. One example is geranial (an isomer of citral) which is a hapten itself with a moderate sensitisation potency, but can be activated to more potent sensitisers via air oxidation (autoxidation) thus acting as a prehapten and also via bioactivation (metabolic activation) thus acting as a prohaptent (118).

Increased understanding of the importance of activation through interaction with the environment that turns non-sensitising compounds into sensitisers has made it important to distinguish between prehaptens and prohaptens. This distinction facilitates discussions by emphasizing the differences in activation mechanisms between the two types of compounds requiring activation to become haptens. It is important to note that prehapten activation, in contrast to bioactivation, can be prevented to a certain extent by avoidance of air exposure during the handling and storage of the chemicals. This concerns the most prominent haptens formed by autoxidation i.e. the hydroperoxides. In bioactivation, hydroperoxides have not been identified as metabolites, but other allergenic oxidation products (in particular aldehydes and epoxides) have been identified as being formed by both activation routes depending on the structure of the compound. One thoroughly studied example is geraniol which forms the aldehyde geranial, epoxy-geraniol, and also epoxy-geranial via both pathways of activation (autoxidation and metabolic oxidation) (119, 120). When haptens are formed by both pathways, the impact on the sensitisation potency depends on the degree of autoxidation in relation to the amount of metabolic oxidation.

Human data on established prehaptens are presented in Table 5-1 and Table 5-2. In Table 5-1 the results from patch testing with air exposed samples of the prehaptens are given. Table 5-2 shows the results from testing with the prehaptens themselves without intended air exposure. In addition to the data given in this chapter, animal data (LLNA) on the pure prehaptens or after controlled air exposure are given in Table 8-2. Possible pro- and prehaptens are identified by SAR analyses in chapter 9.

5.1. Prehaptens

Autoxidation is a free radical chain reaction in which hydrogen atom abstraction in combination with addition of oxygen forms peroxy radicals. The reaction shows selectivity for positions where stable radicals can be formed. So far, all fragrance substances that have been investigated with regard to the influence of autoxidation on the allergenic potential, including identification of formed oxidation products, have oxidisable allylic positions that are able to form hydroperoxides and/or hydrogen peroxide as primary oxidation products upon air exposure. Once the hydroperoxides have been formed outside the skin they form specific antigens and act as skin sensitisers (121). Secondary oxidation products such as aldehydes and epoxides can also be allergenic, thus further increasing the sensitisation potency of the autoxidation mixture (122). The process of photoactivation may also play a role, but further research is required to establish whether this activation route is currently underestimated in importance due to insufficient knowledge of the true haptens in this context.

Most terpenes with oxidisable allylic positions can be expected to autoxidise on air exposure due to their inherent properties. Depending on the stability of the oxidation products that are formed, a difference in the sensitisation potency of the oxidised terpenes can be seen. Oxidation products of commonly used fragrance terpenes (limonene, linalool, geraniol, linalyl acetate) have been identified as potent sensitisers in predictive animal tests (119, 123-128) (see chapter 8). This is also demonstrated for alpha-terpinene (129) and citronellol (AT Karlberg, personal communication 2012). The oxidised fragrance terpenes limonene, linalool and linalyl acetate have been tested in consecutive dermatitis patients and give frequent allergic contact reactions (130-135). Not all oxidised fragrance substances are strong sensitisers, e.g. caryophyllene is readily oxidised but has a low sensitisation potency after autoxidation (136). This is supported by clinical studies showing oxidised caryophyllene to be a less frequent allergen compared to oxidised limonene and oxidised linalool (133). Details are given in Table 5-1 The non-oxidised compounds rarely cause allergic reactions (43-45, 67, 70, 74, 97, 137-139), for details see Table 5-2. As oxidised and non-oxidised fragrance terpenes were not patch tested simultaneously in the same patients, the results are presented in two separate tables (Table 5-1 and Table 5-2).

Oxidised fragrance terpenes with defined content of the major haptens formed after autoxidation have not been commercially available for testing in dermatology clinics. In the published clinical studies testing oxidised fragrance terpenes, the patch test preparations have been obtained specifically for the performed multicentre studies. From 2012, patch test preparations of oxidised limonene and oxidised linalool with defined content of the major allergens in the oxidation mixtures, i.e. the hydroperoxides, are commercially available.

It should be noted that activation of substances via air oxidation results in various haptens that might be the same or cross-reacting with other haptens (allergens). The main allergens after air oxidation of linalool and linalyl acetate are the hydroperoxides. If linalyl acetate is chemically hydrolysed outside the skin it can thereafter be oxidised to the same haptens as seen for linalool. A corresponding example is citronellol and citronellyl acetate. In clinical studies, concomitant reactions to oxidised linalool and oxidised linalyl acetate have been observed (140, 141). Whether these reactions depend on cross-reactivity or are due to exposure to both fragrance substances cannot be elucidated as both have an allergenic effect themselves.

Linalool and linalyl acetate are the main components of lavender oil. They autoxidise on air exposure also when present in the essential oil, and form the same oxidation products found in previous studies of the pure synthetic terpenes. Experimental sensitisation studies showed that air exposure of lavender oil increased the sensitisation potency. Patch test results in dermatitis patients showed a connection between positive reactions to oxidised linalool, linalyl acetate and lavender oil (140).

Air oxidation of prehaptens can be prevented to a certain extent by measures during handling and storage of the ingredients and final products to avoid air exposure, and/or by addition of suitable antioxidants. The autoxidation rate depends not only on the compound itself, but also on its purity. The prevention of autoxidation using antioxidants needs thorough investigation because antioxidants can exert their function by being oxidised instead of the compound that they protect and might thereby be activated to skin sensitising derivatives after oxidation, which is the case for alpha-terpinene from tea tree oil (129). Alpha-Terpinene together with its analogue gamma-terpinene has been suggested as an agent for maintaining the oxidative stability of different matrices, such as food, cosmetics and medicaments (142-144). As antioxidants are now frequently used at elevated concentrations in scented products due to a growing awareness of the problem of autoxidation, there is a risk that sensitisation caused by the antioxidants will rise. One of the most used antioxidants is butylated hydroxytoluene (BHT) which is considered a minimal risk for sensitisation in the concentrations used but nevertheless, with increased concentrations and usage, the risk of sensitisation could increase.

Due to the complexity of scented products, which are mixtures of many different fragrance substances, there are at present no published data identifying the presence of individual hydroperoxides in cosmetic products containing the above fragrance terpenes. However, clinical studies show a clear connection between contact allergy to oxidised limonene and oxidised linalool, and contact allergy to other markers of fragrance contact allergy (130-135); see Table 5-3.

Table 5-1: Contact allergic reactions to the autoxidised fragrance substances limonene, linalool, caryophyllene, myrcene and linalyl acetate in consecutive dermatitis patients.

| INCI name | CAS no | Test conc. (%) | n Positive/n tested (%) | Comments (Ref.) |
|-------------------------------------|--------------------------------------|----------------|---|-----------------|
| D-Limonene (ox.) | 5989-27-5 | 5 | 18/703 (2.6%) | § (130) |
| | | 3 | 28/1172 (1.6%) | |
| | | 2 | 3/362 (0.83%) | |
| D-Limonene (ox.) | 5989-27-5 | 3 | 63/2273 (2.8%) variation between centres: 0.3-6.5% | § (131) |
| D-Limonene (ox.) | 5989-27-5, 5989-54-8, 138-86-3 | 3 | 49/1812 (2.3%) | § (134) |
| L-Limonene (ox.) | | | 36/1812 (2.0%) | |
| D – and/or L- Limonene (ox.) | | | 63/2411 (2.6%) | |
| Linalool (ox.) | 78-70-6 | 2 | 20/1511 (1.3%) variation between centres: 0.4-2.7% | § (133) |
| Caryophyllene (ox.) | 88-44-5 | 3.9 | 2/1511 (0.1%) | |
| Myrcene (ox.) | 123-35-3 | 3 | 1/1511 (0.1%) | |
| Linalool (ox.) | 78-70-6 | 2 | 14/1693 (0.83%) | § (135) |
| | | 4 | 67/2075 (3.2%) | |
| | | 6 | 91/1725 (5.3%) | |
| | | 11 | 72/1004 (7.2%) | |
| Linalool (ox.) | 78-70-6 | 3 | 11/483 (2.3%) | (145) |
| Linalyl acetate (ox.) | 115-95-7 | 6 | 13/1217 (1.1%) | (141) |

Notes: § Bicentric or multicentre studies.
(ox.) Oxidised.

Table 5-2: Contact allergic reactions to limonene, linalool, linalyl acetate and caryophyllene in consecutive dermatitis patient. Please observe that several studies have been performed using the test substances without reporting the autoxidation status but it has been intended to be low. For precise information see the original references.

| INCI name | CAS number | Test conc. (%) | n Positive/n tested (%) | Comments (Ref.) |
|--------------------------|------------|----------------|-------------------------|-----------------|
| Limonene | 138-86-3 | 2 | 0/1200 | (137) |
| Limonene | | | 3/2396 (0.1%) | § (74) |
| DL-Limonene | | | 11/1241 (0.88%) | § (43) |
| Limonene | | | 0/320 | (44) |
| DL-Limonene | | | 3/2396 (0.1%) | § (74) |
| Linalool | 78-70-6 | 30 | 0/179 | (139) |
| | | 20 | 3/1825 (0.2%) | § (45) |
| | | 10 | 2/320 (0.6%) | (44) |
| | | 10 | 4/792 (0.5%) | (138) |
| | | 5 and 1 | 0/100 | (70) |
| Linalool, "stabilised" * | | 10 | 7/2401 (0.3%) | § (74) |
| | 10 | 2/985 (0.2%) | § (43) | |
| Linalyl acetate | 115-95-7 | 1, 5 | 0/100 | (70) |
| | | 10 | 4/1855 (0.2%) | § (67) |
| beta-Caryophyllene | 87-44-5 | 5 | 10/1606 (0.6%) | § (97) |

Notes: § Bicentric or multicentre studies.

(ox.) Oxidised.

* Stabilised: according to the manufacturer contained additional substances aimed at limiting oxidation.

Table 5-3: Concomitant reactions to fragrance markers: Fragrance Mix I and II (FM I, FM II), *Myroxylon pereire* (MP) and to colophonium (coloph.) in the baseline series in patients with positive or negative patch test reactions to oxidised fragrance substances.

| | Total number of pos. and/or neg. reactions | Pos. to FM I | | Pos. to MP | | Pos. to coloph. | | Ref. |
|--|--|--------------|-----|------------|-----|-----------------|-----|--------|
| | | n | % | n | % | n | % | |
| Reactions to ox. D- limonene and/or limonene hydroperoxide fraction | Pos.: 49 | 20 | 41 | 12 | 24 | 12 | 24 | (130)* |
| | Neg.: 2751 | 223 | 8.1 | 142 | 5.2 | 131 | 4.8 | |
| Reactions to ox. D- limonene and/or limonene hydroperoxide fraction ^a | Pos.: 60 | 22 | 37 | 11 | 18 | 13 | 22 | (132)* |
| | Neg.: 729 | 141 | 19 | 71 | 9.7 | 58 | 8 | |

| | | | | | | | | | | |
|---|---|---------------------|----------|----------------------|----------|-------------------|----------|------------------------|----------|--------|
| Reactions to ox. D- limonene and/or ox. L- limonene ^a | Pos. to ox. D- limonene: 41 | 14 | 34 | 11 | 27 | 11 | 27 | (134)* | | |
| | Neg. to ox. D- limonene: 1771 | 113 | 6.4 | 91 | 5.1 | 62 | 3.5 | | | |
| | Pos. to ox. L- limonene: 36 | 11 | 31 | 12 | 33 | 9 | 25 | | | |
| | Neg. to ox. L- limonene: 1776 | 116 | 6.5 | 80 | 4.5 | 64 | 3.6 | | | |
| Reactions to any of ox. linalool, myrcene, caryophyllene | Pos. to any of the tested ox. subst.: 31 | 12 | 39 | 6 | 31 | 12 | 39 | (133)* | | |
| | Neg. to any of the tested ox. subst: 1480 | 93 | 6 | 63 | 4 | 46 | 3 | | | |
| | | Pos. to FM I | | Pos. to FM II | | Pos. to MP | | Pos. to coloph. | | |
| | | n | % | n | % | n | % | n | % | |
| Reactions to ox. linalool | Pos. at test conc. 4%: 30 | 8 | 26.7 | 5 | 16.7 | 10 | 33.3 | 5 | 16.7 | (135)* |
| | Pos. at test conc. 6%: 55 | 12 | 21.8 | 8 | 14.5 | 11 | 20 | 8 | 14.5 | |
| | Pos. at test conc. 11%: 72 | 14 | 19.4 | 9 | 12.5 | 14 | 19.4 | 9 | 12.5 | |
| | Total pos. at any test conc: 75/1004 | n.g. | | n.g. | . | n.g. | | n.g. | . | |
| | Total neg. at any test conc: 929/1004 | 56 | 6.0 | 29 | 3.1 | 45 | 4.8 | 24 | 2.6 | |

Notes: * Bicentric or multicentre studies.
n.g. Not given.
(ox.) Oxidised.

5.2. Prohaptens

Compounds that are bioactivated in the skin and thereby form haptens are referred to as prohaptens. The human skin expresses enzyme systems that are able to metabolise xenobiotics (146), modifying their chemical structure to increase hydrophilicity and allow elimination from the body. Xenobiotic metabolism can be divided into two phases: phase I and phase II. Phase I transformations are known as activation or functionalisation reactions, which normally introduce or unmask hydrophilic functional groups. If the metabolites are sufficiently polar at this point they will be eliminated. However, many phase I products have to undergo subsequent phase II transformations, i.e. conjugation to make them sufficiently water soluble to be eliminated. Although the purpose of xenobiotic metabolism is detoxification, it can also convert relatively harmless compounds into reactive species. Cutaneous enzymes that catalyse phase I transformations include the cytochrome P450 mixed-function oxidase system, alcohol and aldehyde dehydrogenases, monoamine oxidases, flavin-containing monooxygenases and hydrolytic enzymes. Acyltransferases, glutathione S-transferases, UDP-glucuronosyltransferases and sulfotransferases are

examples of phase II enzymes that have been shown to be present in human skin (146). These enzymes are known to catalyse both activating and deactivating biotransformations (147), but the influence of the reactions on the allergenic activity of skin sensitisers has not been studied in detail.

Skin sensitising prohaptens can be recognised and grouped into chemical classes based on knowledge of xenobiotic bioactivation reactions, clinical observations and/or *in vivo* and *in vitro* studies of sensitisation potential and chemical reactivity. Few mechanistic investigations of prohaptens have so far been published. Investigations that are important for the bioactivation of fragrance substances are studies on alkenes, e.g. alpha-terpinene (148-150), the allylic primary alcohols geraniol (120) cinnamyl alcohol (151-155), eugenol and isoeugenol (156).

In order to be able to predict the sensitisation potency of prohaptens, steps of bioactivation have to be included in the predictive tests where intrinsic bioactivating systems are lacking. So far, no such predictive non-animal methods have been developed that take account of this.

When bioactivation occurs, the risk of cross-reactivity also needs to be considered. Cross-reactivity between certain aldehydes and their corresponding alcohols, e.g. cinnamal - cinnamyl alcohol and geraniol - geraniol, due to the metabolic oxidation of the alcohols to the aldehydes in the skin is demonstrated (120, 151-155).

When using derivatives of a fragrance substance, it must be taken into account that the derivative could be metabolically transformed in the skin into the parent or cross-reacting compounds. A prominent example of such bioactivation is the hydrolysis of esters by esterases to the corresponding original alcohols. The metabolic product obtained can act as a hapten or a prohaptent in exactly the same way as the non-esterified parent compound.

Isoeugenol and its derivatives are an important example for this mechanism from which general conclusions may be drawn. As only the use of isoeugenol in fragranced products needs to be indicated on the ingredients list, the additional exposure to isoeugenol through its derivatives should also be taken into account. In a study it was shown that several EDP/EDT/aftershave lotions contained high levels of isoeugenyl acetate and isoeugenol methyl ether (Table 5-4) (157). Isoeugenyl acetate will be hydrolysed by esterases in the skin to generate isoeugenol. The situation may be similar for eugenyl acetate and geranyl acetate, which might be used in fragrance formulations instead of eugenol and geraniol, respectively. Moreover, such derivatives will contribute to exceeding any established 'acceptable dose/area level' of the parent compound, i.e., yield unduly high concentrations on the skin.

Table 5-4: Mean and median content of isoeugenol and its derivatives in the 29 perfume products.

| Fragrance compound INCI Name | Products containing the fragrance | | Content (ppm) | | | |
|---------------------------------|--------------------------------------|----|---------------|------|-------|--------|
| | No. | % | Range | Mean | SD | Median |
| Isoeugenol | 16 | 55 | 27-203 | 71 | 54 | 45 |
| Isoeugenyl acetate | 10 | 34 | 20-4689 | 985 | 1570 | 166 |
| Isoeugenyl methyl ether | 13 | 45 | 65-1755 | 360 | 442.3 | 222 |

5.3. Conclusions

- Many fragrance substances can act as prehapten or prohapten, forming allergens which are more potent than the parent substance by abiotic and/or metabolic activation. Activation can thus increase the risk of sensitisation.
- Fragrance substances of clinical importance known to be prehapten and to form sensitising compounds by air oxidation include limonene, linalool, and linalyl acetate.
- Fragrance substances of clinical importance known to be prohapten and to form sensitising compounds by metabolic transformation include cinnamyl alcohol, eugenol, isoeugenol and isoeugenol acetate.
- Fragrance substances of clinical importance with published data known to be both prehapten and prohapten and to form sensitising compounds by air oxidation (prehapten) and by metabolic transformation include geraniol and alpha-terpinene.
- A fragrance substance that sensitises without activation, but forms more potent sensitising compounds by air oxidation and also by metabolic transformation is, as one example, geranial (one isomer of citral).
- In the case of prehapten, it is possible to prevent activation outside the body to a certain extent by different measures, e.g. prevention of air exposure during handling and storage of the ingredients and the final product, and by the addition of suitable antioxidants. When antioxidants are used, care should be taken that they will not be activated themselves and thereby form new sensitisers.

It should be noted that the possibility to reduce the sensitisation potency by preventing air oxidation is also important for a direct acting hapten or prohapten, if a further activation by air oxidation to more allergenic compounds has been shown.

- In the case of prohapten, the possibility to become activated is inherent to the molecule and activation cannot be avoided by extrinsic measures. Activation processes increase the risk for cross-reactivity between fragrance substances. Cross-reactivity has been shown for certain alcohols and their corresponding aldehydes, i.e. between geraniol and geranial (citral) and between cinnamyl alcohol and cinnamal.

Cross-reactivity is also expected between ester derivatives and their parent alcohols, as the esters will be hydrolysed by esterases in the skin. Esters of important contact allergens that can be activated by hydrolysis in the skin are isoeugenol acetate, eugenyl acetate and geranyl acetate all of which are known to be used as fragrance ingredients.

- Further experimental and clinical research in the area of abiotic and/or metabolic activation of fragrance substances is clearly needed to increase the safety for the consumer. Compounds suspected to act as prehapten and/or prohapten should be considered as allergens, unless it could be demonstrated that they do not become activated by one of the described pathways.

6. Retrieval of evidence and classification of fragrance substances

For a systematic review, a structured approach of identifying, grading and aggregating available information should be used. Regarding the classification of substances as allergens, a number of approaches have been suggested (158-160). The categorisation of skin sensitisers according to sensitising potency has also been proposed (161, 162). For this opinion, these discussions were extended to reconcile different perspectives and to arrive at a strategy that is both consistent and applicable in practice.

By default, positive human evidence (clinical data) overrides negative results obtained in animals. This implies that the observation of a sufficient number of positive clinical cases is more important than potency information derived from animal experiments (LLNA).

Cosmetovigilance information based on consumer complaints only is of limited value in the evaluation of sensitisation risk associated with cosmetic allergens, including fragrances, as it does not identify specific causative substances, and likely to severely under-estimate the frequency of contact dermatitis. An exception is the combination with qualified diagnostic work-up, as in the French REVIDAL/GERDA system (299); however, such data are generally published, thus publicly available, and considered in the present opinion.

6.1. Retrieval of evidence

A systematic search strategy was employed for the retrieval of clinical data, as outlined below. Experimental data are often not published hence the exact definition of the scope considered for the review is necessary and is given below. Additional LLNA data were reviewed, if identified by the search strategy, e.g. in chapter 8.1.2 and, as "additional information", in Annex I of this opinion. This supplemental evidence was, however, not considered for the final categorisation in Table 13-2.

6.1.1. Search strategy for clinical data

Method of literature search:

1. Manual search of the issues of the journal "Contact Dermatitis" (for the 26 "annex substances", which were re-evaluated in the present opinion, starting 1999) up to October 2010, identifying all studies with fragrance substances.
2. PubMed search of CAS numbers identified in the previous opinion, reviews and already identified clinical studies, respectively, and manual screening of identified publications (narrowed for the last 10 years for the 26 "annex substances"), if necessary narrowing the search results by adding "dermatitis" or "allergy". For example, for citral: 5392-40-5 AND (dermatitis or allergy), translated into
 "5392-40-5"[EC/RN Number] AND
 (
 ("dermatitis"[MeSH Terms] OR "dermatitis"[All Fields])
 OR
 ("hypersensitivity"[MeSH Terms] OR "hypersensitivity"[All Fields] OR "allergy"[All Fields] OR "allergy and immunology"[MeSH Terms] OR ("allergy"[All Fields] AND "immunology"[All Fields]) OR "allergy and immunology"[All Fields])
)
3. Manual search of all RIFM reviews published in supplement issues of "Food and Chemical Toxicology"² in the past 20 years. In case of the least evidence on human sensitisation the substances were preliminarily selected and further research initiated.

² Food and Chemical Toxicology, Elsevier Ltd. <http://www.sciencedirect.com/science/journal/02786915>.

4. Consideration of the most important ("top 100") fragrance compounds in terms of volumes used (disregarding functional additives such as solvents) as supplied by the International Fragrance Association IFRA (personal communication 2010).
5. Consideration of fragrance compounds ranking 101 to 200 on the list of use volumes, if they were self-classified by manufacturers as skin sensitisers (R 43).

For the present systematic overview of available clinical data, only original studies were considered, as only these provide direct evidence, while other reviews, partly being based on the same original reports, only served to identify additional literature. In contrast, selected reviews, guidelines and similar publications were used as basis for methodological approaches (e.g., in section 11).

6.1.2. Collection of experimental (LLNA) data

The SCCS requested the International Fragrance Association (IFRA) to submit data on animal tests performed with fragrance substances, to be presented in a structured format. In response, industry submitted first a poster (163) and later a report consisting of LLNA protocol summaries on the 59 fragrance substances in the poster (164). No guinea pig studies were submitted. The SCCS has reviewed and analysed the report and the publications quoted in the report. A summary is given in chapter 8 and full data are given in Annex II. EC3 values on some additional fragrance substances in two published reviews (165, 166) have also been considered. Additional EC3 values may be available in the scientific literature and there may also be other unpublished data.

6.2. Grading of evidence

Assembled evidence has to be graded in two steps: (i) the quality of each single study, and (ii) the strength of evidence underlying the eventual classification as an allergen. Generally, studies (published or not) which are eligible for consideration will contribute to the final overall judgement to different degrees.

- Positive human data, if sufficiently demonstrated (point (i) below), will always over rule experimental (animal), *in vitro* or *in silico* data of similar internal validity, as they provide direct evidence on allergenicity in humans.
- Small study groups will contribute less precise information than larger studies of otherwise similar quality. As a minimum requirement, the size of the study groups and the numbers of events must be stated in the reports.

The following subsections will address special aspects of clinical and experimental studies, respectively.

6.2.1. Quality of a clinical study

Two major types of clinical studies must be distinguished because they provide a different scope of information:

- (i) Case reports or small case series, focusing on patients with positive (test) reactions to the target substance, sometimes including a set of non-exposed, possibly non-diseased "control patients"; these should present a concise summary of all relevant aspects of the patient's history, diagnostic procedures and possibly further outcomes.
- (ii) Clinical series in which results of a group of patients patch tested with the target substance, often combined with other substances, are presented. In the latter type of report, usually only a minority of patients tested show a positive reaction to the test substance. This implies that the majority of patients can be used to illustrate the proportion of irritant, doubtful and negative reactions. The degree of detail on the patients' histories is usually limited in such studies, compared to case reports.

Some of the basic quality criteria in clinical patch testing which should be considered are:

- Adherence to international patch test guidelines (32, 96).
- Material(s) tested should be characterised.
- Total number of patients tested must be given.
- Patient selection should be described.
- Relevance may be demonstrated either on a case-by-case basis, following pertinent guidelines, or in terms of a significant epidemiological association between sensitisation and exposure or valid markers of exposure.

Concerning relevance, it must be noted that while clinical relevance can provide important information (see 4.4.1), it is ideally based on comprehensive knowledge of prior exposures. Since the implementation of labelling 26 fragrances, previous exposure to these can often be ascertained in the assessment of relevance of a positive patch test reaction (44). However, exposure to substances not listed on a product ingredient label is obscure, except in very rare cases where elaborate diagnostics and chemical analyses are feasible (e.g. (167)). Thus, a lack of information on relevance (reported in studies) does not invalidate the impact of diagnosed contact sensitisation.

6.2.2. Quality of an experimental study

International guidelines such as the pertinent OECD guidelines for testing sensitisation have been developed and adopted. Experimental studies following these guidelines are considered as valid. However, a vast number of non-guideline studies are available and should be assessed on a case-by-case basis.

6.2.3. Quality of "other" evidence

Supporting evidence besides human and animal (experimental) data comprises *in vitro* test systems, *in chemico* experiments and structure activity relationships (SARs).

SAR analysis has at present no formal regulatory validation for skin sensitisation, nevertheless it may provide useful indicative information on sensitising potential when no or limited clinical or animal data are available.

SAR studies must consider a possible formation of haptens (allergens) from compounds able to act as prehapten by, e.g. autoxidation outside the body as well as metabolic activation in the skin of compounds able to act as prohapten (122, 168).

6.3. Aggregating evidence for a final conclusion

The criteria listed below are followed as a flow chart to arrive at a conclusion. This implies that if classification into one category is achieved, subsequent categories need not be considered. Based on the above criteria, fragrance substances were selected to be included in the present opinion if classified in one of the categories defined below.

6.3.1. Established contact allergen in humans

To qualify as an *established contact allergen*, the SCCS considers that *at least one* of the following two criteria must be met:

- At least two clinical series fulfilling the quality criteria from two different centres with cases of sensitisation, or at least three separate clinical series from different centres if a study, or studies, do not meet all quality criteria. (→ *sufficient human evidence present*)
or
- Case reports from at least two independent centres describing more than two

patients altogether in whom clinically relevant contact sensitisation had unequivocally been proven (→ *sufficient human evidence present*)
or

- At least one clinical series fulfilling the quality criteria, together with at least one case report of clinically relevant contact sensitisation (→ *sufficient human evidence present*);
or
- Experimentally induced sensitisation (e.g. unequivocally positive human maximisation tests/repeated insult patch test)³ (→ *sufficient human evidence present*).

6.3.2. Established contact allergen in animals

To qualify as an *established contact allergen*, the following criterion must be met:

- At least one positive animal study carried out according to accepted guidelines, providing evidence of a sensitisation potential (→ *sufficient animal evidence present*).

6.3.3. Likely contact allergen, if human, animal and other evidence is considered

To qualify as an *likely contact allergen*, if classification as “established ...” is not applicable, *at least two* of the following criteria must be met:

- Individual cases of allergic patch test reactions not fulfilling the requirements for sufficient evidence (→ *limited human evidence present*)
or
- At least one positive non-guideline animal study, which should be evaluated on a case-by-case basis (→ *limited animal evidence present*)
or
- Other evidence, e.g. results from *in chemico* experiments or *in vitro* tests or from structure-activity considerations based on sufficiently valid results for closely related compounds (→ *other evidence present*).

6.3.4. Possible contact allergen, if human, animal and other evidence is considered

To qualify as a *possible contact allergen*, if classification as “established ...” or as “likely ...” contact allergen is not applicable, *at least one* of the following criteria must be met:

- Individual cases of allergic patch test reactions not fulfilling the requirements for sufficient evidence (→ *limited human evidence present*)
or
- At least one positive non-guideline animal study, which should be evaluated on a case-by-case basis (→ *limited animal evidence present*)
or
- Other evidence, e.g. results from *in chemico* experiments or *in vitro* tests or from structure-activity considerations based on sufficiently valid results for closely related compounds (→ *other evidence present*).

³ It should be noted that the SCCS considers such tests unethical (169. SCCP. Opinion concerning the predictive testing of potentially cutaneous sensitising cosmetic ingredients or mixtures of ingredients adopted by the SCCNFP during the 11th plenary session of 17 February 2000. 2000:).

6.4. Conclusions

The present opinion includes (i) a well-defined search strategy for retrieving pertinent evidence; (ii) a definition of criteria used to evaluate available evidence; and, finally (iii) a set of rules to categorise the substances with regard to the relevant toxicological endpoint, i.e. sensitisation in man, based on the evidence.

7. Reported fragrance allergens from the clinical perspective

In this chapter, clinical evidence regarding sensitisation to individual fragrance chemicals and to natural extracts (essential oils) is tabulated. In this report “single chemicals” refers to chemicals of natural or synthetic origin whose chemical identity is fully known. The term “natural extracts” refers to plant or animal derived mixtures of natural chemicals, for example lavender oil, whose composition may be variable and may or may not have been fully or partly established. Full information, including possible synonyms, structural formulas (in the case of single chemicals only), a short summary of available evidence and further information, e.g. on regulatory status, is presented in Annex I.

7.1. Tabular summary of evaluated individual fragrance chemicals

Regarding nomenclature, INCI names are used wherever possible. If an INCI name is not available, the perfuming name as listed by CosIng is used. Detailed information on the publications identified and considered for this report can be found in Annex I. Several substances are currently banned from the use in cosmetic products by Annex II of the Cosmetics Directive, based on concerns regarding one or more toxicological endpoints. While available clinical evidence regarding this set of substances is listed in Annex I, these substances have not been further evaluated and are thus not included in this chapter.

In this section, a tabular overview on the classification of substances considered is presented in four tables listing:

1. Established contact allergens in humans (→ *sufficient human evidence present*).
2. Substances with positive human data, which are, however, not sufficient to categorise as “established contact allergen in humans” (→ *limited human evidence present*).
3. Substances with negative human data, i.e. patch tests of patients with suspected contact allergy to fragrance ingredients which yielded negative results.
4. Substances eligible for inclusion (see beginning of chapter 6) for which no human data are available.

A critical point in understanding this scheme is that there is publication bias in reporting allergens. This is due to the fact that once a substance has been reported and accepted as a contact allergen in humans, further reports are less likely to be published unless they are part of an epidemiological survey or when there is a novel source of exposure. Moreover, the number of patients displaying positive test reactions obviously not only depends on the underlying prevalence of sensitisation, but also on how often a substance is patch tested. This implies that inclusion of an allergen or allergen mixture in the baseline patch test series (as for Fragrance Mix I and II, *Myroxylon pereirae* and HICC, and partly also other substances/mixtures) will yield the maximum possible number of cases. In contrast, patch testing in “special” series, e.g. as a break-down of single constituents of the respective mix in case of a positive reaction to the latter, or with application only in the case of strongly suspected fragrance intolerance, will mostly result in higher relative numbers than testing the same compound consecutively, but also in lower absolute numbers.

In Table 7-1, the single substances are listed with a semi-quantification of their impact which were categorised as established contact allergens in humans according to the criteria given in chapter 6.3.

Established contact allergens in humans, according to the criteria outlined in chapter 6.3.1, were categorised according to the number of patients reacting positively and to the number of patients tested, based on the publications considered (see annex I for references). The following categories were used:

Opinion on fragrance allergens in cosmetic products

| | |
|------|---|
| + | Up to 10 positive test reactions reported |
| ++ | 11 to 100 |
| +++ | 101 to 1000 |
| ++++ | > 1000 |

If a test allergen has been tested in less than 1,000 patients, "r.t." (rarely tested) is added in the following tables. For this categorisation, absolute numbers of cases of sensitisation, and not the relative frequency of positive patch tests, were used, because relative frequencies depend heavily on the selection of patients for patch testing. Thereby, an important allergen tested routinely, in the baseline series, may yield 1 to 2% positive reactions (usually in several thousand patients), while an allergen tested in a selective fashion (in much fewer patients) may yield an even higher relative frequency. Moreover, case reports/series cannot be interpreted in terms of relative frequencies. The calculation of absolute numbers was based on all available literature, as detailed in the annex I to this opinion, i.e., regarding the 26 substances already listed in Annex III to the Cosmetics Directive includes data already evaluated in the previous opinion.

Table 7-1: Established contact allergens in humans (summary of evaluation as detailed in chapter 6.3). More detailed information forming the basis of this evaluation can be found in Annex I of this opinion.

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Comment: see text |
|---|----------------------------------|--------------------------|
| ACETYLCEDRENE | 32388-55-9 | + |
| AMYL CINNAMAL | 122-40-7 | ++ |
| AMYL CINNAMYL ALCOHOL | 101-85-9 | ++ |
| AMYL SALICYLATE | 2050-08-0 | + |
| trans-ANETHOLE | 4180-23-8 | + (r.t.) |
| ANISYL ALCOHOL | 105-13-5 | + |
| BENZALDEHYDE | 100-52-7 | + |
| BENZYL ALCOHOL | 100-51-6 | ++ |
| BENZYL BENZOATE | 120-51-4 | ++ |
| BENZYL CINNAMATE | 103-41-3 | ++ |
| BENZYL SALICYLATE | 118-58-1 | ++ |
| BUTYLPHENYL METHYLPROPIONAL (Lilial®) | 80-54-6 | ++ |
| CAMPHOR | 76-22-2 / 464-49-3 | + (r.t.) |
| beta-CARYOPHYLLENE (ox.) | 87-44-5 | Non-ox.: +, ox.: + |
| CARVONE | 99-49-0 / 6485-40-1 / 2244-16-8 | + (r.t.) |
| CINNAMAL | 104-55-2 | +++ |
| CINNAMYL ALCOHOL | 104-54-1 | +++ |
| CITRAL | 5392-40-5 | +++ |
| CITRONELLOL | 106-22-9 / 1117-61-9 / 7540-51-4 | ++ |
| COUMARIN | 91-64-5 | +++ |
| (DAMASCENONE) | 23696-85-7 | + (r.t.) |

Opinion on fragrance allergens in cosmetic products

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Comment: see text |
|---|------------------------------------|----------------------------|
| ROSE KETONE-4 | | |
| alpha-DAMASCONE (TMCHB) [#] | 43052-87-5 / 23726-94-5 | ++ |
| cis-beta-DAMASCONE [#] | 23726-92-3 | + |
| delta-DAMASCONE [#] | 57378-68-4 | + |
| DIMETHYLBENZYL CARBINYL ACETATE (DMBCA) | 151-05-3 | + |
| EUGENOL | 97-53-0 | +++ |
| FARNESOL | 4602-84-0 | +++ |
| GERANIOL | 106-24-1 | +++ |
| HEXADECANOLACTONE | 109-29-5 | + (r.t.) |
| HEXAMETHYLINDANOPYRAN | 1222-05-5 | ++ |
| HEXYL CINNAMAL | 101-86-0 | ++ |
| HYDROXYISOHEXYL 3-CYCLOHEXENE CARBOXALDEHYDE (HICC) | 31906-04-4 / 51414-25-6 | ++++ |
| HYDROXYCITRONELLAL | 107-75-5 | +++ |
| ISOEUGENOL | 97-54-1 | +++ |
| alpha-ISOMETHYL IONONE | 127-51-5 | ++ |
| (DL)-LIMONENE | 138-86-3 | ++ (non-ox.); +++ (ox.) |
| LINALOOL | 78-70-6 | ++ (non-ox.) +++ (ox.) |
| LINALYL ACETATE | 115-95-7 | + |
| MENTHOL | 1490-04-6 / 89-78-1 / 2216-51-5 | ++ |
| 6-METHYL COUMARIN [#] | 92-48-8 | ++ (photo-allergy) |
| METHYL 2-OCTYNOATE | 111-12-6 | ++ |
| METHYL SALICYLATE | 119-36-8 | + |
| 3-METHYL-5-(2,2,3-TRIMETHYL-3-CYCLOPENTENYL)PENT-4-EN-2-OL | 67801-20-1 | ++ (r.t.) |
| alpha-PINENE and beta-PINENE | 80-56-8 and 127-91-3, resp. | ++ |
| PROPYLIDENE PHTHALIDE | 17369-59-4 | + (r.t.) |
| SALICYLALDEHYDE | 90-02-8 | ++ |

[#] 76/768/EEC Annex III, part 1

[#] 76/768/EEC Annex III, part 1

Opinion on fragrance allergens in cosmetic products

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Comment: see text |
|---|--|--------------------------|
| alpha-SANTALOL and beta-SANTALOL | 115-71-9 and 77-42-9, resp. | ++ |
| SCLAREOL | 515-03-7 | + |
| TERPINEOL (mixture of isomers) | 8000-41-7 | + |
| alpha-TERPINEOL | 10482-56-1 / 98-55-5 | |
| Terpinolene | 586-62-9 | ++ |
| TETRAMETHYL ACETYLOCTAHYDRONAPHTHALENES | 54464-57-2 / 54464-59-4 / 68155-66-8 / 68155-67-9 | + |
| TRIMETHYL-BENZENEPROPANOL (Majantol) | 103694-68-4 | ++ |
| VANILLIN | 121-33-5 | ++ |

Those substances which were categorised as +++ or more, i.e. those with the most reported cases, were also the top ranking substances in large series of patients tested with the 26 labelled fragrance ingredients ((44, 74) and additionally (170)). Geraniol is an exception, as it was all negative in the Danish study (170), but was still among the top ten in the Dutch and German studies (44, 74), with prevalences of 0.5%-0.6% positives. Geraniol has, in addition, caused many cases of contact allergy in other areas of Europe (49).

The use of absolute numbers allows the pooling of studies with different selection criteria. Limonene and linalool were not tested in their oxidized forms in the three studies (44, 74, 170) and would not have been identified, if only these publications had been the basis of assessment.

It should be noted that oxidised fragrance terpenes with defined content of the major haptens formed after autoxidation have not been commercially available for testing in dermatology clinics. In the published clinical studies testing oxidised fragrance terpenes, the patch test preparations have been obtained specifically for the performed multicentre studies. From 2012, patch test preparations of oxidised limonene and oxidised linalool with defined content of the major allergens in the oxidation mixtures, i.e. the hydroperoxides, are commercially available (see also chapter 5).

Table 7-2 lists those substances which gave rise to a few reported cases of contact sensitisation only, or where results have been reported from just one clinical department. Thus, the level of evidence concerning human data must be regarded as *limited*, according to the definitions given in chapter 6.3.

Opinion on fragrance allergens in cosmetic products

Table 7-2: Fragrance substances with positive human data, which are, however, not sufficient to categorise as “established contact allergen in humans”. More detailed information forming the basis of this evaluation can be found in Annex I of this opinion.

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Comment | Ref. |
|---|-------------------|--|--------------------------|
| AMBRETTOLIDE | 7779-50-2 | 3.4% positive reactions in 178 patients | (171) |
| CARVACROL | 499-75-2 | 2 of 28 patients | (Meynadier, after (172)) |
| CUMINALDEHYDE | 122-03-2 | 3 of 179 patients positive | (139) |
| CYCLOHEXYL ACETATE | 622-45-7 | 0.5% positive of 218 selected patients | (173) |
| CYCLOPENTADECANONE | 502-72-7 | 3 of 178 patients positive | (171) |
| trans-trans-delta-DAMASCONE | 71048-82-3 | 1 positive HRIPT (2/15 with 1%) | (174) |
| 2,3-DIHYDRO-2,2,6-TRIMETHYLBENZALDEHYDE | 116-26-7 | 1 positive HRIPT (5 of 53) | (175). |
| DIMETHYLTETRAHYDRO BENZALDEHYDE | 68737-61-1 | 2.3% positive reactions isomer mixture in 178 patients | (171) |
| ETHYLENE DODECANEDIOATE | 54982-83-1 | 2 / 218 positive PT reactions | (173) |
| ETHYL VANILLIN | 121-32-4 | 1 occupational case | (176) |
| HELIOTROPINE | 120-57-0 | 6 / 1606 consecutive patients positive | (97) |
| HYDROXYCITRONELLOL | 107-74-4 | 6.0% positive PT reactions in 218 patients | (173) |
| ISOAMYL SALICYLATE | 87-20-7 | 1 positive in 179 patients, possibly “excited back syndrome” 0 / 95 in another study with <= 1/10 of above test conc. | (139) (70) |
| ISOLONGIFOLENEKETONE | 33407-62-4 | 1 / 178 patients | (171) |
| METHOXYCITRONELLAL | 3613-30-7 | Positive PT data of unknown validity by Nakayama et al. in 22/137 patients. | (177) |
| METHOXYTRIMETHYLHEPTANOL | 41890-92-0 | 0.9% positive PT | (173) |
| METHYL p-ANISATE | 121-98-2 | 1 / 182 patients positive | (178) |
| METHYL CINNAMATE | 103-26-4 | 6 / 142 patients positive | (179) |
| METHYL DIHYDROJASMONATE | 24851-98-7 | 3 / 1606 patients positive 0 / 100 | (97) (70) |
| METHYLIONANTHEME | 55599-63-8 | 1 case | (180) |
| 5-METHYL-alpha-IONONE | 79-69-6 | 5 / 1606 | (97) |
| METHYL OCTINE CARBONATE | 111-80-8 | 1 case | (181) |

Opinion on fragrance allergens in cosmetic products

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Comment | Ref. |
|---|-------------------|--|----------------|
| MYRCENE | 123-35-3 | 1 / 1511 positive to oxidized myrcene | (133) |
| MYRTENOL | 515-00-4 | 2 HRIPTs with 1 pos. each | (182) |
| NEROL | 106-25-2 | 6.0% positive | (173) |
| Nerolidol (isomer not specified) | 7212-44-4 | Few, unconfirmed pos. cases according to RIFM review | (183) |
| NOPYL ACETATE | 128-51-8 | 2 / 179 positive, possibly "excited back syndrome" | (139) |
| PHENETHYL ALCOHOL | 60-12-8 | 1 / 179; 0 / 100 | (139) (70) |
| PHENYLACETALDEHYDE | 122-78-1 | 1.1% of 182 positive. 1 case | (178) (184) |
| PHENYLPROPANOL | 122-97-4 | 2 / 218 | (173) |
| PHYTOL | 150-86-7 | 1 case in human max. test | (185) |
| RHODINOL | 6812-78-8 | Several pos. HRIPTs, clinical data of uncertain validity | (186) |
| trans-ROSE KETONE-5 | 39872-57-6 | 2 / 22 pos. HRIPT | (187) |

For a number of substances negative patch tests results were obtained, usually in rather small patient samples (max. 313 patients). For some of these substances exposure is substantial, according to data submitted from IFRA. It should be noted that a negative result does not rule out a notable sensitisation prevalence, as the study size has to be larger than, e.g. n=298 to yield a 95% CI which excludes a prevalence of 1% and larger than n=597 to exclude a prevalence of 0.5%.

Opinion on fragrance allergens in cosmetic products

Table 7-3: Fragrance substances with negative human data, i.e. patch tests of patients with suspected contact allergy to fragrance ingredients which yielded negative results.

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Results / Comment | Ref. |
|---|-------------------|--|-------------|
| 6-ACETYL-1,1,2,4,4,7-HEXAMETHYLTETRALINE | 21145-77-7 | 0 / 313 consecutive patients in 2 centres | (70) |
| AMYL CYCLOPENTANONE | 4819-67-4 | 0 / 178 | (171) |
| BENZYL ACETATE | 140-11-4 | 0 / 100 consecutive patients in 1 centre observed | (70) |
| 2-TERT-BUTYL CYCLOHEXYL ACETATE | 88-41-5 | 0 / 313 consecutive patients in 2 centres | (70) |
| 4-tert.-Butylcyclohexyl acetate | 32210-23-4 | 0 / 107 consecutive patients in 1 centre observed | (70) |
| 6-ETHYLIDENEOCTAHYDRO-5,8-METHANO-2H-BENZO-1-PYRAN | 93939-86-7 | 0 / 178 | (171) |
| 3 α ,4,5,6,7,7 α -HEXAHYDRO-4,7-METHANO-1H-INDEN-5(OR 6)-YL ACETATE | 54830-99-8 | 0 / 313 consecutive patients in 2 centres | (70) |
| HEXYL SALICYLATE | 6259-76-3 | 0 / 218 "top 100" substance and classified as R43 | (173) |
| HIBISCOLIDE | 6707-60-4 | 0 / 178 | (171) |
| alpha-IONONE | 127-41-3 | 0 / 205 | (70) |
| beta-IONONE | 79-77-6 | 0 / 205 "top 100" substance | (70) |
| ISOBORNYL ACETATE | 125-12-2 | 0 / 107 "top 100" substance | (70) |
| METHYL ANTHRANILATE | 134-20-3 | 0 / 91 "top 100" substance | (188) |
| METHYL IONONE (mixture of isomers) | 1335-46-2 | 0 / 100 "top 100" substance | (70) |
| OXALIDE | 1725-01-5 | 0 / 178 | (171) |
| TERPINEOL ACETATE (Isomer mixture) | 8007-35-0 | 0 / 106 "top 100" substance | (70) |
| alpha-TERPINYL ACETATE | 80-26-2 | 0 / 179 | (139) |
| TRIMETHYL-PROPYLCYCLOHEXANEPROPANOL | 70788-30-6 | 0 / 178 | (171) |

For yet another subset of substances, no human data were publicly available. However, exposure to these substances is important as they are used in high volumes (this being the sole criterion for inclusion in this list) and, therefore their hazard with regard to contact sensitisation should be examined.

Table 7-4: Fragrance substances lacking human data and used in high volumes according to industry information.

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number |
|---|-------------------|
| ANISALDEHYDE | 123-11-5 |
| BENZYL ACETONE | 2550-26-7 |
| p-tert. -Butyldihydrocinnamaldehyde | 18127-01-0 |
| CITRONELLYL NITRILE | 51566-62-2 |
| CYCLAMEN ALDEHYDE | 103-95-7 |
| alpha-CYCLOHEXYLIDENE BENZENEACETONITRILE | 10461-98-0 |
| DECANAL | 112-31-2 |
| DIHYDROMYRCENOL | 18479-58-8 |
| 2,4-DIMETHYL-3-CYCLOHEXEN-1-CARBOXALDEHYDE | 68039-49-6 |
| 3,7-DIMETHYL-1,6-NONADIEN-3-OL | 10339-55-6 |
| DIPHENYL ETHER | 101-84-8 |
| ETHYL 2-METHYLBUTYRATE | 7452-79-1 |
| 2-ETHYL-4-(2,2,3-TRIMETHYL-3-CYCLOPENTEN-1-YL)-2-BUTEN-1-OL | 28219-61-6 |
| ETHYLENE BRASSYLATE | 105-95-3 |
| EUCALYPTOL | 470-82-6 |
| GERANYL ACETATE | 105-87-3 |
| HEXAHYDRO-METHANOINDENYL PROPIONATE | 68912-13-0 |
| HEXYL ACETATE | 142-92-7 |
| IONONE isomeric mixture | 8013-90-9 |
| ISOAMYL ACETATE | 123-92-2 |
| ISOBERGAMATE # | 68683-20-5 |
| Longifolene | 475-20-7 |
| METHYLENEDIOXYPHENYL METHYLPROPANAL | 1205-17-0 |
| METHYLBENZYL ACETATE | 93-92-5 |
| METHYL DECENOL | 81782-77-6 |
| METHYL beta-NAPHTHYL ETHER | 93-04-9 |
| METHYLUNDECANAL | 110-41-8 |
| OXACYCLOHEXADECENONE | 34902-57-3 |
| PENTADECALACTONE | 106-02-5 |
| PHENETHYL ACETATE | 103-45-7 |
| PHENOXYETHYL ISOBUTYRATE | 103-60-6 |
| PHENYLISOHEXANOL | 55066-48-3 |
| Tetrahydrolinalool | 78-69-3 |
| TETRAHYDRO-METHYL-METHYLPROPYL)-PYRAN-4-OL | 63500-71-0 |

Annex III, part 1

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number |
|--|----------------------|
| TRICHLOROMETHYL PHENYL CARBINYL ACETATE | 90-17-5 |
| TRICYCLODECENYL PROPIONATE | 17511-60-3 |
| TRIMETHYLHEXYL ACETATE | 58430-94-7 |
| gamma-UNDECALACTONE | 104-67-6 |
| VERDYL ACETATE | 2500-83-6/ 5413-60-5 |

7.2. Tabular summary of evaluated natural extracts/essential oils

Natural raw materials in terms of extracts are used in the fragrance and flavour industry for various reasons. Most importantly, several naturally occurring mixtures have a very complex composition and sensory nature which cannot (fully) be achieved by synthetic the demand for perfumes based on natural materials is considerable (189).

The three main methods used to concentrate plant fragrance substances (190); distillation, mechanical separation ("pressing"), and solvent extraction, yield very different extracts. Essential oils are obtained by water steam, water, ethanol, or water/ethanol distillation. Essence oils are essential oils that separate from the aqueous phase in the distillation receiver during the distillative concentration of fruit, usually citrus, juices. Citrus peel oils, apart from distilled lime oil, are prepared in a special way by pressing the peel to release mostly volatile substances from the pericarp in small oil glands, mostly highly volatile terpene hydrocarbons. However, they also contain small amounts of non-volatile compounds such as dyes, waxes and furocoumarins. The method of solvent extraction is generally applied in the separation of heat-labile materials or if an essential oil can only be obtained in very low yield, e.g. from blossoms. It is also used if the non-volatile components are desired for their fixative properties, e.g. in the preparation of resinoids from exudates. The most important extracts are termed: (i) concretes, an extract of fresh plant material with nonpolar solvents, containing not only volatile, but also a large proportion of non-volatile substances such as waxes; and (ii) absolutes, which are prepared by taking up concretes in ethanol; compounds that precipitate on cooling are removed by filtration, yielding a wax-free residue called absolute. Resinoids, used for their fixative properties, are prepared by extracting plant exudates with alcohols or nonpolar solvents. The products are usually highly viscous and thus sometimes diluted, e.g. with phthalates or benzyl benzoate. Oleoresins are concentrates prepared from spices by solvent extraction (189).

An ISO norm exists regarding the nomenclature of aromatic natural raw materials (ISO/DIS 9235 Aromatic raw materials - vocabulary; International Standardisation Organisation, Geneva, Switzerland). This nomenclature has been considered in Annex I, whereas in the present opinion, nomenclature is according to the CosIng database. Concerning extraction processes for many essential oils, ISO standards exist; for detailed information see Annex I to this opinion.

Regarding clinical data in terms of contact allergy to essential oils and natural extracts, the main focus is on general dermatological patients with complaints related to use of cosmetics etc. However, series of cases with occupational exposure to essential oils with occupational allergic contact dermatitis have also been reported (e.g. masseurs, physiotherapists (191, 192), aromatherapists (193-197), beauticians performing massages (198). For further details, e.g. PT results with various essential oils, see Annex I.

In this section, a tabular overview on the classification of substances considered is presented in three tables listing:

1. Extracts identified as *established contact allergens* in humans(→ *sufficient human evidence present*).

2. Extracts with positive human data, which are, however, not sufficient to categorise as *established contact allergen* in humans (→ *limited human evidence present*).
3. Extracts with negative human data, i.e. patch tests of patients with suspected contact allergy to fragrance ingredients which yielded negative results.

In Table 7-5, essential oils with sufficient human evidence to categorise these as *established contact allergens* in humans are presented.

Table 7-5: Natural extracts classified as established contact allergens in humans (summary of evaluation as detailed in chapter 6.3). More detailed information forming the basis of this evaluation can be found in Annex I of this opinion, including variants of botanical nomenclature.

| INCI name (or, if none exists, §perfuming name according to CosIng⁴) in bold; plant part / type of extract (partly indicative) in plain font | CAS number | Comment: see text |
|--|--------------------------------------|--------------------------|
| CANANGA ODORATA and Ylang-ylang oil | 83863-30-3; 8006-81-3 | +++ |
| CEDRUS ATLANTICA BARK OIL | 92201-55-3; 8000-27-9 | ++ |
| CINNAMOMUM CASSIA LEAF OIL CINNAMOMUM ZEYLANICUM BARK OIL | 8007-80-5 84649-98-9 | ++ (r.t.) |
| CITRUS AURANTIUM AMARA FLOWER / PEEL OIL | 8016-38-4; 72968-50-4 | ++ |
| CITRUS BERGAMIA PEEL OIL EXPRESSED [§] | 89957-91-5 | + (r.t.) |
| CITRUS LIMONUM PEEL OIL EXPRESSED [#] | 84929-31-7 | ++ |
| CITRUS SINENSIS (syn.: AURANTIUM DULCIS) PEEL OIL EXPRESSED [§] | 97766-30-8; 8028-48-6 | ++ |
| CYMBOPOGON CITRATUS / SCHOENANTHUS OILS | 89998-14-1; 8007-02-1; 89998-16-3 | ++ |
| EUCALYPTUS SPP. LEAF OIL [§] | 92502-70-0; 8000-48-4 | ++ |
| EUGENIA CARYOPHYLLUS LEAF / FLOWER OIL | 8000-34-8 | +++ |
| EVERNIA FURFURACEA EXTRACT ⁵ (Tree moss) | 90028-67-4 | +++ |
| EVERNIA PRUNASTRI EXTRACT (Oak moss) [#] | 90028-68-5 | +++ |
| JASMINUM GRANDIFLORUM / OFFICINALE | 84776-64-7; 90045-94-6; 8022-96-6 | +++ |
| JUNIPERUS VIRGINIANA | 8000-27-9; 85085-41-2 | ++ |
| LAURUS NOBILIS | 8002-41-3; 8007-48-5; 84603-73-6 | ++ |
| LAVANDULA HYBRIDA | 91722-69-9 | + (r.t.) |
| LAVANDULA OFFICINALIS [§] | 84776-65-8 | ++ |
| MENTHA PIPERITA | 8006-90-4; 84082-70-2 | ++ |
| MENTHA SPICATA | 84696-51-5 | ++ |
| MYROXYLON PEREIRAE (Balsam of Peru) [#] | 8007-00-9 | ++++ |

⁴ <http://ec.europa.eu/consumers/cosmetics/cosing/>

[#] 76/768/EEC Annex III, part 1

[#] 76/768/EEC Annex III, part 1

Opinion on fragrance allergens in cosmetic products

| INCI name (or, if none exists, §perfuming name according to CosIng⁴) in bold; plant part / type of extract (partly indicative) in plain font | CAS number | Comment: see text |
|--|------------------------------------|--------------------------|
| NARCISSUS SPP. | <i>diverse</i> | ++ |
| PELARGONIUM GRAVEOLENS | 90082-51-2; 8000-46-2 | ++ |
| PINUS MUGO/ PUMILA # | 90082-72-7 / 97676-05-6 | ++ |
| POGOSTEMON CABLIN | 8014-09-3; 84238-39-1 | ++ |
| ROSE FLOWER OIL (ROSA SPP.) | <i>Diverse</i> | ++ |
| SANTALUM ALBUM | 84787-70-2; 8006-87-9 | +++ |
| TURPENTINE (oil) # | 8006-64-2; 9005-90-7; 8052-14-0 | ++++ |
| VERBENA absolute # | 8024-12-2 | ++ |

Notes: r.t. Rarely tested.

Table 7-6 lists a number of essential oils, mostly tested in just one clinical department, and thus, or for other reasons, not satisfying the criteria for being categorised as *established contact allergen* in humans (i.e. *limited human evidence present*).

Table 7-6: Natural extracts with positive human data, which are, however, not sufficient to categorise as “established contact allergen in humans”. More detailed information forming the basis of this evaluation can be found in Annex I of this opinion.

| INCI name (or, if none exists, perfuming name according to CosIng) in bold; plant part / type of extract (partly indicative) in plain font | CAS number | Comment | Ref. |
|---|---------------------------|--|-------------|
| ACORUS CALAMUS ROOT OIL | 84775-39-3 | n=7 pos. reactions to “calamus” | (199) |
| CEDRUS DEODARA WOOD OIL | 91771-47-0 | Rudzki 1976/1986 found 3 / 3 positive reactions | (199, 200). |
| CITRUS AURANTIUM AMARA LEAF OIL | 72968-50-4 | Several cases in 2 series from 1 centre | (199, 200) |
| CITRUS TANGERINA ... | 223748-44-5 | 1 case | (201) |
| CYMBOPOGON NARDUS / WINTERIANUS HERB OIL | 89998-15-2; 91771-61-8 | Several cases in 2 series from 1 centre | (199, 200) |
| ILLICIUM VERUM FRUIT OIL | 84650-59-9 | Cases of active sensitisation; 34% consecutive patients pos. to 1% | (202) |

Opinion on fragrance allergens in cosmetic products

| INCI name (or, if none exists, perfuming name according to CosIng) in bold; plant part / type of extract (partly indicative) in plain font | CAS number | Comment | Ref. |
|---|--------------------------|---|-------------|
| LAVANDULA SPICA | 97722-12-8 | Several cases in 2 series from 1 centre | (199, 200) |
| LITSEA CUBEBA | 90063-59-5 | Several cases in 2 series from 1 centre | (199, 200) |
| PELARGONIUM ROSEUM | 90082-55-6 | 2.1% pos. of 1483 patients | (203) |
| ROSMARINUS OFFICINALIS | 84604-14-8 | 3 cases in 2 series from 1 centre | (199, 200) |
| SALVIA spp. | <i>Diverse</i> | Several cases in 2 series from 1 centre | (199, 200) |
| TAGETES PATULA | 91722-29-1 | 1 case (aromatherapist) | (193) |
| THYMUS spp. | 84929-51-1 | 4 / 84 pos | (199) |
| VETIVERIA ZIZANOIDES | 8016-96-4; 84238-29-9 | 1 / 200 and 9 / 86 pos. | (199, 200) |

The final table is an indicative list of natural extracts which lack published human data, but which are of interest: (i) as high-volume exposure; (ii) due to published positive animal experiments; or (iii) because they contain well-known (established) contact allergens.

Table 7-7: Indicative list illustrating natural extracts containing established human allergens or having R43-label or positive LLNA, lacking published human data.

| INCI name (or, if none exists, perfuming name according to CosIng) in bold; plant part / type of extract (partly indicative) in plain font | CAS number | Comment |
|---|-------------------|--|
| CITRUS PARADISI PEEL OIL | 8016-20-4 | high volume substance, classified as R43 |
| CYMBOPOGON MARTINI HERB EXTRACT | 84649-81-0 | Pos. LLNA study by RIFM: EC3 value 9.6% (204). |
| MENTHA ARVENSIS | 68917-18-0 | high volume, classified as R43 |
| OCIMUM BASILICUM | 84775-71-3 | Pos. LLNA study by RIFM: EC3 value < 2.5% (204). |
| PIMENTA RACEMOSA | 85085-61-6 | Contains, among other substances, the established contact allergen eugenol (42-56%) |
| SANTALUM SPICATA | 8024-35-9 | Contains, among other substances, the established contact allergens santalols (75%) and farnesol (10%) |

7.3. Conclusions

- According to the criteria described in chapter 6.3 a total of 54 individual chemicals and 28 natural extracts (essential oils) can be categorised as *established contact allergens* in humans, including all currently regulated substances.
- Of the 54 individual chemicals which are established contact allergens in humans, 12 are considered to be of special concern due to the high number of reported cases, (> 100, i.e. category +++ or ++++ in Table 7-1). These are further considered in chapter 5 (limonene and linalool) and the remainder in chapter 11. In particular one ingredient stands out, hydroxyisohexyl 3-cyclohexene carboxaldehyde, having been the cause of more than 1,500 reported cases since the 1999 opinion (see also chapter 4.3.1, chapter 11.3 and Annex I).
- For an additional 33 individual chemicals (Table 7-2) and 14 natural extracts (Table 7-6), positive patch test results have been reported. However, they do not qualify for the above category, i.e. only *limited human evidence* is present.
- For a number of fragrance substances (n=18, Table 7-3) patch testing did not yield positive results. However, numbers of patients tested are generally too small to rule out the existence of clinical contact sensitisation with sufficient confidence. No clinical evidence has been identified for 39 individual chemicals that have been reported to be frequently used (Table 7-4).
- For the substances (and, if possible, also for the main constituents of the natural mixtures) with limited or no human evidence, additional animal data and/or SAR considerations are taken into account. Aggregated data for these substances are presented in chapter 13.

8. Animal data

8.1. Predictive tests and sensitising potency categories

The animal test methods used in harmonised classification of substances, according to their potential to cause skin sensitisation, are the guinea pig maximisation test (GPMT), the Buehler test⁶ and the local lymph node assay (LLNA)⁷. These methods are used in hazard identification and risk assessment for regulatory purposes under REACH⁸. For registration in REACH, the LLNA is the preferred method for measuring skin sensitisation potential in animals, and justification for the use of other methods needs to be provided. According to the directives on classification and labelling⁹, substances and preparations meeting positive criteria in these tests shall be classified as sensitising and assigned the symbol "Xi" and the risk phrase "R43: May cause sensitisation by skin contact"; or, according to the recent regulation on classification, labelling and packaging (CLP¹⁰) "H317: May cause an allergic skin reaction".

As yet, there is no officially validated *in vitro* test method for skin sensitisation. Therefore, for cosmetic ingredients the LLNA, the GPMT and the Buehler test have also been used in risk assessment for regulatory purposes.

Positive results from the OECD guideline animal tests mentioned above which are sufficient to classify a substance as a skin sensitiser (R43) are:

- GPMT; at least 30% of the animals have a positive response.
- Buehler test; at least 15% of the animals have a positive response.
- LLNA; at least a 3-fold increase in lymph node cell proliferative activity is induced, compared to vehicle-treated controls (stimulation index $SI \geq 3$). For positive LLNAs, an EC3 value is calculated which gives the estimated concentration of a chemical necessary to give a 3-fold increase in proliferative activity compared to vehicle-treated controls.

Further categorisation of substances classified with R43 into three groups according to allergen potency (extreme, strong and moderate) has been proposed by a European Commission expert group on skin sensitisation (161, 205), and proposed also in the ECHA guidance document on application of the CLP criteria (162). Such categorisation is based on EC3 values in the LLNA, on intradermal induction concentration in the GPMT, and topical induction concentration in the Buehler test. The potency categories and their default concentration values based on EC3 values in the LLNA as defined in (161): extreme sensitiser (EC3 value ≤ 0.2); strong sensitiser (EC3 $> 0.2 - \leq 2$); and moderate sensitiser (EC3 value > 2). When LLNA EC3 values are available from more than one study, the lowest value should normally be used. Where multiple animal data sets lead to different categorisation of the same substance, the higher potency category should apply (161, 205).

The potency categorisation of substances based on the LLNA is applied by the SCCP in risk assessment of cosmetic ingredients, particularly hair dye substances (206).

⁶ OECD Guideline for testing of chemicals. Guideline 406: Skin Sensitisation. OECD, Adopted 12 May 1981, updated 17th July 1992.

⁷ OECD Guideline for testing of chemicals. Guideline 429: Skin Sensitisation: Local Lymph Node Assay. OECD, Adopted 22 July 2010.

⁸ Council Regulation (EC) No 440/2008 of 30 May 2008 laying down test methods pursuant to Regulation (EC) No 1907/2006 of the European Parliament and of the Council on the Registration, Evaluation, Authorisation and Restriction of Chemicals (REACH).

⁹ Directives 67/548/EEC and 1999/45/EC.

¹⁰ Regulation No. 1272/2008.

8.1.1. LLNA data

The SCCS requested the International Fragrance Association (IFRA) to submit data on animal tests performed with fragrance substances, to be presented in a structured format. In response, IFRA submitted first a poster (163) and later a report consisting of LLNA protocol summaries on the 59 fragrance substances in the poster (164). No guinea pig studies were submitted. The SCCS has reviewed and analysed the report and the publications quoted in the report.

Table 8-1 displays the EC3 values for fragrance substances in the report submitted by industry (164). EC3 values for some additional fragrance substances in two published reviews (165, 166) have also been included in Table 8-1. Table 8-2 presents LLNA results for oxidised substances. Full data are given in Annex II. Table 8-3 summarises the distribution of fragrance substances, by potency category, according to EC3 values.

Additional EC3 values may be available in the scientific literature. Many more animal experiments may have been performed, but have not been published.

Table 8-1: Summary of local lymph node assay (LLNA) data on 66 fragrance substances, based on a report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009 (164)) and in published reviews by Gerberick et al. 2005 (165) and Kern et al. 2010 (166), respectively. EC3 values (% and M) are given. The order of substances is by decreasing sensitisation potency as assessed by LLNA EC3 values (lowest EC3 value indicating highest potency).

| Substance | CAS no. | EC3 value | | Reference |
|--|------------|-----------|--------|------------|
| | | % | M | |
| Hexyl salicylate | 6259-76-3 | 0.18 | 0.008 | (164, 166) |
| Cinnamal | 104-55-2 | 0.2 | 0.015 | (164) |
| Methyl 2-octynoate | 111-12-6 | <0.5 | <0.032 | (164, 166) |
| Isoeugenol | 97-54-1 | 0.54 | 0.033 | (164) |
| Citral | 5392-40-5 | 1.2 | 0.079 | (164) |
| 2-Hexylidene cyclopentanone | 17373-89-6 | 2.4 | 0.14 | (164) |
| Methyl octine carbonate | 111-80-8 | 2.5 | 0.15 | (164) |
| Peru balsam absolute | 8007-00-9 | 2.5 | n/a | (164) |
| trans-2-Hexenal | 6728-26-3 | 2.6 | 0.26 | (164) |
| Benzyl Salicylate | 118-58-1 | 2.9 | 0.23 | (164, 166) |
| Butylphenyl methylpropional (BMHCA) | 80-54-6 | 2.9 | 0.14 | (164) |
| Phenylacetaldehyde | 122-78-1 | 3 | 0.25 | (164, 165) |
| Allyl phenoxyacetate | 7493-74-5 | 3.1 | 0.16 | (164) |
| Benzylideneacetone | 122-57-6 | 3.7 | 0.25 | (165) |
| 3-Propylideneophthalide | 17369-59-4 | 3.7 | 0.21 | (164, 165) |
| <i>Evernia prunastri</i> extract oak moss | 90028-68-5 | 3.9 | n/a | (164) |
| Balsam oil, Peru (<i>Myroxylon pereirae</i> Klotzsch) | 8007-00-9 | 4 | n/a | (164) |
| Farnesol | 4602-84-0 | 4.1 | 0.18 | (164) |
| p-t-Butyl-dihydrocinnamaldehyde | 18127-01-0 | 4.3 | 0.23 | (164) |
| α-Methyl cinnamic aldehyde | 101-39-3 | 4.5 | 0.31 | (164, 165) |

Opinion on fragrance allergens in cosmetic products

| Substance | CAS no. | EC3 value | | Reference |
|--|-------------|-----------|-------|------------|
| | | % | M | |
| Eugenol | 97-53-0 | 5.3 | 0.32 | (164) |
| Hexyl cinnamal | 101-86-0 | 5.3 | 0.25 | (164) |
| Dihydrocoumarin | 119-84-6 | 5.6 | 0.38 | (165) |
| Geraniol | 106-24-1 | 5.6 | 0.36 | (164) |
| Carvone | 6485-40-1 | 5.7 | 0.38 | (164) |
| Diethyl maleate | 141-05-9 | 5.8 | 0.34 | (165) |
| 2-Methoxy-4-methylphenol | 93-51-6 | 5.8 | 0.42 | (164, 165) |
| Anise alcohol | 105-13-5 | 5.9 | 0.43 | (164, 166) |
| Jasmine absolute (<i>Grandiflorum</i>) | 8022-96-6 | 5.9 | N/a | (164) |
| Dibenzyl ether | 103-50-4 | 6.3 | 0.32 | (164) |
| <i>Cananga odorata</i> leaf/flower oil ylang ylang "extra" | 8006-81-3 | 6.8 | N/a | (164) |
| Isocyclocitral | 1335-66-6 | 7.3 | 0.48 | (164) |
| 2,3-Dihydro-2,2,6-trimethylbenzaldehyde | 116-26-7 | 7.5 | 0.50 | (165) |
| Amyl cinnamal | 122-40-7 | 7.6 | 0.38 | (164) |
| Perillaldehyde p-Mentha-1,8-dien-7-al | 2111-75-3 | 8.1 | 0.54 | (164, 165) |
| p-Isobutyl- α -methyl hydrocinnamaldehyde | 6658-48-6 | 9.5 | 0.46 | (164) |
| d-Limonene* | 5989-27-5 | <10 | <0.73 | (164) |
| Methylundecanal | 110-41-8 | 10 | 0.54 | (165) |
| Acetylcedrene | 32388-55-9 | 13.9 | 0.57 | (166) |
| Methylenedioxyphenyl methylpropanal | 1205-17-0 | 16.4 | 0.85 | (164, 166) |
| Benzyl benzoate | 120-51-4 | 17 | 0.80 | (165) |
| Hydroxyisohexyl 3-cyclohexene carboxaldehyde | 31906-04-4 | 17.1 | 0.81 | (164, 165) |
| Benzyl cinnamate | 103-41-3 | 18.4 | 0.77 | (164, 166) |
| Hydroxycitronellal | 107-75-5 | 19.3 | 1.12 | (164) |
| Cinnamyl alcohol | 104-54-1 | 21 | 1.57 | (165) |
| α -iso-Methylionone | 127-51-5 | 21.8 | 1.06 | (164, 166) |
| Cyklamen aldehyde | 103-95-7 | 22 | 1.64 | (165) |
| 4-Methoxy- α -methyl benzenpropanal | 5462-06-6 | 23.6 | 1.32 | (164) |
| Amyl cinnamyl alcohol | 101-85-9 | ~25 | ~1.22 | (164, 166) |
| Tetramethyl acetyloctahydronaphthalenes (OTNE) | 54464-57-2 | 25.1 | 1.07 | (164) |
| Ethyl acrylate | 140-88-5 | 28 | 2.8 | (165) |
| Linalool* | 78-70-6 | 30 | 1.94 | (165) |
| Trimethylbenzenepropanol Majantol | 103694-68-4 | 30 | ~1.68 | (164) |
| Jasminum Sambac Flower CERA/Extract/Water | 91770-14-8 | 35.4 | N/a | (164) |

| Substance | CAS no. | EC3 value | | Reference |
|---|------------|-----------|-------|------------|
| | | % | M | |
| Citronellol | 106-22-9 | 43.5 | 2.78 | (164, 166) |
| No EC3 value was established; higher concentrations should also have been tested | | | | |
| 6-Methyl-3,5-heptadien-2-one | 1604-28-0 | >5 | >0.40 | (164) |
| <i>Camellia sinensis</i> leaf tea leaf absolute | 84650-60-2 | >5 | N/a | (164) |
| Cinnamyl nitrile | 1885-38-7 | >10 | >0.77 | (164) |
| Menthadiene-7-methyl formate | 68683-20-5 | >10 | >0.51 | (164) |
| <i>Evernia furfuracea</i> extract tree moss absolute | 90028-67-4 | >20 | N/a | (164) |
| Isocyclogeraniol | 68527-77-5 | >25 | >1.62 | (164) |
| 1-Octen-3-yl acetate | 2442-10-6 | >30 | >1.76 | (164) |
| Benzyl alcohol | 100-51-6 | >50 | >4.62 | (164) |
| Coumarin | 91-64-5 | >50 | >3.42 | (164) |
| Vanillin | 121-33-5 | >50 | >3.3 | (164) |
| No EC3 value calculated | | | | |
| Benzaldehyde | 100-52-7 | - | | (165) |

Notes: * Material with low levels of oxidation according to (164)
n/a: Not applicable (mixture of compounds).

M: EC3 based on molar concentration

8.1.2. LLNA data on oxidised fragrance substances

For fragrance substances that can autoxidise upon air exposure, it is also important to investigate the sensitisation potency after air exposure. The oxidised compounds are clinically relevant as they represent what the consumers could come in contact with from perfumes and fragranced products. In Table 8-2 the LLNA data for some of the most commonly used fragrance substances, pure and after autoxidation, are presented. The EC3 values obtained for the pure substances are 5-10 times higher compared to those obtained for the same substances after air exposure. The experimental air exposure simulated air exposure that can take place during normal handling and storage. In the production process, some perfumes are "matured" aerobically, stirring included. During this process, some fragrance substances may be oxidised. It should be noted that, although only a few substances capable of oxidation have so far been investigated, structural alerts indicating possible autoxidation are common among the fragrance substances listed in this document (see chapter 9). It is important to further investigate this issue for increased understanding of the associated risk.

Table 8-2: Local lymph node assay (LLNA) data on four fragrance substances and one essential oil before and after air exposure, comparing the sensitisation potency of the pure (not oxidised) substance with the potency of the oxidised.

| Substance | CAS no. | Doses % (w/v) vehicle: A:OO 4:1* | EC3 value (% w/v) | Reference |
|-----------------------|-----------|--|----------------------|-----------|
| D-Limonene (ox. 10 w) | 5989-27-5 | 1, 5, 25 | 3.0 | (207) |
| D-Limonene (pure) | 5989-27-5 | 25, 50, 100 | 30 | |

| Substance | CAS no. | Doses % (w/v) vehicle: A:OO 4:1* | EC3 value (% w/v) | Reference |
|----------------------------|----------|--|----------------------|-----------|
| Linalool (ox. 10 w) | 78-70-6 | 5, 10, 25 | 9.4 | (127) |
| Linalool (ox. 45 w) | 78-70-6 | 2.5, 10, 25 | 4.8 | |
| Linalool (pure) | 78-70-6 | 25, 50, 100 | 46.2 | |
| Linalyl acetate (ox. 10 w) | 115-95-7 | 0.5, 10, 40 | 3.6 | (128) |
| Linalyl acetate (pure) | 115-95-7 | 10, 30, 100 | 25 | |
| Geraniol (ox. 10 w) | 106-24-1 | 1, 3, 6, 10, 20 | 4.4 | (119) |
| Geraniol (ox. 45 w) | 106-24-1 | 0.5, 1, 3, 6, 10 | 5.8 | |
| Geraniol (pure) | 106-24-1 | 5, 10, 15, 20, 30 | 22.4 | |
| Lavender oil (ox. 10 w) | | 1, 5, 10, 20, 50 | 11 | (140) |
| Lavender oil (ox. 45 w) | | 1, 5, 10, 20, 50 | 4.4 | |
| Lavender oil (not ox.) | | 5, 25, 100 | 36 | |

Notes: Pure: Purified before testing as most commercially available fragrance substances are not pure.

Not ox.: Not purified but used as it was delivered as this is a complex mixture and not a specific substance.

Ox. x w: Oxidised by air exposure during x weeks.

* Acetone:olive oil.

8.2. Methodological considerations

EC3 mean values

In the submitted poster (163) and the report by IFRA (164), the LLNA weighted mean EC3 values ($\mu\text{g}/\text{cm}^2$) are presented. The SCCS considers it is misleading to present EC3 values as mean values from tests performed with different vehicles. It is generally agreed that the lowest EC3 value should be used if there is more than one study fulfilling the OECD guideline requirements (161, 205), and these have been introduced into Table 8-1. The EC3 values in the reviews by Gerberick et al. and Kern et al. (165, 166) were based on single representative experiments with a vehicle described in the OECD guideline 429 (see above), and preferably with acetone:olive oil. EC3 mean values, as in the submission by IFRA, were not presented in these two reviews.

Vehicle

The most frequently used *vehicle* in the submission by IFRA (164) was ethanol:diethyl phthalate (1:3), followed by acetone:olive oil (4:1). In some experiments, antioxidants were mixed with ethanol:diethyl phthalate. The vehicle was not reported in some of the references, and no rationale for using vehicles other than those recommended was given in the report (164). According to the OECD guideline 429 (see above), the recommended vehicles are acetone:olive oil (4:1), N,N-dimethylformamide, methyl ethyl ketone, propylene glycol, and dimethyl sulphoxide, but others may be used if sufficient scientific rationale is provided. It is well known that a difference in the EC3 value can be obtained for the same substance depending on which vehicle is used in the LLNA. Thus as an *additional control*, supplementary to the guideline based LLNA control, a clinically relevant solvent or the commercial formulation in which the test substance is marketed may be used.

Number of doses and animals

According to the OECD guideline 429 (see above), a minimum of three concentrations should be tested. The number of consecutive doses used in the reported data, was generally five, sometimes three and in few experiments two. The SCCS considers that too few concentrations were tested in four studies in which only two concentrations were used. Lower concentrations than those tested should have been used in experiments with five fragrance substances, in which the EC3 value could not be determined. Higher concentrations than those tested should also have been used in experiments with 12 substances, in which the EC3 value could not be determined.

The *number of animals* per dose group was generally four plus a non-exposed control group, sometimes five, and in few experiments six; the minimum according to the OECD guideline being four.

Units for concentrations

In the submission by IFRA (164) the EC3 values are given in weight per area unit ($\mu\text{g}/\text{cm}^2$). The SCCS considers that the EC3 values (%) are the values of primary interest in communicating risk assessment, as EU legislation, OECD guideline 429 and scientific literature refer to EC3 values (%). However, the SCCS recommends that molar (M) EC3 values should be considered, as they give the concentration based on the molecular weight of substances. They have thus been calculated and introduced into Table 8-1.

EC3 values (%) overestimate the intrinsic molecular sensitisation potency for low molecular weight compounds while compounds with a high molecular weight are underestimated. Regarding the differences in molecular weight between the studied fragrance substances, a variation is seen if the ranking list of the sensitisation potency is based on EC3 (%) or EC3 (M) since some substances have a molecular weight twice as high as others.

From comparisons in Table 8-1, we notice that, e.g. hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC) has an EC3 value of 17.1 %, or 0.81 M when the calculation includes its molecular weight, while for trans-2-hexenal the corresponding values are 2.6% and 0.26 M. The example shows that comparing the sensitisation potency between these two substances using the EC3 values in % exaggerates the sensitisation potency of trans-2-hexenal compared to that of HICC. When using the EC3 values in molar concentrations the difference is not so pronounced.

8.3. Summary of animal data by LLNA

The distribution of sensitising potency of fragrance substances compared to other substances, (e.g. biocides, dyes, plastic materials) taken from three references (164-166) as assessed by EC3 values in the LLNA, is shown in Figure 8-1 and Table 8-3.

For 10 substances, no EC3-value could be established. These should have been tested at higher concentrations – some of these would most probably have generated an EC3 value. However, we reported here “No EC3 value established”. 5 substances should have been tested also at lower concentration and in these cases the EC3 value could have been lowered, meaning a more severe potency category could have been achieved. In all, approx 150 experiments were reported in (164), listed in Annex II.

The median EC3 value of evaluable fragrance substances (5.9%) is similar to other substances tested (5.5%). However, very few fragrance substances have low EC3 values (≤ 2).

Substances with an EC3 value ≤ 2 may be categorised as strong or extreme sensitisers. Such potent sensitisers are comparatively rare among fragrance substances assessed in the LLNA. Nevertheless, fragrances are important allergens in humans, which points to repeated skin exposure to less potent sensitisers as a factor strongly determining sensitisation risk.

Opinion on fragrance allergens in cosmetic products

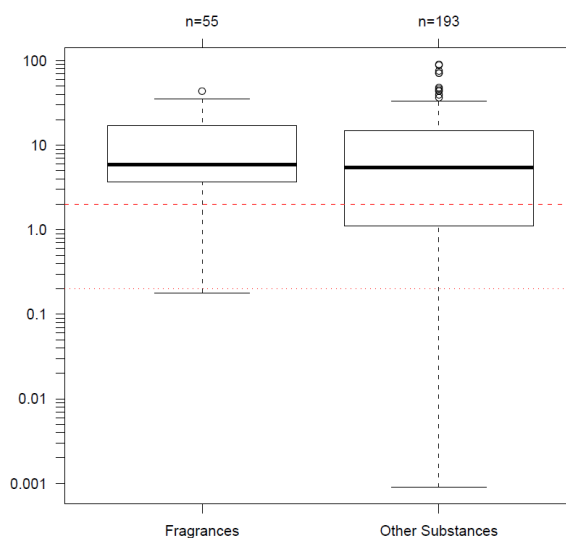


Figure 8-1: The distribution of fragrance chemicals and a variety of other chemicals (e.g. biocides, dyes, plastic materials), taken from the three references (164-166), are depicted as boxplots on a logarithmic scale. The bottom of the box denotes the 1st quartile (25% percentile), the thick line in the box the median, and the top of the box the 3rd quartile (75% percentile). Outliers, i.e. below the 25% and above the 75% percentiles, are shown as whiskers. Beyond the 1.5-fold interquartile range, single values are shown as circles instead of whiskers. The difference in distribution is not significant (Wilcoxon test: $p=0.061$).

Note: EC3 values for the five oxidised fragrances additionally examined (Table 8-2) range from 3.0 to 4.8 (median 4.4) and are lower by a factor of around 7 than EC3 values of the respective non-oxidised material.

Table 8-3: Summary of EC3 values for fragrance substances in Table 8-1 and for other substances, all taken from the three references (164-166). The EC3 value intervals for potency categorisation (161, 205) were used for comparison of fragrances substances vs other substances.

| EC3 value interval | Fragrance substances | | Other substances | |
|------------------------------|----------------------|-----|------------------|-----|
| | n | % | n | % |
| ≤ 0.2 | 2 | 3% | 28 | 11% |
| > 0.2 - ≤ 2 | 3 | 4% | 38 | 15% |
| > 2 | 50 | 71% | 127 | 49% |
| No EC3 value established * | 10 | 14% | 0 | 0% |
| No EC3 value calculated (NC) | 5 | 7% | 69 | 26% |
| All substances | 70 | | 262 | |

Note: * Substances should have been tested also at higher concentrations.

8.4. Conclusions

- In the event that human data are lacking, the LLNA provides important information on skin sensitising potential and potency.
- Animal data on fragrance substances submitted by IFRA (164) and assessed in this opinion were generated exclusively by LLNA. Other guideline methods are, however, also available.
- The vast majority of the submitted (164) and additional (165, 166) fragrance substances tested by the LLNA are skin sensitisers.
- Several studies in the IFRA report (164) were of insufficient quality, not following the OECD guideline.

- Fragrance substances that can be predicted to autoxidise upon air exposure should also be tested after air exposure, as oxidation may significantly increase their sensitising potency.
- It can be concluded that the skin sensitising potency, as assessed by the LLNA, is only one of several factors that are of importance for sensitisation to fragrance substances. This is illustrated by the fact that only a small fraction of sensitising fragrance substances can be categorised as an extreme allergen based on LLNA test results. Therefore, doses from repeated deposition onto skin must be considered a driving force of sensitisation risk.

9. Structure activity relationships (SAR): grouping of substances based on expert judgement

Whether or not a particular chemical will be a sensitiser, and how potent it will be if it is a sensitiser, depends on its ability, either directly or after activation, to react with appropriate proteins in the skin. This fundamental concept was initially demonstrated by Landsteiner and Jacobs in 1936 (208) and subsequently validated by numerous studies with various types of chemicals (some key references: (209-213)). The ability to predict sensitisation potency, or lack of it, depends on being able to predict reactivity to skin proteins. This is the basis of SAR analysis for skin sensitisation. The prediction can often be made based on the chemical structure, recognising structural features (referred to as **structural alerts**) that are associated with reactivity.

The relationships between molecular structure and reactivity that form the basis for structural alerts are based on well established principles of mechanistic organic chemistry (214). Examples of structural alerts are aliphatic aldehydes (alerting to the possibility of sensitisation via a Schiff base reaction with protein amino groups), and α,β -unsaturated carbonyl groups, C=C-CO- (alerting to the possibility of sensitisation via Michael addition of protein thiol groups). Major mechanistic reactivity domains have been discussed in detail by Aptula and Roberts (215). Prediction of the sensitisation potential of compounds that can act via abiotic or metabolic activation (pre- or prohaptens) is more complex compared to that of compounds that act as direct haptens without any activation. The autoxidation patterns can differ due to differences in the stability of the intermediates formed, e.g. it has been shown that autoxidation of the structural isomers linalool and geraniol results in different major haptens/allergens. Moreover, the complexity of the prediction increases further for those compounds that can act both as pre- and prohaptens. In such cases, the impact on the sensitisation potency depends on the degree of abiotic activation (e.g. autoxidation) in relation to the metabolic activation.

These structural alerts can be applied by computerized expert systems, i.e. *in silico* or by estimations made by organic chemists (*in cerebro*) using their experience. When an organic chemist looks at a chemical structure, they recognise parts of the structure that they can associate with reactivity, the type of reactivity (i.e. assign the reaction mechanistic domain), and other features of the molecular structure that will affect the reactivity positively or negatively. Human experts should be aware of the complexities, and how structural modification can alter the reactivity associated with structural alerts, etc. Importantly, they can also recognise where there are unfamiliar structural features whose effects they cannot confidently predict. In such cases they can call for experimental chemistry work (*in chemico*) to be done to ascertain the presence or nature of, and degree of reactivity. *In chemico* methods include organic chemistry experimentation to identify chemical reaction products from oxidation and/or reaction with model nucleophiles, identification of mechanisms of reaction. In so called *in chemico* reactivity methods, the ability of a specific chemical to react with selected peptides is determined so as to predict the sensitisation potential of the chemical under investigation (216, 217). To make *in chemico* reactivity methods able to predict the activity of prohaptens, the addition of horseradish peroxidase and hydrogen peroxide oxidation system has been tested to model the enzymatic oxidation in the skin (218, 219).

Although computerized expert systems are derived from input by human experts, they are less well able to capture the subtleties of structure reactivity relationships, and they sometimes fail to detect aspects of chemistry that are obvious to organic chemists. Human experts should be aware of the complexities, as well as how structural modification can alter the reactivity associated with structural alerts, etc.

The SAR evaluation made in the section below is based on *in cerebro* alerts applied by organic chemists.

Depending on the type of reactivity (the **reaction mechanistic domain**), it is sometimes possible to make a quantitative prediction of potency in the LLNA, which can be used to predict potency in humans relative to related known human sensitisers. These predictions use quantitative mechanistic models (**QMMs**) based on reactivity expressed quantitatively

by model parameters, and sometimes in combination with hydrophobicity. For example, potency of aliphatic aldehydes and ketones (the Schiff base domain) in the LLNA is modelled by a combination of reactivity and hydrophobicity (220), whereas the LLNA potency of DNCB analogues (the S_NAr domain) is well modelled by reactivity alone (221).

QMMs aiming not only to predict the potential to be a sensitizer but also to predict the potency, promise to be a useful tool in non-animal based risk assessment for skin sensitisation. However, in the field of fragrance substances there are major gaps in our present ability to apply QSAR/QMM. This is largely because many of the fragrance substances of interest have the potential to act via abiotic or metabolic activation (pre- and/or prohaptens, i.e. they themselves are only weak or non-sensitizers, but have the potential to be activated to form more potent sensitizers. Resulting sensitization potency will depend on the extent of activation and the nature of the resulting products. It is possible to apply SAR analysis to identify these plausible possibilities, but QSAR modelling for these cases is not yet developed. However, much progress has been made in identifying structural alerts for the various activation mechanisms that have been recognised. This is reviewed by Karlberg et al. (122).

Chemicals with no structural alerts for direct reactivity, or for known activation mechanisms, and no unfamiliar structural features that might be associated with as yet unidentified activation mechanisms, can be predicted to be non-sensitizing. Chemicals that do have alerts for reactivity (direct or via activation) are not necessarily sensitizers – they may be insufficiently reactive and/or insufficiently hydrophobic.

Substances meeting the inclusion criteria (see chapter 6), for which, however, no categorisation as established contact allergen in humans or established contact allergen in animals was possible, have been assessed for structural alerts. The results are presented in four tables based on the prediction made for the actual substance. The following SAR assessments have been used:

- Predicted sensitizer; structural alerts:
Compounds containing structural alerts comprising direct reactive compounds and for compounds that after specific abiotic or metabolic activation (prohaptens and prehaptens) can be predicted to be sensitizers by structural comparison to known allergens.
- Possible sensitizer; structural alerts:
Compounds containing structural alerts that by comparison to known allergens with similar structures were expected to be less reactive and hence less likely to be sensitizing. Also compounds with structural alerts indicating a possible abiotic or metabolic activation (possible prehaptens or prohaptens) but with no structural data available for comparison, were included in this group. Consequently, a possible sensitizer may turn out to be a non sensitizer when tested in vivo.
- Predicted non-sensitizer (NS); no obvious structural alerts
- Not predictable due to insufficient/conflicting data

Table 9-1: Predicted sensitizers.

| Substance (INCI) name | CAS number | Structural alerts |
|---|------------|-------------------------------------|
| p-tert.-Butyldihydrocinnamaldehyde [§] | 18127-01-0 | Schiff base |
| Citronellal | 106-23-0 | Schiff base and possible prehaptent |
| Citronellyl nitrile | 51566-62-2 | Possible prehaptent |
| Decanal | 112-31-2 | Schiff base |
| 3,7-Dimethyl-1,6-nonadien-3-ol | 10339-55-6 | Prehaptent |
| Geranyl acetate | 105-87-3 | Prehaptent and prohaptent |

Opinion on fragrance allergens in cosmetic products

| | | |
|----------------------------------|------------|--|
| Isoamyl salicylate | 87-20-7 | Acyltransfer agent |
| Methyl cinnamate | 103-26-4 | Michael acceptor |
| Methylundecanal | 110-41-8 | Schiff base |
| Myrcene | 123-35-3 | Prehaptent |
| Nerol | 106-25-2 | Prehaptent and prohaptent |
| Nerolidol (isomer not specified) | 7212-44-4 | Possible prehaptent |
| Oxacyclohexadecanone | 34902-57-3 | Michael acceptor |
| Phenethyl salicylate | 87-22-9 | Acyltransfer agent |
| trans-Rose ketone-5 | 39872-57-6 | Michael acceptor and possible prehaptent |

Note: § Classified as R43.

Table 9-2: Possible sensitizers.

| Substance (INCI) name | CAS number | Structural alerts |
|---|-------------------|---|
| Ambrettolide | 7779-50-2 | Possible prehaptent |
| Amylcyclopentanone | 4819-67-4 | Schiff base; the combination of reactivity and hydrophobicity may be enough to confer sensitisation |
| Benzyl acetate | 140-11-4 | Prohaptent via hydrolysis leading to benzyl alcohol |
| Carvacrol | 499-75-2 | Possible prehaptent |
| Cuminaldehyde | 122-03-2 | Schiff base and possible prehaptent |
| alpha-Cyclohexylidene benzeneacetone | 10461-98-0 | Possible Michael acceptor |
| Cyclopentadecanone | 502-72-7 | Schiff base; the combination of reactivity and hydrophobicity may be enough to confer sensitisation |
| trans-beta-Damascone | 23726-91-2 | Possible Michael acceptor |
| trans-trans-delta-Damascone | 71048-82-3 | Possible Michael acceptor and possible prehaptent |
| gamma-Damascone | 35087-49-1 | Possible Michael acceptor and possible prehaptent |
| Dihydromyrcenol | 18479-58-8 | Possible prehaptent |
| 2,3-Dihydro-2,2,6-trimethylbenzaldehyde | 116-26-7 | Possible Michael acceptor and possible prohaptent |
| 2,4-Dimethyl-3-cyclohexen-1-carboxaldehyde § | 68039-49-6 | Schiff base and possible prehaptent |
| Dimethyltetrahydro benzaldehyde | 68737-61-1 | Schiff base and possible prehaptent |
| 6-Ethylideneoctahydro-5,8-methano-2H-benzo-1-pyran | 93939-86-7 | Possible prehaptent |
| 2-Ethyl-4-(2,2,3-trimethyl-3-cyclopenten-1-yl)-2-buten-1-ol | 19-61-6 | Possible prehaptent |
| Ethyl vanillin | 121-32-4 | Complex |
| Heliotropine | 120-57-0 | Possible prohaptent |
| 3a,4,5,6,7,7a-Hexahydro-4,7-methano-1H-inden-5(or 6)-yl | 54830-99-8 | Possible prehaptent |

Opinion on fragrance allergens in cosmetic products

| Substance (INCI) name | CAS number | Structural alerts |
|-------------------------------------|-------------------------|---|
| acetate | | |
| Hexahydro-methanoindenyl propionate | 68912-13-0 | Possible prehapten |
| Ionone isomeric mixture | 8013-90-9 | Possible Michael acceptor and possible prehapten |
| alpha-Ionone | 127-41-3 | Possible Michael acceptor and possible prehapten |
| beta-Ionone | 79-77-6 | Possible Michael acceptor |
| Isobergamate | 68683-20-5 | Possible prehapten |
| Isolongifoleneketone | 33407-62-4 | Schiff base; the combination of reactivity and hydrophobicity may be enough to confer sensitisation |
| Longifolene [§] | 475-20-7 | Possible prehapten |
| Methoxycitronellal | 3613-30-7 | Schiff base |
| Methyl decenol | 81782-77-6 | Possible prehapten |
| Methyl ionone (mixture of isomers) | 1335-46-2 | Possible Michael acceptor and possible prehapten |
| Methylionantheme | 55599-63-8 | Possible Michael acceptor and possible prehapten |
| 5-Methyl-alpha-ionone | 79-69-6 | Possible Michael acceptor and possible prehapten |
| Myrtenol | 515-00-4 | Possible prehapten |
| Nopyl acetate | 128-51-8 | Possible prehapten |
| Phytol | 150-86-7 | Possible prehapten and/or prohaptent |
| Rhodinol | 6812-78-8 | Possible prehapten |
| Terpineol acetate (isomer mixture) | 8007-35-0 | Possible prehapten |
| alpha-Terpinyl acetate | 80-26-2 | Possible prehapten |
| Tricyclodecanyl propionate | 17511-60-3 | Possible prehapten |
| Verdyl acetate | 2500-83-6/ 5413-60-5 | Possible prehapten |

Note: [§] Classified as R43.

Table 9-3: Predicted non-sensitisers with no obvious structural alerts.

| Substance (INCI) name | CAS number | Structural alerts |
|--|------------|--|
| 6-Acetyl-1,1,2,4,4,7-hexamethyltetraline | 21145-77-7 | |
| Benzyl acetone | 2550-26-7 | Schiff base; the combination of reactivity and hydrophobicity may not be enough to confer sensitisation |
| 2-tert.-Butylcyclohexyl acetate | 88-41-5 | |
| 4-tert.-Butylcyclohexyl acetate | 32210-23-4 | |
| Cyclohexyl acetate | 622-45-7 | |
| Diphenyl ether | 101-84-8 | |

Opinion on fragrance allergens in cosmetic products

| Substance (INCI) name | CAS number | Structural alerts |
|--|------------|--|
| Ethyl 2-methylbutyrate | 7452-79-1 | |
| Ethylene dodecanoate | 54982-83-1 | |
| Ethylene brassylate | 105-95-3 | |
| Eucalyptol | 470-82-6 | |
| Hexyl acetate | 142-92-7 | |
| Hibiscolide | 6707-60-4 | |
| Hydroxycitronellol | 107-74-4 | However, dehydration followed by autoxidation could give sensitising impurities |
| Isoamyl acetate | 123-92-2 | |
| Isobornyl acetate | 125-12-2 | |
| Methoxytrimethylheptanol | 41890-92-0 | |
| Methyl p-anisate | 121-98-2 | |
| Methyl anthranilate | 134-20-3 | |
| Methylbenzyl acetate | 93-92-5 | |
| Methyl dihydrojasmonate | 24851-98-7 | Schiff base; the combination of reactivity and hydrophobicity may not be enough to confer sensitisation |
| Oxalide | 1725-01-5 | |
| Pentadecalactone | 106-02-5 | |
| Phenethyl acetate | 103-45-7 | |
| Phenethyl alcohol | 60-12-8 | |
| Phenoxyethyl isobutyrate | 103-60-6 | |
| Phenylisohexanol | 55066-48-3 | |
| Phenylpropanol | 122-97-4 | |
| Tetrahydrolinalool | 78-69-3 | |
| Tetrahydro-methyl-methylpropyl)-pyran-4-ol | 63500-71-0 | |
| Trimethylhexyl acetate | 58430-94-7 | |
| Trimethyl-propylcyclohexanepropanol (tmch) | 70788-30-6 | |
| gamma-Undecalactone | 104-67-6 | |

Table 9-4: Not predictable.

| Substance (INCI) name | CAS number | Structural alerts |
|---|------------|--|
| Anisaldehyde | 123-11-5 | Due to insufficient /conflicting data; structural similarities to benzaldehyde suggest certain activity in man |
| Trichloromethyl phenyl carbonyl acetate | 90-17-5 | Due to insufficient /conflicting data |
| Methyl beta-naphthyl ether | 93-04-9 | Due to insufficient /conflicting data |

9.1. General results

From this work with the included SAR predictions, the following observations can be made.

- For substances for which sufficient experimental/clinical evidence is missing, SAR analyses have been performed to predict a probable or possible risk of allergenic (sensitising) effect. These predictions are based on chemical reactivity and the recognition of structural features in a substance that are in common with the structural features that have been shown to cause sensitisation from other substances. In cases where the SAR analysis indicates a sensitisation potential, the substance should be investigated further to confirm or reject the conclusion drawn from the SAR analysis.
- Prediction of the sensitisation potential of compounds that can act via abiotic or metabolic activation (pre- or prohaptens) becomes more complex compared to that of compounds that act as direct haptens without any activation.
- The complexity of the prediction increases further for those compounds that can act both as prehaptens and prohaptens.
- Prediction of the sensitisation potential of compounds that can act as prehaptens is further complicated by the fact that the autoxidation patterns can differ due to differences in the stability of the intermediates formed, e.g. it has been shown that autoxidation of the structural isomers of linalool and geraniol results in different major haptens/allergens.

9.2. Conclusions

The SAR evaluation made in this section is based on *in cerebro* alerts applied by organic chemists.

- Applying only mechanism-based QSAR (QMM) as a tool in non-animal based risk assessment for skin sensitisation is of limited value for fragrance substances. This is due to major information gaps in the present model when addressing substances that act via abiotic or metabolic activation, and the high incidence of such substances in fragrances.
- Quantitative structure activity relationship (QSAR) models should be further developed, combining, as appropriate, information from *in silico*, *in chemico* and *in vitro* methods.
- SAR, as performed here, is only one consideration in the overall weight of evidence.

10. Exposure

Exposure to fragrance chemicals and other potential allergens is most commonly by direct skin contact. Exposures to fragrance chemicals occur from:

- Personal cosmetic use;
- Detergents and other household products;
- Medicaments;
- Occupation, i.e. personal hygiene, manufacturing ingredient(s), product in work process, plant materials;
- Secondary exposure from another individual (e.g. spouse, child);
- Toys;
- Oral intake;
- Airborne exposure.

Factors that are important for both the induction and elicitation of contact allergy are:

- Dose per unit area;
- Vehicle effects including penetration enhancers;
- Presence of skin irritants;
- Presence of other allergens (combination effects);
- Duration of skin exposure;
- Frequency of applications;
- Anatomical sites of exposure;
- Condition of the skin (barrier function impairment, pre-existing inflammation);
- Occlusion (e.g. in flexures, under clothing and personal protective equipment).

Fragrance mix ingredients are commonly present in cosmetic formulations (71, 222-224). Cosmetics based on natural ingredients may contain fragrance allergens at a higher concentration than other cosmetic products (225). The clinical significance of exposure to natural extracts is difficult to determine as there is often "hidden and variable" exposure to important and potent allergens in natural products.

10.1. Concentrations and quantities used

Consumers are exposed in daily life to fragrance chemicals from a large variety of products, such as cosmetics, toys, detergents and other cleaning products, etc. The fragrance exposure may be via dermal and/or inhalation route. With respect to "Terms of Reference" to the SCCS, only dermal exposure from cosmetics is addressed in this opinion. As cosmetics are the perfumed products most commonly used in daily life, potential fragrance allergens identified by the use of cosmetics also represent the exposures of these chemicals from other product categories. In recent years, it has become a trend to add fragrance chemicals to many other types of consumer products, such as children's toys, toilet paper and nappies, which may contribute significantly to the fragrance exposure of the consumer by the dermal route.

Factors for the fragrance exposure assessment by the dermal route require knowledge on:

- Product types (categorisation of scented products) used by the consumer.
- Market survey (impression of the qualitative and quantitative contents of different allergens in consumer products).

- Hydrolysis, metabolism or oxidation of a fragrance material, which may generate a potential skin allergen.
- Chemicals in the product matrix, which may significantly enhance or reduce dermal absorption of a fragrance material.

Fragrance materials, both defined chemical substances and natural mixtures of chemicals (essential oils), are used in all types of cosmetic products: perfumes, eau de cologne, eau de perfume (EDP), and eau de toilette (EDT), aftershave lotion, deodorants, skin care products, skin cleansers, make-up cosmetics, hair care products, and oral care products, etc. However, some unscented cosmetic products have also reached the market in the last decade. Products containing the highest concentration of fragrance chemicals are perfumes, followed by eau de cologne, eau de perfume (EDP) and eau de toilette (EDT). Concentrations of fragrance chemicals in deodorant products are lower than those in EDT/EDP products, but still significant. Aftershave products also contain relatively high amounts of fragrance chemicals. Other cosmetic products contain relatively low amounts, 0.1-1% of fragrance compound, compared to up to 30% fragrance compound in EDT/EDP (226). The fragrance compound are mixtures of 20 to over 200 synthetic fragrance chemicals or natural fragrance materials (essential oils), selected from over 3,000 fragrance materials (226). For the exposure assessment, levels of fragrance chemicals in cosmetics containing significant amounts of fragrance materials (i.e. EDP/EDT/aftershave/deodorant) should be selected. It may not be possible to detect/measure the amounts of all fragrance chemicals when present in highly diluted form in a cosmetic product such as skin care products, make-up cosmetics etc. On the other hand, if a fragrance is evaluated safe for use when present in significant amounts in a product, it will also be safe for use in other products. Also the analysis of trend of the use of individual fragrance materials should be based on monitoring their contents in fine perfumes and deodorants.

Ninety of the 100 fragrance materials used in annual volumes > 175 tons in perfume formulations are fragrances and the remaining ten are used for other functions such as solvents or antioxidants (IFRA, personal communication 2010).

Among the 26 fragrances currently requiring individual labelling, amyl cinnamal, benzyl benzoate, benzyl salicylate, butyl phenyl methyl propional, citral, citronellol, coumarin, eugenol, geraniol, hexyl cinnamal, hydroxyisohexyl 3-cyclohexene carboxyaldehyde (HICC), alpha-isomethyl ionone, and linalool are used in volumes greater than 175 ton. α -Amylcinnamyl alcohol, anisyl alcohol, benzyl alcohol, benzyl cinnamate, cinnamal, cinnamyl alcohol, farnesol, hydroxycitronellal, isoeugenol, *d*-limonene, methyl-2-octynoate, oak moss (*Evernia prunastri*), tree moss (*Evernia furfuracea*) are used in volumes less than 175 ton.

According to the information from the fragrance industry, 80% of the total fragrance chemical volume is used in cosmetics and 20% in household products.

Since the implementation of the regulation of labelling of 26 fragrance substances in cosmetic products, qualitative information on fragrance exposure from cosmetics is provided in some market surveys performed on cosmetics (Table 10-1, (227)) and (Table 10-2, (228)) and on consumer products including cosmetics (Table 10-3, (229); Table 10-4, (115); and Figure 10-1, (105)). Thus, the implementation of the regulation of fragrance allergens in detergents (Directive 648/2004/EC), similar to that for cosmetics, has also added to the knowledge of fragrance exposure to the consumer. These market surveys revealed that fragrance ingredients which are potent allergens and frequently cause allergies in consumers are used as ingredients in consumer products including cosmetics. The results of these surveys further revealed that limonene and linalool were the most commonly used fragrance chemicals in cosmetics, while anisyl alcohol, cinnamal, α -amylcinnamyl alcohol, oak moss and tree moss were the least used fragrance ingredients in cosmetics and other consumer products. In general, the most potent allergens were also the most infrequently used ingredients. Prior to the regulation of the 26 allergens, analysis of 21 selected fragrance chemicals in deodorants also revealed additional 66 potential allergens in these products on the basis of structure activity relationship (230).

Opinion on fragrance allergens in cosmetic products

Table 10-1: Presence in children's cosmetics of the 26 fragrance substances that are required to be labelled in cosmetics (227).

| Fragrance substance | | % Products labelled to contain the fragrance substance |
|---|------------|--|
| INCI name | CAS number | |
| Amyl cinnamal | 122-40-7 | 8.2 |
| alpha-Amylcinnamyl alcohol | 101-85-9 | 2.9 |
| Anise alcohol | 105-13-5 | 0 |
| Benzyl alcohol | 100-51-6 | 9.6 |
| Benzyl benzoate | 120-51-4 | 9.1 |
| Benzyl cinnamate | 103-41-3 | 2.9 |
| Benzyl salicylate | 118-58-1 | 9.6 |
| Butyl phenyl methyl propional | 80-54-6 | 7.7 |
| Cinnamal | 104-55-2 | 1 |
| Cinnamyl alcohol | 104-54-1 | 6.7 |
| Citral | 5392-40-5 | 8.2 |
| Citronellol | 106-22-9 | 10.5 |
| Coumarin | 91-64-5 | 4.8 |
| Eugenol | 97-53-0 | 7.2 |
| Farnesol | 4602-84-0 | 2.9 |
| Geraniol | 106-24-1 | 12 |
| Hexyl cinnamal | 101-86-0 | 10.1 |
| Hydroxycitronellal | 107-75-5 | 6.3 |
| Hydroxyisohexyl-3-cyclohexene carboxyaldehyde | 31906-04-4 | 5.8 |
| Isoeugenol | 97-54-1 | 0.5 |
| Alpha-isomethyl ionone | 127-51-5 | 5.8 |
| <i>d</i> -Limonene | 5989-27-5 | 23.1 |
| Linalool | 78-70-6 | 21.6 |
| Methyl-2-octynoate | 111-12-6 | 0 |
| <i>Evernia prunastri</i> /oak moss | 90028-68-5 | 0 |
| <i>Evernia furfuracea</i> /tree moss | 90028-67-4 | 0 |

Opinion on fragrance allergens in cosmetic products

Table 10-2: Usage trends in deodorants of fragrance chemicals that are required to be labelled in cosmetics.

| Fragrance substance | | 88 products investigated in 2007 (228) | | | 70 products investigated in 1998 (231) | |
|-------------------------------|------------|--|---|------------|---|-------------|
| INCI name | CAS number | % Products labelled to contain the fragrance | Content in 23 selected products | | Content in all 70 products | |
| | | | % Products found to contain the fragrance | Range(ppm) | % Products found to contain the fragrance | Range (ppm) |
| Amyl cinnamal [▫] | 122-40-7 | 10.2 | 17 | 2.3-165 | 31 | 1-617 |
| alpha-amyl cinnamyl alcohol | 101-85-9 | - | - | - | n.a. | n.a. |
| Anise alcohol | 105-13-5 | 2.3 | 9 | 1, 51 | n.a. | n.a. |
| Benzyl alcohol | 100-51-6 | 17.1 | 26 | 32-166 | 76 | 1-629* |
| Benzyl benzoate | 120-51-4 | 25.0 | 48 | 3-4054 | 71 | 1-1075 |
| Benzyl cinnamate | 103-41-3 | 3.4 | 9 | 74, 143 | n.a. | n.a. |
| Benzyl salicylate | 118-58-1 | 39.8 | 48 | 136-5279 | 49 | 1-18758 |
| Butyl phenyl methyl propional | 80-54-6 | 48.9 | 70 | 1-5455 | 51 | 1-3732 |
| Cinnamal [▫] | 104-55-2 | 1.1 | 4 | 5 | 17 | 1-424 |
| Cinnamyl alcohol [▫] | 104-54-1 | 12.5 | 48 | 2-503 | 39 | 6-1169 |
| Citral [▫] | 5392-40-5 | 26.1 | 44 | 39-554 | n.a. | n.a. |
| Citronellol [▫] | 106-22-9 | 65.9 | 91 | 1-5848 | 81 | 1-5585 |
| Coumarin [▫] | 91-64-5 | 33.0 | 52 | 3.8-1255 | 57 | 1-1411 |
| Eugenol [▫] | 97-53-0 | 27.3 | 30 | 1-514 | 57 | 1-2355 |
| Farnesol [▫] | 4602-84-0 | 14.8 | 39 | 9-1791 | n.a. | n.a. |
| Geraniol [▫] | 106-24-1 | 48.9 | 87 | 1-399 | 76 | 1-1178 |

Opinion on fragrance allergens in cosmetic products

| Fragrance substance | | 88 products investigated in 2007 (228) | | | 70 products investigated in 1998 (231) | |
|---|------------|--|------|------------|--|--------|
| Hexyl cinnamal [▫] | 101-86-0 | 33.0 | 48 | 1-4434 | 71 | 2-1684 |
| Hydroxycitronellal [▫] | 107-75-5 | 27.3 | 70 | 1-1746 | 50 | 1-1023 |
| HICC [◻] | 31906-04-4 | 33.0 | 74 | 1-4431 | 53 | 1-1874 |
| Isoeugenol [▫] | 97-54-1 | 9.1 | 35 | 1-138 | 29 | 1-458 |
| Alpha-isomethyl ionone | 127-51-5 | 46.6 | 65 | 6-2588 | 61 | 1-2765 |
| D-Limonene [◊] | 5989-27-5 | 53.4 | 70 | 1022-11386 | n.a. | n.a. |
| Linalool [◊] | 78-70-6 | 53.4 | 96 | 8-3447 | 97 | 9-1927 |
| Methyl-2-octynoat [◊] | 111-12-6 | 1.1 | - | - | n.a. | n.a. |
| <i>Evernia prunastri</i> [▫] /oak moss | 90028-68-5 | 4.6 | n.a. | n.a. | n.a. | n.a. |
| <i>Evernia furfuracea</i> [▫] /tree moss | 90028-67-4 | 2.3 | n.a. | n.a. | n.a. | n.a. |

Notes: HICC Hydroxyisohexyl-3-cyclohexene carboxyaldehyde.

- Fragrance not detected in any product.

n.a. Not analysed.

* Benzyl alcohol could not be determined in 49% of the products due to interference.

The most common fragrance allergens are contained in the two mixtures, which are used for diagnosing fragrance allergy, called Fragrance Mix I (▫) and Fragrance Mix II (◻), besides the oxidation product of terpens (◊), and tree moss extract are common allergens. Methyl-2-octynoate is an extreme, but rare allergen.

Opinion on fragrance allergens in cosmetic products

Table 10-3: Frequency of occurrence in consumer products of the 26 fragrance allergens that are required to be labelled in cosmetics and detergents (229).

| INCI name of fragrance | PCP (n = 70) | MP (n = 59) | HP (n = 57) | WP (n = 44) | Cos (n = 39) | Deo (n = 17) | Dent (n = 14) | Total (n = 300) |
|------------------------------|-----------------|----------------|----------------|----------------|-----------------|-----------------|------------------|--------------------|
| Linalool | 46 | 47 | 17 | 42 | 26 | 12 | 0 | 190 (63%) |
| Limonene | 34 | 45 | 29 | 43 | 18 | 11 | 9 | 189 (63%) |
| Citronellol | 23 | 24 | 21 | 37 | 25 | 15 | 0 | 145 (48%) |
| Geraniol | 19 | 26 | 15 | 36 | 18 | 12 | 0 | 126 (42%) |
| BPMP | 30 | 27 | 21 | 27 | 13 | 8 | 0 | 126 (42%) |
| Hexyl cinnamal | 37 | 20 | 22 | 22 | 14 | 10 | 0 | 125 (42%) |
| Benzyl salicylate | 23 | 23 | 10 | 31 | 15 | 12 | 0 | 114 (38%) |
| Alpha-isomethyl ionone | 15 | 20 | 7 | 24 | 28 | 10 | 0 | 104 (35%) |
| Coumarin | 12 | 27 | 8 | 23 | 12 | 8 | 0 | 90 (30%) |
| Lyr TM | 17 | 24 | 3 | 24 | 15 | 5 | 0 | 88 (29%) |
| Eugenol | 13 | 26 | 4 | 22 | 6 | 6 | 3 | 80 (27%) |
| Citral | 2 | 28 | 6 | 29 | 7 | 2 | 0 | 74 (25%) |
| Benzyl benzoate | 8 | 9 | 3 | 31 | 11 | 8 | 0 | 70 (23%) |
| Benzyl alcohol | 9 | 8 | 1 | 30 | 9 | 3 | 1 | 61 (20%) |
| Hydroxycitronellal | 5 | 6 | 1 | 30 | 6 | 4 | 0 | 52 (17%) |
| Isoeugenol | 2 | 5 | 0 | 17 | 0 | 3 | 0 | 27 (9%) |
| Cinnamic alcohol | 4 | 2 | 0 | 13 | 4 | 2 | 0 | 25 (8%) |
| Farnesol | 1 | 3 | 0 | 17 | 2 | 0 | 0 | 23 (8%) |
| Amyl cinnamal | 5 | 0 | 3 | 7 | 5 | 2 | 0 | 22 (7%) |
| Cinnamal | 3 | 4 | 0 | 7 | 0 | 0 | 3 | 17 (6%) |
| Evermia prunastri/oak moss | 0 | 3 | 0 | 5 | 5 | 0 | 0 | 13 (4%) |
| Benzyl cinnamate | 2 | 0 | 0 | 8 | 0 | 0 | 0 | 10 (3%) |
| Evermia furfuracea/tree moss | 1 | 5 | 0 | 3 | 0 | 0 | 0 | 9 (3%) |
| Anisyl alcohol | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 (0.3%) |
| Amyl cinnamic alcohol | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| Methyl heptine carbonate | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |

INCI, International Nomenclature of Cosmetic Ingredients; PCP, personal care products; MP, men's products; HP, household products; WP, women's perfumes; Cos, cosmetics; Deo, deodorants; Dent, dental products; BPMP, butyl phenyl methyl propional; LyrTM, hydroxyisohexyl-3-cyclohexene carboxaldehyde.

Table 10-4: Frequency in 516 consumer products of the 26 fragrance substances that are required to be labelled in cosmetics* (115).

| Fragrance substance INCI name | % Product containing the chemical |
|---|-----------------------------------|
| D-Limonene | 48.3 |
| Linalool | 35.8 |
| Butyl phenyl methyl propional | 24.8 |
| Geraniol | 22.1 |
| Alpha-isomethyl ionone | 21.7 |
| Hexyl cinnamal | 21.3 |
| Citronellol | 21.1 |
| Benzyl salicylate | 18.6 |
| Coumarin | 17.0 |
| Eugenol | 15.7 |
| Benzyl alcohol | 15.3 |
| Benzyl benzoate | 14.7 |
| Hydroxyisohexyl-3-cyclohexene carboxyaldehyde | 12.8 |

Opinion on fragrance allergens in cosmetic products

| Fragrance substance INCI name | % Product containing the chemical |
|---------------------------------------|-----------------------------------|
| Citral | 11.6 |
| Hydroxycitronellal | 10.8 |
| Amyl Cinnamal | 7.9 |
| Anise alcohol | 7.0 |
| Cinnamyl alcohol | 6.4 |
| Farnesol | 3.9 |
| Isoeugenol | 3.1 |
| Cinnamal | 2.5 |
| Benzyl cinnamate | 2.3 |
| Amylcinnamyl alcohol | 1.9 |
| Methyl-2-octynoate | 1.0 |
| <i>Evernia prunastri</i> */oak moss | 0.8 |
| <i>Evernia furfuracea</i> */tree moss | 0.4 |

Note: * Consumer Products: Cosmetics and household products with labelling of the 26 fragrance allergens. The content of these fragrances was confirmed by chemical analysis.

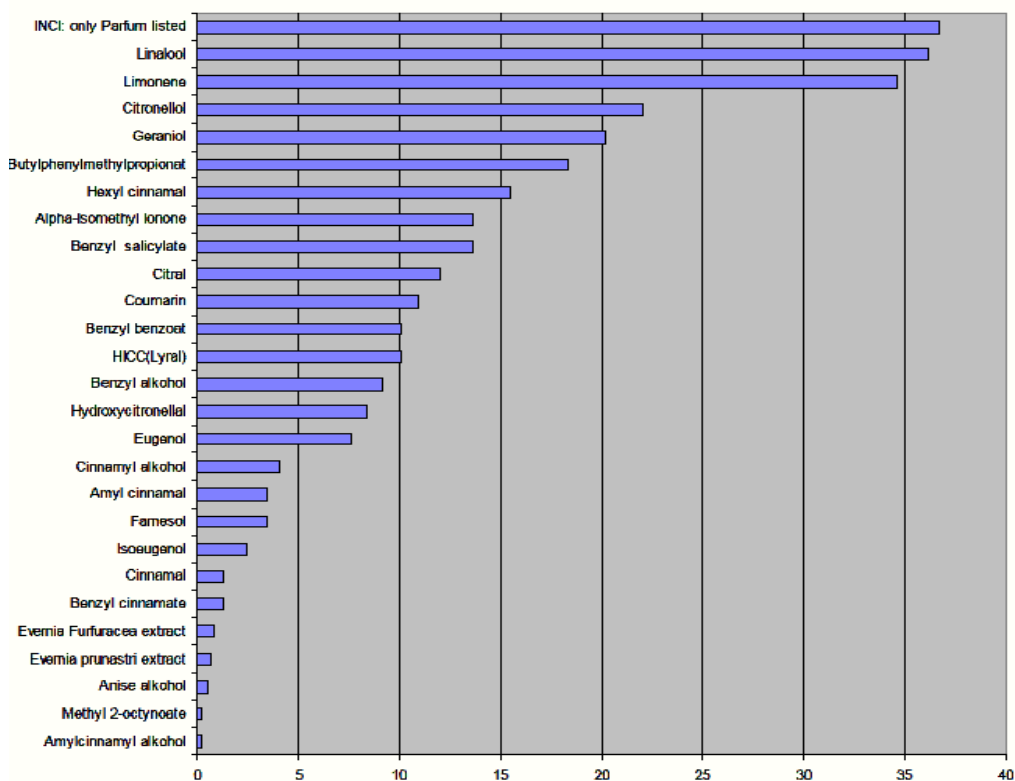


Figure 10-1: Frequency of occurrence in 3,000 consumer products of the 26 fragrance allergens that are required to be labelled in cosmetics and detergents (CVUA Karlsruhe, Germany, 2006/2007), according to (105).

Contents of fragrance substances determined in cosmetic products have been described in several studies, both before and after the regulation of the 26 fragrance allergens. The studies prior to the regulation of the 26 fragrance allergens included many, but not all of these 26 allergens. On the other hand, these studies included some other possible fragrance allergens. The quantitative analysis of fragrance substances has been performed in prestige perfumes (5, 157, 232-234), deodorants (228, 231), children's cosmetics and cosmetic toys (115, 227, 235), products marketed as natural cosmetics (225) and in cosmetics used by patients with contact allergy to fragranced products (35, 71). Quantitative analyses have revealed that the consumer is exposed to most, but not all of the 26 fragrance allergens from the use of cosmetics. However, when fragrance exposure from other consumer products, for example detergents and other household products is also taken into consideration (Table 10-3, Table 10-4, Figure 10-1), (105, 115, 229, 236), exposure to all of the 26 allergens is foreseeable in daily life. Although from the data available, the exposure to α -amylcinnamyl alcohol, cinnamal, methyl-2-octynoate, *Evernia prunastri* (oak moss) and tree moss may appear to be low, these are very strong allergens.

The changes in the use of fragrance chemicals in cosmetic formulations, during last 12 years, i.e. before and after the regulation of the 26 fragrance allergens, is reflected in the studies concerning contents of fragrances substances in popular perfumes (5, 232). As described in Table 10-5, the content of FM I allergens in prestige perfumes was significantly reduced from 1996 to 2003. Whether this is also the case for the perfumes sold as natural cosmetics (Table 10-6) has not yet been investigated.

Table 10-5: Concentration of Fragrance Mix I ingredients in five prestige perfumes before and after the regulation of the 26 fragrance allergens.

| Fragrance INCI name | Concentration in the perfumes before regulation (5) | | | Concentration in the perfumes after regulation (232) | | |
|---------------------|---|---------------|--------------|--|---------------|--------------|
| | In no. of perfumes | Range % (w/w) | Mean % (w/w) | In no. of perfumes | Range % (w/w) | Mean % (w/w) |
| Geraniol* | 5 | 0.072-0.432 | 0.340 | 5 | 0.090-0.236 | 0.156 |
| Cinnamal | 2 | 0.002-0.002 | 0.002 | 0 | - | - |
| Hydroxy-citronellal | 5 | 0.222-0.979 | 0.615 | 5 | 0.015-0.478 | 0.169 |
| Cinnamyl alcohol | 4 | 0.068-0.232 | 0.147 | 0 | - | - |
| Eugenol | 5 | 0.032-0.738 | 0.337 | 2 | 0.001, 0.001 | 0.001 |
| Isoeugenol | 3 | 0.026-0.249 | 0.119 | 2 | 0.001, 0.004 | 0.003 |
| Amyl cinnamal | 1 | 0.019 | 0.019 | 0 | - | - |

Note: * Due to interference by linalyl acetate, concentration of geraniol+linalyl acetate is reported.

Table 10-6: Concentrations of Fragrance Mix I ingredients, hexyl cinnamal and coumarin in 22 perfumes marketed as natural cosmetics investigated in 1996.

| Fragrance | In no. of perfumes | Concentration % (w/w) |
|--------------------|---------------------------|------------------------------|
| Geraniol | 14 | 1.191* |
| Cinnamal | 3 | 0.089, 0.109, 2.101 |
| Hydroxycitronellal | 5 | 0.135-6.044 |
| Cinnamyl alcohol | 8 | 0.035-2.289 |
| Eugenol | 2 | 0.027, 0.139 |
| Isoeugenol | 8 | 0.194-3.039 |
| Amyl cinnamal | 9 | 0.105-7.706 |
| Coumarin | 11 | 0.046-6.043 |

Note: * Quantification was performed in one sample only, due to interference by a very large amount of linalyl acetate in other samples.

The trend in the use of most of the fragrance allergens in deodorants before and after their regulation is reflected by the two studies performed by Rastogi et al. (228, 231). The results of these studies cannot be directly compared, because the study from 1998 included randomly selected deodorants, while selection of the deodorants for the 2007 study was based on the labelling of the presence of known strong fragrance allergens in these products. The number of products analysed in the 1998 study were three times more than those analysed in 2007, but not all of the 26 fragrance allergens were analysed in the 1997 study. However, an indication of the change in the use of the fragrance allergens during 1998-2007 may be obtained by reviewing the results of these two studies. Among the 17 common fragrance substances studied in the two studies, the frequency of use of 16 of these substances in deodorants was reduced in 2007 compared to that in 1998 (Table 10-2). The frequency of use of butyl phenyl methyl propional in deodorants appeared to be unchanged. The contents of benzyl alcohol, benzyl salicylate, cinnamal, cinnamyl alcohol, eugenol, geraniol, isoeugenol and linalool were found to be lower in the deodorants analysed in 2007 compared to those in 1998. Citronellol, coumarin and alpha-isomethylionone contents in the deodorants were similar in both studies, but concentrations of benzyl benzoate, butyl phenyl methyl propional, hexyl cinnamal, hydroxyisohexyl-3-cyclohexene carboxyaldehyde and linalool were much higher in deodorants in 2007 compared to those in 1998. This analysis of trend of use of fragrance allergens in cosmetic products indicates that the regulated fragrance allergens are used less frequently, but exposures from some of the regulated fragrance allergens may be much higher compared to those before regulation.

Table 10-7: Atranol and chloroatranol content in eau de toilette/eau de perfume, investigated in 2004 and in 2007.

| | 2007 Study | 2004 Study |
|---|-------------------|-------------------|
| No. of samples | 22 | 17 |
| Atranol present in no. of samples | 15 (68%) | 12 (70%) |
| Atranol content | ppb (ng/ml) | ppb (ng/ml) |
| Range | n.d.-880 | n.d.-791 |
| Mean±SD | 157±249 | 97±224 |
| Median | 47 | 20 |
| Chloroatranol present in no. of samples | 9 (41%)* | 14 (82%) |
| Atranol content | ppb (ng/ml) | Ppb (ng/ml) |
| Range | 0.9-208 | 1-175 |
| Mean±SD | 63±73 | 36±51 |
| Median | 22 | 10 |

Notes: n.d. Not detected.

* $P < 0.05$ (chi-square test).

SD: Standard deviation.

Atranol (CAS no. 526-37-4) and chloroatranol (CAS no. 57074-21-2), constituents of oak moss and tree moss have been shown to be very potent fragrance allergens (237, 238). The EC Scientific Committee on Consumer Products (SCCP) recommended that atranol and chloroatranol should not be present in cosmetic products (239). Two other commonly used fragrance chemicals, isoeugenol (240) and hydroxyisohexyl-3-cyclohexene carboxyaldehyde (HICC) (71), have also been shown to be important contact allergens. The contents of atranol, chloroatranol, isoeugenol and hydroxyisohexyl-3-cyclohexene carboxyaldehyde in fine fragrances was determined for the exposure assessment of these fragrances (233). The results revealed that isoeugenol was present in 56%, HICC in 72%, atranol in 59%, and chloroatranol in 36% of the 22 eau de toilette/eau de parfum products. The concentrations of isoeugenol were, in all products, below 0.02% which is the maximum concentration recommended by the fragrance industry. HICC reached a maximum concentration of 0.2%, which is 10-fold higher than the maximum tolerable concentration considered safe by the EC Scientific Committee (241). The concentrations of atranol and chloroatranol in the products investigated in 2007 were comparable to those found in similar products in 2004 (Table 10-7, (233, 234). A significant decrease in the frequency of the presence of chloroatranol in the products was found in 2007 (Table 10-7).

10.2. Global exposure (household and occupational exposures)

Fragrances are used in cosmetics that the consumer applies to themselves, as described in the previous section. In addition, exposure to fragrance substances is possible by a number of other exposure routes briefly outlined in this section.

Topical pharmaceutical products

In a study from Belgium, 370 of the 3,280 topical products marketed in Belgium have been found to contain one or more of 66 fragrance substances (242). This publication also contains a description of causative fragrance allergens in 127 patients reacting to 48 specific topical products. In a broader sense, exposure of the patient by extracts used in aromatherapy falls in this category as well.

Childrens products and toys

Children's products may contain fragrance allergens and high levels may be present (235). It has been stated that children may become sensitised to fragrance chemicals used by their mothers (243).

Clothing

Washed fabrics have been reported to contain fragrances (244). Odour-neutralising agents are sometimes used for shoe insoles. In one case, an insole containing cinnamon, has been reported to lead to plantar vesicular contact dermatitis due to contact sensitisation to FM I and, in the breakdown, to cinnamal and cinnamyl alcohol (245).

Cleaning agents and other household products

Contact dermatitis from geraniol in washing-up liquid has been reported (246). Terpenes are used as solvents and cleansing agents (e.g. limonene) (247) and have been reported as cause of hand dermatitis (248, 249). In an analysis of 59 household products the most common fragrance allergens were limonene (78%), linalool (61%) and citronellol (47%) (250). In a review of 301 cosmetic and detergent consumer products in Sweden, in half of the cosmetics and one-third of the detergents, one or more of the 26 fragrances requiring labelling were identified (251). In the UK, a review of 300 consumer products showed that linalool and limonene were present in 63% of products. Dental products contained on average 1.1 fragrance substances that are presently required to be labelled and women's perfumes contained 12 of these fragrance substances (Table 4-1 and Table 4-3) (229).

Candles

The dermal hand transfer of three fragrance materials (cinnamic aldehyde, d-limonene and eugenol) from scented candles was determined in ten subjects (i.e. 20 hands) after grasping scented candles for five consecutive 20 second exposures/grasps. The total mean residues of cinnamal and eugenol transferred per grasp from the candles to the hands were 0.255 µg/cm(2) and 0.279 µg/cm(2), respectively (252).

Food

Food causing cheilitis or bullous stomatitis (e.g. due to cinnamal (253)) or lichen planus-like lesions (e.g. due to cinnamal (254)) or contact gingivitis (e.g. due to eugenol (255)) has been reported. Moreover, food containing fragrance allergens, e.g. citrus oil terpenes (256) may cause allergic contact dermatitis by handling this food.

Occupational exposure

In a number of occupations, contact allergy to fragrances is more common than in others, including geriatric nurses, masseurs and physiotherapists, metal furnace operators and potters/glass makers, according to a multifactorial analysis (90). Moreover, hairdressers, beauty therapists and aroma therapists are examples of occupations where there is occupational exposure to fragrance-containing cosmetic and other products. Cleaners are exposed to fragrance-containing household products (e.g. detergents). Cooks and bakers are exposed to flavour chemicals and spices. Healthcare workers are also at risk of acquiring fragrance contact allergy. "Odour maskers" may contain important fragrance allergens (89, 90, 257-259). Occupational exposure and

occupational ACD to fragrances have been described in perfume bottlers (260). Industrial use of a powder masking the vinyl smell of car seats, containing cinnamal, causing occupational ACD has been reported (259).

A number of fragrance chemicals are also used as biocides (see Commission Regulation (EC) No 1451/2007 of 4 December 2007 on the second phase of the 10-year work programme referred to in Article 16(2) of Directive 98/8/EC of the European Parliament and of the Council concerning the placing of biocidal products on the market, published 11.12.2007 EN Official Journal of the European Union L 325/3 –L325/65), see Table 10-8 below.

Table 10-8: Parts of Annex I to (EC) No 1451/2007 (see above): "Active substances identified as existing", if use is 'perfuming' or 'masking' according to CosIng.

| Biocide | EINECS | CAS number | Biocidal product group |
|--|---------------|-------------------|-------------------------------|
| Linalool | 201-134-4 | 78-70-6 | 19 |
| Geraniol | 203-377-1 | 106-24-1 | 18, 19 |
| Benzyl benzoate | 204-402-9 | 120-51-4 | 2, 18 |
| Eugenol | 202-589-1 | 97-53-0 | Not given |
| Farnesol | 225-004-1 | 4602-84-0 | Not given |
| (R)-p-mentha-1,8-diene | 227-813-5 | 5989-27-5 | 12 |
| Citriodiol/mixture of cis- and trans-p-menthane-3,8 diol | 255-953-7 | 42822-86-6 | 1, 2, 19 |
| Citral | 226-394-6 | 5392-40-5 | Not given |
| Pine ext. | 304-455-9 | 94266-48-5 | 10 |
| TANACETUM CINERARIIFOLIUM FLOWER EXTRACT | 289-699-3 | 89997-63-7 | 18 |
| Citrus oils (main component: limonene) | several | various | |
| Clove oil (main component: eugenol (83.8 %), caryophyllene (12.4 %)) | / | 8000-34-8 | |

Product groups(According to Biocide Directive 98/8/EC)

- 1 Human hygiene biocidal products
- 2 Private area and public health area disinfectants and other biocidal products
- 3 Veterinary hygiene biocidal products
- 10 Masonry preservatives
- 12 Slimicides
- 18 Insecticides, acaricides and products to control other arthropods
- 19 Repellents and attractants

The above illustrates that the consumer is exposed to fragrance substances from a wide variety of cosmetic products, other consumer products, pharmaceuticals and occupational exposures.

All these exposures are of importance in the context of contact allergy as it is not the source of exposure that is critical for both induction and elicitation, but the cumulative dose per unit area.

10.3. Exposures related to particular anatomical sites

Contact allergy to fragrances most often causes dermatitis of the hands, face and axillae. Axillary involvement has been shown to be statistically related to fragrance allergy (9). It is recognised that the axillary skin is a problematic area as it is moist, occluded and is easily irritated. Moreover, facial eczema is a common manifestation of fragrance allergy (3, 47). There is an association between fragrance allergy and hand eczema or aggravation of hand eczema (13-15). Vehicles may influence elicitation capacity of an allergen and the presence of detergents (surfactants) as in hand cleaning products may increase the clinical response by a factor of 4-6 (261). Men using wet shaving as opposed to electric razors have an increased risk of being fragrance allergic (17), most likely due to microtraumata and to the presence of surface active substances in shaving foam.

In use tests, the upper arm has been shown to be more sensitive than the forehead and lower arm (262). The axillae, neck and face are more sensitive than the upper arms (10). The threshold of elicitation may vary depending on the volatility of the substance (263). A cumulative effect of exposures occurs so that repeating exposures cause elicitation in more individuals (264).

Patients appear to become sensitised to fragrances primarily from deodorants and perfumes and to a lesser extent from other cosmetic types (74). Allergic contact dermatitis may develop where a perfume has been applied (behind ears, neck, upper chest, antecubital fossae, wrists and the axillae bilaterally (265). Following this, eczema may appear, or be worsened by, the use of a variety of product types including other cosmetics, household products, industrial products and flavours.

The association between contact allergy to fragrance ingredients and certain anatomical sites, which mirrors exposure to fragrance-containing products on these anatomical sites, has been described in several publications (266, 267), see above. However, due to the potential confounding effect of other factors, at least on some anatomical sites, an adjusted analysis will provide a more valid impression of the association between certain anatomical sites and contact allergy to fragrance ingredients. As an adjusted, multifactorial analysis relies on: (i) a substantial number of observations (patients tested); and (ii) an outcome prevalence not too close to 0%, such an approach has, hitherto, been limited to FM I.

In a paper published 2001, data from the IVDK in terms of patch test reactions to FM I and relevant clinical and demographic information of the patients tested (n=57,779) was studied by Poisson regression analysis (90). Risk was quantified by the prevalence ratio, which can be interpreted as an estimate of relative risk, i.e. the factor by which the risk of being sensitised to FM I (in this example) is to be multiplied (RR > 1: elevated risk; or RR < 1: reduced risk) if a certain "risk factor" is present, compared to those patients in whom this risk factor is not present (the reference category) (general aspects of such analyses are discussed in (268)). In the analysis, potential risk factors and confounders, respectively, including occupation, year of patch testing (to address a possible time trend), sex, age, past or current atopic dermatitis, in addition to anatomical site. The relevant part of Table 3 of (90) is reproduced below.

Table 10-9: Result of a Poisson regression analysis of patients tested with the Fragrance Mix between January 1992 and December 1998, considering two alternative outcomes – part I: non-occupational factors

| Attribute | Prevalence (%) | At least + (11.5%) | | At least ++ (4.0%) | |
|--------------|----------------|--------------------|--------------|--------------------|--------------|
| | | PR | 95% CI | PR | 95% CI |
| Age: | | | | | |
| ≤30 | 26.7 | 1.00 | Reference | 1.00 | Reference |
| >30–44 | 23.8 | 1.42 | 1.31 to 1.53 | 1.61 | 1.40 to 1.84 |
| >44–58 | 25.6 | 1.67 | 1.55 to 1.80 | 1.90 | 1.66 to 2.16 |
| >58 | 23.9 | 1.93 | 1.77 to 2.10 | 2.07 | 1.79 to 2.39 |
| Sex (female) | 64.5 | 1.29 | 1.21 to 1.37 | 1.18 | 1.07 to 1.31 |
| Main site:* | | | | | |
| Trunk | 2.9 | 1.00 | Reference | 1.00 | Reference |
| Hands | 29.9 | 1.24 | 1.07 to 1.46 | 1.28 | 0.98 to 1.67 |
| Arm | 3.8 | 1.23 | 1.01 to 1.49 | 1.19 | 0.86 to 1.65 |
| Face | 15.2 | 1.20 | 1.03 to 1.42 | 1.13 | 0.86 to 1.48 |
| Neck | 1.4 | 1.39 | 1.10 to 1.75 | 1.31 | 0.88 to 1.94 |
| Feet | 2.8 | 1.26 | 1.02 to 1.55 | 1.19 | 0.84 to 1.68 |
| Leg | 8.7 | 1.59 | 1.36 to 1.89 | 1.50 | 1.14 to 1.99 |
| Axilla | 0.9 | 2.77 | 2.20 to 3.46 | 2.73 | 1.87 to 4.00 |
| Other site | 8.9 | 0.66 | 0.55 to 0.80 | 0.48 | 0.35 to 0.67 |

*Additionally controlled for several more sites—none of these associated with a significantly increased or decreased risk.

Compared to the trunk, which was arbitrarily chosen as the reference category, all other anatomical sites are associated with an increased risk of being sensitised to FM I (significantly if the lower limit of 95% CI is > 1). Most evidently, dermatitis of the axilla(e) is strongly associated with contact allergy to FM I, presumably due to the application of deodorants. Furthermore, the part of the table shown above illustrates a strong, positive age gradient, i.e. the older patients are, the more likely they are to be sensitised to FM I, the risk being almost double when comparing the oldest with the youngest age group. This observation is in concordance with a bivariate (unadjusted) association between age and contact allergy to FM I found in another study (89). This association is presumably the result of life long exposures and cumulative risk.

In a similar analysis of *Myroxylon pereirae* resin, published in 2002 (269): (i) an even stronger age gradient; and (ii) no particular association to axillary dermatitis (included in the “other” category) was found (Table 10-10).

Table 10-10: Association between selected risk factors and positive patch test to *Myroxylon pereirae* resin. For full model see (269). Risk quantified with the prevalence ratio (PR) with accompanying 95% confidence interval (CI).

| Factor | PR | 95% CI |
|------------------------------------|-----------|---------------|
| Atopic dermatitis, past or present | 1.02 | (0.95-1.10) |
| Female sex | 1.13 | (1.06-1.20) |
| <i>Site</i> | | |
| Trunk | 1.00 | (reference) |
| Hand or Arm | 1.03 | (0.94-1.12) |
| Foot or Leg | 1.76 | (1.61-1.92) |
| Head or Neck | 0.94 | (0.86-1.03) |
| "Other" site | 0.72 | (0.64-0.81) |
| Missing site | 1.07 | (0.97-1.19) |
| <i>Age</i> | | |
| 30 years and younger | 1.00 | (reference) |
| 31 to 44 | 1.92 | (1.73-2.12) |
| 45 to 58 | 2.87 | (2.61-3.16) |
| 58 or older | 3.85 | (3.49-4.25) |

10.4. Conclusion

There are various modes of exposure to fragrances, including not only products used for their scent, such as perfumes and eau de toilette, after shaves, and deodorants, but also types of products where scent is an added feature, such as other cosmetic categories (including wipes), topical pharmaceuticals, household products, and products encountered in the occupational setting.

Consumer exposure can change over time, both qualitatively and quantitatively.

Different routes of exposure are reflected by certain anatomical sites affected: deodorants are associated with axillary dermatitis, the axillary skin being particularly vulnerable to sensitisation due to occlusion, maceration and irritation. However, while sensitisation and initial disease may follow a distinct pattern, later less specific exposures, e.g. via hand creams, cleaning lotions etc. may be sufficient to cause allergic contact dermatitis.

11. Dose-response relationships and thresholds

The dose-response relationship between exposure to contact allergens and induction of allergy, i.e. sensitisation, is well established in animal models and by experiments in healthy volunteers (270). It seems that not only the dose per unit area of allergen (271), but also the number of exposures, i.e. the accumulated dose, is of importance for the risk of induction of contact allergy (272). The induction of contact allergy is an immunological process (type IV-allergy), which is without any clinical symptoms. In the case of continued exposure or re-exposure with a sufficient dose of allergen, elicitation will occur. Elicitation is an inflammatory response (eczema) with clinical symptoms of erythema, induration and in some cases vesicles. Studies of the elicitation response are normally done in patients with an allergy to the substance in question. Different provocation models exist (see chapter 11.2.1). Elicitation experiments in healthy human volunteers following the induction have only rarely been performed (273) and may be considered a less valid model than patient studies. The reason is that following experimental induction, the level of sensitivity may not be at the same level as in a real life situation and that individuals who have actually acquired the disease are a more relevant endpoint to study.

Knowledge of the dose-response relationship provides an opportunity to establish levels of exposure which are safe for the majority of individuals. In the following chapter, the use of different data and models for the establishment of such safe levels in relation to fragrance ingredients are explored. The focus will be on those chemicals, which have been identified in chapter 7.1 as established contact allergens in humans and which have already given rise to a significant number of published cases (category 3 or more): cinnamal, cinnamyl alcohol, citral, coumarin, eugenol, farnesol, geraniol, hydroxycitronellal, isoeugenol. Limonene and linalool are considered in chapter 5 as their ability to cause sensitisation depends on air oxidation, and hydroxyisohexyl 3-cyclohexene carboxaldehyde is considered in chapter 4.3.2 and 11.4.

11.1. Induction

A model for dermal sensitisation quantitative risk assessment (QRA) has been developed and implemented by the fragrance industry. This model relies on thresholds, no effect or low-effect levels, established in healthy human volunteers and/or in animal experiments, mainly the local lymph node assay (LLNA) (see chapter 8.1). A set of safety factors are applied for inter-individual differences, for vehicle effects and for use considerations, stated to give rise to a safety margin from 10 to 1000 (274). In this way, a so-called "acceptable exposure level" is derived. The exposure to an allergen in different types of products should be below this level. The restrictions, which have been introduced by the fragrance industry based on the QRA model, are given in Table 11-1 for some important product categories.

The IFRA guidelines give concentration limits for 11 product categories (http://www.ifraorg.org/en-us/standards_1, last accessed 2011-11-02), three of which are mentioned in Table 11-1. These three products have the lowest concentrations except for lip products, which give a slightly lower concentration limit.

Table 11-1: Current IFRA restrictions based on induction experiments.

| Fragrance chemicals | IFRA guideline ¹ | | |
|---------------------------------|-----------------------------|----------------|-------------|
| | Deodorant (%) | Hand cream (%) | Perfume (%) |
| Cinnamal | 0.02 | 0.05 | 0.05 |
| Cinnamyl alcohol | 0.1 | 0.4 | 0.4 |
| Citral | 0.05 | 0.3 | 0.6 |
| Coumarin | 0.13 | 0.8 | 1.6 |
| Eugenol | 0.2 | 0.5 | 0.5 |
| Farnesol | 0.11 | 0.6 | 1.2 |
| Geraniol | 0.4 | 2.8 | 5.3 |
| Hydroxycitronellal ² | 0.2 | 1.0 | 1.0 |
| Isoeugenol ² | 0.01 | 0.02 | 0.02 |

Notes: 1) Exposure per mg/cm²/day is based on 8.5 mg/cm²/day for deodorants, 2.2 for perfumes and 4.2 for hand creams as it is these exposure levels that are used by the IFRA.

2) Cosmetic Directive Annex III: Hydroxycitronellal restricted to 1% in all products and isoeugenol to 0.02% in all products.

The SCCP evaluated this methodology (275) as well as its application to three model fragrance substances.

It was, among other things, concluded that:

“The data provided show that the application of the dermal sensitisation QRA approach would allow increased exposures to allergens already known to cause allergic contact dermatitis in consumers. The model has not been validated and no strategy of validation has been suggested. There is no confidence that the levels of skin sensitisers identified by the dermal sensitisation QRA are safe for the consumer.”

and that:

“Identification of safe levels of exposure to existing substances known to cause allergic contact dermatitis in the consumer should be based on clinical data and/or elicitation low-effect levels. Currently, these are the only methods which have proven efficient in reducing/preventing existing problems of sensitisation/allergic contact dermatitis in the consumer.”

11.2. Elicitation

11.2.1. General considerations

A response in terms of elicitation of allergic contact dermatitis by application of the (suspected) allergen under standardised conditions is the outcome of interest of the routine diagnostic procedure for suspected contact allergy, the patch test. While the patch test procedure is largely standardised, it is optimised as a diagnostic tool for contact allergy. Thus exposure conditions are not comparable to actual exposures occurring in the daily life or working environment of the patient, which often involve long-term, repeated and low-dose contact with the allergen. Here, procedures such as the repeated open application test (ROAT) or provocative use test are often used, because they reflect actual exposure much better and can be used, for instance, to validate the current clinical relevance of a positive PT reaction.

Generally, exposure of a sensitised patient to a set of graded doses (quantity/area) of the suspected allergen, i.e. threshold testing, will allow not only quantitative diagnosis of the presence or absence of specific contact sensitisation but will additionally provide evidence on the intensity (degree) of sensitisation. This may have important individual consequences in terms of everyday or occupational exposures being capable (or not) of eliciting allergic contact dermatitis. However, beyond the individual perspective, clinical dose-response data collected from sensitised individuals provide a valuable estimate of the usual doses/unit area resulting in a positive, allergic response in a certain proportion of sensitised persons, e.g. 10, 50 or 90%. Maximum concentration levels can be derived, which are safe in terms of eliciting allergic reactions in only a defined low percentage of sensitised persons. As such data will always be based on small samples, the precision of the estimate should be considered, and therefore results are preferably given with confidence intervals.

A statistically significant relationship between threshold concentrations in the ROAT and patch test has been found, on analysing results from different allergens (see Table 11-2) (276), but the dose of allergen per unit area per application needed to elicit a reaction in the two study methods is not the same. A translation factor between the two methods has been suggested for non-volatile substances: $ED_{xx}(ROAT) = 0.0296 * ED_{xx}(\text{patch test})$ based on testing nickel and methyl dibromo glutaronitrile (276). Based on this the eliciting dose per application in an open test is 33 times lower than in the patch test. In practice it means that the cumulative dose in a ROAT (in $\mu\text{g}/\text{cm}^2$) in two weeks with two applications per day (total 28 applications) will be almost identical to the eliciting patch test dose (in $\mu\text{g}/\text{cm}^2$) for a given number of responders (see Figure 11-1). For a given cut-off point the elicitation dose determined by patch testing will be higher than determined by ROATs.

Table 11-2: Spearman's rank correlation between the threshold concentration in the patch test and the repeated open application test for three allergens.

| Allergen | Number of patients | Correlation coefficient | P-value |
|----------|--------------------|-------------------------|---------|
| Nickel | 18 | 0.45 | 0.033 |
| MDBGN | 15 | 0.76 | 0.0021 |
| HICC | 16 | 0.59 | 0.011 |

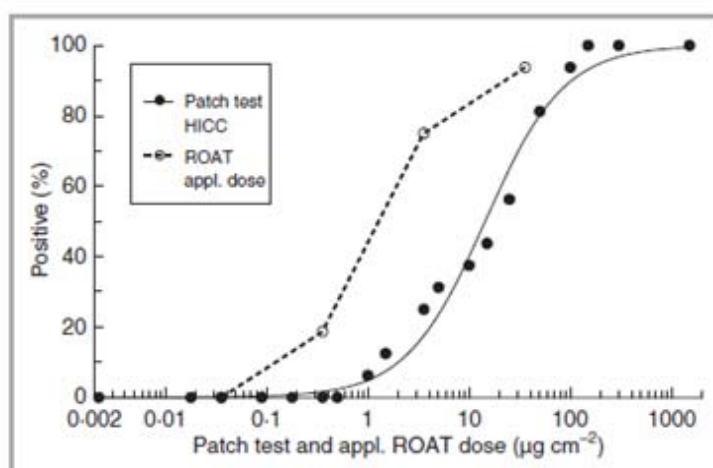


Figure 11-1: The fitted dose-response curve for patch test (solid line) is seen to be displaced to the right compared to the observed response from repeated open applications of the same allergen (HICC). It means that a smaller dose per application is needed to elicit a response than by one single occluded application as in the patch test.

In the translation between methods, evaporation needs to be taken into consideration for volatile substances. The experience, based on a study of the fragrance ingredient HICC and using the results from the literature on isoeugenol, is that if the same equation is used as for non-volatile substances, the response in the ROAT will be overestimated by a factor 3 to 4. Thus, the translation factor would be 0.1060 instead of 0.0296, but this needs to be confirmed by other fragrance allergens. This implies that for the fragrance ingredients tested, the eliciting dose per application in a ROAT was 9.4 times lower than the patch test compared to a 33 times lower dose for non-volatile substances (276). This needs to be confirmed by studying other fragrance allergens. Thus, according to these experiments, the dose ($\mu\text{g}/\text{cm}^2$) eliciting a response in threshold patch testing will be at most 33 times higher than established in the ROAT if an identical vehicle is used.

Volatility effects in skin sensitisation

The potency of volatile skin sensitisers can be underestimated, to an extent depending on how rapidly it evaporates, by assays such as the LLNA in which the test substance is applied topically to exposed healthy skin without occlusion. Such sensitisers present a greater sensitisation risk to consumers when the skin is occluded by clothing and/or compromised, than when healthy non-occluded skin is exposed.

Volatility at physiological temperature, say 40°C , is represented by the vapour pressure p_{40} at that temperature. This is related to the boiling point T_B by the Clapeyron-Clausius equation, which can be written (277):

$$\text{Log}(p_{40}) = - (T_B - 40)\text{Tr}/2.303RT$$

Where p is in atmospheres, T_B is in $^\circ\text{C}$, R is the gas constant, Tr is the Trouton constant (also defined as the molar entropy of vaporisation, and equal to $22 \text{ cal}\cdot\text{deg}^{-1}$ for many organic compounds) and T is physiological temperature in degrees absolute (= 313 for 40°C).

It has been shown, in experiments where evaporation from a glass slide is measured under simulated LLNA conditions, that 2-hexenal ($T_B = 146\text{-}149^\circ\text{C}$, $p_{40} = 17 \text{ mmHg}$) evaporates rapidly, less than 20% remaining after 5 minutes, whereas with cinnamal ($T_B = 248^\circ\text{C}$, $p_{40} = 0.5 \text{ mmHg}$), more than 90% remains after 1 hour (278). In agreement with these findings, cinnamal fits a QSAR relating LLNA EC3 to reactivity, whereas the EC3 for 2-hexenal is higher (lower potency) than predicted from its reactivity.

The above is only a partial rationalisation, since different solubilities in different vehicles will influence the tendency to evaporate, according to Henry's law.

11.2.2. Studies on specific fragrance ingredients

Studies concerning chloroatranol/atranol, cinnamal, hydroxycitronellal, hydroxyisohexyl 3-cyclohexenecarboxaldehyde and isoeugenol have been identified. These are summarised in Annex III.

Overview of results

In four studies dummy deodorants spiked with a single fragrance allergen in realistic use concentrations have been used to study elicitation responses, unscented deodorants were used as control products in paired designs. The deodorants were used by patients sensitised to the fragrance allergen in question as well as a healthy control group

Opinion on fragrance allergens in cosmetic products

(without fragrance allergy) (102,103,104,279). Between 76 and 100% of the sensitised individuals reacted to the deodorants spiked with allergen, isoeugenol, cinnamal, hydroxycitronellal and hydroxyisohexyl 3-cyclohexene carboxaldehyde, and none of the controls (Table 11-4).

Table 11-3: Overview of results of deodorant provocation investigations with different allergens. Frequency in % of test groups, which reacted at different doses of allergen applied in a roll-on deodorant in the axilla, is given in the table.

| Dose in ppm in deodorant | Isoeugenol | Cinnamal (1) | Cinnamal (2) | Hydroxycitronellal | HICC |
|---|--------------------|--------------------|--------------|--------------------|---------------------|
| 0 | 0 | 0 | 0 | 0 | 0 |
| 63 | 23 | | | | |
| 100 | | | 11 | | |
| 200 | 69 | | | | 64 |
| 320 | | 25 | 55 | 57 | |
| 600 | | | | | 85 |
| 630 | 76 | | | | |
| 1000 | | 75 | 88 | 71 | |
| 1800 | | | | | 100 |
| 3200 | | 100 | | 100 | |
| No. test persons | 13 | 8 | 9 | 7 | 14 |
| No. of control persons | 10 | 20 | | 7 | 10 |
| % control persons, who reacted | 0 | 0 | | 0 | 0 |
| Exposure according to study should be: | < 63 ppm | <100 ppm | | <320 ppm | < 200 ppm |
| Reference | (279) | (103) | | (104) | (102) |

Note: HICC hydroxyisohexyl 3-cyclohexene carboxaldehyde.

Eleven studies concerning dose-response results of the five allergens listed above were identified, including the above mentioned studies of deodorants. An overview of the results of the studies concerning thresholds is given in Table 11-4. In Annex III the details of each study are given.

Table 11-4: Overview of threshold results from clinical studies.

“Observed” means that the proportion was actually observed in the study while “estimated” means that the value is derived from a fitted curve, i.e. is interpolated.

| Chloroatranol | | | |
|--------------------------|--------------------------|----------|-------|
| ROAT | | | Ref. |
| In ethanol 92 % positive | 0.025 µg/cm ² | observed | (238) |
| In ethanol 100% positive | 0.125 µg/cm ² | observed | (238) |
| PATCH TEST | | | |

Opinion on fragrance allergens in cosmetic products

| | | | |
|--|--|--------------|-------|
| ED10% | 0.0004 µg/cm ² | estimated | (238) |
| ED50% | 0.0045 µg/cm ² | estimated | (238) |
| Cinnamal | | | |
| ROAT | | | |
| In ethanol no effect | 0.02% | observed | (101) |
| In ethanol 44 % positive | 0.1% | observed | (101) |
| In ethanol 72 % positive | 0.8% | observed | (101) |
| Deodorant matrix 11% positive | 0.26 µg/cm ² (0.01%) | observed | (103) |
| Deodorant matrix 41% positive | 0.84 µg/cm ² (0.032%) | observed | (103) |
| Deodorant matrix 82% positive | 2.63 µg/cm ² (0.1%) | observed | (103) |
| PATCH TEST | | | |
| ED50% | 96 µg/cm ² | estimated | (101) |
| No effect level | 0.4 µg/cm ² (0.01%) | observed | (101) |
| No effect level | NG (0.002%) | observed | (103) |
| HICC | | | |
| ROAT | | | |
| In a cream base ED10% | 4.9 µg/cm ² | interpolated | (105) |
| In a perfume (ethanol) ED10% | 1.2 µg/cm ² | interpolated | (105) |
| In ethanol 61% positive | 15.3 µg/cm ² (3.4-22.2) | observed | (224) |
| In ethanol 89% positive | 126.2 µg/cm ² (40.5-226.2) | observed | (224) |
| In ethanol/water no response | 0.0357 µg/cm ² | observed | (263) |
| In ethanol/water ED10% | 0.064 µg/cm ² | estimated | (263) |
| In deodorant matrix between 64% to 100% positive | 0.79 µg/cm ² (median) | observed | (102) |
| PATCH TEST | | | |
| ED10% (95% CI) | 0.662 µg/cm ² (0.052-2.35) | estimated | (263) |
| ED10% | 0.75 µg/cm ² | estimated | (102) |
| ED10% | 0.9 µg/cm ² 29 (7-69) ppm | estimated | (224) |
| ED50% (95% CI) | 11.1 µg/cm ² (3.41- 33.1) | estimated | (263) |
| ED50% (95% CI) | 18.3 µg/cm ² (3.41- 33.1) | estimated | (102) |
| ED50% (95% CI) | 20 µg/cm ² 662 (350-1250) ppm | estimated | (224) |
| No effect level | <0.0022 µg/cm ² | observed | (263) |
| Hydroxycitronellal | | | |
| ROAT | | | |
| Deodorant matrix 57 % positive | 0.94 µg/cm ² (0.032%) | observed | (104) |
| Deodorant matrix 71 % positive | 2.94 µg/cm ² (0.1%) | observed | (104) |
| Deodorant matrix 100 % positive | 9.40 µg/cm ² (0.32%) | observed | (104) |
| PATCH TEST | | | |

| | | | |
|--------------------------------|--|-----------|-------|
| No effect level | <0.00012 % (=0.036 µg/cm ²)* (*calculated) | observed | (104) |
| Isoeugenol | | | |
| ROAT | | | |
| in ethanol 63% positive | 5.6 µg/cm ² | observed | (100) |
| in ethanol 42% positive | 2.2 µg/cm ² | observed | (264) |
| in ethanol 67% positive | 9.0 µg/cm ² | observed | (264) |
| Deodorant matrix 23 % positive | 0.167 µg/cm ² | observed | (279) |
| Deodorant matrix 69 % positive | 0.53 µg/cm ² | observed | (279) |
| Deodorant matrix 77 % positive | 1.67 µg/cm ² | observed | (279) |
| PATCH TEST | | | |
| ED50% (in petrolatum) | 32 µg/cm ² | estimated | (100) |
| No effect (in ethanol) | <0.0005% (0.15 µg/cm ²) | observed | (264) |
| No effect (in petrolatum) | <0.4 µg/cm ² | observed | (100) |

Summary of results for specific fragrance ingredients

Chloroatranol (constituent of *Evernia prunastri*)

In ROAT a dose of 0.025 µg/cm² to 0.125 µg/cm² in ethanol elicited reactions in 92% to 100% of sensitised subjects.

In patch testing the ED10% was 0.0004 µg/cm².

Cinnamal

In ROAT a dose of 0.26 µg/cm² gave a response in 11% when applied as deodorant in the axilla and 82% responded to 2.63 µg/cm².

The ED50 in patch testing was 96 µg/cm².

HICC

In ROAT a dose of 0.0357 µg/cm² gave no response, while the dose that elicited a reaction in 10% of the sensitised test group (in ethanol) ranged from 0.064 µg/cm² to 1.2 µg/cm². The dose in a cream base was 4.9 µg/cm².

In ROAT a dose of 15.3 µg/cm² to 126.2 µg/cm² in ethanol elicited reactions in 61% to 89% of sensitised subjects.

The ED10 in patch testing ranged from 0.66-0.9 µg/cm².

Hydroxycitronellal

In ROAT a dose of 0.94 µg/cm² gave a response in 57% when applied in a deodorant in the axilla and 100% responded to 9.40 µg/cm².

The no-effect level in patch testing was below 0.036 µg/cm².

Isoeugenol

In ROAT a dose of 2.2 µg/cm² a response in 42% and 9.0 µg/cm² in 67%, when applied in ethanol on the arm. With a deodorant applied to the skin of the axillary, a dose of 0.167 µg/cm² caused a response in 23% and 77% reacted to 1.67 µg/cm².

The ED50 in patch testing was 32 µg/cm².

The no-effect in patch testing was below 0.15 µg/cm².

Elicitation levels have been studied for cinnamal, isoeugenol and hydroxycitronellal which are established contact allergens in humans and which already have given rise to a significant number of cases (> 100, see chapter 7). Further HICC has been studied extensively, but is considered in a separate section (chapter 11.3) of this opinion. It is however not possible to derive a safe threshold directly from the data of cinnamal, isoeugenol and hydroxycitronellal. The main reasons are that many of the test subjects reacted to all the tested doses in ROAT, which is a simulation of every day exposures. Thus it was not possible to determine the dose only eliciting responses in a few, e.g. 10% of the subjects and that only a limited number of exposure scenarios were studied.

The studies have covered few product types: hydro-alcoholic products, e.g. perfumes and deodorant roll-on matrix. The vehicle is one of many factors which influence the thresholds of allergic reactions. Also the presence of irritants and other allergens can influence the elicitation level. This means that the currently available studies do not cover all the relevant exposure scenarios. However, taking into account that dose-response investigations in sensitised patients are very complex to perform, it is not likely that much more data will become available in the near future. It is therefore necessary to exploit the full pool of elicitation data, also covering chemicals other than fragrance ingredients, to derive a more general threshold which could be used when no or insufficient data exist to set a specific threshold for a substance of concern.

General thresholds

The methodology of the different experiments has varied to some extent as different anatomical sites of exposure have been employed, different vehicles, exposure periods and cut-off points. The reason is that the studies have been performed to investigate various clinical and scientific aspects of allergic contact reactions and not for formal regulatory requirements. Some studies are small and for this reason the precision of the estimates of thresholds is limited. In spite of this, the results of the various experiments are reasonably uniform, except for chloroatranol which had very low threshold reactions, and show that low concentrations may elicit allergic reactions.

The reasonably uniform data generated on the above fragrance ingredients are in agreement with a recent "meta-analysis" of dose-response data of different allergens, incorporating some of the same studies as mentioned above, but also other allergens, such as preservatives and metals. The ED10 at patch testing varied by a factor of 7 from the lowest to the highest value and the median was 0.82 µg/cm² if the three outliers formaldehyde (1997), nickel (1999) and methyl dibromo glutaronitrile (2004) were left out and 0.84 µg/cm² if included (see Table 11-6 and Figure 11-2 below: (280)). An explanation of these results could be that thresholds in elicitation is less dependent on the antigenic properties of the individual substance (inherent potency) than thresholds of induction and more on the level of sensitivity of the individual, i.e. the level of T-cell clones able to recognise the antigen, which is not present in naïve not-sensitised, individuals. This seems plausible, based on both the recent clinical evidence (280) and guinea pig QSAR evidence (281). It provides the basis for a general approach in establishing safe thresholds for substances of concern.

The consequences of a limit of 0.8 µg/cm² for the product types most important for fragrance allergy are calculated below.

The calculation is based on:

- The generally safe exposure level, which is the median ED10 value (the dose which will elicit allergic contact dermatitis in 10% of sensitised eczema patients) under patch test conditions: 0.8 µg/cm² (280).

- Exposure doses and exposure areas from SCCS notes of guidance 7th revision (282) [Tables 2 and 3] and Technical dossier Quantitative Risk Assessment from RIFM (274).

Equation:

Safe concentration in product = (Generally safe exposure level (0.8 µg/cm²)/daily exposure to product (µg/cm²/day)) x 100 (for %).

Table 11-5: Concentration limits in different product types based on 0.8 µg/cm² allergen as a 'generally safe exposure level', if specific dose-response data are unavailable.

| | Estimated daily exposure level (g) (Table 3 SCCS NoG) | Mean exposed skin surface (cm²) (Table 2 SCCS NoG) | Exposure /cm²/day in grams | Exposure /cm²/day in µg (1g= 1x10⁶ µg) | Concentration limit in product % in product: (GEL/daily exposure) x 100 |
|---------------------------------------|--|--|--|--|--|
| Body lotion | 7.82 g | 15,670 cm ² | 0.000499 | 499 | 0.16% |
| Face cream | 1.54 g | 565 cm ² | 0.002725 | 2725 | 0.03% |
| Hand cream | 2.16 g | 860 | 0.002511 | 2511 | 0.03% |
| Deodorant aerosol spray ethanol based | 1.43 g | 200 cm ² | 0.007150 | 7150 | 0.01% |
| Perfume spray | not given | ? | 0.00221 ¹⁾ | 2210 | 0.04% |

Note: 1) 2.21 mg/cm²/day from Technical dossier Quantitative Risk Assessment.

The estimated daily use of the various product categories in Table 11-5 are based on the SCCS Notes of Guidance (see above), except for perfume, for which no value is given. This value is taken from the Technical Dossier on Quantitative Risk Assessment from RIFM.

Generally the estimated use of different products is higher in the IFRA/RIFM assessments than in SCCS Notes of Guidance.

Table 11-6: Overview of dose-response studies and thresholds for eight allergens, after (280).

ED₁₀ patch test values from each of the 16 selected studies with 95 % confidence intervals with the allergens chromium (283), MCI/MI (Kathon TM CG) (284), nickel (285), methyl dibromo glutaronitrile (MDBGN) (286), hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC) (102, 224, 263), isoeugenol (264, 279) and formaldehyde (287). The shaded values were considered as outliers.

| Study | Number of patients | ED₁₀ (µg/cm²) | 95 % interval |
|--------------|---------------------------|--|----------------------|
| MCI/MI | 12 | 1.05 | 0.17–2.27 |
| Formaldehyde | 20 | 20.1 | 4.09–43.9 |
| Nickel 1997 | 24 | 1.58 | 0.32–4.04 |
| Nickel 1998 | 19 | 0.8 | 0.078–2.59 |

Opinion on fragrance allergens in cosmetic products

| Study | Number of patients | ED ₁₀ (µg/cm ²) | 95 % interval |
|-----------------|--------------------|--|---------------|
| Nickel 1999 | 26 | 7.49 | 2.42–14.5 |
| Nickel 2005 | 13 | 0.74 | 0.066–2.38 |
| Nickel 2007 | 20 | 0.82 | 0.13–2.37 |
| Cobalt 2005 | 11 | 0.44 | 0.033–1.3 |
| Chromium | 17 | 1.04 | 0.0033–5.55 |
| Isoeugenol 2001 | 24 | 1.48 | 0.22–4.74 |
| Isoeugenol 2005 | 13 | 0.23 | 0.0073–1.32 |
| HICC 2003 | 18 | 0.85 | 0.062–3.26 |
| HICC 2007 | 14 | 1.17 | 0.043–5.05 |
| HICC 2009 | 17 | 0.66 | 0.052–2.35 |
| MDBGN 2004 | 19 | 0.025 | 0.00021–0.19 |
| MDBGN 2008 | 18 | 0.50 | 0.052–1.69 |

Note: The ED₁₀ value is the concentration which elicits an allergic reaction in 10% of a group of sensitised individuals under patch test conditions.

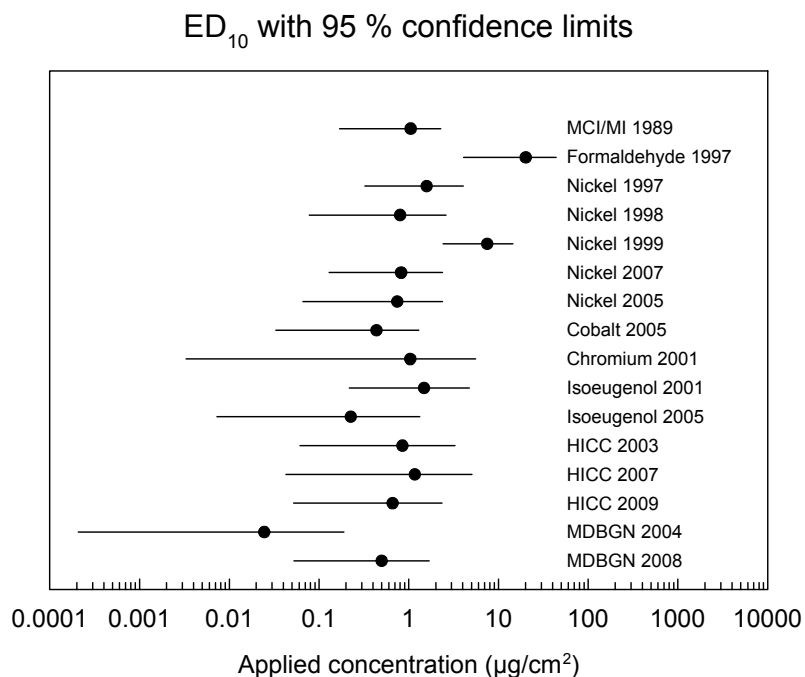


Figure 11-2: The threshold data with 95% confidence intervals from Table 11-6 presented graphically, after (280).

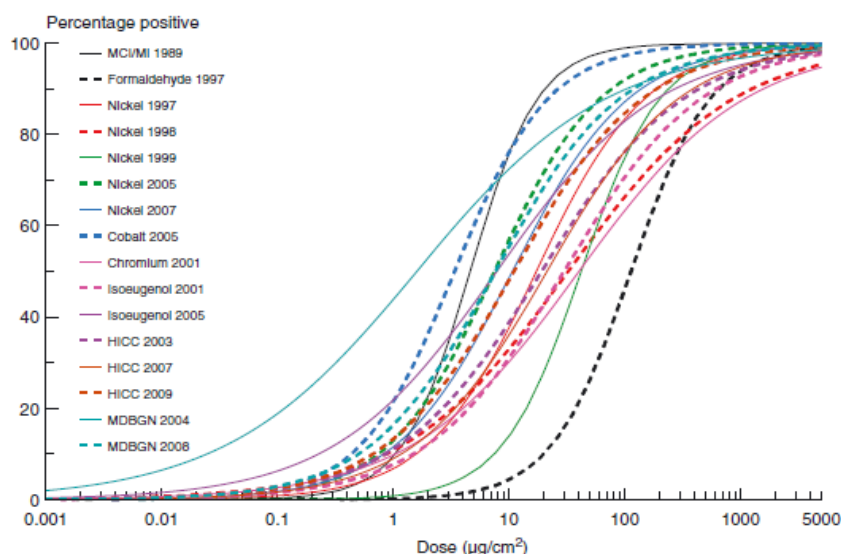


Figure 11-3: The fitted dose-response curves from the studies in Table 11-6, which are the basis for estimation of the ED10 value, after (280).

The meta-analysis above has shown that the median elicitation dose by patch testing for 10% of sensitised individuals was $0.8 \mu\text{g}/\text{cm}^2$. In the model data for the fragrance substances isoeugenol and HICC was included. The two studies on isoeugenol and the three studies on HICC gave an average ED10 value of $0.85 \mu\text{g}/\text{cm}^2$ and $0.89 \mu\text{g}/\text{cm}^2$ with a range 0.23-1.48. This means that even if the model was used for these substances individually the result would be very similar to the general threshold value.

The data from cinnamal and hydroxycitronellal studies was not incorporated in the model because: (i) serial dilution patch testing was done in petrolatum for cinnamal, making the dosing less exact; (ii) and only seven patients participated in the hydroxycitronellal study, while a criteria for inclusion in the model was ten participants (280).

According to the above calculations, a limit of $0.8 \mu\text{g}/\text{cm}^2$ for the product types of most importance for fragrance allergy corresponds to concentrations of 100 to 400 ppm (0.01-0.04%) for deodorants, perfume spray, hand and face lotions. For body lotion the general threshold was 0.16%. However, it does not seem meaningful in the context of contact allergy to distinguish between different types of creams, as a body cream would be applied with the hands and the relevant parameter in contact allergy is dose per area skin and not total dose.

A general threshold would have to take into consideration the uncertainties in quantification of exposure and safe thresholds as well as the possibilities of aggregate exposures and exposure to chemically similar substances. Therefore in setting one general threshold the product category carrying the highest risk of sensitisation and elicitation, which is deodorants, was chosen to drive the generation of the threshold. This means that a threshold of $0.8 \mu\text{g}/\text{cm}^2$ is equal to 0.01% or 100 ppm (see Table Table 11-1 and the related text), the lowest of the threshold values derived.

The approach taken by the SCCS is based on scientific evidence published in peer-reviewed journals (283)(284)(285)(286)(102, 224, 263)(264, 279)(287) in the past 20 years. The meta-analysis deriving the general threshold limit at $0.8 \mu\text{g}/\text{cm}^2$ limit has been published (280) in a peer-reviewed journal. The use of threshold limits based on elicitation data is a well established methodology which has been applied (with success) in EU to prevent further cases of induction and elicitation (primary and secondary prevention) in the case of nickel allergy, chromium in cement, chromium in shoes in

Germany, dimethyl fumurate in consumer items and also in part in IFRA guidelines e.g. concerning HICC.

The elicitation threshold model is based on 16 studies of 8 allergens, two of which are fragrance ingredients. It includes data from moderate to extreme allergens with a median EC3 value of 1.2.

The 11 fragrance allergens to which the limit is suggested to apply range from extreme to moderate with median EC3 value of 4.8, although in the case of coumarin an EC3 value could not be established.

Thus in general the potency profile of the fragrance substances of concern is not very different from those included in the model to provide the suggested general safe threshold.

The approach is targeting the relevant end-point, namely, allergic contact dermatitis. The mere consideration of potency of the allergen, according to the LLNA (EC3), is insufficient in identifying the size of the problems of contact allergy/allergic contact dermatitis. Additional information is needed from clinical and epidemiological studies, exposure assessment and dose-elicitation studies. For instance, the elicitation thresholds of e.g. HICC (EC3: 17.1) and isoeugenol (EC3: 0.54) are very similar (0.85 µg/cm² and 0.89 µg/cm², respectively) despite very different potencies. Both are frequent causes of contact allergy.

It should be noted that the general threshold is only suggested to be used for substances of concern if no specific data of sufficient quality exist to set an individual safe threshold. In cases where specific data of sufficient quality are available, these data should be used to set an individual safe threshold.

The general threshold is indicative of a safe level for the majority of sensitised individuals, but does not preclude that the most sensitive subset of the population may react upon exposure to the allergen. These levels are based on patch tests and take no account of anatomical sites of exposure, frequency of exposure or vehicle effects. Therefore, any limitations in exposures are not substitutes for providing information to the consumer about the presence of a substance in a product as a certain fraction of sensitised individuals will still need to avoid specific exposures.

Based on experience, limitations in exposure based on elicitation thresholds will, apart from helping the sensitised consumer, also significantly reduce the risk of induction. This is the case for nickel allergy, where the restrictions in the EU nickel directive are based on elicitation threshold, leading to a significant reduction in new cases of sensitisation in young women (288) and in a reduction in morbidity, i.e. elicitation (289). Another example is restriction of chromium VI in cement (290).

It is not possible to provide a safe threshold for natural extracts of concern, as no specific investigations exist, and the model providing the general use concentration limit (0.01%) has been based on chemicals only.

The SCCP concluded in 2004 that Chloroatranol and atranol, the main allergenic constituents of *Evernia prunastri* and *Evernia furfuracea*, should not be present in consumer products because they are extremely potent allergens (239). The persistently high frequency of contact allergy to *Evernia prunastri* and *Evernia furfuracea* noted in eczema patients does point to a persisting problem with exposure to the allergenic constituents.

11.3. Hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC)

Hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC) has been the most frequently reported individual fragrance chemical causing allergy since the 1999 opinion on fragrance allergy. In total, reports of about 1500 cases have been published in the scientific literature (see chapter 7.1 and Annex I to this opinion), while the second most

frequently reported individual chemical was cinnamal with around 350 published cases. Only a minority of the cases seen by clinicians is published and only a (small) proportion of those with allergic contact dermatitis seeks or has the possibility to seek medical attention.

Natural extracts such as *Myroxylon pereirae* and turpentine (oil) have been more frequently reported, but while HICC is a synthetic fragrance chemical, where the only source of exposure is fragrances, the natural extracts are used in many other contexts than fragrances/cosmetics.

Of patients tested by the Danish monitoring network of dermatologists 2.4% were found to be allergic to HICC in 2005-2008 (with no decreasing trend from 2003 to 2007 (291)) (for more studies see chapter 4.3.2); in 70% of the cases the reaction was of current relevance, i.e. causing disease (69). This is in agreement with the results of a recent German study with HICC, where 48 out of 51 patients (94.1%) with a positive patch test reaction to HICC also reacted in a repeated open application test, simulating normal use conditions of cosmetics containing HICC (105). In a Danish study 69% of 14 HICC allergic individuals developed allergic contact dermatitis from use of cosmetics containing HICC in realistic amounts (102).

On the basis of the high frequency of allergy to HICC, in 2003 the Scientific Committee on Cosmetic Products and Non-Food Products (SCCNFP) recommended 0.02% (200 ppm) as maximum amount of HICC in cosmetic products (292). This has not been implemented and no restrictions apply in the Cosmetic Directive.

The fragrance industry, via the International Fragrance Association (IFRA), has its own safety guidelines. Up until 2003 HICC was used without any restriction; in 2003 a limit of 1.5% HICC in any kind of product was introduced. In 2008 this was changed according to the new risk assessment model (QRA) applied by the fragrance industry to different levels in 11 different product types derived from the QRA (see 11.1). Limits from 0.11% in lip products to 1.5% in hair styling products were set. In 2009 a further lowering was made of the limits by industry with the following reasoning: "The industry firmly believes and continues to support thresholds based on induction rather than elicitation. However, given the exceptional situation in Europe, the fragrance industry elected to take further restrictive action on this material" (293). An overview of the IFRA restrictions is given in the table below.

Table 11-7: Restriction for HICC independent of the QRA according to (293).

| IFRA QRA Category | Product type that drives the category | Consumer exposure level 2003–2008 (%) | IFRA Standard July 2008 (%) | IFRA Standard July 2009 (%) |
|--------------------------|--|--|------------------------------------|------------------------------------|
| Category 1 | Lip products | 1.5 | 0.11 | 0.02 |
| Category 2 | Deodorants/antiperspirants | 1.5 | 0.15 | 0.02 |
| Category 3 | Hydroalcoholics for shaved skin | 1.5 | 0.60 | 0.2 |
| Category 4 | Hydroalcoholics for unshaved skin | 1.5 | 1.5 | 0.2 |
| Category 5 | Hand cream | 1.5 | 1.0 | 0.2 |
| Category 6 | Mouthwash | 1.5 | 1.5 | Not applicable* |
| Category 7 | Intimate wipes | 1.5 | 0.3 | 0.02 |

Opinion on fragrance allergens in cosmetic products

| | | | | |
|-------------|--------------------------------|-----|----------------|----------------|
| Category 8 | Hair styling aids | 1.5 | 1.5 | 0.2 |
| Category 9 | Rinse-off hair conditioners | 1.5 | 1.5% | 0.2% |
| Category 10 | Hard surface cleaners | 1.5 | 1.5% | 0.2% |
| Category 11 | Incidental or non-skin contact | 15 | Not restricted | Not restricted |

Note: HICC Hydroxyisohexyl 3-cyclohexene carboxaldehyde.

QRA Quantitative risk assessment.

* Not applicable because HICC is not approved for flavour use.

As an update since the presentation of the pre-consultation version of the opinion, surveillance data on HICC from two European countries have become available, covering the period 2002-2011 (IVDK/Germany (294)) and 2003-2011 (Danish contact dermatitis group (295)), respectively. The first analysis identified a slight decrease, which was considered "not overwhelming in absolute terms", namely, from 2.3% in 2002 to 2.1% in 2011 (crude prevalences, Figure 11-4). Thus, despite statistical significance, the decrease is too slight to be interpreted as relevant improvement. In the Danish study, some fluctuation around a mean prevalence of about 2.5% was noted, but no trend (Figure 11-5). It is reported that 74% of the positive reactions were regarded as clinically relevant.

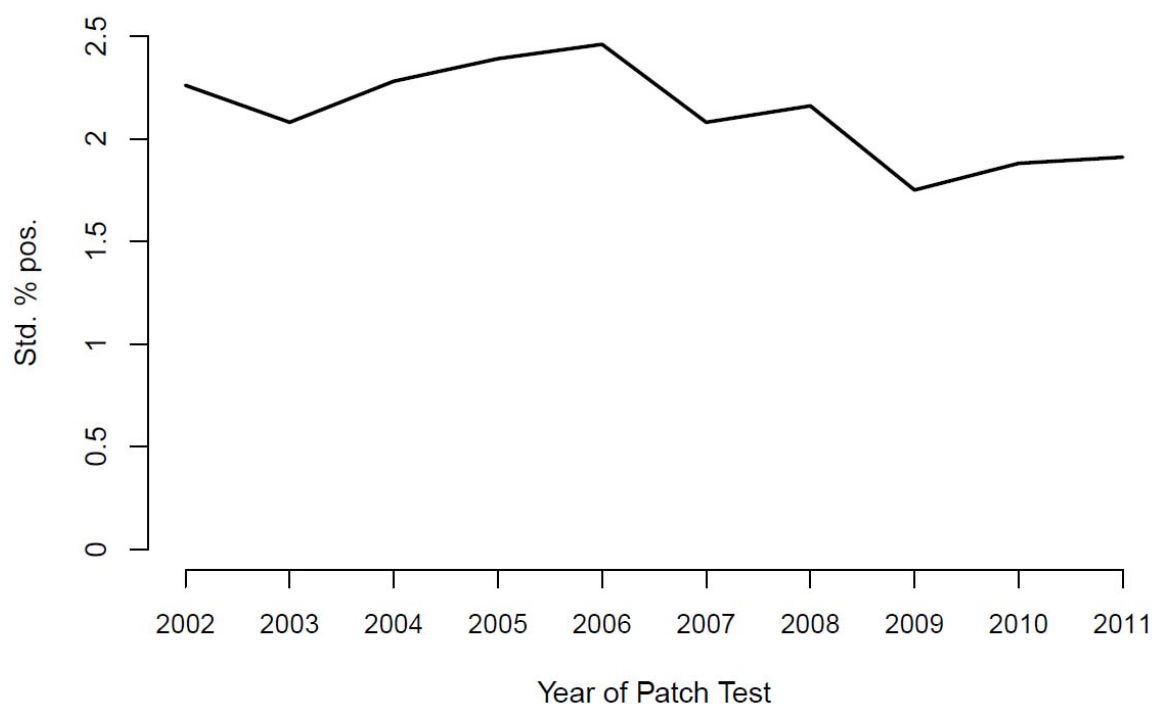


Figure 11-4: Time trend of hydroxyisohexyl 3-cyclohexene carboxaldehyde sensitisation prevalence [standardised prevalence of positives (%)] during 2002-2011. The decrease over time is statistically significant, after **(294)**.

Opinion on fragrance allergens in cosmetic products

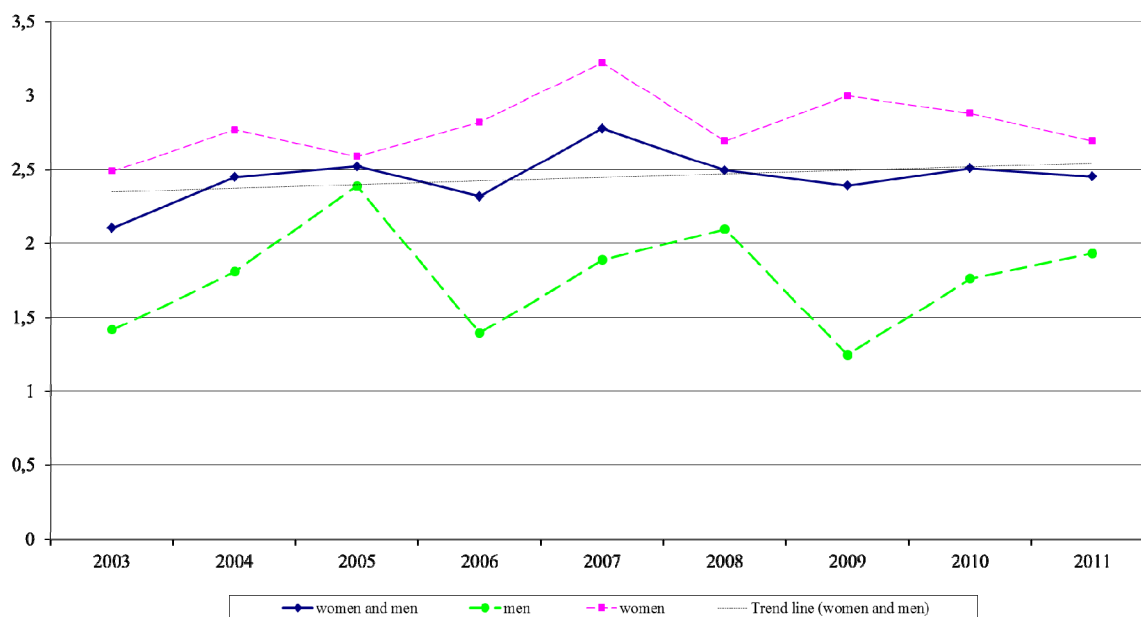


Figure 11-5: Prevalence of positive patch test reactions to hydroxyisohexyl 3-cyclohexene carboxaldehyde over time in 37 860 subjects tested by the Danish Contact Dermatitis Group (295).

11.4. Conclusion

- A dose-response relationship between exposure to contact allergens and induction of allergy (sensitisation) as well as elicitation is well established. This means that in principle, thresholds can be identified which are safe for the consumer.
- A model for dermal sensitisation quantitative risk assessment has been developed (QRA) and implemented by the fragrance industry. This model relies on thresholds, no effect or low-effect levels, established in healthy human volunteers and/or in animal experiments. The SCCP has previously reviewed this methodology and concluded that: "There is no confidence that the levels of skin sensitisers identified by the dermal sensitisation QRA are safe for the consumer."
- Elicitation data can provide thresholds indicative for the safe use of those substances which have already caused significant problems in the consumer. In this context, "safe use" means that the thresholds will protect the majority of consumers from allergic contact dermatitis, but does not preclude that the most sensitive subset of the population may react upon exposure to the allergen.
- Furthermore, based on experience from intervention studies, such thresholds will also be sufficiently low to protect (most of) the non-sensitised consumers from developing contact allergy.
- Elicitation levels have been studied specifically for the fragrance chemicals cinnamal, hydroxycitronellal and isoeugenol. These studies, however, are not adequate to derive safe thresholds for the individual substances directly from the data.
- In the absence of adequate substance specific data it is possible to use a general threshold. Based on a statistical analysis of the available data in the scientific literature, a threshold of $0.8 \mu\text{g}/\text{cm}^2$ was derived. This corresponds to 0.01% (100 ppm) limit in cosmetic products indicative for safe use.

- It is not possible to provide a safe threshold for natural extracts of concern, as no specific investigations exist and the model providing the general threshold (0.01%) has been based on individual chemicals only. However the maximum use concentration applies to the identified chemicals both if added as chemicals or as an identified constituent of a natural ingredient. This will also reduce the risk of sensitisation and elicitation from natural extracts.
- For substances for which there are no clinical data of concern, models such as the dermal sensitisation QRA approach may, after refinement and validation, be used to suggest a safe level of exposure prior to incorporation into products. However, aggregated exposures must be incorporated in the dermal sensitisation QRA model.
- HICC has for more than 10 years been recognized as an important allergen with more cases documented in the scientific literature than for any other fragrance chemical in this period. HICC has been shown to be a significant cause of disease as many of those with contact allergy to HICC had also reactions to cosmetics, which contained or were likely to contain HICC. Since 2003 attempts have been made by the fragrance industry to contain the outbreak of HICC allergy, but with no convincing success so far. Recent voluntary restrictions (recommendations to lower use concentrations, at least for some product types, to the level recommended by the SCCS in 2003) are not reflected in available evidence and are considered insufficient. The SCCS considers that the number of cases of HICC allergy documented over the last decade is exceptionally high and that continued exposure to HICC by the consumer is not considered safe, even at concentrations as low as 200 ppm. Therefore, HICC should not be used in consumer products in order to prevent further cases of contact allergy to HICC and to limit the consequences to those who already have become sensitized.
- The SCCP concluded in 2004 that chloroatranol and atranol, the main allergenic constituents of *Evernia prunastri* and *Evernia furfuracea*, should not be present in consumer products because they are extremely potent allergens. The persistently high frequency of contact allergy to *Evernia prunastri* and *Evernia furfuracea* noted in eczema patients does point to a persisting problem with exposure to the allergenic constituents, despite efforts to reduce the allergen content (296).

12. Data gaps and research needed

In the course of working on this opinion, the following points are highlighted as important data gaps, ordered by research area:

12.1. Clinical and epidemiological research

- Clinical data on more fragrance substances are needed to assess more fully the epidemiology of fragrance contact allergy and pin-point the culprit substances for induction and elicitation of contact allergy in man.
- Data from a broader range of EU countries on the clinical and epidemiological picture of fragrance contact allergy is needed, as difference in exposure and use habits are expected across Europe.
- A co-ordinated strategy for data collection should be developed.
- Very little is known about susceptible groups of the population, e.g. up to 10% of the European population carry mutations, which impair the skin barrier and which seem to increase the risk of fragrance allergy. Data are needed to qualify and quantify the increase in risk of susceptible groups in order to provide a better protection of all consumers.
- Aberrant enzyme activity in certain individuals, often related to genetic enzyme polymorphisms, may give an increased or reduced risk of sensitisation to prohaptens (that need enzymatic activation) in certain individuals or populations. More research into the role of relevant traits is needed.
- Dose-response data from clinical studies are available for only a few allergens. To establish individual safe levels such data are required for all established allergens of concern and covering an appropriate range of product types. This would also consolidate the basis of the use of a general threshold for safe use of fragrance allergens.
- Data on human exposure to fragrances from the use of different product categories is very scarce and therefore does not provide an optimal basis of risk assessment, e.g. exposure data on use for perfume/eau de cologne are lacking.
- Most experimental studies are done on individual fragrance ingredients, while exposure to allergens in cosmetic products is usually to mixtures of allergens. The risk of sensitisation and elicitation may depend on the mixture of substances, but very few studies on this exist. It is necessary to improve the knowledge base on cocktail effects on sensitisation/elicitation to improve the basis of risk assessment and management.
- Screening in dermatitis patients should be performed with air exposed samples of such fragrance substances that in experimental studies have been demonstrated to act as pre-haptens, i.e. autoxidise and form oxidation mixtures containing allergenic oxidation products.
- Patch testing should if possible, be performed with the isolated true haptens formed from pre-haptens and pro-haptens to increase the possibility to diagnose allergy from these type of substances.
- There is a need for more experimental research to further establish the impact of the behaviour of fragrance substances when applied on the skin (including factors such as volatility, autoxidation, skin penetration, reactivity in skin and bioactivation).

12.2. Non-human studies

- Several studies in the industry submission (164) were of insufficient quality, not following the OECD guidelines.
- In some cases it was found that either very few concentrations points had been used in LLNAs, or concentrations were insufficient for achieving a 3-fold increase of the SI.

A sufficient number of doses (concentrations) should be applied in LLNAs (at least 5) so that interpolation (for deriving an EC3 value) can rely on more than two or three actual data points to be more reliable. SCCS therefore suggests a change in the OECD guideline 429. (It is important to remember that the production of unreliable data is a waste of animals.) Moreover, the maximum concentration should be high enough to achieve a > 3-fold increase in SI, as far as this is possible with the substance/vehicle combination chosen.

- Data on experimental results are often not published, but available only on file in the companies having performed the tests. Access to such results would be important for the scientific community, e.g. in the context of REACH, or independently, either to the public domain, or to a Public Trustee.
- The OECD guideline 429 recommends several vehicles. It is well known that a difference in the EC3 value can be obtained for the same substance depending on which vehicle is used in the LLNA. Thus, as an additional control, supplementary to the guideline based LLNA control, a clinically relevant solvent or the commercial formulation in which the test substance is marketed may be used.
- As long as no validated *in vitro* method exists, more research is needed. Until one or more method(s) have been decided to fulfil the requirements for substituting *in vivo* testing, the *in vivo* testing for prediction of skin sensitisation has to be used.
- Applying only mechanism-based QSAR (QMM) as a tool in non-animal based risk assessment for skin sensitisation is of limited value for fragrance substances. This is due to major information gaps in the present model when addressing substances that act via abiotic or metabolic activation, and the high incidence of such substances in fragrances. Therefore, further experimental and clinical research in the area of abiotic and/or metabolic activation of fragrance substances is needed to increase the safety for the consumer, i.e. experimental studies which include air oxidation and bioactivation.
- Further experimental investigations of the sensitisation potential of fragrance substances are needed to determine the impact of the volatility of the substance as well as the effect of the vehicle on skin penetration/absorption and reactivity.
- From a clinical perspective it is important for the individual who is sensitised to one fragrance substance to know if they must also avoid other fragrance substances that can cause allergic contact dermatitis due to cross-reactivity with the original sensitiser. Prediction of risks for cross-reactivity requires sound application of theoretical principles in combination with well-designed experimental studies. This is a field that has not been studied very much so far and needs to be focused on much more in the future.
- Quantitative structure activity relationship (QSAR) models should be further developed, combining, as appropriate, information from *in silico*, *in chemico* and *in vitro* methods as possible. Prediction of different activation pathways should be included.
- Effect estimates such as proportions of sensitised humans or animals, or mean stimulation indices, EC3 values and other derivations should ideally be accompanied by an interval estimate (confidence interval) to address precision (297).

13. Opinion

Contact allergy to fragrances is a common, significant and relevant problem in Europe. The studies since the SCCNFP opinion on fragrance allergy in consumers in 1999 (SCCNFP/0017/98) (SCCNFP 1999) have confirmed that the 26 fragrance allergens, identified by the SCCNFP, are still relevant fragrance allergens for consumers because of their exposure from cosmetic products. Additional exposure to many of these 26 fragrance allergens also occurs from the use of other consumer products, such as detergents, toys, etc. Some of these fragrance substances are also used as preservatives.

The overall trend of fragrance contact allergy appears to have been stable for the last 10 years, as some causes of fragrance allergy have decreased and others increased. From the few population-based studies, it can be estimated that the frequency of contact allergy to fragrance ingredients in the general population in Europe is 1-3%. This is based on the limited testing with eight common fragrance allergens (FM I) out of the approximately 2500 fragrance ingredients listed in CosIng and indicative of the substances that may be present in fragrance compounds. However, the real prevalence of contact allergy to fragrance substances may be higher if the testing were to be performed with the full spectrum of fragrance allergens, including oxidised substances, where relevant.

Among eczema patients in the European population, around 16% are sensitised to fragrance ingredients. The disease can be severe and generalised, with a significant impairment of quality of life and potential consequences for fitness for work.

Contact sensitisation, and its clinical manifestation, allergic contact dermatitis, can be prevented if the exposure to known contact allergens is reduced or abolished (primary prevention). Experiences so far, have indicated that not all substances that later turned out to be significant contact allergens after human exposure, were predicted by experimental studies, e.g. the preservative methyl dibromo glutaronitrile and the fragrance chemical HICC. Thus, a significant exposure of the population may occur before a substance is established as an important contact allergen in man.

Elicitation of allergic contact dermatitis occurs when a consumer sensitised to a certain substance is re-exposed to the substance in question. Prevention at this stage, termed secondary prevention, can be achieved if use of the allergen in products is eliminated or reduced to a tolerable level (general prevention), or if the patients succeed in avoiding all sources of exposure (individual prevention). Ingredient listing of individual fragrance allergens has been shown to be an important tool to enable consumers with an identified allergy to reduce/avoid relevant exposures. Moreover, ingredient listing is also of great importance to ensure that an adequate diagnosis of fragrance contact allergy can be made without undue delay. If the information given on the presence of fragrance allergens is incomplete, diagnosis of fragrance contact allergy may be missed.

The SCCNFP, in its 1999 opinion, identified 26 fragrance allergens for which information should be provided to consumers concerning their presence in cosmetic products. This was implemented in the European Cosmetics legislation (298) as ingredient labelling of these 26 fragrance substances (Annex III, entries 67-92). However, safe use concentrations for these substances in cosmetic products have not yet been determined and much new evidence concerning fragrance allergy has been published since 1999. The present opinion updates the SCCNFP opinion with a systematic and critical review of the scientific literature up to October 2010. This review addresses the issue of contact allergy to fragrance substances, including natural extracts and updates the list of fragrance allergens relevant to consumers. Clinical, epidemiological and experimental studies were evaluated, as well as modelling studies performed, to establish lists of: (i) established fragrance allergens; (ii) likely fragrance allergens; and (iii) possible fragrance allergens. The review also includes fragrances, which on modification by oxidation or by enzyme mediated processes, can produce allergens. Available dose-response data have been

examined to answer whether safe thresholds can be established for the most frequent fragrance allergens.

13.1. Question 1

Does the SCCS still consider that the fragrance allergens currently listed in Annex III, entries 67-92, for labelling purposes represent those fragrance ingredients that the consumer needs to be made aware of when present in cosmetic products?

In order to answer this question, the SCCS has used clinical and epidemiological data to identify known fragrance allergens. These were categorised as *established contact allergens in humans* (see Table 13-1).

Where sufficient animal evidence was present, these substances were categorised as established contact allergens in animals (Table 13-2). For a number of other fragrance substances, combinations of limited clinical data together with SAR considerations have been applied to indicate likely fragrance allergens in man (Table 13-3). Finally, SAR has also been applied to substances that lack human data to identify fragrance chemicals that have the structural potential to be contact allergens. Substances with insufficient human data were also considered as possible fragrance allergens. For these further tests (experimental/clinical data) are required (Table 13-4).

Table 13-1: Established contact allergens in humans.

For categorisation of importance (+ to +++) see chapter 7.1. Allergens of special concern are substances where between 100 and 1,000 cases (+++) and more than 1,000 (++++) have been published. These are set in bold. Fragrance substances identified as allergens in the 1999 opinion of SCCNFP (1) are marked with an asterisk.

"ox." = oxidised; "non-ox." = non-oxidised; "r.t." = rarely tested (see chapter 7)

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Human evidence: see text |
|--|--------------------|--------------------------|
| Individual chemicals | | |
| ACETYLCEDRENE | 32388-55-9 | + |
| AMYL CINNAMAL* | 122-40-7 | ++ |
| AMYL CINNAMYL ALCOHOL* | 101-85-9 | ++ |
| AMYL SALICYLATE | 2050-08-0 | + |
| trans-ANETHOLE | 4180-23-8 | + (r.t.) |
| ANISE ALCOHOL* | 105-13-5 | + |
| BENZALDEHYDE | 100-52-7 | + |
| BENZYL ALCOHOL* | 100-51-6 | ++ |
| BENZYL BENZOATE* | 120-51-4 | ++ |
| BENZYL CINNAMATE* | 103-41-3 | ++ |
| BENZYL SALICYLATE* | 118-58-1 | ++ |
| BUTYLPHENYL METHYLPROPIONAL * | 80-54-6 | ++ |
| CAMPHOR | 76-22-2 / 464-49-3 | + (r.t.) |
| beta-CARYOPHYLLENE (ox.) | 87-44-5 | Non-ox.: +, ox.: + |

Opinion on fragrance allergens in cosmetic products

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Human evidence: see text |
|--|--|--|
| CARVONE | 99-49-0 / 6485-40-1 / 2244-16-8 | + (r.t.) |
| CINNAMAL* | 104-55-2 | +++ |
| CINNAMYL ALCOHOL* | 104-54-1 | +++ |
| CITRAL* | 5392-40-5 | +++ |
| CITRONELLOL* | 106-22-9 / 1117-61-9 / 7540-51-4 | ++ |
| COUMARIN* | 91-64-5 | +++ |
| (DAMASCENONE) ROSE KETONE-4 | 23696-85-7 | + (r.t.) |
| alpha-DAMASCONE (TMCHB) | 43052-87-5 / 23726-94-5 | ++ |
| cis-beta-DAMASCONE | 23726-92-3 | + |
| delta-DAMASCONE | 57378-68-4 | + |
| DIMETHYLBENZYL CARBINYL ACETATE (DMBCA) | 151-05-3 | + |
| EUGENOL* | 97-53-0 | +++ |
| FARNESOL* | 4602-84-0 | ++ - +++ |
| GERANIOL* | 106-24-1 | +++ |
| HEXADECANOLACTONE | 109-29-5 | + (r.t.) |
| HEXAMETHYLINDANOPYRAN | 1222-05-5 | ++ |
| HEXYL CINNAMAL* | 101-86-0 | ++ |
| HYDROXYISOHEXYL CARBOXALDEHYDE (HICC)* | 3-CYCLOHEXENE 31906-04-4 / 51414-25-6 | ++++ |
| HYDROXYCITRONELLAL* | 107-75-5 | +++ |
| ISOEUGENOL* | 97-54-1 | +++ |
| alpha-ISOMETHYL IONONE* | 127-51-5 | ++ |
| (DL)-LIMONENE* | 138-86-3 | ++ (non-ox.); +++ (ox.) |
| LINALOOL* | 78-70-6 | ++ (non-ox.) +++ (ox.) |
| LINALYL ACETATE | 115-95-7 | + (non-ox.) ++ (ox.) |
| MENTHOL | 1490-04-6 / 89-78-1 / 2216-51-5 | ++ |
| 6-METHYL COUMARIN | 92-48-8 | ++ |
| METHYL 2-OCTYNOATE* | 111-12-6 | ++ |
| METHYL SALICYLATE | 119-36-8 | + |
| 3-METHYL-5-(2,2,3-TRIMETHYL-3- | 67801-20-1 | ++ (r.t.) |

Opinion on fragrance allergens in cosmetic products

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Human evidence: see text |
|--|---|---------------------------------|
| CYCLOPENTENYL)PENT-4-EN-2-OL | | |
| alpha-PINENE and beta-PINENE | 80-56-8 and 127-91-3, resp. | ++ |
| PROPYLIDENE PHTHALIDE | 17369-59-4 | + (r.t.) |
| SALICYLALDEHYDE | 90-02-8 | ++ |
| alpha-SANTALOL and beta-SANTALOL | 115-71-9 and 77-42-9, resp. | ++ |
| SCLAREOL | 515-03-7 | + |
| TERPINEOL (mixture of isomers) | 8000-41-7 | + |
| alpha-TERPINEOL | 10482-56-1 / 98-55-5 | |
| Terpinolene | 586-62-9 | + |
| TETRAMETHYL ACETYLOCTAHYDRONAPHTHALENES | 54464-57-2 / 54464-59-4 / 68155-66-8 / 68155-67-9 | + |
| TRIMETHYL-BENZENEPROPANOL (Majantol) | 103694-68-4 | ++ |
| VANILLIN | 121-33-5 | ++ |
| Natural extracts | | |
| <i>CANANGA ODORATA and Ylang-ylang oil</i> | 83863-30-3; 8006-81-3 | +++ |
| <i>CEDRUS ATLANTICA BARK OIL</i> | 92201-55-3; 8000-27-9 | ++ |
| <i>CINNAMOMUM CASSIA LEAF OIL</i> <i>CINNAMOMUM ZEYLANICUM BARK OIL</i> | 8007-80-5 84649-98-9 | ++ (r.t.) |
| <i>CITRUS AURANTIUM AMARA FLOWER / PEEL OIL</i> | 8016-38-4; 72968-50-4 | ++ |
| <i>CITRUS BERGAMIA PEEL OIL EXPRESSED</i> | 89957-91-5 | + (r.t.) |
| <i>CITRUS LIMONUM PEEL OIL EXPRESSED</i> | 84929-31-7 | ++ |
| <i>CITRUS SINENSIS (syn.: AURANTIUM DULCIS) PEEL OIL EXPRESSED</i> | 97766-30-8; 8028-48-6 | ++ |
| <i>CYMBOPOGON CITRATUS / SCHOENANTHUS OILS</i> | 89998-14-1; 8007-02-1; 89998-16-3 | ++ |
| <i>EUCALYPTUS SPP. LEAF OIL</i> | 92502-70-0; 8000-48-4 | ++ |
| <i>EUGENIA CARYOPHYLLUS LEAF / FLOWER OIL</i> | 8000-34-8 | +++ |
| <i>EVERNIA FURFURACEA EXTRACT*</i> | 90028-67-4 | +++ |
| <i>EVERNIA PRUNASTRI EXTRACT*</i> | 90028-68-5 | +++ |
| <i>JASMINUM GRANDIFLORUM / OFFICINALE</i> | 84776-64-7; 90045-94-6; 8022-96-6 | +++ |
| <i>JUNIPERUS VIRGINIANA</i> | 8000-27-9; | ++ |

Opinion on fragrance allergens in cosmetic products

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Human evidence: see text |
|---|--|---------------------------------|
| | 85085-41-2 | |
| <i>LAURUS NOBILIS</i> | 8002-41-3; 8007-48-5; 84603-73-6 | ++ |
| <i>LAVANDULA HYBRIDA</i> | 91722-69-9 | + (r.t.) |
| <i>LAVANDULA OFFICINALIS</i> | 84776-65-8 | ++ |
| <i>MENTHA PIPERITA</i> | 8006-90-4; 84082-70-2 | ++ |
| <i>MENTHA SPICATA</i> | 84696-51-5 | ++ |
| MYROXYLON PEREIRAE | 8007-00-9; | ++++ |
| <i>NARCISSUS SPP.</i> | diverse | ++ |
| <i>PELARGONIUM GRAVEOLENS</i> | 90082-51-2; 8000-46-2 | ++ |
| <i>PINUS MUGO/PUMILA</i> | 90082-72-7 / 97676-05-6 | ++ |
| <i>POGOSTEMON CABLIN</i> | 8014-09-3; 84238-39-1 | ++ |
| <i>ROSE FLOWER OIL (ROSA SPP.)</i> | Diverse | ++ |
| SANTALUM ALBUM | 84787-70-2; 8006-87-9 | +++ |
| TURPENTINE (oil) | 8006-64-2; 9005-90-7; 8052-14-0 | ++++ |
| VERBENA ABSOLUTE | 8024-12-2 | ++ |

Table 13-2: Fragrance substances categorised as established contact allergens in animals.

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Human evidence: see text | EC 3 value (min; %) |
|---|-------------------|---------------------------------|----------------------------|
| Individual chemicals | | | |
| Allyl phenoxyacetate | 7493-74-5 | none | 3.1 |
| p-tert. -Butyldihydrocinnamaldehyde | 18127-01-0 | none | 4.3 |
| CYCLAMEN ALDEHYDE | 103-95-7 | none | 22 |
| Dibenzyl ether | 103-50-4 | none | 6.3 |
| 2,3-DIHYDRO-2,2,6-TRIMETHYLBENZALDEHYDE | 116-26-7 | limited | 7.5 |
| trans-2-Hexenal | 6728-26-3 | none | 2.6 |
| 2-Hexylidene cyclopentanone | 17373-89-6 | none | 2.4 |
| HEXYL SALICYLATE | 6259-76-3 | negative | 0.18 |
| p-Isobutyl- α -methyl hydrocinnamaldehyde | 6658-48-6 | none | 9.5 |

Opinion on fragrance allergens in cosmetic products

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Human evidence: see text | EC 3 value (min; %) |
|---|-------------------|---------------------------------|----------------------------|
| Isocyclocitral | 1335-66-6 | none | 7.3 |
| α -Methyl cinnamic aldehyde | 101-39-3 | none | 4.5 |
| METHYLENEDIOXYPHENYL METHYLPROPANAL | 1205-17-0 | none | 16.4 |
| METHYLUNDECANAL | 110-41-8 | none | 10 |
| 2-Methoxy-4-methylphenol | 93-51-6 | none | 5.8 |
| 4-Methoxy- α -methyl benzenpropanal | 5462-06-6 | none | 23.6 |
| METHYL OCTINE CARBONATE | 111-80-8 | limited | 2.5 |
| Perillaldehyde p-Mentha-1,8-dien-7-al | 2111-75-3 | none | 8.1 |
| PHENYLACETALDEHYDE | 122-78-1 | limited | 3 |
| Natural extracts | | | |
| Jasminum Sambac Flower CERA / Extract / Water | 91770-14-8 | none | 35.4 |

Table 13-3: Fragrance substances categorised as likely contact allergens by combination of evidence.

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Human evidence: see text | EC 3 value (min; %) | SAR |
|---|-------------------|---------------------------------|----------------------------|------------|
| AMBRETTOLIDE | 7779-50-2 | limited | none | + |
| CARVACROL | 499-75-2 | limited | none | + |
| Citrus paradisi § | 8016-20-4 | none | R43 | n.a. |
| CUMINALDEHYDE | 122-03-2 | limited | none | + |
| CYCLOPENTADECANONE | 502-72-7 | limited | none | + |
| trans-trans-delta-DAMASCONE | 71048-82-3 | limited | none | + |
| 2,4-dimethyl-3-cyclohexen-1-carboxaldehyde § | 68039-49-6 | none | R43 | + |
| DIMETHYLTETRAHYDRO BENZALDEHYDE | 68737-61-1 | limited | none | + |
| ETHYL VANILLIN | 121-32-4 | limited | none | + |
| HELIOTROPINE | 120-57-0 | limited | none | + |
| ISOAMYL SALICYLATE | 87-20-7 | limited | none | ++ |
| ISOLONGIFOLENEKETONE | 33407-62-4 | limited | none | + |
| Longifolene § | 475-20-7 | none | R43 | + |
| Mentha arvensis § | 68917-18-0 | none | R43 | n.a. |
| METHOXYCITRONELLAL | 3613-30-7 | limited | none | + |
| METHYL CINNAMATE | 103-26-4 | limited | none | ++ |
| METHYLIONANTHEME | 55599-63-8 | limited | none | + |

Opinion on fragrance allergens in cosmetic products

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Human evidence: see text | EC 3 value (min; %) | SAR |
|--|------------|--------------------------|---------------------|-----|
| 5-METHYL-alpha-IONONE | 79-69-6 | limited | none | + |
| MYRCENE | 123-35-3 | limited | none | ++ |
| MYRTENOL | 515-00-4 | limited | none | + |
| NEROL | 106-25-2 | limited | none | ++ |
| Nerolidol (isomer not specified) | 7212-44-4 | limited | none | ++ |
| NOPYL ACETATE | 128-51-8 | limited | none | + |
| PHYTOL | 150-86-7 | limited | none | + |
| RHODINOL | 6812-78-8 | limited | none | + |
| trans-ROSE KETONE-5 | 39872-57-6 | limited | none | ++ |

§ Substances/natural mixtures were classified as R43, according to the submission by IFRA. The evidence on which this classification was based was not available to the SCCS, so the validity of classification cannot be assessed. Nevertheless, the four substances/substance mixtures should be treated as *likely contact allergens*.

n.a.: not applicable (natural mixture)

Table 13-4: Fragrance substances categorised as possible contact allergens.

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Human evidence: see text | EC 3 value (min; %) | SAR |
|---|------------|--------------------------|---------------------|-----|
| Individual chemicals | | | | |
| CYCLOHEXYL ACETATE | 622-45-7 | limited | none | 0 |
| ETHYLENE DODECANEDIOATE | 54982-83-1 | limited | none | 0 |
| HYDROXYCITRONELLOL | 107-74-4 | limited | none | 0 |
| METHOXYTRIMETHYLHEPTANOL | 41890-92-0 | limited | none | 0 |
| METHYL p-ANISATE | 121-98-2 | limited | none | 0 |
| METHYL DIHYDROJASMONATE | 24851-98-7 | limited | none | 0 |
| PHENETHYL ALCOHOL | 60-12-8 | limited | none | 0 |
| PHENYLPROPANOL | 122-97-4 | limited | none | 0 |
| AMYL CYCLOPENTANONE | 4819-67-4 | negative | none | + |
| BENZYL ACETATE | 140-11-4 | negative | none | + |
| 6-ETHYLIDENEOCTAHYDRO-5,8-METHANO-2H-BENZO-1-PYRAN | 93939-86-7 | negative | none | + |
| 3 α ,4,5,6,7,7 α -HEXAHYDRO-4,7-METHANO-1H-INDEN-5(OR 6)-YL ACETATE | 54830-99-8 | negative | none | + |
| alpha-IONONE | 127-41-3 | negative | none | + |
| beta-IONONE | 79-77-6 | negative | none | + |
| METHYL IONONE (mixture of | 1335-46-2 | negative | none | + |

Opinion on fragrance allergens in cosmetic products

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Human evidence: see text | EC 3 value (min; %) | SAR |
|---|------------------------|---------------------------------|----------------------------|------------|
| isomers) | | | | |
| TERPINEOL ACETATE (Isomer mixture) | 8007-35-0 | negative | none | + |
| alpha-TERPINYL ACETATE | 80-26-2 | negative | none | + |
| CITRONELLYL NITRILE | 51566-62-2 | none | none | ++ |
| alpha-CYCLOHEXYLIDENE BENZENEACETONITRILE | 10461-98-0 | none | none | + |
| DECANAL | 112-31-2 | none | none | ++ |
| DIHYDROMYRCENOL | 18479-58-8 | none | none | + |
| 3,7-DIMETHYL-1,6-NONADIEN-3-OL | 10339-55-6 | none | none | ++ |
| 2-ETHYL-4-(2,2,3-TRIMETHYL-3-CYCLOPENTEN-1-YL)-2-BUTEN-1-OL | 28219-61-6 | none | none | + |
| GERANYL ACETATE | 105-87-3 | none | none | ++ |
| HEXAHYDRO-METHANOINDENYL PROPIONATE | 68912-13-0 | none | none | + |
| IONONE isomeric mixture | 8013-90-9 | none | none | + |
| ISOBERGAMATE | 68683-20-5 | none | none | + |
| METHYL DECENOL | 81782-77-6 | none | none | + |
| TRICYCLODECENYL PROPIONATE | 17511-60-3 | none | none | + |
| OXACYCLOHEXADECENONE | 34902-57-3 | none | none | ++ |
| VERDYL ACETATE | 2500-83-6/ 5413-60-5 | none | none | + |
| trans-beta-Damascone | 23726-91-2 | none | none | + |
| gamma-Damascone | 35087-49-1 | none | none | + |
| Citronellal | 106-23-0 | none | none | ++ |
| Phenethyl salicylate | 87-22-9 | none | none | ++ |
| Natural extracts | | | | |
| ACORUS CALAMUS ROOT OIL | 84775-39-3 | Limited | none | |
| CEDRUS DEODARA WOOD OIL | 91771-47-0 | Limited | none | |
| CITRUS AURANTIUM AMARA LEAF OIL | 72968-50-4 | Limited | none | |
| CITRUS TANGERINA ... | 223748-44-5 | Limited | none | |
| CYMBOPOGON NARDUS / WINTERIANUS HERB OIL | 89998-15-2; 91771-61-8 | Limited | none | |
| ILLICIAM VERUM FRUIT OIL | 84650-59-9 | Limited | none | |
| LAVANDULA SPICA | 97722-12-8 | Limited | none | |

Opinion on fragrance allergens in cosmetic products

| INCI name (or, if none exists, perfuming name according to CosIng) | CAS number | Human evidence: see text | EC 3 value (min; %) | SAR |
|---|-----------------------|---------------------------------|----------------------------|------------|
| LITSEA CUBEBA | 90063-59-5 | Limited | none | |
| PELARGONIUM ROSEUM | 90082-55-6 | Limited | none | |
| SALVIA spp. | Diverse | Limited | none | |
| TAGETES PATULA | 91722-29-1 | Limited | none | |
| THYMUS spp. | 84929-51-1 | Limited | none | |
| VETIVERIA ZIZANOIDES | 8016-96-4; 84238-29-9 | Limited | none | |

Regarding the above categorisation of fragrance substances, the following aspects need to be considered when interpreting an outcome other than established contact allergen in humans:

- If human evidence is negative, there is still a potential sensitisation risk, as in this set of substances the number of (consecutive) patients tested was low, i.e. up to a few hundred.
- If EC3 values are given as higher (>) than a certain value (see 8.3), an exact EC3 could not be established, as the substance had been tested in too low concentration(s). In these cases, the substances have not been categorised as 'established contact allergen in animals'.
- For SAR, the categories of prediction are: non-sensitiser (0); possible-sensitiser (+); predicted sensitiser (++); and not predictable (n.p.). (For details see Table 9-3 and Table 9-4). SAR predictions are only considered when human and animal data are limited or missing.
- Several substances are currently banned from the use in cosmetic products by Annex II of the Cosmetics Directive, based on concerns regarding one or more toxicological endpoints. While available clinical evidence regarding this set of substances is listed in Annex I to this opinion, these substances have not further been evaluated.

Fragrance ingredients listed in Table 13-1 clearly have caused disease in man, and based on the clinical experience alone, these 82 substances were classified as established contact allergens in humans, 54 individual chemicals and 28 natural extracts (mixtures of chemicals), including all 26 fragrance allergens identified by SCCNFP in 1999. For a number of other substances, no patch test data were available, but positive animal data, obtained by a validated guideline method (LLNA) addressing hazard, indicate that a – yet not quantified – risk for humans is very likely to exist, given sufficient exposure. In other cases only in a relatively small number of patients has been tested positively ('limited human evidence'). Here, combination with SAR analyses corroborates the conclusion that these substances, too, are sufficiently qualified to be regarded as 'likely fragrance allergens'.

Of those 82 substances identified as established contact allergens in humans, 12 chemicals (listed in Table 13-5) and eight natural extracts are considered of special concern as they have given rise to at least 100 reported cases. These substances pose a particularly high risk of sensitisation to the consumer and are further considered in the answer of question 2. One substance, hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC), was shown to be the cause of allergic contact dermatitis in more than 1500 reported cases since 1999. The number of cases is only those reported in scientific publications, and therefore the actual number of cases is severely under-estimated.

Table 13-5: Established fragrance contact allergens of special concern (single chemicals only).

| |
|---|
| Cinnamal |
| Cinnamyl Alcohol* |
| Citral |
| Coumarin |
| Eugenol* |
| Farnesol* |
| Geraniol* |
| Hydroxycitronellal |
| Hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC) |
| Isoeugenol* |
| Limonene (oxidised) |
| Linalool* (oxidised) |

*including their respective esters

The established contact allergens in animals (Table 13-2) and the likely contact allergens, identified based on a combination of limited evidence from man together with positive SAR predictions (Table 13-3), are predicted to cause disease in man given sufficient exposure.

Information on the presence of all the substances given in Table 13-1, Table 13-2 and Table 13-3 in cosmetic products is important in order to enable aimed testing of patients with contact dermatitis and to diagnose fragrance allergy without delay. Further, this information is important to the sensitised consumer as it will enable them to avoid cosmetic products, which they may not tolerate.

Substances given in Table 13-4 are possible contact allergens and further data are required to judge if these are contact allergens in humans and give rise to contact allergy in consumers.

Conclusions - Question 1

The studies since the SCCNFP Opinion on fragrance allergy in consumers (1) have confirmed that the fragrance allergens currently listed in Annex III, entries 67-92 are still relevant fragrance allergens for the consumers from their exposure to cosmetic products.

The review of the clinical and experimental data shows that many more fragrance substances than those identified in the SCCNFP opinion of 1999 have been shown to be sensitisers in humans. A comprehensive list of established contact allergens in humans is given in Table 13-1.

Moreover, animal experiments indicate that additional fragrance substances can be expected to be contact allergens in humans, although human evidence is currently lacking.

Additionally, limited human and/or animal evidence together with structure activity relationship analysis suggests that other fragrance ingredients may be a cause of concern with regard to their potential of causing contact allergy in humans.

Ingredient listing is important in clinical practice for the management of patients who are allergic to one or more of the listed fragrance chemicals. It is also important for the

patients in order to avoid future exposure to fragrance contact allergens which they may not tolerate.

The SCCS considers that those substances itemised in Table 13-1, Table 13-2 and Table 13-3 represent those fragrance ingredients that the consumer should be made aware of when present in cosmetic products.

Substances known to be transformed (e.g. hydrolysis of esters) to known contact allergens should be treated as equivalent to these known contact allergens. The combined concentration of the alcohol and its ester must be considered regarding exposure. Important indicative, but not exhaustive, examples include isoeugenol and its esters, geraniol and its esters, eugenol and its esters, and linalool and its esters.

13.2. Question 2

Can the SCCS establish any threshold for their safe use based on the available scientific data?

Dose-response relationships exist between exposure to contact allergens and the proportion of consumers who will become sensitised to an allergen (i.e. induction), as well as the proportion who will suffer from allergic contact dermatitis (elicitation). For a number of recognised contact allergens in man, dose-elicitation studies on sensitised individuals are available. These studies indicate that it is in principle possible to derive exposure levels that the majority of sensitised individuals will tolerate. The SCCS considers that thresholds based on elicitation levels in sensitised individuals will be sufficiently low to protect both the majority of sensitised individuals as well as most of the non-sensitised consumers from developing contact allergy and limit the risk of induction.

Among the established chemical fragrance allergens, 12 were identified as posing a high risk of sensitisation to the consumer (Table 13-5), i.e. more than 100 reported cases. For these substances, limitation of exposure would help to protect sensitised consumers from developing allergic contact dermatitis.

In cases where specific data of sufficient quality on threshold levels for a particular allergen are available, these data should be used to set an individual safe threshold. However, when such quality data are not available and a substance has been identified to pose a high risk of sensitisation to the consumer, a general threshold limit can be applied.

Dose-response studies have been performed with only four of these fragrance substances (HICC, isoeugenol, cinnamal and hydroxycitronellal). In addition, such a study has also been performed on chloroatranol, a potent allergen in *Evernia prunastri* and *Evernia furfuracea*. These studies, however, are not adequate to derive safe thresholds for the individual substances directly from the data.

If no adequate data are available, for substances posing a high risk to the consumer (like the 12 listed in Table 13-5), the use of a general threshold may be considered. A threshold of 0.8 µg/cm² has been derived based on a statistical analysis of the available data in the scientific literature, including two fragrance allergens. This corresponds to 0.01% (100 ppm) limit in cosmetic products indicative for safe use. This approximation may hold for weak to strong allergens. However, some strong and extreme sensitisers may require lower individual thresholds. As an example, chloroatranol, present in the natural product *Evernia prunastri* and in *Evernia furfuracea*, has been shown to have an elicitation threshold of 0.0004 µg/cm² under experimental conditions similar to those yielding above results. On the other hand, for very weak sensitisers, this generic threshold may be too conservative.

The model providing the general threshold of 100 ppm has been based on single substances only and no general safe level for the natural extracts of concern can be

identified, but the maximum use concentration applies to the identified fragrance allergens also when present in the natural extract.

Hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC) has been the most frequently reported chemical causing fragrance allergy since the 1999 opinion on fragrance allergy. In total, reports of more than 1500 cases have been published in the scientific literature (see chapter 7.1 and Annex I), which will severely underestimate the actual prevalence in the population. HICC has been shown to be a significant cause of disease as many of those with contact allergy to HICC had also reactions to cosmetics, which contained or were likely to contain HICC. The SCCP concluded in 2003 that 200 ppm of HICC would be tolerated by the majority of sensitised individuals and this level of exposure would have a low potential to induce sensitisation (241). Since 2003 attempts have been made by the fragrance industry to contain the outbreak of HICC allergy, but with no convincing success so far. Recent voluntary restrictions (recommendations to lower use concentrations, at least for some product types, to the level recommended by the SCCS in 2003) are not reflected in available evidence and are considered insufficient. The SCCS considers that the number of cases of HICC allergy documented over the last decade is exceptionally high and that continued exposure to HICC by the consumer is not considered safe, even at concentrations as low as 200 ppm. Chloroatranol and atranol are the main allergenic components of *Evernia prunastri* and *Evernia furfuracea*. The SCCS concluded in 2004 (239) that these should not be present in cosmetic products, due to their exceptionally high sensitisation potential. Attempts to effectively reduce the content of these compounds in "oak moss abs." (300) have largely failed to reduce contact allergy to *Evernia prunastri* and *Evernia furfuracea* and the data presented in this opinion show that the number of cases remains high.

Conclusions - Question 2

There are two components to the safety of fragrance ingredients in terms of contact allergy. First, the need to eliminate or reduce induction of contact allergy (primary prevention), which, when it occurs, is life long. Secondly, the need to eliminate or reduce elicitation reactions (secondary prevention) on the skin of those individuals who are already sensitised. Human dose elicitation experiments have hitherto been performed only for a very small number of substances. It is unlikely that more of these studies will be performed due to experimental and subject recruitment difficulties.

For individual substances, no levels that could be considered safe for the majority of consumers could be established from the available data.

The dose elicitation studies available indicate that a general level of exposure of up to 0.8 µg/cm² (0.01%) may be tolerated by most consumers with contact allergy to fragrance allergens. The SCCS considers that this level of exposure could be efficient in limiting elicitation unless there is substance specific data, either experimental or clinical, to the contrary.

Such a threshold based on elicitation levels in sensitised individuals will be sufficiently low to protect both sensitised individuals as well as most of the non-sensitised consumers from developing contact allergy.

The SCCS is of the opinion that for substances identified as posing a high risk to the consumer and for which no individual thresholds could be derived (Table 13-5), the general threshold of 0.01% would limit the problem of fragrance allergy in the consumer significantly.

It was not possible to provide a safe threshold for natural extracts of concern, as no specific investigations exist and the model providing the general threshold (0.01%) has been based on individual chemicals only. However the SCCS considers that the maximum use concentration applies to the above identified fragrance allergens also when present in the natural extract. This will also reduce the risk of sensitisation and elicitation from natural extracts.

It is important to stress that this general threshold, although limiting the problem, does not preclude that the most sensitive segment of the population may react upon exposure to these levels. Hence, this threshold does not remove the necessity for providing information to the consumer concerning the presence of the fragrance substance in cosmetics.

In the case of hydroxyisohexyl 3-cyclohexene carboxaldehyde, in 2003 the SCCP suggested that levels of up to 200 ppm would be tolerated by the majority of sensitised individuals. Recent voluntary restrictions (recommendations to lower use concentrations, at least for some product types, to the level recommended by the SCCS in 2003) are not reflected in available evidence and are considered insufficient. The SCCS considers that the number of cases of HICC allergy documented over the last decade is exceptionally high and that continued exposure to HICC by the consumer is not considered safe, even at concentrations as low as 200 ppm. Therefore, HICC should not be used in consumer products in order to prevent further cases of contact allergy to HICC and to limit the consequences to those who already have become sensitized. The SCCP concluded in 2004 that chloroatranol and atranol, the main allergenic constituents of *Evernia prunastri* and *Evernia furfuracea*, should not be present in products for the consumer. The persistently high frequency of contact allergy to *Evernia prunastri* and *Evernia furfuracea* noted in eczema patients does point to a persisting problem with exposure to allergenic constituents, despite efforts to reduce the allergen content (296). The SCCS is of the opinion that the presence of the two constituents, chloroatranol and atranol, in cosmetic products are not safe.

13.3. Question 3

Can the SCCS identify substances where processes (e.g. metabolism, oxidation and hydrolysis) may lead to cross-reactivity and new allergens which are relevant for the protection of the consumer?

Many fragrance substances can act as prehapten or prohaptens, forming allergens which are more potent than the parent substance by abiotic and/or metabolic activation, and thus increasing the risk of sensitisation.

Experimental and clinical studies have shown that there are fragrance substances that act as prehapten, i.e. their sensitisation potency is markedly increased by air exposure due to oxidation (autoxidation). Non/low-sensitising compounds are thereby transformed into more potent sensitisers. Limonene, linalool, linalyl acetate, alpha-terpinene and geraniol have all been identified as prehapten. These fragrance substances are common in scented cosmetics as well as in household products. The clinical studies show that the exposure to allergens formed due to autoxidation causes significant contact allergy in consumers. Patch testing with oxidised limonene and oxidised linalool shows that these substances rank among the most common contact allergens.

In the SAR analyses performed in this work by the SCCS, fragrance compounds with structural alerts that indicate that they are possible prehapten have been identified (Table 9-1, Table 9-2). In such cases further thorough investigations are needed. It is also important to investigate the stability of the primary oxidation products (the hydroperoxides) formed from various structures of fragrance compounds. The stability of these compounds can have great impact on the sensitisation potency of the oxidised compound as they are strong sensitisers. However, the secondary oxidation products (aldehydes and epoxides) can also be important sensitisers depending on the overall structure of the compound as was demonstrated for oxidised geraniol.

Air oxidation of prehapten can be prevented to a certain extent by measures during handling and storage of the ingredients and final products to avoid air exposure, and/or by addition of suitable antioxidants. The autoxidation rate depends not only on the compound itself, but also on its purity. The prevention of autoxidation using antioxidants

needs thorough investigation because antioxidants can exert their function by being oxidised instead of the compound that they protect and might thereby be activated to skin sensitising derivatives after oxidation. As antioxidants are now frequently used at elevated concentrations in scented products due to a growing awareness of the problem of autoxidation, there is a risk that sensitisation caused by the antioxidants will rise. One of the most used antioxidants is butylated hydroxytoluene (BHT) which is considered a minimal risk for sensitisation in the concentrations used but nevertheless, with increased concentrations and usage, the risk of sensitisation could increase.

It should be noted that, to decrease the risk for sensitisation in the population, the possibility to reduce the sensitisation potency by preventing autoxidation is important also for a direct acting hapten or prohaptent, if a further activation by air oxidation to more allergenic compounds has been shown.

Based on the clinical data, oxidised limonene and oxidised linalool are allergens of high concern (Table 13-5) which pose a high risk of sensitisation to the consumer. For these substances the presence of the oxidised fraction represented by the peroxide content should not be higher than 10 ppm. Alternatively, the suggested general threshold dose/area of 0.8 µg/cm² (100 ppm in cosmetic products) could be applicable to the total oxidised fraction, i.e. not only peroxides but also secondary oxidation products such as aldehydes and epoxides.

Compounds that are bioactivated by metabolising enzymes to haptens are referred to as prohaptens. Established prohaptens of clinical importance are cinnamyl alcohol, geranial, geraniol, eugenol, isoeugenol and alpha-terpinene.

Table 13-6: Known prehaptens and prohaptens.

| Fragrance substance | Activation by air oxidation | Bioactivation (oxidation) | Bioactivation (hydrolysis) |
|----------------------------|------------------------------------|----------------------------------|-----------------------------------|
| Cinnamyl alcohol | | x | |
| Eugenol | | x | |
| Eugenyl acetate | | x | x |
| Geranial | x | x | |
| Geraniol | x | x | |
| Geranyl acetate | x | x | x |
| Isoeugenol | | x | |
| Isoeugenyl acetate | | x | x |
| Limonene | x | | |
| Linalool | x | | |
| Linalyl acetate | x | | |
| alpha-Terpinene. | x | x | |

When bioactivation occurs, the risk of cross-reactivity should be considered. An increased complexity in the cross-reactivity pattern is obtained when a compound could act both as a prehaptent and a prohaptent. For instance, it is known that cinnamyl alcohol and cinnamal can cross-react due to the formation of common sensitising substances. The same applies to geraniol and citral.

In case derivatives of a fragrance substance are used, it must be taken into account that the derivative could be transformed into the parent or a cross-reacting compound. For such derivatives the same rules as for the corresponding parents should apply, unless the

stability of the derivative has been demonstrated. In particular, hydrolysis of esters to the corresponding alcohols can cause cross-reactions. Acetate esters of eugenol, isoeugenol and geraniol are frequently used in cosmetics.

To be able to predict the sensitisation potency of prohaptens, steps of bioactivation have to be included in the predictive tests.

Activation of individual compounds to various haptens increases the risks of cross-reactivity between chemicals and also causes difficulties in prediction of these risks. Prediction of risks requires sound application of theoretical principles in combination with well designed experimental studies. Based on the acquired knowledge, qualified suggestions using structure activity relationship (SAR) regarding many fragrance substances have been made (Table 9-1 to Table 9-3). However, as the stability of formed oxidation products (mainly hydroperoxides) is important for the sensitisation potency, the SAR hypotheses must be followed by experimental investigations for the actual compounds.

Conclusions - Question 3

Many fragrance substances can act as prehaptens or prohaptens, forming allergens which are more potent than the parent substance by abiotic and/or metabolic activation. Activation can thus increase the risk of sensitisation. Fragrances with published data showing the formation of sensitising compounds by autoxidation, bioactivation or both include the following (see also Table 13-6).

Fragrance substances of clinical importance known to be prehaptens and to form sensitising compounds by air oxidation are limonene, linalool, and linalyl acetate.

Fragrance substances of clinical importance known to be prohaptens and to form sensitising compounds by metabolic transformation are cinnamyl alcohol, eugenol, isoeugenol and isoeugenyl acetate.

Fragrance substances of clinical importance with published data known to be both prehaptens and prohaptens and to form sensitising compounds by air oxidation (prehaptens) and by metabolic transformation are geraniol and alpha-terpinene.

A fragrance substance that sensitises without activation but forms more potent sensitising compounds by air oxidation and also by metabolic transformation is geraniol (one isomer of citral).

In the case of prehaptens, it is possible to prevent activation outside the body to a certain extent by different measures, e.g. prevention of air exposure during handling and storage of the ingredients and the final product and by the addition of suitable antioxidants. When antioxidants are used, care should be taken that they will not be activated themselves and thereby form new sensitisers.

The possibility to reduce the sensitisation potency by preventing air oxidation is important also for a direct acting hapten or prohaptens, if a further activation by air oxidation to more allergenic compounds has been shown.

In the case of prohaptens, the possibility to become activated is inherent to the molecule and activation cannot be avoided by extrinsic measures. Activation processes increase the risk for cross-reactivity between fragrance substances. Cross-reactivity has been shown for certain alcohols and their corresponding aldehydes, i.e. between geraniol and geraniol (citral) and between cinnamyl alcohol and cinnamal.

Cross-reactivity is also expected between ester derivatives and their parent alcohols, as the esters will be hydrolysed by esterases in the skin. Esters of important contact allergens that can be activated by hydrolysis in the skin are isoeugenyl acetate, eugenyl acetate and geranyl acetate which all are known to be used as fragrance ingredients.

The substances presented above are based on current knowledge and should be seen as indicative and illustrative of the general problem. As substances with structural alerts for acting as pro- and/or prehapten are quite common among the fragrance substances listed (see Tables 9-1 and 9-2), the possibility for activation to generate new potent allergens should be considered.

The SCCS is of the opinion that substances known to be transformed (e.g. by oxidation either via air oxidation or via bioactivation) to known contact allergens should be treated as equivalent to these contact allergens, i.e. the same restrictions and other regulatory requirements should apply, unless specific data exist that allow for an individual assessment. Important indicative examples include limonene, linalool, linalyl acetate, geraniol, geranial, alpha-terpinene, eugenol, isoeugenol and cinnamyl alcohol.

14. List of abbreviations

| | |
|-------------|--|
| ACD | Allergic contact dermatitis |
| alc. | Alcohol (as vehicle) |
| CI | Confidence interval |
| CLP | Classification, labelling and packaging |
| coloph. | Colophonium |
| DCs | Dendritic cells |
| EC | European Commission |
| ESSCA | European Surveillance System on Contact Allergies |
| EDT | Eau de toilette |
| EDP | Eau de perfume |
| EU | European Union |
| FM | Fragrance mix |
| GC | Gas chromatography |
| GPMT | Guinea pig maximisation test |
| HICC | Hydroxyisohexyl 3-cyclohexene carboxaldehyde |
| HRIPT | Human repeat insult patch test |
| IFRA | International Fragrance Association (www.ifraorg.org) |
| IVDK | Information Network of Departments of Dermatology (www.ivdk.gwdg.de) |
| INCI | International Nomenclature on Cosmetic Ingredients |
| LCs | Langerhans cells |
| LLNA | Local lymph node assay |
| MPR | <i>Myroxylon pereirae</i> resin |
| NACDG | North American Contact Dermatitis Group |
| OECD | Organization of Economic Co-operation and Development |
| ox. | oxidised |
| pet. | Petrolatum (as vehicle) |
| ppm | parts per million (10000 ppm = 1%) |
| PPV | Positive predictive value |
| PR | Prevalence ratio |
| PT(ed)(ing) | Patch test(ed) (ing) |
| QMM | Quantitative mechanistic model |
| QRA | Quantitative risk assessment |
| (Q)SAR | (Quantitative) structure activity relationship |
| REACH | Registration, Evaluation, Authorisation and restriction of CHemicals |
| RIFM | Research Institute for Fragrance Materials (www.rifm.org/) |

Opinion on fragrance allergens in cosmetic products

| | |
|--------|---|
| ROAT | Repeated open application test |
| SC | Single constituents (of one of the fragrance mixes) |
| SCCS | Scientific Committee on Consumer Safety |
| SCCNFP | Scientific Committee on Cosmetic Products and Non-Food Products |
| SCCP | Scientific Committee on Consumer Products |
| UK | United Kingdom |
| US(A) | United States (of America) |
| UV | Ultraviolet |

15. References

1. SCCNFP. The Scientific Committee on Cosmetic Products and Non-Food Products intended for Consumers: Opinion concerning Fragrance Allergy in Consumers. A Review of the Problem. Analysis of the Need for appropriate Consumer Information and Identification of Consumer Allergens, adopted 8 December 1999. *SCCNFP/0017/98 Final* 1999:
2. Rustemeyer T, van Hoogstraten I M W, von Blomberg M E, Scheper R J. Mechanisms in Allergic Contact Dermatitis. In: Frosch P J, Menné T, Lepoittevin J P, eds. *Contact Dermatitis*. Heidelberg: Springer, 2006:
3. Johansen J D, Andersen T F, Kjoller M, Veien N, Avnstorp C, Andersen K E, Menne T. Identification of risk products for fragrance contact allergy: a case-referent study based on patients' histories. *Am J Contact Dermat* 1998; 9: 80-86.
4. Marks J G, Belsito D V, DeLeo V A, et al. North American Contact Dermatitis Group patch test results for the detection of delayed-type hypersensitivity to topical allergens. *J Am Acad Dermatol* 1998; 38: 911-918.
5. Johansen J D, Rastogi S C, Menné T. Contact allergy to popular perfumes: assessed by patch test, use test and chemical analysis. *Br J Dermatol* 1996; 135: 419-422.
6. Johansen J D, Rastogi S C, Andersen K E, Menne T. Content and reactivity to product perfumes in fragrance mix positive and negative eczema patients. A study of perfumes used in toiletries and skin-care products. *Contact Dermatitis* 1997; 36: 291-296.
7. de Groot A C, Frosch P J. Adverse reactions to fragrances. A clinical review. *Contact Dermatitis* 1997; 36: 57-86.
8. Cronin E. Contact Dermatitis. *Churchill Livingstone, Edinburgh* 1980:
9. Johansen J D, Andersen T F, Veien N, Avnstorp C, Andersen K E, Menne T. Patch testing with markers of fragrance contact allergy. Do clinical tests correspond to patients' self-reported problems? *Acta Derm Venereol* 1997; 77: 149-153.
10. Johansen J D, Rastogi S C, Bruze M, Andersen K E, Frosch P, Dreier B, Lepoittevin J P, White I, Menne T. Deodorants: a clinical provocation study in fragrance-sensitive individuals. *Contact Dermatitis* 1998; 39: 161-165.
11. Lammintausta K, Kalimo K, Havu V K. Occurrence of contact allergy and hand eczemas in hospital wet work. *Contact Dermatitis* 1982; 8: 84-90.
12. Meding B. Epidemiology of Hand Eczema in an Industrial City. *Acta Dermatol Venerol (Stockh) Suppl* 1990; 153: 2-43.
13. Heydorn S, Johansen J D, Andersen K E, Bruze M, Svedman C, White I R, Basketter D A, Menne T. Fragrance allergy in patients with hand eczema - a clinical study. *Contact Dermatitis* 2003; 48: 317-323.
14. Buckley D A, Rycroft R J, White I R, McFadden J P. Contact allergy to individual fragrance mix constituents in relation to primary site of dermatitis. *Contact Dermatitis* 2000; 43: 304-305.
15. Heydorn S, Menne T, Johansen J D. Fragrance allergy and hand eczema - a review. *Contact Dermatitis* 2003; 48: 59-66.
16. Wöhrl S, Hemmer W, Focke M, Götz M, Jarisch R. The significance of fragrance mix, balsam of Peru, colophony and propolis as screening tools in the detection of fragrance allergy. *Br J Dermatol* 2001; 145: 268-273.

17. Edman B. The influence of shaving method on perfume allergy. *Contact Dermatitis* 1994; 31: 291-292.
18. Heydorn S, Menne T, Andersen K E, Bruze M, Svedman C, White I R, Basketter D A. Citral a fragrance allergen and irritant. *Contact Dermatitis* 2003; 49: 32-36.
19. Rothenborg H W, Menne T, Sjolín K E. Temperature dependent primary irritant dermatitis from lemon perfume. *Contact Dermatitis* 1977; 3: 37-48.
20. Tanaka S, Matsumoto Y, Dlova N, Ostlere L S, Goldsmith P C, Rycroft R J, Basketter D A, White I R, Banerjee P, McFadden J P. Immediate contact reactions to fragrance mix constituents and Myroxylon pereirae resin. *Contact Dermatitis* 2004; 51: 20-21.
21. Hausen B M. Contact allergy to balsam of Peru. II. Patch test results in 102 patients with selected balsam of Peru constituents. *Am J Contact Dermat* 2001; 12: 93-102.
22. Katsarou A, Armenaka M, Ale I, Koufou V, Kalogeromitros D. Frequency of immediate reactions to the European standard series. *Contact Dermatitis* 1999; 41: 276-279.
23. Nakayama H. Perfume allergy and cosmetic dermatitis (in Japanese). *Jpn J Dermatol* 1974; 84: 659-667.
24. Nakayama H, Harada R, Toda M. Pigmented cosmetic dermatitis. *Int J Dermatol* 1981; 15: 673-675.
25. Cronin E. Photosensitivity to musk ambrette. *Contact Dermatitis* 1984; 11: 88-92.
26. Darvay A, White I R, Rycroft R J, Jones A B, Hawk J L, McFadden J P. Photoallergic contact dermatitis is uncommon. *Br J Dermatol* 2001; 145: 597-601.
27. Wang L, Sterling B, Don P. Berloque dermatitis induced by "Florida water". *Cutis* 2002; 70: 29-30.
28. Elberling J, Linneberg A, Dirksen A, Johansen J D, Frolund L, Madsen F, Nielsen N H, Mosbech H. Mucosal symptoms elicited by fragrance products in a population-based sample in relation to atopy and bronchial hyper-reactivity. *Clin Exp Allergy* 2005; 35: 75-81.
29. Kumar P, Caradonna-Graham V M, Gupta S, Cai X, Rao P N, Thompson J. Inhalation challenge effects of perfume scent strips in patients with asthma. *Ann Allergy Asthma Immunol* 1995; 75: 429-433.
30. Millqvist E, Bende M, Lowhagen O. Sensory hyperreactivity--a possible mechanism underlying cough and asthma-like symptoms. *Allergy* 1998; 53: 1208-1212.
31. Elberling J, Linneberg A, Mosbech H, Dirksen A, Frolund L, Madsen F, Nielsen N H, Johansen J D. A link between skin and airways regarding sensitivity to fragrance products? *Br J Dermatol* 2004; 151: 1197-1203.
32. Lindberg M, Matura M. Patch Testing. In: Johansen J D, Frosch P, Lepoittevin J P, eds. *Contact Dermatitis*. Heidelberg etc., : Springer, 2011: 439-464.
33. Basketter D. Diagnostic patch testing - does it have a wider relevance? *Contact Dermatitis* 2012; 67: 1-2.
34. Larsen W G. Perfume Dermatitis. A Study of 20 Patients. *Arch Dermatol* 1977; 113: 623-626.
35. Frosch P J, Pirker C, Rastogi S C, Andersen K E, Bruze M, Svedman C, Goossens A, White I R, Uter W, Arnau E G, Lepoittevin J P, Menne T, Johansen J D. Patch testing with a new fragrance mix detects additional patients sensitive to perfumes and missed by the current fragrance mix. *Contact Dermatitis* 2005; 52: 207-215.

36. Frosch P J, Rastogi S C, Pirker C, Brinkmeier T, Andersen K E, Bruze M, Svedman C, Goossens A, White I R, Uter W, Arnau E G, Lepoittevin J P, Johansen J D, Menne T. Patch testing with a new fragrance mix - reactivity to the individual constituents and chemical detection in relevant cosmetic products. *Contact Dermatitis* 2005; 52: 216-225.
37. Bruze M, Andersen K E, Goossens A. Recommendation to include fragrance mix 2 and hydroxyisohexyl 3-cyclohexene carboxaldehyde (LyrAl) in the European baseline patch test series. *Contact Dermatitis* 2008; 58: 129-133.
38. Lindberg M, Edman B, Fischer T, Stenberg B. Time trends in Swedish patch test data from 1992 to 2000. A multi-centre study based on age- and sex-adjusted results of the Swedish standard series. *Contact Dermatitis* 2007; 56: 205-210.
39. Temesvari E, Nemeth I, Baló-Banga M J, Husz S, Kohanka V, Somos Z, Judak R, Remenyik E V, Szegedi A, Nebenfuhrer L, Meszaros C, Horvath A. Multicentre study of fragrance allergy in Hungary. Immediate and late type reactions. *Contact Dermatitis* 2002; 46: 325-330.
40. Machovcova A, Dastychova E, Kostalova D, Vojtechovska A, Reslova J, Smejkalova D, Vaneckova J, Vocilkova A. Common contact sensitizers in the Czech Republic. Patch test results in 12,058 patients with suspected contact dermatitis*. *Contact Dermatitis* 2005; 53: 162-166.
41. Lunder T, Kansky A. Increase in contact allergy to fragrances: patch-test results 1989-1998. *Contact Dermatitis* 2000; 43: 107-109.
42. Schnuch A, Lessmann H, Geier J, Frosch P J, Uter W. Contact allergy to fragrances: frequencies of sensitization from 1996 to 2002. Results of the IVDK*. *Contact Dermatitis* 2004; 50: 65-76.
43. Uter W, Geier J, Frosch P J, Schnuch A. Contact allergy to fragrances: current patch test results (2005 to 2008) from the IVDK network. *Contact Dermatitis* 2010; 63: 254-261.
44. van Oosten E J, Schuttelaar M L, Coenraads P J. Clinical relevance of positive patch test reactions to the 26 EU-labelled fragrances. *Contact Dermatitis* 2009; 61: 217-223.
45. deGroot A C, Coenraads P J, Bruynzeel D P, Jagtman B A, van Ginkel C J W, Noz K, van der Valk P G M, Pavel S, Vink J, Weyland J W. Routine patch testing with fragrance chemicals in The Netherlands. *Contact Dermatitis* 2000; 42: 184-185.
46. Hendriks S A, van Ginkel C J. Evaluation of the fragrance mix in the European standard series. *Contact Dermatitis* 1999; 41: 161-162.
47. Nardelli A, Carbonez A, Ottoy W, Drieghe J, Goossens A. Frequency of and trends in fragrance allergy over a 15-year period. *Contact Dermatitis* 2008; 58: 134-141.
48. Brites M M, Goncalo M, Figueiredo A. Contact allergy to fragrance mix--a 10-year study. *Contact Dermatitis* 2000; 43: 181-182.
49. Cuesta L, Silvestre J F, Toledo F, Lucas A, Perez-Crespo M, Ballester I. Fragrance contact allergy: a 4-year retrospective study. *Contact Dermatitis* 2010; 63: 77-84.
50. Katsarma G, Gawkrödger D J. Suspected fragrance allergy requires extended patch testing to individual fragrance allergens. *Contact Dermatitis* 1999; 41: 193-197.
51. Buckley D A, Basketter D A, Kan-King-Yu D, White I R, White J L, McFadden J P. Atopy and contact allergy to fragrance: allergic reactions to the fragrance mix I (the Larsen mix). *Contact Dermatitis* 2008; 59: 220-225.
52. Thyssen J P, Carlsen B C, Menne T, Johansen J D. Trends of contact allergy to fragrance mix I and Myroxylon pereirae among Danish eczema patients tested between 1985 and 2007. *Contact Dermatitis* 2008; 59: 238-244.

53. Uter W, Hegewald J, Aberer W, Ayala F, Bircher A J, Brasch J, Coenraads P J, Schuttelaar M L, Elsner P, Fartasch M, Mahler V, Belloni Fortina A, Frosch P J, Fuchs T, Johansen J D, Menne T, Jolanki R, Krecisz B, Kiec-Swierczynska M, Larese F, Orton D, Peserico A, Rantanen T, Schnuch A. The European standard series in 9 European countries, 2002/2003 - First results of the European Surveillance System on Contact Allergies. *Contact Dermatitis* 2005; 53: 136-145.
54. Hegewald J, Uter W, Aberer W, Ayala F, Beliauskiene A, Belloni Fortina A, Bircher A, Brasch J, Chowdhury M M, Coenraads P J, Schuttelaar M-L, Elsner P, English J, Fartasch M, Mahler V, Frosch P J, Fuchs T, Gawkrödger D J, Giménez-Arnau A M, Green C M, Johansen J D, Menné T, Jolanki R, King C M, Krecisz B, Kiec-Swierczynska M, Larese F, Ormerod A D, Orton D, Peserico A, Rantanen T, Rustemeyer T, Sansom J E, Statham B N, Corradin M T, Wallnofer W, Wilkinson M, Schnuch A. The European Surveillance System of Contact Allergies (ESSCA): results of patch testing the standard series, 2004. *J Eur Acad Dermatol Venereol* 2008; 22: 174-181.
55. Uter W, Räsmsch C, Aberer W, Ayala F, Balato A, Beliauskiene A, Fortina A B, Bircher A, Brasch J, Chowdhury M M, Coenraads P J, Schuttelaar M L, Cooper S, Corradin M T, Elsner P, English J S, Fartasch M, Mahler V, Frosch P J, Fuchs T, Gawkrödger D J, Gimenez-Arnau A M, Green C M, Horne H L, Jolanki R, King C M, Krecisz B, Kiec-Swierczynska M, Ormerod A D, Orton D I, Peserico A, Rantanen T, Rustemeyer T, Sansom J E, Simon D, Statham B N, Wilkinson M, Schnuch A. The European baseline series in 10 European Countries, 2005/2006--results of the European Surveillance System on Contact Allergies (ESSCA). *Contact Dermatitis* 2009; 61: 31-38.
56. An S, Lee A Y, Lee C H, Kim D W, Hahm J H, Kim K J, Moon K C, Won Y H, Ro Y S, Eun H C. Fragrance contact dermatitis in Korea: a joint study. *Contact Dermatitis* 2005; 53: 320-323.
57. Hussain I, Rani Z, Rashid T, Haroon T S. Suitability of the European standard series of patch test allergens in Pakistani patients. *Contact Dermatitis* 2002; 46: 50-51.
58. Gupta N, Shenoj S D, Balachandran C. Fragrance sensitivity in allergic contact dermatitis. *Contact Dermatitis* 1999; 40: 53-54.
59. Freireich-Astman M, David M, Trattner A. Standard patch test results in patients with contact dermatitis in Israel: age and sex differences. *Contact Dermatitis* 2007; 56: 103-107.
60. Lazarov A. European Standard Series patch test results from a contact dermatitis clinic in Israel during the 7-year period from 1998 to 2004. *Contact Dermatitis* 2006; 55: 73-76.
61. Kashani M N, Gorouhi F, Behnia F, Nazemi M J, Dowlati Y, Firooz A. Allergic contact dermatitis in Iran. *Contact Dermatitis* 2005; 52: 154-158.
62. Akyol A, Boyvat A, Peksari Y, Gurgey E. Contact sensitivity to standard series allergens in 1038 patients with contact dermatitis in Turkey. *Contact Dermatitis* 2005; 52: 333-337.
63. Lu X, Li L F, Wang W, Wang J. A clinical and patch test study of patients with positive patch test reactions to fragrance mix in China. *Contact Dermatitis* 2005; 52: 188-191.
64. Belsito D V, Fowler J F, Jr., Sasseville D, Marks J G, Jr., De Leo V A, Storrs F J. Delayed-type hypersensitivity to fragrance materials in a select North American population. *Dermatitis* 2006; 17: 23-28.
65. Zug K A, Warshaw E M, Fowler J F, Jr., Maibach H I, Belsito D L, Pratt M D, Sasseville D, Storrs F J, Taylor J S, Mathias C G, Deleo V A, Rietschel R L, Marks J. Patch-test results of the North American Contact Dermatitis Group 2005-2006. *Dermatitis* 2009; 20: 149-160.

66. Bruynzeel D P, Diepgen T L, Andersen K E, Brandao F M, Bruze M, Frosch P J, Goossens A, Lahti A, Mahler V, Maibach H I, Menne T, Wilkinson J D. Monitoring the European standard series in 10 centres 1996-2000. *Contact Dermatitis* 2005; 53: 146-149.
67. Frosch P J, Johansen J D, Menne T, Pirker C, Rastogi S C, Andersen K E, Bruze M, Goossens A, Lepoittevin J P, White I R. Further important sensitizers in patients sensitive to fragrances. I. Reactivity to 14 frequently used chemicals. *Contact Dermatitis* 2002; 47: 78-85.
68. Krautheim A, Uter W, Frosch P, Schnuch A, Geier J. Patch testing with fragrance mix II: results of the IVDK 2005-2008. *Contact Dermatitis* 2010; 63: 262-269.
69. Heisterberg M V, Andersen K E, Avnstorp C, al. e. Fragrance mix II in the baseline series contributes significantly to detection of fragrance allergy. *Contact Dermatitis* 2010: (accepted):
70. Frosch P J, Pilz B, Andersen K E, Burrows D, Camarasa J G, et al. Patch testing with fragrances: results of a multicenter study of the European Environmental and Contact Dermatitis Research Group with 48 frequently used constituents of perfumes. *Contact Dermatitis* 1995; 33: 333-342.
71. Frosch P J, Johansen J D, Menne T, Rastogi S C, Bruze M, Andersen K E, Lepoittevin J P, Gimenez Arnau E, Pirker C, Goossens A, White I R. Lyral is an important sensitizer in patients sensitive to fragrances. *Br J Dermatol* 1999; 141: 1076-1083.
72. Beliauskienė A, Valiukeviciene S, Uter W, Schnuch A. The European baseline series in Lithuania: results of patch testing in consecutive adult patients. *Journal of the European Academy of Dermatology and Venereology : JEADV* 2011; 25: 59-63.
73. Geier J, Brasch J, Schnuch A, Lessmann H, Pirker C, Frosch P J. Lyral has been included in the patch test standard series in Germany. *Contact Dermatitis* 2002; 46: 295-297.
74. Schnuch A, Uter W, Geier J, Lessmann H, Frosch P J. Sensitization to 26 fragrances to be labelled according to current European regulation. Results of the IVDK and review of the literature. *Contact Dermatitis* 2007; 57: 1-10.
75. Api A M. Only Peru Balsam extracts or distillates are used in perfumery. *Contact Dermatitis* 2006; 54: 179.
76. Avalos-Peralta P, Garcia-Bravo B, Camacho F M. Sensitivity to Myroxylon pereirae resin (balsam of Peru). A study of 50 cases. *Contact Dermatitis* 2005; 52: 304-306.
77. Cachao P, Menezes Brandao F, Carmo M, Frazao S, Silva M. Allergy to oil of turpentine in Portugal. *Contact Dermatitis* 1986; 14: 205-208.
78. Lear J T, Heagerty A H M, Tan B B, et al. Transient re-emergence of oil of turpentine allergy in the pottery industry. *Contact Dermatitis* 1996; 34: 169-172.
79. Treudler R, Richter G, Geier J, Schnuch A, Orfanos C E, Tebbe B. Increase in sensitization to oil of turpentine: recent data from a multicenter study on 45,005 patients from the German-Austrian Information Network of Departments of Dermatology (IVDK). *Contact Dermatitis* 2000; 42: 68-73.
80. Schäfer T, Böhler E, Ruhdorfer S, Weigl L, Wessner D, Filipiak B, Wichmann H E, Ring J. Epidemiology of contact allergy in adults. *Allergy* 2001; 56: 1192-1196.
81. Meding B, Swanbeck G. Occupational hand eczema in an industrial city. *Contact Dermatitis* 1990; 22: 13-23.
82. Nielsen N H, Menné T. Allergic contact sensitization in an unselected Danish population - the Glostrup allergy study, Denmark. *Acta Dermatol Venerol (Stockh)* 1992; 72: 456-460.

83. Meneghini C L, Sertoli A, Nava C, Angelini G, Francalani S, Foti C, Moroni P. Irritant contact dermatitis of the hands in housewives. In: Elsner P, Maibach H I, eds. *Irritant Dermatitis New Clinical and Experimental Aspects Curr Probl Dermatol*. Basel: Karger, 1995: 41-48.
84. Seidenari S, Manzini B M, Danese P, Motolese A. Patch and prick test study of 593 healthy subjects. *Contact Dermatitis* 1990; 23: 162-167.
85. Mørtz C G, Bindslev-Jensen C, Lauritsen J, Andersen K E. Allergic contact sensitization in 8th grade school children in Odense, Denmark. . *Abstract presented at the Jadassohn Centenary Congress, London 9-12 Oct 1996* 1996:
86. Guin J D, Berry V K. Perfume sensitivity in adult females. A study of contact sensitivity to a perfume mix in two groups of student nurses. *J Am Acad Dermatol* 1980; 3: 299-302.
87. Nielsen N H, Linneberg A, Menne T, Madsen F, Frolund L, Dirksen A, Jorgensen T. Allergic contact sensitization in an adult Danish population: two cross-sectional surveys eight years apart (the Copenhagen Allergy Study). *Acta Derm Venereol* 2001; 81: 31-34.
88. Thyssen J P, Linneberg A, Menne T, Nielsen N H, Johansen J D. The prevalence and morbidity of sensitization to fragrance mix I in the general population. *Br J Dermatol* 2009; 161: 95-101.
89. Buckley D A, Rycroft R J, White I R, McFadden J P. The frequency of fragrance allergy in patch-tested patients increases with their age. *Br J Dermatol* 2003; 149: 986-989.
90. Uter W, Schnuch A, Geier J, Pfahlberg A, Gefeller O. Association between occupation and contact allergy to the fragrance mix: a multifactorial analysis of national surveillance data. *Occup Environ Med* 2001; 58: 392-398.
91. Dotterud L K, Smith-Sivertsen T. Allergic contact sensitization in the general adult population: a population-based study from Northern Norway. *Contact Dermatitis* 2007; 56: 10-15.
92. Smith-Sivertsen T, Dotterud L K, Lund E. Nickel allergy and its relationship with local nickel pollution, ear piercing, and atopic dermatitis: a population-based study from Norway. *J Am Acad Dermatol* 1999; 40: 726-735.
93. White J M, Gilmour N J, Jeffries D, Duangdeeden I, Kullavanijaya P, Basketter D A, McFadden J P. A general population from Thailand: incidence of common allergens with emphasis on para-phenylenediamine. *Clin Exp Allergy* 2007; 37: 1848-1853.
94. Bruze M. What is a relevant contact allergy? *Contact Dermatitis* 1990; 23: 224-225.
95. Ale I, Maibach H I. Clinical Relevance in Allergic Contact Dermatitis. An algorithmic approach. *Derm Beruf Umwelt* 1995; 43: 119-121.
96. Wahlberg J E, Lindberg M. Patch Testing. In: Frosch P J, Menné T, Lepoittevin J P, eds. *Contact Dermatitis*. Berlin: Springer, 2006: 365-390.
97. Frosch P J, Johansen J D, Menne T, Pirker C, Rastogi S C, Andersen K E, Bruze M, Goossens A, Lepoittevin J P, White I R. Further important sensitizers in patients sensitive to fragrances. II. Reactivity to essential oils. *Contact Dermatitis* 2002; 47: 279-287.
98. Rothenborg H W, Hjorth N. Allergy to perfumes from toilet soaps and detergents in patients with dermatitis. *Arch Dermatol* 1968; 97: 417-421.
99. Hannuksela M, Kousa M, Pirila V. Allergy to ingredients of vehicles. *Contact Dermatitis* 1976; 2: 105-110.
100. Johansen J D, Andersen K E, Menné T. Quantitative aspects of isoeugenol contact allergy assessed by use and patch tests. *Contact Dermatitis* 1996; 34: 414-418.

101. Johansen J D, Andersen K E, Rastogi S C, Menne T. Threshold responses in cinnamic-aldehyde-sensitive subjects: results and methodological aspects. *Contact Dermatitis* 1996; 34: 165-171.
102. Jorgensen P H, Jensen C D, Rastogi S, Andersen K E, Johansen J D. Experimental elicitation with hydroxyisohexyl-3-cyclohexene carboxaldehyde-containing deodorants. *Contact Dermatitis* 2007; 56: 146-150.
103. Bruze M, Johansen J D, Andersen K E, Frosch P, Lepoittevin J P, Rastogi S, Wakelin S, White I, Menne T. Deodorants: an experimental provocation study with cinnamic aldehyde. *J Am Acad Dermatol* 2003; 48: 194-200.
104. Svedman C, Bruze M, Johansen J D, Andersen K E, Goossens A, Frosch P J, Lepoittevin J P, Rastogi S, White I R, Menne T. Deodorants: an experimental provocation study with hydroxycitronellal. *Contact Dermatitis* 2003; 48: 217-223.
105. Schnuch A, Uter W, Dickel H, Szliska C, Schliemann S, Eben R, Rueff F, Gimenez-Arnau A, Löffler H, Aberer W, Frambach Y, Worm M, Niebuhr M, Hillen U, Martin V, Jappe U, Frosch P J, Mahler V. Quantitative patch and repeated open application testing in hydroxyisohexyl 3-cyclohexene carboxaldehyde sensitive-patients. *Contact Dermatitis* 2009; 61: 152-162.
106. Uter W, Geier J, Schnuch A, Frosch P J. Patch test results with patients' own perfumes, deodorants and shaving lotions: results of the IVDK 1998-2002. *J Eur Acad Dermatol Venereol* 2007; 21: 374-379.
107. Basra M K, Fenech R, Gatt R M, Salek M S, Finlay A Y. The Dermatology Life Quality Index 1994-2007: a comprehensive review of validation data and clinical results. *Br J Dermatol* 2008; 159: 997-1035.
108. Skoet R, Zachariae R, Agner T. Contact dermatitis and quality of life: a structured review of the literature. *Br J Dermatol* 2003; 149: 452-456.
109. Moberg C, Alderling M, Meding B. Hand eczema and quality of life: a population-based study. *Br J Dermatol* 2009; 161: 397-403.
110. Meding B, Swanbeck G. Consequences of having hand eczema. *Contact Dermatitis* 1990; 23: 6-14.
111. Agner T, Andersen K E, Brandao F M, Bruynzeel D P, Bruze M, Frosch P, Goncalo M, Goossens A, Le Coz C J, Rustemeyer T, White I R, Diepgen T. Contact sensitisation in hand eczema patients-relation to subdiagnosis, severity and quality of life: a multi-centre study. *Contact Dermatitis* 2009; 61: 291-296.
112. Lysdal S H, Johansen J D. Fragrance contact allergic patients: strategies for use of cosmetic products and perceived impact on life situation. *Contact Dermatitis* 2009; 61: 320-324.
113. Meding B, Wrangsjo K, Jarvholm B. Fifteen-year follow-up of hand eczema: predictive factors. *J Invest Dermatol* 2005; 124: 893-897.
114. Hald M, Agner T, Blands J, Ravn H, Johansen J D. Allergens associated with severe symptoms of hand eczema and a poor prognosis. *Contact Dermatitis* 2009; 61: 101-108.
115. Wijnhoven S W P, Ezendam J, Schuur A G, van Loveren H, van Engelen J G M. *Allergens in consumer products. RIVM Reprot 320025001*. Bilthoven: Institute for Public Health and the Environment, 2008.
116. Schnuch A, Aberer W, Agathos M, Becker D, Brasch J, Elsner P, Frosch P J, Fuchs T, Geier J, Hillen U, Löffler H, Mahler V, Richter G, Szliska C. Durchführung des Epikutantests mit Kontaktallergenen. Leitlinien der Deutschen Dermatologischen Gesellschaft; Deutschen Gesellschaft für Allergie und klinische Immunologie. *J Dtsch Dermatol Ges* 2008; 6: 770-775.

117. Scheinmann P L. The foul side of fragrance-free products: What every clinician should know about managing patients with fragrance allergy. *J Am Acad Dermatol* 1999; 41: 1020-1024.
118. Hagvall L, Backtorp C, Norrby P O, Karlberg A T, Borje A. Experimental and theoretical investigations of the autoxidation of geranial: a dioxolane hydroperoxide identified as a skin sensitizer. *Chemical research in toxicology* 2011; 24: 1507-1515.
119. Hagvall L, Backtorp C, Svensson S, Nyman G, Borje A, Karlberg A T. Fragrance compound geraniol forms contact allergens on air exposure. Identification and quantification of oxidation products and effect on skin sensitization. *Chem Res Toxicol* 2007; 20: 807-814.
120. Hagvall L, Baron J M, Borje A, Weidolf L, Merk H, Karlberg A T. Cytochrome P450-mediated activation of the fragrance compound geraniol forms potent contact allergens. *Toxicol Appl Pharmacol* 2008; 233: 308-313.
121. Brared Christensson J, Matura M, Backtorp C, Borje A, Nilsson J L, Karlberg A T. Hydroperoxides form specific antigens in contact allergy. *Contact Dermatitis* 2006; 55: 230-237.
122. Karlberg A T, Bergstrom M A, Borje A, Luthman K, Nilsson J L. Allergic contact dermatitis--formation, structural requirements, and reactivity of skin sensitizers. *Chem Res Toxicol* 2008; 21: 53-69.
123. Karlberg A T, Boman A, Melin B. Animal experiments on the allergenicity of d-limonene--the citrus solvent. *Ann Occup Hyg* 1991; 35: 419-426.
124. Karlberg A T, Magnusson K, Nilsson U. Air oxidation of d-limonene (the citrus solvent) creates potent allergens. *Contact Dermatitis* 1992; 26: 332-340.
125. Karlberg A T, Shao L P, Nilsson U, Gafvert E, Nilsson J L. Hydroperoxides in oxidized d-limonene identified as potent contact allergens. *Arch Dermatol Res* 1994; 286: 97-103.
126. Sköld M, Börje A, Matura M, Karlberg A T. Studies on the autoxidation and sensitizing capacity of the fragrance chemical linalool, identifying a linalool hydroperoxide. *Contact Dermatitis* 2002; 46: 267-272.
127. Sköld M, Börje A, Harambasic E, Karlberg A T. Contact allergens formed on air exposure of linalool. Identification and quantification of primary and secondary oxidation products and the effect on skin sensitization. *Chem Res Toxicol* 2004; 17: 1697-1705.
128. Sköld M, Hagvall L, Karlberg A T. Autoxidation of linalyl acetate, the main component of lavender oil, creates potent contact allergens. *Contact Dermatitis* 2008; 58: 9-14.
129. Rudback J, Bergstrom M A, Borje A, Nilsson U, Karlberg A T. alpha-Terpinene, an antioxidant in tea tree oil, autoxidizes rapidly to skin allergens on air exposure. *Chem Res Toxicol* 2012; 25: 713-721.
130. Karlberg A T, Dooms-Gossens A. Contact allergy to oxidized d-limonene among dermatitis patients. *Contact Dermatitis* 1997; 36: 201-206.
131. Matura M, Goossens A, Bordalo O, Garcia-Bravo B, Magnusson K, Wrangsjo K, Karlberg A T. Oxidized citrus oil (R-limonene): a frequent skin sensitizer in Europe. *J Am Acad Dermatol* 2002; 47: 709-714.
132. Matura M, Goossens A, Bordalo O, Garcia-Bravo B, Magnusson K, Wrangsjo K, Karlberg A T. Patch testing with oxidized R-(+)-limonene and its hydroperoxide fraction. *Contact Dermatitis* 2003; 49: 15-21.
133. Matura M, Skold M, Borje A, Andersen K E, Bruze M, Frosch P, Goossens A, Johansen J D, Svedman C, White I R, Karlberg A T. Selected oxidized fragrance terpenes are common contact allergens. *Contact Dermatitis* 2005; 52: 320-328.

134. Matura M, Skold M, Borje A, Andersen K E, Bruze M, Frosch P, Goossens A, Johansen J D, Svedman C, White I R, Karlberg A T. Not only oxidized R-(+)- but also S-(-)-limonene is a common cause of contact allergy in dermatitis patients in Europe. *Contact Dermatitis* 2006: 55: 274-279.
135. Christensson J B, Matura M, Gruvberger B, Bruze M, Karlberg A T. Linalool--a significant contact sensitizer after air exposure. *Contact Dermatitis* 2010: 62: 32-41.
136. Sköld M, Karlberg A T, Matura M, Börje A. The fragrance chemical beta-caryophyllene-air oxidation and skin sensitization. *Food Chem Toxicol* 2006: 44: 538-545.
137. Santucci B, Cristaudo A, Cannistraci C, Picardo M. Contact dermatitis to fragrances. *Contact Dermatitis* 1987: 16: 93-95.
138. Fregert S, Hjorth N. Results of Standard Patch Tests with Substances Abandoned. *Contact Dermatitis Newsletter* 1969: 5: 85-86.
139. de Groot A C, Liem D H, Nater J P, van Ketel W G. Patch tests with fragrance materials and preservatives. *Contact Dermatitis* 1985: 12: 87-92.
140. Hagvall L, Skold M, Brared-Christensson J, Borje A, Karlberg A T. Lavender oil lacks natural protection against autoxidation, forming strong contact allergens on air exposure. *Contact Dermatitis* 2008: 59: 143-150.
141. Berglund V. *Master Thesis University of Gothenburg*. 2011.
142. Ruberto G, Baratta M T, Deans S G, Dorman H J. Antioxidant and antimicrobial activity of *Foeniculum vulgare* and *Crithmum maritimum* essential oils. *Planta Med* 2000: 66: 687-693.
143. Kim H J, Chen F, Wu C, Wang X, Chung H Y, Jin Z. Evaluation of antioxidant activity of Australian tea tree (*Melaleuca alternifolia*) oil and its components. *J Agric Food Chem* 2004: 52: 2849-2854.
144. Foti M C, Ingold K U. Mechanism of inhibition of lipid peroxidation by gamma-terpinene, an unusual and potentially useful hydrocarbon antioxidant. *J Agric Food Chem* 2003: 51: 2758-2765.
145. Buckley D A. Allergy to oxidized linalool in the UK. *Contact Dermatitis* 2011: 64: 240-241.
146. Smith C K, Hotchkiss S A. Enzymes and mechanisms of xenobiotic metabolism. In: editor?) w i, eds. *Allergic Contact Dermatitis Chemical and Metabolic Mechanisms*. Taylor and Francis, London and New York, 2001: 45-87.
147. Kalgutkar A S, Gardner I, Obach R S, Shaffer C L, Callegari E, Henne K R, Mutlib A E, Dalvie D K, Lee J S, Nakai Y, O'Donnell J P, Boer J, Harriman S P. A comprehensive listing of bioactivation pathways of organic functional groups. *Curr Drug Metab* 2005: 6: 161-225.
148. Nilsson A M, Bergstrom M A, Luthman K, Nilsson J L, Karlberg A T. A conjugated diene identified as a prohaptens: contact allergenic activity and chemical reactivity of proposed epoxide metabolites. *Chem Res Toxicol* 2005: 18: 308-316.
149. Bergström M A, Luthman K, Nilsson J L, Karlberg A T. Conjugated dienes as prohaptens in contact allergy: in vivo and in vitro studies of structure-activity relationships, sensitizing capacity, and metabolic activation. *Chem Res Toxicol* 2006: 19: 760-769.
150. Bergström M A, Ott H, Carlsson A, Neis M, Zwadlo-Klarwasser G, Jonsson C A, Merk H F, Karlberg A T, Baron J M. A skin-like cytochrome P450 cocktail activates prohaptens to contact allergenic metabolites. *J Invest Dermatol* 2007: 127: 1145-1153.

151. Basketter D A. Skin Sensitization to Cinnamic Alcohol: The Role of Skin Metabolism. *Acta Derm Venereol* 1992; 72: 264-265.
152. Smith C K, Moore C A, Elahi E N, Smart A T, Hotchkiss S A. Human skin absorption and metabolism of the contact allergens, cinnamic aldehyde, and cinnamic alcohol. *Toxicol Appl Pharmacol* 2000; 168: 189-199.
153. Cheung C, Hotchkiss S A, Pease C K. Cinnamic compound metabolism in human skin and the role metabolism may play in determining relative sensitisation potency. *J Dermatol Sci* 2003; 31: 9-19.
154. Elahi E N, Wright Z, Hinselwood D, Hotchkiss S A, Basketter D A, Pease C K. Protein binding and metabolism influence the relative skin sensitization potential of cinnamic compounds. *Chem Res Toxicol* 2004; 17: 301-310.
155. Ott H, Wiederholt T, Bergstrom M A, Heise R, Skazik C, Czaja K, Marquardt Y, Karlberg A T, Merk H F, Baron J M. High-resolution transcriptional profiling of chemical-stimulated dendritic cells identifies immunogenic contact allergens, but not prohaptens. *Skin Pharmacol Physiol* 2010; 23: 213-224.
156. Bertrand F, Basketter D A, Roberts D W, Lepoittevin J P. Skin sensitization to eugenol and isoeugenol in mice: possible metabolic pathways involving ortho-quinone and quinone methide intermediates. *Chemical research in toxicology* 1997; 10: 335-343.
157. Rastogi S C, Johansen J D. Significant exposures to isoeugenol derivatives in perfumes. *Contact Dermatitis* 2008; 58: 278-281.
158. Flyvholm M A, Andersen K E, Baranski B, Sarlo K. *Criteria for classification of skin- and airway-sensitizing substances in the work and general environments*. Regional Office for Europe: WHO, 1996.
159. Basketter D A, Flyvholm M A, Menne T. Classification criteria for skin-sensitizing chemicals: a commentary. *Contact Dermatitis* 1999; 40: 175-182.
160. Schnuch A, Lessmann H, Schulz K H, Becker D, Diepgen T L, Drexler H, Erdmann S, Fartasch M, Greim H, Kricke-Helling P, Merget R, Merk H, Nowak D, Rothe A, Stropp G, Uter W, Wallenstein G. When should a substance be designated as sensitizing for the skin ('Sh') or for the airways ('Sa')? *Hum Exp Toxicol* 2002; 21: 439-444.
161. Basketter D A, Andersen K E, Liden C, Van Loveren H, Boman A, Kimber I, Alanko K, Berggren E. Evaluation of the skin sensitizing potency of chemicals by using the existing methods and considerations of relevance for elicitation. *Contact Dermatitis* 2005; 52: 39-43.
162. Anonymous. *Guidance to Regulation (EC) No 1272/2008 on classification, labelling and packaging (CLP) of substances and mixtures*. ECHA Reference: ECHA-11-G-06-EN. Date: 04/2011 http://echa.europa.eu/documents/10162/17217/clp_en.pdf 2011.
163. Lalko J, Api A M, Politano V T, Letizia C. Quantitative risk assessment for dermal sensitization to fragrance ingredients: The utility of LLNA data in the weight of evidence approach to identifying thresholds. *46th Congress of the European Societies of Toxicology, September 13-16 2009, Dresden, Germany* 2009:
164. RIFM. Local lymph node assay (LLNA) protocol summaries: Data presented at the 46th Congress of the European Societies of Toxicology. *Research Institute for Fragrance Materials, Inc* 2009:
165. Gerberick G F, Kern P S, Schlatter H, Dearman R J, Kimber I, Patlewicz G Y, Basketter D A. Compilation of historical local lymph node data for evaluation of skin sensitization alternative methods *Dermatitis* 2005; 16: 157-202.
166. Kern P S, Gerberick G F, Ryan C A, Kimber I, Aptula A, Basketter D A. Local lymph node data for the evaluation of skin sensitization alternatives: a second compilation. *Dermatitis* 2010; 21: 8-32.

167. Arnau E G, Andersen K E, Bruze M, Frosch P J, Johansen J D, Menne T, Rastogi S C, White I R, Lepoittevin J P. Identification of Lilial as a fragrance sensitizer in a perfume by bioassay-guided chemical fractionation and structure-activity relationships. *Contact Dermatitis* 2000; 43: 351-358.
168. Lepoittevin J P. Metabolism versus chemical transformation or pro- versus prehapten? *Contact Dermatitis* 2006; 54: 73-74.
169. SCCP. Opinion concerning the predictive testing of potentially cutaneous sensitizing cosmetic ingredients or mixtures of ingredients adopted by the SCCNFP during the 11th plenary session of 17 February 2000. 2000:
170. Heisterberg M V, Menne T, Johansen J D. Contact allergy to the 26 specific fragrance ingredients to be declared on cosmetic products in accordance with the EU cosmetics directive. *Contact Dermatitis* 2011; 65: 266-275.
171. Larsen W, Nakayama H, Fischer T, Elsner P, Frosch P, Burrows D, Jordan W, Shaw S, Wilkinson J, Marks J, Jr., Sugawara M, Nethercott M, Nethercott J. Fragrance contact dermatitis: a worldwide multicenter investigation (Part II). *Contact Dermatitis* 2001; 44: 344-346.
172. Andersen A. Final report on the safety assessment of sodium p-chloro-m-cresol, p-chloro-m-cresol, chlorothymol, mixed cresols, m-cresol, o-cresol, p-cresol, isopropyl cresols, thymol, o-cymen-5-ol, and carvacrol. *Int J Toxicol* 2006; 25 Suppl 1: 29-127.
173. Larsen W, Nakayama H, Fischer T, Elsner P, Frosch P, Burrows D, Jordan W, Shaw S, Wilkinson J, Marks J, Sugawara M, Nethercott M, Nethercott J. Fragrance contact dermatitis - a worldwide multicenter investigation (Part III). *Contact Dermatitis* 2002; 46: 141-144.
174. Lapczynski A, Lalko J, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on trans,trans-delta-damascone. *Food Chem Toxicol* 2007; 45 Suppl 1: S211-215.
175. Letizia C S, Cocchiara J, Wellington G A, Funk C, Api A M. Food and chemical toxicology. *Food Chem Toxicol* 2000; 38 Suppl 3: S1-236.
176. Mitchell D M, Beck M H. Contact allergy to benzyl alcohol in a cutting oil reodorant. *Contact Dermatitis* 1988; 18: 301-302.
177. Mitchell J C. Contact hypersensitivity to some perfume materials. *Contact Dermatitis* 1975; 1: 196-199.
178. Malten K E, van Ketel W G, Nater J P, Liem D H. Reactions in selected patients to 22 fragrance materials. *Contact Dermatitis* 1984; 11: 1-10.
179. Mitchell J C, Calnan C D, Clendenning W E, Cronin E, Hjorth N, Magnusson B, Maibach H I, Meneghini C L, Wilkinson D S. Patch testing with some components of balsam of Peru. *Contact Dermatitis* 1976; 2: 57-58.
180. Bernaola G, Escayol P, Fernandez E, de Corres L F. Contact dermatitis from methylionone fragrance. *Contact Dermatitis* 1989; 20: 71-72.
181. English J S, Rycroft R J. Allergic contact dermatitis from methyl heptene and methyl octene carbonates. *Contact Dermatitis* 1988; 18: 174-175.
182. Bhatia S P, McGinty D, Letizia C S, Api A M. Fragrance material review on myrtenol. *Food Chem Toxicol* 2008; 46 Suppl 11: S237-240.
183. Lapczynski A, Bhatia S P, Letizia C S, Api A M. Fragrance material review on nerolidol (isomer unspecified). *Food Chem Toxicol* 2008; 46 Suppl 11: S247-250.
184. Sanchez-Politta S, Campanelli A, Pashe-Koo F, Saurat J H, Piletta P. Allergic contact dermatitis to phenylacetaldehyde: a forgotten allergen? *Contact Dermatitis* 2007; 56: 171-172.

185. McGinty D, Letizia C S, Api A M. Fragrance material review on phytol. *Food Chem Toxicol* 2010: 48 Suppl 3: S59-63.
186. Lapczynski A, Bhatia S P, Letizia C S, Api A M. Fragrance material review on rhodinol. *Food Chem Toxicol* 2008: 46 Suppl 11: S259-262.
187. Lapczynski A, Lalko J, McGinty D, Bhatia S P, Letizia C S, Api A M. Fragrance material review on alpha-isodamascone. *Food Chem Toxicol* 2007: 45 Suppl 1: S267-271.
188. Trattner A, David M. Patch testing with fine fragrances: comparison with fragrance mix, balsam of Peru and a fragrance series. *Contact Dermatitis* 2003: 49: 287-289.
189. Surburg H, Panten J. *Common fragrance and flavor materials: preparation, properties and uses*. Weinheim: Wiley-VCH, 2006.
190. Schmidt E. Production of Essential Oils. In: Husnu Can Baser K, Buchbauer G, eds. *Handbook of Essential Oils - Science, Technology, and Applications*. Boca Raton: CRC Press, 2010: 88-95.
191. Trattner A, David M, Lazarov A. Occupational contact dermatitis due to essential oils. *Contact Dermatitis* 2008: 58: 282-284.
192. Jung P, Sesztak-Greinecker G, Wantke F, Gotz M, Jarisch R, Hemmer W. Mechanical irritation triggering allergic contact dermatitis from essential oils in a masseur. *Contact Dermatitis* 2006: 54: 297-299.
193. Bilsland D, Strong A. Allergic contact dermatitis from the essential oil of French marigold (*Tagetes patula*) in an aromatherapist. *Contact Dermatitis* 1990: 23: 55-56.
194. Cockayne S E, Gawkrödger D J. Occupational contact dermatitis in an aromatherapist. *Contact Dermatitis* 1997: 37: 306-307.
195. Boonchai W, Iamtharachai P, Sunthonpalin P. Occupational allergic contact dermatitis from essential oils in aromatherapists. *Contact Dermatitis* 2007: 56: 181-182.
196. Keane F M, Smith H R, White I R, Rycroft R J. Occupational allergic contact dermatitis in two aromatherapists. *Contact Dermatitis* 2000: 43: 49-51.
197. Selvaag E, Holm J O, Thune P. Allergic contact dermatitis in an aroma therapist with multiple sensitizations to essential oils. *Contact Dermatitis* 1995: 33: 354-355.
198. Romaguera C, Vilaplana J. Occupational contact dermatitis from ylang-ylang oil. *Contact Dermatitis* 2000: 43: 251.
199. Rudzki E, Grzywa Z. Allergy to perfume mixture. *Contact Dermatitis* 1986: 15: 115-116.
200. Rudzki E, Grzywa Z, Bruo W S. Sensitivity to 35 essential oils. *Contact Dermatitis* 1976: 2: 196-200.
201. Vilaplana J, Romaguera C. Contact dermatitis from the essential oil of tangerine in fragrance. *Contact Dermatitis* 2002: 46: 108.
202. Rudzki E, Grzywa Z. Sensitizing and irritating properties of star anise oil. *Contact Dermatitis* 1976: 2: 305-308.
203. Sugiura M, Hayakawa R, Kato Y, Sugiura K, Hashimoto R. Results of patch testing with lavender oil in Japan. *Contact Dermatitis* 2000: 43: 157-160.
204. Lalko J, Api A M. Investigation of the dermal sensitization potential of various essential oils in the local lymph node assay. *Food Chem Toxicol* 2006: 44: 739-746.
205. SCCP. *Memorandum Classification and categorization of skin sensitizers and grading of test reactions (SCCP/0919/05)*. Scientific Committee for on Consumer Protection, adopted 20 September 2005. 2005.

206. SCCP. *Memorandum on Hair Dye Substances and their Skin Sensitising Properties. Scientific Committee on Consumer Protection, adopted 19 December 2006.* 2006.
207. Christensson J B, Johansson S, Hagvall L, Jonsson C, Borje A, Karlberg A T. Limonene hydroperoxide analogues differ in allergenic activity. *Contact Dermatitis* 2008; 59: 344-352.
208. Landsteiner K, Jacobs J. Studies on the sensitization of animals with simple chemical compounds. *J Exp Med* 1936; 64: 625-629.
209. Ridriguez E, Towers G H N, Mitchell J C. Biological aspects of sesquiterpene lactones. *Phytochemistry* 1976; 15: 1573-1580.
210. Roberts D W, Goodwin B F J, Williams D L, Jones K, Johnson A W, Alderson J C E. Correlation between skin sensitization potential and chemical reactivity for nitrobenzyl compounds. *Food Chem Toxicol* 1984; 21: 811-813.
211. Dupuis G, Benezra C. *Allergic contact dermatitis to simple chemicals: a molecular approach* New York: Marcel Dekker, 1982.
212. Smith C K, Hotchkiss S A. *Allergic Contact Dermatitis, Chemical and Metabolic Mechanisms.* London: Taylor and Francis, 2001.
213. Roberts D W, Lepoittevin J P. Hapten-Protein Interactions. In: Lepoittevin J P, Basketter D, Goossens A, Karlberg A T, eds. *Allergic Contact Dermatitis The Molecular Basis.* Heidelberg: Springer, 1998:
214. Sykes P. *A guidebook to mechanism in organic chemistry* Edinburgh: Pearson, 1961.
215. Aptula A O, Roberts D W. Mechanistic applicability domains for nonanimal-based prediction of toxicological end points: general principles and application to reactive toxicity. *Chem Res Toxicol* 2006; 19: 1097-1105.
216. Gerberick G F, Vassallo J D, Bailey R E, Chaney J G, Morrall S W, Lepoittevin J P. Development of a peptide reactivity assay for screening contact allergens. *Toxicological sciences : an official journal of the Society of Toxicology* 2004; 81: 332-343.
217. Natsch A, Gfeller H, Rothaupt M, Ellis G. Utility and limitations of a peptide reactivity assay to predict fragrance allergens in vitro. *Toxicology in vitro : an international journal published in association with BIBRA* 2007; 21: 1220-1226.
218. Gerberick G F, Troutman J A, Foertsch L M, Vassallo J D, Quijano M, Dobson R L, Goebel C, Lepoittevin J P. Investigation of peptide reactivity of pro-hapten skin sensitizers using a peroxidase-peroxide oxidation system. *Toxicological sciences : an official journal of the Society of Toxicology* 2009; 112: 164-174.
219. Troutman J A, Foertsch L M, Kern P S, Dai H J, Quijano M, Dobson R L M, Lalko J F, Lepoittevin J-P, Gerberick G F. The incorporation of lysine into the peroxidase peptide reactivity assay for skin sensitization assessments. *Toxicol Sci* 2011; 122: 422-436.
220. Roberts D W, Aptula A O, Patlewicz G. Mechanistic applicability domains for non-animal based prediction of toxicological endpoints. QSAR analysis of the schiff base applicability domain for skin sensitization. *Chem Res Toxicol* 2006; 19: 1228-1233.
221. Roberts D W, Aptula A O, Patlewicz G Y. Chemistry-Based Risk Assessment for Skin Sensitization: Quantitative Mechanistic Modeling for the S(N)Ar Domain. *Chem Res Toxicol* 2011:
222. Johansen J D. Contact allergy to fragrances: clinical and experimental investigations of the fragrance mix and its ingredients. *Contact Dermatitis* 2002; 46 (suppl. 3): 4-31.
223. Fenn R S. Aroma chemical usage trends in modern perfumery. *Perfumer Flavorist* 1989; 14: 1-10.

224. Johansen J D, Frosch P J, Svedman C, Andersen K E, Bruze M, Pirker C, Menne T. Hydroxyisohexyl 3-cyclohexene carboxaldehyde- known as Lyral: quantitative aspects and risk assessment of an important fragrance allergen. *Contact Dermatitis* 2003; 48: 310-316.
225. Rastogi S C, Johansen J D, Menne T. Natural ingredients based cosmetics. Content of selected fragrance sensitizers. *Contact Dermatitis* 1996; 34: 423-426.
226. Müller P M, Lamparsky D. *Perfumes: Art Science and Technology*. London: Elsevier Applied Science, 1991.
227. Poulsen P B, Schmidt A. *A survey and health assessment of cosmetic products for children. Survey of Chemical Substances in Consumer Products, No. 88*. Copenhagen: Danish Environmental Protection Agency, 2007.
228. Rastogi S C, Jensen G H, Johansen J D. *Survey and risk assessment of chemical substances in deodorants. Survey of Chemical Substances in Consumer Products, No. 86*. Copenhagen: Danish Environmental Protection Agency, 2007.
229. Buckley D A. Fragrance ingredient labelling in products on sale in the U.K. *Br J Dermatol* 2007; 157: 295-300.
230. Rastogi S C, Lepoittevin J P, Johansen J D, Frosch P J, Menne T, Bruze M, Dreier B, Andersen K E, White I R. Fragrances and other materials in deodorants: search for potentially sensitizing molecules using combined GC-MS and structure activity relationship (SAR) analysis. *Contact Dermatitis* 1998; 39: 293-303.
231. Rastogi S C, Johansen J D, Frosch P, Menne T, Bruze M, Lepoittevin J P, Dreier B, Andersen K E, White I R. Deodorants on the European market: quantitative chemical analysis of 21 fragrances. *Contact Dermatitis* 1998; 38: 29-35.
232. Rastogi S C, Menne T, Johansen J D. The composition of fine fragrances is changing. *Contact Dermatitis* 2003; 48: 130-132.
233. Rastogi S C, Johansen J D, Bossi R. Selected important fragrance sensitizers in perfumes--current exposures. *Contact Dermatitis* 2007; 56: 201-204.
234. Rastogi S C, Bossi R, Johansen J D, Menne T, Bernard G, Gimenez-Arnau E, Lepoittevin J P. Content of oak moss allergens atranol and chloroatranol in perfumes and similar products. *Contact Dermatitis* 2004; 50: 367-370.
235. Rastogi S C, Johansen J D, Menne T, Frosch P, Bruze M, Andersen K E, Lepoittevin J P, Wakelin S, White I R. Contents of fragrance allergens in children's cosmetics and cosmetic-toys. *Contact Dermatitis* 1999; 41: 84-88.
236. Rastogi S C. *Contents of selected fragrance materials in cleaning products and other consumer products. Survey of chemical compounds in consumer products, No. 8*. Copenhagen: Danish Environmental Protection Agency, 2002.
237. Bernard G, Gimenez-Arnau E, Rastogi S C, Heydorn S, Johansen J D, Menne T, Goossens A, Andersen K, Lepoittevin J P. Contact allergy to oak moss: search for sensitizing molecules using combined bioassay-guided chemical fractionation, GC-MS, and structure-activity relationship analysis. *Arch Dermatol Res* 2003; 295: 229-235.
238. Johansen J D, Andersen K E, Svedman C, Bruze M, Bernard G, Gimenez-Arnau E, Rastogi S C, Lepoittevin J P, Menne T. Chloroatranol, an extremely potent allergen hidden in perfumes: a dose-response elicitation study. *Contact Dermatitis* 2003; 49: 180-184.
239. SCCP. *Opinion on Atranol and Chloroatranol present in natural extracts (e.g. oak moss and tree moss extract)*. Scientific Committee on Consumer Products, adopted 7 December 2004. 2004.

240. White I R, Johansen J D, Arnau E G, Lepoittevin J P, Rastogi S, Bruze M, Andersen K E, Frosch P J, Goossens A, Menne T. Isoeugenol is an important contact allergen: can it be safely replaced with isoeugenyl acetate? *Contact Dermatitis* 1999; 41: 272-275.
241. SCCP. *Opinion on Hydroxyisohexyl 3-cyclohexene carboxaldehyde (sensitisation only)*. Scientific Committee on Consumer Products. Adopted 7 December 2004. 2004.
242. Nardelli A, D'Hooghe E, Drieghe J, Dooms M, Goossens A. Allergic contact dermatitis from fragrance components in specific topical pharmaceutical products in Belgium. *Contact Dermatitis* 2009; 60: 303-313.
243. Fisher A A. Cosmetic dermatitis in childhood. *Cutis* 1995; 55: 15-16.
244. Corea N V, Basketter D A, Clapp C, Van Asten A, Marty J P, Pons-Guiraud A, Laverdet C. Fragrance allergy: assessing the risk from washed fabrics. *Contact Dermatitis* 2006; 55: 48-53.
245. Hartmann K, Hunzelmann N. Allergic contact dermatitis from cinnamon as an odour-neutralizing agent in shoe insoles. *Contact Dermatitis* 2004; 50: 253-254.
246. Murphy L A, White I R. Contact dermatitis from geraniol in washing-up liquid. *Contact Dermatitis* 2003; 49: 52.
247. Foti C, Zambonin C G, Conserva A, Casulli C, D'Accolti L, Angelini G. Occupational contact dermatitis to a limonene-based solvent in a histopathology technician. *Contact Dermatitis* 2007; 56: 109-112.
248. Topham E J, Wakelin S H. D-Limonene contact dermatitis from hand cleansers. *Contact Dermatitis* 2003; 49: 108-109.
249. Wakelin S H, McFadden J P, Leonard J N, Rycroft R J. Allergic contact dermatitis from d-limonene in a laboratory technician. *Contact Dermatitis* 1998; 38: 164-165.
250. Rastogi S C, Heydorn S, Johansen J D, Basketter D A. Fragrance chemicals in domestic and occupational products. *Contact Dermatitis* 2001; 45: 221-225.
251. Yazar K, Johnsson S, Lind M L, Boman A, Liden C. Preservatives and fragrances in selected consumer-available cosmetics and detergents. *Contact Dermatitis* 2011; 64: 265-272.
252. Api A M, Bredbenner A, McGowen M, Niemiera D, Parker L, Renskers K, Selim S, Sgaramella R, Signorelli R, Tedrow S, Troy W. Skin contact transfer of three fragrance residues from candles to human hands. *Regul Toxicol Pharmacol* 2007; 48: 279-283.
253. Nadiminti H, Ehrlich A, Udey M C. Oral erosions as a manifestation of allergic contact sensitivity to cinnamon mints. *Contact Dermatitis* 2005; 52: 46-47.
254. Hoskyn J, Guin J D. Contact allergy to cinnamal in a patient with oral lichen planus. *Contact Dermatitis* 2005; 52: 160-161.
255. Silvestre J F, Albares M P, Blanes M, Pascual J C, Pastor N. Allergic contact gingivitis due to eugenol present in a restorative dental material. *Contact Dermatitis* 2005; 52: 341.
256. Guarneri F, Barbuzza O, Vaccaro M, Galtieri G. Allergic contact dermatitis and asthma caused by limonene in a labourer handling citrus fruits. *Contact Dermatitis* 2008; 58: 315-316.
257. Wallenhammar L M, Ortengren U, Adreasson H, Barregard L, Björkner B, Karlsson S, et al. Contact allergy and hand eczema in Swedish dentists. *Contact Dermatitis* 2000; 43: 192-199.
258. Geier J, Lessmann H, Schnuch A, Uter W. Contact sensitizations in metalworkers with occupational dermatitis exposed to water-based metalworking fluids: results of the research project "FaSt". *Int Arch Occup Environ Health* 2004; 77: 543-551.

259. Decapite T J, Anderson B E. Allergic contact dermatitis from cinnamic aldehyde found in an industrial odour-masking agent. *Contact Dermatitis* 2004; 51: 312-313.
260. Schubert H J. Skin diseases in workers at a perfume factory. *Contact Dermatitis* 2006; 55: 81-83.
261. Heydorn S, Andersen K E, Johansen J D, Menne T. A stronger patch test elicitation reaction to the allergen hydroxycitronellal plus the irritant sodium lauryl sulfate. *Contact Dermatitis* 2003; 49: 133-139.
262. Hannuksela M. Sensitivity of Various Skin Sites in the Repeated Open Application Test. *Am J Contact Dermatitis* 1991; 2: 102-104.
263. Fischer L A, Menné T, Avnstorp C, Kasting G B, Johansen J D. Hydroxyisohexyl 3-cyclohexene carboxaldehyde allergy: relationship between patch test and repeated open application test thresholds. *Br J Dermatol* 2009; 161: 560-567.
264. Andersen K E, Johansen J D, Bruze M, Frosch P J, Goossens A, Lepoittevin J P, Rastogi S, White I, Menne T. The time-dose-response relationship for elicitation of contact dermatitis in isoeugenol allergic individuals. *Toxicol Appl Pharmacol* 2001; 170: 166-171.
265. DeGroot A C, Frosch P J. Adverse reactions to fragrances. A clinical review. *Contact Dermatitis* 1997; 36: 57-86.
266. Schnuch A, Geier J, Uter W, Frosch P J. Another look on allergies to fragrances: frequencies of sensitisation to the fragrance mix and its constituents. Results from the IVDK. *Exog Dermatol* 2002; 1: 231-237.
267. Schnuch A, Uter W, Geier J, Lessmann H, Frosch P J. Contact allergy to farnesol in 2021 consecutively patch tested patients. Results of the IVDK. *Contact Dermatitis* 2004; 50: 117-121.
268. Uter W, Schnuch A, Gefeller O. Guidelines for the descriptive presentation and statistical analysis of contact allergy data. *Contact Dermatitis* 2004; 51: 47-56.
269. Uter W, Gefeller O, Geier J, Lessmann H, Pfahlberg A, Schnuch A. *Untersuchungen zur Abhängigkeit der Sensibilisierung gegen wichtige Allergene von arbeitsbedingten sowie individuellen Faktoren. Schriftenreihe der Bundesanstalt für Arbeitsschutz und Arbeitsmedizin, Forschung, Fb 949.* Bremerhaven: 2002.
270. van Loveren H, Cockshott A, Gebel T, Gundert-Remy U, de Jong W H, Matheson J, McGarry H, Musset L, Selgrade M K, Vickers C. Skin sensitization in chemical risk assessment: report of a WHO/IPCS international workshop focusing on dose-response assessment. *Regulatory toxicology and pharmacology : RTP* 2008; 50: 155-199.
271. Kimber I, Dearman R J, Basketter D A, Ryan C A, Gerberick G F, McNamee P M, Lalko J, Api A M. Dose metrics in the acquisition of skin sensitization: thresholds and importance of dose per unit area. *Regulatory toxicology and pharmacology : RTP* 2008; 52: 39-45.
272. Paramasivan P, Lai C, Pickard C, Ardern-Jones M, Healy E, Friedmann PS. Repeated low-dose skin exposure is an effective sensitizing stimulus, a factor to be taken into account in predicting sensitization risk. *Br J Dermatol.* 2010; 162: 594-7
273. Suskind R R. The hydroxycitronellal story: What can we learn from it? In: Frosch P J, Johansen J D, White I R, eds. *Fragrances Beneficial and adverse effects.* Berlin, Heidelberg, New York: Springer, 1988: 159-165.
274. Api A M, Basketter D, Cadby P A, Cano M-F, Ellis G, Gerberick F, Griem P, McNamee P M, Ryan C A, Safford B. *Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients Technical Dossier. June 22, 2006 QRA Expert Group.* http://www.ifraorg.org/en-us/search/tags_21261 (last accessed 2011-11-27). 2006.

275. SCCP. Opinion on Dermal Sensitisation Quantitative Risk Assessment (Citral, Farnesol and Phenylacetaldehyde). Scientific Committee for on Consumer Protection, adopted 24 June 2008. 2008:
276. Fischer L A, Voelund A, Andersen K E, Menne T, Johansen J D. The dose-response relationship between the patch test and ROAT and the potential use for regulatory purposes. *Contact Dermatitis* 2009; 61: 201-208.
277. Roberts D W. QSAR for upper-respiratory tract irritation. *Chem Biol Interact* 1986; 57: 325-345.
278. Roberts D W, Natsch A. High throughput kinetic profiling approach for covalent binding to peptides: application to skin sensitization potency of Michael acceptor electrophiles. *Chem Res Toxicol* 2009; 22: 592-603.
279. Bruze M, Johansen J D, Andersen K E, Frosch P, Goossens A, Lepoittevin J P, Rastogi S C, White I, Menne T. Deodorants: an experimental provocation study with isoeugenol. *Contact Dermatitis* 2005; 52: 260-267.
280. Fischer L A, Menne T, Voelund A, Johansen J D. Can exposure limitations for well-known contact allergens be simplified? An analysis of dose-response patch test data. *Contact Dermatitis* 2011; 64: 337-342.
281. Franot C, Roberts D W, Basketter D A, Benezra C, Lepoittevin J P. Structure-activity relationships for contact allergenic potential of gamma,gamma-dimethyl-gamma-butyrolactone derivatives. 2. Quantitative structure-skin sensitization relationships for alpha-substituted-alpha-methyl-gamma,gamma-dimethyl-gamma-butyrolactone s. *Chem Res Toxicol* 1994; 7: 307-312.
282. SCCS. *Notes of Guidance for the Testing of Cosmetic Ingredients and their Safety Evaluation, 7th revision*. Scientific Committee for Consumer Safety, adopted 14 December 2010. 2010.
283. Basketter D, Horev L, Slodovnik D, Merimes S, Trattner A, Ingber A. Investigation of the threshold for allergic reactivity to chromium. *Contact Dermatitis* 2001; 44: 70-74.
284. Pasche F, Hunziker N. Sensitization to Kathon CG in Geneva and Switzerland. *Contact Dermatitis* 1989; 20: 115-119.
285. Fischer L A, Johansen J D, Menne T. Nickel allergy: relationship between patch test and repeated open application test thresholds. *Br J Dermatol* 2007; 157: 723-729.
286. Fischer L A, Johansen J D, Menne T. Methyl dibromoglutaronitrile allergy: relationship between patch test and repeated open application test thresholds. *Br J Dermatol* 2008; 159: 1138-1143.
287. Flyvholm M A, Hall B M, Agner T, et al. Threshold for occluded formaldehyde patch test in formaldehyde-sensitive patients. Relationship to repeated open application test with a product containing formaldehyde releaser. *Contact Dermatitis* 1997; 36: 26-33.
288. Thyssen J P, Johansen J D, Menne T, Nielsen N H, Linneberg A. Nickel allergy in Danish women before and after nickel regulation. *The New England journal of medicine* 2009; 360: 2259-2260.
289. Thyssen J P, Linneberg A, Menne T, Nielsen N H, Johansen J D. The association between hand eczema and nickel allergy has weakened among young women in the general population following the Danish nickel regulation: results from two cross-sectional studies. *Contact Dermatitis* 2009; 61: 342-348.
290. Zachariae C O, Agner T, Menné T. Chromium allergy in consecutive patients in a country where ferrous sulfate has been added since 1981. *Contact Dermatitis* 1996; 35: 83-85.

291. Braendstrup P, Johansen J D. Hydroxyisohexyl 3-cyclohexene carboxaldehyde (Lyrall) is still a frequent allergen. *Contact Dermatitis* 2008; 59: 187-188.
292. SCCNFP. *Opinion on hydroxyisohexyl 3-cyclohexene carboxaldehyde. The Scientific Committee on Cosmetic products and Non-Food Products Intended for Consumers, adopted 9 December 2003.* 2003.
293. Api A M, Vey M. A new IFRA Standard on the fragrance ingredient, hydroxyisohexyl 3-cyclohexene carboxaldehyde. *Contact Dermatitis* 2010; 62: 254-255.
294. Schnuch A, Geier J, Uter W. Is hydroxyisohexyl 3-cyclohexene carboxaldehyde sensitisation declining in central Europe? *Contact Dermatitis* 2012; 67: 47-49.
295. Heisterberg M V, Laurberg G, Veien N, Menné T, Avnstorp C, Kaaber K, Andersen K A, Sommerlund M, Danielsen A, Andersen B, Kristensen B, Kristensen O, Nielsen N H, Thormann J, Vissing S, Johansen J D. Prevalence of allergic contact dermatitis caused by hydroxyisohexyl 3-cyclohexene carboxaldehyde has not changed in Denmark. *Contact Dermatitis* 2012; 67: 49-51.
296. Nardelli A, Gimenez-Arnau E, Bernard G, Lepoittevin J P, Goossens A. Is a low content in atranol/chloroatranol safe in oak moss-sensitized individuals? *Contact Dermatitis* 2009; 60: 91-95.
297. Anonymous. *OECD Guidelines for the Testing of Chemicals / Section 4: Health Effects. Test No. 429: Skin Sensitisation (Local Lymph Node Assay).* Paris: OECD, 2002.
298. Anonymous. 76/768/EEC - Council Directive 76/768/EEC of 27 July 1976 on the approximation of the laws of the Member States relating to cosmetic products. *Official Journal L* 1976; 262, 27/09/1976: 169.
299. Vigan M. Contact dermatitis sentinel network by GERDA. *Nouv Dermatol* 1996; 15: 677-678.
300. SCCP. *Opinion on Oak moss/Tree moss (sensitisation only) Scientific Committee on Consumer Products, adopted 15 April 2008.* 2008.

Annex I - Catalogue of fragrance allergens

Contents

| | |
|--|-----|
| Single chemicals | 142 |
| Catalogue of single chemicals evaluated | 146 |
| Natural extracts / essential oils | 237 |
| Catalogue of natural extracts / essential oils evaluated | 238 |
| References | 277 |

Single chemicals

Often, results with the single constituents of the FM I or, yet more rarely, FM II, are presented in one paper. As the main ordering of this annex is by allergen, core information on these studies is presented in a tabular format and referenced by a unique acronym in the single sections, to avoid redundancy. Regarding nomenclature, terms which are often not officially an INCI Name but Perfuming Name as listed by CosIng are used. "Current Regulation" refers to the EU Cosmetics Directive only.

Table 55: Background information on studies reporting results with (all) single constituents of the FM I (**amyl cinnamal, cinnamyl alcohol, cinnamal, eugenol, geraniol, hydroxycitronellal, isoeugenol, EVERNIA PRUNASTRI**)

| Reference | Country | Study period, Patients | Comments by reviewers |
|--------------------|--|--|---|
| Larsen 2002 c (1) | 7 industrial countries worldwide | Prior to 2002 n=218 patients with known contact allergy to fragrance ingredients | Test concentrations identified as non-irritating in serial dilution testing in 20 healthy volunteers |
| Utrecht 1999 (2) | Utrecht, The Netherlands | 1994-1998 n=757 patients with suspected ACD to cosmetics | All patients tested with FM I and single constituents |
| Sheffield 1999 (3) | UK | 1994-1995 n=744, 40 of these positive to FM I and tested with single constituents | |
| IVDK 2007 (4) | Germany + one centre in Austria and Switzerland each | 01/2003 - 12/2004, n=1658 to 21325, see text, consecutive patients | |
| Hungary 2002 (5) | Hungary, multicentre study, | 1998-1999, n=3604 patients | recruitment not clear, presumably consecutive patients |
| Groningen 2009 (6) | Groningen, The Netherlands | 04/2005-06/2007 n=320 | patients selected according to history or site suspicious of contact allergy to fragrance ingredients |
| IVDK 2010 (7) | Germany, Switzerland and one centre in | 2005-2008 n=36961 tested with FM I, n=4167 with FM II and | |

| | | | |
|--|---------|--------|--|
| | Austria | all SC | |
|--|---------|--------|--|

Table 56: Results of PTing with single constituents of the FM I in patients positive to the FM I (as percent)

| N(pos) to FM I, ref. | <i>Evernia prun.</i> | Isoeu g. | Hydroxy citron. | Cinna mal | Cinnamy l alcohol | Eugen ol | Gera niol | Alpha- amyl cinnam al |
|-------------------------|----------------------|----------|-----------------|-----------|-------------------|----------|-----------|-----------------------|
| N=160 (5) | 13.1% | 14.8 % | 2.5% | 8.1% | 20.6% | 8.8% | 7.5% | 5.0% |
| N= 991 (8) | 18.4% | 11.2 % | 10.1% | 6.1% | 6.1% | 6.6% | 4.6% | 2.4% |
| N=50 (2) | 19.6% | 14.3 % | 8.9% | 8.9% | 7.1% | 5.4% | 2.7% | 0% |
| n=40 Sheffield 1999 (3) | 30% | 20% | 2.5% | 12.5 % | 10% | 5% | 0% | 0% |
| N=226 Coimbra 2000 (9) | 22.1% | 19.9 % | 6.6% | 13.3 % | 7.9% | 14.6 % | 8.4% | 4.4% |
| N=655 IVDK 2010 (7) | 29.8% | 18.0 % | 12.8% | 11.6 % | 9.6% | 6.7% | 4.7% | 2.8% |

Table 57: Background information on studies reporting results with (all) single constituents of the FM II (**citronellol, citral, coumarin, hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC), Farnesol, alpha-Hexyl-cinnamic aldehyde**)

| Reference | Country | Study period, Patients | Comments by reviewers |
|--------------------|--|--|---|
| IVDK 2007 (4) | Germany + one centre in Austria and Switzerland each | 01/2003 - 12/2004, n=1658 to 21325, see text, consecutive patients | |
| EU 2005 (10) | 6 European centres | 10/2002 - 06/2003, n=1701 | Applied in consecutive patients |
| Groningen 2009 (6) | Groningen, The Netherlands | 04/2005-06/2007 n=320 | patients selected according to history or site suspicious of contact allergy to fragrance ingredients |

| | | | |
|-----------------|--|--|--|
| IVDK 2010b (11) | Germany, Switzerland and one centre in Austria | 2005-2008 n=35633 tested with FM II, n=2217 with all SC | |
|-----------------|--|--|--|

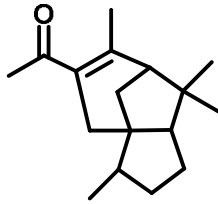
Table 58: Background information on studies reporting results with several fragrance compounds not, or only partly, corresponding to mixes (later created) or with essential oils

| Reference | Country | Study period, Patients |
|--------------------|--|---|
| deGroot 2000 (12) | The Netherlands (multicentre) | 09/1998-04/1999 n=1825 consecutive patients |
| An 2005 (13) | South Korea (multicentre) | 04/2002 – 06/2003 n=422 consecutive patients |
| Sugiura 2000 (14) | Nagoya, Japan | 1990-1998 n=1483 patients with suspected cosmetic dermatitis |
| Frosch 1995 (15) | 11 European depts. | Prior to 1995 n=1069 consecutive patients |
| Frosch 2002 a (16) | 6 European depts. | 10/1997-10/1998 n=1855 consecutive patients |
| Frosch 2002 b (17) | 6 European depts. | Prior to 2002 n=1606 consecutive patients |
| Coimbra 2000 (9) | Portugal | 07/1989-06/1999 n=226 with FM I SC n=67 also with other fragrances |
| Larsen 1977 (18) | US | 1977 n=20 "perfume-sensitive patients" |
| Larsen 2001 (19) | worldwide multicentre | ? (prior to 2001) n=178 patients with known contact allergy to fragrance ingredients |
| Belsito 2006 (20) | North American (5 US, 1 Canadian) depts. | 2003 n=1603 patients |
| NACDG 2009 (21) | US and Canada | 2005-2006 n= 4454 patients |
| Wöhrl 2001 (22) | "FAZ" clinic Vienna | 1997-2000 n=747 of 2660 consecutive patients tested with special series |
| EECDRG 1995 (15) | European, multicentre | Different fragrances, tested in 2 concentrations, in sets of about 100 patients each in different centres |
| Goossens 1997 (23) | Leuven, Belgium | 1978-1987 n=111 "Japanese perfume series" (highly selected patients) |

Opinion on fragrance allergens in cosmetic products

| Reference | Country | Study period, Patients |
|----------------------------|--|--|
| Malten 1984 (24) | Dutch multicentre | N=182 patients with suspected cosmetic dermatitis tested with 22 fragrance compounds |
| DeGroot 1985 (25) | Dutch | N=179 patients with suspected cosmetic dermatitis tested with 16 fragrance compounds |
| Rudzki 1976 (26) | Warsaw, Poland | N=200 consecutive patients |
| Rudzki 1986 (27) | Warsaw, Poland | N=86 patients of 299 (of 5315) patients with positive reaction to FM I tested with essential oils series |
| Santucci 1987 (28) | Rome, Italy | N=1500 consecutive patients; n=63 reacting positively to FM I re-tested with extended fragrance series |
| Nakayama 1974 (after (29)) | Japan | N=183 patients with cosmetic dermatitis |
| IVDK 2010c (30) | Germany, Switzerland and one centre in Austria | 15682 patients tested with at least one essential oil in different test series |
| Trattner/David (31) | Tel Aviv, Israel | N=641 consecutive patients |

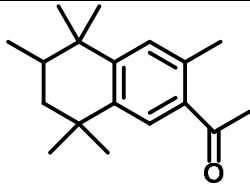
Catalogue of single chemicals evaluated

| | |
|--|---|
| ACETYLCEDRENE |  |
| CAS # 32388-55-9 | |
| EC # 251-020-3 | |
| 1-[(3R,3aR,7R,8aS)-2,3,4,7,8,8a-Hexahydro-3,6,8,8-tetramethyl-1H-3a,7-methanoazulen-5-yl]-ethanone | |
| Other names 1-(2,3,4,7,8,8a-Hexahydro-3,6,8,8-tetramethyl-1H-3a,7-methanoazulen-5-yl)-, [3R-(3α,3αβ,7β,8α)]-Ethanone; 1H-3a,7-Methanoazulene, Ethanone deriv.; Acetyl-α-cedrene; Lixetone; Vertofix | |

Current regulation: /

Clinical data:
In the Frosch 2002 a study, a total of 0.2% had positive PT reactions (16). In the Frosch 1995 dose-finding pilot study, 1 positive reaction to 1% and none to 5% "Vertofix ®" in pet., tested in 100 consecutive patients in Stockholm, were observed (15). In a case report, a 28-year-old patient with axillary dermatitis after using 2 different deodorants tested positive not only to HICC, but also to acetyl cedrene (tested 10.8% in diisopropylene glycol (20 healthy controls negative) (32). In this case report it is stated that "Acetyl cedrene (Vertofix Coeur) is a complex reaction mixture of which a principal constituent is methyl cedryl ketone".

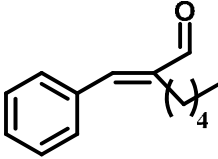
Additional information:
Acetyl cedrene (Vertofix®, IFF) is a complex mixture obtained from cedar wood oil by the acetylation of terpenes. The principal component of acetyl cedrene is methyl cedryl ketone (CAS 32388-55-9). It is a "top 100" substance (IFRA, pers. comm.2010)

| | |
|--|---|
| 6-ACETYL-1,1,2,4,4,7-HEXAMETHYLTETRALINE |  |
| CAS # 21145-77-7 | |
| EC # 216-133-4 / 244-240-6 | |
| 1-(5,6,7,8-Tetrahydro-3,5,5,6,8,8-hexamethyl-2-naphthalenyl)-ethanone | |
| AHMT (perfume), AHTN, Extralide, Fixolide, Musk tonalid, NSC 19550, Tentarome, Tetralide, Tonalid, Tonalide. | |

Current regulation: Annex III, part 1, entry 182

Clinical data:
In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and 5% "Tonalide ®" in pet., tested in 313 consecutive patients in Bordeaux and London, were observed (15).

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

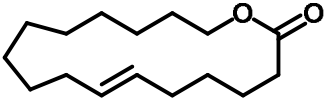
| | |
|---|---|
| AMYL CINNAMAL |  |
| CAS # 122-40-7 | |
| EC # 204-541-5 | |
| 2-(Phenylmethylene)-heptanal | |
| Cinnamaldehyde, α -amyl- (4CI); Cinnamaldehyde, α -pentyl- (6CI,7CI,8CI); 2-(Phenylmethylene)heptanal; 2-Benzylideneheptanal; Amylcinnamaldehyde; Amylcinnamic acid aldehyde; Amylcinnamic aldehyde; Flomine; Jasminal; Jasminaldehyde; Jasmine aldehyde; NSC 6649; Pentylcinnamaldehyde; α -Amyl- β -phenylacrolein; α -Amylcinnamal; α -Amylcinnamaldehyde; α -Pentylcinnamaldehyde | |

Current regulation: Annex III, part 1, entry 67

Clinical data:
In the "background information" section of the 1999 opinion (33), amyl cinnamal (synonymous: alpha amyl cinnamaldehyde) has been classified as frequently reported contact allergen because it has been identified as a cause of allergic reactions in persons with eczema from cosmetic products.

Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded $n=4$, i.e., 0.2% (95% CI: 0.1 – 0.5%) positive reactions to this compound (1% pet.) in 2062 consecutively PTed patients (4). In the Groningen 2009 study, no positive reactions to this allergen, tested at 2% pet., were observed (6). The Larsen 2001 study yielded 2.3% positive reactions in 178 patients with known contact allergy to fragrance ingredients (test concentration: 5% pet.) (19). In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded $n=2$ (0.3%) positive reactions to amyl cinnamal (22). The IVDK 2010 study, 0.26% (95% CI: 0 – 0.60%) of 1214 consecutively tested patients reacted to the compound, while 0.61% (95% CI: 0.36 – 0.86%) of 4375 of patients tested in a more aimed manner, partly as break-down testing to the FM I, had a positive PT reaction (7).

Additional information:
It is a "top 100" substance and classified as R43 (IFRA, pers. comm. 2010).

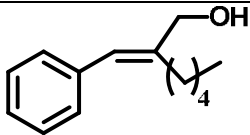
| | |
|---|---|
| AMBRETTOLIDE |  |
| CAS # 7779-50-2 | |
| EC # 231-929-1 | |
| Oxacycloheptadec-7-en-2-one | |
| 1-Oxa-7-cycloheptadecen-2-one; 16-Hydroxy-6-hexadecenoic acid lactone; 16-Hydroxy-6-hexadecenoic acid ω -lactone | |

Current regulation: /

Clinical data:
The Larsen 2001 study, using omega-6-hexadecenlactone (HDL, 5% pet.) as test concentration, diagnosed 3.4% positive reactions in 178 patients with known contact allergy to fragrance ingredients (19).

Additional information:

Ambrettolide is 1 of 2 components of Ambrette seed oil (obtained from *Hibiscus abelmoschus* L., *Malvaceae*) responsible for the musk odour. In Surburg/Panten, the compound has the chemical name (Z)-7-hexadecen-16-olide (or Hexadec-7-en-16-olide according to CosIng), CAS 123-69-3 (34).

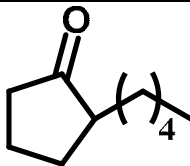
| | |
|---|---|
| AMYL CINNAMYL ALCOHOL |  |
| CAS # 101-85-9 | |
| EC # 202-982-8 | |
| 2-(Phenylmethylene)-heptan-1-ol, | |
| 2-Benzylidene- (6CI,8CI)1-heptanol; 2-Amyl-3-phenyl-2-propen-1-ol; 2-Benzylidene-1-heptanol; 2-Pentyl-3-phenyl-2-propen-1-ol; Buxinol; α-Amylcinnamic alcohol; α-Amylcinnamyl alcohol | |

Current regulation: Annex II, Part 1, entry 74

Clinical data:
In the "background information" section of the 1999 opinion, amyl cinnamyl alcohol is mentioned to cross-react with amyl cinnamal. Moreover, this compound has been identified as a cause of allergic reactions in a notable number of persons with eczema from the use of cosmetic products (33).

Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded 0.4% (95% CI: 0.1 – 0.7%) positive reactions in 1977 consecutively PTed patients (4). The IVDK 2010 study, 0.79% (95% CI: 0.54 – 1.04%; percentages standardised for age and sex) of 5650 patients PTed reacted to the compound (7). In the Groningen 2009 study, 0.6% (95% CI: 0.1 – 2.2%) had positive reactions to this allergen (6).

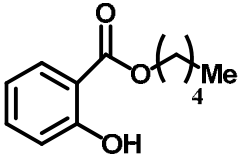
Additional information: A RIFM review is available (35) where selected clinical studies published until 1994 were considered.

| | |
|---|---|
| AMYL CYCLOPENTANONE |  |
| CAS # 4819-67-4 | |
| EC # 225-392-2 | |
| 2-Pentylcyclopentanone | |
| 2-Pentyl-1-cyclopentanone; 2-Pentylcyclopentanone; 2-Pentylcyclopenten-1-one; 2-n-Amylcyclopentanone; 2-n-Pentyl cyclopentanone; Delphone | |

Current regulation: /

Clinical data:
In the Larsen 2001 study, none of 178 patients with contact allergy to fragrance ingredients reacted positively to this ingredient, PTed at 5% pet. (19).

Additional information: /

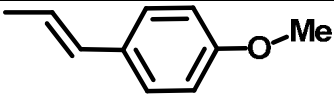
| | |
|--|---|
| AMYL SALICYLATE |  |
| CAS # 2050-08-0 | |
| EC # 218-080-2 | |
| Pentyl-2-hydroxybenzoate | |
| Amyl ester salicylic acid, (4CI); Pentyl ester salicylic acid, (6CI,8CI); 2-Hydroxybenzoic acid pentyl ester; Amyl salicylate; NSC 403668; NSC 44877; NSC 46125; Pentyl salicylate | |

Current regulation: /

Clinical data:

In the Frosch 2002 a study, a total of n=3 (0.2%) had positive PT reactions (16). In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% amyl salicylate and 1 positive reaction to 5% amyl salicylate were observed in 100 consecutive patients patch tested in Stockholm (15).

Additional information:
A RIFM review is available (36). It is a "top 100" substance (IFRA, pers. comm.2010)

| | |
|--|---|
| trans-ANETHOLE |  |
| CAS # 4180-23-8 | |
| EC # 224-052-0 / 203-205-5 | |
| 1-Methoxy-4-(1E)-1-propen-1-yl-benzene | |
| (E)-p-Propenyl-anisole (8CI); (E)-1-Methoxy-4-(1-propenyl)-benzene; 1-Methoxy-4-(1E)-1-propenyl-benzene (9CI); (E)-1-(4-Methoxyphenyl)propene; (E)-1-p-Methoxyphenylpropene; (E)-Anethol; (E)-Anethole (REACH, EINECS); E-Anethole (INCI); 1-Methoxy-4-[(1E)-1-propenyl]benzene; (E)-1-Methoxy-4-(1-propenyl)-benzene (CosIng); NSC 209529; trans-1-(4-Methoxyphenyl)-1-propene; trans-1-(p-Methoxyphenyl)-1-propene; trans-1-(p-Methoxyphenyl)propene; trans-1-p-Anisylpropene; trans-4-(1-Propenyl)anisole; trans-Anethol; trans-Anethole; trans-p-Anethole; trans-p-Methoxy-β-methylstyrene | |

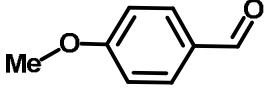
Current regulation: /

Clinical data:

A case of a 64 year old patient, who developed severe cheilitis and a loss of taste has been described (37). Both were reversible after the cessation of use of previous toothpastes. The patch test was strongly positive to anethole (isoform not given) 5% pet.; this was found an ingredient of the causative toothpaste. Two cases of occupational allergic contact dermatitis occurring in a traditional cake factory due to anise oil have been described, both testing (strongly) positive to anise oil (5% o.o.) and anethole (5% pet.) (38).

Additional information:
It is a "top 100" substance (IFRA, pers. comm.2010). trans-Anethole can be purified from star anise oil (34, 39), see 3.2., and is the main component of anise, star anise

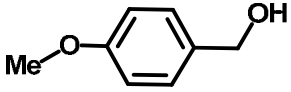
and fennel oils (38)

| | |
|--|---|
| ANISALDEHYDE |  |
| CAS # 123-11-5 | |
| EC # 204-602-6 | |
| 4-Methoxy-benzaldehyde | |
| p-Methoxybenzaldehyde; p-Anisaldehyde; 4-Anisaldehyde; Aubepine; Cratagine; NSC 5590; Obepin; p-Anisic aldehyde; Anisic aldehyde; p-Formylanisole. | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

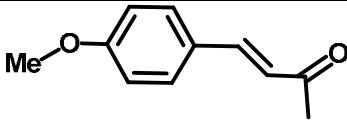
| | |
|---|---|
| ANISYL ALCOHOL |  |
| CAS # 105-13-5 | |
| EC # 203-273-6 | |
| 4-Methoxy-benzenemethanol | |
| p-Methoxy-benzyl alcohol (8CI); (4-Methoxyphenyl)methyl alcohol; 4-(Hydroxymethyl)anisole; 4-(Methoxyphenyl)methanol; 4-Methoxy- α -hydroxytoluene; 4-Methoxybenzenemethanol; 4-Methoxybenzyl alcohol; Anise alcohol; Anisic alcohol; NSC 2151; [4-(Methoxy)phenyl]methanol; p-(Methoxyphenyl)methanol; p-Anisalcohol; p-Anisyl alcohol; p-Methoxybenzyl alcohol | |

Current regulation: Annex III, part 1, n° 80

Clinical data:
In the "background information" section of the 1999 opinion, anisyl alcohol is classified as "less frequently reported allergen"; 2 studies were identified where 3 and 4 cases, respectively, with cosmetic dermatitis due to contact allergy to anisyl alcohol had been reported (33).

Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded n=1, i.e., 0.1% (95% CI: 0.00 – 0.3%) positive reactions in 2004 consecutively PTed patients, patch test concentration: 1% pet. (4). Similar results were obtained in the following period, with n=1 (and n=3 irritant and n=6 doubtful) reactions in 986 patients tested with 1% in pet. (30). In the Groningen 2009 study, no positive reactions to this allergen, tested at 5% pet., were observed in 320 patients (6). This test concentration has been regarded as relatively high by Hostynek and Maibach (40). The test concentration of Anisyl Alcohol has been further validated by Bruze et al. and 10% in pet was recommended as a non-irritant concentration for routine investigations (40a).

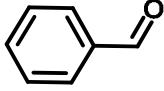
Additional information: /

| | |
|---|---|
| ANISYLIDENE ACETONE |  |
| CAS # 943-88-4 | |
| EC # 213-404-9 | |
| 4-(4-Methoxyphenyl)-3-Buten-2-one | |
| 1-(p-Methoxyphenyl)-1-buten-3-one; 4-(4-Methoxyphenyl)-3-buten-2-one; 4-(p-Methoxyphenyl)-3-buten-2-one; 4-Methoxybenzalacetone; 4-Methoxybenzylideneacetone; 4-Methoxystyryl methyl ketone; 4'-Methoxybenzylideneacetone; Anisalacetone; Methyl p-methoxystyryl ketone; NSC 31752; NSC 7946; p-Anisalacetone; p-Methoxybenzalacetone; p-Methoxybenzylideneacetone; p-Methoxystyryl methyl ketone | |

Current regulation: Annex III, part 1, n° 443

Clinical data:
In the Malten 1984 study, 1.1% of 182 patients displayed a positive PT reaction to anisylidene acetone 2% pet. (24)

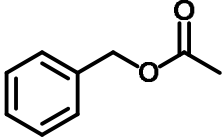
Additional information: /

| | |
|--|---|
| BENZALDEHYDE |  |
| CAS # 100-52-7 | |
| EC # 202-860-4 | |
| Benzaldehyde | |
| Artificial Almond Oil; Benzaldehyde FFC; Benzenecarbonal; Benzenecarboxaldehyde; Benzoic acid aldehyde; Benzoic aldehyde; NSC 7917; Phenylformaldehyde; Phenylmethanal | |

Current regulation: /

Clinical data:
 In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded n=3 (0.4%) positive reactions to benzaldehyde 5% pet. (22). The IVDK 2010 study, 6 weak positive reactions were observed, i.e., 0.16% (95% CI: 0.03 – 0.29%; percentages standardised for age and sex) of 2820 patients PTed reacted to the compound (7). A review is available in the Int. J. Toxicol. (41). In the case of a 19 year old pastry maker, Seite-Bellezza et al. report on immediate reactions to MP, cinnamal and benzaldehyde (tested at 5% pet.) subsiding after a few hours, in line with the patient's history (42).

Additional information: /

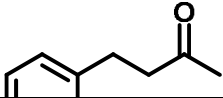
| | |
|--|---|
| BENZYL ACETATE |  |
| CAS #140-11-4 | |
| EC # 205-399-7 / 202-940-9 | |
| Benzyl acetate | |
| Benzyl ester acetic acid; Benzyl alcohol, acetate (6CI); (Acetoxymethyl)benzene; Benzyl ethanoate; NSC 4550; Phenylmethyl acetate; Methyl Phenylacetate; α-Acetoxytoluene ; Methyl alpha-Toluate | |

Current regulation: /

Clinical data:

In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and 5% benzyl acetate in pet., tested in 100 consecutive patients in Odense, DK, were observed (15). Benzyl acetate is a component of several natural mixtures, for example a major constituent of Narcissus abs., and a minor constituent of Jasmine abs. (17).

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

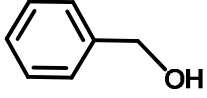
| | |
|-----------------------|---|
| BENZYL ACETONE |  |
| CAS # 2550-26-7 | |

| | |
|--|--|
| EC # 219-847-4 | |
| 4-Phenyl-2-butanone | |
| 4-Phenylbutan-2-one (REACH, EINECS); Benzylacetone; Methyl 2-phenylethyl ketone; Methyl phenethyl ketone; NSC 44829; NSC 813M; Phenethyl methyl ketone; 1-Phenyl-3-butanone; 2-Phenylethyl methyl ketone | |

Current regulation: /

Clinical data: /

Additional information:
It is a "top 100" substance (IFRA, pers. comm.2010). A RIFM review is available (43).

| | |
|---|---|
| BENZYL ALCOHOL |  |
| CAS # 100-51-6 | |
| EC # 202-859-9 | |
| Phenylmethanol | |
| Benzyl alcohol; (Hydroxymethyl)benzene; Benzenecarbinol; Benzylic alcohol; NSC 8044; Phenylcarbinol; Benzenemethanol; Phenylmethyl alcohol; Sunmorl BK 20; TB 13G; α-Hydroxytoluene; α-Toluenol | |
| Current regulation: Annex III, part 1, n° 45; Annex VI, part1, n ° 34 | |

Clinical data:
In the "background information" section of the 1999 opinion, benzyl alcohol is classified as allergen frequently causing allergic reactions. It has been found to cause allergic reactions in 1.2 to 15% of patients with eczema from cosmetic products (33). A CIR expert panel review is available in the Int. J. Toxicol. (44).

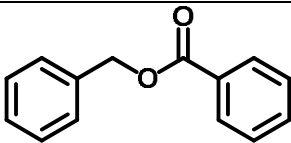
Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded 0.3% (95% CI: 0.1 – 0.7%) positive reactions in 2166 consecutively PTed patients (4). In the Groningen 2009 study, n=1, i.e. 0.3% (95% CI: 0.01 – 1.7%) had positive reactions to this allergen (6).

Both in terms of case reports (45-47) and clinical epidemiology data (0.22 % [95% CI: 0.16 – 0.28%] positive tested with benzyl alcohol in the context of a "topical drugs" series, n=26448 (7)) the relevance of this alternative exposure is highlighted. In a study from Alicante, Spain, 86 selected patients were tested with benzyl alcohol, yielding 2 positive reactions (48).

After application of saline soaks preserved with benzyl alcohol onto his stasis dermatitis, a 53 year old patient developed a rash, which was, according to test results obtained by J. D. Guin and J. Goodman, at least partly due to an immediate hypersensitivity to benzyl alcohol, as verified by an intense urticarial reaction at the test site lasting several days (49). According to 2 cases reported by A. A. Fisher, PT-proven, relevant delayed type hypersensitivity is not associated with immediate reactions in scratch or intradermal tests (50). D. W. Shaw describes a patient with allergic contact dermatitis caused by benzyl alcohol in a hearing aid impression material and in topical medications (51). Another contribution points to covert exposures to benzyl alcohol even in products labelled "fragrance free" (52) probably because benzyl alcohol is used as preservative, or an essential oil containing benzyl alcohol is used as cosmetic ingredient.

Additional information:

Benzyl alcohol is a component of several natural mixtures, including Myroxylon pereirae resin, which have been used for extraction, but is nowadays synthesised (53). It is permitted in certain foodstuffs (liquors: < 100 mg/l, sweets and cakes: < 250 mg/kg) under the coding "E 1519" (http://www.zusatzstoffe-online.de/zusatzstoffe/317.e1519_benzylalkohol.html, last accessed 2009-11-27). In addition to being a fragrance compound (which may be used, even in relatively high concentration, to scent topical medications (54)), benzyl alcohol is used as antioxidant in topical therapeutics or cosmetics. The German "Rote Liste" (<http://www.rote-liste.de>, last accessed 2009-11-11), for instance, lists 205 specialties containing benzyl alcohol. Benzyl alcohol may be used up to 1.0% as a preservative in cosmetic products according to the Cosmetic Directive 76/768/EEC

| | |
|---|---|
| BENZYL BENZOATE |  |
| CAS # 120-51-4 | |
| EC # 204-402-9 | |
| Benzyl benzoate | |
| Benzyl ester benzoic acid; Ascabin; Ascabiol; Benylate; Benzyl benzenecarboxylate; Benzyl benzoate; Benzyl phenylformate; Benzylets; Colebenz; NSC 8081; Nicca Sunsolt LM 7EX; Novoscabin; Pelemol B66; Peruscabin; Phenylmethyl benzoate; Scabagen; Scabanca; Scabcare BB; Scabide; Scabiozon; Scobenol; Vanzoate; Venzonate | |

Current regulation: Annex III, part 1, n° 85

Clinical data:

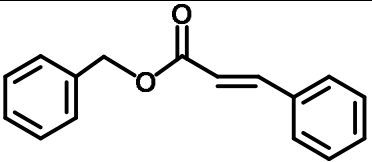
In the "background information" section of the 1999 opinion, benzyl benzoate is classified as "less frequently reported allergen"; in several studies, only single cases had been reported in each, except for patients sensitive to MP (33).

Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded n=1, i.e., 0.1% (95% CI: 0.00 – 0.3%) positive reactions in 2003 consecutively PTed patients, test concentration 1% pet. (4). In the subsequent period (2005-2008), n=1062 patients were tested in the IVDK 2010 study, with no positive reactions (7). In the Groningen 2009 study, no positive reactions to this allergen, tested at 5% pet., were observed in 320 patients (6). Thus, the pooled proportion of positive patch test reactions is 1 / 3385 (0.03%, exact upper 1-sided 95% CI: 0.14%)

Additional information:

Benzyl benzoate naturally occurs in MP resin and ylang-ylang oil. Nowadays it is synthesised and used for a variety of purposes (53). These include use as a scabicide (one brand specialty on the German market, using a concentration of 10% for children and 25% for adults), possibly with some differences among European countries. In France, a combination of benzyl benzoate 10% and sulfiram 2% is reported to be used most often (55). Hausen et al. review the older literature and mention a study identifying 1 sensitised patient in 73 patients treated for scabies (details not given) (53). According to the mandatory factsheet (see PDF "benzylbenzoate_infosheet_DE.pdf") dermatitis after anti-scabies treatment is "rare", in a range between 1:1000 and 1:10000.

It is a "top 100" substance (IFRA, pers. comm.2010).

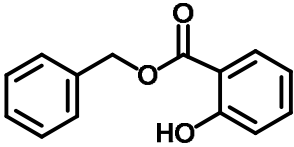
| | |
|---|---|
| BENZYL CINNAMATE |  |
| CAS # 103-41-3 | |
| EC # 203-109-3 | |
| Benzyl 3-phenylprop-2-enoate | |
| Benzyl ester cinnamic acid; 3-phenyl-phenylmethyl ester 2-propenoic acid; 3-Phenyl-2-propenoic acid benzyl ester; Benzyl 3-phenylpropenoate; Benzyl γ -phenylacrylate; Cinnamein; NSC 11780; NSC 44403 | |

Current regulation: Annex III, part 1, n° 81

Clinical data:
 In the "background information" section of the 1999 opinion, benzyl cinnamate (synonymous: benzyl 3-phenyl-2-propenoate, cinnamein) is classified as "less frequently reported allergen"; one study of patients with contact allergy to cosmetic products was identified and further a study where benzyl cinnamate associated with contact sensitisation to MP (33).

Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded 0.3% (95% CI: 0.1 – 0.6%) positive reactions in 2042 consecutively P_Ted patients, test concentration 5% pet. (4). The IVDK 2010 study, n=4 weak positive were observed, amounting to 0.12% (95% CI: 0 – 0.25%; percentages standardised for age and sex) of 2872 patients P_Ted reacted to the compound (7). In the Groningen 2009 study, no positive reactions to this allergen, using the same test concentration, were observed in 320 patients (6). In the Wöhrl 2001 study, P_Ting 747 patients with suspected contact allergy to fragrance ingredients yielded n=3 (0.4%) positive reactions (22).

Additional information: A RIFM review is available (56).

| | |
|--|---|
| BENZYL SALICYLATE |  |
| CAS # 118-58-1 | |
| EC # 204-262-9 | |
| Benzyl 2-hydroxybenzoate | |
| Salicylic acid, Benzyl ester; Benzoic acid, 2-Hydroxy-, phenylmethyl ester; Benzyl o-hydroxybenzoate; NSC 6647 | |

Current regulation: Annex III, part 1, n° 75

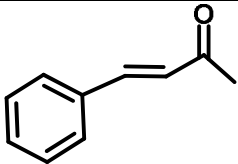
Clinical data:

In the "background information" section of the 1999 opinion (33), benzyl salicylate is classified among the frequent allergens, with 0.2 to 10% of patients with eczema from cosmetic products testing positively. In one study, benzyl salicylate accounted for 75% of reactions to commercial products (33).

Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded n=2, i.e. 0.1% (95% CI: 0.01 – 0.4%) positive reactions in 2041 consecutively PTed patients (test concentration 1% pet.) (4). The IVDK 2010 study, 2 of 3775 patients PTed reacted weakly positive to the compound (7). In the Groningen 2009 study, n=1, i.e. 0.3% (95% CI: 0.01 – 1.7%) had positive reactions to this allergen, tested at 2% pet. (6). In the deGroot 2000 study, 10 of 1825 consecutive patients tested positive to benzyl salicylate (2% pet.), of these, 3 were not detected by the FM I (12). In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded n=3 (0.4%) positive reactions (22). Trattner/David found 2 positive cases in 641 consecutive eczema patients (31). In a study from Alicante, Spain, 86 selected patients were tested with benzyl salicylate, yielding 2 positive reactions (48).

Additional information:

It is a "top 100" substance and classified as R43 (IFRA, pers. comm.2010). A RIFM review is available, including internal results on, e.g. HRIPT, and a review of LLNA results, where benzyl salicylate is classified as "weak" allergen (57).

| | |
|--|---|
| BENZYLIDENEACETONE |  |
| CAS # 122-57-6 | |
| EC # 204-555-1 | |
| 4-Phenyl-3-buten-2-one | |
| 4-Phenylbut-3-en-2-one; 2-Butenone, 4-Phenyl- (2CI); Ketone, Methyl styryl (7CI); 1-Phenyl-1-buten-3-one; 2-Phenylethenyl methyl ketone; 2-Phenylvinyl methyl ketone; 4-Phenyl-3-buten-2-one; 4-Phenyl-3-butene-2-one; 4-Phenylbutenone; Acetocinnamone; Benzalacetone; Benzylideneacetone; Methyl 2-phenylvinyl ketone; Methyl phenylvinyl ketone; Methyl styryl ketone; Methyl β-styryl ketone; NSC 5605; Styryl methyl ketone | |

Current regulation: Annex II, n° 356

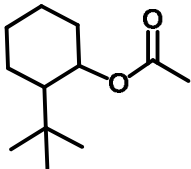
Clinical

data:

In the Malten 1984 study, none of 182 patients displayed a positive PT reaction to

benzylidene acetone 0.5% pet. (24).

Additional information: /

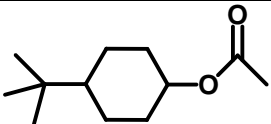
| | |
|---|---|
| 2-TERT-BUTYLCYCLOHEXYL ACETATE |  |
| CAS # 88-41-5 | |
| EC # 201-828-7 | |
| 2-(1,1-dimethylethyl)cyclohexyl acetate | |
| Cyclohexanol, 2-(1,1-dimethylethyl)-, acetate ; Cyclohexanol, 2-Tert-butyl-, acetate; 2-Tert-Butylcyclohexanol acetate; Verdox; o-Tert-Butylcyclohexyl acetate | |

Current regulation: /

Clinical data:
In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and 5% "Verdox ®" in pet., tested in 313 consecutive patients in Bordeaux and London, were observed (15)

Additional information:

It is a "top 100" substance (IFRA, pers. comm.2010). A RIFM review is available (58).

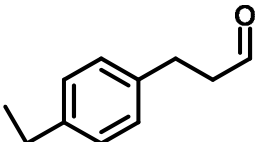
| | |
|--|---|
| 4-TERT-BUTYLCYCLOHEXYL ACETATE |  |
| CAS # 32210-23-4 | |
| EC # 250-954-9 | |
| 4-(1,1-Dimethylethyl)cyclohexyl acetate | |
| Boisinol A 464D; Cyclohexanol, 4-tert-Butyl-, acetate; Cyclohexanol, 4-(1,1-Dimethylethyl)-, acetate; 4-(1,1-Dimethylethyl)cyclohexyl acetate; 4-tert-Butylcyclohexanol acetate; Dorisyl; Madeflor; NSC 163103; Oryclone, Oryclone special, Oryclon extra; p-t-BCHA; p-tert-Butylcyclohexyl acetate; para-tert-Butylcyclohexyl acetate; PTBCHA; Velvetone; Verbeniax; Vertenex; Vertinate; Vertopol; Ylanate | |

Current regulation: /

Clinical data:
In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and 5% "Vertenex ®" in pet., tested in 107 consecutive patients in High Wycombe, were observed (15).

Additional information:

It is a "top 100" substance (IFRA, pers. comm.2010). A RIFM review is available (59).

| | |
|---|---|
| p-tert -Butyldihydrocinnamaldehyde |  |
| CAS # 18127-01-0 | |
| EC # 242-016-2 | |

| | |
|--|--------------------------------------|
| 4-(1,1-Dimethylethyl)-benzenepropanal | |
| p-tert-Butyl-hydrocinnamaldehyde; Butylphenyl)propanal; Butyldihydrocinnamaldehyde | 3-(4-tert- Bourgeonal; p-tert- |

Current regulation: III/155

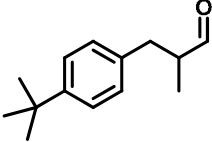
Clinical data: /

Additional

information:

It is a "top 200" substance and classified as R43 (IFRA, pers. comm.2010)

http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details_v2&id=39132

| | |
|---|---|
| BUTYLPHENYL METHYLPROPIONAL (Lilial®) |  |
| CAS # 80-54-6 | |
| EC # 201-289-8 | |
| 3-(4-tert-Butylphenyl)-2-methylpropanal | |
| <p>p-t-Butyl-alpha-methylhydrocinnamic aldehyde; 2-(4-tert-Butylbenzyl)propionaldehyde (REACH, EINECS); 4-(1,1-Dimethylethyl)-alpha-methyl-benzenepropanal; Hydrocinnamaldehyde, p-tert-Butyl-alpha-methyl-; (±)-2-Methyl-3-(4-tert-butylphenyl)propanal; 2-Methyl-3-(4-tert-butylphenyl)propanal; 2-[(4-tert-Butylphenyl)methyl]propanal; 3-(4-tert-Butylphenyl)-2-methylpropanal; 3-(p-tert-Butylphenyl)-2-methylpropionaldehyde; 3-(p-tert-Butylphenyl)isobutylaldehyde; 4-(1,1-Dimethylethyl)-alpha-methylbenzenepropanal; 4-tert-Butyl-alpha-methylhydrocinnamic aldehyde; Lilestralis; Lilial; Lysmeral; NSC 22275; lilestral; p-tert-Butyl-alpha-methylhydrocinnamaldehyde; p-tert-Butyl-alpha-methylhydrocinnamic aldehyde; pt-Bucinal; alpha-Methyl-p-tert-butylhydrocinnamaldehyde; beta-Lilial</p> | |

Current regulation: Annex III, part 1, n° 83

Clinical data:

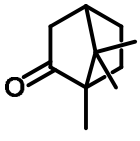
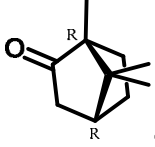
In the "background information" section of the 1999 opinion, lilial is classified as "less frequently reported allergen"; with 2 cases of contact allergy reported in 1 study of 176 eczema patients and 1 case with contact allergy to Lilial from a deodorant; a number of other reported positive cases were considered to possibly have been false positive (33).

Since the last SCCNFP-opinion of 1999, the Frosch 2002a study yielded 0.2% positive reactions to Lilial® (10% pet.) among the 1855 consecutive patients tested (16). The IVDK 2007 study yielded 0.4% (95% CI: 0.2 – 0.8%) positive reactions in 2004 patients consecutively tested (4). The IVDK 2010 study, 0.62% (95% CI: 0.04 – 1.21%; percentages standardised for age and sex) of 1947 patients PTed reacted to the compound (7). In the Groningen 2009 study, n=2, i.e. 0.6% (95% CI: 0.1 – 2.2%) had positive reactions to this allergen, tested at only 1% pet. (6). In the deGroot 2000 study, 9 of 1825 consecutively tested patients had a positive reaction to lilial® (5%

pet.) (12). Lilial® has been identified as constituent of perfumes used by a patient, causing ACD (60).

Additional information:

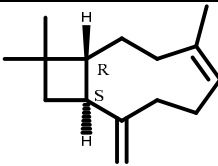
It is a "top 100" substance and classified as R43 (IFRA, pers. comm.2010).

| | |
|---|---|
| CAMPHOR |  76-22-2 |
| CAS # 76-22-2 / 464-49-3 | |
| EC # 207-355-2 / 200-945-0 | |
| 1,7,7-Trimethyl-bicyclo[2.2.1]heptan-2-one (76-22-2) (1R,4R)-1,7,7-Trimethyl-bicyclo[2.2.1]heptan-2-one (464-49-3) | |
| 76-22-2: DL-Bornan-2-one (REACH, EINECS); 2-Bornanone; Bornan-2-one, INCI name according to CAS; CAMPHOR/DL-bornan-2-one; Camphor; (±)-Camphor; DL-Camphor; 1,7,7-Trimethylnorcamphor; 2-Camphanone; Alphanon; Borneo camphor; Root bark oil; Spirit of camphor |  464-49-3 |
| 464-49-3: (1R)-1,7,7-Trimethyl-bicyclo[2.2.1]heptan-2-one; (1R,4R)-(+)- Camphor; (+)-2-Bornanone; (+)-Camphor; (1R)-(+)-Camphor; (1R)-Camphor; (1R,4R)-(+)-Camphor; (R)-(+)-Camphor; (R)-Camphor; Camphor; D-Camphor; D-(+)-Camphor; Alcanfor; Japanese camphor. | |

Current regulation: /

Clinical data:
From the UK, a case of allergic contact dermatitis after application of Earex ® ear drops due to rectified camphor oil (tested 10% pet.) was reported (61). Application of a liquid rubefacient of Asian origin caused allergic contact dermatitis in a 58-year-old patient, according to the positive PT result with 10% camphor ("alcaonfor") in pet. due to this ingredient (62). In the US, a case of contact dermatitis due to "Vics VapoRub" has been reported (63).

Additional information: It is a "top 100" substance (IFRA, pers. comm. 2010).

| | |
|---|---|
| beta-CARYOPHYLLENE |  |
| CAS # 87-44-5 | |
| EC # 201-746-1 | |
| (1R,4E,9S)-4,11,11-Trimethyl-8-methylene-bicyclo[7.2.0]undec-4-ene | |
| (E)-(1R,9S)-(-)-4,11,11-Trimethyl-8-methylene-bicyclo[7.2.0]undec-4-ene; [1R-(1R*,4E,9S*)]-4,11,11-Trimethyl-8-methylene-bicyclo[7.2.0]undec-4-ene; (-)-(E)-Caryophyllene; (-)-Caryophyllene; (-)-E-Caryophyllene; (-)-trans-Caryophyllene; (-)-β-Caryophyllene; (E)-Caryophyllene; Caryophyllene; Caryophyllene B; NSC 11906; l-Caryophyllene; trans-Caryophyllene; β-Caryophyllen; β-Caryophyllene; (-)-β-Caryophyllene | |

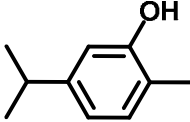
Current regulation: /

Clinical data:

In the Frosch 2002 b study, 0.6% positive reactions to caryophyllene (5% pet.) in 1606 consecutive were observed (17).

Additional information:

beta-Caryophyllene autoxidizes at air exposure. As the primary oxidation products, the hydroperoxides, are very unstable and immediately form epoxides with low sensitizing capacity, the increase in allergenic activity caused by autoxidation is comparably low (64). A multicenter study identified 0.5% positive reactions to oxidized *beta*-caryophyllene (3.0% pet.) in 1511 consecutive patients (65). Of these, 2 patients (0.1%) reacted to the major oxidation product (caryophyllene oxide) (3.9% pet.).

| | |
|---|---|
| CARVACROL |  |
| CAS # 499-75-2 | |
| EC # 207-889-6 | |
| 2-Methyl-5-(1-methylethyl)-phenol | |
| 2-Hydroxy-1-methyl-4-(1-methylethyl)benzene; 2-Hydroxy-p-cymene; 2-Methyl-5-(1-methylethyl)phenol; 2-Methyl-5-isopropylphenol; 3-Isopropyl-6-methylphenol; 5-Isopropyl-2-methylphenol; 5-Isopropyl-o-cresol; 6-Methyl-3-isopropylphenol; Antioxine; Dentol; Isopropyl o-cresol; Isothymol; NSC 6188; p-Cymen-2-ol | |

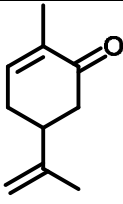
Current regulation: /

Clinical data:

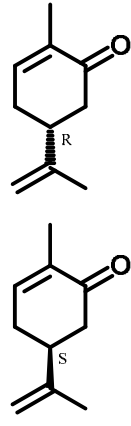
The DeGroot 1985 study identified 2 (1.1%) positive reactions among 179 patients using a 5% PT preparation of this compound – these reactions may have been at least partly due to an “excited back syndrome” and are thus of limited evidence (25). Meynadier et al. ¹¹ patch tested 28 patients with contact allergy to fragrance ingredients using 2% carvacrol in pet. Positive reactions were observed in 3 of 28 patients (after (66)).

Additional information:

Carvacrol is derived from p-cymene by sulfonation followed by alkali fusion. Carvacrol can also be derived from savory, thyme, marjoram, oregano, lovage root, and Spanish origanum oil (66). Carvacrol is a flavor ingredient that can be found in alcoholic beverages, baked goods, chewing gum, condiment relish, frozen dairy, gelatin pudding, non-alcoholic beverages, and soft candy at concentrations from 0.1 to 28.54 ppm (RIFM 2001, according to (66)).

| | |
|---|---|
| CARVONE |  |
| CAS # 99-49-0 / 6485-40-1 / 2244-16-8 | |
| EC # 202-759-5 / 229-352-5 / 218-827-2 | |
| 2-Methyl-5-(1-methylethenyl)-2-cyclohexen-1-one (99-49-0) | |
| | 99-49-0 |

¹¹ Meynadier, J. M., J. Meynadier, J. L. Peyron, and L. Peyron. 1986. Clinical forms of skin manifestations in allergy to perfume. *Ann. Dermatol. Venerol.* 113:31–39.

| | |
|---|---|
| <p>(5R)-2-Methyl-5-(1-methylethenyl)-2-cyclohexen-1-one (6485-40-1)</p> <p>(5S)-2-Methyl-5-(1-methylethenyl)-2-cyclohexen-1-one (2244-16-8)</p> |  <p>6485-40-1</p> <p>2244-16-8</p> |
| <p>99-49-0: p-Mentha-6,8-dien-2-one; (±)-Carvone; 2-Methyl-5-isopropenyl-2-cyclohexenone; 5-Isopropyl-2-methyl-2-cyclohexen-1-one; Carvone; DL-Carvone; Karvon; Limonen-6-one; NSC 6275; p-Mentha-1(6),8-dien-2-one</p> <p>6485-40-1: R)-(-)-p-Mentha-6,8-dien-2-on); (-)-(5R)-Carvone; (-)-(R)-Carvone; (-)-Carvone; (-)-p-Mentha-6,8-dien-2-one; (4R)-(-)-Carvone; (R)-(-)-Carvone; (R)-Carvone; L(-)-Carvone; L-Carvone; l-1-Methyl-4-isopropenyl-6-cyclohexen-2-one; l-Carvone</p> <p>2244-16-8: (S)-(+)-p-Mentha-6,8-dien-2-one; (+)-Carvone; (S)-(+)-Carvone; (S)-(+)-p-Mentha-6,8-dien-2-one; (S)-Carvone; (+)-Carvone; D-(+)-Carvone; D-Carvone; Talent; d-1-Methyl-4-isopropenyl-6-cyclohexen-2-one; (S)-2-Methyl-5-(1-methylvinyl)cyclohex-2-en-1-one; d-Carvone</p> | |

Current regulation: /

Clinical data:

Cases of allergic contact cheilitis due to L-carvone in toothpastes have been reported (67-69). In an earlier study, 15 of 541 (2.8%) of consecutive PT patients tested also with L-Carvone (5% pet.) exhibited positive reactions, which were (i) associated with positive PT results to *Compositae* mix and (ii) mostly were not considered clinically relevant. Upon re-testing with lower concentrations (2% and 1% pet.) only 2 of 8 patients thus tested were positive (70).

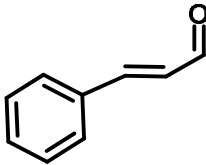
"Carvone has occasionally been reported as an allergen, usually in flavourings. Isomers of carvone have been either a mint or a rye flavour and aroma. We report a woman with positive patch-test reactions to carvone (newly added to the North American Contact Dermatitis Group standard series) and dermatitis on the head. She had used a hair conditioner with a "mint" scent, and the dermatitis resolved when she discontinued using this product. While the manufacturer would not confirm carvone as an ingredient, the clinical course, patch-test results, and ingredient list strongly suggest that this was a relevant allergen in this case of allergic contact dermatitis"¹²

Additional information:

D-Carvone occurs in caraway seed oil and dill oil in a concentration of up to 60%. L-Carvone is a component of the oil from *Mentha spicata* (spearmint).

R-Carvone is identified as a secondary oxidation product in autoxidized limonene (71). However, it is not a major allergen in this oxidation mixture and only one of 30 patients with known contact allergy to oxidized R- limonene reacted when tested with carvone (3% pet.) (72). Experimental findings in guinea pigs show no cross reactivity between R- and S carvone, but both enantiomers were found to be equally strong sensitizers (73).

¹² <http://www.ncbi.nlm.nih.gov/pubmed/20233552>

| | |
|---|---|
| CINNAMAL |  |
| CAS # 104-55-2 | |
| EC # 203-213-9 | |
| 3-Phenyl-2-propenal | |
| Cinnamaldehyde; 3-Phenyl-2-propen-1-al; 3-Phenyl-2-propenaldehyde; 3-Phenylacrolein; 3-Phenylacrylaldehyde; 3-Phenylpropenal; Abion CA; Benzylideneacetaldehyde; Cassia aldehyde; Cinnacure; Cinnamal; Cinnamic aldehyde; Cinnamite; Cinnamyl aldehyde; NSC 16935; NSC 40346; Phenylacrolein; Zimtaldehyde; β -Phenylacrolein | |

Current regulation: Annex III, part 1, n° 76

Clinical data:
In the "background information" section of the previous opinion (33), cinnamal, one of the 8 constituents of the FM I, is classified as frequent allergen, causing allergic reactions in a notable persons with eczema from cosmetic products in several studies (33).

Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded 1.0% (95% CI: 0.6 – 1.6%) positive reactions in 2063 consecutively PTed patients (4). In the Groningen 2009 study, 1.6% (95% CI: 0.5 – 3.6%) had positive reactions to cinnamal (6). In a study by the North American Contact Dermatitis Group, no significant trend of cinnamal contact sensitisation in the consecutive patients analysed was observed between 1984 (5.9% pos.) and 2000 (3.6% pos.); tested at 1% pet. (74). In the An 2005 study, 7 of 422 consecutive patients, i.e., 1.7%, had positive reaction (13). The Belsito 2006 study (20) yielded 1.7% positive reactions. In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded 1.9% positive reactions (22). The NACDG study found 3.1% positive reactions in 4435 patients tested (21). The IVDK 2010 study, 1.43% (95% CI: 0.67 – 2.18%) of 1214 consecutively tested patients reacted to the compound, while 2.64% (95% CI: 2.16 – 3.13%) of 4527 of patients tested in a more aimed manner, partly as break-down testing to the FM I, had a positive PT reaction (7). In a study from Alicante, Spain, 86 selected patients were patch tested with an extended fragrance series; n=7 reacted positively to cinnamal (48).

While, in addition to typical ACD due to contact sensitisation, immediate reactions to some fragrance compounds (and MPR, see below) are observed not infrequently, such immediate type reactions may rarely be very severe (anaphylaxis) and possibly immunologically mediated, as illustrated by the case of a 42 year old nurse with anaphylaxis (maximum grade of contact urticaria syndrome) 20 min after application of cinnamal (75). Following industrial use as "odour masking" agent, cinnamal caused occupational ACD in an exposed worker (76).

Additional information:

A specific RIFM review is available (77); another RIFM review addresses several cinnamic compounds (78).

| | |
|-------------------------------|---|
| CINNAMYL ALCOHOL |  |
| CAS # 104-54-1 | |
| EC # 203-212-3 | |
| 3-Phenyl-2-propen-1-ol | |

| | |
|---|--|
| Cinnamyl alcohol; 1-Phenyl-3-hydroxy-1-propene; 1-Phenylprop-1-en-3-ol; 3-Hydroxy-1-phenylprop-1-ene; 3-Phenyl-2-propenol; 3-Phenylallyl alcohol; Cinnamic alcohol; NSC 623440; NSC 8775; Styrene; Styryl alcohol; Styryl carbinol; γ -Phenylallyl alcohol | |
|---|--|

Current regulation: Annex III, part 1, n° 69

Clinical data:

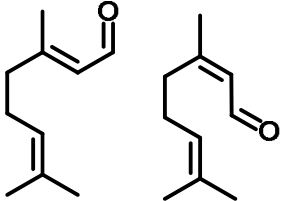
In the "background information" section of the previous opinion (33), cinnamyl alcohol, one of the 8 constituents of the FM I, is classified as frequent allergen, causing allergic reactions in a notable persons with eczema from cosmetic products (33).

Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded 0.6% (95% CI: 0.3 – 1.1%) positive reactions in 2063 consecutively PTed patients (4). In the Groningen 2009 study, 2.5% (95% CI: 1.1 – 4.9%) had positive reactions to cinnamyl alcohol, tested at 2% pet., i.e., twice the commonly used concentration (6). As test concentrations of up to 5% are apparently non-irritating (de Groot et al. after (33)), the latter data can be regarded as valid. In the An 2005 study, 13 of 422 consecutive patients, i.e., 3.1%, had positive reaction (13) (test concentration 2%). In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded 1.5% positive reactions (22). The IVDK 2010 study, 0.73% (95% CI: 0.17 – 1.30%) of 1214 consecutively tested patients reacted to the compound, while 2.36% (95% CI: 1.89 – 2.83%) of 4502 of patients tested in a more aimed manner, partly as break-down testing to the FM I, had a positive PT reaction (7). In a study from Alicante, Spain, 86 selected patients were patch tested with an extended fragrance series; n=12 reacted positively to cinnamyl alcohol (48).

Additional information:

In a recent experimental study protein-cinnamal adducts were detected in skin homogenates treated with cinnamal and cinnamyl alcohol but not with alpha-amyl cinnamal. This suggests that there is a common hapten involved in cinnamal and cinnamyl alcohol sensitization, in line with the observation of a marked concordance upon patch testing (7, 79), and that metabolic activation (to cinnamal) is involved in the latter. Conversely, there does not appear to be a common hapten for cinnamal and alpha-amyl cinnamal (80), again in line with the observations in the IVDK 2010 study (7).

A RIFM review is available (81)

| | |
|--|--|
| CITRAL |  <p>Citral = isomeric mixture of Geranial and Neral</p> |
| CAS # 5392-40-5 | |
| EC # 226-394-6 | |
| 3,7-Dimethyl-2,6-octadienal | |
| 3,7-Dimethyl-2,6-octadien-1-al; Citral; Citral PQ Extra; Lemarome N; Lemsyn GB; NSC 6170 | |

Current regulation: Annex III, part 1, n° 70

Clinical data:

In the "background information" section of the previous opinion (33), citral is classified as frequent allergen, causing about 1% allergic reactions in consecutive PT patients, and being a proven cause of contact allergic reactions in 2.6% patients with eczema from

cosmetic products (33).

Since the last SCCNFP-opinion of 1999, the Frosch 2002 a study yielded 1.1% positive (and 1.3% doubtful) reactions among the 1855 consecutive patients tested (16). In a study on 586 consecutive patients with hand eczema it has been noted that citral (2% pet.) not only caused (mostly weak) positive PT reactions, but far more often irritant reactions (n=82 vs. n=28). It was hypothesised that this very property could contribute to citral's sensitising potential (82). In the EU 2005 study, 12 of 1701 patients (0.7%, 95% CI: 0.4 – 1.2%) reacted positively to 2% citral in pet. (10). The IVDK 2007 study yielded 0.6% (95% CI: 0.3 – 1.1%) positive reactions in 2021 consecutively PTed patients; 10 of 13 citral positive patients also reacted positively to geraniol (4). In the Groningen 2009 study, 0.6% (95% CI: 0.1 – 2.2%) had positive reactions to this allergen (6). In the deGroot 2000 multicentre study, 19 of 1825 consecutive patients tested positively to citral (2% pet.), 4 of whom did not react positively to the FM I (12). In the An 2005 study, 5 of 422 consecutive patients, i.e., 1.2%, had positive reaction (13) (test concentration 2%). In the Malten 1984 study, neral at 1% in pet. yielded 2.6% positive reactions in 182 patients (24). In a study from Alicante, Spain, 86 selected patients were tested with citral, yielding 2 positive reactions (48).

Citral in a lip salve has been reported to have caused longstanding, recurrent allergic contact cheilitis in a 30 year old female patient, diagnosed by a strong positive reaction to the FM II, followed by a strong positive reaction to citral (83).

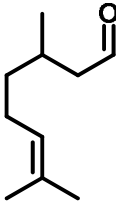
Additional information:

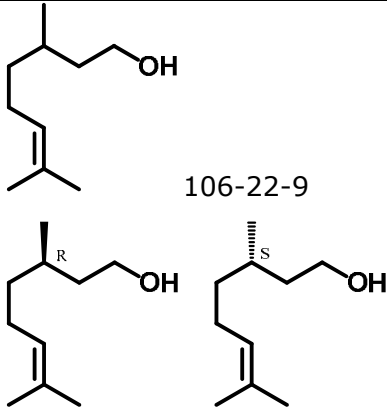
Citral is the mixture of two isomers: cis-citral (neral) and trans-citral (geranial).

Geranial forms oxidation product with increased sensitizing capacity both via spontaneous autoxidization at air exposure and via metabolic oxidation (Hagvall L. Thesis 2009: <http://hdl.handle.net/2077/18951>).

Geranial and neral have been identified as secondary oxidation products when geraniol autoxidizes (84). They have also been identified as metabolites of geraniol (85). This explains the simultaneous reactions to geraniol and citral seen by (4).

It is a "top 100" substance and classified as R43 (IFRA, pers. comm.2010).

| | |
|--|---|
| CITRONELLAL |  |
| CAS # 106-23-0 | |
| EC # 203-376-6 | |
| 3,7-Dimethyl-6-octenal | |
| (±)-Citronellal; 2,3-Dihydrocitral; 3,7-Dimethyloct-6-en-1-al; Citronellal; NSC 46106; Rhodinal; dl-Citronellal; β-Citronellal | |
| Current regulation: / | |
| Clinical / | data: |
| Additional information: A compound of essential oils of citrus fruits, namely grapefruit, but also contained in "citronella oil" and oil of Melissa. | |

| | |
|---|--|
| CITRONELLOL |  |
| CAS # 106-22-9 / 1117-61-9 / 7540-51-4 | |
| EC # 247-737-6 / 214-250-5 / 231-415-7 | |
| 3,7-Dimethyl-6-octen-1-ol (106-22-9); (3R)-3,7-Dimethyl-6-octen-1-ol (1117-61-9); (3S)-3,7-Dimethyl-6-octen-1-ol (7540-51-4) | |
| 106-22-9: (±)-3,7-Dimethyl-6-octen-1-ol; (±)-Citronellol; (±)-β-Citronellol; 2,3-Dihydrogeraniol; 2,6-Dimethyl-2-octen-8-ol; Cephrol; Citronellol; Citronello 950; DL-Citronellol; Dihydrogeraniol; NSC 8779; Rodinol; dl-Citronellol; β-Citronellol | |
| 1117-61-9: (R)-3,7-Dimethyl-6-octen-1-ol; (R)-(+)-3,7-Dimethyl-6-octen-1-ol; (+)-(R)-Citronellol; (+)-Citronellol; (+)-β-Citronellol; (3R)-(+)-β-Citronellol; (R)-(+)-Citronellol; (R)-(+)-β-Citronellol; (R)-Citronellol; (R)-β-Citronellol; D-Citronellol; d-Citronellol | |
| 7540-51-4: (-)-3,7-Dimethyl-6-octen-1-ol; (-)-(S)-Citronellol; (-)-Citronellol; (-)-β-Citronellol; (S)-(-)-Citronellol; (S)-(-)-β-Citronellol; (S)-3,7-Dimethyl-6-octen-1-ol; (S)-Citronellol; (S)-β-Citronellol; L-Citronellol; l-Citronellol | |
| Current regulation: Annex III, part 1, n° 86 | |

Clinical data:

In the "background information" section of the 1999 opinion, citronellol is classified as "less frequently reported allergen"; with few cases of contact allergy reported in the literature (33).

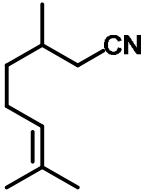
Since the last SCCNFP-opinion of 1999, in the Larsen 2002 c study, „DL citronello" (5% in pet.) elicited positive PT reactions in 8.7% of the patients (1). In 1855 consecutive patients of the Frosch 2002 a study, 0.4% positive reactions were noted (16). In the EU 2005 study, 4 of 1701 patients (0.2%, 95% CI: 0.06 – 0.6%) reacted positively to 1%

citronellol in pet.; at the same concentration, n=23 doubtful or irritant reactions were observed (10). The IVDK 2007 study yielded 0.5% (95% CI: 0.2 – 0.9%) positive reactions in 2003 patients consecutively PTed (4). In the Groningen 2009 study, n=1, i.e. 0.3% (95% CI: 0.01 – 1.7%) had positive reactions to this allergen, tested at only 2% pet. (6). The Larsen 2001 study yielded 5.6% positive reactions to l-citronellol (5% pet.) in 178 patients with known contact allergy to fragrance ingredients (19).

Additional information:

Citronellol autoxidizes spontaneously in contact with air in the same way as linalool forming allergenic primary oxidation products, hydroperoxides (AT Karlberg, personal communication, 2011).

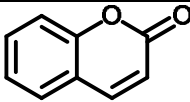
RIFM reviews have been published regarding L-citronellol (86), D-citronellol (87) and DL-citronellol (88). Another review is available by Hostynek and Maibach (89). It is a "top 100" substance and classified as R43 (IFRA, pers. comm.2010).

| | |
|--|--|
| CITRONELLYL NITRILE |  |
| CAS # 51566-62-2 | |
| EC # 257-288-8 | |
| 3,7-Dimethyl-6-octenenitrile | |
| 3,7-Dimethyl-6-octenenitrile (REACH, EINECS, INCI); Agrunitril; Agrunitrile; Citronellyl nitrile | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010)

| | |
|---|---|
| COUMARIN |  |
| CAS # 91-64-5 | |
| EC # 202-086-7 | |
| 2H-1-Benzopyran-2-one | |
| 1,2-Benzopyrone; 2-Chromenone; 2-Propenoic acid, 3-(2-hydroxyphenyl)-, δ-lactone; 5,6-Benzo-2-pyrone; Benzo-α-pyrone; Coumarinic anhydride; NSC 8774; Rattex; Tonka bean camphor; cis-o-Coumarinic acid lactone; o-Hydroxycinnamic acid lactone | |

Current regulation: Annex III, part1, n° 77

Clinical data:

In the "background information" section of the previous opinion (33), coumarin is classified as frequent allergen, causing allergic reactions in about 0.4 – 0.8% in consecutive PT patients, and causing contact allergic reactions in 0.8-10% of patients with eczema from cosmetic products (33).

Since the last SCCNFP-opinion of 1999, in the Frosch 2002 a study, 0.3% positive PT reactions to consecutive patients were noted (16). In the EU 2005 study, none of the

1701 patients reacted positively to 5% coumarin in pet., while 7 doubtful or irritant reactions were observed (10). The IVDK 2007 study yielded 0.4% (95% CI: 0.2 – 0.8%) positive reactions in 2020 consecutively P Ted patients (4). In the Groningen 2009 study, 0.6% (95% CI: 0.1 – 2.2%) had positive reactions to this allergen (6). In the deGroot 2000 study, 13 of 1825 consecutive patients reacted positively to coumarin (5% pet.) (12).

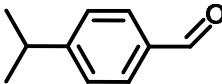
V. Mutterer et al. present the case of a 44 year old patient in whom coumarin was identified as culprit allergen by controlled ROAT testing with 1%, after having caused dermatitis by the use of a deodorant containing coumarin at 0.23% and an EdT (90).

Additional information:

Coumarin is found in several plant families, including *Melilotus* and *Galium*, e.g., *Galium odoratum* (sweet woodruff), however, also in oil of lavender, lovage and others (53).

Researchers from INSERM and "Rhodia Organique, Lyon , France" observed that pure coumarin is not an allergen in the LLNA, however, commercially available materials, containing "contaminants" (3,4-dihydrocoumarin, 6-chlorocoumarin and 6,12-epoxy-6H,12H-dibenzo[b,f][1,5] dioxocin, were identified as weak and moderate sensitisers, resp. (91).

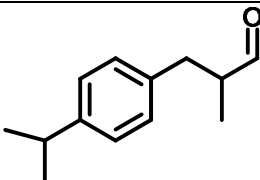
Coumarin is a "top 100" substance and classified as R43 (IFRA, pers. comm.2010).

| | |
|--|---|
| CUMINALDEHYDE |  |
| CAS # 122-03-2 | |
| EC # 204-516-9 | |
| 4-(1-Methylethyl)-benzaldehyde | |
| 4-Isopropylbenzaldehyde; p-Isopropylbenzaldehyde; 4-(Propan-2-yl)benzaldehyde; 4-Isopropylphenylcarboxaldehyde; Cumaldehyde; Cumeric aldehyde; Cuminal; Cuminaldehyde; Cuminic aldehyde; Cuminyl aldehyde; NSC 4886; p-Cumic aldehyde; p-Isopropylbenzaldehyde; p-Isopropylbenzenecarboxaldehyde | |

Current regulation: /

Clinical data:
The DeGroot 1985 study identified 3 (1.7%) positive reactions among 179 patients using a 15% PT preparation of cuminaldehyde (25).

Additional information: ...

| | |
|--|---|
| CYCLAMEN ALDEHYDE |  |
| CAS # 103-95-7 | |
| EC # 203-161-7 | |
| α-Methyl-4-(1-methylethyl)-benzenepropanal | |
| p-Isopropyl-α-methyl-hydrocinnamaldehyde; 2-Methyl-3-(4-isopropylphenyl)propionaldehyde; 2-Methyl-3-(p-isopropylphenyl)propionaldehyde; 3-(4-Isopropylphenyl)-2-methylpropanal; 3-(p-Isopropylphenyl)-2- | |

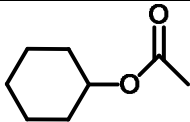
| | | |
|---|--|--|
| methylpropionaldehyde; methylpropionaldehyde(REACH, EINECS); methylhydrocinnamic aldehyde; Cyclamal; Cyclamen aldehyde; Cyclosal; Cyclosal perfume; Cymal; p-Isopropyl- α - methylhydrocinnamaldehyde; methylethyl)benzenepropanal; isopropylhydrocinnamaldehyde | 3-p-Cumenyl-2- 4-Isopropyl- α - α -Methyl-4-(1- α -Methyl-p- | |
|---|--|--|

Current regulation: ...

Clinical
/

data:

Additional information: It is a "top 100" substance and classified as R43 (IFRA, pers. comm.2010).

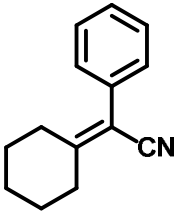
| | |
|---|---|
| CYCLOHEXYL ACETATE |  |
| CAS # 622-45-7 | |
| EC # 210-736-6 | |
| Cyclohexyletanoat | |
| Acetic acid cyclohexanyl ester; Acetoxycyclohexane; Cyclohexyl acetate; NSC 8772 | |

Current regulation: /

Clinical data:

In the Larsen 2002 c study, 0.5% positive reactions among 218 patients with know contact allergy to fragrance ingredients were observed (1).

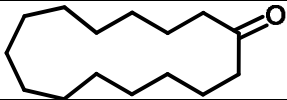
Additional information: A RIFM review is available (92).

| | |
|--|---|
| <i>alpha</i>-CYCLOHEXYLIDENE BENZENEACETONITRILE |  |
| CAS # 10461-98-0 | |
| EC # 423-740-1 | |
| α-Cyclohexylidenebenzeneacetonitrile | |
| α -Cyclohexylidene-benzeneacetonitrile (REACH); Δ 1 α - Phenyl- α -Cyclohexaneacetonitrile; 2-Cyclohexylidene-2- phenylacetonitrile; NSC 408284; Peonile (REACH) | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

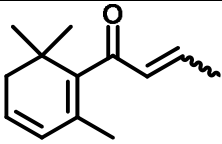
| | |
|---------------------------|---|
| CYCLOPENTADECANONE |  |
| CAS # 502-72-7 | |

| | |
|---|--|
| EC # 207-951-2 | |
| Cyclopentadecanone | |
| CPE 218; Exaltone; NSC 63900; Normuscon; Normuscone | |

Current regulation: /

Clinical data:
In the Larsen 2001 study, n=3, i.e., 1.7% positive reactions were observed to the compound, tested 5% pet., in 178 patients with known contact allergy to fragrance ingredients (19).

Additional information: ...

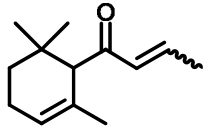
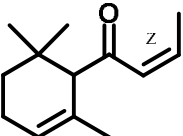
| | |
|--|---|
| DAMASCENONE |  |
| ROSE KETONE-4 (Not officially an INCI Name but Perfuming Name; Damascenone as such is not listed in CosIng) | |
| CAS # 23696-85-7 | |
| EC # 245-833-2 | |
| 1-(2,6,6-Trimethyl-1,3-cyclohexadien-1-yl)-2-buten-1-one | |
| 1-(2,6,6-Trimethyl-1,3-cyclohexadienyl)-2-buten-1-one; 1-Crotonoyl-2,6,6-trimethyl-1,3-cyclohexadiene; 2,6,6-Trimethyl-1-(2-butenoyl)-1,3-cyclohexadiene; 2,6,6-Trimethyl-1-crotonyl-1,3-cyclohexadiene; Rose ketone # 4 | |

Current regulation: Annex III, part1, n° 160 (max. conc. 0.02%)

Clinical data: /

Additional information:

RIFM reviews are available (93, 94), quoting 1 negative, and 2 positive (2 of 37, 1 of 50 volunteers) HRIPTs with damascenone based on 2 LLNA, the EC3 values were calculated as 1.24% and 1.22%, respectively (94).

| | | |
|---|---|------------|
| alpha-DAMASCONE (TMCHB) |  | 43052-87-5 |
| CAS # 43052-87-5 / 23726-94-5 | | |
| EC # x / 245-845-8 |  | 23726-94-5 |
| 1-(2,6,6-Trimethylcyclohex-2-en-1-yl)but-2-enone (43052-87-5); (2Z)-1-(2,6,6-Trimethyl-2-cyclohexen-1-yl)-2-buten-1-one (23726-94-5) | | |
| 43052-87-5: 2,6,6-Trimethyl-1-crotonyl-2-cyclohexene; α -Damascone | | |
| 23726-94-5: (Z)-1-(2,6,6-Trimethyl-2-cyclohexen-1-yl)-2-buten-1-one; (Z)- α -Damascone; cis- α -Damascone | | |

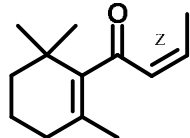
Current regulation: Annex III, part1, n° 157 (max. conc. 0.02%)

Clinical data:
In the Frosch 2002 b study, n=8 (0.5%) mostly strong positive PT reactions to consecutive patients were noted using a mixture of alpha and beta damascene, 0.1% pet. each (17). In human sensitisation experiments, after epicutaneous induction with 30% 1-(2,6,6-trimethylcyclohex-2-en-1-yl)but-2-enone (TMCHB, CAS # 43052-87-5) with adjuvant, to enhance response to this weak sensitiser, 8 of 30 patients were elicited by a challenge with 3% TMCHB 2 weeks later (95).

Additional information:

The former CAS # refers to alpha-Damascone or 1-(2,6,6-Trimethyl-2-cyclohexen-1-yl)-2-Buten-1-one. The latter CAS # refers to the identified ingredient cis-alpha-Damascone or (Z)-1-(2,6,6-Trimethyl-2-cyclohexen-1-yl)-2-buten-1-one, the content of which is restricted (SCCS-opinion 0392/00).

A RIFM review is available on alpha-damascone (96), quoting a number of partly positive HRIPT and other human studies, as well as different animal experiments. In 1 LLNA reported, an EC3 value of 3.3% was found. Another RIFM review is available for cis-alpha-damascone (97), supplying, however, no data on sensitisation.

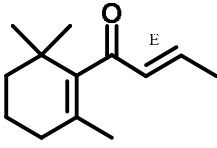
| | |
|---|---|
| cis-beta-DAMASCONE |  |
| CAS # 23726-92-3 | |
| EC # 245-843-7 | |
| (2Z)-1-(2,6,6-Trimethyl-1-cyclohexen-1-yl)-2-buten-1-one | |
| (Z)-1-(2,6,6-Trimethyl-1-cyclohexen-1-yl)-2-buten-1-one; (Z)- β -Damascone | |

Current regulation: Annex III, part 1, n° 162 (max. conc. 0.02%)

Clinical data:
Regarding results of the Frosch 2002 b study, see under alpha-damascone.

Additional information:

A RIFM review is available (98), citing several negative and one positive HRIPTs, and a number of – mostly positive – animal experiments.

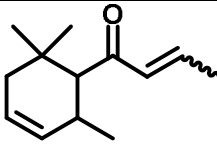
| | |
|---|---|
| <i>trans-beta-DAMASCONE</i> |  |
| CAS # 23726-91-2 | |
| EC # 245-842-1 | |
| (2E)-1-(2,6,6-Trimethyl-1-cyclohexen-1-yl)-2-buten-1-one | |
| (E)-1-(2,6,6-Trimethyl-1-cyclohexen-1-yl)-2-buten-1-one; (E)- β -Damascone; Damascone beta; trans-2,6,6-Trimethyl-1-crotonylcyclohex-1-ene; trans- β -Damascone; β -Damascone | |

Current regulation: Annex III, part 1, n° 158 (max. conc. 0.02%)

Clinical data: /

Additional information:

A RIFM review is available (99), citing 2 negative HRIPT and 1 negative maximisation test, and a number of positive animal experiments (the EC3 value, based on 1 LLNA, was found to be 2.4%).

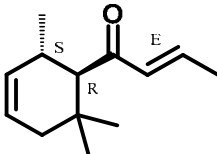
| | |
|--|---|
| <i>delta-DAMASCONE</i> |  |
| CAS # 57378-68-4 | |
| EC # 260-709-8 | |
| 1-(2,6,6-Trimethyl-3-cyclohexen-1-yl)-2-buten-1-one | |
| δ -Damascone | |

Current regulation: Annex III, part 1, n° 161 (max. conc. 0.02%)

Clinical data: /

Additional information:

A RIFM review is available (100), citing several positive HRIPT and 1 negative HRIPT. Cross sensitisation to alpha- and beta-damascone was demonstrated in 3 sensitised subjects. 2 LLNA studies are reported on, yielding EC3 values of 5.19% and 9.6%, resp.

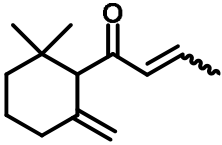
| | |
|---|---|
| <i>trans-trans-delta-DAMASCONE</i> |  |
| CAS # 71048-82-3 | |
| EC # 275-156-8 | |
| (2E)-rel-1-[(1R,2S)-2,6,6-Trimethyl-3-cyclohexen-1-yl]-2-buten-1-one | |
| [1 α (E),2 β]-1-(2,6,6-Trimethyl-3-cyclohexen-1-yl)-2-buten-1-one; trans- δ -Damascone; δ -Damascone; trans, trans- δ -Damascone | |

Current regulation: Annex III, part 1, n° 165 (max. conc. 0.02%)

Clinical data: /

Additional information:

A RIFM review is available (101), citing 1 positive HRIPT (2/15 with 1%).

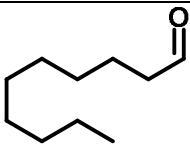
| | |
|---|---|
| <i>gamma-DAMASCONE</i> |  |
| CAS # 35087-49-1 | |
| EC # 481-910-9 | |
| 1-(2,2-Dimethyl-6-methylenecyclohexyl)-2-buten-1-one | |
| γ -Damascone | |

Current regulation: /

Clinical data: /

Additional information:

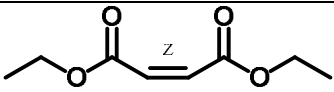
A RIFM review is available (102), citing 1 positive Buehler test and 1 LLNA study yielding an EC3 value of 4.6%

| | |
|---|---|
| <i>DECANAL</i> |  |
| CAS # 112-31-2 | |
| EC # 203-957-4 | |
| n-Decanal | |
| Capraldehyde; Capric aldehyde; Caprinaldehyde; Caprinic aldehyde; Decaldehyde; Decanaldehyde; Decyl aldehyde; Decylic aldehyde; NSC 6087; n-Decaldehyde; n-Decyl aldehyde | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|--|---|
| <i>DIETHYL MALEATE</i> |  |
| CAS # 141-05-9 | |
| EC # 205-451-9 | |
| (Z)-Diethyl but-2-enedioate | |
| 2-Butenedioic acid (Z)-, diethyl ester; 2-Butenedioic acid (Z)-, diethyl ester; Maleic acid, diethyl ester; (Z)-2-Butenedioic acid diethyl ester; Diethyl (Z)-2-butenedioate; Ethyl maleate; Staflex DEM | |

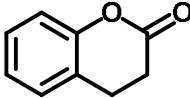
Current regulation: Annex II, n° 426

Clinical

data:

In the Malten 1984 study, 3.2% of 182 patients displayed a positive PT reaction to diethyl maleate 0.1% pet. (24). In this study, it has been noted that "in the max. test and clinically this is a strong sensitiser having caused patch test sensitisation (42%)"

Additional information: /

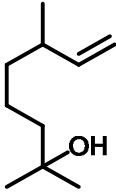
| | |
|---|---|
| DIHYDROCOUMARIN |  |
| CAS # 119-84-6 | |
| EC # 204-354-9 | |
| 3,4-Dihydro-2H-1-benzopyran-2-one | |
| Hydrocoumarin; Hydrocinnamic acid, o-hydroxy-, δ -lactone; 2-Chromanone; 3,4-Dihydro-1H-benzopyran-2-one; 3,4-Dihydrocoumarin; Dihydrocoumarin; Melilotin; Melilotin (coumarin); Melilotol | |

Current regulation: Annex II, n° 427

Clinical data:

In the Malten 1984 study, 3.7% of 182 patients displayed a positive PT reaction to dihydrocoumarine 5% pet. (24).

Additional information: /

| | |
|--|---|
| DIHYDROMYRCENOL |  |
| CAS # 18479-58-8 | |
| EC # 242-362-4 | |
| (±)-2,6-Dimethyloct-7-en-2-ol | |
| 1,1,5-Trimethyl-6-heptenol; 2,6-Dimethyl-7-octen-2-ol; 3,7-Dimethyl-1-octen-7-ol; 2,6-Dimethyl-7-octen-2-ol (INCI) | |

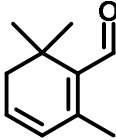
Current regulation: /

Clinical data: /

Additional information:

A RIFM review is available (103), listing 2 negative HRIPTs and 1 negative human maximisation test.

It is a "top 100" substance (IFRA, pers. comm.2010).

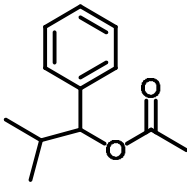
| | |
|---|--|
| 2,3-DIHYDRO-2,2,6-TRIMETHYLBENZALDEHYDE |  |
| CAS # 116-26-7 | |
| EC # 204-133-7 | |
| 2,6,6-Trimethyl-1,3-cyclohexadiene-1-carboxaldehyde | |
| 2,2,6-Trimethyl-4,6-cyclohexadien-1-aldehyde; 2,6,6-Trimethyl-1,3-cyclohexadiene-1-aldehyde; Safranal | |

Current regulation: /

Clinical data: /

Additional information:

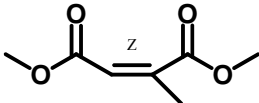
A RIFM review quotes one positive HRIPT (5 of 53) and one negative HRIPT (0 of 54) (93).

| | |
|---|---|
| DIMETHYLBENZYL CARBINYL ACETATE (DMBCA) |  |
| CAS # 151-05-3 | |
| EC # 205-781-3 | |
| 2-Methyl-1-phenylpropyl acetate | |
| Benzeneethanol, α,α-dimethyl-, acetate; Phenethyl alcohol, α,α-dimethyl-, acetate; 1,1-Dimethyl-2-phenylethyl acetate; 2-Methyl-1-phenyl-2-propyl acetate; 2-Methyl-1-phenylpropan-2-yl acetate; Benzyl dimethylcarbinol acetate; Benzyl dimethylcarbinyl acetate; Dimethylbenzylcarbinol acetate; Dimethylbenzylcarbonyl acetate; NSC 46123; α,α-Dimethylphenethyl acetate | |

Current regulation: /

Clinical data:
In the Frosch 2002 a study, 0.2% positive PT reactions to consecutive patients were noted (16). In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and one to 5% DMBCA in pet., tested in 313 consecutive patients in Bordeaux and London, were observed (15).

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

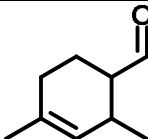
| | |
|---|---|
| DIMETHYL CITRACONATE |  |
| CAS # 617-54-9 | |
| EC # | |
| (2Z)-Diethyl-2-methyl-but-2-enedioate | |
| (2Z)-2-methyl-2-Butenedioic acid, dimethyl ester; 2-Butenedioic acid, 2-methyl-, dimethyl ester, (Z)-; Citraconic acid, dimethyl ester; Dimethyl methylmaleate; Methylmaleic acid, dimethyl ester | |

Current regulation: Annex II, n° 431

Clinical data:

In the Malten 1984 study, 3.7% of 182 patients displayed a positive PT reaction to dimethylcitraconate 12% pet. (24). In this paper, a human maximisation test positive in "4/44" is quoted.

Additional information: ...

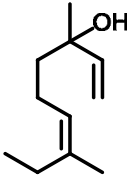
| | |
|---|--|
| 2,4-DIMETHYL-3-CYCLOHEXEN-1-CARBOXALDEHYDE |  |
| CAS # 68039-49-6 | |
| EC # 268-264-1 | |
| 2,4-Dimethyl-cyclohex-3-ene-1-carboxaldehyde | |
| (Z)-Vertocitral C; 2,4-Dimethyl-3-cyclohexene-1-carboxaldehyde; 2,4-Dimethyl-3-cyclohexenecarboxaldehyde; 2,4-Dimethyl-3-cyclohexenylcarbaldehyde; Cyclal C; Ligustral; Tricyclal; Triplal; Tripral; Zestover | |

Current regulation: /

Clinical data: /

Additional information:

It is a "top 100" substance and classified as R43 (IFRA, pers. comm.2010).

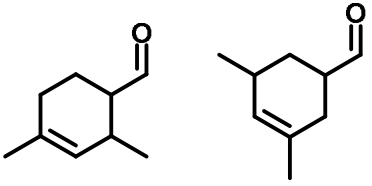
| | |
|--|---|
| 3,7-DIMETHYL-1,6-NONADIEN-3-OL |  |
| CAS # 10339-55-6 | |
| EC # 233-732-6 | |
| (7Z)-3,7-Dimethyl-1,6-nonadien-3-ol | |
| Ethyl linalool; Methyl linalool | |

Current regulation: /

Clinical data: /

Additional information:

It is a "top 100" substance (IFRA, pers. comm.2010). A RIFM review is available (104), citing 1 negative human maximisation test (n=25).

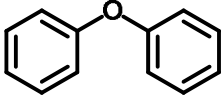
| | |
|--|---|
| DIMETHYLTETRAHYDRO BENZALDEHYDE |  |
| CAS # 68737-61-1 | |
| EC # 272-113-5 | |
| 2,4-Dimethyl-cyclohex-3-ene-1-carboxaldehyde 3,5-Dimethyl-cyclohex-3-ene-1-carboxaldehyde | |
| Hivertal; Vertocitral | |

Current regulation: /

Clinical data:

In the Larsen 2001 study, 2.3% positive PT reactions were observed with the isomer mixture, tested 5% pet., in 178 patients with known contact allergy to fragrance ingredients (19).

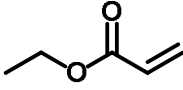
Additional information: /

| | |
|--|--|
| DIPHENYL ETHER |  |
| CAS # 101-84-8 | |
| EC # 202-981-2 | |
| Phenyl ether | |
| 1,1'-oxybis-Benzene; Barrel Therm 330; Benzene, phenoxy-; Biphenyl oxide; Chemcryl JK-EB; Diphenyl ether; Diphenyl oxide; NSC 19311; Oxybisbenzene; Phenoxybenzene; Phenyl oxide | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|---|---|
| ETHYL ACRYLATE |  |
| CAS # 140-88-5 | |
| EC # 205-438-8 | |
| Ethyl 2-propenoate | |
| Acrylic acid ethyl ester (6CI,8CI); 2-Propenoic acid ethyl ester; Ethyl 2-propenoate; Ethyl acrylate; Ethyl acrylic ester; Ethyl propenoate; NSC 8263 | |

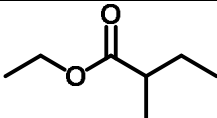
Current regulation: Annex II, n° 435

Clinical data:

In the Malten 1984 study, n=1 (0.5%) of 182 patients displayed a positive PT reaction to ethyl acrylate 1% pet. (24). In the NACDG 2009 multicentre study, 0.9% of

consecutive patients (n=4428) had a positive PT reaction (21).

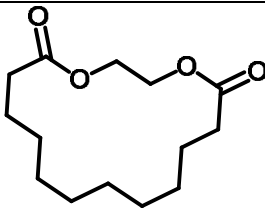
Additional information: /

| | |
|---|---|
| ETHYL 2-METHYLBUTYRATE |  |
| CAS # 7452-79-1 | |
| EC # 231-225-4 | |
| Ethyl 2-methylbutyrate | |
| Butyric acid, 2-methyl-, ethyl ester (6CI,7CI,8CI); (±)-Ethyl 2-methylbutanoate; 2-Methylbutanoic acid ethyl ester; 2-Methylbutyric acid ethyl ester; Ethyl 2-methylbutanoate; Ethyl 2-methylbutyrate; Ethyl α-methylbutyrate; NSC 1103 | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

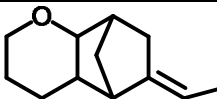
| | |
|--|---|
| ETHYLENE DODECANEDIOATE |  |
| CAS # 54982-83-1 | |
| EC # 259-423-6 | |
| 1,4-Dioxacyclohexadecane-5,16-dione | |
| Cyclic ethylene dodecanedioate; Ethylene dodecanedioate; Musk 144; Musk C-14 | |

Current regulation: /

Clinical data:

In the Larsen 2002 c study on 218 patients with known contact allergy to fragrance ingredients, this compound caused 0.9% positive PT reactions at 5% pet. (1).

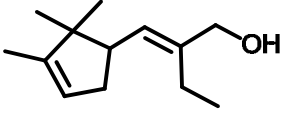
Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|---|---|
| 6-ETHYLIDENEOCTAHYDRO-5,8-METHANO-2H-BENZO-1-PYRAN |  |
| CAS # 93939-86-7 | |
| EC # 300-376-9 | |
| 6-Ethylideneoctahydro-5,8-methano-2H-1-benzopyran | |
| | |

Current regulation: /

Clinical data:
In the Larsen 2001 study, no positive PT reactions were observed with this compound, tested 5% pet., in 178 patients with known contact allergy to fragrance ingredients (19).

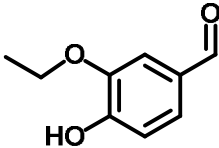
Additional information: /

| | |
|---|---|
| 2-ETHYL-4-(2,2,3-TRIMETHYL-3-CYCLOPENTEN-1-YL)-2-BUTEN-1-OL |  |
| CAS # 28219-61-6 | |
| EC # 248-908-8 | |
| 2-Ethyl-4-(2,2,3-trimethyl-3-cyclopenten-1-yl)-2-buten-1-ol | |
| 2-Ethyl-4-(2,2,3-trimethyl-3-cyclopenten-1-yl)-2-buten-1-ol; 2-Ethyl-4-(2',2',3-trimethylcyclopent-3'-enyl)but-2-enol; Bacdanol; Bangalol; Dartanol; Finanol; Levosandol; Radjanol; Sanjinol | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

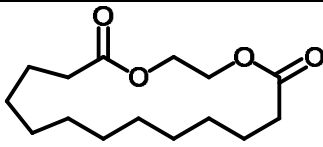
| | |
|--|---|
| ETHYL VANILLIN |  |
| CAS # 121-32-4 | |
| EC # 204-464-7 | |
| 3-Ethoxy-4-hydroxybenzaldehyde | |
| 2-Ethoxy-4-formylphenol; 3-Ethoxy-4-hydroxybenzaldehyde; 3-Ethylvanillin; 4-Hydroxy-3-ethoxybenzaldehyde; Arovanillon; Bourbonal; Ethavan; Ethovan; Ethylprotal; Ethylvanillin; NSC 1803; NSC 67240; Protocatechuic aldehyde ethyl ether; Quantrovanil; Rhodiarome; Vanillal; Vaniom | |

Current regulation: /

Clinical data:

The case of a 28-year-old metal grinder with allergic contact dermatitis to a "cutting oil reodorant" has been reported, who tested positively not only to the cutting fluid, the reodorant, but also to several ingredients of the latter product, including "Vanillal S 10026", 5% pet. (105).

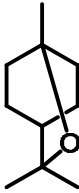
Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|---|---|
| ETHYLENE BRASSYLATE |  |
| CAS # 105-95-3 | |
| EC # 203-347-8 | |
| 1,4-Dioxacycloheptadecane-5,17-dione | |
| Tridecanedioic acid, cyclic ethylene ester; Ethylene glycol, cyclic tridecanedioate; Astratone; Cyclic ethylene glycol tridecanedioate; Cyclic ethylene tridecanedioate; Emeressence 1150; Ethylene brassylate; Musk T; NSC 46155 | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|---|---|
| EUCALYPTOL |  |
| CAS # 470-82-6 | |
| EC # 207-431-5 | |
| 1,3,3-Trimethyl-2-Oxabicyclo[2.2.2]octane | |
| 1,8-Epoxy-p-menthane; 1,3,3-Trimethyl-2-oxabicyclo[2.2.2]octane; 1,8-Cineol; 1,8-Cineole; 1,8-Epoxy-p-menthane; 2-Oxa-1,3,3-trimethylbicyclo[2.2.2]octane; Cajeputol; Cineol; Cineole; Eucalyptol; Eucalyptole; Eucalytol; Eucapur; Eukalyptol; NSC | |

Opinion on fragrance allergens in cosmetic products

| | |
|-------------------------|--|
| 6171; Terpan; p-Cineole | |
|-------------------------|--|

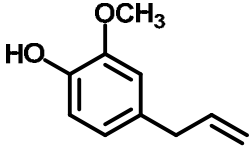
Current regulation: /

Clinical data: /

Additional information:

It is a "top 100" substance (IFRA, pers. comm.2010).

See also ***EUCALYPTUS SPP. LEAF OIL***; eucalyptol is the major ingredient there (up to 85%), but found in significant quantities also in a number of other essential oils (see 3.2).

| | |
|--|---|
| EUGENOL |  |
| CAS # 97-53-0 | |
| EC # 202-589-1 | |
| 2-Methoxy-4-(2-propen-1-yl)-phenol | |
| Other names: 4-Allyl-2-methoxy-phenol; 1-Allyl-4-hydroxy-3-methoxybenzene; 2-Hydroxy-5-allylanisole; 2-Methoxy-1-hydroxy-4-allylbenzene; 2-Methoxy-4-(2-propenyl)phenol; 2-Methoxy-4-(2'-propenyl)phenol; 2-Methoxy-4-[2-allyl]phenol; 2-Methoxy-4-allylphenol; 3-(3-Methoxy-4-hydroxyphenyl)propene; 3-(4-Hydroxy-3-methoxyphenyl)-1-propene; 4-Allyl-1-hydroxy-2-methoxybenzene; 4-Allyl-2-methoxyphenol; 4-Allylguaiacol; 4-Hydroxy-3-methoxyallylbenzene; Allylguaiacol; Bioxeda; Caryophylllic acid; Dentogum; Eugenic acid; Eugenol; NSC 209525; NSC 8895; p-Allylguaiacol; p-Eugenol | |

Current regulation: Annex III, part 1, n° 71

Clinical data:

In the "background information" section of the previous opinion (33), eugenol, one of the 8 components of the FM I, is classified as frequent allergen, causing allergic reactions in about 1.2% in consecutive PT patients and accounting for 4 to 16% of reactions to the FM I. Allergic reactions had been observed in 0.7 – 20% of patients with eczema from cosmetic products (33).

Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded 0.5% (95% CI: 0.3 – 1.0%) positive reactions in 2065 consecutively PTed patients (4). In the Groningen 2009 study, 1.3% (95% CI: 0.3 – 3.2%) had positive reactions to eugenol, tested at 2% pet., i.e., twice the commonly used concentration (6). F. Giusti et al. examined 1754 consecutive patients tested with eugenol 1% pet. in addition to the baseline series, 09/1998 - 01/2000. 21 patients (1.2%) reacted positively to eugenol (106). In the An 2005 study, 8 of 422 consecutive patients, i.e., 1.9%, had positive reaction (13) (test concentration 2%). In the Wöhrle 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded 2.5% positive reactions (22). The IVDK 2010 study, 0.44% (95% CI: 0.04 – 0.84%) of 1214 consecutively tested patients reacted to the compound, while 1.57% (95% CI: 1.19 – 1.95%) of 4801 of patients tested in a more aimed manner, partly as break-down testing to the FM I, had a positive PT reaction (7). In a study from Alicante, Spain, 86 selected patients were patch tested with an extended fragrance series; n=12 reacted positively to eugenol (48).

Moreover, eugenol is capable of inducing immediate type reactions of the airways, as illustrated by the well-documented case of a 30 year old hairdresser who developed severe occupational bronchial asthma due to eugenol (107). A case of urticaria after dental treatment with eugenol-containing material was reported from India (108); however, occasional cases are also reported from Europe (109). Occupational exposure to eugenol / zinc oxide type dental restorative material, which is apparently less frequently used nowadays, may lead to occupational sensitisation to eugenol, as illustrated by a case report (110).

Additional information:

Eugenol is the main component (80-95%) of clove oil, but also found in citronella oil, pimento leaf oil and cinnamon bark oil (see section 3.2).

It is a "top 100" substance and classified as R43 (IFRA, pers. comm.2010).

FARNESOL

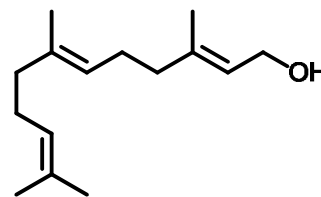
CAS # 4602-84-0

EC # 225-004-1

3,7,11-Trimethyl-2,6,10-Dodecatrien-1-ol

Farnesol; 3,7,11-Trimethyl-2,6,10-dodecen-1-ol; FCI 119a; Farnesyl alcohol; NSC 60597; Nikkosome

Current regulation: Annex III, part 1, n° 82



Clinical data:

In the “background information” section of the 1999 opinion, farnesol is classified as “less frequently reported allergen”; in 1 study of patients with cosmetic dermatitis 2 cases with contact allergy to farnesol had been reported; in other studies, positive reactions were seen in patients with positive PT reactions to MPR (33).

Since the last SCCNFP-opinion of 1999, farnesol is used not only for its scent, but also for its (slight) antimicrobial activity, useful, for instance, in deodorants. Thus, axillary dermatitis is a relatively typical presentation (111). In a multicentre study based on 1997/98 PT data, 0.5% positive reactions in consecutive patients were noted (Frosch 2002 a (16)). Farnesol is included in the FM II. In the original publication on single constituents of the FM II, 6 of 1701 consecutive patients reacted positively to farnesol 5%, ie., 0.35% (95% CI: 0.13 – 0.77%) (10). In a study on consecutive patients tested in 2003, 38 of 4238 patients had positive reactions to farnesol 5% pet. (0.9%, 95% CI: 0.6 – 1.2%) (4)(IVDK 2007). (A paper on farnesol previously published by the IVDK (112) presents results included in this later analysis.) In a series from Nagoya, Japan, 1.1% positive reactions in 1483 patients with suspected cosmetic dermatitis were observed (tested at 5% pet.) (14). In the Groningen 2009 study, 0.9% (95% CI: 0.2 – 2.7%) had positive reactions (6).

Additional information:

“Farnesol is an acyclic primary sesquiterpene alcohol found in essential oils such as lemongrass, citronella, tuberose blossom, sandalwood and orange blossom” (23). A RIFM review is available (113).

| | |
|---|--|
| GERANIOL | |
| CAS # 106-24-1 | |
| EC # 203-377-1 | |
| (2E)-3,7-Dimethyl-2,6-octadien-1-ol | |
| (E)-3,7-Dimethyl-2,6-octadien-1-ol; (E)-Geraniol; (E)-Nerol; 3,7-Dimethyl-trans-2,6-octadien-1-ol; Geraniol; Geranyl alcohol; Lemonol; MosquitoSafe; NSC 9279; trans-3,7-Dimethyl-2,6-octadien-1-ol; trans-Geraniol; β-Geraniol | |

Current regulation: Annex III, part 1, n° 78

Clinical data:

In the “background information” section of the previous opinion (33), geraniol, one of the 8 components of the FM I, is classified as frequent allergen, causing allergic reactions in about 0.4% in consecutive PT patients and accounting for 3 to 7% of reactions to the FM I. Allergic reactions had been observed in 1.2 – 30% of patients with

eczema from cosmetic products (33).

Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded 0.5% (95% CI: 0.2 – 0.9%) positive reactions in 2063 consecutively PTed patients (4). In the Groningen 2009 study, 0.6% (95% CI: 0.1 – 2.2%) had positive reactions to this allergen, tested at 2%, i.e. twice the usual concentration (6). In a series from Nagoya, Japan, 0.3% positive reactions in 1483 patients with suspected cosmetic dermatitis were observed (tested at the unusually high concentration of 5% pet.) (14). In the Wöhrle 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded n=7 (0.9%) positive reactions (22). The IVDK 2010 study, 0.39% (95% CI: 0.10 – 0.69%) of 1214 consecutively tested patients reacted to the compound, while 0.87% (95% CI: 0.63 – 1.10%) of 5695 of patients tested in a more aimed manner, partly as breakdown testing to the FM I, had a positive PT reaction (7). In a study from Alicante, Spain, 86 selected patients were patch tested with an extended fragrance series; n=17 reacted positively to geraniol (48).

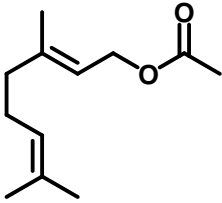
The fact that geraniol also occurs in food flavourings, and can elicit signs and symptoms of manifest contact sensitisation, is illustrated by the case of a 19 year old Japanese woman with cheilitis due to geraniol, improving after avoidance of respective foodstuff (114). A 20 year old Japanese woman with urticaria at the site of application of cosmetics with generalisation (contact urticaria syndrome grade 2), which A. Yamamoto et al. diagnosed as immediate type hypersensitivity to geraniol (without CA) (115).

Additional information:

Geraniol is a component of Palmarosa oil (CYMBOPOGON MARTINI see below), geranium oil (about 40%), citronella oil (30-40%), rose oil, lavender oil, and jasmine oil. It is sensitive to heat which induces autooxidation and isomeric with linalool (53).

Geraniol forms oxidation product with increased sensitizing capacity both via spontaneous autoxidation at air exposure and via metabolic oxidation. Geraniol and neral together with hydroperoxide have been identified as oxidation products when geraniol autoxidizes (84). Geraniol and neral were also identified as metabolites of geraniol (85). This explains the simultaneous reactions to geraniol and citral seen by (4).

A review is available by Hostynek and Maibach (116) and by RIFM (117). It is a "top 100" substance and classified as R43 (IFRA, pers. comm.2010).

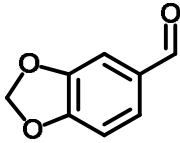
| | |
|---|---|
| GERANYL ACETATE |  |
| CAS # 105-87-3 | |
| EC # 203-341-5 | |
| (2E)-1-Acetate-3,7-dimethyl-2,6-octadien-1-ol | |
| (E)-Acetat-3,7-dimethyl-2,6-Octadien-1-ol; Geraniol acetate; (E)-3,7-Dimethyl-2,6-octadien-1-ol acetate; (E)-3,7-Dimethyl-2,6-octadienyl acetate; Acetic acid (2E)-3,7-dimethyl-2,6-octadienyl ester; Acetic acid geraniol ester; Bay pine (oyster) oil; Geranyl acetate; Geranyl ethanoate; NSC 2584; trans-1-Acetoxy-3,7-dimethyl-2,6-octadiene; trans-3,7-Dimethyl-2,6-octadien-1-yl acetate; trans-Geranyl acetate; β-Geranyl acetate | |

Current regulation: /

Clinical data: /

Additional information:

It is a "top 100" substance (IFRA, pers. comm.2010).

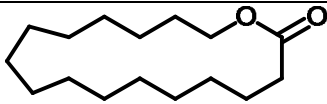
| | |
|--|---|
| HELIOTROPINE |  |
| CAS # 120-57-0 | |
| EC # 204-409-7 | |
| 1,3-Benzodioxole-5-carboxaldehyde | |
| Piperonal; 2H-Benzo[3,4-d]-1,3-dioxolan-5-ylformaldehyde; 3,4-(Methylenedioxy)benzaldehyde; 3,4- Dihydroxybenzaldehyde methylene ketal; 3,4- Dimethylenedioxybenzaldehyde; 5-Formyl-1,3- benzodioxolane; 5-Formyl-1,3-benzodioxole; 5- Formylbenzodioxole; Benzo[1,3]dioxole-5-carbaldehyde; Benzo[d][1,3]dioxole-5-carboxaldehyde; Heliotropin; Heliotropin; Heliotropine; NSC 26826; Piperonaldehyde; Piperonylaldehyde; Protocatechuic aldehyde methylene ether | |

Current regulation: /

Clinical data:

In the Frosch 2002 b study, n=2 (0.2%) positive reactions to "piperonal" (1% pet.) and n=6 (0.4%) to "piperonal" (5% pet.), respectively, in 1606 consecutive were observed (17). In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and 5% heliotropine in pet., tested in 106 consecutive patients in Barcelona, were observed (15).

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

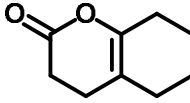
| | |
|---|---|
| HEXADECANOLACTONE |  |
| CAS # 109-29-5 | |
| EC # 203-662-0 | |
| Oxacycloheptadecan-2-one | |
| o-Lactone-16-hydroxy-hexadecanoic acid; 1,16- Hexadecanolide; 16-Hexadecanolactone; Cyclohexadecanolide; Dihydroambrettolide; Hexadecanoic acid, 16-Hydroxy-, o-lactone; Hexadecanolactone; Hexadecanolide; Juniperic acid lactone; NSC 33546 | |

Current regulation: /

Clinical data:

In the Larsen 2001 study, 1 of 178 patients with previously diagnosed contact allergy to fragrance ingredients had a positive PT reaction to this compound, tested 5% pet. (19). In the An 2005 study, 6 of 422 consecutive patients, i.e., 1.4%, had positive reactions to 5% "hexadecanolide" (13).

Additional information: /

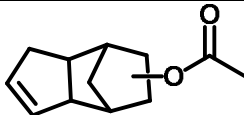
| | |
|--|---|
| HEXAHYDROCOUMARIN |  |
| CAS # 700-82-3 | |
| EC # 211-851-4 | |
| 3,4,5,6,7,8-Hexahydro-2H-1-benzopyran-2-one | |
| 3,4,5,6,7,8-Hexahydro-coumarin; δ -Lactone-2-hydroxy-1-cyclohexene-1-propanoic acid; 3,4,5,6,7,8-Hexahydrocoumarin; Hexahydrocoumarin; Δ -1,6-2-Oxabicyclo(4.4.0)decen-3-one | |

Current regulation: Annex II, n° 1135

Clinical data: /

Additional information:

A RIFM review is available (93), p. S115 ff, citing a number of positive human sensitisation experiments.

| | |
|--|--|
| 3α,4,5,6,7,7α-HEXAHYDRO-4,7-METHANO-1H-INDEN-5(OR 6)-YL ACETATE |  |
| CAS # 54830-99-8 | |
| EC # 259-367-2 | |
| 3α,4,5,6,7,7α-Hexahydro-4,7-methano-1H-indenol Acetate | |
| Acetoxidyhydrodicyclopentadiene; Cyclacet; Dicyclopentenyl acetate; Dicylat; Tricyclo[5.2.1.0 ^{2,6}]dec-3-enyl acetate; Tricyclodecenyyl acetate | |

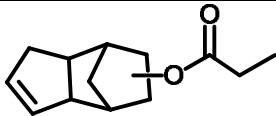
Current regulation: /

Clinical data:

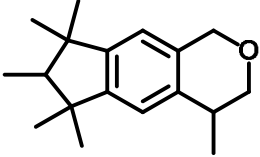
In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and 1 to 5% "Cyclacet ®" in pet., tested in 313 consecutive patients in Bordeaux and London, were observed (15).

Additional information:

Produced by IFF under the brand name "Cyclacet" (<http://www.iff.com/Ingredients.nsf/0/1C9F2CB39EB1EF6480256993002FBC14>, last accessed 2010-07-08).

| | |
|--|---|
| HEXAHYDRO-METHANOINDENYL PROPIONATE |  |
| CAS # 68912-13-0 | |
| EC # 272-805-7 | |
| 3α,4,5,6,7,7α-Hexahydro-4,7-methano-1H-indenol propanoate | |

| | |
|---|--|
| 3a,4,5,6,7,7a-Hexahydro-4,7-methano-1H-indenyl propionate (Mixture of Isomers); Dicyclopentadiene propionate; tricyclodecanyl propionate; Tricyclo[5.2.1.0 ^{2,6}]dec-3-enyl propionate; Verdyl propionate | |
| Current regulation: / | |
| Clinical data: / | |
| Additional information: It is a "top 100" substance (IFRA, pers. comm.2010). | |

| | |
|--|---|
| HEXAMETHYLINDANOPYRAN |  |
| CAS # 1222-05-5 | |
| EC # 214-946-9 | |
| 1,3,4,6,7,8-Hexahydro-4,6,6,7,8,8-hexamethylcyclopenta[γ]-2-benzopyran | |
| 1,3,4,6,7,8-Hexahydro-4,6,6,7,8,8-hexamethylcyclopenta[γ]-2-benzopyrane; 1,3,4,6,7,8-Hexahydro-4,6,6,8,8,8-hexamethylcyclopenta-2-benzopyran; Abbalide; Galaxolide; Galaxolide 50; Galaxolide 50BB; Galaxolide 50IPM; Galaxolide White; HHCB; Pearlide | |

Current regulation: /

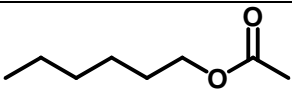
Clinical data:

In the Frosch 2002 a study, n=3 (0.2%) had positive reactions to the compound, tested 10% in isopropyl myristate (with 1 patient reacting positively to the diluent) (16). The Larsen 2001 study, testing with HHCB 7% pet., found 3.4% positive reactions in 178 patients with known contact allergy to fragrance ingredients (19). In the An 2005 study, 5 of 422 consecutive patients, i.e., 1.2%, had a positive reaction to "Galaxolide 50", tested at 5% (13) (test concentration 2% pet.). The DeGroot 1985 study identified 3 (1.7%) positive reactions among 179 patients using a 25% PT preparation of HHCB (25). In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and 5% "Galaxolide 50 ®" in pet., tested in 100 consecutive patients in Stockholm, were observed (15).

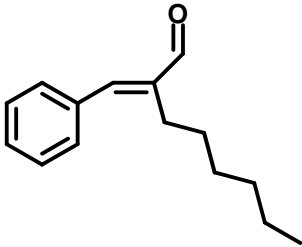
Additional information:

[0403/00 - Opinion concerning Hexahydro-hexamethyl-cyclopenta\(γ\)-2-benzopyran \(HHCB\)](#)

[0610/02 - Opinion on Hexahydro-hexamethyl-Cyclopenta \(γ\)-2-Benzopyran \(HHCB\)](#) (no restrictions) It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|--|---|
| HEXYL ACETATE |  |
| CAS # 142-92-7 | |
| EC # 205-572-7 | |
| Hexyl ethanoate | |
| Acetic acid, hexyl ester, Hexyl alcohol, acetate; 1-Hexyl acetate; Exceed 600; Hexyl acetate; Hexyl ester acetic | |

| | |
|--|--|
| acid;; NSC 7323; n-Hexyl acetate; n-Hexyl ethanoate | |
| Current regulation: / | |
| Clinical data: / | |
| Additional information: It is a "top 100" substance (IFRA, pers. comm.2010). | |

| | |
|--|---|
| HEXYL CINNAMAL |  |
| CAS # 101-86-0 | |
| EC # 202-983-3 | |
| α-Hexyl-cinnamaldehyde | |
| 2-(Phenylmethylene)octanal; 2-Hexyl-3-phenyl-2-propenal; 2-Hexylcinnamaldehyde; Hexyl cinnamic aldehyde; NSC 406799; NSC 46150; α-Hexylcinnamaldehyde; α-Hexylcinnamic aldehyde; α-Hexylcinnamyl aldehyde; α-n-Hexyl-β-phenylacrolein; α-n-Hexylcinnamaldehyde | |
| Current regulation: Annex III, part 1, n° 87 | |

Clinical data:

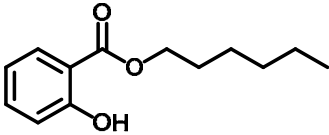
In the "background information" section of the 1999 opinion, hexyl cinnamal (synonymous: alpha-hexyl cinnamal, AHCA) is classified as "less frequently reported allergen"; 2 studies with 1 case and 1 study with 7 cases of contact allergy to this compound in patients with eczema from cosmetic products were found (33).

Since the last SCCNFP-opinion of 1999, in the Frosch 2002 a study, 0.3% positive PT reactions to consecutive patients were noted (16). In the subsequent EU 2005 study, 2 of 1701 patients had positive reactions to AHCA, and n=16 doubtful or irritant to AHCA at 10% in pet. (10). The IVDK 2007 study yielded n=3, i.e. 0.2% (95% CI: 0.03 – 0.4%) positive reactions in 2019 consecutively PTed patients, using 10% pet. as test concentration (4). In the Groningen 2009 study, 0.6% (95% CI: 0.1 – 2.2%) had positive reactions to this allergen, using a lower test concentration of 5% pet. (6).

Additional information:

It is a "top 100" substance and classified as R43 (IFRA, pers. comm.2010).

Hexyl cinnamal is regarded as "a recommended positive control for skin sensitization testing", e.g., in the context of the LLNA (118).

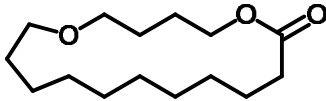
| | |
|---|---|
| HEXYL SALICYLATE |  |
| CAS # 6259-76-3 | |
| EC # 228-408-6 | |
| Hexyl-2-hydroxybenzoate | |
| Salicylic acid, hexyl ester; 1-Hexyl salicylate; Hexyl salicylate; n-Hexyl salicylate | |
| Current regulation: / | |

Clinical data:

None of the 218 patients with known contact allergy to fragrance ingredients reacted positively to this compound (tested at 5% in pet.) in the Larsen 2002 c study (1).

Additional information:

In a RIFM review, 2 human sensitisation experiments are mentioned which yielded no evidence of sensitising potential (HRIPT, n=103, maximisation test, n=22) (119). It is a "top 100" substance and classified as R43 (IFRA, pers. comm.2010).

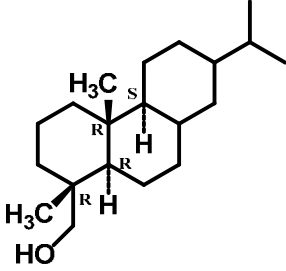
| | |
|---|---|
| HIBISCOLIDE |  |
| CAS # 6707-60-4 | |
| EC # 229-755-6 | |
| 1,6-Dioxacycloheptadecan-7-one | |
| Undecanoic acid, 11-(4-hydroxybutoxy)-, o-lactone; 12-Oxa-1,16-hexadecanolide; Cervolide; Musk 781; NSC 34741; 12-Oxahexadecan-16-olide | |

Current regulation: /

Clinical data:

None of the 178 patients with known contact allergy to fragrance ingredients reacted positively to "12-oxahexadecanolide" (tested at 5% in pet.) in the Larsen 2001 study (19).

Additional information: /

| | |
|--|---|
| HYDROABIETYL ALCOHOL, when used as a fragrance ingredient |  |
| CAS # 13393-93-6 | |
| EC # 236-476-3 | |
| (1R,4aR,4βS,10aR)-Tetradecahydro-1,4a-dimethyl-7-(1-methylethyl)-1-Phenanthrenemethanol | |
| Tetradecahydro-1,4a-dimethyl-7-(1-methylethyl)-phenanthrenemethanol; Tetrahydroabietyl alcohol | 1- |


Current regulation: AnnexII, n° 440

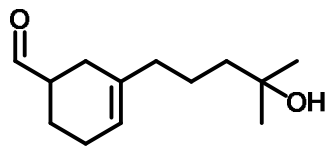
Clinical data:

In the deGroot 2000 study, 17 of 1825 consecutively tested patients had positive reactions to hydroabietyl alcohol (10% pet.) (12).

Additional information:

Commercial hydroabietyl alcohol consists of di- and tetrahydroabietyl alcohol together with non-modified colophony (120)

| | | |
|------------------------|----------------------|---|
| HYDROXYISOHEXYL | 3-CYCLOHEXENE |  |
|------------------------|----------------------|---|

| | |
|--|---|
| CARBOXALDEHYDE (HICC) regioisomers |  51414-25-6 |
| CAS # 31906-04-4 / 51414-25-6 | |
| EC # 250-863-4 / 257-187-9 | |
| 4-(4-Hydroxy-4-methylpentyl)-3-cyclohexene-1-carboxaldehyde (31906-04-4) | |
| 3-(4-Hydroxy-4-methylpentyl)-3-cyclohexene-1-carboxaldehyde (51414-25-6) | |
| 31906-04-4: 4-(4-Hydroxy-4-methylpentyl)-3-cyclohexenecarboxaldehyde; 4-(4-Methyl-4-hydroxyamyl)cyclohex-3-ene carboxaldehyde; Lyrall | |

Current regulation: Annex III, part 1, n° 79

Clinical data:

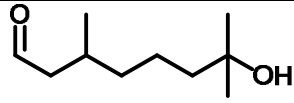
In the "background information" section of the previous opinion (33) HICC is classified as frequent allergen, causing allergic reactions in about 2.8% in consecutive PT patients, two thirds of these being relevant (33).

Since the last SCCNFP-opinion of 1999, in the Frosch 2002 a study, 2.7% of the 1855 consecutive patients reacted positively to HICC (5% pet.) (16). In the EU 2005 study, 28 of 1701 patients (1.7%, 95% CI: 1.1 – 2.4%) reacted positively to 5% HICC in pet. (10). In 21325 patients PTed consecutively in the IVDK 2007 study, 2.4% (95% CI: 2.2 – 2.6%) positive reactions were noted to 5% HICC in pet. (4). Similar to other studies, HICC was the most common single fragrance allergen among 320 patients tested in the Groningen 2009 study, with 3.1% (95% CI: 1.5 – 5.7%) positive reactions despite testing with a lower concentration of 2% pet. (6). In the An 2005 study, 7 of 422 consecutive patients, i.e., 1.7%, had positive reaction (13). The Belsito 2006 study (20) yielded a relatively low prevalence of 0.4% (7 of 1603; exact 95% CI (recalculated): 0.17 – 0.90%) positive reactions with 5% HICC in pet. and even less with lower test concentrations; possible reasons for the much lower prevalence were discussed. The IVDK 2010 study, 2.36% (95% CI: 2.19 - 2.53%) of 37270 consecutively tested patients reacted to HICC (7). In a study from Alicante, Spain, 86 selected patients were patch tested with an extended fragrance series; n=8 reacted positively to HICC (48).

Further clinical data with a focus on quantitative dose-response (see also section 4.3), is discussed in (121).

Among the early case reports, S.A. Hendriks reported the case of a 20 year old patient developing axillary dermatitis after 5 months use of a deodorant containing HICC (122).

Additional information: /

| | |
|--|---|
| HYDROXYCITRONELLAL |  |
| CAS # 107-75-5 | |
| EC # 203-518-7 | |
| 7-Hydroxy-3,7-dimethyl-octanal | |
| (±)-Hydroxycitronellal; 3,7-Dimethyl-7-hydroxyoctanal; 7-Hydroxy-3,7-dimethyloctanal; 7-Hydroxycitronellal; Citronellal hydrate; Citronellal, hydroxy-; Cyclalial; Cyclosia; Cyclosia base; Fixol; Hydroxycitronellal; Laurine; Lilyl aldehyde; Muguet synthetic; Muguetine principle; NSC | |

Opinion on fragrance allergens in cosmetic products

406740; Phixia

Current regulation: Annex III, part 1, n° 72

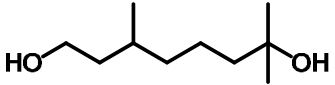
Clinical data:

In the "background information" section of the previous opinion (33), hydroxycitronellal, one of the 8 components of the FM I, is classified as frequent allergen, causing allergic reactions in about 0.75% in consecutive PT patients and accounting for 6 to 16% of reactions to the FM I. Allergic reactions had been observed in 10 – 45% of patients with eczema from cosmetic products (33).

Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded 1.3% (95% CI: 0.9 – 1.9%) positive reactions in 2063 consecutively PTed patients (4). In the Groningen 2009 study, 2.2% (95% CI: 0.9 – 4.5%) had positive reactions to this compound, tested at 2% pet., i.e., twice the commonly used concentration (6). The Sugiura 2000 study observed 1% positive PT reactions (test concentration 5% pet.) in 1483 patients tested for suspected cosmetic dermatitis (14). In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded 1.5% positive reactions (22). The IVDK 2010 study, 1.17% (95% CI: 0.48 – 1.85%) of 1214 consecutively tested patients reacted to the compound, while 2.95% (95% CI: 2.43 – 3.47%) of 4359 of patients tested in a more aimed manner, partly as break-down testing to the FM I, had a positive PT reaction (7). In a study from Alicante, Spain, 86 selected patients were tested with hydroxycitronellal, yielding 6 positive reactions (48).

Additional information:

Hydroxycitronellal is a synthetic fragrance, which only recently has been found in a few essential oils, e.g., of a *Narcissus* species and in essential oils of pepper (53)

| | |
|---|---|
| HYDROXYCITRONELLOL |  |
| CAS # 107-74-4 | |
| EC # 203-517-1 | |
| 3,7-Dimethyl-7-octanediol | |
| 2,6-Dimethyl-2,8-octanediol; 3,7-Dimethyl-1,7-octanediol; 3,7-Dimethyloctan-1,7-diol; Citronellol, hydroxy-; Hydroxyciol; Hydroxycitronellol; NSC 406140; NSC 67886 | |

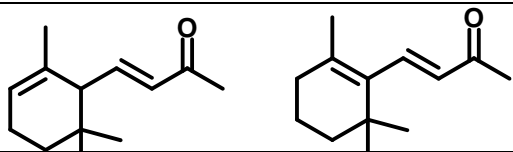
Current regulation: /

Clinical data:

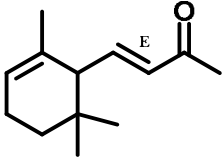
This compound elicited 6.0% positive PT reactions in 218 fragrance sensitive individuals (Larsen 2002 c, (1)).

Additional information:

A RIFM review is available, reporting results of a human induction study (maximisation test) in 25 volunteers, yielding no evidence of sensitisation (123).

| | |
|--------------------------------|--|
| IONONE isomeric mixture |  |
| CAS # 8013-90-9 | |
| EC # 232-396-8 | |

| | |
|---|--|
| Ionone | |
| Irisone, mixture of alpha- and beta ionone | |
| Current regulation: / | |
| Clinical data: / (see single isomers) | |
| Additional information: | |
| It is a "top 100" substance, further specified with "mixed isomers" (IFRA, pers. comm.2010). | |
| INCI: "MIXED IONONES", with CAS # 14901-07-6 / 6901-97-9 / 8013-90-9 (http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=35383 , last accessed 2010-07-13). | |
| A RIFM review is available on "ionone" (124), quoting negative human and experimental results. | |

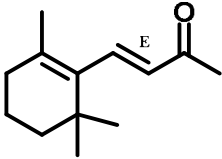
| | |
|---|--|
| alpha-IONONE |  |
| CAS # 127-41-3 | |
| EC # 204-841-6 | |
| (3E)-4-(2,6,6-Trimethyl-2-cyclohexen-1-yl)-3-Buten-2-one | |
| (E)-4-(2,6,6-Trimethyl-2-cyclohexen-1-yl)-3-Buten-2-one; (5E)-Ionone; (E)-4-(2,6,6-Trimethyl-2-cyclohexen-1-yl)-3-buten-2-one; (E)- α -Ionone; (\pm)-trans- α -Ionone; (\pm)- α -Ionone; 4-(2,6,6-Trimethyl-2-cyclohexen-1-yl)-3-buten-2-one; 4-(2,6,6-Trimethyl-2-cyclohexenyl)-3-buten-2-one; trans- α -Ionone; α -Cyclocitrylideneacetone; α -Ionone | |

Current regulation: /

Clinical data:

In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and 5% alpha-ionone in pet., tested in 205 consecutive patients, were observed (15).

Additional information: A RIFM review is available (125).

| | |
|---|---|
| beta-IONONE |  |
| CAS # 79-77-6 | |
| EC # 201-224-3 | |
| (3E)-4-(2,6,6-Trimethyl-1-cyclohexen-1-yl)-3-buten-2-one | |
| (E)-4-(2,6,6-Trimethyl-1-cyclohexen-1-yl)-3-buten-2-one; (E)- β -Ionone; Ionone beta; trans- β -Ionone; β -Ionone | |

Current regulation: /

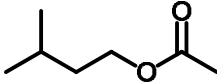
Clinical data:

In the Frosch 1995 dose finding pilot study, no positive reaction to 1% and 5% beta-

ionone in pet., tested in 205 consecutive patients, were observed (15).

Additional information:

It is a "top 100" substance (IFRA, pers. comm.2010). A RIFM review is available (126).

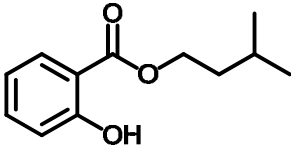
| | |
|---|--|
| ISOAMYL ACETATE |  |
| CAS # 123-92-2 | |
| EC # 204-662-3 | |
| 3-Methylbutyl acetate | |
| <p>1-Butanol, 3-methyl-, acetate; Acetic acid, isoamyl ester; Isopentyl alcohol, acetate; 3-Methyl-1-butanol acetate; 3-Methyl-1-butyl acetate; 3-Methylbutyl acetate; 3-Methylbutyl ethanoate; Acetic acid 3-methyl-1-butyl ester; Acetic acid 3-methylbutyl ester; Acetic acid isopentyl ester; Banana oil; Isoamyl acetate; Isoamyl alcohol acetate; Isoamyl ethanoate; Isopentyl acetate; Isopentyl ethanoate; NSC 9260; Pear oil; i-Amyl acetate; iso-Amyl acetate; iso-Pentyl acetate</p> | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

In CosIng, it is listed as "solvent"
<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=76810>,
 last accessed 2010-07-13)

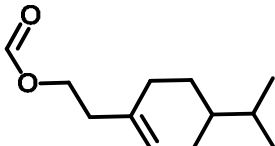
| | |
|--|---|
| ISOAMYL SALICYLATE |  |
| CAS # 87-20-7 | |
| EC # 201-730-4 | |
| 3-Methylbutyl-2hydroxybenzoate | |
| Isopentyl 2-Hydroxybenzoate; Isopentyl salicylate; Salicylic acid, isopentyl ester (6CI,8CI); Isopentyl alcohol, salicylate; 3-Methylbutyl salicylate; Isoamyl o-hydroxybenzoate; Isoamyl salicylate; Isopentyl salicylate; NSC 7952 | |

Current regulation: /

Clinical data:

The DeGroot 1985 study identified 1 (0.6%) positive reactions among 179 patients using a 50% PT preparation of this compound – this reaction may have been due to an “excited back syndrome” and is thus a limited evidence (25). In the Frosch 1995 dose finding pilot study, no positive reaction to 1% and 5% isoamyl salicylate in pet., tested in 95 consecutive patients, were observed (15).

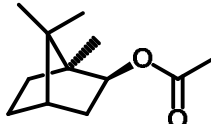
Additional information: A RIFM review is available (127).

| | |
|--|---|
| ISOBERGAMATE |  |
| CAS # 68683-20-5 | |
| EC # 272-066-0 | |
| 4-(Isopropyl)cyclohexadiene-1-ethyl formate | |
| Structure is incompletely defined 4-(1-Methylethyl)-1,?-cyclohexadiene-1-ethyl formate 4-(Isopropyl)cyclohexadiene-1-ethyl methanoate; menthadienyl formate; Menthadiene-7-methyl formate | |

Current regulation: Annex III, part 1, n° 170

Clinical data: /

Additional information: A RIFM review is available (128).

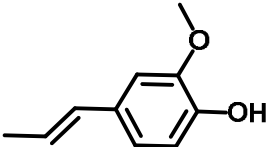
| | |
|---|---|
| ISOBORNYL ACETATE |  |
| CAS # 125-12-2 | |
| EC # 204-727-6 | |
| (1R,2R,4R)-1,7,7-trimethyl-Bicyclo[2.2.1]hept-2-yl acetate | |
| Bicyclo[2.2.1]heptan-2-ol, 1,7,7-Trimethyl-, acetate, (1R,2R,4R)-rel- ; Bicyclo[2.2.1]heptan-2-ol, 1,7,7-trimethyl-, acetate, exo-; Isoborneol, acetate; (±)-Isobornyl acetate; Isobornyl acetate; NSC 62486; Pichtosin; Pichtosine; exo-Bornyl acetate | |

Current regulation: /

Clinical data:

In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and 5% isobornyl acetate in pet., tested in 107 consecutive patients in High Wycombe, were observed (15).

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|---|---|
| ISOEUGENOL |  |
| CAS # 97-54-1 | |
| EC # 202-590-7 | |
| 2-Methoxy-4-(1-propen-1-yl)-phenol | |
| Phenol, 2-methoxy-4-(1-propenyl)- ; Phenol, 2-methoxy-4-propenyl-; 1-(3-Methoxy-4-hydroxyphenyl)-1-propene; 2-Methoxy-4-(1-propenyl)phenol; 2-Methoxy-4-propenylphenol; 3-Methoxy-4-hydroxy-1-propenylbenzene; 4-Hydroxy-3-methoxy-1-propenylbenzene; 4-Hydroxy-3-methoxy- β -methylstyrene; 4-Propenylguaiacol; Isoeugenol; NSC 6769 | |

Current regulation: Annex III, part 1, n° 73

Clinical data:

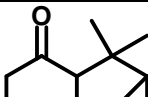
In the "background information" section of the previous opinion (33), isoeugenol, one of the 8 components of the FM I, is classified as frequent allergen, causing allergic reactions in about 1.9% in consecutive PT patients and accounting for 6 to 22% of reactions to the FM I. Allergic reactions had been observed in 2 – 25% of patients with eczema from cosmetic products (33).

Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded 1.3% (95% CI: 0.8 – 1.8%) positive reactions in 2063 consecutively PTed patients (4). In the Groningen 2009 study, 1.3% (95% CI: 0.3 – 3.2%) had positive reactions to isoeugenol, tested at 2% pet., i.e., twice the commonly used concentration (6). In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded 5.4% positive reactions (22). At St Johns Institute of Dermatology in London 3636 subjects were patch tested with isoeugenol 2001-2005, 97 of whom were positive. Year-on-year incidence showed an increasing trend, with an overall incidence of 2.67% (129). The IVDK 2010 study, 1.62% (95% CI: 0.87 – 2.38%) of 1214 consecutively tested patients reacted to the compound, while 3.41% (95% CI: 2.90 – 3.92%) of 5747 of patients tested in a more aimed manner, partly as break-down testing to the FM I, had a positive PT reaction (7). In a study from Alicante, Spain, 86 selected patients were patch tested with an extended fragrance series; n=11 reacted positively to isoeugenol (48).

Additional information:

Isoeugenol occurs in a cis- (CAS 5912-86-7) and a trans-isomers (CAS 5932-68-3), the latter dominating in trade products (82-88%) (53).

Isoeugenyl methyl ether caused 7.3% positive reactions in the Larsen 2002 c study (1). A number of derivatives of isoeugenol, such as isoeugenyl acetate, transisoeugenol, isoeugenyl benzoate, isoeugenyl phenylacetate, isoeugenyl methyl ether and benzyl isoeugenyl have been examined in 2261 consecutive patients; a varying proportion of positive patch test reactions and a varying proportion of concomitant reactions with isoeugenol have been observed (130). In an earlier study, 5 of 7 patients positive to isoeugenol also displayed positive reactions to isoeugenol acetate (1.2% eth.) (131) (see also section 5 and 6).

| | |
|-----------------------------|---|
| ISOLONGIFOLENEKETONE |  |
| CAS # 33407-62-4 | |

| | |
|---|--|
| EC # 245-890-3 | |
| 1,3,4,6,7,8a-Hexahydro-1,1,5,5-tetramethyl-2H-2,4a-methanonaphthalen-8(5H)-one | |
| Hexahydro-1,1,5,5-tetramethyl-2H-2,4a-methanonaphthalen-8(5H)-one | |

Current regulation: /

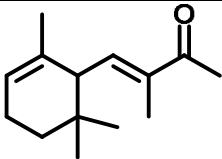
Clinical data:

The Larsen 2001 study identified 1 in 178 patients with known contact allergy to fragrance ingredients who reacted positively in the PT (5% pet.) (19).

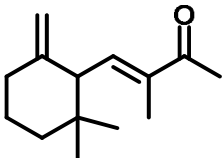
Additional information:

Not listed in CosIng under this CAS #. Other CAS # reported in RIFM review ¹³:

- 29461-14-1 CosIng: INCI name "ISOLONGIFOLENE KETONE EXO";
- 23787-90-8 CosIng: INCI name "ISOLONGIFOLANONE";
- 29461-13-0: CosIng: INCI name "HEXAHYDRO-TETRAMETHYLMETHANONAPHTHALEN-8-ONE".

| | |
|--|--|
| <i>alpha-ISOMETHYL IONONE</i> |  |
| CAS 127-51-5 | |
| EC 204-846-3 | |
| 3-Methyl-4-(2,6,6-trimethyl-2-cyclohexen-1-yl)-3-buten-2-one | |
| 4-(2,6,6-Trimethyl-2-cyclohexen-1-yl)-3-methyl-3-buten-2-one; Cetone Alpha; Isomethyl- α -ionone; NSC 66432; α -Cetone | |

Current regulation: Annex III, part 1, n° 90

| | |
|----------------------------------|---|
| <i>gamma-Methylionone</i> |  |
| CAS 7388-22-9 | |
| EC / | |
| | |

According to CosIng, "alpha-ISOMETHYL IONONE" (CAS # 127-51-5) and "gamma-Methylionone" (CAS # 7388-22-99) are synonyms, with one CAS number, and one preferred chemical name. The substance(s) are accordingly treated in the 1999 opinion (33) as one. As this treatment is also found in the literature, both substances are reviewed together.

¹³ Opdyke, D. L. J.; Letizia, C. **Monographs on fragrance raw materials. Isolongifolanone.** Food and Chemical Toxicology (1983), 21(6), 859

Clinical data:

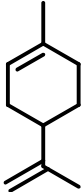
In the "background information" section of the 1999 opinion, "gamma-methylionone" is classified as "less frequently reported allergen"; 1 study with 2 cases and 2 studies with 1 case were found among patients with eczema from cosmetic products (33).

The IVDK 2007 study yielded n=1, i.e. 0.1% (95% CI: 0.00 – 0.2%) positive reactions in 2004 consecutively PTed patients (4). In the subsequent period (2005-2008), n=986 patients were tested in the IVDK 2010 study, with no positive reactions (7). In the Groningen 2009 study, n=2, i.e. 0.6% (95% CI: 0.1 – 2.2%) had positive reactions to this allergen, tested at only 1% pet. (6). In a Korean study with 422 consecutive patients, 2.1% reacted positively to "alpha isomethyl ionone (gamma-methylionone), CAS # 127-51-5", tested 5% pet. (13)

Additional information:

It is a "top 100" substance (IFRA, pers. comm.2010) under the label of "alpha-ISOMETHYL IONONE (CAS # 127-51-5)".

A RIFM review is available, listing 4 human sensitisation experiments employing different study protocols – all yielding negative results (132). Another review is available by Hostynek and Maibach (133), both referring to "alpha-ISOMETHYL IONONE (CAS # 127-51-5)".

| | |
|--|---|
| (DL)-LIMONENE |  |
| CAS # 138-86-3 | |
| EC # 231-732-0 | |
| 1-Methyl-4-(1-methylethenyl)-cyclohexene | |
| <p>p-Mentha-1,8-diene; (±)-Dipentene; (±)-Limonene; (±)-α-Limonene; 1,8-p-Menthadiene; 1-Methyl-4-(1-methylethenyl)cyclohexene; 1-Methyl-4-isopropenyl-1-cyclohexene; 1-Methyl-4-isopropenylcyclohexene; 1-Methyl-p-isopropenyl-1-cyclohexene; 4-Isopropenyl-1-methyl-1-cyclohexene; 4-Isopropenyl-1-methylcyclohexene; Cajeputen; Cajeputene; Cinen; Cinene; DL-Limonene; Dipenten; Dipentene; Eulimen; Flavor orange; Goldflush II; Kautschin; Limonen; Limonene; NSC 21446; NSC 844; Nesol; Orange X; Orange flavor; PC 560; Roti-Histol; SF 001; dl-Limonene; α-Limonene</p> | |
| Current regulation: Annex III, part1, n° 88, 167, 168 | |

Clinical data:

In the “background information” section of the 1999 opinion, d-limonene (CAS 5989-27-5) is classified as “less frequently reported allergen in relation to cosmetic exposure”; with contact allergy to oxidised limonene not infrequently reported in the literature (33).

Since 1999, several studies have been performed using limonene where the oxidation state is not given, but intended to be low. In one study, 0.6% positive reactions to limonene (3% pet.) were observed in 1606 consecutive patients (17). The IVDK 2007 study yielded n=3, i.e. 0.1% (95% CI: 0.03 – 0.4%) positive reactions in 2396 patients consecutively P_Ted with limonene (2% pet.) (4). The IVDK 2010 study, 0.28% (95% CI: 0 – 0.57%; percentages standardised for age and sex) of 1241 patients P_Ted with dipentene reacted to the compound (7). In the Groningen 2009 study, no positive reactions to this allergen, tested at 2% pet., were observed in 320 patients (6).

Regarding selected case reports, a case of a 40 year old citrus fruit picker with work related hand dermatitis and bronchial asthma has been described, who tested extreme positive to DL-limonene (2% pet.), and, less extremely, to citronellol and to the biocide dichlorophene (134). Moreover, limonene is used as a solvent in technical applications and cleaning and can lead to allergic contact dermatitis (e.g., a histopathology technicians (135, 136) or a painter and decorator (137)). In “water-free” hand cleansers it is reported to be used in concentrations around 10 – 20% (137). Wax polishes may contain dipentene and have caused one reported case of occupational ACD in a car mechanic (138). Another case of occupational ACD from dipentene in honing oil has been reported (139). In a case series from Sweden, 2 of 105 car mechanics patch tested for occupational contact dermatitis had positive reactions to oxidised *d*-limonene (5% pet.) (140).

Additional information:

Limonene is a monocyclic monoterpene existing in two enantiomers: (R)-(+)-limonene (CAS 5989-27-5) and (S)-(-)-limonene (CAS 5989-54-8). Racemic limonene is known as dipentene.

The allergenicity of limonene is closely related to oxidation (71, 72, 141, 142). It has been demonstrated that both enantiomers, R-(+)- and S-(-)-limonene spontaneously autoxidize, and that the primary oxidation products formed, the hydroperoxides, are strong and clinically relevant contact allergens. Among 2411 consecutive patients in a multi-centre European study, 63 (2.6 % [95%CI: 2.0-3.3]) reacted to oxidised (R)-(+)-and/or (S)-(-)-limonene (3.0% pet.) (72). In other multi-studies also, a considerable proportion of patients showed

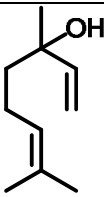
positive patch test reactions to oxidised R-(+)- limonene, e.g.,

- between 0.3% and 5.1% of subgroups of 2800 patients in Stockholm and Leuven, depending on test concentration, oxidation state and department(141),
- between 0.3% and 6.5% in 4 different departments in altogether 2273 patients (72, 143).

The primary oxidation products are the major allergens forming specific antigens (Bråred-Christensson J, Matura M, Bäcktorp C, Börje A, Nilsson JLG, Karlberg A-T. Hydroperoxides form specific antigens in contact allergy. Contact Dermatitis 2006; 55: 230-237.).

Current IFRA standards emphasise "a peroxide value of less than 20 millimoles peroxides per litre, determined according to the FMA method" (<http://www.ifraorg.org/Home/Code,+Standards+Compliance/IFRA+Standards/page.aspx/56>, last accessed 2009-11-11). For a more general discussion see section 5.

There is no scientific rationale for the difference in peroxide value allowed for limonene (20 millimoles peroxides per litre) compared to linalool (10 millimoles peroxides per litre). Specific values for hydroperoxides, which are allergens, would be desirable.

| | |
|--|--|
| LINALOOL |  |
| CAS # 78-70-6 (isomeric mixture) | |
| EC # 201-134-4; 245-083-6 See: http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=27933 | |
| 3,7-Dimethyl-1,6-octadien-3-ol | |
| (±)-Linalool; 2,6-Dimethyl-2,7-octadien-6-ol; 2-Methyl-1-prenyl-3-buten-2-ol; 3,7-Dimethyl-1,6-octadiene-3-ol; 3,7-Dimethyl-3-hydroxy-1,6-octadiene; L 260-2; Linalol; Linalool; Linalyl alcohol; Linanol; NSC 3789; dl-Linalool; β-Linalool | |

Current regulation: Annex III, part 1, n° 84

Clinical data:
In the "background information" section of the 1999 opinion, linalool in non-oxidized form is classified as "less frequently reported allergen"; with 4 cases of contact allergy reported in 2 studies on patients with eczema from cosmetic products (33).

Since the last SCCNFP-opinion of 1999, studies have been performed on contact allergy to linalool, oxidation state not given, but intended to be low. In the Larsen 2002 c study, none of the 218 patients with known contact allergy to fragrance ingredients had a positive reaction to linalool 5% pet., as prepared specially for this study (1). The IVDK 2007 study yielded 0.3% (95% CI: 0.1 – 0.6%) positive reactions in 2401 patients consecutively tested with stabilised linalool (10% pet.) (4). The IVDK 2010 study, 1 patient had a weak, and another a ++ reaction among the n=985 patients tested with 10% linalool (stabilised) in pet. (7). In the Groningen 2009 study, n=2, i.e. 0.6% (95% CI: 0.1 – 2.2%) had positive reactions to this allergen (6). The deGroot 2000 study with 1825 consecutively tested patients yielded 3 positive reactions to linalool (12). The DeGroot 1985 study found no positive reactions among 179 patients using a 30 % PT preparation of linalool (25).

Additional information:

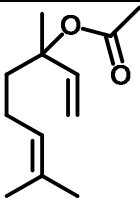
The allergenicity of linalool is closely related to oxidation and the primary oxidation products, the hydroperoxides, are the main allergens (144). In a clinical study 2002-

2003 in 6 European centres including 1511 consecutive patients, 1.3% showed a positive reaction to oxidized linalool (2.0% pet.) and 1.1% to the hydroperoxide fraction (65). A recent dose-response study in Sweden including 3400 patients in two test centres showed a positive reaction in 5.3% of the 1725 patients tested with oxidized linalool 6% pet. (145).

A review by RIFM is available both regarding linalool (146) and linalool "and related esters" (147). Another review is available by Hostynek and Maibach (148).

It is a "top 100" substance (IFRA, pers. comm.2010).

Additional CAS numbers exist for the single isomers: CAS # 126-90-9 (S-isomer), CAS # 126-91-0 (R-isomer); however, in the studies reviewed the isomeric mixture has been used throughout.

| | |
|--|---|
| LINALYL ACETATE |  |
| CAS # 115-95-7 | |
| EC # 204-116-4 | |
| 3,7-Dimethyl-1,6-octadien-3-yl acetat | |
| 1,6-Octadien-3-ol, 3,7-dimethyl-, acetate; Linalool acetate K; (±)-Linaloyl acetate; (±)-Linalyl acetate; 1,5-Dimethyl-1-vinyl-4-hexenyl acetate; 3,7-Dimethyl-1,6-octadien-3-yl acetate; 3-Acetoxy-3,7-dimethyl-1,6-octadiene; Acetic acid linalool ester; Bergamiol; Bergamol; Bergamot mint oil; Linalyl acetate; NSC 2138; dl-Linalool acetate | |

Current regulation: /

Clinical data:
In 100 patients tested in Odense, DK, in the early 90s, no positive reactions were observed with 1 and 5% linalyl acetate in pet. (15). In the Frosch 2002 a study, testing with linalyl acetate (10% pet.), 0.2% positive PT reactions to consecutive patients were noted (16). Similarly, the RIFM review mentioned quotes a number of studies where no allergic reactions to this compound had been observed, with the exception of one positive reaction in a Dutch study in 1988(149).

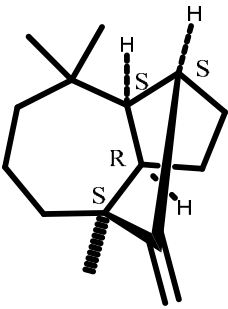
Additional information:

This is the main component of lavender oil (30%), also part of bergamot oil, neroli oil, peppermint oil, lemon oil and jasmine oil (53).

Linalyl acetate autoxidizes spontaneously at air exposure and the major allergens, the hydroperoxides, are the primary oxidation products (150). The pattern of autoxidation is similar to that for linalool and as the acetate can be metabolically hydrolysed to the corresponding alcohol cross reactions to allergens from oxidized linalool should be possible. This was indicated in a study of lavender oil and oxidised linalyl acetate which elicited positive PT reactions in some patients with known contact allergy to oxidised linalool (n=3) (151).

A RIFM review is available reporting 7 human sensitisation experiments yielding few or no cases of sensitisation (152).

It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|--|---|
| Longifolene |  |
| CAS # 475-20-7 | |
| EC # 207-491-2 | |
| (1S,3aR,4S,8aS)-Decahydro-4,8,8-trimethyl-9-methylene-1,4-methanoazulene | |
| 1,4-Methanoazulene, decahydro-4,8,8-trimethyl-9-methylene-, (1S,3aR,4S,8aS)-(+)-; 1,4-Methanoazulene, decahydro-4,8,8-trimethyl-9-methylene-, [1S-(1a,3aβ,4a,8aβ)]-; (+)-Longifolene; Junipen; Junipene; Kuromatsuen; Kuromatsuene; Longifolen; NSC 150808; d-Longifolene; α-Longifolene | |

Current regulation: /

Clinical data: /

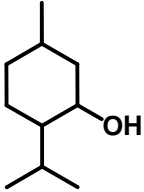
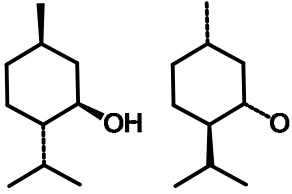
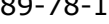
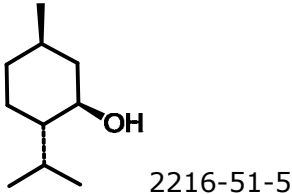
Additional information:

It is a "top 200" substance and classified as R43 (IFRA, pers. comm.2010)

http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details_v2&id=77412

This substance is listed in the Register of Flavouring Substances pursuant to Article 3(1) of Regulation EC No. 2232/96 (28 Oct 1996) that lays down a procedure for flavouring substances used or intended for use in or on foodstuffs. Adopted February 23, 1999.

A RIFM review is available citing one negative human maximisation test (n=25) with 10% pet. (153).

| | |
|---|---|
| MENTHOL |  |
| CAS # 1490-04-6 / 89-78-1 / 2216-51-5 | |
| EC # 216-074-4 / 239-388-3 / 218-690-9 | |
| 5-Methyl-2-(1-methylethyl)-cyclohexanol (1490-04-6) (1R,2S,5R)-rel-5-Methyl-2-(1-methylethyl)-cyclohexanol (89-78-1) (1R,2S,5R)-5-Methyl-2-(1-methylethyl)-cyclohexanol (2216-51-5) | |
| Other names: |  |
| 1490-04-6: Menthol; 1-Methyl-4-isopropyl-3-cyclohexanol; 2-Isopropyl-5-methylcyclohexan-1-ol; 2-Isopropyl-5-methylcyclohexanol; 3-Hydroxy-p-menthane; 5-Methyl-2-(1-methylethyl)cyclohexanol; 5-Methyl-2-isopropylcyclohexanol; Menthyl alcohol; p-Menthan-3-ol |  |
| 89-78-1: (1a,2β,5a)-5-Methyl-2-(1-methylethyl)-cyclohexanol; cis-1,3,trans-1,4-Menthol; dl-Menthol; (1R,2S,5R)-rel-5-Methyl-2-(1-methylethyl)cyclohexanol; (±)-Menthol; DL-Menthol; Fisherman's Friend Lozenges; Hexahydrothymol; Menthacamphor; Menthol; Menthomenthol; NSC 2603; Peppermint camphor; Racementhol; Therapeutic Mineral Ice; Thymomenthol; rac-Menthol |  |

2216-51-5: (1R,2S,5R)-5-Methyl-2-(1-methylethyl)-cyclohexanol; [1R-(1 α ,2 β ,5 α)]-5-Methyl-2-(1-methylethyl)-cyclohexanol; (1R,3R,4S)-(-)-Menthol; (-)-Menthol; (-)-Menthyl alcohol; (-)-trans-p-Methan-cis-3-ol; (1R)-(-)-Menthol; (1R,2S,5R)-(-)-Menthol; (1R,2S,5R)-2-Isopropyl-5-methylcyclohexan-1-ol; (1R,2S,5R)-2-Isopropyl-5-methylcyclohexanol; (R)-(-)-Menthol; 1R-Menthol; L-Menthol; L-Mentholum; Levomenthol; NSC 62788; l(-)-Menthol; l-Menthol

Current regulation: /

Clinical data:

Among 512 patients referred from a dental department for diagnostic work-up of various intraoral symptoms and complaints within 4 years, 10 patients had positive (+ to +++) PT reactions to menthol 5% pet. at D4, mostly reporting dramatic improvement after cessation of use of peppermint-containing oral products (154). In 63 patients positive to the FM I, 1 had a positive PT reaction to menthol, 5% pet., in the Santucci 1987 study (28). The IVDK 2010 study, 1 of 1147 patients tested with 1% menthol in pet. had a weak positive reaction to menthol (7).

A case of contact allergy to "peppermint and menthol" in a transdermal therapeutic system with flurbiprofen for lumbar pain has been described (155). Moreover, a case of rhinitis caused by different menthol-containing products, diagnostically proven by repeatedly positive urticarial reactions after application of 2% menthol in pet. or 5% peppermint oil in pet., has been reported (156). "A case of asthma due to menthol is reported in a 40-year-old woman with no history of asthma or any other allergy. During the last two years, the patient had presented dyspnoea, wheezing and nasal symptoms when exposed to mentholated products such as toothpaste and candies. The aetiology was suggested by the history of exposure and diagnosis was established by skin tests and bronchial challenge with menthol. The patient achieved control of symptoms by avoiding menthol and its derivatives." (157).

Additional information:

Menthol is an ingredient of several essential oils, like peppermint oil, and has been identified as causative allergen in case reports listed above.

Four stereoisomeric forms are known. Natural menthol occurs as L-form (CAS 2216-51-5), trade products are DL-menthol (CAS 1490-04-6). D-form: CAS 89-78-1, racemic: CAS 15356-70-4. Sensitive to light, air and heat (53).

L-menthol and menthol (isomer not specified) are "top 100" substances (IFRA, pers. comm.2010). RIFM reviews are available regarding "menthol" (158), D-menthol (159), L-menthol (160), DL-menthol (161) and menthol, racemic (162). A CIR expert panel review is available (163).

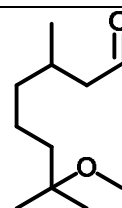
METHOXYCITRONELLAL

CAS # 3613-30-7

EC # 222-784-5

7-Methoxy-3,7-dimethyl-octanal

7-Methoxy-3,7-dimethyloctanal; 7-Methoxy-6,7-dihydrocitronellal; 7-Methoxycitronellal; Methoxycitronellal; Methoxydihydrocitronellal

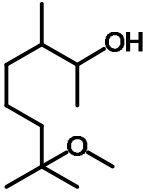


Current regulation: /

Clinical data:

Nakayama et al. found 1974 (after (29)) 12 "strong positive" and 10 "weak positive" reactions to methoxycitronellal (unknown test concentration), with cross-reactions to hydroxycitronellal (proportion not given), in 183 patients.

Additional information: /

| | |
|--|---|
| METHOXYTRIMETHYLHEPTANOL |  |
| CAS # 41890-92-0 | |
| EC # 255-574-7 | |
| 7-Methoxy-3,7-dimethyl-2-octanol | |
| 3,7-Dimethyl-7-methoxy-2-octanol; Dihydromethoxyelgenol; Elesant; Osyrol | |

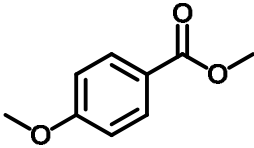
Current regulation: /

Clinical data:

In the Larsen 2002 c study, 0.9% of the patients with known contact allergy to fragrance ingredients had a positive PT reaction to this ingredient not reported as allergen previously (1).

Additional information:

A RIFM review is available (128) citing 1 negative maximisation test (n=27).

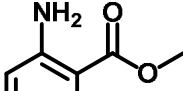
| | |
|---|---|
| METHYL p-ANISATE |  |
| CAS # 121-98-2 | |
| EC # 204-513-2 | |
| Methyl-4-methoxybenzoate | |
| p-Anisic acid, methyl ester; 4-(Methoxycarbonyl)anisole; 4-Methoxybenzoic acid methyl ester; Methyl p-anisate; Methyl p-methoxybenzoate; NSC 7324; p-Methoxybenzoic acid methyl ester | |

Current regulation: /

Clinical data:

In the Malten 1984 study, n=1 (0.5%) of 182 patients displayed a positive PT reaction to methyl anisate 4% pet. (24).

Additional information: /

| | |
|----------------------------|---|
| METHYL ANTHRANILATE |  |
| CAS # 134-20-3 | |

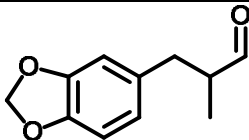
| | |
|--|--|
| EC # 205-132-4 | |
| Methyl 2-aminobenzoate | |
| Anthranilic acid, methyl ester; 2-(Methoxycarbonyl)aniline; 2-Aminobenzoic acid methyl ester; 2-Carbomethoxyaniline; Bird Shield; Grain 96-1; Methyl 2-aminobenzoate; Methyl 6-aminobenzoate; Methyl anthranilate; Methyl o-aminobenzoate; NSC 3109; ReJex-iT; Rejex-iT AP 50; Rejex-iT TP 40; Sunarome UVA; [2-(Methoxycarbonyl)phenyl]amine; o-(Methoxycarbonyl)aniline; o-Aminobenzoic acid methyl ester; o-Carbomethoxyaniline | |

Current regulation: /

Clinical data:

In 91 Israeli patients with a positive or doubtful reaction to FMI or MP methyl anthranilate was tested (conc. not given), with a negative result (164).

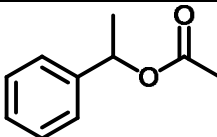
Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|---|--|
| METHYLENEDIOXYPHENYL METHYLPROPANAL |  |
| CAS # 1205-17-0 | |
| EC # 214-881-6 | |
| 3-(1,3-Benzodioxol-5-yl)-2-methylpropanal | |
| Hydrocinnamaldehyde, α -methyl-3,4-(methylenedioxy)-; 2-Methyl-3-(3,4-methylenedioxyphenyl)propanal; 2-Methyl-3-(3,4-methylenedioxyphenyl)propionaldehyde; 3-(3,4-Methylenedioxyphenyl)-2-methylpropanal; Heliobouquet; Heliofresh; Heliogan; Helional; Helipropanal; NSC 22282; Tropional; α -Methyl-1,3-benzodioxole-5-propanal; α -Methyl-3,4-(methylenedioxy)hydrocinnamaldehyde | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

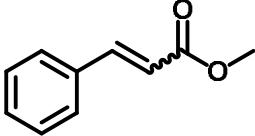
| | |
|--|---|
| METHYLBENZYL ACETATE |  |
| CAS # 93-92-5 | |
| EC # 202-288-5 | |
| 1-Phenylethyl acetate | |
| Benzenemethanol, α -methyl-, acetate ; Benzyl alcohol, α -methyl-, acetate ; (\pm)-Styrallyl acetate; (\pm)- α -Methylbenzyl acetate; (\pm)- α -Phenethyl acetate; 1-Acetoxy-1-phenylethane; 1-Phenylethyl acetate; Gardeniol II; Gardenol; Methyl phenyl carbinyl acetate; Methylphenylcarbinol acetate; NSC 2397; Styrallyl acetate; | |

Styrylallyl acetate; dl-1-Phenylethyl acetate; sec-Phenethyl acetate; sec-Phenylethyl acetate; α -Methylbenzenemethanol acetate; α -Methylbenzyl acetate; α -Methylbenzyl alcohol, acetate; α -Phenethyl acetate; α -Phenylethyl acetate

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|---|---|
| METHYL CINNAMATE |  |
| CAS # 103-26-4 | |
| EC # 203-093-8 | |
| Methyl 3-phenylprop-2-enoate | |
| 3-Phenyl-2-propenoic acid methyl ester; Cinnamic acid, methyl ester; 3-Phenyl-2-propenoic acid methyl ester; 3-Phenylacrylic acid methyl ester; Methyl 3-phenyl-2-propenoate; Methyl 3-phenylacrylate; Methyl 3-phenylpropenoate; Methyl cinnamate; Methyl cinnamylate; NSC 9411; SemaSORB 9815 | |

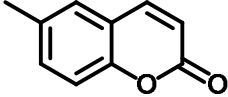
Current regulation: /

Clinical data:

Patch tests with some components of Peru balsam were carried out at 8 worldwide centers in 142 patients who had previously reacted to 25% MP. Reactions to methyl cinnamate (dose and vehicle not reported) were observed in 6 of 142 patients (no further details reported) (165).

Additional information:

A RIFM review is available (166), reviewing, e.g., a number of animal studies with conflicting results. See also under Myroxylon pereirae.

| | |
|--|---|
| 6-METHYL COUMARIN |  |
| CAS # 92-48-8 | |
| EC # 202-158-8 | |
| 6-Methylchromen-2-one | |
| Coumarin, 6-methyl-; 6-MC; 6-Methyl-2H-1-benzopyran-2-one; 6-Methyl-2H-chromen-2-one; 6-Methylbenzopyrone; 6-Methylcoumarin; 6-Methylcoumarinic anhydride; NSC 5870; Toncarine | |

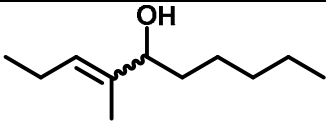
Current regulation: Annex III, part 1, n° 46

Clinical data:

Two of 24 white volunteers developed a photoallergic reaction after single epicutaneous exposure with 5% methyl coumarin in ethanol and UV-A radiation (16 J/cm²). After a photomaximisation test, 6 of 10 subjects developed photocontact allergic reactions

(167). Cardoso et al. report on 2 photoallergic patch test reactions to this substance, which were apparently clinically relevant, in 83 Portuguese patients tested (168). Similar results (2 of 76 patients with positive photopatchtest) were reported from New York (169).

Additional information: /

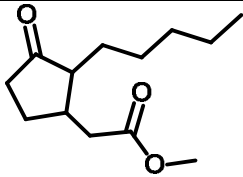
| | |
|------------------------------|---|
| METHYL DECENOL |  |
| CAS # 81782-77-6 | |
| EC # 279-815-0 | |
| 4-Methyl-3-decen-5-ol | |

Current regulation: /

Clinical data: /

Additional information:

A RIFM review is available (170), reporting 1 negative HRIPT (n=50). It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|---|---|
| METHYL DIHYDROJASMONATE |  |
| CAS # 24851-98-7 | |
| EC # 246-495-9 | |
| Methyl 2-(3-oxo-2-pentylcyclopentyl) acetate | |

Cyclopentaneacetic acid, 3-oxo-2-pentyl-, methyl ester; Kharismal; MDJ; Methyl (3-oxo-2-pentylcyclopentyl)acetate; Methyl 3-oxo-2-pentylcyclopentane ethanoate; Hedione

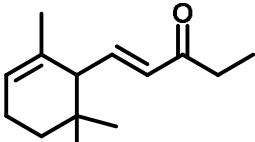
Current regulation: /

Clinical data:

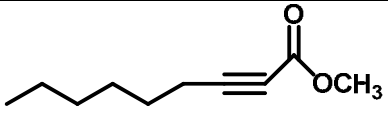
In the Frosch 2002 b study, 3 of 1606 consecutive patients (0.2%) showed positive reactions to hedione (5% pet.) (17). In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and 5% hedione in pet., tested in 100 consecutive patients in Belfast, were observed (15).

Additional information:

It is a "top 100" substance (IFRA, pers. comm.2010). An older RIFM review exists (128) citing 1 negative human maximisation test (n=25).

| | |
|---|---|
| METHYL IONONE (mixture of isomers) |  |
| CAS # 1335-46-2 | |
| EC # 215-635-0 | |

| | |
|---|--|
| 1-(2,6,6-Trimethyl-1-cyclohex-2-enyl)pent-1-en-3-one | |
| 6-Methylionone | |
| Current regulation: / | |
| Clinical data: | |
| See METHYLIONANTHEME for one clinical case report. Regarding methyl ionone gamma, the Frosch 1995 dose-finding pilot study found no positive reaction to 1% and 5% of this substance in pet., tested in 100 consecutive patients in Belfast (15). | |
| Additional information: | |
| It is a "top 100" substance (IFRA, pers. comm.2010). A RIFM review is available (171). | |

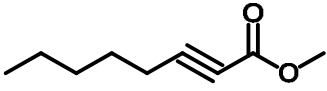
| | |
|--------------------------------|---|
| METHYL OCTINE CARBONATE |  |
| CAS # 111-80-8 | |
| EC # | |
| Methyl 2-octynoate | |
| Methyl 2-Nonynoate, MOC | |

Current regulation: Annex III, part 1, n°173

Clinical data:

English and Rycroft reported a case of a 19-year-old laboratory technician working in the fragrance industry, who developed hand dermatitis after contact with methyl heptene and methyl octane carbonates; patch testing was strongly positive to both compounds at 1% in MEK (172).

Additional information: /

| | |
|---|---|
| METHYL 2-OCTYNOATE |  |
| CAS # 111-12-6 | |
| EC # 203-836-6 | |
| Methyl oct-2-ynoate | |
| M2O; Methyl heptin carbonate; Folione; Methyl hept-1-yne-1-carboxylate; Methyl pentylacetylenecarboxylate; NSC 72098; Vert de violette artificiel | |

Current regulation: Annex III, part 1, n° 89

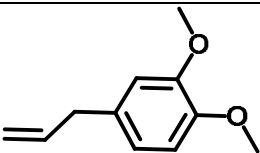
Clinical data:

In the "background information" section of the 1999 opinion, methyl 2-octynoate is classified as "less frequently reported allergen"; with only single cases of reported contact allergy, but the observation of this compound being a strong sensitizer according to IFRA (33), as also reported by Hostynek and Maibach (173)

Since the last SCCNFP-opinion of 1999, the IVDK 2007 study yielded 0.3% (95% CI: 0.1 – 0.49%) positive reactions in 2401 consecutively PTed patients (1% pet.) (4). The IVDK 2010 study, n=1 weak positive reaction was observed in 988 patients tested with the compound (7). In the Groningen 2009 study, n=1, i.e. 0.3% (95% CI: 0.01 – 1.7%)

had positive reactions to this allergen, tested at only 2% pet. (6). In a previous case report of a fragrance laboratory assistant with work-related ACD both methyl heptin and methyl octin carbonate had been found sensitizers – probably due to their very similar chemical structure (172). In a recent bi-centric study with 350 eczema patients who were consecutively tested with 1% and 2% M20 in pet.; 0.8% positive reactions were observed. However, in 3 additional cases active sensitization, with first reactions appearing 2 to 4 weeks after the patch test, and prompt reactions in the 2 cases repeat-patch tested, was observed (174).

Additional information: /

| | |
|--|---|
| METHYL EUGENOL |  |
| CAS # 93-15-2 | |
| EC # 202-223-0 | |
| 1,2-Dimethoxy-4-(prop-2-enyl)benzene | |
| 4-Allylveratrole; Eugenyl methyl ether extra; 1,2-Dimethoxy-4-allylbenzene; 1,3,4-Eugenol methyl ether; 1-(3,4-Dimethoxyphenyl)-2-propene; 1-Allyl-3,4-dimethoxybenzene; 3,4-Dimethoxy-1-(2-propenyl)benzene; 3,4-Dimethoxyallylbenzene; 3-(3,4-Dimethoxyphenyl)propene; 4-Allyl-1,2-dimethoxybenzene; Benzene, 4-allyl-1,2-dimethoxy-; Chavibetol methyl ether; Ent 21040; Eugenol methyl ether; Eugenyl methyl ether; Methyl eugenol ether; Methyl eugenyl ether; Methylchavibetol; NSC 209528; NSC 8900; O-Methyleugenol; Veratrole methyl ether; Veratrole, 4-allyl- | |

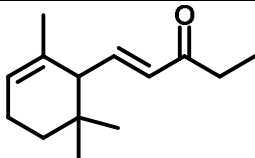
Current regulation: Annex II, 451

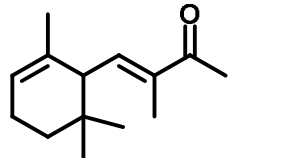
Clinical data:

In a previous study by Larsen et al (2002 c), 1.8% of patients with contact allergy to fragrance ingredients reacted positively to this compound (1).

Additional information:

Quote from the SCCS-opinion [0373/00](#): "Methyleugenol should not be intentionally added as a cosmetic ingredient. However, when fragrance compounds containing methyleugenol naturally present in essential oils are used as components in cosmetic products, the highest concentration of methyleugenol in the finished products must not exceed 0.01 % in fine fragrance, 0.004 % in eau de toilette, 0.002 % in a fragrance cream, 0.0002 % in other leave-on products and in oral hygiene products, and 0.001% in rinse-off products." (The reason is genotoxicity and carcinogenicity).

| | |
|---|---|
| METHYLIONANTHEME |  |
| CAS # 55599-63-8 | |
| EC # | |
| (1E)-2-Methyl-1-(2,6,6-trimethyl-2-cyclohexen-1-yl)-1-penten-3-one mixt. with (3E)-3-methyl-4-(2,6,6-trimethyl-2-cyclohexen-1-yl)-3-buten-2-one | |

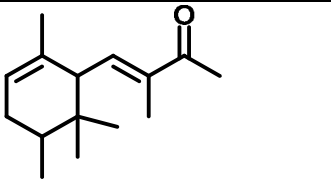
| | |
|--|---|
| 8-Methyl- α -ionone-10-methyl- α -ionone mixt.; Iralia Mixture |  |
|--|---|

Current regulation: ...

Clinical data:
One case of ACD has been reported, caused by an E.d.C. (175).

Additional information:

Patented by GIVAUDAN SA 1933, is composed of isomeric n-methylionones and iso-methylionones. Methylionone has CAS # 1335-94-0 (not in CosIng) and 1335-46-2 (METHYL α -IONONE ISOMERS); other names: Methyl- α -cyclocitrilydenacetone; Iralia; Isoaldeine (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.detail&id=41456>, last accessed 2010-07-14).

| | |
|---|--|
| 5-METHYL-α-IONONE |  |
| CAS # 79-69-6 | |
| EC # 201-219-6 | |
| 4-(2,5,6,6-Tetramethyl-2-cyclohexen-1-yl)-3-buten-2-one | |
| Methyl- α -Ionone; 6-Methyl- α -ionone; α -Irone | |

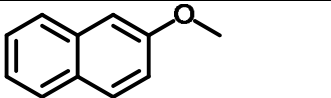
Current regulation: /

Clinical data:

In the Frosch 2002 b study, 5 of 1606 consecutive patients (0.3%) showed positive reactions to α -irone (10% pet.) (17).

Additional information:

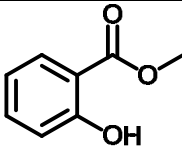
A RIFM review is available (176), citing a (negative) human maximisation test and the study results quoted.

| | |
|---|---|
| METHYL beta-NAPHTHYL ETHER |  |
| CAS # 93-04-9 | |
| EC # 202-213-6 | |
| 2-Methoxynaphthalene | |
| beta-Naphthyl methyl ether; methyl 2-naphthyl ether; Nerolin (old); NSC 4171; Yara yara; β -Methoxynaphthalene; β -Naphthol methyl ether; β -Naphthyl methyl ether; 2-Methoxynaphthalene; Methyl β -naphthyl ether; 2-Naphthol methyl ether; 2-Naphthyl methyl ether; 6-Methoxy-2-naphthalene | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|--|---|
| METHYL SALICYLATE |  |
| CAS # 119-36-8 | |
| EC # 204-317-7 | |
| Methyl 2-hydroxybenzoate | |
| Other names: Salicylic acid, methyl ester; 2-(Methoxycarbonyl)phenol; 2-Carbomethoxyphenol; 2-Hydroxybenzoic acid methyl ester; Analgit; Anthrapole ND; Ben Gay; Exagien; Flucarmit; Methyl ester of 2-hydroxy benzoic acid; Methyl o-hydroxybenzoate; Methyl salicylate; NSC 8204; Wintergreen oil; o-Hydroxybenzoic acid methyl ester; "Oil of wintergreen" | |

Current regulation: /

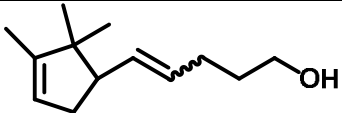
Clinical data:

The deGroot 2000 study yielded 7 positive reactions to methyl salicylate (2% pet.) in 1825 consecutive patients (12).

A case of ACD following the application of a compress bandage containing methyl salicylate has been reported, using 2% "o.o." as PT concentration; the dose per area of methyl salicylate in the occlusive bandage was not reported (177). A similar case was reported in 1977, positive to 2% methyl salicylate in olive oil, with elicitation of pruritus and erythema after oral ingestion of acetyl salicylic acid (178).

Additional information:

A RIFM review is available (179) providing an overview on 3 human sensitisation experiments (e.g., the HRIPT) which were all negative, and clinical data. In a number of older PT studies, positive test results were seen in 6 of 4600, 3 of 183, 3 of 241, 17 of 585, 1 of 70, all employing a test concentration of 2%, usually in pet., according to above review. Methyl salicylate may occur in topical analgesic (OTC) medications, in Germany, for instance, in "Camphopin® Salbe" („Rote Liste 2010").

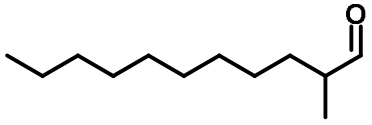
| | |
|---|---|
| 3-METHYL-5-(2,2,3-TRIMETHYL-3-CYCLOPENTENYL)PENT-4-EN-2-OL |  |
| CAS # 67801-20-1 | |
| EC # 267-140-4 | |
| 3-Methyl-5-(2,2,3-trimethyl-1-cyclopent-3-enyl)pent-4-en-2-ol | |
| 3-Methyl-5-(2,2,3-trimethyl-3-cyclopenten-1-yl)-4-penten-2-ol; 3-Methyl-5-(2,2,3-trimethylcyclopent-3-enyl)pent-4-en-2-ol; Ebanol | |

Current regulation: /

Clinical data:

In the Larsen 2001 study, 1 of 178 patients with known contact allergy to fragrance ingredients exhibited a positive PT reaction to "MTCP", tested 5% pet. (19). In the An 2005 study, 12 of 422 consecutive patients, i.e., 2.8%, had positive reactions to "ebanol", tested at 5% (13).

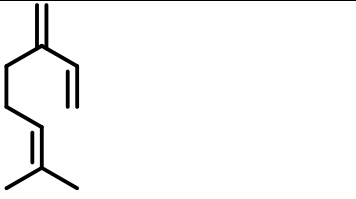
Additional information: /

| | |
|---|---|
| METHYLUNDECANAL |  |
| CAS # 110-41-8 | |
| EC # 203-765-0 | |
| 2-Methylundecanal | |
| Aldehyde c-12 mna; undecenal, 2-methyl-; 2-Methyl-1-undecanal; Aldehyde M.N.A.; Methyl n-nonyl acetaldehyde; Methylnonylacetaldehyde; NSC 46127 | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|--|---|
| MYRCENE |  |
| CAS # 123-35-3 | |
| EC # 204-622-5 | |
| 7-Methyl-3-methylideneocta-1,6-diene | |
| 2-Methyl-6-methylene-2,7-octadiene; 7-Methyl-3-methylene-1,6-octadiene; NSC 406264; β -Geraniolene; β -Myrcene | |

Current regulation: /

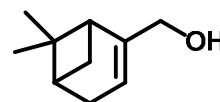
Clinical data:

In a clinical study in 6 European centres, including 1511 consecutive patients, 1 patient had a positive reaction to oxidized myrcene (65).

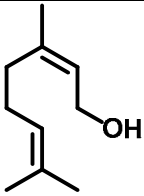
Additional information:

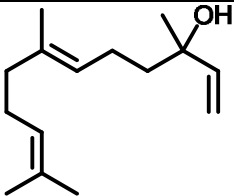
Myrcene autoxidizes spontaneously and rapidly at air exposure. In experimental studies on beta-myrcene an EC3 value of 4.3% was seen for a sample air-exposed 10 weeks (Sköld M. Contact allergy to autoxidized fragrance terpenes (180).

| |
|-----------------|
| MYRTENOL |
| CAS # 515-00-4 |
| EC # 208-193-5 |

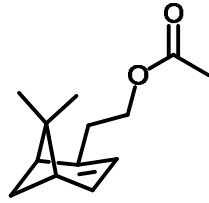


| |
|--|
| (7,7-Dimethyl-4-bicyclo[3.1.1]hept-3-enyl)methanol |
| (-)-Pin-2-ene-10-ol; 2-Pinen-10-ol; (6,6-Dimethylbicyclo[3.1.1]hept-2-en-2-yl)methanol; (±)-Myrtenol; 6,6-Dimethyl-2-(hydroxymethyl)bicyclo[3.1.1]hept-2-ene; NSC 408846; α-Pinene-10-ol |
| Current regulation: / |
| Clinical data: / |
| Additional information: A RIFM review exists (181), citing 2 of 3 HRIPT studies with 1 case of sensitisation to myrtenol each. |

| | |
|---|---|
| NEROL |  |
| CAS # 106-25-2 | |
| EC # 203-378-7 | |
| (2Z)-3,7-Dimethylocta-2,6-dien-1-ol | |
| 2,6-Octadien-1-ol, 3,7-dimethyl-, (Z)-; (Z)-3,7-Dimethyl-2,6-octadien-1-ol; (Z)-Geraniol; (Z)-Nerol; 2-cis-3,7-Dimethyl-2,6-octadien-1-ol; 3,7-Dimethyl-cis-2,6-octadien-1-ol; Nerol 900; Neryl alcohol; cis-3,7-Dimethyl-2,6-octadien-1-ol; cis-Geraniol; β-Nerol; cis-geraniol - i.e., isomeric to geraniol | |
| Current regulation: / | |
| Clinical data: In the Larsen 2002 c study, 6.0% of the fragrance sensitive patients reacted positively to 5% in pet. (1). | |
| Additional information: A RIFM review is available (182) citing (negative) human sensitisation experiments, an older study from Japan and the Larsen 2002 c study (see above). Regarding autoxidation studies – see geraniol. | |

| | |
|--|---|
| Nerolidol (isomer not specified) |  |
| CAS # 7212-44-4 | |
| EC # 230-597-5 | |
| 3,7,11-Trimethyl-1,6,10-odecatrien-3-ol | |
| Nerolidol; (±)-Nerolidol; FCI 119b; Nerodilol | |
| Current regulation: / | |
| Clinical data: / | |
| Additional information: | |

RIFM review is available (183) citing the occurrence of "3 positive reactions in 2273 patients". Another RIFM review is available on cis-nerolidol (184), mentioning that no data on this compound are available.

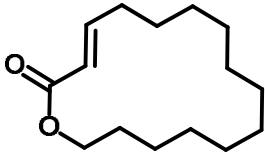
| | |
|---|---|
| NOPYL ACETATE |  |
| CAS # 128-51-8 | |
| EC # 204-891-9 | |
| 2-(7,7-Dimethyl-4-bicyclo[3.1.1]hept-3-enyl)ethyl acetate | |
| 2-Norpinene-2-ethanol, 6,6-Dimethyl-, acetate; Bicyclo[3.1.1]hept-2-ene-2-ethanol, 6,6-dimethyl-, acetate; 2-(6,6-Dimethylbicyclo[3.1.1]hept-2-en-2-yl)ethyl acetate; 7,7-Dimethylbicyclo[3.1.1]hept-2-ene-2-ethanol acetate; Citroviol; NSC 1286; NSC 404963; Nopol acetate; Nopyl acetate | |

Current regulation: /

Clinical data:

The DeGroot 1985 study identified 2 (1.1%) positive reactions among 179 patients using a 25% PT preparation of this compound – reactions may have at least partly been due to an "excited back syndrome" and thus a limited evidence (25).

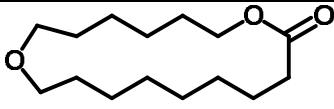
Additional information: /

| | |
|--|---|
| OXACYCLOHEXADECENONE |  |
| CAS # 34902-57-3 | |
| EC # 609-040-9 | |
| (3E)-Oxacyclohexadec-3-en-2-one | |
| Globalide; Oxacyclohexadecen-2-one | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|--|---|
| OXALIDE |  |
| CAS # 1725-01-5 | |
| EC # 217-033-3 | |
| 1,8-Dioxacycloheptadecan-9-one | |
| Nonanoic acid, 9-[(6-hydroxyhexyl)oxy]-, o-lactone; 10-Oxa-16-hexadecanolide; Oxalide; Oxalide T | |

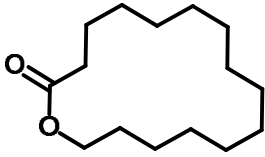
Current regulation: /

Clinical data:

In the Larsen 2001 study, none of 178 patients with known contact allergy to fragrance ingredients exhibited a positive PT reaction to "10-oxahexadecanolide", tested 5% pet. (19).

Additional information:

A RIFM review is available (128), citing a negative maximisation test (n=29).

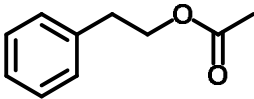
| | |
|---|---|
| PENTADECALACTONE |  |
| CAS # 106-02-5 | |
| EC # 203-354-6 | |
| 1-Oxacyclohexadecan-2-one | |
| Pentadecanoic acid, 15-hydroxy-, ξ -lactone; 1,15-Pentadecanolide; 15-Hydroxypentadecanoic acid lactone; 15-Pentadecanolide; 15-Pentadodecanolactone; 2-Pentadecalone; CPE 215; Cyclopentadecanolide; Exaltolide; Macrolide Supra; Muskalactone; NSC 36763; Pentadecalactone; Pentadecanolactone; Pentadecanolide; Pentalide; Thibetolide; cpd Supra; ω -Pentadecalactone; angelica lactone; hexaltolide | |

Current regulation: /

Clinical data: /

Additional information:

It is a "top 100" substance (IFRA, pers. comm.2010). The substance has been used for clinical olfactory testing in the 60ies under the name of exaltolide.

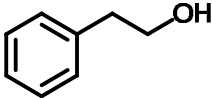
| | |
|---|---|
| PHENETHYL ACETATE |  |
| CAS # 103-45-7 | |
| EC # 203-113-5 | |
| 2-Phenylethyl acetate | |
| Acetic acid, phenethyl ester ; Phenethyl alcohol, acetate; 2-Phenethyl acetate; 2-Phenylethyl acetate; Benzylcarbiny acetate; NSC 71927; Phenethyl acetate; Phenylethyl ethanoate; β -Phenethyl acetate; β -Phenylethanol acetate; β -Phenylethyl acetate | |

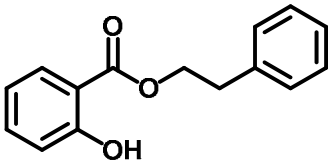
Current regulation: /

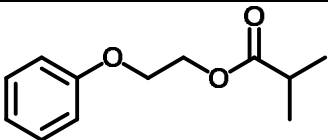
Clinical data: /

Additional information:

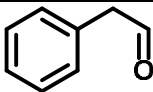
It is a "top 100" substance (IFRA, pers. comm.2010). Exposure via plants (*Tanacetum parthenium*) is possible (185).

| | |
|---|---|
| PHENETHYL ALCOHOL |  |
| CAS # 60-12-8 | |
| EC # 200-456-2 | |
| 2-Phenylethanol | |
| Phenethyl alcohol; (2-Hydroxyethyl)benzene; 2-Phenethanol; 2-Phenethyl alcohol; 2-Phenyl-1-ethanol; 2-Phenylethyl alcohol; Benzyl carbinol; Ethanol, 2-phenyl-; NSC 406252; PEA; Phenethanol; Phenethylol; Phenylethanol; Phenylethyl alcohol; β -(Hydroxyethyl)benzene; β -PEA; β -Phenethanol; β -Phenethyl alcohol; β -Phenethylol; β -Phenylethanol; β -Phenylethyl alcohol | |
| Current regulation: / | |
| Clinical data: | |
| The DeGroot 1985 study identified 1 (0.6%) positive reactions among 179 patients using a 25% PT preparation of phenylethyl alcohol (25). In the Frosch 1995 dose-finding pilot study, no positive reaction to this compound, tested 1% pet. in 100 consecutive patients in Odense, DK, was observed (15). | |
| Additional information: It is a "top 100" substance (IFRA, pers. comm.2010). | |

| | |
|--|---|
| PHENETHYL SALICYLATE |  |
| CAS # 87-22-9 | |
| EC # 201-732-5 | |
| 2-Phenylethyl 2-hydroxybenzoate | |
| Salicylic acid, phenethyl ester; 2-Phenylethyl salicylate; Benzylcarbinyl salicylate; NSC 72035; Phenethyl salicylate | |
| Current regulation: / | |
| Clinical data: / | |
| Additional information: | |
| A RIFM review exists (186), quoting a negative human maximisation test and a number of animal experiments, including cross-sensitisation experiments with benzyl salicylate. One LLNA study is reported yielding an EC3 value of 2.1%. | |

| | |
|--|---|
| PHENOXYETHYL ISOBUTYRATE |  |
| CAS # 103-60-6 | |
| EC # 203-127-1 | |
| 2-Phenoxyethyl 2-methylpropanoate | |
| Isobutyric acid, 2-phenoxyethyl ester; Ethanol, 2-phenoxy-, isobutyrate; 2-Phenoxyethyl isobutyrate; NSC 227210; NSC | |

| | |
|---|--|
| 406209; Phenoxyethyl isobutyrate; β -Phenoxyethyl isobutyrate | |
| Current regulation: / | |
| Clinical data: / | |
| Additional information: It is a "top 100" substance (IFRA, pers. comm.2010). | |

| | |
|---|---|
| PHENYLACETALDEHYDE |  |
| CAS # 122-78-1 | |
| EC # 204-574-5 | |
| 2-Phenylacetaldehyde | |
| Benzylcarboxaldehyde; Hyacinthin; NSC 406309; Phenacetaldehyde; Phenylacetaldehyde; Phenylacetic aldehyde; Phenylethanal; α -Phenylacetaldehyde; α -Tolualdehyde; α -Toluic aldehyde | |

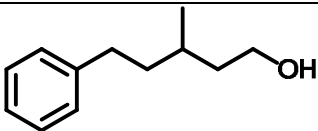
Current regulation: /

Clinical data:

In the Malten 1984 study, 1.1% of 182 patients displayed a positive PT reaction to phenylacetaldehyde 2% pet. (24). In a case report, Sanchez-Politta et al. describe a 26-year-old worker in a perfume factory, who suffered from a spill of pure phenylacetaldehyde and became sensitised, as proven by positive patch tests with 0.5%, 1% and 2% (10 healthy controls negative) (187).

Additional information:

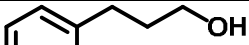
SCCS opinion: [1153/08 - Opinion on "Dermal Sensitization Quantitative Risk Assessment" \(QRA: Citral, farnesol and phenylacetaldehyde\)](#)

| | |
|--|---|
| PHENYLISOHEXANOL |  |
| CAS # 55066-48-3 | |
| EC # 259-461-3 | |
| 3-Methyl-5-phenylpentan-1-ol | |
| 3-Methyl-5-phenyl-1-pentanol; 3-Methyl-5-phenylpentanol; 5-Phenyl-3-methylpentanol; Mefrosol; Phenoxanol | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|-----------------------|---|
| PHENYLPROPANOL |  |
|-----------------------|---|

Opinion on fragrance allergens in cosmetic products

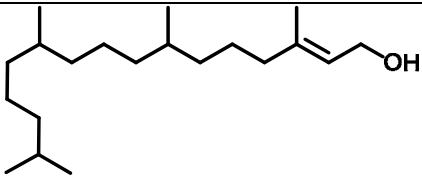
| |
|---|
| CAS # 122-97-4 |
| EC # 204-587-6 |
| 3-Phenylpropan-1-ol |
| (3-Hydroxypropyl)benzene; 1-Hydroxy-3-phenylpropane; 3-Benzenepropanol; 3-Hydroxy-1-phenylpropane; 3-Phenyl-1-propanol; 3-Phenyl-n-propanol; 3-Phenylpropanol; 3-Phenylpropyl alcohol; Dihydrocinnamyl alcohol; Hydrocinnamic alcohol; Hydrocinnamyl alcohol; NSC 16942; γ -Phenylpropanol; γ -Phenylpropyl alcohol; Phenethyl Carbinol |

Current regulation: /

Clinical data:

The Larsen 2002 c study yielded 0.9% positive reactions in 218 patients with contact allergy to fragrance ingredients (1).

Additional information: ...

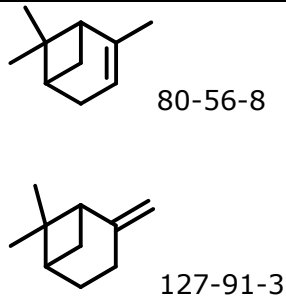
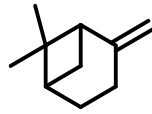
| | |
|--|---|
| PHYTOL |  |
| CAS # 150-86-7 | |
| EC # 205-776-6 | |
| (E,7R,11R)-3,7,11,15-tetramethylhexadec-2-en-1-ol | |
| Phytol; (7R,11R,2E)-Phytol; (E)-Phytol; (E,R,R)-Phytol; 3,7,11,15-Tetramethylhexadec-2-en-1-ol; trans-Phytol | |

Current regulation: /

Clinical data: /

Additional information:

Phytol is a main constituent of Jasmin abs. with 7.4% reported content (17). In a human maximization study involving 25 subjects, there was one case of contact sensitization to 10% phytol (6900 $\mu\text{g}/\text{cm}^2$), applied in petrolatum, as reported in a RIFM review (188).

| | |
|---|---|
| <i>alpha-PINENE and beta-PINENE</i> |  |
| CAS # 80-56-8 (alpha-Pinene); CAS # 127-91-3 (beta-Pinene) | |
| EC # 201-291-9 (alpha-Pinene; according to CAS service: 219-445-9); EC # 204-872-5 (beta-Pinene; according to CAS service: 245-424-9) | |
| 2,6,6-Trimethylbicyclo[3.1.1]hept-2-ene (80-56-8) |  |
| 6,6-Dimethyl- 2-methylenebicyclo[3.1.1]heptane (127-91-3) | |

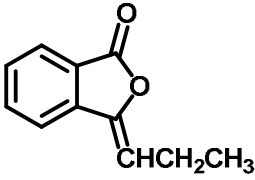
| | |
|---|--|
| <p>80-56-8: 2-Pinene; (\pm)-2-Pinene; (\pm)-α-Pinene; Acintene A; NSC 7727; PC 500; PC 500 (terpene); Sylvapine A; α-Pinene</p> <p>127-91-3: 2(10)-Pinene ; (\pm)-2(10)-Pinene; (\pm)-6,6-Dimethyl-2-methylenebicyclo[3.1.1]heptane; (\pm)-β-Pinene; 6,6-Dimethyl-2-methylenebicyclo[3.1.1]heptane; NSC 21447; NSC 406265; NSC 59190; Nopinene; Nopinene; PC 600; PC 600 (pesticide); Pseudopinene; Pseudopinene; Terebenthene; β-Pinene</p> | |
| <p>Current regulation: Annex III, part 1, n° 130 (Peroxide value less than 10 mmoles/L in substance)</p> | |

Clinical data:

In 63 patients positive to the FM I, 2 had a positive PT reaction to beta-pinene (and none to alpha-pinene 5% pet.), 1% pet., in the Santucci 1987 study (28). A clinical series from Portugal, addressing contact allergy to oil of turpentine diagnosed in 30 patients, used a series with pure terpenes. A total of 17 of 30 patients reacted positively to alpha-pinene, and 2 to beta-pinene (189). In a series of 24 patients with occupational contact dermatitis from the pottery industry, Lear et al. found 14 to be sensitised to "Indonesian oil of turpentine" and 8 to alpha-pinene (190).

A case report from Zacher and Ippen on 2 patients with allergic contact dermatitis due to bergamot oil (191) describes positive patch test reactions to alpha-pinene and beta-pinene in one, a worker in a perfume factory.

Additional information: /

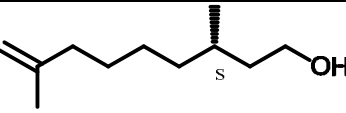
| | |
|--|---|
| PROPYLIDENE PHTHALIDE |  |
| CAS # 17369-59-4 | |
| EC # 241-402-8 | |
| 3-Propylidene-2-benzofuran-1-one | |
| 3-Propylidene-1(3H)-isobenzofuranone; Propylidenephthalide; Celeriax; Propylidenephthalide | 3- |

Current regulation: Annex III, part 1, n° 175

Clinical data:

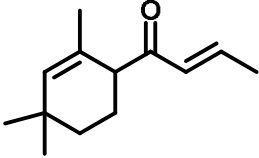
In the Malten 1984 study, 2.6% of 182 patients displayed a positive PT reaction to ethyl acrylate 1% pet. (24). In this paper, "3/25" positive results in human maximisation tests are listed.

Additional information: /

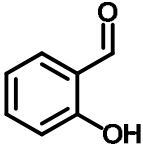
| | |
|---|---|
| RHODINOL |  |
| CAS # 6812-78-8 | |
| EC # 229-887-4 | |
| (3S)-3,7-Dimethyloct-7-en-1-ol | |
| Rhodinol; (-)-Rhodinol; α -citronellol; (-)- α -Citronellol; (S)- | |

Opinion on fragrance allergens in cosmetic products

| | |
|--|--|
| α -Citronellol | |
| Current regulation: / | |
| Clinical data: / (see below) | |
| Additional information: | |
| <p>A RIFM review exists citing a positive HRIPT with several cases of sensitisation, 5 of these proven upon re-challenge, and a negative human maximisation test (192). In a previous RIFM review (128), a Japanese clinical study (source not accessible) is cited: "In patch tests using cosmetics ingredients and fragrance materials on patients with eczema and dermatitis, 5% rhodinol (vehicle not specified) produced one sensitization reaction in 202 patients (Itoh et al., 1988¹⁴)"</p> | |

| | |
|---|---|
| trans-ROSE KETONE-5 |  |
| CAS # 39872-57-6 | |
| EC # 254-663-8 | |
| (2E)-1-(2,4,4-Trimethylcyclohex-2-en-1-yl)but-2-en-1-one | |
| alpha-Isodamascone; trans-2,4,4-Trimethyl-1-crotonyl-2-cyclohexene; (E)-1-(2,4,4-Trimethyl-2-cyclohexen-1-yl)-2-buten-1-one | |
| Current regulation: Annex III, part 1, n° 159 (max. conc. 0.02%) | |

| | |
|--|--|
| Clinical data: / | |
| Additional information: | |
| <p>A RIFM review is available (193) quoting 2 HRIPT studies: one with 0.2% concentration in DEP in 103 volunteers, and negative result, one with 2% concentration, sensitising 2 of 22 volunteers.</p> | |

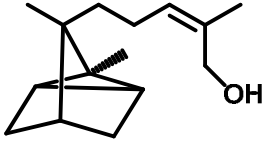
| | |
|---|---|
| SALICYLALDEHYDE |  |
| CAS # 90-02-8 | |
| EC # 201-961-0 | |
| 2-Hydroxybenzaldehyde | |
| Salicylaldehyde; 2-Formylphenol; NSC 112278; NSC 49178; NSC 83559; NSC 83560; NSC 83561; NSC 83562; NSC 97202; Salicylal; Salicylic aldehyde; o-Formylphenol; o-Hydroxybenzaldehyde | |
| Current regulation: / | |

Clinical data:

¹⁴ Itoh M., Hosono K., Kantoh H., Kinoshita M., Yamada K., Kurosaka R. and Nishimura M. (1988) Patch test results with cosmetic ingredients conducted between 1978-1986. *Nippon Koshohen Kagakkaishi* 12 (1), 27-41.

In a series of 40 of 744 consecutive patients PTed with an extended fragrance series (Sheffield 1999), 1 positive reaction to salicylaldehyde was observed (3). In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded n=1 (0.1%) positive reaction to salicylaldehyde 2% pet. (22). The IVDK 2010 study, 0.48% (95% CI: 0.18 – 0.79%; percentages standardised for age and sex) of 2729 patients PTed reacted to the compound (7). An earlier study by Bruze and Zimerson points to possible cross-reactivity between salicylaldehyde and “simple methylol phenols” occurring in synthetic resins based on phenol and formaldehyde (194). Among 24 patients sensitised to resorcinol by application of a wart remover, 2 positive reactions to salicylaldehyde were observed (195).

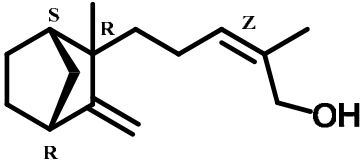
Additional information: Along with other derivates of salicylic acid, salicylaldehyde is found in the bark of several trees, such as willow or aspen, and can cause allergic contact dermatitis by this exposure (196).

| | |
|--|---|
| <i>alpha-SANTALOL</i> |  |
| CAS # 115-71-9 | |
| EC # 204-102-8 | |
| (R Z)- 5-(2,3-dimethyltricyclo[2.2.1.0^{2,6}]hept-3-yl)-2-methylPent-2-en-1-ol | |
| 2-Penten-1-ol, 5-(2,3-dimethyltricyclo[2.2.1.0 ^{2,6}]hept-3-yl)-2-methyl-, [R(Z)]-; 2-Penten-1-ol, 5-(2,3-dimethyltricyclo[2.2.1.0 ^{2,6}]hept-3-yl)-2-methyl-, stereoisomer; α -Santalol; Tricyclo[2.2.1.0 ^{2,6}]heptane, 2-penten-1-ol deriv.; (+)-(Z)- α -Santalol; (+)- α -Santalol; (Z)- α -Santalol; Sandal; Santalol a; cis- α -Santalol; d- α -Santalol | |

Current regulation: /

Clinical data: / (see beta-santalol)

Additional information:
Following a precautionary principle, both isoforms – often not differentiated in reports – are considered as one and considered as established contact allergen in humans.

| | |
|--|--|
| <i>beta-SANTALOL</i> |  |
| CAS # 77-42-9 | |
| EC # 201-027-2 | |
| (2Z)-2-Methyl-5-[(1S,2R,4R)-2-methyl-3-methylenebicyclo[2.2.1]hept-2-yl]pent-2-en-1-ol | |
| 2-Methyl-5-(2-methyl-3-methylene-2-norbornyl)-2-penten-1-ol; [1S-[1 α ,2 α (Z),4 α]]-2-Methyl-5-(2-methyl-3-methylenebicyclo[2.2.1]hept-2-yl)-2-penten-1-ol; β -Santalol; (-)-(Z)- β -Santalol; (-)- β -Santalol; Santalol b; cis- β -Santalol | |

Current regulation: /

Clinical

data:

A RIFM review is available for alpha-santalol (197) and on "santalol" (CAS # 11031-45-1 (198)). The former review cites a Japanese study: "Between April 1979 and August 1990, a total of 3123 male and female patients were patch tested to 2% santalol (.alpha. or .beta. not specified) in petrolatum. Reactions were observed in 47/3123 (1.5%) of the patients. The incidence of positive reactions from 1979 to 1990 was 1.5%. The rate of reactions observed was higher during the earlier period of the patch testing than the later stage (Utsumi et al., 1992)¹⁵." In another Japanese study cited by the RIFM review "... patch tests were conducted with 0.05–0.5% santalol (specified as santalol 1) in a base cream or in 99% ethanol. Patches consisted of a piece of 1 cm² lint with a 2 cm² cellophane disc placed on the lint and then covered with a 4 cm² plaster. Patches were applied to the back, the forearm, and the inside of the upper arm for 24–48 h. Reactions were observed in 15 patients and questionable reactions were observed in 10 patients out of the total 427 participating. A second sample of santalol (specified as santalol 2) was tested on 214 patients. Reactions were observed in three patients and questionable reactions were observed in six patients (Takenaka et al., 1986)¹⁶." Moreover, "The Mid-Japan Contact Dermatitis Research group (MJDCRG) conducted a 6-year (1976–1981) patch test study on facial dermatoses patients with various fragrance materials. During the year 1979, a total of 327 patients were tested with a mixture of .alpha. and .beta. santalol at concentrations of 10%, 2%, and 1% in white petrolatum. Reactions were observed in 1.5%, 0.6% and 0.6% of the 327 patients tested at concentrations 10%, 2%, and 1%, respectively (MJDCRG, 1984)¹⁷."

The Goossens 1997 study found 5 of 111 patients positive to "santalol 10% pet." (isoform not specified) – all sensitised to other fragrance allergens as well (23). In the Larsen 2001 study, patch testing with "2-methyl-5-(2,3-dimethyl tricyclo[2.2.1.0(2,6)]hept-3-yl-2 pentenol(.alpha.-form) and 2-methyl-5-(2-methyl-3-methylenebicyclo[2.2.1]hept-3-yl-2-penten-1-ol(beta-form) 5% pet." (no CAS numbers given) yielded a total of 2 positive reactions among the 178 patients with known contact allergy to fragrance ingredients (19).

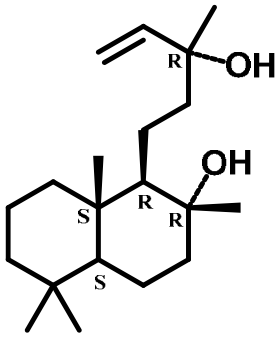
Additional information: "There is no one CAS number for the mixture. The alpha form has a CAS No. 115-71-9 and the beta form is 37172-32-0 (this # is trans-.beta.-santalol). There was no reported use of these materials in the last two IFRA Surveys (8 years total)" (A.M. Api, pers. comm., 2010).

Following a precautionary principle, both isoforms – often not differentiated in reports – are considered as one and considered as established contact allergen in humans

¹⁵ Utsumi, M., Sugai, T., Shoji, A., Watanabe, K., Asoh, S., Hashimoto, Y., 1992. Incidence of positive reactions to sandalwood oil and its related fragrance materials in patch tests and a case of contact allergy to natural and synthetic sandalwood oil in a museum worker. *Skin Research* 34, 209–213

¹⁶ Takenaka, T., Hasegawa, E., Takenaka, U., Saito, F., Odaka, T., 1986. Fundamental studies of safe compound perfumes for cosmetics Part 1. The primary irritation of compound materials to the skin. *Unknown Source*, 313–329.

¹⁷ Mid-Japan Contact Dermatitis Research Group, 1984. Determination of suitable concentrations for patch testing of various fragrance materials. A summary of group study conducted over a 6-year period. *Journal of Dermatology*, 11(1), 31–35.

| | |
|--|---|
| SCLAREOL |  |
| CAS # 515-03-7 | |
| EC # 208-194-0 | |
| (1R,2R,8aS)-1-[(3R)-3-Hydroxy-3-methylpent-4-enyl]-2,5,5,8a-tetramethyl-3,4,4a,6,7,8-hexahydro-1H-naphthalen-2-ol | |
| (α R,1R,2R,4aS,8aS)- α -Ethenyldecahydro-2-hydroxy- α ,2,5,5,8a-pentamethyl-1-naphthalenepropanol; [1R-[1 α (R*),2 β ,4 α ,8a α]] - α -ethenyldecahydro-2-hydroxy- α ,2,5,5,8a-pentamethyl-1 Naphthalenepropanol; (13R)-Labd-14-ene-8,13-diol; Sclareol; (-)-Sclareol; [1R-[1.alpha.(R*),2.beta.,4a.beta.,8a.alpha.]]-2-hydroxy-.alpha.,2,5,5,8a-pentamethyl-.alpha.-vinyldecahydronaphthalene-1-propan-1-ol | |

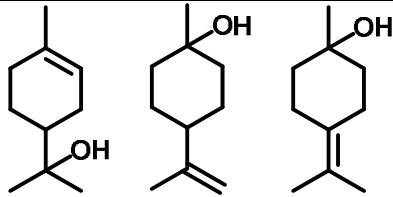
Current regulation: /

Clinical data: /

Additional information:

An older RIFM review exists (128), reporting several human maximisation tests with different samples of sclareol, yielding partly positive, partly negative results. A more recent RIFM review is available (199), citing no clinical data, but several maximisation studies, one of which was positive in a few volunteers, which was apparently due to an impurity.

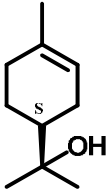
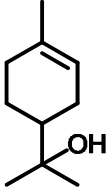
[0986/06 - Opinion on Sclareol \(sensitisation only\)](http://ec.europa.eu/health/ph_risk/committees/04_sccp/docs/sccp_o_056.pdf)
(http://ec.europa.eu/health/ph_risk/committees/04_sccp/docs/sccp_o_056.pdf)

| | |
|---|---|
| TERPINEOL |  |
| CAS # 8000-41-7 | |
| EC # 232-268-1 | |
| Mixtures of isomers | |
| Terpineol 318, mixture of terpineol isomers alfa, beta, gamma | alfa gamma beta |

Current regulation: /

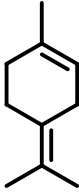
Clinical data:
A RIFM review is available (200), citing negative human induction studies and one clinical study "Takenaka 1986", finding 4 of 312 patients with 0.05% to 0.5% terpineol in a cream base and in ethanol, resp., and 2 negative clinical studies of limited size. In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and 5% terpineol in pet., tested in 100 consecutive patients in Belfast, were observed (15).

Additional information: It is a "top 100" substance (IFRA, pers. comm. 2010).

| | |
|--|---|
| alpha-TERPINEOL |  10482-56-1 |
| CAS # 10482-56-1 / 98-55-5 | |
| EC # 233-986-8 / 202-680-6 |  98-55-5 |
| 2-[(1S)-4-Methyl-1-cyclohex-3-enyl]propan-2-ol (10482-56-1) | |
| 2-(4-Methyl-1-cyclohex-3-enyl)propan-2-ol (98-55-5) | |
| 10482-56-1: (S)-(-)-p-Menth-1-en-8-ol; (-)-α-Terpineol; (S)-(-)-Terpineol; (S)-(-)-α-Terpineol; (S)-α-Terpineol; l-α-Terpineol | |
| 98-55-5: p-Menth-1-en-8-ol; (±)-α-Terpineol; 1,1-Dimethyl-1-(4-methylcyclohex-3-enyl)methanol; 1-p-Menthen-8-ol; 2-(4-Methyl-3-cyclohexenyl)-2-propanol; 4-(2-Hydroxy-2-propyl)-1-methylcyclohexene; 8-Hydroxy-p-menth-1-ene; NSC 21449; NSC 403665; PC 593; Pine Oil 593; Terpineol 350; dl-α-Terpineol; α,α,4-Trimethyl-3-cyclohexene-1-methanol; α-Terpineol | |
| Current regulation: / | |

Clinical data:
A RIFM review is available (201) specifically on (-)-alpha-terpineol stating that "no data is available" regarding skin sensitisation. Another RIFM review is available on alpha-terpineol (202). In the Frosch 2002 b study, 1 of 1606 consecutive patients showed a positive reaction, but 11 patients doubtful reactions to alpha-terpineol (5% pet.) (17). The DeGroot 1985 study identified no positive reactions among 179 patients using a 15% PT preparation of terpineol (mixed isomers) (25). In 63 patients positive to the FM I, 2 had a positive PT reaction to alpha terpineol, 5% pet., in the Santucci 1987 study (28). A clinical series from Portugal, addressing contact allergy to oil of turpentine diagnosed in 30 patients, used a series with pure terpenes. A total of 3 of 30 patients reacted positively to alpha-terpineol (189)

Additional information: see also terpineol (mixture of isomers). Comments on turpentine under pinene.

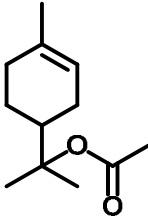
| | |
|--|---|
| Terpinolene |  |
| CAS # 586-62-9 | |
| EC # 209-578-0 | |
| 1-Methyl-4-propan-2-ylidenecyclohexene | |
| p-Mentha-1,4(8)-diene; 1-Methyl-4-(1-methylethylidene)-cyclohexene; 4-Isopropylidene-1-methylcyclohexene; Isoterpinene; Nofmer TP; Terpinolen; Terpinolene; α-Terpinolene; δ-Terpinene | |

Current regulation: Annex III, part 1, n° 133 (Peroxide value less than 10 mmoles/L in substance)

Clinical data:
A 49-year-old machine cleaner developed occupational contact dermatitis due to the cleaner, which gave a positive patch test result at 1:10 000 in water. Of the ingredients identified by chromatography, only .delta.-3-carene and terpinolene, tested 5% pet.,

gave a positive result (negative in 10 controls) (203). Eleven patients sensitised to tea tree oil showed positive reactions to alpha-terpinene, terpinolene and ascaridol (204).

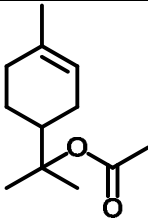
Additional information: It is a "top 100" substance (IFRA, pers. comm. 2010)

| | |
|---|---|
| TERPINEOL ACETATE (Isomer mixture) |  |
| CAS # 8007-35-0 | |
| EC # 232-357-5 | |
| 4-Methyl-1-propan-2-yl-1-cyclohex-2-enyl acetate | |
| Terpinyl acetate | |

Current regulation: /

Clinical data:
In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and 5% terpinyl acetate in pet., tested in 106 consecutive patients in Barcelona, were observed (15)

Additional information: It is a "top 100" substance (IFRA, pers. comm. 2010)

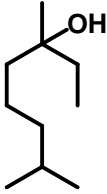
| | |
|---|--|
| alpha-TERPINYL ACETATE |  |
| CAS # 80-26-2 | |
| EC # 201-265-7 | |
| 2-(4-Methyl-1-cyclohex-3-enyl)propan-2-yl acetate | |
| 3-Cyclohexene-1-methanol, $\alpha,\alpha,4$ -trimethyl-, acetate; p-Menth-1-en-8-ol, acetate; (\pm)- α -Terpineol acetate; (\pm)- α -Terpinyl acetate; 2-(4-Methyl-3-cyclohexen-1-yl)-2-propyl acetate; Terpinyl acetate; α -Terpineol acetate; p-Menth-1-en-8-yl acetate; 1-Methyl-1-(4-methylcyclohex-3-enyl)ethyl ethanoate; (\pm)-.alpha.,.alpha.,4-trimethylcyclohex-3-ene-1-methyl acetate | |

Current regulation: /

Clinical data:

The DeGroot 1985 study identified no positive reactions among 179 patients using a 10% PT preparation of "terpinyl acetate" (25).

Additional information: It is a "top 100" substance (IFRA, pers. comm. 2010)

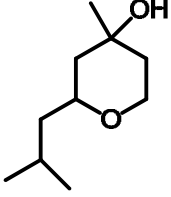
| | |
|---|---|
| Tetrahydrolinalool |  |
| CAS # 78-69-3 | |
| EC # 201-133-9 | |
| 3,7-Dimethyloctan-3-ol | |
| 2,6-Dimethyl-6-octanol; 3,7-Dimethyloctan-3-ol; Linalool tetrahydride; NSC 128151; Tetrahydrolinalool | |

Current regulation: /

Clinical data:
/

Additional information: It is a "top 100" substance (IFRA, pers. comm. 2010). A RIFM

review is available (205) quoting 1 negative human maximisation test.

| | |
|---|---|
| TETRAHYDRO-METHYL-METHYLPROPYL)-PYRAN-4-OL |  |
| CAS # 63500-71-0 | |
| EC # 405-040-6 | |
| 4- Methyl-2-(2-methylpropyl)tetrahydro-2H-4-pyranol | |
| 2-(2-Methylpropyl)-4-hydroxy-4-methyltetrahydropyran; 2-Isobutyl-4-hydroxy-4-methyltetrahydropyran; 2-Isobutyl-4-methyltetrahydropyran-4-ol; 4-Hydroxy-4-methyl-2-(2-methylpropyl)tetrahydropyran; Florosa; Rozanol | |

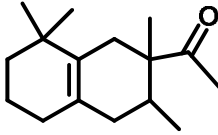
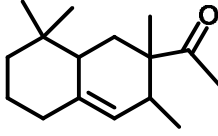
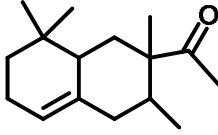
Current regulation: /

Clinical

data:

/

Additional information: It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|--|---|
| TETRAMETHYL ACETYLOCTAHYDRONAPHTHALENES |  |
| CAS # 54464-57-2 / 54464-59-4 / 68155-66-8 / 68155-67-9 | |
| EC # 259-174-3 / 259-175-9 / 268-978-3 / 268-979-9 | |
| 1-(1,2,3,4,5,6,7,8-Octahydro-2,3,8,8-tetramethyl-2-naphthalenyl)-ethanone (54464-57-2) | |
| 1-(1,2,3,4,5,6,7,8-Octahydro-2,3,5,5-tetramethyl-2-naphthalenyl)-ethanone (54464-59-4) | |
| 1-(1,2,3,5,6,7,8,8a-Octahydro-2,3,8,8-tetramethyl-2-naphthalenyl)-ethanone (68155-66-8) |  |
| 1-(1,2,3,4,6,7,8,8a-Octahydro-2,3,8,8-tetramethyl-2-naphthalenyl)-ethanone (68155-67-9) | |
| 54464-57-2: 1-(1,2,3,4,5,6,7,8-Octahydro-2,3,8,8-tetramethyl-2-naphthalenyl)ethanone; 1',2',3',4',5',6',7',8'-Octahydro-2',3',8',8'-tetramethyl-2'-acetonaphthone; 7-Acetyl-1,2,3,4,5,6,7,8-octahydro-1,1,6,7-tetramethylnaphthalene; Amberonne; Ambralux; Iso Ambois Super; Iso-E Super; Isocyclemon E; OTNE; Orbitone |  |

Current regulation: /

Clinical

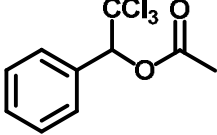
data:

In the Frosch 2002 a study, 0.2% of 1855 consecutive patients reacted to the compound (brand name mentioned: „Iso E. Super“, 5% pet.) (16). In the Frosch 1995 dose-finding pilot study, 1 positive reaction both to 1% and 5% "Iso E Super ®" in pet., tested in 313 consecutive patients in Bordeaux and London, were observed (15). The Larsen 2001 study yielded 1.7% positive reactions (5% pet.) in 178 patients with known contact allergy to fragrance ingredients (19).

Additional information: According to CosIng: "Mixture of isomers: 1-(1,2,3,4,5,6,7,8-Octahydro-2,3,8,8-tetramethyl-2-naphthyl)ethan-1-one; 1-(1,2,3,4,5,6,7,8-Octahydro-2,3,5,5-tetramethyl-2-naphthyl)ethan-1-one; 1-(1,2,3,5,6,7,8,8a-Octahydro-2,3,8,8-tetramethyl-2-naphthyl)ethan-1-one (68155-67-9); 1-(1,2,3,4,6,7,8,8a-Octahydro-2,3,8,8-tetramethyl-2-naphthyl)ethan-1-one (68155-66-8) "

(<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=40504>, last accessed 2009-11-11).

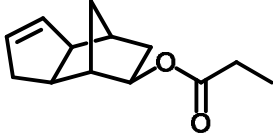
It is a "top 100" substance (IFRA, pers. comm. 2010)

| | |
|--|---|
| TRICHLOROMETHYL PHENYL CARBINYL ACETATE |  |
| CAS # 90-17-5 | |
| EC # 201-972-0 | |
| 2,2,2-Trichloro-1-phenylethyl acetate | |
| Benzenemethanol, α -(trichloromethyl)-, acetate; Benzyl alcohol, α -(trichloromethyl)-, acetate (Trichloromethyl)phenylcarbinyl acetate; (\pm)- α -(Trichloromethyl)benzyl acetate; 2-Acetoxy-1,1,1-trichloro-2-phenylethane; Crystal rose; NSC 165582; Rosacetol; Rosephenone; Rosetone; Rosone; α -(Trichloromethyl)benzyl acetate | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm. 2010)

| | |
|--|---|
| TRICYCLODECENYL PROPIONATE |  |
| CAS # 17511-60-3 | |
| EC # 241-514-7 | |
| 3α,4,5,6,7,7α-Hexahydro-4,7-methano-1H-inden-6-yl propionate | |
| 4,7-Methano-1H-inden-6-ol, 3 α ,4,5,6,7,7 α -Hexahydro-, propanoate; 4,7-Methanoinden-6-ol, 3 α ,4,5,6,7,7 α -Hexahydro-, propionate; Cyclaprop; Florocyclene; Greenyl propionate; Tricyclo(5.2.1.02,6)dec-3-en-8-yl propionate. | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm. 2010).

| | |
|---|---|
| 3-(5,5,6-TRIMETHYLBICYCLO[2.2.1]HEPT-2-YL)-CYCLOHEXAN-1-OL |  |
|---|---|

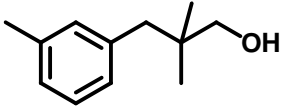
Opinion on fragrance allergens in cosmetic products

| | |
|---|--|
| CAS # 3407-42-9 | |
| EC # 222-294-1 | |
| 3-(5,5,6-Trimethyl-6-bicyclo[2.2.1]heptanyl)cyclohexan-1-ol | |
| 3-(5,5,6-Trimethyl-2-norbornyl)-cyclohexanol; 3-(5,5,6-Trimethylbicyclo[2.2.1]hept-2-yl)cyclohexan-1-ol; Hydroxy-1-(5-isocamphyl)cyclohexane; Sandela | |

Current regulation: /

Clinical data: /

Additional information: part of "synthetic sandalwood oil".

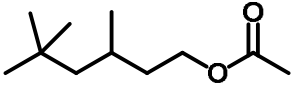
| | |
|--|---|
| TRIMETHYL-BENZENEPROPANOL (Majantol) |  |
| CAS # 103694-68-4 | |
| EC # 403-140-4 | |
| 2,2-Dimethyl-3-(3-methylphenyl)propan-1-ol | |
| 2,2-Dimethyl-3-(3-tolyl)propan-1-ol; 3-(2,2-Dimethyl-3-hydroxypropyl)toluene | |

Current regulation: /

Clinical data:

In the Larsen 2002 c study, majantol (conc. not given, elsewhere reported as 5% pet.) caused positive PT reactions in 3.2% of patients with known contact allergy to fragrance ingredients. In a later study by the IVDK, 0.5% (95% CI: 0.3 – 0.7%) consecutive patients displayed a positive reaction to majantol 5% pet. (206). In the IVDK 2010 study, majantol was tested both in n=2189 consecutive patients, yielding 0.36 % (95% CI: 0.12–0.60%) positive reactions, and in the context in a special series, applied in an aimed fashion to n=4972 patients, yielding 0.76% (95% CI: 0.49–1.03%) (standardised) positive reactions (7). In a recent study from Copenhagen, DK, 6 of 722 patients tested with this compound were found positive, 2 of these to material used earlier provided by Symrise, 4 to material by Allmiral/Hermal/Trolab used later instead. There was no significant difference between these proportions obtained with batches of majantol from different production processes (207).

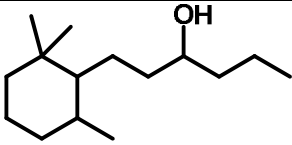
Additional information: /

| | |
|---|---|
| TRIMETHYLHEXYL ACETATE |  |
| CAS # 58430-94-7 | |
| EC # 261-245-9 | |
| 3,5,5-Trimethylhexyl acetate | |
| 1-Hexanol, 3,5,5-trimethyl-, acetate; Vanoris; neononyl acetate | |

Current regulation: /

Clinical data: /

Additional information: It is a "top 100" substance (IFRA, pers. comm. 2010)

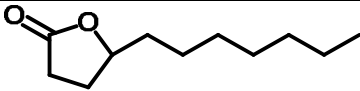
| | |
|---|---|
| TRIMETHYL-PROPYLCYCLOHEXANEPROPANOL (TMCH) |  |
| CAS # 70788-30-6 | |
| EC # 274-892-7 | |
| 1-(2,2,6-Trimethylcyclohexyl)hexan-3-ol | |
| Other names: 2,2,6-Trimethyl-alpha-propylcyclohexanepropanol (REACH, EINECS); cyclohexanepropanol; Finotimber; Timberol | .alpha.-Propyl-2,2,6-trimethyl-6-(2,2,6-Trimethylcyclohexyl)-4-hexanol; |

Current regulation: /

Clinical data:

In the Larsen 2001 study, none of 178 patients with contact allergy to fragrance ingredients reacted positively to this ingredient, PTed at 5% pet. (19).

Additional information: /

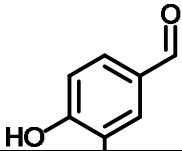
| | |
|---|---|
| gamma-UNDECALACTONE |  |
| CAS # 104-67-6 | |
| EC # 203-225-4 | |
| 5-Heptyltetrahydrofuran-2-one | |
| Undecanoic acid, 4-hydroxy-, γ -lactone; (RS)- γ -Undecalactone; (\pm)- γ -Undecalactone; 4-Hydroxyundecanoic acid lactone; 4-Undecanolide; 5-Heptyldihydro-2(3H)-furanone; NSC 406421; NSC 46118; NSC 76413; Neutralizing agent 350120-1; Peach lactone; Peche Pure; Persicol; γ -(n-Heptyl)- γ -butyrolactone; γ -Heptyl- γ -butyrolactone; γ -Heptylbutyrolactone; γ -Undecalactone; γ -Undecanolactone; γ -Undecanolide; γ -n-Heptylbutyrolactone | |

Current regulation: /

Clinical data: /

Additional information:

It is a "top 100" substance (IFRA, pers. comm. 2010)

| | |
|-----------------|---|
| VANILLIN |  |
| CAS # 121-33-5 | |
| EC # 204-465-2 | |

| | |
|---|--|
| 4-Hydroxy-3-methoxybenzaldehyde | |
| 2-Methoxy-4-formylphenol; 3-Methoxy-4-hydroxybenzaldehyde; 4-Formyl-2-methoxyphenol; 4-Hydroxy-5-methoxybenzaldehyde; 4-Hydroxy-m-anisaldehyde; H 0264; Lioxin; NSC 15351; NSC 403658; NSC 48383; Rhovanil; Vanillaldehyde; Vanillic aldehyde; Vanillum; m-Methoxy-p-hydroxybenzaldehyde; p-Hydroxy-m-methoxybenzaldehyde; p-Vanillin | |

Current regulation: /

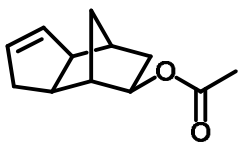
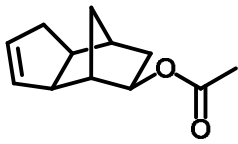
Clinical data:

In a series of 40 of 744 consecutive patients PTed with an extended fragrance series (Sheffield 1999), 1 positive reaction to vanillin was observed (3). In the Wöhrle 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded n=1 (0.1%) positive reaction to vanillin 10 % pet. (22). The IVDK 2010 study, n=10, i.e., 0.19% (95% CI: 0.07 - 0.32%; percentages standardised for age and sex) of 4377 patients PTed reacted to the compound, tested 10% pet. (7). In n=102 patients with a positive reaction to MPR, 19 compounds of this natural mixture were tested, among these, vanillin, to which none reacted positively (208). In 21 patients with contact allergy to propolis, 2 also reacted to vanillin (10% pet.) (209).

A 13-year-old girl with recurrent (peri-)cheilitis after application of a vanilla lip salve tested strongly positive to this salve (as is), "Vanilla 10% pet." (unclear, whether natural extract or vanillin) and MPR (210). Trattner/David identified 1 / 641 consecutive patients with positive reaction to vanillin (31).

Additional information:

Naturally occurring in the fruit of *Vanilla planifolia* after a fermentation process, in styrax, clove oil, potatoes, wood, including Myroxylon pereirae resin, and other material (53). Nowadays, vanillin is synthesised from eugenol, guajakol and lignin residues from paper production, however, not fully achieving the subtle scent and taste of the natural material (53). It is a "top 100" substance and classified as R43 (IFRA, pers. comm. 2010).

| | |
|--|---|
| VERDYL ACETATE | |
| CAS # 2500-83-6/ 5413-60-5 |  |
| EC # 219-700-4 / 226-501-6 | |
| 3a,4,5,6,7,7a-Hexahydro-4,7-methanoinden-6-yl acetat (2500-83-6) | 2500-83-6 |
| 3a,4,5,6,7,7a-Hexahydro-4,7-methano-1H-inden-5-yl acetat (5413-60-5) |  |
| 2500-83-6: 4,7-Methano-1H-inden-5-ol, 3a,4,5,6,7,7a-hexahydro-, acetate; 4,7-Methanoinden-5-ol, 3a,4,5,6,7,7a-hexahydro-, acetate; NSC 142428; NSC 94573 | 5413-60-5 |
| 5413-60-5: 4,7-Methano-1H-inden-6-ol, 3a,4,5,6,7,7a-hexahydro-, acetate; 4,7-Methanoinden-6-ol, 3a,4,5,6,7,7a-hexahydro-, acetate; 4,7-Methano-3a,4,5,6,7,7a-hexahydroinden-6-yl acetate; 8-Acetoxytricyclo[5.2.1.0 ^{2,6}]dec-3-ene; Greenyl acetate; | |

| | |
|-------------------------------------|--|
| Herbaflorat; Jasmacyclene; NSC 6598 | |
|-------------------------------------|--|

Current regulation: /

Clinical data: /

Additional information:

In CosIng, both above CAS numbers are listed under "verdyl acetate" (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41289>, last accessed 2010-07-19).

In the CAS, there are 2 separate entries; moreover, there are 2 separate RIFM reviews:

- # 2500-83-6: Other names: Tricyclo[5.2.1.0^{2,6}]dec-4-en-8-yl acetate (REACH, EINECS, INCI Name according to CAS); 3a,4,5,6,7,7a-Hexahydro-4,7-methanoinden-6-yl Acetate; Tricyclodecen-4-yl 8-Acetate. It is a "top 100" substance (IFRA, pers. comm. 2010). A RIFM review is available, stating that "no data is available" regarding the skin sensitising properties of the substance (211).
- # 5413-60-5: Other names: 3a,4,5,6,7,7a-hexahydro-4,7-methanoinden-6-yl acetate (REACH, EINECS, INCI Name according to CAS), 4,7-Methano-3a,4,5,6,7,7a-hexahydroinden-6-yl acetate; 4,7-Methanoinden-6-ol, 3a,4,5,6,7,7a-hexahydro-, acetate; 8-Acetoxytricyclo[5.2.1.0^{2,6}]dec-3-ene; Tricyclodecanyl acetate; Greenyl acetate; Herbaflorat; Jasmacyclene; NSC 6598; Verdyl acetate. It is a "top 100" substance (IFRA, pers. comm. 2010). A RIFM review is available (212), citing 2 negative human maximisation tests and 1 negative HRIPT.

Natural extracts / essential oils

Natural raw materials in terms of extracts are used in the fragrance and flavour industry for various reasons. Most importantly, several naturally occurring mixtures have a very complex composition and sensory nature which cannot (fully) be achieved by synthetic material. Moreover, several compounds cannot be synthesised at a competitive price, and the demand for perfumes based on natural materials is considerable (34).

The three main methods used to concentrate plant fragrance substances as essential oils comprise steam distillation, mechanical processes from the epicarp of Citrus fruits ("pressing") and dry distillation. An Essential oil is „obtained by steam distillation with addition of water in the still (hydrodistillation) or without addition of water in the still (directly by steam“)(213). Essential oil of fruit juice is „obtained by from a fruit juice during its concentration or during UHT (flash pasteurization) treatment“ (213). Cold pressed essential oil is „obtained by mechanical processes from the epicarp of the fruit of a Citrus, at ambient temperature“(213). Citrus peel oils, apart from distilled Citrus oils, are produced with various methods (214). The oil consists of a high volume of volatile terpenes, mostly monoterpenes but also contains small amounts of non-volatile compounds such as dyes, waxes and furocoumarines.

The method of solvent extraction is generally applied in the separation of heat-labile materials or if an essential oil can only be obtained in very low yield, e.g. from blossoms. It is also used if the non-volatile components are desired for their fixative properties, e.g. in the preparation of resinoids from exudates. The most important extracts are termed: (i) concrete: an extract „obtained from a fresh plant natural raw material by extraction with a solvent“¹⁸, containing not only volatile, but also a large proportion of non-volatile substances such as waxes; and (ii) absolute: „product, obtained by extraction with ethanol from a concrete, a floral pomade, a resinoid or a supercritical fluid extract. The ethanolic solution is generally cooled down and filtered in order to eliminate the «waxes»; the ethanol is then eliminated by distillation“¹⁹. Resinoids, used for their fixative properties, are „obtained from a dry plant natural raw material by extraction with a solvent“²⁰. The products are usually highly viscous and thus might sometimes be diluted, e.g. with phthalates or benzyl benzoate. Oleoresins are extracts „of spice or aromatic herb“ by „treating a natural raw material with a solvent, then, after filtration if necessary, the solvent is eliminated“²¹.

Regarding clinical data in terms of contact allergy to fragrance ingredients, the main focus of case report or clinical studies regarding essential oils and natural extracts, respectively, is on general dermatological patients with complaints related to use of cosmetics etc. However, series of cases with occupational exposure to essential oils with occupational allergic contact dermatitis have also been reported (e.g., masseurs,

¹⁸ ISO/DIS 9235

¹⁹ ISO/DIS 9235

²⁰ ISO/DIS 9235

²¹ ISO/DIS 9235

physiotherapists (215, 216), aromatherapists (217-221), beauticians doing massages (222); for further details, e.g., PT results with various essential oils, see original case reports. "Current Regulation" refers to the EU Cosmetics Directive only.

Catalogue of natural extracts / essential oils evaluated

ACORUS CALAMUS ROOT OIL

Calamus Oil; "Sweet Flag Oil"

CAS 84775-39-3; EC 283-869-0

(*Acorus calamus*, ext. = INCI name)

Current regulation: /

Clinical data:

The Rudzki 1976 study found no positive reaction in 200 patients to "calamus" essential oil, 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=7 (8.1%) positive reactions to "calamus" essential oil 2% pet. (27).

Additional information:

ISO 4720:2009 nomenclature: *Acorus calamus* L. (sweet flag calamus). *Acorus Calamus Root Oil* is an essential oil obtained from the rhizomes of the calamus, *Acorus calamus* L., Araceae. It contains beta-asarone (up to 96%, depending on ploidy, and with this, origin (34)), calamene (about 4%), calamol (about 3%) alpha-asarone (about 1%), camphene (about 1%) and some beta-pinene and asaronaldehyde (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41330>, last accessed 2010-01-29). Use is restricted due to potential toxicity of beta-asarone (34).

CANANGA ODORATA and Ylang-ylang oil

Ylang-ylang and cananga oils are essential oils that are obtained from two subspecies of the cananga tree (34). In the INCI nomenclature, both are not differentiated.

CANANGA ODORATA FLOWER EXTRACT

CAS 83863-30-3; EC 281-092-1
(ylang-ylang, ext.) INCI name:
CANANGA ODORATA EXTRACT

CANANGA ODORATA FLOWER OIL

CAS 8006-81-3, 68606-83-7; EC / (oils,
ylang-ylang) INCI name: CANANGA
ODORATA OIL

Current regulation: ...

Clinical data:

Ylang-ylang oil

ISO 4720:2009 nomenclature: *Cananga odorata* (Lam.) Hook. f. et Thomson *forma*

genuina)

In the Larsen 2002 c study, “synthetic ylang-ylang oil” caused 6.4% positive reactions in 218 patients with known contact allergy to fragrance ingredients (1). In a Japanese study, M. Sugawara et al. noted a significant decline of the proportion of patients reacting positively to “ylang-ylang oil 5% pet.” from 1971 to 1989, the overall number in patients with cosmetic dermatitis amounting to 176 of 1438 (12.2%, 95% CI: 10.6 – 14.0%) (223). In the Frosch 2002 b study, two fractions of Ylang-Ylang oil (I and II) were separately tested, each at 10% pet. Fraction I yielded 2.6%, fraction II 2.5% positive test reactions (no data on concomitant reactivity given) (17). The deGroot 2000 study, with 1825 consecutively tested patients, found 18 positive PT reactions to “ylang-ylang oil”, tested at 4% in pet. (12). The Sugiura 2000 study with 1483 patients with suspected cosmetic dermatitis observed 0.8% positive PT reactions with ylang-ylang oil (5% pet.) (14). The Coimbra 2000 study found in 67 patients with positive reaction to the FM I who were tested with ylang-ylang oil (2% pet.) 13.4% positive reactions (9). The Belsito 2006 study (20) yielded 0.6% positive reactions to ylang-ylang oil. The subsequent NACDG 2009 study identified 1.5% positive reactions in 4434 patients PTed with 2% “ylang-ylang oil” (21). The IVDK 2010c study found 2.5% positive reactions in 3175 consecutively tested patients, and 3.9% in 2155 patients tested in the context of a special series (30). In a study from Alicante, Spain, 86 selected patients were patch tested with an extended fragrance series; n=12 reacted positively to ylang-ylang oil and 3 to “cananga oil” (48).

Cananga oil

ISO 4720:2009 nomenclature: *Cananga odorata* (Lam.) Hook. f. et Thomson *forma macrophylla*. For Oil of cananga (*Cananga odorata* (Lam.) Hook. f. et Thomson, *forma macrophylla*) an ISO standard exists: ISO 3523:2002. Cananga oil is produced by steam distillation of the flowers of *Cananga odorata* (DC.) Hook f. et Thomson subsp. *macrophylla* (*Annonaceae*). The composition resembles that of “ylang-ylang III”, but with a higher content of caryophyllene (30-40%). Cananga oil originates almost exclusively in Java; annual production about 50 t. The oil is used mainly in perfuming soaps where it is more stable than ylang-ylang oils due to its lower ester content (34).

Sugiura et al. (2000) found 1.1% positive reactions to “cananga oil”, tested 5% pet. (14). Cananga oil (2% pet.) mentioned in the same Portuguese study already cited (9) yielded 10.4% positive reactions. In the An 2005 study, 5 of 422 consecutive patients, i.e., 1.2%, had positive reactions to cananga odorata oil tested at 2% concentration (13).

Studies with both oils

The Goossens 1997 study found 3 of 111 patients positive to “ylang-ylang oil 5% pet.”, and 4 to “cananga oil 15% pet.” – all sensitised to other fragrance allergens (23). The Rudzki 1976 study found 1 positive reaction in 200 patients to “cananga” and 4 to “ylang-ylang” essential oil, both tested 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=10 (11.6%) positive reactions to “cananga” and n=8 (9.3%) to “ylang-ylang” essential oils, each tested at 2% pet. (27). Nakayama et al. found 1974 (after (29)) 11 “strong positive” and 15 “weak positive” reactions to “Cananga oil” and 9 and 16, resp., to “Ylang-ylang oil” (unknown test concentration) in 183 patients.

A number of case reports highlight the possibility of occupational contact and sensitisation, e.g. (222, 224).

Additional information:

Ylang-ylang oil

The composition of this essential oil is defined by a standard: ISO 3063:2004. Ylang-ylang oils are obtained by steam distillation of freshly picked blossoms of *Cananga odorata* (DC.) Hook f. et Thomson subsp. *genuina* (*Annonaceae*). The oil is produced mainly in Madagascar and the Comoro islands. Four fractions are collected at progressively longer distillation times and are known as "extra", "I", "II" and "III". The composition of the various oil fractions depends on the duration of distillation. The first fraction has the highest content of strongly odiferous constituents such as p-cresyl methyl ether (5-16%), methyl benzoate (4-9%), (-)-linalool (7-24%), benzyl acetate (5.5-17.5%), and geranyl acetate (2.5-14%). The other fractions contain increasing amounts of sesquiterpene hydrocarbons such as caryophyllene, germacrene-D, and (E,E)-alpha-farnesene (> 70% in "ylang-ylang III"). Components such as p-cresol, eugenol and isoeugenol are important for odour, although they are present only in low concentration (34). According to (30) the maximum observed concentration in ylang-ylang I and II are (in %): germacrene-D (28); (E,E)-alpha-farnesene (21); caryophyllene (17); linalool (I: 19.0; II: 9.5); benzyl benzoate (8.0); farnesol (4.0); benzyl salicylate (4.0); (E,E)-farnesyl acetate (3.5); geraniol (2.5); isoeugenol (0.8); benzyl alcohol (0.5); eugenol (0.5); p-cresyl methyl ether (I: 5.0; II 3.5); methyl benzoate (I: 5.5; II: 3.5); benzyl acetate (I: 10.0; II: 5.0); geranyl acetate (I: 15.0; II: 12.0).

CEDRUS ATLANTICA BARK OIL

CAS 92201-55-3; EC 295-985-9
(*Cedrus atlantica*, ext. = INCI) /
8000-27-9; EC / (Oils,
cedarwood) INCI name: CEDRUS
ATLANTICA OIL

Cedarwood oil

Current regulation: /

Clinical data:

In the Wöhrle 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded n=5 (0.7%) positive reactions to cedarwood oil 10% pet. (22). (The exact origin of "cedarwood oil" in this study is not clear.) The IVDK 2010 c study identified 0.8% positive reactions in 6223 patients tested in the context of a special series with a cedarwood oil tagged with CAS # 8000-27-9 (30).

Additional information:

Cedrus Atlantica Bark Oil is the volatile oil obtained from the bark of *Cedrus atlantica*, *Pinaceae*

(<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=55309>, last accessed 2010-01-05). The main odiferous component is alpha-atlantone [32207-08-2] (39)

Nomenclature also used: *Cedrus atlantica* wood oil (*Cedrus atlantica* (Endl.) G.Manetti ex Carrière)²²

See also *Juniperus virginiana*.

²² ISO 4720:2009 nomenclature

CEDRUS DEODARA WOOD OILCAS 91771-47-0; EC 294-939-5 (*Cedrus deodara*, ext.)*Cedarwood oil*

Current regulation: /

Clinical data:

The Rudzki 1976 study found 3 positive reactions in 200 patients to "cedarwood" essential oil 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=3 (3.5%) positive reactions to "Himalayan cedarwood" essential oil 2% pet. (27). (The labelling in the latter report points to *Cedrus deodara* as source of "cedarwood oil" in these 2 Polish studies.)

Additional information:

Cedrus Deodara Wood Oil, Himalayan cedarwood oil (*Cedrus deodara* (Roxb. ex D. Don) G. Don)²³, is the volatile oil obtained by steam distillation of the stumps of the Deodar Cedar, *Cedrus deodara*, *Pinaceae* (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=55311>, last accessed 2010-01-29).

Several other conifer species are called cedars, and the corresponding oils vary considerably in composition. These include Cedar leaf oil (Thuja oil) produced by steam distillation of fresh leaves and branch ends of *Thuja occidentalis* L. (*Cupressaceae*) from North America, containing a minimum of 60% thujone [8007-20-3] [90131-58-1] (34). Texas cedarwood oil is produced by steam distillation of chopped wood of *Juniperus mexicana* Schiede (*Cupressaceae*), containing alpha-cedrene (15-25%), thujopsene (25-35%), cedrol 20% minimum [8000-27-9] [91722-61-1] (34). Chinese cedarwood oil is similar to Texas cedarwood oil, obtained by steam distillation of *Cupressus funebris* Endl., *Cupressaceae* (*Chamaecyparis funebirs* (Endl.) France), which is a weeping cypress [8000-27-9] [85085-29-6] (34).

CINNAMOMUM CASSIA LEAF OIL

94961-46-6 [invalid] / 8007-80-5; EC / (Oils, cassia) INCI name: CINNAMONUM CASSIA OIL

*Cassia Oil; Cassia leaf Oil; Cinnamon Oil Chinense***CINNAMOMUM ZEYLANICUM BARK OIL**CAS 84649-98-9; EC 284-635-0 (*Cinnamomum zeylanicum*, ext. = INCI)*Cimmamon Bark Oil Ceylon; Cinnamon Oil Ceylon*

Current regulation: /

Clinical data:

The Rudzki 1976 study found 2 positive reactions in 200 patients to "cassia" essential oil, 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=24

²³ ISO 4720:2009 nomenclature

(27.9%) positive reactions to "cassia" essential oil 2% pet. (27).

A 32 year old Spanish physiotherapist developed vesicular hand dermatitis after using a "balsam from ash extract" cream. PTing revealed positive reactions to this cream, the FM I, eugenol, and 2 components of the cream: "cinnamon oil" (0.5% pet.) and clove oil (1% pet.) (225).

Additional information:

ISO 4720:2009 nomenclature: *Cinnamomum tsumu* Helms, syn. *Cinnamomum cassia* auct. and *Cinnamomum zeylanicum* Blume syn. *Cinnamomum verum* J. Presl, respectively. Cassia oil (Chinese cinnamon oil) is obtained by steam distillation of the leaves, twigs, and bark of *Cinnamomum aromaticum* Nees (*C. cassia* Blume, *Lauraceae*). In contrast to cinnamon bark oil (see below), cassia oil contains a considerable amount of 2-methoxycinnamal (3-15%), in addition to its main constituent, cinnamal (70-88%). Cassia oil is predominantly used in flavouring soft drinks, with an annual production of a few hundred tons (34). For Oil of cassia, Chinese type (*Cinnamomum aromaticum* Nees, syn. *Cinnamomum cassia* Nees ex Blume) an ISO standard exists: ISO 3216:1997

Cinnamomum Zeylanicum Bark Oil is the volatile oil expressed from the bark of the Ceylon Cinnamon, *Cinnamomum zeylanicum*, *Lauraceae*. It contains mainly cinnamaldehyde (34), e.g. 50-60%, and lesser quantities of eugenol (4-8%), phellandrene

(<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=75370>, last accessed 2009-11-16). For Oil of cinnamon leaf, Sri Lanka type (*Cinnamomum zeylanicum* Blume) an ISO standard exists: ISO 3524:2003

Cinnamomum Cassia Leaf Oil is the volatile oil obtained by steam distillation from the leaves and twigs of the Chinese Cinnamon, *Cinnamomum cassia* (L.), *Lauraceae*. It contains 80% eugenol (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=75368>, last accessed 2009-11-16). The cinnamon leaf oil produced by steam distillation of the leaves of *Cinnamomum zeylanicum* Blume (*C. verum* J.S. Presl) similarly has a content of 70-83% eugenol (34).

Considering the content of well-known allergenic compounds, the essential oil is considered an Established contact allergen in humans,

CITRUS AURANTIUM AMARA FLOWER OIL CAS 8016-38-4, 68916-04-1; EC / (Oils, neroli) /

Neroli oil

CITRUS AURANTIUM AMARA PEEL OIL 72968-50-4; EC 277-143-2 (Orange, sour, ext.)

"Bitter Orange Oil"

INCI names: CITRUS AURANTIUM AMARA ...

Current regulation: /

Clinical data:

The Coimbra 2000 study found in 67 patients with positive reaction to the FM I who were tested with "neroli oil" (2% pet.) 6.6% positive reactions (9). The Rudzki 1976 study found 3 positive reactions in 200 patients to "bitter orange" essential oil 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=2 (2.3%)

positive reactions to "bitter orange" essential oil 2% pet. (27). The IVDK 2010 c study identified 0.7% positive reactions in 6220 patients tested in the context of a special series (30)

Additional information:

ISO 4720:2009 nomenclature: *Citrus aurantium* L., syn. *Citrus amara* Link, syn. *Citrus bigaradia* Loisel, syn. *Citrus vulgaris* Risso. For Oil of neroli (*Citrus aurantium* L. spp. *aurantium*, syn. *Citrus aurantium* L. spp. *amara* var. *pumilia*) an ISO standard exists: ISO 3517:2002. Citrus Aurantium Peel Oil Expressed is an essential oil expressed from the fresh epicarps of the Sour Orange, *Citrus aurantium*, Rutaceae. It contains D-limonene (about 90%), citral, decanaldehyde, methyl anthranilate, linalool, terpineol (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41394>, last accessed 2010-01-29). The aldehyde content is lower and the ester content (e.g., linalyl and geranyl acetate) is higher than in sweet orange oil (34). It is predominantly used for flavouring alcoholic beverages. According to (30) the maximum observed concentration in neroli oil are (in %): linalool (44); limonene (18); β -pinene (17); linalyl acetate (15); *trans*- β -ocimene (8); geranyl acetate (5); *trans*-nerolidol (5); (*E,E*)-farnesol (4); myrcene (4); farnesol (4,0); geraniol (3,5); citral (0,3) (30).

CITRUS AURANTIUM AMARA LEAF OIL

72968-50-4; EC 277-143-2 (Orange, sour, ext.)

Petitgrain oil Paraguay / ... bigarade

Current regulation: /

Clinical data:

The Rudzki 1976 study found 1 positive reaction in 200 patients to "Petitgrain bigarade" and "Petitgrain Paraguay" essential oil each, both tested at 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=7 (8.1%) positive reactions to "Petitgrain bigarade" and n=4 (4.6%) to "Petitgrain Paraguay" essential oil each, both tested at 2% pet. (27).

Additional information:

ISO 4720:2009 nomenclature: *Citrus sinensis* L. Pers. X *Citrus aurantium* L. ssp. *amara* var. *pumilia*. Petitgrain oils in general are steam distilled from the leaves of citrus trees. Citrus Aurantium Leaf Oil is an essential oil obtained from the leaves of the Sour Orange, *Citrus aurantium*, Rutaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41392>, last accessed 2010-02-10). Petitgrain oil Paraguay is obtained from an acclimatised variety of the bitter orange tree. Main constituents are linalool (15-30%) and linalyl acetate (40-60%). A number of trace constituents contribute essentially to the odour (34). Petitgrain oil bigarade is derived from the same species of tree grown in France, Italy, Spain and North Africa (34). For Oil of bitter orange petitgrain, cultivated (*Citrus aurantium* L.) an ISO standard exists: ISO 8901:2003.

Considering the content of well-known allergenic compounds, the essential oil is regarded as an established contact allergen in humans

CITRUS BERGAMIA PEEL OIL EXPRESSED

CAS 89957-91-5, 8007-75-8; EC

| | |
|--|--|
| | 289-612-9 (<i>Bergamot, ext.</i>) |
| <i>Bergamot Oil, Bergamot Orange Oil</i> | INCI: CITRUS AURANTIUM BERGAMIA EXTRACT |

Current regulation: /

Clinical data:

The Rudzki 1976 study found 3 positive reactions in 200 patients to "Bergamot" essential oil 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found no positive reaction to "Bergamot" essential oil 2% pet. (27). In 63 patients positive to the FM I, 2 had a positive PT reaction to bergamot oil, 2% pet., in the Santucci 1987 study (28). A case report from Zacher and Ippen describes 2 patients with allergic contact dermatitis due to bergamot oil (191), one a worker in a perfume factory, the other sensitised by non-occupational use of cosmetics.

Additional information:

ISO 4720:2009 nomenclature: *Citrus bergamia* (Risso et Poit.), syn. *Citrus aurantium* L. subsp. *bergamia* (Wight et Arnott) Engler. Citrus Bergamia Peel Oil Expressed is an essential oil expressed from the epicarps of the Bergamot, *Citrus bergamia* risso, Rutaceae. It contains 35-45% L-linalyl acetate, about 6% linalool, D-limonene, DL-limonene and bergaptene (http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=se_arch.details&id=41398, last accessed 2009-11-27). According to Surburg/Panten: linalyl acetate 22-36%, linalool 3-15%, geranial 0.25-0.5%, citral 1%, with a relatively low terpene content of 25-50% (34, 39). Bergaptene content by HPLC is 0.18-0.38% (34). Annual production from Italy, Brazil, Spain and Ivory Coast is 100 to 150 t. For Oil of bergamot [*Citrus aurantium* L. subsp. *bergamia* (Wight et Arnott) Engler], Italian type an ISO standard exists: ISO 3520:1998.

| | |
|--|--|
| CITRUS LIMONUM PEEL OIL EXPRESSED | CAS 84929-31-7, 8008-56-8; EC 284-515-8 (<i>Lemon, ext.</i>) |
| <i>Lemon oil</i> | INCI names: CITRUS MEDICA LIMONUM ... |

Current regulation: /

Clinical data:

The Coimbra 2000 study found in 67 patients with positive reaction to the FM I who were tested with "lemon oil" (2% pet.) 4.5% positive reactions (9). In the Wöhrle 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded n=2 (0.3%) positive reactions to "lemon oil" 2% pet. (22).

The Rudzki 1976 study found 1 positive reaction in 200 patients to "lemon" essential oil 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=2 (2.3%) positive reactions to "lemon" essential oil 2% pet. (27). The IVDK 2010 c study identified 0.3% positive reactions in 6467 patients tested in the context of a special series (30).

Additional information:

ISO 4720:2009 nomenclature: *Citrus limon* (L.) Burm. f. According to (30) the maximum observed concentration in lemon oil are (in %): limonene (80); β -pinene (16.5); γ -terpinene (12); citral (3.0); geranial (2.0); neral (1.2); β -bisabolene (0.9); geranyl

acetate (0.7); neryl acetate (0.6); linalool (0,3); geraniol (0,2) (30). An ISO standard exists for Oil of lemon [Citrus limon (L.) Burm. f.], obtained by expression: ISO 855:2003. The composition of lemon oil depends on the variety of lemon and the country of origin, see table from (34).

Table 3. Specifications for qualities of lemon oils of different origins

| Parameter | Type | | Mediterranean | | Equatorial |
|--------------------------------|-----------------|---------------|---------------|---------------|---------------------|
| | American Origin | | Italy | Spain | Ivory coast, Brazil |
| | Coast | Desert | | | |
| d_{20}^{20} | 0.851–0.857 | 0.849–0.854 | 0.850–0.858 | 0.849–0.858 | 0.845–0.854 |
| n_D^{20} | 1.4370–1.4760 | 1.4370–1.4760 | 1.4370–1.4760 | 1.4370–1.4760 | 1.4370–1.4790 |
| α_D^{20} | +57° to +65°6' | +67° to +78° | +57° to +66° | +57° to +66° | +57° to +70° |
| Composition by GC [area %] | | | | | |
| β -Pinene | 9–14 | 10–13 | 10–16.5 | 10–16.5 | 7–16 |
| Limonene | 63–70 | 70–80 | 60–68 | 60–70 | 59–75 |
| γ -Terpinene | 8.3–9.5 | 6.5–8 | 8–12 | 8–12.8–12 | 6–12 |
| Neral | 0.6–0.9 | 0.3–0.6 | 0.6–1.2 | 0.4–1 | 0.2–1.2 |
| Geraniol | 1.0–2 | 0.5–0.9 | 0.8–2 | 0.6–2 | 0.5–2 |
| Evaporation residue [weight %] | | | | | |
| | 1.75–3.9 | | 1.5–3.9 | 1.5–3.9 | 1.5–4 |
| Carbonyl value | | | | | |
| | 8–14 | 6.25–12 | 11–17 | 11–17 | 6–17 |
| CD value | min. 0.2 | min. 0.2 | 0.45–0.9 | 0.4–0.9 | 0.2–0.96 |

CITRUS PARADISI PEEL OIL

Grapefruit oil, expressed

CAS 8016-20-4 ; EC /

INCI: CITRUS GRANDIS OIL

Current regulation: II/358 R1

Clinical data: /

Additional information:

Citrus Paradisi Peel Oil is the volatile oil expressed from the peel of the Grapefruit, Citrus *paradisi*, Rutaceae
http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details_v2&id=55434

It is a "top 200" substance and classified as R43 (IFRA, pers. comm.2010)

CITRUS SINENSIS (syn.: AURANTIUM DULCIS) CAS 97766-30-8, 8008-57-9, EC
 PEEL OIL EXPRESSED 307-891-8 (Orange, sweet,

(Sweet) Orange oil

Valencia, ext. = INCI) / 8028-48-6; EC 232-433-8 (Orange, sweet, ext.)

INCI names: CITRUS AURANTIUM DULCIS ...

Current regulation: /

Clinical data:

The Coimbra 2000 study found in 67 patients with positive reaction to the FM I who were tested with "orange oil" (2% pet.) 4.5% positive reactions (9). In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded n=1 (0.1%) positive reactions to orange oil 2% pet. (22). The Rudzki 1976 study found 1 positive reaction in 200 patients to "sweet orange" essential oil, 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=3 (3.5%) positive reactions to "sweet orange" essential oil 2% pet. (27). In the Frosch 1995 dose-finding pilot study, neither positive nor irritant reaction to 1% and 5% "orange oil Bras." in pet., tested in 205 consecutive patients in Dortmund and Göttingen, were observed (15). The IVDK 2010 c study identified 0.2% positive reactions in 6246 patients tested in the context of a special series (30).

Additional information:

ISO 4720:2009 nomenclature: *Citrus sinensis* (L.) Osbeck. For Oil of sweet orange (*Citrus sinensis* (L.) Osbeck), CAS 8008-57-9, obtained by mechanical treatment, an ISO norm exists: ISO 3140:2005. The oils have a high terpene hydrocarbon content (> 90%, mainly (+)-limonene. Important for aroma are aldehydes, mainly decanal and citral, and aliphatic and terpenoid esters. The sesquiterpene aldehydes alpha-sinensal [17909-77-2] and beta-sinensal [6066-88-8] contribute particularly to the special sweet aroma (34). According to (30) the maximum observed concentration in sweet orange oil are (in %): *limonene* (95.0); *linalool* (0.7); n-decanal (0.7); *citral* (0.3); alpha-sinensal (0.05); beta-sinensal (0.06) (30). Worldwide production is more than 30000 tons / year. Main uses comprise the flavouring of beverages and confectioneries and perfuming E.d.C, soaps and household products.

For the latter uses relevant here, both "Orange peel oil, sweet (*Citrus sinensis* (L.) Osbeck) (8008-57-9)", "Orange peel, sweet, extract (*Citrus sinensis* L. Osbeck) (8028-48-6)" and "Orange, sweet, Valencia, ext. (97766-30-8)" are among the top 100 used fragrance materials and classified as R43 (IFRA, pers. comm. 2010).

ORANGE OIL TERPENES (CAS # 68647-72-3) are a "top 100 mixture of substances and classified as R43 (IFRA, pers. comm.2010). Other names: ORANGE, SWEET, TERPENES (REACH); Terpenes and Terpenoids, sweet orange-oil (REACH). The CAS entry refers to a group of substances "Terpenes and Terpenoids, sweet orange-oil" (REACH).

CITRUS TANGERINA ...

CAS 223748-44-5; EC /

Oil of tangerine

[no info in CAS database]

Current regulation: /

Clinical data:

In a 17 year old girl, the perfume used for 3 months caused ACD due to the ingredient "oil of tangerine", with a strong positive PT reaction (to 2% or 10% in pet.; 50 controls

negative) (226).

Additional information:

Citrus Tangerina Peel Oil is the volatile oil expressed from the peel of the ripe fruit the Tangerine, *Citrus Tangerina*, *Rutaceae* (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=55441>, last accessed 2010-01-29); (*Citrus tangerina* Tanaka).

CORIANDRUM SATIVUM HERB OIL

CAS 84775-50-8; EC 283-880-0
(Coriander, ext.)

Coriander oil

INCI: CORIANDRUM SATIVUM
EXTRACT

Current regulation: /

Clinical data:

The Rudzki 1976 study found 2 positive reactions in 200 patients to "coriander" essential oil, 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=3 (3.5%) positive reactions to "coriander" essential oil 2% pet. (27).

Additional information:

Coriander Sativum Herb Oil is an essential oil obtained from the herbs of the Coriander, *Coriandrum sativum* L., *Umbelliferae* (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=39388>, last accessed 2010-01-29). The main component of coriander oil is linalool (by GC: 65-78%) and mono- and polyunsaturated fatty aldehydes contributing to the particular aroma. In contrast to the seed oil, coriander leaf oil contains these aldehydes as main constituents, e.g. 2-deccanal and 2-dodecanal (34). For Oil of coriander fruits (*Coriandrum sativum* L.) an ISO standard exists: ISO 3516:1997.

CYMBOPOGON OILS

Cymbopogon oils are produced from several aromatic grasses that belong to the genus *Cymbopogon* Speng. (*Poaceae*). The oils are obtained by steam distillation of the aerial parts of the plants (34).

The composition of the essential oil derived from *Cymbopogon flexuosus* (Nees ex Steudel) J.F. Watson is defined by a standard: ISO 4718:2004, as is the oil derived from *Cymbopogon citratus*: 3217:1974.

CYMBOPOGON CITRATUS LEAF OIL

Cymbopogon citratus (DC.) Stapf.²⁴

CAS 89998-14-1; EC 289-752-0
(*Cymbopogon citratus*, ext. =
INCI)

²⁴ ISO 4720:2009 nomenclature

Lemon Grass Oil; Indian Verbena Oil; Indian Melissa Oil

CYMOPOGON SCHOENANTHUS OIL

Cymbopogon flexuosus (Nees ex Steudel) J.F. Watson²⁵

CAS 8007-02-1; EC 289-754-1 (oils, lemongrass) / 89998-16-3; EC 289-752-0 (Cymbopogon Schoenanthus, ext. = INCI)

Lemon Grass Oil

Current regulation: /

Clinical data:

The Frosch 2002 b study on 1606 consecutive patients reported 1.6% positive reactions to "lemongrass oil (East India), CAS 8007-02-1", PTed at 2% pet. (17). In a series of 40 of 744 consecutive patients PTed with an extended fragrance series (Sheffield 1999), 3 positive reactions to lemongrass oil were observed (3). In the Wöhrle 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded n=6 (0.8%) positive reactions to lemongrass oil 2% pet. (22). The IVDK 2010 c study identified 0.6% positive reactions in 2435 consecutively tested patients and 2.3% positive reactions in 8445 patients tested in the context of a special series (30).

Additional information:

Cymbopogon Citratus Leaf Oil is an essential oil obtained from the leaves of the Lemon Grass, *Cymbopogon citratus* (DC., ex Nees), *Poaceae*. It contains citral (75-85%), methylheptenone, citronellal, geraniol, limonene (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=39457>, last accessed 2009-11-12). According to Surburg/Panten, by GC: neral (31-40%), geranial (40-50%) (34).

Indian lemongrass oil is obtained by the so-called Indian variety of lemongrass, *Cymbopogon flexuosus* (Nees ex Steud.) Stapf. Content by GC: 25-35% neral, 35-47% geranial (34).

Cymbopogon Schoenanthus Oil is the volatile oil obtained by the steam distillation of fresh Lemon Grass, *Cymbopogon schoenanthus* (L.), *Poaceae* (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=75419>, last accessed 2009-11-12). According to (30) the maximum observed concentration in lemongrass oil are (in %): citral (85.0); geraniol (7.0); limonene (4.0); geranyl acetate (2.2); caryophyllene (1.6); trans-isocitral (1.4); 6-methyl 5-hepten-2-one (1.3); caryophyllene oxide (1.2); 4-nonanone (1); citronellol (0.8); eugenol (0.3); linalool (0.2) (also according to (227))

In a LLNA study by RIFM, the lemongrass oil as used was reported to contain 68.8% citral, 6.7% limonene, 6.1% geraniol, 2.2% geranyl acetate, 1.6% caryophyllene, 1.4% trans-isocitral, 1.3% 6-methyl 5-hepten-2-one, 1.2% caryophyllene oxide and 1% 4-nonanone, according to analyses of the supplier. The EC3 value was calculated to be 6.5% (227).

CYMOPOGON MARTINI HERB EXTRACT

CAS 84649-81-0; EC 283-461-2 (Cymbopogon Martini, ext)

²⁵ ISO 4720:2009 nomenclature

INCI: CYMBOPOGON MARTINI OIL*Palmarosa oil*

Current regulation: /

Clinical data: /

Additional information:

ISO 4720:2009 nomenclature: *Cymbopogon martini* (Roxb.) Will. Watson var. *motia* and var. *sofia*. Cymbopogon Martini Herb Extract is an extract obtained from the herbs of the plant, *Cymbopogon martini*, Gramineae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=39460>, last accessed 2009-11-24), namely, by steam distillation of wild or cultivated *Cymbopogon martini* (Roxb.) J.F. Wats., collected when in blossom (34). The main constituent is geraniol (72-94%) (34).

In a LLNA study by RIFM, the palmarosa oil as used was reported to contain 79.4% geraniol, 9.4% geranyl acetate and 1.9% caryophyllene, according to analyses of the supplier. The EC3 value was calculated to be 9.6% (227).

CYMBOPOGON NARDUS HERB OILCAS 89998-15-2; EC 289-753-6 (*Cymbopogon nardus*, ext. = INCI)*Citronella Oil (Sri Lanka)***CYMBOPOGON WINTERIANUS HERB OIL**CAS 91771-61-8; EC 294-954-7 (*Cymbopogon Winterianus*, ext. = INCI)*Citronella Oil (Java)*

Current regulation: ...

Clinical data:

The Rudzki 1976 study found 5 positive reactions in 200 patients to "citronella" essential oil 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=1 (1.1%) positive reactions to "citronella" essential oil 2% pet. (27).

Additional information:

ISO 4720:2009 nomenclature: *Cymbopogon nardus* (L.) W. Watson var. *lenabatu* Stapf. and *Cymbopogon winterianus* Jowitt, respectively. Cymbopogon Nardus Herb Oil is an essential oil obtained from the herbs of the plant, *Cymbopogon* (syn: *Andropogon*) *nardus* (L.), Gramineae. The Ceylon citronella oil contains geraniol (about 60%), citronellal (about 15%), camphene, limonene, linalool, borneol. According to Surburg/Panten, the Sri Lankan oil contains citronellal (3-6%), borneol (4-7%), citronellol (3-8.5%), geraniol 15-23% and methyl isoeugenol (7.11%) (34).

The Java citronella oil contains 25-50% citronellal, 25-45% geraniol (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.h.details&id=39469>, last accessed 2009-11-24). Cymbopogon Winterianus Herb Oil as a synonym for Java citronella oil is obtained from the herbs of the plant, *Cymbopogon winterianus*, Gramineae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=sea>

[rch.details&id=39472](#), last accessed 2009-11-24). This oil, produced in Taiwan and Java, contains citronellal (31-40%), geraniol (20-25%), citronellol (8.5-14%), geranyl acetate (2.5-5.5%), citronellyl acetate (2-4%) and many minor components. Annual worldwide production is currently at around 1000 t (34). For Oil of citronella, Sri Lankan type (*Cymbopogon nardus* (L.) W. Watson var. *lenabatu* Stapf.) an ISO standard exists: ISO 3849:2003, for Oil of citronella, Java type the ISO 3848:2001.

In a LLNA study by RIFM, the citronella oil as used was reported to contain 36.6% citronellal, 20.6% geraniol, 4.1% limonene, 3.7% geranyl acetate, 3.0% citronellyl acetate, 2.6% elemol, 2.2% beta-bourbonene, 1.9% delta-cadiene, 1.6% isopugenol I, 1.4% germacrene D and eugenol and linalol at < 1%, according to analyses of the supplier. The EC3 value was calculated as > 50 % (227).

Considering the content of well-known allergenic compounds, this essential oil is regarded as established contact allergen in humans.

EUCALYPTUS SPP. LEAF OIL

CAS 92502-70-0; EC 296-357-7
(*Eucalyptus*, ext. = INCI)

Eucalyptus Oil

CAS 8000-48-4; EC / (Oils,
eucalyptus) INCI: *EUCALYPTUS*
GLOBULUS OIL

Current regulation: /

Clinical data:

In a study with 218 fragrance sensitive patients, 1.8% reacted positively to 10% eucalyptus oil (pet.) (1). In a series of 40 of 744 consecutive patients PTed with an extended fragrance series (Sheffield 1999), 1 positive reaction to "eucalyptus oil" was observed (3). In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded n=4 (0.6%) positive reactions to eucalyptus oil 2% pet. (22). The Rudzki 1976 study found 3 positive reactions in 200 patients to "eucalyptus" essential oil 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=1 (1.1%) positive reactions to "Eucalyptus" essential oil 2% pet. (27). The IVDK 2010 c study identified 0.2% positive reactions in 6680 patients tested in the context of a special series (30).

In a professional athlete, the use of an "analgesic and anti-inflammatory cream" over 2 years lead to ACD, which was attributed to eucalyptol (eucalyptus oil, 1% pet., 25 controls negative), the sole ingredient of the cream eliciting a positive PT reaction (228)

Additional information:

ISO 4720:2009 nomenclature: *Eucalyptus globulus* Labill. Eucalyptus oils are produced from plants belonging to the genus *Eucalyptus* (*Myrtaceae*), which includes about 500 species in Australia, the country of origin, alone. At present, few of the oils, which are used to characterise species, are commercially important (34). Some species are rich in 1,8-cineole (80-85% content). Other species contain less cineole, but 10-22% alpha-pinene. *E. citriodora* predominantly contains citronellal (min. 75% by GC), with some citronellol and isopulegol (5-10% each) (34). *E. dives* contains (-)-piperitone and 15-25% alpha-phellandrene (34). According to (30) the maximum observed concentration in eucalyptus oil are (in %): 1,8-cineole (58; 70-80 after rectification); α -pinene (22); limonene (8); para-cymene (5); trans-pinocarveol (5); aromadendrene (10); globulol (2.5) [the latter 2 components only traces after rectification] (30).

For Crude or rectified oils of *Eucalyptus globulus* (*Eucalyptus globulus* Labill.) an ISO standard exists: ISO 770:2002.

It is a "top 100" substance and classified as R43 (IFRA, pers. comm.2010).

| | |
|---|-----------------------------------|
| EUGENIA CARYOPHYLLUS LEAF / FLOWER OIL | CAS 8000-34-8; EC / (Oils, clove) |
| <i>Clove oil</i> | INCI: EUGENIA CARYOPHYLLUS OIL |

Current regulation: /

Clinical data:

In the Larsen 2002 c study, 19.3% of patients with known contact allergy to fragrance ingredients reacted positively to "clove bud oil" (10 % pet.) (1). In a series of 40 of 744 consecutive patients PTed with an extended fragrance series (Sheffield 1999), 2 positive reactions to "clove oil" were observed (3). The Coimbra 2000 study found in 67 patients with positive reaction to the FM I who were tested with clove oil (2% pet.) 13.4% positive reactions (9). In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded 1.6% positive reactions 2% pet. (22). The Rudzki 1976 study found 2 positive reactions in 200 patients to "clove" essential oil, 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=12 (13.3%) positive reactions to "clove" essential oil 2% pet. (27). The IVDK 2010 c study identified 1.5% positive reactions 6893 patients tested in the context of a special series (30).

A 32 year old Spanish physiotherapists developed vesicular hand dermatitis after using a "balsam from ash extract" cream. PTing revealed positive reactions to this cream, the FM I, eugenol, and 2 components of the cream: cinnamon oil (0.5% pet.) and clove oil (1% pet.) (225).

Additional information:

ISO 4720:2009 nomenclature: *Syzygium aromaticum* (L.) Merr. & L. M. Perry syn. *Eugenia caryophyllus* (Spreng.) Bullock & S. G. Harrison. Standards regarding the composition of clove oil are available: ISO 3141:1997, ISO 3142:1997, ISO 3143:1997. Clove oils are produced from the clove tree *Syzygium aromaticum* (L.) Merr. et L.M. Perry [*Eugenia caryophyllus* (Spreng.) Bullock ex S.G. Harrison. The content of clove bud, clove leaf and clove stem oil has, with little variation, been determined by GC as 75-92% eugenol, 2-17% caryophyllene and 0.2-15% eugenyl acetate – the latter compound found in particularly high concentration in bud oil (34). According to another source, the following maximum content (%) has been observed regarding the constituents listed: eugenol (92,0);

caryophyllene (17); eugenyl acetate (15); isoeugenol (0.5) (30).

In a LLNA study by RIFM, the clove leaf oil as used was reported to contain 85.3% eugenol, 9.9% caryophyllene and 2.2% alpha humulene, according to analyses of the supplier. The EC3 value was calculated to be 7.1% (227).

EVERNIA FURFURACEA LICHEN EXTRACT

CAS 90028-67-4; EC 289-860-8
(*Evernia furfuracea*, ext. = INCI)

Tree moss extract

Current regulation: /

Clinical data:

The Larsen 1977 study in 20 "perfume-sensitive patients" yielded n=6 positive reactions with "treemoss abs. in benzyl benzoate, 5% petrolatum" (18). In the IVDK 2007 study, 2.7% (95% CI: 2.0 – 3.6%) of 1658 consecutive patients had a positive reaction to "tree moss absolute" (4). In the Groningen 2009 study, 2.5% (95% CI: 1.1 – 4.9%) had positive reactions to the allergen, tested at 2%, i.e., twice the commonly used concentration, and not in pet., but in diethylphthalate (6). The IVDK 2010 study, 6.02% (95% CI: 4.90 – 7.14%; percentages standardised for age and sex) of 1947 patients PTed reacted to the compound (7).

Additional information:

Syn.: *Pseudoevernia furfuracea* (L.) Zopf (53). The lichen grows on the bark of pine and fir trees. The extraction process with carbohydrate solvents yields a "concrete" (2-5% yield) which, in a next step eliminating waxy compounds, is extracted with warm alcohol and subsequent cooling, yielding an "absolute" (40-60% yield) (53).

EVERNIA PRUNASTRI

CAS 90028-68-5; EC 289-861-3
(*Evernia prunastri*, ext. = INCI)

Oak moss abs.

Current regulation: Annex III, part 1, n° 91

Clinical data:
In the "background information" section of the 1999 opinion, oak moss extract is classified as "most frequently reported allergen"; in consecutive PT patients, about 2.8% positive reactions had been reported (33). 'The German MAK commission has labelled oak moss extract as 'sensitising to the skin' (229).

Since the last SCCNFP-opinion of 1999, a "polymer based method" was developed to reduce the natural content of these two compounds from around 1 - several percent to < 75 ppm for atranol and < 25 ppm for chloratranol. However, PTing 14 subjects with previous positive PT reactions to the "oak moss" allergen preparation with the modified *Evernia prunastri* material still elicited positive reactions in 8/14 subjects; thus, the reduction in allergen content was deemed unsafe for the consumer (230). In a study of 885 consecutive eczema patients tested in Gentofte, Denmark, 3.2% had a positive or follicular patch test response to oak moss absolute. Two types of oak moss absolute were tested, one contaminated by resin acids and one without any detectable resin acids. There was no difference in reactivity between the two types of oak moss absolute

(231). The IVDK 2007 study yielded 2.2% (95% CI: 1.6 – 3.0%) positive reactions in 2063 consecutively tested patients (4). In the Groningen 2009 study, 1.9% (95% CI: 0.7 – 4.0%) had positive reactions to oak moss, tested at 2% pet., i.e., twice the commonly used concentration (6). In the An 2005 study, 6 of 422 consecutive patients, i.e., 1.4%, had positive reaction (13) (test concentration 2% pet.). In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded 5.0% positive reactions (22). The IVDK 2010 study, 1.81% (95% CI: 1.07 – 2.56%) of 1213 consecutively tested patients reacted to the compound, while 5.59% (95% CI: 4.90 – 6.27%) of 4482 of patients tested in a more aimed manner, partly as breakdown testing to the FM I, had a positive PT reaction (7). In a study from Alicante, Spain, 86 selected patients were tested with *E. prunastri* extract, yielding 2 positive reactions (48).

L. Kanerva et al. report on a 41 year old female hairdresser in whom oak moss abs. contained in a perming solution (concentration in the product unknown) was unequivocally identified as allergen causing (i) occupational hand dermatitis and (ii) scalp dermatitis after application to the own hair (232). Another case of occupational hand dermatitis in a grinding engineer was, at least partly, attributable to contact sensitisation to "oak moss resin" contained in a soluble oil (233).

Additional information:

Source: *Evernia prunastri* (Oak moss) (*Evernia prunastri* var. *prunastri* L. Ach). Oak moss is extracted as described above. Chloratranol and atranol are the degradation products of chloratranorin and atranorin, resp., which are recognised as the main sensitisers in *Evernia prunastri* extracts.

ILLICIIUM VERUM FRUIT OIL

CAS 84650-59-9, 8007-70-3; EC 283-518-1

"Anise Oil", Star anise oil

(Star anise, *Illicium verum*, ext. = INCI)

Current regulation: /

Clinical data:

In a study involving 100 consecutive patients, Rudzki and Grzywa found (i) a relatively high frequency of active sensitisation to star anise oil (n=5) tested with 0.5, 1 and 2% concentration (most likely in yellow petrolatum, as the other allergens in this series). Later patch testing with constituents of this essential oil (1%) in 3 patients yielded positive results to anethole in 3 cases, and to alpha-pinene and safrole in the 1 case tested to these substances. 34% of the consecutive patients reacted positively to star anise oil at 1%, which was considered as (marginally) non-irritating PT concentration (234).

Additional information:

ISO 4720:2009 nomenclature: *Illicium verum* Hook. f. *Illicium Verum* Fruit Oil is an essential oil distilled from the fruits of the Star Anise, *Illicium verum*, Illiciaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=40297>, last accessed 2010-01-29). The main component is trans-anethole (86-93%), which can be purified from star anise oil. Main uses are alcoholic beverages, food flavouring and oral care products (34, 39). For Oil of star anise, Chinese type (*Illicium verum* Hook. f.) an ISO standard exists: ISO 11016:1999.

JASMINUM GRANDIFLORUM FLOWER EXTRACT CAS 84776-64-7; EC 283-993-5
(Jasmine, *Jasminum grandiflorum*, ext. = INCI)

Jasmine abs.

JASMINUM OFFICINALE FLOWER OIL CAS 90045-94-6; EC 289-960-1
(Jasmine, *Jasminum officinale*, ext. = INCI)

JASMINUM OFFICINALE OIL CAS 8022-96-6; EC / (Oils, jasmine) INCI: JASMINUM OFFICINALE OIL

Current regulation: /

Clinical data:

In the Frosch 2002 b study, a total of 1.2% of 1606 consecutive patients had a positive PT to "jasmine absolute", tested 5% in pet. (17). The deGroot 2000 study yielded 13 positive reactions to "jasmine, synthetic" in 1825 consecutively tested patients (12). In the early Larsen 1977 study, 18 of 20 "perfume sensitive patients" reacted to "Jasmin synthetic" 10% pet. (18), while 7 reacted to "Jasmin absolute" (10% pet.) – all of these also positive to the synthetic fragrance. The Sugiura 2000 study set in Nagoya, Japan, yielded 1% positive PT reactions in 1483 patients PTed for suspected cosmetic dermatitis, using 5% pet. as test concentration (14). The Larsen 2001 study in 178 patients with known contact allergy to fragrance ingredients found 16.9% positive reactions to jasmine absolute (10% pet.) (19). In the An 2005 study, 5 of 422 consecutive patients, i.e., 1.2%, had a positive reaction to Jasmine officinale oil (Jasmine absolute, Egyptian), tested at 2% (13). In the NACDG 2009 study, 1.1% of 4447 patients tested with "Jasmine absolute 2% pet." were found PT-positive (21). The Belsito 2006 study (20) yielded 0.4% positive reactions to "jasmine absolute". The Goossens 1997 study found 5 of 111 patients positive to "jasmine absolute" (10% pet.)– all sensitised to other fragrance allergens (23). In 63 patients positive to the FM I, 13 had positive PT reactions to "jasmine absolute", 2% pet., and 12 to "jasmine synthetic", 2% pet. in the Santucci 1987 study – the amount of concomitant reactivity was not examined (28). Nakayama et al. found 1974 (after (29)) 19 "strong positive" and 25 "weak positive" reactions to "jasmin oil" (unknown test concentration) in 183 patients. The IVDK 2010 c study identified 1.5% positive reactions in 3668 consecutively tested patients and 1.2% positive reactions in 982 patients tested in the context of a special series (30). In a study from Alicante, Spain, 86 selected patients were tested with jasmine absolute, yielding 3 positive reactions, and with "Jasmine synthetic", also resulting in 3 positive reactions (48).

Additional information:

Jasminum Grandiflorum Flower Extract is an extract obtained from the flowers of the Spanish Jasmine, *Jasminum grandiflorum* L., Oleaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=39752>, last accessed 2009-11-12).

Jasminum Officinale Flower Oil is an essential oil obtained by molecular distillation of the flowers from the Jasmine, *Jasminum officinale* L., Oleaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=39754>, last accessed 2009-11-25).

Jasminum Officinale Oil is the volatile oil obtained from the flowers of the Jasmine, *Jasminum officinale* L., Oleaceae

(<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=34776>, last accessed 2010-01-05); this latter extract is used by Almirall/Hermal/Trolab for the preparation of a PT allergen.

Jasmine absolute is obtained by solvent extraction, via concrete, from the flowers of *J. grandiflorum* (L.) Aiton from China and India. The main volatile compound is benzyl acetate, however, minor compounds such as indole [120-72-9], cis-jasmone [488-10-8] and methyl jasmonate [1211-29-6] contribute to the typical jasmine fragrance (34). Reported compounds include the following (maximum observed concentration given in parentheses): benzyl acetate (28); benzyl benzoate (24.0); phytol acetate (9); isophytol (8.5); phytol (7.4); linalool (7.0); eugenol (4.0); squalene (4); indole (3.5); benzyl alcohol (2.5); cis-jasmone (2.5); methyl linolenate (2.0); methyl palmitate (1.4); p-cresol (1.0); cis-3-hexenyl benzoate (1.0); benzyl salicylate (0.4); jasmin lactone (0.9); methyl jasmonate (0.7); isoeugenol (0.4) ((30), also according to (17))

JUNIPERUS VIRGINIANA OIL

CAS 8000-27-9; EC / (Oils, cedarwood) [this also refers to *Cedrus atlantica* ...] / 85085-41-2; EC 285-370-3 (Juniper, *Juniperus virginiana*, ext. = INCI)

JUNIPERUS VIRGINIANA WOOD OIL

CAS 85085-41-2; EC 285-370-3

Cedar Wood Oil (Virginian)

Current regulation: /

Clinical data:

In the Frosch 2002 b study, a total of 0.6% of 1606 consecutive patients had a positive PT to "cedarwood oil (Moroccan and Chinese 1:1)", tested 10% in pet. (17). After application of Penaten-baby™ oil as immersion oil for dermatoscopy a patient developed multiple patches of eczema at the application sites. Investigation revealed that the oil was kept in a bottle previously used for *Juniperus virginiana* oil, to which contact sensitisation was verified by patch testing (235).

Additional information:

ISO 4720:2009 nomenclature: *Juniperus virginiana* L.. *Juniperus Virginiana* Oil is the volatile oil obtained from the fruits and leaves of the Red Cedar, *Juniperus virginiana* L., Cupressaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=78070>, last accessed 2010-01-05)

Juniperus Virginiana Wood Oil is an essential oil obtained from the wood and twigs of the Red Cedar, *Juniperus virginiana* L., Cupressaceae. It contains chiefly (alpha and beta) cedrene and cedral (cedar camphor), cuparene, thujopsene, widdrol (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=39767>, last accessed 2009-11-12)(235). According to Surburg/Panten by GC: alpha-cedrene 22-35%, thujopsene 10-25%, cedrol 16-25% (34).

See also *Cedrus atlantica*. According to (30) the maximum observed concentration in cedar wood oil are (in %): α-cedrene (32); thujopsene (25); cedrol (25); β-cedrene (6); widdrol (5) and cuparene (traces) (30).

For Oil of cedarwood, Virginian (*Juniperus virginiana* L.) an ISO standard is available: ISO 4724:2004. For Oil of cedarwood, Texas (*Juniperus mexicana* Schiede) an ISO standard exists: ISO 4725:2004.

LAURUS NOBILIS OIL

CAS 8002-41-3; EC / (Oils, laurel)
 INCI: LAURUS NOBILIS OIL /
 8007-48-5; EC / (Oils, sweet
 bay)/ 84603-73-6; EC 283-272-5
 (Laurus nobilis, ext.) INCI:
 LAURUS NOBILIS EXTRACT

Laurel oil

Current regulation: Annex II, n° 359 (seed oil)

Clinical data:

In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded n=4 (0.6%) positive reactions to "laurel oil" 2% pet. (22).

After sensitisation by a one-time occlusive application a 36 year old Turkish patient developed widespread allergic contact dermatitis 3 days after massage with olive oil containing *Laurus nobilis* oil; sensitisation was proven by a strong positive reaction to the commercial test preparation and the massage oil previously used (236). Topical application of laurel oil for knee arthropathy led to an erythema exudativum multiforme-like rash on the legs of a 63 year old patient; interestingly, laurel oil yielded a "target like" strongly positive PT reaction in this case (237). In an earlier Turkish case with a similar history, the EEM-like appearance was lacking; however, a very intense, edematous reaction was noted (238). In a series of 40 of 744 consecutive patients PTed with an extended fragrance series (Sheffield 1999), 2 positive reactions to "laurel oil" were observed (3). The IVDK 2010 c study identified 1.0% positive reactions in 6297 patients tested in the context of a special series (30).

Additional information:

ISO 4720:2009 nomenclature: *Laurus nobilis* L. Laurel leaf oil is obtained by steam distillation of leaves from *Laurus nobilis* L. (Lauraceae), an evergreen cultivated primarily in the Mediterranean countries. The main components are 1,8-cineole (30-70%), linalool (about 10%) and eugenol (34). According to (30) the maximum observed concentration in laurel oil are (in %): 1,8-cineole (70); β -caryophyllene (11); linalool (11); limonene (5.0); eugenol (2.0); geraniol (0.3) (30).

LAVANDULA HYBRIDA HERB OIL

CAS 91722-69-9; EC 294-470-6
 (Lavender, *Lavandula hybrida*,
 ext. = INCI)

Lavandin Oil

Current regulation: /

Clinical data:

The Rudzki 1976 study found 1 positive reaction in 200 patients to "lavandin" essential oil, 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=4 (4.6%) positive reactions to "lavandin" essential oil 2% pet. (27). In the Frosch 1995 dose-finding pilot study, no positive reaction to 1% and 5% lavandin oil in pet., tested in 205 consecutive patients in Dortmund and Göttingen, and just 1 irritant reaction to

the higher concentration, were observed (15).

Additional information:

ISO 4720:2009 nomenclature: *Lavandula angustifolia* Mill. x *Lavandula latifolia* Medik. Lavandula Hybrida Herb Oil is an essential oil distilled from the flowering herbs of the Lavandin, *Lavandula hybrida*, *Labiatae* (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=39789>, last accessed 2010-01-29. Nomenclature according to Surburg/Panten: *Lavandula x intermedia* Lois, which is a hybrid of lavender and spike (see below) (34). The oils from the most important variants, abrial and grosso, contain linalool (24-38%), linalyl acetate (20-38%) as well as 1,8-cineole (4-11%), and camphor (6-11%) (34). A third variant is called super because of its high concentration of linalyl acetate (35-47%), more closely resembling lavender oil (34). For Oil of lavandin Grosso (*Lavandula angustifolia* Mill. x *Lavandula latifolia* Medik.), French type an ISO standard exists: ISO 8902:2009, for Oil of lavandin Abrial (*Lavandula angustifolia* Miller x *Lavandula latifolia* Medikus), French type a different ISO standard: ISO 3054:2001.

It is a "top 100" substance (IFRA, pers. comm.2010)

Considering the content of well-known allergenic compounds, this essential oil is regarded as established contact allergen in humans.

LAVANDULA OFFICINALIS FLOWER OIL

CAS 84776-65-8, 8000-28-0; EC 283-994-0 (*Lavender, Lavandula angustifolia angustifolia*, ext. = INCI)

Lavender oil

Current regulation: /

Clinical data:

In a large series from Nagoya, Japan, 1483 patients were tested with lavender oil 20% in pet., with overall 3.7% positive reactions from 1990 to 1998. However, within this period, a sharp increase was noted in 1997 and 1998, which as attributed to changed exposure by M. Sugiura et al. (14). On the individual level, relevance of positive reactions remained unclear in about half of the cases. The Coimbra 2000 study found in 67 patients with positive reaction to the FM I who were tested with "lavender absolute" (2% pet.) 6.6% positive reactions (9). In the An 2005 study, 5 of 422 consecutive patients, i.e., 1.2%, had positive reactions to "Lavandula augustifolia oil" (Lavender absolute) 2% (13). The Goossens 1997 study found 4 of 111 patients positive to "lavender oil 20% pet."- all of them sensitised to other fragrance allergens (23). The Rudzki 1976 study found no positive reaction in 200 patients to "lavender" essential oil, 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=3 (3.5%) positive reactions to "lavender" essential oil 2% pet. (27). Nakayama et al. found 1974 (after (29)) 6 "strong positive" reactions to "Lavender oil" (unknown test concentration) in 183 patients. In a study from Alicante, Spain, 86 selected patients were tested with "lavender absolute", yielding 2 positive reactions (48).

R. Goiriz et al. report on a case of photo contact allergy (10 controls negative) in a 45 year old woman developing after application of a ketoprofen-containing topical gel ("Fastum")(239). A physiotherapist developed acute, recurrent dermatitis after use of "Difflam® gel", scented with lavender oil. Both the gel and lavender oil (2% pet.) tested positive; avoidance resulted in clearing (240). In a study on 218 patients with known

contact allergy to fragrance ingredients, Larsen (2002 c) found positive reactions to 10% lavender oil (pet.) in 2.8% of these (1). A case of vulvovaginitis with spread and affecting the dominant hand applying various tea tree and lavender oil creams was reported by S. Varma; the PT with 10% lavender oil abs. in pet. (50 controls negative) was positive (241). In two cases, facial "pillow dermatitis" due to lavender oil, applied to the pillows, developed, confirmed by positive PT to lavender abs. (2% pet.) (242).

Additional information:

ISO 4720:2009 nomenclature: *Lavandula angustifolia* Mill. *Lavandula officinalis* Flower Oil is an essential oil obtained from the fresh flowering tops of the Lavender, *Lavandula officinalis* (syn: *L. vera*), *Labiatae*. It contains 30-40% esters calculated as linalyl acetate, linalool, pinene, limonene, geraniol, some eucalyptol (cineol) (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=40370>, last accessed 2009-11-09). According to Surburg/Panten, lavender oil is obtained by steam distillation of freshly cut flowering tops of *Lavandula angustifolia* Mill. (Lamiaceae). Main constituents according to GC are linalyl acetate (25-45%), cis-ocimene (4-10%), trans-ocimene (1.5-6%), 1,8-cineole ($\leq 1\%$) camphor ($\leq 0.5\%$), linalool (25-38%), 1-terpinen-4-ol (2-6%) and lavandulyl acetate [25905-14-0] ($\geq 2\%$) (34).

In addition to distillation, both *Lavandula officinalis* and Lavandin are also solvent extracted, yielding concretes and, after ethanol extraction, absolutes, which are said to have a longer-lasting odour (34).

For Oil of lavender (*Lavandula angustifolia* Mill.) an ISO standard exists: ISO 3515:2002.

LAVANDULA SPICA HERB OIL

CAS 97722-12-8; EC 307-762-6
(Lavender, *Lavandula spica*, ext.
= INCI

"Spike Oil"

Current regulation: ...

Clinical data:

The Rudzki 1976 study found 1 positive reaction in 200 patients to "spike" essential oil, 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=8 (9.3%) positive reactions to "spike" essential oil 2% pet. (27).

Additional information:

ISO 4720:2009 nomenclature: *Lavandula latifolia* Medik. *Lavandula Spica* Herb Oil is an essential oil distilled from the flowering herbs of the Spikenard, *Lavandula spica* (syn: *Lavandula latifolia*), *Labiatae*. It contains eucalyptol (35%), camphor, linalool, borneol, terpineol, D-camphene and sesquiterpenes (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=40372>, last accessed 2010-01-29). According to Surburg/Panten, Spanish spike lavender oil is steam distilled from the flowering tops of *Lavandula latifolia* Medik.. The main components are linalool (34-50%), 1,8-cineole (16-39%) and camphor (8-16%) (34). For Oil of spike lavender (*Lavandula latifolia* (L.f.) Medikus), Spanish type an ISO standard exists: ISO 4719:1999

Considering the content of well-known allergenic compounds, this essential oil is regarded as established contact allergen in humans.

LITSEA CUBEBA FRUIT EXTRACT

CAS 90063-59-5, 68855-99-2; EC 290-018-7 (*Litsea cubeba*, ext.)
 INCI: LITSEA CUBEBA OIL

Current regulation: ...

Clinical data:

The Rudzki 1976 study found 3 positive reaction in 200 patients to "Litsea cubeba" essential oil 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=7 (8.1%) positive reactions to this essential oil 2% pet. (27).

Additional information:

ISO 4720:2009 nomenclature: *Litsea cubeba* (Lour) Pers. Litsea Cubeba Fruit Extract is an extract obtained from the fruits of the plant, *Litsea cubeba*, Lauraceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=40036>, last accessed 2009-11-24. The content by GC is: neral (25-33%), geranial (38-45%) – i.e. about ¾ citral, for which the extract had previously served as a raw material (34); direct use for perfuming is limited to household products (39). For Oil of Litsea cubeba (*Litsea cubeba* Pers.) an ISO standard exists: ISO 3214:2000.

In a LLNA study by RIFM, the "Litsea cubeba oil" as used was reported to contain 85.7% citral, 2.9% limonene, 1.7% linalool, 1.4% citronellal and < 1% caryophyllene and methyl heptanone, according to analyses of the supplier. The EC3 value was calculated as 8.4 % (227).

Considering the content of well-known allergenic compounds, this essential oil is regarded as established contact allergen in humans.

MENTHA ARVENSIS LEAF OIL

CAS 68917-18-0 ; EC /

Cornmint oil

INCI: MENTHA ARVENSIS OIL

Current regulation: /

Clinical data: /

Additional information:

Mentha Arvensis Leaf Oil is the oil derived from the leaves of the Horse Mint, *Mentha arvensis* L., Labiatae (http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details_v2&id=57860)

It is a "top 200" substance and classified as R43 (IFRA, pers. comm.2010)

MENTHA PIPERITA OIL

CAS 8006-90-4; EC / (Oils, peppermint) INCI: MENTHA

PIPERITA OIL / 84082-70-2; EC
282-015-4 (Peppermint, ext.) INCI
names: MENTHA PIPERITA ...

Peppermint oil

Current regulation: /

Clinical data:

In the Frosch 2002 b study, 0.6% of 1606 consecutive patients reacted positively to "peppermint oil (American)", tested 2% in pet. (17). In a series of 40 of 744 consecutive patients PTed with an extended fragrance series (Sheffield 1999), 2 positive reactions to "peppermint oil" were observed (3). In the Wöhrl 2001 study, PTing 747 patients with suspected contact allergy to fragrance ingredients yielded n=1 (0.1%) positive reactions to peppermint oil 2% pet. (22). Among 512 patients referred from a dental department for diagnostic work-up of various intraoral symptoms and complaints within 4 years, 6 patients had positive (+ to +++) PT reactions to "peppermint oil" 1% pet. at D4, mostly combined with positive reactions to menthol (see above) and reporting dramatic improvement after cessation of use of peppermint-containing oral products (154). The Rudzki 1976 study found 1 positive reaction in 200 patients to "Peppermint" essential oil, 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=6 (6.9%) positive reactions to "peppermint" essential oil 2% pet. (27). In 63 patients positive to the FM I, 3 had positive PT reactions to peppermint oil, 2% pet., in the Santucci 1987 study (28). The IVDK 2010 c study identified 0.6% positive reactions in 6546 patients tested in the context of a special series (30).

An unusual case of "baboon-like" allergic contact dermatitis of the vulva after drinking excessive amounts of a herbal tea containing, among other ingredients, peppermint. While the PT reaction to peppermint oil was only weak to doubtful, dramatic improvement after cessation and prompt relapse after repeat ingestion proved the diagnosis (243). Recurrent foot and lower leg dermatitis after the application of a "foot spray" (containing peppermint oil) was diagnosed as allergic contact dermatitis due to this ingredient in a 59 year old golf player (244). In another case, ACD after application of a transdermal system for the treatment of lumbar pain was attributed to CA to peppermint oil (2% pet.) and its main ingredient menthol (1% pet.) (155). In a patient with toothpaste-induced cheilitis, not only *M. piperita*, but also *M. arvensis*, but not *M. spicata* or *cardica* extracts (all tested 1% pet.), as well as natural and synthetic menthol caused positive PT reactions (245).

Additional information:

ISO 4720:2009 nomenclature: *Mentha x piperita* L. A standard by ISO exists for Oil of peppermint (*Mentha x piperita* L.): ISO 856:2006. A review by the Cosmetic Ingredient Review Expert Panel, Washington, DC on the "Final report on the safety assessment of *Mentha Piperita* (Peppermint) Oil, *Mentha Piperita* (Peppermint) Leaf Extract, *Mentha Piperita* (Peppermint) Leaf, and *Mentha Piperita* (Peppermint) Leaf Water" is available (163), stating that "Peppermint Oil is used at a concentration of < or = 3% in rinse-off formulations and < or = 0.2% in leave-on formulations. Peppermint Oil is composed primarily of menthol and menthone. Other possible constituents include pulegone, menthofuran, and limone. According to Surburg/Panten: (-)-menthol (34-46%), (-)-menthone (15-27%), (-)-menthyl acetate (2.5-7%) and menthofuran [17957-94-7] (0.5-6%) (34). According to (30) the maximum observed concentration in peppermint oil are (in %): (-)-menthol (49); (-)-menthone (28); (-)-menthyl acetate (8); mentofuran (8); isomenthone (8); neo menthol (6); pulegone (3.5); limonene (3.0); linalool (0.4) (30). Most of the safety test data concern Peppermint Oil. The oil is considered to present the "worst case scenario" because of its many constituents, so data on the oil were considered relevant to the entire group of ingredients. ... Repeated

intradermal dosing with Peppermint Oil produced moderate and severe reactions in rabbits" concluding that "with the limitation that the concentration of pulegone in these ingredients should not exceed 1%, it was concluded that Mentha Piperita (Peppermint) Oil, Mentha Piperita (Peppermint) Extract, Mentha Piperita (Peppermint) Leaves, Mentha Piperita (Peppermint) Water are safe as used in cosmetic formulations".

MENTHA SPICATA HERB OIL

CAS 84696-51-5, 8008-79-5; EC 283-656-2 (Spearmint, ext.)

Spearmint oil

INCI: MENTHA VIRIDIS EXTRACT

Current regulation: /

Clinical data:

In the Frosch 2002 b study, 0.8% of 1606 consecutive patients reacted positively to "spearmint oil (American)", tested 2% in pet. (17). The CAS # quoted (8008-79-5) refers, according to CosIng, to MENTHA VIRIDIS LEAF OIL, the volatile oil obtained from the dried tops and leaves of the Garden Mint, Mentha viridis L., Labiatae. The Larsen 2001 study diagnosed 5.0% positive reactions in 178 patients with known contact allergy to fragrance ingredients, using this oil at 5% pet. test concentration (19). In the An 2005 study, 6 of 422 consecutive patients, i.e., 1.4%, had positive reactions to "Mentha viridis oil" 5% (13). PT results with toothpaste ingredients were positive in 7 patients, of whom 4 had strong positive reactions to spearmint (246).

Additional information:

ISO 4720:2009 nomenclature: Mentha spicata L. Mentha Spicata Oil is an essential oil obtained from the herbs of the Spearmint, Mentha spicata L., Labiatae (syn: Mentha viridis L., Labiatae). It contains carvone (more than 50%), limonene, pinene (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=40394>, last accessed 2009-11-11). According to Surburg/Panten, the content is limonene (9-16.5%), (-)-carvone (60-70%), menthone (0-0.2%) and viridiflorol (0-0.5%) (34). Exposure by toothpastes, and subsequent contact allergic reaction of the lips or the oral mucosa, have been reported (e.g., (247, 248)). L-Carvone is a component of the oil from Mentha spicata (spearmint) (53) and had been tested with positive results in "toothpaste cases", even at a concentration as low as 0.067% (68).

For Oil of spearmint -- Part 1: Native type (Mentha spicata L.) an ISO standard exists: ISO 3033-1:2005, for Oil of spearmint -- Part 2: Chinese type (80 % and 60 %) (Mentha viridis L. var. crispa Benth.), redistilled oil: ISO 3033-2:2005, for Oil of spearmint -- Part 3: Indian type (Mentha spicata L.), redistilled oil: ISO 3033-3:2005 and for Oil of spearmint -- Part 4: Scotch variety (Mentha x gracilis Sole): ISO 3033-4:2005.

MYROXYLON PEREIRAE RESIN

CAS 8007-00-9; EC 232-352-8 (Balsams, Peru)

Balsam of Peru

INCI: MYROXYLON PEREIRAE / Balsams, Peru

Current regulation: Annex III, part1, n° 154

Clinical data:

This natural mixture has been employed as screening agent in Baseline series worldwide for many decades. Hence, a wealth of data is available; table 3.2 – 1 summarises results of the past 10 years.

Additional information:

ISO 4720:2009 nomenclature: *Myroxylon pereirae* (Royle) Klotzsch, syn. *Myroxylon balsamum* var. *pereirae* (Royle) Harms. *Myroxylon pereirae* resin (MPR, Balsamum peruvianum) is harvested from the balsam of Peru tree, *Myroxylon balsamicum* (L.) HARMS var. *pereirae* (ROYLE) HARMS, synonymous *Myroxylon pereirae* (ROYLE) KLOTZSCH (249) after thermal stress, almost exclusively in El Salvador. Main constituents of the pleasantly, vanilla-like smelling dark brown liquid are benzyl esters of cinnamic and benzoic acid (35 – 75%), up to 30% cinnamic acid, up to about 10% benzoic acid, approximately 5% alpha- and beta-nerolidol, benzyl alcohol and mostly less than 1% cinnamyl alcohol, benzyl ferulate and -isoferulate, cinnamic acid amyl ester, coniferyl alcohol, coniferyl benzoate, eugenol, isoeugenol, farnesol, vanillin, and several trace constituents (250-253). The composition of MPR varies with the origin and other factors; moreover, MPR is sometimes blended with other natural mixtures such as turpentine, styrax or colophonium (249).

MPR can be used to improve taste or smell in gargling solutions, cosmetic products such as soaps, shampoo or lipsticks, as well as sweets, tobacco and beverages (249, 254). According to EU legislation and IFRA guidelines MPR should not be used in products intended for skin contact; however, extracts and distillates of MPR may be used in a concentration of < 0,4% (IFRA-Guidelines, www.ifraorg.org (255)). E. Temesvári et al. report on the interesting case of severe ACD with subsequent hypopigmentation after a “temporary henna tattoo”, which was, unexpectedly, not due to p-phenylene diamine, but to the oil used to disperse the pigment, which presumably contained allergens also included in the FM I and MPR, both of which were extreme positive on a later PT (256).

In addition to delayed type hypersensitivity reactions, MPR (and some of his constituents such as benzoic acid (257)) are capable of eliciting (non-immunological) urticarial immediate reactions (258-260). In one case, the immediate reaction to MPR (and to FM I) at the test site spread systemically in terms of a generalised urticaria, while no delayed type reactions were observed to the PT (261). Generally, there is apparently no association of immediate reactions to MPR (and cinnamal or cinnamyl alcohol) and contact sensitisation to these compounds (262). In animal experiments the sensitising potency of MPR was clearly established (250), with coniferyl benzoate identified as single compound with the most marked potency (252). However, due to the limited chemical stability of this compound is unclear whether other, more stable compounds are, in fact, more important allergens, such as cinnamic acid and (iso-) ferulic acid esters or oxidised constituents of the resin fraction (263).

Table 3.2.2 – 1: Results with contact allergy to fragrance ingredients screening agents reported since 1999 in patients patch tested for suspected allergic contact dermatitis: **Myroxylon pereirae resin** (Balsam of Peru) 1). If not given in the publication, the confidence interval (CI) was calculated from the absolute numbers by the reviewers.

| Country | Population | Years | No. tested | Crude % positive (95% CI) § |
|--------------------------|----------------------|-------------------|------------|-----------------------------|
| Tel Aviv, Israel (264) # | Consecutive patients | 1999-2000 | 943 | 6.6 % (5.1 – 8.4) § |
| South Korea (13) | Consecutive patients | 04/2002 – 06/2003 | 422 | 7.3% (5.1 – 10:3%) § |

Opinion on fragrance allergens in cosmetic products

| | | | | |
|---|------------------------------------|-----------|-------|---------------------|
| Tel Aviv, Israel (265) | Consecutive patients | 1998-2004 | 2156 | 3.6 (2.9 – 4.5) § % |
| Manipal, India (266) | Dermatitis patients | 1989-1998 | 1780 | n=17 |
| Tehran, Iran (267) | Consecutive patients | 2002-2004 | 250 | 2.4 (0.9 – 5.2) § % |
| Sevilla, Spain (268) | Consecutive patients | 2002-2004 | 863 | 5.8 (4.3 – 7.6) § % |
| Ankara, Turkey (269) | Consecutive patients | 1992-2004 | 1038 | 2.1 (1.3 – 3.2) § % |
| Vienna, Austria (22) | Consecutive patients of one clinic | 1997-2000 | 2660 | 5.4% (4.6 – 6.3%) § |
| Czech Republic (270) | Consecutive patients | 1997-2001 | 12058 | 7.3% (6.8 – 7.8) § |
| Copenhagen, Denmark (271) | Consecutive patients | 1985-2007 | 16173 | 3.9 (3.6 – 4.2) § % |
| Sweden (272) | Consecutive patients | 2000 | 3790 | 6.5% |
| 9 European countries (273) § | Consecutive patients | 2002-2003 | 9672 | 6.1 % |
| Germany, 3 Swiss + 1 Austrian Dept. (7) | Consecutive patients | 2005-2008 | 36919 | 8.0% (7.7 – 8.3%) |
| 10 depts. From 7 EU countries (274) * | Consecutive patients | 1996-2000 | 26210 | 6.0 % |
| USA (Canada) (20) | Probably consecutive patients | 2003 | 1603 | 6.6% |
| NACDG 2009 (21) | Consecutive patients | 2005-2006 | 4449 | 11.9% |

§ Calculated by reviewers, where possible (if actual numbers were given)

Probably included in (265)

\$ > 5-fold difference between departments

* About 4-fold difference between departments

NARCISSUS SPP. EXTRACT / OIL

CAS: diverse

Narcissus abs.

Current regulation: /

Clinical data:

In the Frosch 2002 b study, 1.3% positive reactions to "narcissus absolute" (2% pet.) were observed in 1606 consecutive (17). The extract used by the PT allergen provider Almirall/Hermal/Trolab has the CAS number 90064-25-8. The IVDK 2010 c study identified 0.5% positive reactions in 2445 consecutively tested patients and 0.6% positive reactions in 809 patients tested in the context of a special series (30).

Additional information:

Commonly used: *Narcissus poeticus* L. According to (30) the maximum observed concentration in Narcissus abs. are (in %): α -terpineol (23.7); trans-Isoeugenol methyl ether (20); benzyl benzoate (20); coumarin (5.7); benzyl alcohol (4.0); Δ^3 -carene (3.4); cinnamyl alcohol (2.5); phenylethyl alcohol (2.2); ethyl palmitate (2.2); phenylpropyl acetate (1.7); 1,8-cineole (1.5); caryophyllene (1.0); benzyl acetate (0.7); isoeugenol (0.5); farnesol (0.3) (also according to (17)) (30).

OCIMUM BASILICUM HERB OIL

CAS 84775-71-3; EC 283-900-8 (*Ocimum basilicum*, ext. = INCI)

Basil Oil (sweet)

Current regulation: /

Clinical data:

/

Additional information:

ISO 4720:2009 nomenclature: *Ocimum basilicum* L. For Oil of basil, methyl chavicol type (*Ocimum basilicum* L.) an ISO standard exists: ISO 11043:1998. *Ocimum Basilicum* Herb Oil is an essential oil obtained from the herbs of the Sweet Basil, *Ocimum basilicum* L., *Labiatae*. (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=40474>, last accessed 2009-11-24). The chemical composition varies greatly with the origin (34):

- Basil oil of the methylchavicol type (Réunion type) is extracted from flowering tops or whole plants from Réunion, Comores, Madagascar, but also other countries such as Egypt. Mainly used for seasoning food. Content by GC: methylchavicol 75-87%, linalool 0.5-3%
- Basil oil, linalool type is produced mainly in the Mediterranean area. Content by GC: Linalool 45-62%, methylchavicol trace to 30%, eugenol 2-15%
- Indian Basil oil is produced exclusively in India. Content by GC: methylchavicol trace to 70%, linalool 25%.

In a LLNA study by RIFM, the basil oil as used was reported to contain 51% linalool, 10.4% eugenol, 7.7% cineol, 3.7% bergamotene, 2.7% germacrene D, 2.7% cadinol and 1.3% cadinene, according to analyses of the supplier. The EC3 value was calculated to be < 2.5% (227).

PELARGONIUM GRAVEOLENS FLOWER OIL

CAS 90082-51-2; EC 290-140-0 (*Pelargonium graveolens*, ext. = INCI) / 8000-46-2; EC / (Oils, geranium) INCI: GERANIUM

Geranium Oil Bourbon

 Current regulation: /

Clinical data:

The Coimbra 2000 study found in 67 patients with positive reaction to the FM I who were tested with "geranium oil Bourbon" (2% pet.) 7.4% positive reactions (9). In the Larsen 2001 study, 8.4% positive reactions were observed in 178 patients with known contact allergy to fragrance ingredients ("geranium oil Bourbon", 10% pet.) (19). The Goossens 1997 study found 3 of 111 patients positive to "geranium oil 20% pet." – all sensitised to other fragrance allergens (23). The Rudzki 1976 study found 3 positive reactions in 200 patients to "geranium" essential oil 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=2 (2.3%) positive reactions to "geranium" essential oil 2% pet. (27). Nakayama et al. found 1974 (after (29)) 3 "strong positive" reactions to "Geranium oil" (unknown test concentration) in 183 patients, Trattner/David 1 / 641 consecutive patients positive to "Geranium oil" (31). In a study from Alicante, Spain, 86 selected patients were patch tested with an extended fragrance series; n=8 reacted positively to geranium oil bourbon (48).

Additional information:

ISO 4720:2009 nomenclature: *Pelargonium x ssp.* For Oil of geranium (*Pelargonium X ssp.*) an ISO standard exists: ISO 4731:2006 *Pelargonium Graveolens Flower Oil* is the volatile oil obtained from the flowers of the Bourbon Geranium, *Pelargonium graveolens* L. Hér. Ex Aiton, *P. roseum* Willdenow (and other nondefined hybrids that have developed in different regions of the world) Geraniaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=57527>, last accessed 2009-11-16)(34). The Bourbon type (Réunion, Madagascar) is more valuable than the North African and Chinese products, and differs in characteristic components: (-)-6,9-guaiadiene [36577-33-0] 5-9% in the Bourbon type, and 10-epi-gamma-eudesmol [15051-81-7] 3-6% in the African type, in addition to the main components (-)-citronellol, isomenthone, formates and tiglates. Chinese oil is similar to Bourbon oil, however, it contains more citronellol (32-43%) and lower amounts of linalool (2-4.5%) and geraniol (5-12%) (34).

In a LLNA study by RIFM, the geranium oil as used was reported to contain 41.1% citronellol, 9.8% 2,6-guiadine, 6.2% isomethone, 4.9% geraniol, 2.2% cis-rose oxide, 2.1% linalool, 1.5% geranyl formate, 1.3% phenyl ethyl tiglate, 1.0% trans-rose oxide, and geranyl tiglate and alpha-pinene at < 1%, according to analyses of the supplier. The EC3 value was calculated to be > 50% (227).

PELARGONIUM ROSEUM LEAF OIL

CAS 90082-55-6; EC 290-144-2
(*Pelargonium roseum*, ext. =
INCI)

Geranium Oil; Rose Geranium Oil

 Current regulation: /

Clinical data:

In the Sugiura 2000 study, among 1483 patients with suspected cosmetic dermatitis, 2.1% positive PT reactions to "geranium oil" (tested 20% in pet.) were observed (14).

Additional information:

Pelargonium Roseum Leaf Oil is an essential oil obtained from the leaves of the plant, Pelargonium roseum, Geraniaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=40565>, last accessed 2009-11-16).

PIMENTA RACEMOSA LEAF/FRUIT OIL

CAS 85085-61-6; EC 285-385-5

Bay oil (34)

Current regulation: /

Clinical data:
/

Additional information:

ISO 4720:2009 nomenclature: *Pimenta racemosa* (Mill.) J.W. Moore. For Oil of bay [*Pimenta racemosa* (Mill.) J.W. Moore] an ISO standard exists: ISO 3045:2004 Pimenta Racemosa Leaf/Fruit Oil is an essential oil obtained from the fruits of the plant, *Pimenta racemosa*, Myrtaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41014>, last accessed 2010-02-10).

Steam distillation of the leaves of *Pimenta racemosa* (Mill.) J.W. Moore (Myrtaceae) yields bay oil, which consists of myrcene (20-30%), eugenol (42-56%) and chavicol (8-13%) (34).

Considering the content of well-known allergenic compounds, this essential oil is regarded as established contact allergen in humans.

Pinus mugo leaf and twig oil and extract

CAS 90082-72-7, 8000-26-8; EC 290-163-6

Dwarf pine needle oil
(German: Latschenkiefernöl)

Current regulation: Annex III, part 1, 109

Clinical data:

In the Frosch 2002 b study, 0.7% positive reactions to dwarf pine needle oil (2% pet.) were observed in 1606 consecutive (17). The Rudzki 1976 study found 4 positive reactions in 200 patients to "Pine needle" essential oil, 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=3 (3.5%) positive reactions to "pine needle" essential oil 2% pet. (27).

Additional information:

ISO 4720:2009 nomenclature: *Pinus mugo* Turra syn. *Pinus montana* Mill.) Pinus Mugo Twig Oil is an essential obtained from the twigs of the Pine, *Pinus mugo*, Pinaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41476&back=1>, last accessed 2010-03-09). Pinus Mugo Twig Leaf Extract is an extract obtained from the twigs leaves of the Pine, *Pinus mugo*, Pinaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41476&back=1>).

[h.details&id=41473&back=1](#), last accessed 2010-03-09).

Dwarf pine needle oil is obtained from *Pinus mugo* Turra subsp. *mugo* and subsp. *pumilio* (Haenke) Franco (34). For Oil of dwarf pine (*Pinus mugo* Turra) an ISO standard exists: ISO 21093:2003. American pine oils contain almost no 3-carene or camphene (34).

PINUS PUMILA TWIG LEAF EXTRACT / OIL

CAS 97676-05-6; EC 307-681-6
(*Pine, Pinus pumila, ext. = INCI*)

Dwarf pine needle oil

Current regulation: Annex III, part 1, 114

Clinical data: /

Additional information:

Pinus Pumila Twig Leaf Extract obtained from the twigs leaves of the Pine, *Pinus pumila*, Pinaceae

(<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41483&back=1>, last accessed 2009-11-12), *Pinus Pumila* Twig Leaf Oil is the essential oil obtained from the twigs leaves of the Pine, *Pinus pumila*, Pinaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41484&back=1>, last accessed 2009-11-12). Main constituents are alpha-pinene (60-70%) and beta-pinene (20-25%). (34) Occurrence from Siberia to Japan, classified as Endangered Species

Considering the content of well-known allergenic compounds, this essential oil is regarded as established contact allergen in humans.

POGOSTEMON CABLIN OIL

CAS 8014-09-3; EC / (*Oils, patchouli*) / 84238-39-1; EC 282-493-4 (*Patchouli, ext.*)

Patchouli oil

INCI: POGOSTEMON CABLIN / *Patchouli, ext.*

Current regulation: /

Clinical data:

In the Frosch 2002 b study, 0.8% positive reactions to patchouli oil (10% pet.) in 1606 consecutive were observed (17). Nakayama et al. found 1974 (after (29)) 3 "strong positive" and 8 "weak positive" reactions to "Patchouli oil" (unknown test concentration) in 183 patients. The IVDK 2010 c study identified 0.6% positive reactions in 2446 consecutively tested patients and 1.4% positive reactions in 828 patients tested in the context of a special series (30).

Additional information:

ISO 4720:2009 nomenclature: *Pogostemon cablin* (Blanco) Benth. syn. *Mentha cablin* Blanco. An ISO standard is available for Oil of patchouli (*Pogostemon cablin* (Blanco) Benth.): ISO 3757:2002. *Pogostemon Cablin* Leaf Oil is an essential oil obtained from the fermented leaves of the Patchouli, *Pogostemon cablin* (syn: *Pogostemon patchouli*),

Labiatae (Lamiaceae (34)). It contains patchouli alcohol, beta-patchoulene, azulene, eugenol, sesquiterpenes (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=40927>, last accessed 2009-11-12). Although the sesquiterpene alcohol (-)-patchoulol [5986-55-0] is the main component of patchouli oil (27-35%), the compound largely contributing to the characteristic odour is norpatchoulol [41429-52-1] (0.35-1%). Other constituents include (+)-alpha-bulnesene [6391-11-0] (13-21%), (-)-alpha-guajene [3691-12-1] (11-16%), (-)-β-patchoulene [514-51-2] (1.8-3.5%) and (-)-seychellene [20085-93-2] (1-3%) (34). According to (30) the maximum observed concentration in patchouli oil are (in %): (-)-patchoulol (35); (+)-alpha-lulnesene (21); (-)-alpha-guajene (16); β-pinene (6); (-)-β-patchoulene (3.5); (-)-seychellene (3); pogostol (2.5); α-pinene (2.5); norpatchoulol (1) (30).

It is a "top 100" substance (IFRA, pers. comm.2010).

| | |
|------------------------------------|--|
| ROSE FLOWER OIL (ROSA SPP.) | CAS 8007-01-0; EC / (Oils, rose) |
| ROSA ALBA FLOWER EXTRACT | CAS 93334-48-6; EC 297-122-1 (Rose, <i>Rosa alba</i> , ext. = INCI) |
| ROSA CANINA FLOWER OIL | CAS 84696-47-9; EC 283-652-0 (Rose, <i>Rosa canina</i> , ext.) INCI: ROSA CANINA |
| ROSA CENTIFOLIA FLOWER OIL | CAS 84604-12-6, EC 283-289-8 (Rose, <i>Rosa centifolia</i> , ext.) INCI: ROSA CENTIFOLIA / Rose, <i>Rosa centifolia</i> , ext. |
| ROSA DAMASCENA FLOWER OIL | CAS 90106-38-0; EC 290-260-3 (Rose, <i>Rosa Damascena</i> , ext. = INCI) |
| ROSA GALLICA FLOWER OIL | CAS 84604-13-7; EC 283-290-3 (Rose, <i>Rosa Gallica</i> , ext.) INCI: ROSA GALLICA |
| ROSA MOSCHATA OIL | -- |
| ROSA RUGOSA FLOWER OIL | CAS 92347-25-6; EC 296-213-3 (Rose, <i>Rosa rugosa</i> , ext.) |

Current regulation: /

Clinical data:

In the Sugiura 2000 study, 1483 patients with suspected cosmetic dermatitis were PTed with "rose oil Bulgaria" (2% pet.), yielding 0.4% positive reactions (14); Trattner/David found 2 / 641 consecutive patients positive to "Rose oil (Bulgarian)" (31). The Bulgarian rose oil usually corresponds to *Rosa Damascena* Flower Oil (http://en.wikipedia.org/wiki/Rose_oil, last accessed 2009-11-16). The Coimbra 2000 study found in 67 patients with positive reaction to the FM I who were tested with "rose Bulgarian oil" (2% pet.) 4.5% positive reactions (9). One case of contact allergy to "Bulgarian rose oil (2 % pet.)" – and geraniol – in a 48-year-old female with ACD after application of "Eau de Rochas" E.d.C. was diagnosed, among 326 patients with suspected contact allergy to fragrance ingredients had tested negative (275). However, other rose oils are also used (and capable of eliciting ACD) as illustrated by the case of a 27 year old woman who developed ACD after using "Rose Absolute Eau ® eau de

parfum", a "non-scented" body lotion and a number of other topicals. PTing revealed a number of (previously) relevant reaction, including "Rose centifolia" (5% alc.) and "Rose oil Bulgarian" (2% pet.) essential oil preparations (276). In the An 2005 study, 5 of 422 consecutive patients, i.e., 1.2%, had positive reactions to "Rose oil Bulgarian", tested at 2% concentration (13). Nakayama et al. found 1974 (after (29)) 4 "strong positive" reactions to "Rose oil Bulgarian" (unknown test concentration) in 183 patients. In a study from Alicante, Spain, 86 selected patients were tested with rose oil absolute, yielding 6 positive reactions (48).

Additional information:

ISO 4720:2009 nomenclature: *Rosa x damascena* Mill. and *Rosa sertata* X *Rosa rugosa*. Rose Flower Oil is the volatile oil obtained from the flowers of *Rosa* spp. , rosaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=59362>, last accessed 2009-11-16). "Rose oil, meaning either rose otto (attar of rose, attar of roses) or rose absolute, is the essential oil extracted from the petals of various types of rose. Rose ottos are extracted through steam distillation, while rose absolutes are obtained through solvent extraction or supercritical carbon dioxide extraction, with the absolute being used more commonly in perfumery" (http://en.wikipedia.org/wiki/Rose_oil, last accessed 2009-11-17) There are several more specifically named flower extracts used for masking or perfuming:

- Rosa Alba Flower Extract is an extract obtained from the flowers of the Rose, *Rosa alba* L., Rosaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=40969>, last accessed 2009-11-16).
- Rosa Canina Flower Oil is the volatile oil obtained from the flowers of the Hip Rose, *Rosa canina* L., Rosaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=59263>, last accessed 2009-11-16).
- Rosa Centifolia Flower Oil is the volatile oil obtained from the flowers of the Cabbage Rose, *Rosa centifolia* (L.), Rosaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=79757>, last accessed 2009-11-16).
- Rosa Damascena Flower Oil is the volatile oil obtained from the flowers of the Damask Rose, *Rosa damascena*, Rosaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=79760>, last accessed 2009-11-16).
- Rosa Gallica Flower Oil is the volatile oil obtained from the flowers of the French Rose, *Rosa gallica* L., Rosaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=59346>, last accessed 2009-11-16).
- Rosa Moschata Oil is the oil obtained from the Musk Rose, *Rosa moschata*, Rosaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=79761>, last accessed 2009-11-16).
- Rosa Rugosa Flower Oil is the volatile oil obtained from the flowers of the Rose, *Rosa rubiginosa* L., Rosaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=83588>, last accessed 2009-11-16).

Apparently, the *Rosa Damascena* and the *Rosa centifolia* are the species most commonly used for extraction of essential rose oils, the former mostly grown in Bulgaria, Turkey, Russia, India and China, the latter more commonly in Morocco, France and Egypt (276). Main constituents by GC are: citronellol (20-49%), geraniol (6-23%), nerol (3-12%) and phenylethyl alcohol (up to 3.5%) (34).

For Oil of rose (*Rosa x damascena* Miller) an ISO standard exists: ISO 9842:2003.

ROSMARINUS OFFICINALIS FLOWER OILCAS 84604-14-8; EC 283-291-9
(Rosemary, ext.)

"Rosemary Oil"

INCI: ROSMARINUM OFFICINALIS
/ Rosemary, ext.

Current regulation: /

Clinical data:

The Rudzki 1976 study found no positive reaction in 200 patients to "rosemary" essential oil, 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=3 (3.5%) positive reactions to "rosemary" essential oil 2% pet. (27).

Additional information:

ISO 4720:2009 nomenclature: *Rosmarinus officinalis* L. Rosmarinus Officinalis Flower Oil is an essential oil obtained from the leaves and fresh flowering tops of the Rosemary, *Rosmarinus officinalis* L., Lamiaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=40978>, last accessed 2010-01-29). Major constituents are: 1,8-cineole (17-55%), alpha-pinene (9-26%), camphor (5-22%) and verbenone [18309-32-5] as traces in North African oils, but between 0.7 and 2.5% in Spanish oils (34). For Oil of rosemary (*Rosmarinus officinalis* L.) an ISO standard exists: ISO 1342:2000.

Considering the content of well-known allergenic compounds, this essential oil is regarded as established contact allergen in humans.

SALVIA spp. HERB OIL

Sage oil

SALVIA OFFICINALIS LAVANDULIFOLIA HERB OILCAS 97952-71-1; EC 308-365-0
(Sage, *Salvia officinalis*
lavandulifolia, ext. = INCI)**SALVIA LAVANDULIFOLIA HERB OIL**CAS 90106-49-3; EC 290-272-9
(Sage, *Salvia lavandulifolia*, ext.
= INCI)**SALVIA SCLAREA FLOWER OIL**CAS 84775-83-7; EC 283-911-8
(Sage, *Salvia sclarea*, ext.) INCI:
SALVIA SCLAREA / Sage, *Salvia*
sclarea, ext.**SALVIA HISPANICA HERB OIL**CAS 93384-40-8; EC 297-250-8
(Sage, *Salvia hispanica*, ext. =
INCI)

Current regulation: /

Clinical data:

The Rudzki 1976 study found 1 positive reaction in 200 patients to "Clary sage", 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=4 (4.6%) positive reactions to "clary sage" essential oil 2% pet. (27).

Additional information:

ISO 4720:2009 nomenclature: *Salvia officinalis* L. *Salvia Officinalis Lavandulifolia* Herb Oil is an essential oil obtained from the herbs of the Sage, *Salvia officinalis* L. spp. *lavandulifolia*, *Lamiaceae*, Syn. Dalmatian sage (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41084>, last accessed 2010-01-29).

Salvia Lavandulifolia Herb Oil is an essential oil obtained from the herbs of the Sage, *Salvia lavandulifolia*, *Lamiaceae* (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=40987>, last accessed 2010-01-29).

Salvia Sclarea Flower Oil is an essential oil obtained from the flowers and foliage of the Clary Sage, *Salvia sclarea* L., *Lamiaceae* (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41086>, last accessed 2010-01-29).

Salvia Hispanica Herb Oil is an essential oil obtained from the herbs of the Spanish Sage, *Salvia hispanica* L., *Lamiaceae* (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=40985>, last accessed 2010-01-29).

Clary sage oil is obtained by steam distillation of flowering tops and foliage of cultivated *Salvia sclarea* L. (*Lamiaceae*). Main constituents are linalyl acetate (56-78%) and linalool (6.5-24%) (34). Dalmatian sage oil is steam distilled from partially dried leaves of *S. officinalis* L. (*Lamiaceae*). The content by GC is: alpha-thujone (18-43%), beta-thujone (3-8.5%), 1,8-cineole (5.5-13%), camphor (3-8.5%) as main constituents (34). Spanish sage oil does not contain thujone, but mainly camphor (15-36%) and 1,8-cineole (11-30%), and is used mainly in pharmaceutical preparations and technical perfumery (34). For Oil of sage, Spanish (*Salvia lavandulifolia* Vahl) an ISO standard exists: ISO 3526:2005, for Oil of Dalmatian sage (*Salvia officinalis* L.): ISO 9909:1997.

Considering the content of well-known allergenic compounds, this essential oil is regarded as established contact allergen in humans.

SANTALUM ALBUM WOOD OIL

CAS 84787-70-2; EC 284-111-1
(Sandalwood, ext.) INCI:
SANTALUM ALBUM / Sandalwood,
ext.

Sandalwood oil ([East] India)

SANTALUM ALBUM OIL

CAS 8006-87-9; EC / (Oils,
sandalwood)

Sandalwood oil ([East] India)

Current regulation: /

Clinical data:

In the Sugiura 2000 study, 1483 patients with suspected cosmetic dermatitis were PTed with "sandalwood oil" (2% pet.), yielding 0.8% positive reactions (14). In the Frosch 2002 b study, "sandalwood oil (East India)" is mentioned with a CAS # 8015-65-4, which, however, is attributed to AMYRIS BALSAMIFERA BARK OIL, see above. Assuming that this CAS # is erroneous, study results are considered to be valid for *S. album* wood oil, tested at 2% and 10% concentration, yielding 0.4% and 0.9% positive reactions,

respectively (17). Out of 6 of 15 patients with a positive reaction to the higher concentration no clinical relevance was found, compared to 2 of 7 patients positive to the lower concentration (17). The Coimbra 2000 study found in 67 patients with positive reaction to the FM I who were tested with "sandalwood oil" (2% pet.) 6.6% positive reactions (9). In the An 2005 study, 10 of 422 consecutive patients, i.e., 2.4%, had positive reactions to "Santalum album oil" 2% (13). The Goossens 1997 study found 4 of 111 patients positive to "sandalwood oil 10% pet." – all sensitised to other fragrance allergens (23). The Rudzki 1976 study found no positive reaction in 200 patients to "sandalwood", 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=2 (2.3%) positive reactions to "sandalwood" essential oil 2% pet. (27). In 63 patients positive to the FM I, 1 had a positive PT reaction to sandalwood oil, 2% pet., in the Santucci 1987 study (28). Nakayama et al. found 1974 (after (29)) 6 "strong positive" and 8 "weak positive" reactions to "Sandalwood oil" (unknown test concentration) in 183 patients. The IVDK 2010 c study identified 1.3% positive reactions in 3671 consecutively tested patients and 1.8% positive reactions in 1002 patients tested in the context of a special series (30). In a study from Alicante, Spain, 86 selected patients were tested with sandalwood oil, yielding 2 positive reactions (48).

Additional information:

ISO 4720:2009 nomenclature: *Santalum album* L. *Santalum Album* Oil is the volatile oil obtained from the heartwood of the Sandalwood, *Santalum album* L., Santalaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=80209>, last accessed 2009-11-26).

Santalum Album Wood Oil is an essential oil obtained from the wood of the Sandalwood, *Santalum album* L., Santalaceae. It contains 75% santalol isomers (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41092>, last accessed 2009-11-12), typically up to 55% .alpha.-santalol and up to 24% .beta.-santalol (30). East Indian sandalwood oil consists almost exclusively of closely related sesquiterpenoids; by far the main constituents are the alcohols alpha-santalol [115-71-9] (41-55%) and cis-beta-santalol [77-42-9] (16-24%), the latter being mainly responsible for the specific odour (34, 39).

An ISO standard regarding the composition of "*Santalum album* oil" is available: ISO 3518:2002. "Sandalwoods" are labelled as *Amyris balsamifera*, *Eremophila mitchelli*, *Fusanus acuminatus* (= *Santalum acuminatum*), *Santalum album*, *S. austrocaledonicum*, *S. latifolium*, *S. spicatum* and *S. yasi*. The majority of currently available trade oils, reportedly from *S. album*, contained approximately 50-70% santalols (Z-alpha and Z-beta), as analysed with gas chromatography-mass spectrometry (GC-MS) (277). A review on the toxicological properties of "*Santalum album* oil" is available (278).

AMYRIS BALSAMIFERA BARK OIL (*Sandalwood oil (Caribbean)*), CAS 8015-65-4; EC / (Oils, amyris) / 90320-49-3; EC 90320-49-3 (*Amyris balsamifera*, ext. = INCI name) is used as a cheap substitute for East Indian Sandalwood in perfumes and cosmetics. Originally cultivated primarily in Haiti where it was known as 'candle wood' and used as a torch by locals due to the tree's high oil content (<http://www.amphora-retail.com/sandalwood-amyris-essential-10ml-p-107.html>, last accessed 2009-11-12). The major components are sesquiterpenoids such as valerianol, elemol, β -eudesmol and epi-gamma-eudesmol (39). For Oil of amyris (*Amyris balsamifera* L.) an ISO standard exists: ISO 3525:2008. *Amyris Balsamifera* Bark Oil is the volatile oil distilled from the bark of the tree, *Amyris balsamifera*, Rutaceae (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=74455>, last accessed 2009-11-12).

SANTALUM SPICATA WOOD OILCAS 8024-35-9; EC 296-618-5
(Sandalwood oil, Western
Australia)*Sandalwood oil (Australia)*

Current regulation: /

Clinical data:

In clinical studies, mostly *S. album* wood oil had been used (see above); in a number of studies this is not clear.

Additional information:

ISO 4720:2009 nomenclature: *Santalum spicatum* (R.Br.) A. DC, syn. *Eucarya spicata* (R.Br.) Sprag & Summ. For Oil of Australian sandalwood (*Santalum spicatum* (R.Br.) A.DC.) an ISO standard exists: ISO 22769:2009. Santalum Spicata Wood Oil is an essential oil obtained from the wood of the Australian Sandalwood, *Santalum spicata*, Santalaceae. It contains 75% santalols and 10% farnesol (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41093>, last accessed 2009-11-12). This oil also contains santalols as main constituents but differs somewhat in the remaining composition. Today, it makes up a considerable part of the sandalwood oil market (34).

Considering the content of well-known allergenic compounds (santalols), this essential oil is regarded as established contact allergen in humans.

TAGETES PATULA FLOWER OILCAS 91722-29-1; EC 294-431-3
(*Tagetes patula*, ext. = INCI)*"Marigold Oil; Tagetes Oil"*

Current regulation: /

Clinical data:

In an aromatherapist, an essential oil solvent-extracted from *Tagetes patula*, patch tested at 1.5% in grapeseed oil (vehicle negative, 7 controls negative to essential oils) resulted in a +++ reaction, in accordance with a work-related bilateral hand dermatitis (217).

Additional information:

Tagetes Patula Flower Oil is an essential oil obtained by hydrodistillation of the flowers of the *Tagetes*, *Tagetes patula* L., *Compositae*. It contains mainly D-limonene, ocimene, 2,6-dimethyloct-7-en-4-one (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41506>, last accessed 2010-01-28). According to Surburg/Panten, tagetes oil is steam distilled from the flowering plants of *Tagetes minuta* L. (*T. glandulifera* Schrank., *Asteraceae*). Main components comprise cis-ocimene, dihydrotagetone, tagetone, and cis- and trans-ocimenone (34, 39).

THYMUS spp. HERB OIL

THYMUS VULGARIS HERB OIL

CAS 84929-51-1, 8007-46-3; EC 284-535-7 (Thyme, *Thymus vulgaris*, ext.)

"Thyme oil"

INCI: THYMUS VULGARIS / Thyme, *Thymus vulgaris*, ext.

Current regulation: /

Clinical data:

The Rudzki 1976 study found no positive reaction in 200 patients to "thyme" essential oil, 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=4 (4.6%) positive reactions to "thyme" essential oil 2% pet. (27). In 63 patients positive to the FM I, none had a positive PT reaction to thymol, 1% pet., in the Santucci 1987 study (28).

Additional information:

ISO 4720:2009 nomenclature: *Thymus vulgaris* L. *Thymus vulgaris* Herb Oil is an essential oil obtained from the herbs of the Thyme, *Thymus vulgaris* L., Lamiaceae. It contains 20-40% thymol and carvacrol, cymene, pinene, linalool, bornyl acetate (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41133>, last accessed 2010-01-29).

Other species are used for extraction, e.g., *Thymus Mastichina* (CAS 84837-14-9), *Thymus Serpillum* (CAS 84776-98-7), *Thymus Zygis* (CAS 85085-75-2), according to CosIng. The main constituent is thymol (37-56%) (34). For Oil of thyme containing thymol, Spanish type [*Thymus zygis* (Loefl.) L.] an ISO standard exists: ISO 14715:2010, for Oil of Spanish wild marjoram (*Thymus mastichina* L.): ISO 4728:2003.

TURPENTINE (oil)

CAS 8006-64-2 / 9005-90-7 / 8052-14-0; EC 232-350-7 / 232-688-5 / -

Current regulation: III/124 ; III/125 ; III/126

Clinical data:

Oil of turpentine has been patch tested in a number of baseline series, i.e., in consecutive patients, although not included in the European Baseline series.

In a series of 24 patients with occupational contact dermatitis from the pottery industry, Lear at al. found 14 to be sensitised to "Indonesian oil of turpentine" and 8 to alpha-pinene (190)

Table 3.2.2 – 2: Overview of results with **Oil of turpentine** in patients patch tested for suspected allergic contact dermatitis. If not given in the publication, the confidence interval (CI) was calculated from the absolute numbers by the SCCS.

| Country | Population | Years | No. tested | Crude % positive (95% CI) [§] |
|---------|------------|-------|------------|--|
|---------|------------|-------|------------|--|

Opinion on fragrance allergens in cosmetic products

| | | | | |
|---|-------------------------------------|-------------------------|-------|---|
| Lisbon, Portugal (189); virtually no .delta.-3-carene | Consecutive patients | 1979-1983 | 4316 | 2.3 % (1.9 – 2.8) § |
| Birmingham, UK (190) | Potters with occup. dermatitis hand | 6 months; prior to 1996 | 24 | 14 / 24 pos. to "Indonesian turpentine" |
| Austria/Germany (IVDK) (279) | Consecutive patients | 1992-1995 | 27658 | 0.47 % (0.39 – 0.55) § |
| Austria/Germany (IVDK) (280) | Consecutive patients | 1996-2002 | 59478 | Annual prevalences 1.6 to 4.4 % |
| Augsburg/Germany (281) | Population sample | 1998 | 1141 | 1.2% (on population level!) |
| Europe (ESSCA) (273) | Consecutive patients | 2002/03 | 3767 | 1.6 % |
| Austria/Germany/Switzerland (IVDK) (7) | Consecutive patients | 2005-2008 | 37163 | 1.8 % |

Additional information:

ISO 4720:2009 nomenclature: *Pinus pinaster* Aiton and *Pinus massoniana* Lamb.
Turpentine, oil: Any of the volatile predominately terpenic fractions or distillates resulting from the solvent extraction of, gum collection from, or pulping of softwoods. Turpentine is a mixture of terpene hydrocarbons obtained from various species of *Pinus* http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.de tails_v2&id=41521

The composition of oil of turpentine varies with its origin, in particular, the content of .delta.-3-carene, one of its main allergenic compounds (189, 279). Similarly, the peroxide degree may vary. The main constituents are .alpha.-pinene (50-72%), .beta.-pinene (6-15%), carenes (< 0.1-17%), camphene (up to 1%), dipentene (0.5-5%), along with a number of other substances (279).

It is a "top 200" substance and classified as R43 (IFRA, pers. comm.2010)

Verbena absolute (*Lippia citriodora* Kunth.) CAS 8024-12-2, 84961-67-1; EC /)

Current regulation: Annex III, part 1, n° 206

Clinical data: /

Additional information:

ISO 4720:2009 nomenclature: *Aloysia citriodora* Palau syn. *Lippia citriodora* Kunth syn. *Aloysia triphylla* (L' Hér.) Kuntze. An older RIFM review is available citing several positive human maximisation studies both with "Verbena absolute" and "Verbena oil" (128).

VETIVERIA ZIZANOIDES ROOT OIL

CAS 8016-96-4; EC / (Oils, vetiver) / 84238-29-9; EC 282-490-8 (Vetiveria zizanioides, ext. = INCI)

"Vetiver oil; khas khas oil"

Current regulation: ...

Clinical data:

The Rudzki 1976 study found 1 positive reaction in 200 patients to "vetiver" essential oil, 2% pet. (26). The later Rudzki 1986 study in 86 FM I positive patients found n=9 (10.4%) positive reactions to "vetiver" essential oil 2% pet. (27).

Additional information:

ISO 4720:2009 nomenclature: *Vetiveria zizanioides* (L.) Nash. *Vetiveria Zizanioides* Root Oil is an essential oil distilled from the dried roots of the grass *Vetiveria zizanioides* (L.) Nash *Poaceae* (<http://ec.europa.eu/consumers/cosmetics/cosing/index.cfm?fuseaction=search.details&id=41293>, last accessed 2010-01-29). Vetiver oil has a high sesquiterpene content. The ketones alpha-vetivone [15764-04-2] (6-12%) and beta-vetivone [18444-79-6] (4-10%), which usually form more than 10% of the oil, as well as khusimol [16223-63-5] (24-36%) and isovelencenol [22387-74-2] (12-24%) are the main constituents (in Bourbon oil, i.e., from Réunion) (34). For Oil of vetiver (*Vetiveria zizanioides* (L.) Nash) an ISO standard exists: ISO 4716:2002.

Acknowledgement: We thank Erich Schmidt for critically reviewing the nomenclature of natural extracts and for providing ISO terminology.

References

1. Larsen W, Nakayama H, Fischer T, Elsner P, Frosch P, Burrows D, Jordan W, Shaw S, Wilkinson J, Marks J, Sugawara M, Nethercott M, Nethercott J. Fragrance contact dermatitis - a worldwide multicenter investigation (Part III). *Contact Dermatitis* 2002; 46: 141-144.
2. Hendriks S A, van Ginkel C J. Evaluation of the fragrance mix in the European standard series. *Contact Dermatitis* 1999; 41: 161-162.
3. Katsarma G, Gawkrödger D J. Suspected fragrance allergy requires extended patch testing to individual fragrance allergens. *Contact Dermatitis* 1999; 41: 193-197.
4. Schnuch A, Uter W, Geier J, Lessmann H, Frosch P J. Sensitization to 26 fragrances to be labelled according to current European regulation. Results of the IVDK and review of the literature. *Contact Dermatitis* 2007; 57: 1-10.
5. Temesvari E, Nemeth I, Balo-Banga M J, Husz S, Kohanka V, Somos Z, Judak R, Remenyik E V, Szegedi A, Nebenfuhrer L, Meszaros C, Horvath A. Multicentre study of fragrance allergy in Hungary. Immediate and late type reactions. *Contact Dermatitis* 2002; 46: 325-330.
6. van Oosten E J, Schuttelaar M L, Coenraads P J. Clinical relevance of positive patch test reactions to the 26 EU-labelled fragrances. *Contact Dermatitis* 2009; 61: 217-223.
7. Uter W, Geier J, Frosch P J, Schnuch A. Contact allergy to fragrances: current patch test results (2005 to 2008) from the IVDK network. *Contact Dermatitis* 2010; 63: 254-261.
8. Schnuch A, Geier J, Uter W, Frosch P J. Another look on allergies to fragrances: frequencies of sensitisation to the fragrance mix and its constituents. Results from the IVDK. *Exog Dermatol* 2002; 1: 231-237.
9. Brites M M, Goncalo M, Figueiredo A. Contact allergy to fragrance mix--a 10-year study. *Contact Dermatitis* 2000; 43: 181-182.
10. Frosch P J, Rastogi S C, Pirker C, Brinkmeier T, Andersen K E, Bruze M, Svedman C, Goossens A, White I R, Uter W, Arnau E G, Lepoittevin J P, Johansen J D, Menne T. Patch testing with a new fragrance mix - reactivity to the individual constituents and chemical detection in relevant cosmetic products. *Contact Dermatitis* 2005; 52: 216-225.
11. Krautheim A, Uter W, Frosch P, Schnuch A, Geier J. Patch testing with fragrance mix II: results of the IVDK 2005-2008. *Contact Dermatitis* 2010; 63: 262-269.
12. deGroot A C, Coenraads P J, Bruynzeel D P, Jagtman B A, van_Ginkel C J W, Noz K, van_der_Valk P G M, Pavel S, Vink J, Weyland J W. Routine patch testing with fragrance chemicals in The Netherlands. *Contact Dermatitis* 2000; 42: 184-185.
13. An S, Lee A Y, Lee C H, Kim D W, Hahm J H, Kim K J, Moon K C, Won Y H, Ro Y S, Eun H C. Fragrance contact dermatitis in Korea: a joint study. *Contact Dermatitis* 2005; 53: 320-323.
14. Sugiura M, Hayakawa R, Kato Y, Sugiura K, Hashimoto R. Results of patch testing with lavender oil in Japan. *Contact Dermatitis* 2000; 43: 157-160.
15. Frosch P J, Pilz B, Andersen K E, Burrows D, Camarasa J G, et al. Patch testing with fragrances: results of a multicenter study of the European Environmental and Contact Dermatitis Research Group with 48 frequently used constituents of perfumes. *Contact Dermatitis* 1995; 33: 333-342.

16. Frosch P J, Johansen J D, Menne T, Pirker C, Rastogi S C, Andersen K E, Bruze M, Goossens A, Lepoittevin J P, White I R. Further important sensitizers in patients sensitive to fragrances. I. Reactivity to 14 frequently used chemicals. *Contact Dermatitis* 2002; 47: 78-85.
17. Frosch P J, Johansen J D, Menne T, Pirker C, Rastogi S C, Andersen K E, Bruze M, Goossens A, Lepoittevin J P, White I R. Further important sensitizers in patients sensitive to fragrances. II. Reactivity to essential oils. *Contact Dermatitis* 2002; 47: 279-287.
18. Larsen W G. Perfume Dermatitis. A Study of 20 Patients. *Arch Dermatol* 1977; 113: 623-626.
19. Larsen W, Nakayama H, Fischer T, Elsner P, Frosch P, Burrows D, Jordan W, Shaw S, Wilkinson J, Marks J, Jr., Sugawara M, Nethercott M, Nethercott J. Fragrance contact dermatitis: a worldwide multicenter investigation (Part II). *Contact Dermatitis* 2001; 44: 344-346.
20. Belsito D V, Fowler J F, Jr., Sasseville D, Marks J G, Jr., De Leo V A, Storrs F J. Delayed-type hypersensitivity to fragrance materials in a select North American population. *Dermatitis* 2006; 17: 23-28.
21. Zug K A, Warshaw E M, Fowler J F, Jr., Maibach H I, Belsito D L, Pratt M D, Sasseville D, Storrs F J, Taylor J S, Mathias C G, Deleo V A, Rietschel R L, Marks J. Patch-test results of the North American Contact Dermatitis Group 2005-2006. *Dermatitis* 2009; 20: 149-160.
22. Wöhrl S, Hemmer W, Focke M, Götz M, Jarisch R. The significance of fragrance mix, balsam of Peru, colophony and propolis as screening tools in the detection of fragrance allergy. *Br J Dermatol* 2001; 145: 268-273.
23. Goossens A, Merckx L. Allergic Contact Dermatitis from farnesol in a deodorant. *Contact Dermatitis* 1997; 37: 179-180.
24. Malten K E, van Ketel W G, Nater J P, Liem D H. Reactions in selected patients to 22 fragrance materials. *Contact Dermatitis* 1984; 11: 1-10.
25. de Groot A C, Liem D H, Nater J P, van Ketel W G. Patch tests with fragrance materials and preservatives. *Contact Dermatitis* 1985; 12: 87-92.
26. Rudzki E, Grzywa Z, Bruo W S. Sensitivity to 35 essential oils. *Contact Dermatitis* 1976; 2: 196-200.
27. Rudzki E, Grzywa Z. Allergy to perfume mixture. *Contact Dermatitis* 1986; 15: 115-116.
28. Santucci B, Cristaudo A, Cannistraci C, Picardo M. Contact dermatitis to fragrances. *Contact Dermatitis* 1987; 16: 93-95.
29. Mitchell J C. Contact hypersensitivity to some perfume materials. *Contact Dermatitis* 1975; 1: 196-199.
30. Uter W, Schmidt E, Geier J, Lessmann H, Schnuch A, Frosch P J. Contact allergy to essential oils: current patch test results (2000-2008) from the IVDK network. *Contact Dermatitis* 2010; 63: 277-283.
31. Trattner A, David M. Patch testing with fine fragrances: comparison with fragrance mix, balsam of Peru and a fragrance series. *Contact Dermatitis* 2003; 49: 287-289.
32. Handley J, Burrows D. Allergic contact dermatitis from the synthetic fragrances Lylal and acetyl cedrene in separate underarm deodorant preparations. *Contact Dermatitis* 1994; 31: 288-290.

33. SCCNFP. The Scientific Committee on Cosmetic Products and Non-Food Products intended for Consumers: Opinion concerning Fragrance Allergy in Consumers. A Review of the Problem. Analysis of the Need for appropriate Consumer Information and Identification of Consumer Allergens, adopted 8 December 1999. *SCCNFP/0017/98 Final* 1999:
34. Surburg H, Panten J. *Common fragrance and flavor materials: preparation, properties and uses*. Weinheim: Wiley-VCH, 2006.
35. Bhatia S P, Wellington G A, Cocchiara J, Lalko J, Letizia C S, Api A M. Fragrance material review on alpha-amylcinnamyl alcohol. *Food Chem Toxicol* 2007; 45 Suppl 1: S32-39.
36. Lapczynski A, McGinty D, Jones L, Bhatia S P, Letizia C S, Api A M. Fragrance material review on pentyl salicylate. *Food Chem Toxicol* 2007; 45 Suppl 1: S460-466.
37. Franks A. Contact allergy to anethole in toothpaste associated with loss of taste. *Contact Dermatitis* 1998; 38: 354-355.
38. Garcia-Bravo B, Perez Bernal A, Garcia-Hernandez M J, Camacho F. Occupational contact dermatitis from anethole in food handlers. *Contact Dermatitis* 1997; 37: 38.
39. Fahlbusch K-G, Hammerschmidt F-J, Panten J, Pickenhagen W, Schatkowski D, Bauer K, Garbe D, Surburg H. Flavors and Fragrances. In: Wiley-VCH, eds. *Ullmann's Encyclopedia of Industrial Chemistry*. Weinheim: Wiley-VCH, 2002:
40. Hostynek J J, Maibach H I. Is there evidence that anisyl alcohol causes allergic contact dermatitis? *Exog Dermatol* 2003; 2: 230-233.
- 40a. Bruze M, Svedman C, Andersen KE, Bruynzeel D, Goossens A, Johansen JD, Matura M, Orton D, Vigan M; ESCD. Patch test concentrations (doses in mg/cm²) for the 12 non-mix fragrance substances regulated by European legislation. *Contact Dermatitis* 2012; 66: 131-136
41. Andersen A. Final report on the safety assessment of benzaldehyde. *Int J Toxicol* 2006; 25 Suppl 1: 11-27.
42. Seite-Bellezza D, el Sayed F, Bazex J. Contact urticaria from cinnamic aldehyde and benzaldehyde in a confectioner. *Contact Dermatitis* 1994; 31: 272-273.
43. Opdyke D L, Letizia C. Monographs on fragrance raw materials. *Food Chem Toxicol* 1983; 21: 645-667.
44. Nair B. Final report on the safety assessment of Benzyl Alcohol, Benzoic Acid, and Sodium Benzoate. *Int J Toxicol* 2001; 20 Suppl 3: 23-50.
45. Sestini S, Mori M, Francalanci S. Allergic contact dermatitis from benzyl alcohol in multiple medicaments. *Contact Dermatitis* 2004; 50: 316-317.
46. Podda M, Zollner T, Grundmann-Kollmann M, Kaufmann R, Boehncke W H. Allergic contact dermatitis from benzyl alcohol during topical antimycotic treatment. *Contact Dermatitis* 1999; 41: 302-303.
47. Shoji A. Allergic reaction to benzyl alcohol in an antimycotic preparation. *Contact Dermatitis* 1983; 9: 510.
48. Cuesta L, Silvestre J F, Toledo F, Lucas A, Perez-Crespo M, Ballester I. Fragrance contact allergy: a 4-year retrospective study. *Contact Dermatitis* 2010; 63: 77-84.
49. Guin J D, Goodman J. Contact urticaria from benzyl alcohol presenting as intolerance to saline soaks. *Contact Dermatitis* 2001; 45: 182-183.

50. Fisher A A. Allergic paraben and benzyl alcohol hypersensitivity relationship of the "delayed" and "immediate" varieties. *Contact Dermatitis* 1975: 1: 281-284.
51. Shaw D W. Allergic contact dermatitis to benzyl alcohol in a hearing aid impression material. *Am J Contact Dermat* 1999: 10: 228-232.
52. Jacob S E, Barron G S. Benzyl alcohol: a covert fragrance. *Dermatitis* 2007: 18: 232-233.
53. Hausen B M, Brinkmann J, Dohn W. *Lexikon der Kontaktallergene (6. Erg.-Lieferung)*. Landsberg am Lech: Ecomed, 1998.
54. Corazza M, Manovani L, Maranini C, Virgili A. Allergic Contact Dermatitis from benzyl alcohol. *Contact Dermatitis* 1996: 34: 74.
55. Buffet M, Dupin N. Current treatments for scabies. *Fundam Clin Pharmacol* 2003: 17: 217-225.
56. Bhatia S P, Wellington G A, Cocchiara J, Lalko J, Letizia C S, Api A M. Fragrance material review on benzyl cinnamate. *Food Chem Toxicol* 2007: 45 Suppl 1: S40-48.
57. Lapczynski A, McGinty D, Jones L, Bhatia S, Letizia C S, Api A M. Fragrance material review on benzyl salicylate. *Food Chem Toxicol* 2007: 45 Suppl 1: S362-380.
58. Bhatia S P, Jones L, Letizia C S, Api A M. Fragrance material review on 2-tert-butylcyclohexyl acetate. *Food Chem Toxicol* 2008: 46 Suppl 12: S44-47.
59. Bhatia S P, Jones L, Letizia C S, Api A M. Fragrance material review on 4-tert-butylcyclohexyl acetate. *Food Chem Toxicol* 2008: 46 Suppl 12: S36-41.
60. Arnau E G, Andersen K E, Bruze M, Frosch P J, Johansen J D, Menne T, Rastogi S C, White I R, Lepoittevin J P. Identification of Lilial as a fragrance sensitizer in a perfume by bioassay-guided chemical fractionation and structure-activity relationships. *Contact Dermatitis* 2000: 43: 351-358.
61. Stevenson O E, Finch T M. Allergic contact dermatitis from rectified camphor oil in Earex ear drops. *Contact Dermatitis* 2003: 49: 51.
62. Vilaplana J, Romaguera C, Campderros L. [Contact dermatitis by camphor present in a flushing solution]. *Actas Dermosifiliogr* 2007: 98: 345-346.
63. Noiles K., M. P. Contact dermatitis to Vicks VapoRub. *Dermatitis* 2010: 21: 167-169.
64. Sköld M, Karlberg A T, Matura M, Börje A. The fragrance chemical beta-caryophyllene-air oxidation and skin sensitization. *Food Chem Toxicol* 2006: 44: 538-545.
65. Matura M, Skold M, Borje A, Andersen K E, Bruze M, Frosch P, Goossens A, Johansen J D, Svedman C, White I R, Karlberg A T. Selected oxidized fragrance terpenes are common contact allergens. *Contact Dermatitis* 2005: 52: 320-328.
66. Andersen A. Final report on the safety assessment of sodium p-chloro-m-cresol, p-chloro-m-cresol, chlorothymol, mixed cresols, m-cresol, o-cresol, p-cresol, isopropyl cresols, thymol, o-cymen-5-ol, and carvacrol. *Int J Toxicol* 2006: 25 Suppl 1: 29-127.
67. Corazza M, Levratti A, Virgili A. Allergic contact cheilitis due to carvone in toothpastes. *Contact Dermatitis* 2002: 46: 366-367.
68. Worm M, Jeep S, Sterry W, Zuberbier T. Perioral contact dermatitis caused by L-carvone in toothpaste. *Contact Dermatitis* 1998: 38: 338.
69. Hausen B M. Zahnpasta-Allergie. *Dtsch Med Wochenschr* 1984: 109: 300-302.

70. Paulsen E, Andersen K E, Carlsen L, et al. Carvone: an overlooked contact allergen cross-reacting with sesquiterpene lactones? *Contact Dermatitis* 1993; 29: 138-143.
71. Karlberg A T, Magnusson K, Nilsson U. Air oxidation of d-limonene (the citrus solvent) creates potent allergens. *Contact Dermatitis* 1992; 26: 332-340.
72. Matura M, Goossens A, Bordalo O, Garcia-Bravo B, Magnusson K, Wrangsjö K, Karlberg A T. Patch testing with oxidized R-(+)-limonene and its hydroperoxide fraction. *Contact Dermatitis* 2003; 49: 15-21.
73. Nilsson A M, Gafvert E, Salvador L, Luthman K, Bruze M, Gruvberger B, Nilsson J L, Karlberg A T. Mechanism of the antigen formation of carvone and related alpha, beta-unsaturated ketones. *Contact Dermatitis* 2001; 44: 347-356.
74. Nguyen S H, Dang T P, MacPherson C, Maibach H, Maibach H I. Prevalence of patch test results from 1970 to 2002 in a multi-centre population in North America (NACDG). *Contact Dermatitis* 2008; 58: 101-106.
75. Diba V C, Statham B N. Contact urticaria from cinnamal leading to anaphylaxis. *Contact Dermatitis* 2003; 48: 119.
76. Decapite T J, Anderson B E. Allergic contact dermatitis from cinnamic aldehyde found in an industrial odour-masking agent. *Contact Dermatitis* 2004; 51: 312-313.
77. Cocchiara J, Letizia C S, Lalko J, Lapczynski A, Api A M. Fragrance material review on cinnamaldehyde. *Food Chem Toxicol* 2005; 43: 867-923.
78. Bickers D, Calow P, Greim H, Hanifin J M, Rogers A E, Saurat J H, Sipes I G, Smith R L, Tagami H. A toxicologic and dermatologic assessment of cinnamyl alcohol, cinnamaldehyde and cinnamic acid when used as fragrance ingredients. *Food Chem Toxicol* 2005; 43: 799-836.
79. Buckley D A, Basketter D A, Smith Pease C K, Rycroft R J, White I R, McFadden J P. Simultaneous sensitivity to fragrances. *Br J Dermatol* 2006; 154: 885-888.
80. Elahi E N, Wright Z, Hinselwood D, Hotchkiss S A, Basketter D A, Pease C K. Protein binding and metabolism influence the relative skin sensitization potential of cinnamic compounds. *Chem Res Toxicol* 2004; 17: 301-310.
81. Letizia C S, Cocchiara J, Lalko J, Lapczynski A, Api A M. Fragrance material review on cinnamyl alcohol. *Food Chem Toxicol* 2005; 43: 837-866.
82. Heydorn S, Menne T, Andersen K E, Bruze M, Svedman C, White I R, Basketter D A. Citral a fragrance allergen and irritant. *Contact Dermatitis* 2003; 49: 32-36.
83. Hindle E, Ashworth J, Beck M H. Chelitis from contact allergy to citral in lip salve. *Contact Dermatitis* 2007; 57: 125-126.
84. Hagvall L, Backtorp C, Svensson S, Nyman G, Borje A, Karlberg A T. Fragrance compound geraniol forms contact allergens on air exposure. Identification and quantification of oxidation products and effect on skin sensitization. *Chem Res Toxicol* 2007; 20: 807-814.
85. Hagvall L, Baron J M, Borje A, Weidolf L, Merk H, Karlberg A T. Cytochrome P450-mediated activation of the fragrance compound geraniol forms potent contact allergens. *Toxicol Appl Pharmacol* 2008; 233: 308-313.
86. Lapczynski A, Bhatia S P, Letizia C S, Api A M. Fragrance material review on l-citronellol. *Food Chem Toxicol* 2008; 46 Suppl 11: S110-113.
87. Lapczynski A, Letizia C S, Api A M. Fragrance material review on (+)-(R)-citronellol. *Food Chem Toxicol* 2008; 46 Suppl 11: S114-116.

88. Lapczynski A, Bhatia S P, Letizia C S, Api A M. Fragrance material review on dl-citronellol. *Food Chem Toxicol* 2008: 46 Suppl 11: S103-109.
89. Hostynek J J, Maibach H I. Sensitization Potential of Citronellol. *Exog Dermatol* 2004: 3: 307-312.
90. Mutterer V, Gimenez Arnau E, Lepoittevin J P, Johansen J D, Frosch P J, Menne T, Andersen K E, Bruze M, Rastogi S C, White I R. Identification of coumarin as the sensitizer in a patient sensitive to her own perfume but negative to the fragrance mix. *Contact Dermatitis* 1999: 40: 196-199.
91. Vocanson M, Goujon C, Chabeau G, Castelain M, Valeyrie M, Floc'h F, Maliverney C, Gard A, Nicolas J F. The skin allergenic properties of chemicals may depend on contaminants--evidence from studies on coumarin. *Int Arch Allergy Immunol* 2006: 140: 231-238.
92. Bhatia S P, Letizia C S, Api A M. Fragrance material review on cyclohexyl acetate. *Food Chem Toxicol* 2008: 46 Suppl 12: S52-55.
93. Letizia C S, Cocchiara J, Wellington G A, Funk C, Api A M. Food and chemical toxicology. *Food Chem Toxicol* 2000: 38 Suppl 3: S1-236.
94. Lapczynski A, Lalko J, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on damascenone. *Food Chem Toxicol* 2007: 45 Suppl 1: S172-178.
95. Takanami I, Nakayama H. TMCHB: a possible alternative to DNCB in skin testing for immune competence. *Contact Dermatitis* 1988: 19: 81-83.
96. Lapczynski A, Lalko J, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on alpha-damascone. *Food Chem Toxicol* 2007: 45 Suppl 1: S179-187.
97. Lapczynski A, Lalko J, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on cis-alpha-damascone. *Food Chem Toxicol* 2007: 45 Suppl 1: S188-191.
98. Lalko J, Lapczynski A, Letizia C S, Api A M. Fragrance material review on cis-beta-damascone. *Food Chem Toxicol* 2007: 45 Suppl 1: S192-198.
99. Lapczynski A, Lalko J, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on trans-beta-damascone. *Food Chem Toxicol* 2007: 45 Suppl 1: S199-204.
100. Lalko J, Lapczynski A, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on delta-damascone. *Food Chem Toxicol* 2007: 45 Suppl 1: S205-210.
101. Lapczynski A, Lalko J, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on trans,trans-delta-damascone. *Food Chem Toxicol* 2007: 45 Suppl 1: S211-215.
102. Lalko J, Lapczynski A, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on gamma-damascone. *Food Chem Toxicol* 2007: 45 (Suppl. 1): S216-S220.
103. McGinty D, Letizia C S, Api A M. Fragrance material review on dihydromyrcenol. *Food Chem Toxicol* 2010: 48 Suppl 3: S70-75.
104. McGinty D, Letizia C S, Api A M. Fragrance material review on 3,7-dimethyl-1,6-nonadien-3-ol. *Food Chem Toxicol* 2010: 48 Suppl 3: S52-55.
105. Mitchell D M, Beck M H. Contact allergy to benzyl alcohol in a cutting oil reodorant. *Contact Dermatitis* 1988: 18: 301-302.

106. Giusti F, Porcaro V, Seidenari S. Evaluation of eugenol allergy in a patch-test population. *Contact Dermatitis* 2001; 44: 37-38.
107. Quirce S, Fernandez-Nieto M, del Pozo V, Sastre B, Sastre J. Occupational asthma and rhinitis caused by eugenol in a hairdresser. *Allergy* 2008; 63: 137-138.
108. Bhalla M, Thami G P. Acute urticaria due to dental eugenol. *Allergy* 2003; 58: 158.
109. Sarrami N, Pemberton M N, Thornhill M H, Theaker E D. Adverse reactions associated with the use of eugenol in dentistry. *Br Dent J* 2002; 193: 257-259.
110. Kanerva L, Estlander T, Jolanki R. Dental nurse's occupational allergic contact dermatitis from eugenol used as a restorative dental material with polymethylmethacrylate. *Contact Dermatitis* 1998; 38: 339-340.
111. Hemmer W, Focke M, Leitner B, Gotz M, Jarisch R. Axillary dermatitis from farnesol in a deodorant. *Contact Dermatitis* 2000; 42: 168-169.
112. Schnuch A, Uter W, Geier J, Lessmann H, Frosch P J. Contact allergy to farnesol in 2021 consecutively patch tested patients. Results of the IVDK. *Contact Dermatitis* 2004; 50: 117-121.
113. Lapczynski A, Bhatia S P, Letizia C S, Api A M. Fragrance material review on farnesol. *Food Chem Toxicol* 2008; 46 Suppl 11: S149-156.
114. Tamagawa-Mineoka R, Katoh N, Kishimoto S. Allergic contact cheilitis due to geraniol in food. *Contact Dermatitis* 2007; 56: 242-243.
115. Yamamoto A, Morita A, Tsuji T, Suzuki K, Matsunaga K. Contact urticaria from geraniol. *Contact Dermatitis* 2002; 46: 52.
116. Hostynek J J, Maibach H I. Is there evidence that geraniol causes allergic contact dermatitis? *Exog Dermatol* 2004; 3: 318-331.
117. Lapczynski A, Bhatia S P, Foxenberg R J, Letizia C S, Api A M. Fragrance material review on geraniol. *Food Chem Toxicol* 2008; 46 Suppl 11: S160-170.
118. Dearman R J, Wright Z M, Basketter D A, Ryan C A, Gerberick G F, Kimber I. The suitability of hexyl cinnamic aldehyde as a calibrant for the murine local lymph node assay. *Contact Dermatitis* 2001; 44: 357-361.
119. Lapczynski A, Jones L, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on hexyl salicylate. *Food Chem Toxicol* 2007; 45 Suppl 1: S410-417.
120. Karlberg A T. Contact allergy to colophony. Chemical identifications of allergens, sensitization experiments and clinical experiences. *Acta Dermatol Venerol (Stockh) Suppl* 1988; 139: 1-43.
121. Schnuch A, Uter W, Dickel H, Szliska C, Schliemann S, Eben R, Rueff F, Gimenez-Arnau A, Loffler H, Aberer W, Frambach Y, Worm M, Niebuhr M, Hillen U, Martin V, Jappe U, Frosch P J, Mahler V. Quantitative patch and repeated open application testing in hydroxyisohexyl 3-cyclohexene carboxaldehyde sensitive-patients. *Contact Dermatitis* 2009; 61: 152-162.
122. Hendriks S A, Bousema M T, van Ginkel C J. Allergic contact dermatitis from the fragrance ingredient Lyrall in underarm deodorant. *Contact Dermatitis* 1999; 41: 119.
123. Lapczynski A, Bhatia S P, Letizia C S, Api A M. Fragrance material review on hydroxycitronellol. *Food Chem Toxicol* 2008; 46 Suppl 11: S179-181.
124. Lalko J, Lapczynski A, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on ionone. *Food Chem Toxicol* 2007; 45 Suppl 1: S251-257.

125. Lalko J, Lapczynski A, Politano V T, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on alpha-ionone. *Food Chem Toxicol* 2007: 45 Suppl 1: S235-240.
126. Lalko J, Lapczynski A, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on beta-ionone. *Food Chem Toxicol* 2007: 45 Suppl 1: S241-247.
127. Lapczynski A, Jones L, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on isoamyl salicylate. *Food Chem Toxicol* 2007: 45 Suppl 1: S418-423.
128. Ford R A, Api A M, Letizia C S. Monographs on fragrance raw materials. *Food Chem Toxicol* 1992: 30 Suppl: 1S-138S.
129. White J M, White I R, Glendinning A, Fleming J, Jefferies D, Basketter D A, McFadden J P, Buckley D A. Frequency of allergic contact dermatitis to isoeugenol is increasing: a review of 3636 patients tested from 2001 to 2005. *Br J Dermatol* 2007: 157: 580-582.
130. Tanaka S, Royds C, Buckley D, Basketter D A, Goossens A, Bruze M, Svedman C, Menne T, Johansen J D, White I R, McFadden J P. Contact allergy to isoeugenol and its derivatives: problems with allergen substitution. *Contact Dermatitis* 2004: 51: 288-291.
131. White I R, Johansen J D, Arnau E G, Lepoittevin J P, Rastogi S, Bruze M, Andersen K E, Frosch P J, Goossens A, Menne T. Isoeugenol is an important contact allergen: can it be safely replaced with isoeugenyl acetate? *Contact Dermatitis* 1999: 41: 272-275.
132. Lapczynski A, Lalko J, Politano V T, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on alpha-iso-methylionone. *Food Chem Toxicol* 2007: 45 Suppl 1: S280-289.
133. Hostynek J J, Maibach H I. Is there evidence that alpha-isomethylionone causes allergic contact dermatitis? *Exog Dermatol* 2004: 3: 121-125.
134. Guarneri F, Barbuzza O, Vaccaro M, Galtieri G. Allergic contact dermatitis and asthma caused by limonene in a labourer handling citrus fruits. *Contact Dermatitis* 2008: 58: 315-316.
135. Foti C, Zambonin C G, Conserva A, Casulli C, D'Accolti L, Angelini G. Occupational contact dermatitis to a limonene-based solvent in a histopathology technician. *Contact Dermatitis* 2007: 56: 109-112.
136. Wakelin S H, McFadden J P, Leonard J N, Rycroft R J. Allergic contact dermatitis from d-limonene in a laboratory technician. *Contact Dermatitis* 1998: 38: 164-165.
137. Topham E J, Wakelin S H. D-Limonene contact dermatitis from hand cleansers. *Contact Dermatitis* 2003: 49: 108-109.
138. Martins C, Goncalo M, Goncalo S. Allergic contact dermatitis from dipentene in wax polish. *Contact Dermatitis* 1995: 33: 126-127.
139. Rycroft R J. Allergic contact dermatitis from dipentene in honing oil. *Contact Dermatitis* 1980: 6: 325-329.
140. Meding B, Barregard L, Marcus K. Hand eczema in car mechanics. *Contact Dermatitis* 1994: 30: 129-134.
141. Karlberg A T, Dooms-Gossens A. Contact allergy to oxidized d-limonene among dermatitis patients. *Contact Dermatitis* 1997: 36: 201-206.
142. Matura M, Skold M, Borje A, Andersen K E, Bruze M, Frosch P, Goossens A, Johansen J D, Svedman C, White I R, Karlberg A T. Not only oxidized R-(+)- but

- also S-(-)-limonene is a common cause of contact allergy in dermatitis patients in Europe. *Contact Dermatitis* 2006: 55: 274-279.
143. Matura M, Goossens A, Bordalo O, Garcia-Bravo B, Magnusson K, Wrangsjo K, Karlberg A T. Oxidized citrus oil (R-limonene): a frequent skin sensitizer in Europe. *J Am Acad Dermatol* 2002: 47: 709-714.
 144. Sköld M, Börje A, Harambasic E, Karlberg A T. Contact allergens formed on air exposure of linalool. Identification and quantification of primary and secondary oxidation products and the effect on skin sensitization. *Chem Res Toxicol* 2004: 17: 1697-1705.
 145. Christensson J B, Matura M, Gruvberger B, Bruze M, Karlberg A T. Linalool--a significant contact sensitizer after air exposure. *Contact Dermatitis* 2010: 62: 32-41.
 146. Letizia C S, Cocchiara J, Lalko J, Api A M. Fragrance material review on linalool. *Food Chem Toxicol* 2003: 41: 943-964.
 147. Bickers D, Calow P, Greim H, Hanifin J M, Rogers A E, Saurat J H, Sipes I G, Smith R L, Tagami H. A toxicologic and dermatologic assessment of linalool and related esters when used as fragrance ingredients. *Food Chem Toxicol* 2003: 41: 919-942.
 148. Hostynek J J, Maibach H I. Is there evidence that linalool causes allergic contact dermatitis? *Exog Dermatol* 2003: 2: 223-229.
 149. de Groot A C, Bruynzeel D P, Bos J D, der Meeren H L v, van Joost T, Jagtman B A, Weyland J W. The allergens in cosmetics. *Arch Dermatol* 1988: 124: 1525-1529.
 150. Sköld M, Hagvall L, Karlberg A T. Autoxidation of linalyl acetate, the main component of lavender oil, creates potent contact allergens. *Contact Dermatitis* 2008: 58: 9-14.
 151. Hagvall L, Skold M, Brared-Christensson J, Borje A, Karlberg A T. Lavender oil lacks natural protection against autoxidation, forming strong contact allergens on air exposure. *Contact Dermatitis* 2008: 59: 143-150.
 152. Letizia C S, Cocchiara J, Lalko J, Api A M. Fragrance material review on linalyl acetate. *Food Chem Toxicol* 2003: 41: 965-976.
 153. Ford R A, Letizia C S, Api A M. Longifolene. *Food Chem Tox* 1992: 30(Suppl.): 67S-68S.
 154. Morton C A, Garioch J, Todd P, et al. Contact sensitivity to menthol and peppermint in patients with intra-oral symptoms. *Contact Dermatitis* 1995: 32: 281-284.
 155. Foti C, Conserva A, Antelmi A, Lospalluti L, Angelini G. Contact dermatitis from peppermint and menthol in a local action transcutaneous patch. *Contact Dermatitis* 2003: 49: 312-313.
 156. Andersson M, Hindsen M. Rhinitis because of toothpaste and other menthol-containing products. *Allergy* 2007: 62: 336-337.
 157. dos Santos M A, Santos Galvao C E, Morato Castro F. Menthol-induced asthma: a case report. *J Investig Allergol Clin Immunol* 2001: 11: 56-58.
 158. Bhatia S P, McGinty D, Letizia C S, Api A M. Fragrance material review on menthol. *Food Chem Toxicol* 2008: 46 Suppl 11: S209-214.
 159. Bhatia S P, McGinty D, Letizia C S, Api A M. Fragrance material review on d-menthol. *Food Chem Toxicol* 2008: 46 Suppl 11: S215-217.

160. Bhatia S P, McGinty D, Letizia C S, Api A M. Fragrance material review on l-menthol. *Food Chem Toxicol* 2008: 46 Suppl 11: S218-223.
161. Bhatia S P, McGinty D, Letizia C S, Api A M. Fragrance material review on d,l-menthol. *Food Chem Toxicol* 2008: 46 Suppl 11: S224-227.
162. Bhatia S P, McGinty D, Letizia C S, Api A M. Fragrance material review on menthol racemic. *Food Chem Toxicol* 2008: 46 Suppl 11: S228-233.
163. Nair B. Final report on the safety assessment of Mentha Piperita (Peppermint) Oil, Mentha Piperita (Peppermint) Leaf Extract, Mentha Piperita (Peppermint) Leaf, and Mentha Piperita (Peppermint) Leaf Water. *Int J Toxicol* 2001: 20 Suppl 3: 61-73.
164. Trattner A, David M. Patch testing with fine fragrances: comparison with fragrance mix, balsam of Peru and a fragrance series. *Contact Dermatitis* 2003: 49: 287-289.
165. Mitchell J C, Calnan C D, Clendenning W E, Cronin E, Hjorth N, Magnusson B, Maibach H I, Meneghini C L, Wilkinson D S. Patch testing with some components of balsam of Peru. *Contact Dermatitis* 1976: 2: 57-58.
166. Bhatia S P, Wellington G A, Cocchiara J, Lalko J, Letizia C S, Api A M. Fragrance material review on methyl cinnamate. *Food Chem Toxicol* 2007: 45 Suppl 1: S113-119.
167. Kaidbey K H, Kligman A M. Photocontact allergy to 6-methylcoumarin. *Contact Dermatitis* 1978: 4: 277-282.
168. Cardoso J C, Canelas M M, Goncalo M, Figueiredo A. Photopatch testing with an extended series of photoallergens: a 5-year study. *Contact Dermatitis* 2009: 60: 325-329.
169. Victor F C, Cohen D E, Soter N A. A 20-year analysis of previous and emerging allergens that elicit photoallergic contact dermatitis. *J Am Acad Dermatol* 2010: 62: 605-610.
170. McGinty D, Letizia C S, Api A M. Fragrance material review on 4-methyl-3-decen-5-ol. *Food Chem Toxicol* 2010: 48 Suppl 3: S93-96.
171. Lalko J, Lapczynski A, McGinty D, Bhatia S, Letizia C S, Api A M. Fragrance material review on methyl ionone (mixture of isomers). *Food Chem Toxicol* 2007: 45 Suppl 1: S300-307.
172. English J S, Rycroft R J. Allergic contact dermatitis from methyl heptine and methyl octine carbonates. *Contact Dermatitis* 1988: 18: 174-175.
173. Hostynek J J, Maibach H I. Is there evidence that methyl heptine carbonate causes allergic contact dermatitis? *Cutan Ocul Toxicol* 2006: 25: 259-271.
174. Heisterberg M V, Vigan M, Johansen J D. Active sensitization and contact allergy to methyl 2-octynoate. *Contact Dermatitis* 2010: 62: 97-101.
175. Bernaola G, Escayol P, Fernandez E, de Corres L F. Contact dermatitis from methylionone fragrance. *Contact Dermatitis* 1989: 20: 71-72.
176. Lalko J, Lapczynski A, McGinty D, Bhatia S P, Letizia C S, Api A M. Fragrance material review on alpha-irone. *Food Chem Toxicol* 2007: 45 Suppl 1: S272-275.
177. Oiso N, Fukai K, Ishii M. Allergic contact dermatitis due to methyl salicylate in a compress. *Contact Dermatitis* 2004: 51: 34-35.
178. Hindson C. Contact eczema from methyl salicylate reproduced by oral aspirin (acetyl salicylic acid). *Contact Dermatitis* 1977: 3: 348-349.

179. Lapczynski A, Jones L, McGinty D, Bhatia S P, Letizia C S, Api A M. Fragrance material review on methyl salicylate. *Food Chem Toxicol* 2007: 45 Suppl 1: S428-452.
180. Sköld M. *Contact allergy to autoxidized fragrance terpenes*. Thesis University of Gothenburg 2005.
181. Bhatia S P, McGinty D, Letizia C S, Api A M. Fragrance material review on myrtenol. *Food Chem Toxicol* 2008: 46 Suppl 11: S237-240.
182. Lapczynski A, Foxenberg R J, Bhatia S P, Letizia C S, Api A M. Fragrance material review on nerol. *Food Chem Toxicol* 2008: 46 Suppl 11: S241-244.
183. Lapczynski A, Bhatia S P, Letizia C S, Api A M. Fragrance material review on nerolidol (isomer unspecified). *Food Chem Toxicol* 2008: 46 Suppl 11: S247-250.
184. Lapczynski A, Letizia C S, Api A M. Fragrance material review on cis-nerolidol. *Food Chem Toxicol* 2008: 46 Suppl 11: S245-246.
185. Christensen L P, Jakobsen H B, Paulsen E, Hodal L, Andersen K E. Airborne Compositae dermatitis: monoterpenes and no parthenolide are released from flowering *Tanacetum parthenium* (feverfew) plants. *Arch Dermatol Res* 1999: 291: 425-431.
186. Lapczynski A, McGinty D, Jones L, Bhatia S, Letizia C S, Api A M. Fragrance material review on phenethyl salicylate. *Food Chem Toxicol* 2007: 45 Suppl 1: S467-471.
187. Sanchez-Politta S, Campanelli A, Pashe-Koo F, Saurat J H, Piletta P. Allergic contact dermatitis to phenylacetaldehyde: a forgotten allergen? *Contact Dermatitis* 2007: 56: 171-172.
188. McGinty D, Letizia C S, Api A M. Fragrance material review on phytol. *Food Chem Toxicol* 2010: 48 Suppl 3: S59-63.
189. Cachao P, Menezes Brandao F, Carmo M, Frazao S, Silva M. Allergy to oil of turpentine in Portugal. *Contact Dermatitis* 1986: 14: 205-208.
190. Lear J T, Heagerty A H M, Tan B B, et al. Transient re-emergence of oil of turpentine allergy in the pottery industry. *Contact Dermatitis* 1996: 34: 169-172.
191. Zacher K D, Ippen H. Kontaktekzem durch Bergamottöl. *Derm Beruf Umwelt* 1984: 32: 95-97.
192. Lapczynski A, Bhatia S P, Letizia C S, Api A M. Fragrance material review on rhodinol. *Food Chem Toxicol* 2008: 46 Suppl 11: S259-262.
193. Lapczynski A, Lalko J, McGinty D, Bhatia S P, Letizia C S, Api A M. Fragrance material review on alpha-isodamascone. *Food Chem Toxicol* 2007: 45 Suppl 1: S267-271.
194. Bruze M, Zimerson E. Cross-reaction patterns in patients with contact allergy to simple methylol phenols. *Contact Dermatitis* 1997: 37: 82-86.
195. Barbaud A, Reichert-Penetrat S, Trechot P, Granel F, Schmutz J L. [Sensitization to resorcinol in a prescription verrucide preparation: unusual systemic clinical features and prevalence]. *Ann Dermatol Venereol* 2001: 128: 615-618.
196. Aalto-Korte K, Valimaa J, Henriks-Eckerman M L, Jolanki R. Allergic contact dermatitis from salicyl alcohol and salicylaldehyde in aspen bark (*Populus tremula*). *Contact Dermatitis* 2005: 52: 93-95.
197. Bhatia S P, McGinty D, Letizia C S, Api A M. Fragrance material review on alpha-santalol. *Food Chem Toxicol* 2008: 46 Suppl 11: S267-269.

198. Bhatia S P, McGinty D, Letizia C S, Api A M. Fragrance material review on santalol. *Food Chem Toxicol* 2008: 46 Suppl 11: S263-266.
199. Bhatia S P, McGinty D, Letizia C S, Api A M. Fragrance material review on sclareol. *Food Chem Toxicol* 2008: 46 Suppl 11: S270-274.
200. Bhatia S P, McGinty D, Foxenberg R J, Letizia C S, Api A M. Fragrance material review on terpineol. *Food Chem Toxicol* 2008: 46 Suppl 11: S275-279.
201. Bhatia S P, Letizia C S, Api A M. Fragrance material review on (-)-alpha-terpineol. *Food Chem Toxicol* 2008: 46 Suppl 11: S204-205.
202. Bhatia S P, Letizia C S, Api A M. Fragrance material review on alpha-terpineol. *Food Chem Toxicol* 2008: 46 Suppl 11: S280-285.
203. Castelain P Y, Camoin J P, Jouglard J. Contact dermatitis to terpene derivatives in a machine cleaner. *Contact Dermatitis* 1980: 6: 358-360.
204. Hausen B M, Reichling J, Harkenthal M. Degradation products of monoterpenes are the sensitizing agents in tea tree oil. *Am J Contact Dermat* 1999: 10: 68-77.
205. Lapczynski A, Foxenberg R J, Bhatia S P, Letizia C S, Api A M. Fragrance material review on tetrahydrolinalool. *Food Chem Toxicol* 2008: 46 Suppl 11: S286-288.
206. Schnuch A, Geier J, Uter W, Frosch P J. Majantol--a new important fragrance allergen. *Contact Dermatitis* 2007: 57: 48-50.
207. Heisterberg M V, Johansen J D. Contact allergy to trimethyl-benzenepropanol (Majantol). *Contact Dermatitis* 2009: 61: 360-361.
208. Hausen B M. Contact allergy to balsam of Peru. II. Patch test results in 102 patients with selected balsam of Peru constituents. *Am J Contact Dermat* 2001: 12: 93-102.
209. Rudzki E, Grzywa Z. Dermatitis from propolis. *Contact Dermatitis* 1983: 9: 40-45.
210. Ferguson J E, Beck M H. Contact sensitivity to vanilla in a lip salve. *Contact Dermatitis* 1995: 33: 352.
211. Bhatia S P, Letizia C S, Api A M. Fragrance material review on tricyclo[5.2.1.0^{2,6}]dec-4-en-8-yl acetate. *Food Chem Toxicol* 2008: 46 Suppl 12: S100-101.
212. Bhatia S P, Jones L, Letizia C S, Api A M. Fragrance material review on tricyclodecenyyl acetate. *Food Chem Toxicol* 2008: 46 Suppl 12: S93-96.
213. Anonymous. *ISO/DIS 9235 Aromatic raw materials - vocabulary*. Geneva, Switzerland: International Standardisation Organisation, 2010.
214. Schmidt E. Production of Essential Oils. In: Husnu Can Baser K, Buchbauer G, eds. *Handbook of Essential Oils - Science, Technology, and Applications*. Boca Raton: CRC Press, 2010: 88-95.
215. Trattner A, David M, Lazarov A. Occupational contact dermatitis due to essential oils. *Contact Dermatitis* 2008: 58: 282-284.
216. Jung P, Sesztak-Greinecker G, Wantke F, Gotz M, Jarisch R, Hemmer W. Mechanical irritation triggering allergic contact dermatitis from essential oils in a masseur. *Contact Dermatitis* 2006: 54: 297-299.
217. Bilslund D, Strong A. Allergic contact dermatitis from the essential oil of French marigold (*Tagetes patula*) in an aromatherapist. *Contact Dermatitis* 1990: 23: 55-56.
218. Cockayne S E, Gawkrödger D J. Occupational contact dermatitis in an aromatherapist. *Contact Dermatitis* 1997: 37: 306-307.

219. Boonchai W, Iamtharachai P, Sunthonpalin P. Occupational allergic contact dermatitis from essential oils in aromatherapists. *Contact Dermatitis* 2007: 56: 181-182.
220. Keane F M, Smith H R, White I R, Rycroft R J. Occupational allergic contact dermatitis in two aromatherapists. *Contact Dermatitis* 2000: 43: 49-51.
221. Selvaag E, Holm J O, Thune P. Allergic contact dermatitis in an aroma therapist with multiple sensitizations to essential oils. *Contact Dermatitis* 1995: 33: 354-355.
222. Romaguera C, Vilaplana J. Occupational contact dermatitis from ylang-ylang oil. *Contact Dermatitis* 2000: 43: 251.
223. Sugawara M, Nakayama H, Watanabe S. Contact hypersensitivity to ylang-ylang oil. *Contact Dermatitis* 1990: 23: 248-249.
224. Kanerva L, Estlander T, Jolanki R. Occupational allergic contact dermatitis caused by ylang-ylang oil. *Contact Dermatitis* 1995: 33: 198-199.
225. Sanchez-Perez J, Garcia-Diez A. Occupational allergic contact dermatitis from eugenol, oil of cinnamon and oil of cloves in a physiotherapist. *Contact Dermatitis* 1999: 41: 346-347.
226. Vilaplana J, Romaguera C. Contact dermatitis from the essential oil of tangerine in fragrance. *Contact Dermatitis* 2002: 46: 108.
227. Lalko J, Api A M. Investigation of the dermal sensitization potential of various essential oils in the local lymph node assay. *Food Chem Toxicol* 2006: 44: 739-746.
228. Vilaplana J, Romaguera C. Allergic contact dermatitis due to eucalyptol in an anti-inflammatory cream. *Contact Dermatitis* 2000: 43: 118.
229. Commission M. *List of MAK and BAT Values 2010 (Report No. 46)*. Weinheim: Wiley-VCH, 2011.
230. Nardelli A, Gimenez-Arnau E, Bernard G, Lepoittevin J P, Goossens A. Is a low content in atranol/chloroatranol safe in oak moss-sensitized individuals? *Contact Dermatitis* 2009: 60: 91-95.
231. Johansen J D, Heydorn S, Menne T. Oak moss extracts in the diagnosis of fragrance contact allergy. *Contact Dermatitis* 2002: 46: 157-161.
232. Kanerva L, Jolanki R, Estlander T. Hairdresser's dermatitis caused by oak moss in permanent waving solution. *Contact Dermatitis* 1999: 41: 55-56.
233. Owen C M, August P J, Beck M H. Contact allergy to oak moss resin in a soluble oil. *Contact Dermatitis* 2000: 43: 112.
234. Rudzki E, Grzywa Z. Sensitizing and irritating properties of star anise oil. *Contact Dermatitis* 1976: 2: 305-308.
235. Franz H, Frank R, Rytter M, Haustein U F. Allergic contact dermatitis due to cedarwood oil after dermatoscopy. *Contact Dermatitis* 1998: 38: 182-183.
236. Adisen E, Önder M. Allergic contact dermatitis from *Laurus nobilis* oil induced by massage. *Contact Dermatitis* 2007: 56: 360-361.
237. Athanasiadis G I, Pfab F, Klein A, Braun-Falco M, Ring J, Ollert M. Erythema multiforme due to contact with laurel oil. *Contact Dermatitis* 2007: 57: 116-118.
238. Özden M G, Öztas P, Öztas M O, Önder M. Allergic contact dermatitis from *Laurus nobilis* (laurel) oil. *Contact Dermatitis* 2001: 45: 178.

239. Goiriz R, Delgado-Jimenez Y, Sanchez-Perez J, Garcia-Diez A. Photoallergic contact dermatitis from lavender oil in topical ketoprofen. *Contact Dermatitis* 2007; 57: 381-382.
240. Rademaker M. Allergic contact dermatitis from lavender fragrance in Diffiam gel. *Contact Dermatitis* 1994: 31:
241. Varma S, Blackford S, Statham B N, Blackwell A. Combined contact allergy to tea tree oil and lavender oil complicating chronic vulvovaginitis. *Contact Dermatitis* 2000: 42: 309-310.
242. Coulson I H, Khan A S. Facial 'pillow' dermatitis due to lavender oil allergy. *Contact Dermatitis* 1999: 41: 111.
243. Vermaat H, van Meurs T, Rustemeyer T, Bruynzeel D P, Kirtschig G. Vulval allergic contact dermatitis due to peppermint oil in herbal tea. *Contact Dermatitis* 2008: 58: 364-365.
244. Kalavala M, Hughes T M, Goodwin R G, Anstey A V, Stone N M. Allergic contact dermatitis to peppermint foot spray. *Contact Dermatitis* 2007: 57: 57-58.
245. Wilkinson S M, Beck M H. Allergic contact dermatitis from menthol in peppermint. *Contact Dermatitis* 1994: 30: 42.
246. Andersen K E. Contact allergy to toothpaste flavors. *Contact Dermatitis* 1978: 4: 195-198.
247. Clayton R, Orton D. Contact allergy to spearmint oil in a patient with oral lichen planus. *Contact Dermatitis* 2004: 51: 314-315.
248. Skrebova N, Brocks K, Karlsmark T. Allergic contact cheilitis from spearmint oil. *Contact Dermatitis* 1998: 39: 35.
249. Hänsel R, Keller K, Rimpler H, Schneider G. *Hagers Handbuch der pharmazeutischen Praxis. Drogen E - O*. Berlin, 894-902: Springer, 1993.
250. Hausen B M, Wollenweber E. Propolis allergy. (III). Sensitization studies with minor constituents. *Contact Dermatitis* 1988: 19: 296-303.
251. Hausen B M, Evers P, Stüwe T H, et al. Propolis allergy (IV) Studies with further sensitizers from propolis and constituents common to propolis, poplar buds and balsam of Peru. *Contact Dermatitis* 1992: 26: 34-44.
252. Hausen B M, Simatupang T, Bruhn G, Evers P, König W A. Identification of new allergenic constituents and proof of evidence for coniferyl benzoate in Balsam of Peru. *Am J Contact Dermat* 1995: 6: 199-208.
253. Hjorth N. Eczematous allergy to balsams, allied perfumes and flavouring agents. *Acta Derm Venereol* 1961: 41 (Suppl. 46): 1-216.
254. Wurm G. *Hagers Handbuch der pharmazeutischen Praxis. Waren und Dienste*. Berlin, 644-689: Springer, 1990.
255. Api A M. Only Peru Balsam extracts or distillates are used in perfumery. *Contact Dermatitis* 2006: 54: 179.
256. Temesvari E, Podanyi B, Ponyai G, Nemeth I. Fragrance sensitization caused by temporary henna tattoo. *Contact Dermatitis* 2002: 47: 240.
257. Lammintausta K, Maibach H I, Wilson D. Mechanisms of subjective (sensory) irritation. Propensity to non- immunologic contact urticaria and objective irritation in stingers. *Derm Beruf Umwelt* 1988: 36: 45-49.
258. Forsbeck M, Skog E. Immediate reactions to patch tests with balsam of Peru. *Contact Dermatitis* 1977: 3: 201-205.

259. Katsarou A, Armenaka M, Ale I, Koufou V, Kalogeromitros D. Frequency of immediate reactions to the European standard series. *Contact Dermatitis* 1999; 41: 276-279.
260. Temesvari E, Soos G, Podanyi B, Kovacs I, Nemeth I. Contact urticaria provoked by balsam of Peru. *Contact Dermatitis* 1978; 4: 65-68.
261. Cancian M, Fortina A B, Peserico A. Contact urticaria syndrome from constituents of balsam of Peru and fragrance mix in a patient with chronic urticaria. *Contact Dermatitis* 1999; 41: 300.
262. Tanaka S, Matsumoto Y, Dlova N, Ostlere L S, Goldsmith P C, Rycroft R J, Basketter D A, White I R, Banerjee P, McFadden J P. Immediate contact reactions to fragrance mix constituents and Myroxylon pereirae resin. *Contact Dermatitis* 2004; 51: 20-21.
263. Uter W, Lessmann H. Kontaktallergene. In: Schulze-Werninghaus G, Fuchs T, Bachert C, Wahn U, eds. *Manuale allergologicum*. Deisenhofen: Dustri, 2008: 237-308.
264. Freireich-Astman M, David M, Trattner A. Standard patch test results in patients with contact dermatitis in Israel: age and sex differences. *Contact Dermatitis* 2007; 56: 103-107.
265. Lazarov A. European Standard Series patch test results from a contact dermatitis clinic in Israel during the 7-year period from 1998 to 2004. *Contact Dermatitis* 2006; 55: 73-76.
266. Gupta N, Sheno S D, Balachandran C. Fragrance sensitivity in allergic contact dermatitis. *Contact Dermatitis* 1999; 40: 53-54.
267. Kashani M N, Gorouhi F, Behnia F, Nazemi M J, Dowlati Y, Firooz A. Allergic contact dermatitis in Iran. *Contact Dermatitis* 2005; 52: 154-158.
268. Avalos-Peralta P, Garcia-Bravo B, Camacho F M. Sensitivity to Myroxylon pereirae resin (balsam of Peru). A study of 50 cases. *Contact Dermatitis* 2005; 52: 304-306.
269. Akyol A, Boyvat A, Peksari Y, Gurgey E. Contact sensitivity to standard series allergens in 1038 patients with contact dermatitis in Turkey. *Contact Dermatitis* 2005; 52: 333-337.
270. Machovcova A, Dastychova E, Kostalova D, Vojtechovska A, Reslova J, Smejkalova D, Vaneckova J, Vocilkova A. Common contact sensitizers in the Czech Republic. Patch test results in 12,058 patients with suspected contact dermatitis*. *Contact Dermatitis* 2005; 53: 162-166.
271. Thyssen J P, Carlsen B C, Menne T, Johansen J D. Trends of contact allergy to fragrance mix I and Myroxylon pereirae among Danish eczema patients tested between 1985 and 2007. *Contact Dermatitis* 2008; 59: 238-244.
272. Lindberg M, Edman B, Fischer T, Stenberg B. Time trends in Swedish patch test data from 1992 to 2000. A multi-centre study based on age- and sex-adjusted results of the Swedish standard series. *Contact Dermatitis* 2007; 56: 205-210.
273. Uter W, Hegewald J, Aberer W, Ayala F, Bircher A J, Brasch J, Coenraads P J, Schuttelaar M L, Elsner P, Fartasch M, Mahler V, Belloni Fortina A, Frosch P J, Fuchs T, Johansen J D, Menne T, Jolanki R, Krecisz B, Kiec-Swierczynska M, Larese F, Orton D, Peserico A, Rantanen T, Schnuch A. The European standard series in 9 European countries, 2002/2003 - First results of the European Surveillance System on Contact Allergies. *Contact Dermatitis* 2005; 53: 136-145.
274. Bruynzeel D P, Diepgen T L, Andersen K E, Brandao F M, Bruze M, Frosch P J, Goossens A, Lahti A, Mahler V, Maibach H I, Menne T, Wilkinson J D. Monitoring

- the European standard series in 10 centres 1996-2000. *Contact Dermatitis* 2005: 53: 146-149.
275. Vilaplana J, Romaguera C, Grimalt F. Contact dermatitis from geraniol in Bulgarian rose oil. *Contact Dermatitis* 1991: 24: 301.
276. Nardelli A, Thijs L, Janssen K, Goossens A. Rosa centifolia in a 'non-scented' moisturizing body lotion as a cause of allergic contact dermatitis. *Contact Dermatitis* 2009: 61: 306-309.
277. Howes M J, Simmonds M S, Kite G C. Evaluation of the quality of sandalwood essential oils by gas chromatography-mass spectrometry. *J Chromatogr A* 2004: 1028: 307-312.
278. Burdock G A, Carabin I G. Safety assessment of sandalwood oil (*Santalum album* L.). *Food Chem Toxicol* 2008: 46: 421-432.
279. Treudler R, Richter G, Geier J, Schnuch A, Orfanos C E, Tebbe B. Increase in sensitization to oil of turpentine: recent data from a multicenter study on 45,005 patients from the German-Austrian Information Network of Departments of Dermatology (IVDK). *Contact Dermatitis* 2000: 42: 68-73.
280. Schnuch A, Lessmann H, Geier J, Frosch P J, Uter W. Contact allergy to fragrances: frequencies of sensitization from 1996 to 2002. Results of the IVDK*. *Contact Dermatitis* 2004: 50: 65-76.
281. Schäfer T, Böhler E, Ruhdorfer S, Weigl L, Wessner D, Filipiak B, Wichmann H E, Ring J. Epidemiology of contact allergy in adults. *Allergy* 2001: 56: 1192-1196.

Opinion on fragrance allergens in cosmetic products

Annex II - Animal Data

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| Substance | | Vehicle (AOO=acetone:olive oil; DEP=diethyl phthalate; DMF=dimethyl formamide; DMSO=dimethyl sulphoxide; EtOH=ethanol; MEK=methyl ethyl ketone) | Conc. in vehicle (%, generally w/v) | No. animals per dose group | EC3 value * | | | | Comment (deviation from OECD 429 etc) | Reference |
|---------------------------------|------------|---|---|--|-------------|--------------------|------|---|--|-----------|
| INCI name (<i>other name</i>) | CAS no. | | | | % | µg/cm ² | M | lowest for the substance (%) | | |
| <i>Allyl phenoxyacetate</i> | 7493-74-5 | 1:3 EtOH:DEP | 0.5, 1.0, 2.5, 5.0, 10.0 | 4 | 3.1 | 775 | 0.16 | 3.1 | RIFM, 2007a | |
| Amyl cinnamal | 122-40-7 | 1:3 EtOH:DEP | 1.0, 2.5, 5.0, 10.0, 25.0 | 4 | 7.6 | 1900 | 0.38 | 7.6 | RIFM, 2006a | |
| Amyl cinnamal | 122-40-7 | 4:1 AOO | - | 4 | 10.6 | 2650 | 0.52 | Elahi gives ref to Basketter et al 1999, but no data on the substance is found. It is not known if Elahi, Aptula and Roberts quote the same experiment | Elahi et al., 2004 | |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | | | |
|----------------------|----------|--------------|----------------------------|---|------|--------|-------|------|---|------------------------|
| Amyl cinnamal | 122-40-7 | - | - | - | 11 | 2750 | 0.54 | | Aptula gives ref to Kimber et al 2003, but no LLNA data on the substance is found. It is not known if Elahi, Aptula and Roberts quote the same experiment; original reference is not given. | Aptula et al., 2007 |
| Amyl cinnamal | 122-40-7 | - | - | - | 11 | 2750 | 0.54 | | Original ref not given. | Roberts et al., 2007 |
| Amylcinnamyl alcohol | 101-85-9 | 1:3 EtOH:DEP | 1.0, 2.5, 5.0, 10.0, 25.0 | 4 | > 25 | >6250 | >1.22 | > 25 | Should have been tested at higher concentrations | RIFM, 2004a |
| Anise alcohol | 105-13-5 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | 5.9 | 1475 | 0.43 | 5.9 | | RIFM, 2005a |
| Benzaldehyde | 100-52-7 | - | - | - | - | - | - | | No data in the ref | Roberts et al., 2007 |
| Benzaldehyde | 100-52-7 | - | - | - | - | - | - | | No data in the ref (poster abstract) | Basketter et al., 2003 |
| Benzyl alcohol | 100-51-6 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | > 50 | >12500 | >4.62 | > 50 | Should have been tested at higher concentrations | RIFM, 2005b |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | | | |
|---|------------|--------------|----------------------------|---|------|--------|-------|------|--|-------------|
| Benzy l benzoate | 120-51-4 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | > 50 | >12500 | >2.36 | > 50 | Should have been tested at higher concentrations | RIFM, 2005c |
| Benzy l cinnamate | 103-41-3 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | 18.4 | 4600 | 0.77 | 18.4 | | RIFM, 2005d |
| Benzy l salicylate | 118-58-1 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | 2.9 | 725 | 0.13 | 2.9 | | RIFM, 2005e |
| <i>p-tert-Butyl-dihydrocinnamaldehyde</i> | 18127-01-0 | 1:3 EtOH:DEP | 1.0, 2.5, 5.0, 10.0, 25.0 | 4 | 4.3 | 1075 | 0.23 | 4.3 | | RIFM, 2007b |
| Butylphenyl methylpropional (BMHCA) | 80-54-6 | EtOH | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 2.9 | 725 | 0.14 | 2.9 | | RIFM, 2001a |
| Butylphenyl methylpropional (BMHCA) | 80-54-6 | DEP | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 4.1 | 1025 | 0.20 | | | RIFM, 2001b |
| Butylphenyl methylpropional (BMHCA) | 80-54-6 | 1:3 EtOH:DEP | 0.3, 1.0, 3.0, 10.0, 30.0 | 4 | 13.9 | 3475 | 0.68 | | | RIFM, 2001c |
| Butylphenyl methylpropional (BMHCA) | 80-54-6 | 1:3 DEP:EtOH | 0.3, 1.0, 3.0, 10.0, 30.0 | 4 | 8.8 | 2200 | 0.43 | | | RIFM, 2001d |
| Butylphenyl methylpropional (BMHCA) | 80-54-6 | 4:1 AOO | 1.0, 2.5, 5.0, 10.0, 25.0 | 4 | 16.8 | 4200 | 0.82 | | | RIFM, 2001e |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | | |
|---|------------|---|----------------------------|---|-------|-------|-------|-------|---|
| Butylphenyl methylpropional (BMHCA) | 80-54-6 | 4:1 AOO | 1, 2.5, 10, 25, 50 | 4 | 18.7 | 4675 | 0.92 | | Basketter et al., 2001 |
| Camellia sinensis leaf Tea Leaf Absolute | 84650-60-2 | DMF | 0.5, 1.0, 2.5, 5.0, 10.0 | 4 | > 5.0 | >1250 | N/a | > 5.0 | Should have been tested at higher concentrations RIFM, 2005m |
| Cananga odorata leaf / flower oil Ylang Ylang Extra | 8006-81-3 | 1:3 EtOH:DEP | 0.5, 1.0, 2.5, 5.0, 10.0 | 4 | 6.8 | 1700 | N/a | 6.8 | RIFM, 2007f |
| Carvone | 6485-40-1 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | 10.7 | 2675 | 0.71 | | RIFM, 2007c |
| Carvone | 6485-40-1 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | 5.7 | 1425 | 0.38 | 5.7 | RIFM, 2007d |
| Carvone | 6485-40-1 | 4:1 AOO | 6.0, 12, 20 | 4 | 13 | 3250 | 0.86 | | Nilsson et al., 2005 |
| Cinnamal | 104-55-2 | 3:1 EtOH:DEP | 0.1, 0.3, 1.0, 3.0, 10.0 | 4 | 0.2 | 50 | 0.015 | 0.2 | RIFM, 2003a |
| Cinnamal | 104-55-2 | 0.1% α -tocopherol in 3:1 EtOH:DEP | 0.1, 0.3, 1.0, 3.0, 10.0 | 4 | 0.2 | 50 | 0.015 | | RIFM, 2003b |
| Cinnamal | 104-55-2 | 2.0% α -tocopherol in 3:1 EtOH:DEP | 0.1, 0.3, 1.0, 3.0, 10.0 | 4 | 0.6 | 150 | 0.045 | | RIFM, 2003c |
| Cinnamal | 104-55-2 | 0.3% antioxidant mix (equal parts BHT, tocopherol and eugenol) in 3:1 | 0.1, 0.3, 1.0, 3.0, 10.0 | 4 | 0.7 | 175 | 0.053 | | RIFM, 2003d |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

EtOH:DEP

| | | | | | | | | | |
|----------|----------|--|---------------------------|---|-----|-----|-------|---|---------------------------|
| Cinnamal | 104-55-2 | 0.1% Trolox C in 3:1 EtOH:DEP | 0.1, 0.3, 1.0, 3.0, 10.0 | 4 | 0.7 | 175 | 0.053 | | RIFM, 2003e |
| Cinnamal | 104-55-2 | 2.0% α -tocopherol in 3:1 EtOH:DEP | 0.1, 0.3, 1.0, 3.0, 10.0 | 4 | 0.8 | 200 | 0.060 | | RIFM, 2003f |
| Cinnamal | 104-55-2 | 3:1 EtOH:DEP | 0.1, 0.3, 1.0, 3.0, 10.0 | 4 | 0.9 | 225 | 0.068 | | RIFM, 2003g |
| Cinnamal | 104-55-2 | 0.1% α -tocopherol in 3:1 EtOH:DEP | 0.1, 0.3, 1.0, 3.0, 10.0 | 4 | 1.1 | 275 | 0.083 | | RIFM, 2003h |
| Cinnamal | 104-55-2 | 0.3% antioxidant mix (equal parts BHT, tocopherol and eugenol) in 3:1 EtOH:DEP | 0.1, 0.3, 1.0, 3.0, 10.0 | 4 | 1.3 | 325 | 0.098 | | RIFM, 2003i |
| Cinnamal | 104-55-2 | 0.1% Trolox C in 3:1 EtOH:DEP | 0.1, 0.3, 1.0, 3.0, 10.0 | 4 | 1.4 | 350 | 0.11 | | RIFM, 2003j |
| Cinnamal | 104-55-2 | - | - | - | - | - | - | No data in the ref (poster abstract) | Basketter et al., 2002 |
| Cinnamal | 104-55-2 | 4:1 AOO | 0.5, 1, 2.5, 5, 10 | 4 | 3.1 | 775 | 0.23 | | Basketter et al., 2001 |
| Cinnamal | 104-55-2 | 4:1 AOO | - | 4 | 1.3 | 325 | 0.10 | | Elahi et al., 2004 |
| Cinnamal | 104-55-2 | 4:1 AOO | 1, 2.5 | - | 1.4 | 348 | 0.11 | Too few concentrations tested; few details given in ref | Smith and Hotchkiss, 2001 |
| Cinnamal | 104-55-2 | 4:1 AOO | 1.0, 2.5, 5.0, 10.0, 25.0 | 4 | 1.7 | 425 | 0.13 | | Wright et al., 1995 |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | | |
|-------------------------|-----------|---|---|---|------|-------|-------|------|---|
| Cinnamal | 104-55-2 | MEK | 1.0, 2.5, 5.0, 10.0, 25.0 | 4 | 1.1 | 275 | 0.083 | | Wright et al., 1996 |
| Cinnamal | 104-55-2 | DMF | 0.1, 0.25, 0.5, 1.0, 2.5, 5.0, 10.0, 25.0 | 4 | 0.5 | 125 | 0.038 | | Wright et al., 1997 |
| Cinnamal | 104-55-2 | propylene glycol | 1.0, 2.5, 5.0, 10.0, 25.0 | 4 | 1.4 | 350 | 0.11 | | Wright et al., 1998 |
| Cinnamal | 104-55-2 | DMSO | 0.1, 0.25, 0.5, 1.0, 2.5, 5.0, 10.0, 25.0 | 4 | 0.9 | 225 | 0.068 | | Wright et al., 1999 |
| Cinnamal | 104-55-2 | 90:10 EtOH:water | 1.0, 2.5, 5.0, 10.0, 25.0 | 4 | 1.6 | 400 | 0.12 | | Wright et al., 2000 |
| Cinnamal | 104-55-2 | 50:50 EtOH:water | 1.0, 2.5, 5.0, 10.0, 25.0 | 4 | 1.2 | 300 | 0.091 | | Wright et al., 2001 |
| Cinnamyl alcohol | 104-54-1 | - | - | - | - | - | - | - | No data in the ref (poster abstract) Basketter et al., 2002 |
| <i>Cinnamyl nitrile</i> | 1885-38-7 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | > 10 | >2500 | >0.77 | > 10 | Report: systemic toxicity at 25% and 50%. Should have been tested at higher concentrations RIFM, 2005f |
| Citral | 5392-40-5 | 1:3 EtOH:DEP | 0.4, 2.0, 4.0, 8.0, 20.0 | 4 | 1.2 | 300 | 0.079 | 1.2 | RIFM, 2004b |
| Citral | 5392-40-5 | 0.1% α -tocopherol in 3:1 EtOH:DEP | 0.3, 1.0, 3.0, 10.0, 30.0 | 4 | 1.5 | 375 | 0.099 | | RIFM, 2003k |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | | |
|-------------|-----------|--|----------------------------|---|------|-------|------|--------------------------------------|------------------------|
| Citral | 5392-40-5 | 0.3% antioxidant mix (equal parts BHT, tocopherol and eugenol) in 3:1 EtOH:DEP | 0.3, 1.0, 3.0, 10.0, 30.0 | 4 | 2.1 | 525 | 0.14 | | RIFM, 2003l |
| Citral | 5392-40-5 | 0.1% Trolox C in 3:1 EtOH:DEP | 0.3, 1.0, 3.0, 10.0, 30.0 | 4 | 3.7 | 925 | 0.24 | | RIFM, 2003m |
| Citral | 5392-40-5 | 3:1 EtOH:DEP | 0.3, 1.0, 3.0, 10.0, 30.0 | 4 | 4.6 | 1150 | 0.30 | | RIFM, 2003n |
| Citral | 5392-40-5 | 0.3% antioxidant mix (equal parts BHT, tocopherol and eugenol) in 3:1 EtOH:DEP | 0.3, 1.0, 3.0, 10.0, 30.0 | 4 | 4.6 | 1150 | 0.30 | | RIFM, 2003o |
| Citral | 5392-40-5 | 3:1 EtOH:DEP | 0.3, 1.0, 3.0, 10.0, 30.0 | 4 | 5.3 | 1325 | 0.35 | | RIFM, 2003p |
| Citral | 5392-40-5 | 0.1% Trolox C in 3:1 EtOH:DEP | 0.3, 1.0, 3.0, 10.0, 30.0 | 4 | 5.8 | 1400 | 0.38 | | RIFM, 2003q |
| Citral | 5392-40-5 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | 6.3 | 1575 | 0.41 | | RIFM, 2003r |
| Citral | 5392-40-5 | 0.1% α -tocopherol in 3:1 EtOH:DEP | 0.3, 1.0, 3.0, 10.0, 30.0 | 4 | 6.8 | 1700 | 0.44 | | RIFM, 2003s |
| Citral | 5392-40-5 | - | - | - | - | - | - | No data in the ref (poster abstract) | Basketter et al., 2002 |
| Citronellol | 106-22-9 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | 43.5 | 10875 | 2.78 | 43.5 | RIFM, 2004c |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | | | |
|--|------------|--------------|----------------------------|---|------|--------|-------|------|--|------------------------|
| Coumarin | 91-64-5 | DMF | 10, 25, 50 | 4 | >50 | >12500 | >3.42 | >50 | Should have been tested at higher concentrations | Vocanson et al., 2006 |
| <i>Dibenzyl ether</i> | 103-50-4 | 1:3 EtOH:DEP | 1.0, 2.5, 5.0, 10.0, 25.0 | 4 | 6.3 | 1575 | 0.32 | 6.3 | | RIFM, 2007e |
| Eugenol | 97-53-0 | 3:1 EtOH:DEP | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 5.3 | 1325 | 0.32 | 5.3 | | RIFM, 2001f |
| Eugenol | 97-53-0 | 1:3 EtOH:DEP | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 10.5 | 2625 | 0.64 | | | RIFM, 2001g |
| Eugenol | 97-53-0 | EtOH | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 10.7 | 2675 | 0.65 | | | RIFM, 2001h |
| Eugenol | 97-53-0 | DEP | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 15.1 | 3775 | 0.92 | | | RIFM, 2001i |
| Eugenol | 97-53-0 | 4:1 AOO | 2.5, 5.0, 10.0, 25.0, 50.0 | - | 11.9 | 2975 | 0.72 | | | Basketter et al., 1999 |
| Eugenol | 97-53-0 | - | - | - | - | - | - | | No data in the ref (poster abstract) | Basketter et al., 2003 |
| Evernia furfuracea extract <i>Treemoss absolute</i> | 90028-67-4 | 1:3 EtOH:DEP | 5.0, 10.0, 20 | 4 | > 20 | >5000 | N/a | > 20 | Should have been tested at higher concentrations | RIFM, 2004k |
| Evernia furfuracea extract <i>Treemoss absolute</i> | 90028-67-4 | 1:3 EtOH:DEP | 10.0, 25.0 | 4 | > 25 | >6250 | N/a | | Too few concentrations tested | RIFM, 2004d |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | | | |
|---|------------|--------------|-------------------------------|---|------|------|------|------|--|----------------------|
| Evernia prunastri extract <i>Oakmoss</i> | 90028-68-5 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | 3.88 | 970 | N/a | 3.88 | | RIFM, 2004j |
| | | | | | | | | | | |
| Farnesol | 4602-84-0 | 4:1 AOO | 5.0, 10.0, 25.0 | 4 | 5.5 | 1375 | 0.25 | | Should also have been tested at lower concentrations | RIFM, 2004d |
| Farnesol | 4602-84-0 | 4:1 AOO | 5.0, 10.0, 25.0 | 4 | 4.1 | 1025 | 0.18 | 4.1 | Should also have been tested at lower concentrations | RIFM, 2004d |
| | | | | | | | | | | |
| Geraniol | 106-24-1 | EtOH | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 5.6 | 1400 | 0.36 | 5.6 | | RIFM, 2001j |
| Geraniol | 106-24-1 | 3:1 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | 11.4 | 2850 | 0.74 | | | RIFM, 2003t |
| Geraniol | 106-24-1 | DEP | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 11.8 | 2950 | 0.76 | | | RIFM, 2001k |
| Geraniol | 106-24-1 | 1:3 EtOH:DEP | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 20.4 | 5100 | 1.32 | | | RIFM, 2001l |
| Geraniol | 106-24-1 | 3:1 EtOH:DEP | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 25.8 | 6450 | 1.67 | | | RIFM, 2001m |
| Geraniol | 106-24-1 | - | - | - | 26 | 6500 | 1.69 | | | Roberts et al., 2007 |
| | | | | | | | | | | |
| <i>trans-2-Hexenal</i> | 6728-26-3 | 1:3 EtOH:DEP | 0.5, 1.0, 2.5, 5, 10 | 4 | 2.6 | 650 | 0.26 | 2.6 | | RIFM, 2005g |
| <i>trans-2-Hexenal</i> | 6728-26-3 | - | - | - | 5.5 | 1375 | 0.56 | | | Roberts et al., 2007 |
| | | | | | | | | | | |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | | |
|------------------------------------|------------|-------------------|----------------------------|-------|----------|-----------|-----------|--------------------------------------|---------------------------------------|
| Hexyl cinnamal | 101-86-0 | generally 4:1 AOO | | -0162 | 5.3-14.7 | 1325-3675 | 0.25-0.68 | 5.3 | "numerous accounts in the literature" |
| <i>2-Hexylidene cyclopentanone</i> | 17373-89-6 | 1:3 EtOH:DEP | 0.1, 0.5, 1.0, 2.5, 5.0 | 5 | 2.4 | 600 | 0.14 | 2.4 | RIFM, 2008a |
| Hexyl salicylate | 6259-76-3 | 1:3 EtOH:DEP | 0.05, 0.25, 0.5, 1.0, 2.5 | 4 | 0.18 | 45 | 0.008 | 0.18 | RIFM, 2006b |
| Hydroxycitronellal | 107-75-5 | 1:3 EtOH:DEP | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 19.3 | 4825 | 1.12 | 19.3 | RIFM, 2001n |
| Hydroxycitronellal | 107-75-5 | DEP | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 19.7 | 4925 | 1.14 | | RIFM, 2001o |
| Hydroxycitronellal | 107-75-5 | 3:1 EtOH:DEP | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 22.2 | 5550 | 1.29 | | RIFM, 2001p |
| Hydroxycitronellal | 107-75-5 | EtOH | 1.0, 3.0, 10.0, 30.0, 50.0 | 4 | 26.4 | 6600 | 1.53 | | RIFM, 2001q |
| Hydroxycitronellal | 107-75-5 | AOO | 25, 50, 100 | - | - | - | - | EC3 value not given | Ashby et al., 1995 |
| Hydroxycitronellal | 107-75-5 | 4:1 AOO | 2.5, 5, 10, 25, 50 | 4 | 33.0 | 8250 | 1.92 | | Basketter et al., 2001 |
| Hydroxycitronellal | 107-75-5 | - | - | - | - | - | - | No data in the ref (poster abstract) | Basketter et al., 2002 |
| Hydroxycitronellal | 107-75-5 | - | - | - | 25.25 | 6313 | 1.47 | | Estrada et al., 2003 |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | | | |
|--|------------|--------------|--------------------------|---|------|-------|-------|------|---|---------------------------|
| Hydroxycitronellal | 107-75-5 | 4:1 AOO | 10, 25 | - | 23 | 5750 | 1.34 | | Too few concentrations tested; few details given in ref | Smith and Hotchkiss, 2001 |
| Hydroxyisohexyl 3-cyclohexene carboxaldehyde | 31906-04-4 | 4:1 AOO | 1.0, 2.5, 5, 10, 25, 50 | 4 | 17.1 | 4275 | 0.81 | 17.1 | | RIFM, 2001r |
| <i>p</i> -Isobutyl- α -methyl hydrocinnamaldehyde | 6658-48-6 | 70% EtOH | 10.0, 25.0, 50.0, 100.0 | 4 | 9.5 | 2375 | 0.46 | 9.5 | Should also have been tested at lower concentrations | RIFM, 2001w |
| <i>Isocyclocitral</i> | 1335-66-6 | 1:3 EtOH:DEP | 0.5, 1.0, 2.5, 5.0, 10.0 | 4 | 7.3 | 1825 | 0.48 | 7.3 | | RIFM, 2006c |
| <i>Isocyclogeraniol</i> | 68527-77-5 | 1:3 EtOH:DEP | 5.0, 10.0, 25.0, 50.0 | 4 | > 25 | >6250 | >1.62 | > 25 | Should have been tested at higher concentrations | RIFM, 2005h |
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.5, 5.0 | 6 | 0.54 | 145 | 0.033 | 0.54 | Too few concentrations tested | RIFM, 2001s |
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.5, 1.0, 5.0 | 5 | 0.6 | 150 | 0.037 | | | RIFM, 2002a |
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.5, 1.0, 5.0 | 5 | 0.76 | 191 | 0.046 | | | RIFM, 2002b |
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.5, 1.0, 5.0 | 5 | 0.79 | 199 | 0.048 | | | RIFM, 2002c |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | |
|------------|---------|---------|--------------------------|--------|------|-----|-------|--|
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.5, 1.0, 5.0 | 5 | 1.19 | 296 | 0.072 | RIFM, 2001t |
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.5, 1.0, 5.0 | 5 | 1.28 | 320 | 0.078 | RIFM, 2004e |
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.25, 0.5, 1.0, 2.5, 5.0 | 6 | 1.54 | 385 | 0.094 | RIFM, 2001u |
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.5, 1.0, 5.0 | 5 | 1.95 | 488 | 0.119 | RIFM, 2001v |
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.25, 0.5, 1.0, 2.5, 5.0 | | 3.3 | 825 | 0.20 | Basketter et al., 1999 |
| Isoeugenol | 97-54-1 | - | - | - | - | - | - | No data in the ref (poster abstract) Basketter et al., 2002 |
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.25, 0.5, 1.0, 2.5, 5.0 | 4 or 5 | 1.3 | 325 | 0.079 | Loveless et al., 1996 |
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.25, 0.5, 1.0, 2.5, 5.0 | 4 or 5 | 3.3 | 825 | 0.20 | Loveless et al., 1996 |
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.25, 0.5, 1.0, 2.5, 5.0 | 4 or 5 | 1.8 | 450 | 0.11 | Loveless et al., 1996 |
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.25, 0.5, 1.0, 2.5, 5.0 | 4 or 5 | 3.1 | 775 | 0.19 | Loveless et al., 1996 |
| Isoeugenol | 97-54-1 | 4:1 AOO | 0.25, 0.5, 1.0, 2.5, 5.0 | 4 or 5 | 1.6 | 400 | 0.097 | Loveless et al., 1996 |
| Isoeugenol | 97-54-1 | AOO | 0.5, 1.0, 2.5, 5.0, 10.0 | 4 | 1.0 | 250 | 0.061 | Wright et al., 2001 |
| Isoeugenol | 97-54-1 | MEK | 0.5, 1.0, 2.5, 5.0, 10.0 | 4 | 1.0 | 250 | 0.061 | Wright et al., 2001 |
| Isoeugenol | 97-54-1 | DMF | 0.5, 1.0, 2.5, 5.0, 10.0 | 4 | 1.4 | 350 | 0.085 | Wright et al., 2001 |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | | | |
|---|------------|------------------|----------------------------------|---|------|-------|-------|------|--|-----------------------|
| Isoeugenol | 97-54-1 | propylene glycol | 0.5, 1.0, 2.5, 5.0, 10.0 | 4 | 2.5 | 625 | 0.15 | | | Wright et al., 2001 |
| Isoeugenol | 97-54-1 | DMSO | 0.5, 1.0, 2.5, 5.0, 10.0 | 4 | 0.9 | 225 | 0.055 | | | Wright et al., 2001 |
| Isoeugenol | 97-54-1 | 90:10 EtOH:water | 0.5, 1.0, 2.5, 5.0, 10.0 | 4 | 1.8 | 450 | 0.11 | | | Wright et al., 2001 |
| Isoeugenol | 97-54-1 | 50:50 EtOH:water | 0.5, 1.0, 2.5, 5.0, 10.0 | 4 | 4.9 | 1225 | 0.30 | | | Wright et al., 2001 |
| | | | | | | | | | | |
| <i>Jasmine absolute (Grandiflorum)</i> | 8022-96-6 | 1:3 EtOH:DEP | 1.0, 2.5, 5.0, 10.0, 25.0 | 4 | 5.9 | 1475 | N/a | 5.9 | | RIFM, 2006d |
| | | | | | | | | | | |
| Jasminum Sambac Flower CERA / Extract / Water | 91770-14-8 | 1:3 EtOH:DEP | 10.0, 25.0, 50.0, 75.0, 100.0 | 4 | 35.4 | 9100 | N/a | 35.4 | | RIFM, 2006e |
| | | | | | | | | | | |
| <i>d-Limonene**</i> | 5989-27-5 | EtOH | 10.0, 20.0, 50.0, 75.0, 100.0 | 4 | < 10 | < 250 | <0.73 | < 10 | Should also have been tested at lower concentrations | RIFM, 2004l |
| <i>d-Limonene**</i> | 5989-27-5 | 3:1 EtOH:DEP | 10.0, 20.0, 50.0, 75.0, 100.0 | 4 | 22.0 | 5500 | 1.61 | | | RIFM, 2004m |
| <i>d-Limonene**</i> | 5989-27-5 | 1:3 EtOH:DEP | 10.0, 20.0, 50.0, 75.0, 100.0 | 4 | 38.0 | 9500 | 2.79 | | | RIFM, 2004n |
| <i>d-Limonene**</i> | 5989-27-5 | DEP | 10.0, 20.0, 50.0, 75.0, 100.0 | 4 | 63.0 | 15.75 | 4.62 | | | RIFM, 2004o |
| <i>d-Limonene**</i> | 5989-27-5 | 4:1 AOO | 25, 50, 100 | 4 | 68.5 | 17125 | 5.03 | | | Warbrick et al., 2001 |
| | | | | | | | | | | |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | | | |
|---|------------|--------------|-------------------------------|---|------|--------|-------|-------|--|---------------------------|
| Linalool** | 78-70-6 | - | - | - | - | - | - | - | No data in the ref (poster abstract) | Basketter et al., 2002 |
| <i>Menthadiene-7-methyl formate</i> | 68683-20-5 | 1:3 EtOH:DEP | 0.5, 1.0, 2.5, 5.0, 10.0 | 5 | > 10 | > 2500 | >0.51 | > 10 | Should have been tested at higher concentrations | RIFM, 2008c |
| <i>4-Methoxy-α-methyl benzenopropanal</i> | 5462-06-6 | 1:3 EtOH:DEP | 0.5, 1.0, 2.5, 5.0, 10.0 | 5 | 23.6 | 5900 | 1.32 | 23.63 | | RIFM, 2004f |
| <i>α-Methyl cinnamic aldehyde</i> | 101-39-3 | - | - | - | 4.5 | 1125 | 0.31 | 4.5 | | Roberts et al., 2007 |
| Methylenedioxyphenyl methylpropanal | 1205-17-0 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | 16.4 | 4100 | 0.85 | 16.4 | | RIFM, 2005i |
| <i>6-Methyl-3,5-heptadien-2-one</i> | 1604-28-0 | 1:3 EtOH:DEP | 0.5, 1.0, 2.5, 5.0, 10.0 | 5 | > 5 | > 1250 | >0.40 | > 5 | Should have been tested at higher concentrations | RIFM, 2008d |
| <i>α-iso-Methylionone</i> | 127-51-5 | 1:3 EtOH:DEP | 10.0, 25.0, 50.0, 75.0, 100.0 | 4 | 21.8 | 5450 | 1.06 | 21.8 | | RIFM, 2005j |
| <i>Methyl octine carbonate</i> | 111-80-8 | - | - | - | 2.5 | 635 | 0.15 | 2.5 | | Roberts et al., 2007 |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | | | |
|--|-----------|--------------|----------------------------|---|-------|--------|--------|-------|--|------------------------|
| Methyl 2-octynoate | 111-12-6 | 1:3 EtOH:DEP | 0.5, 1.0, 2.0, 5.0, 10.0 | 4 | < 0.5 | < 125 | <0.032 | < 0.5 | Should also have been tested at lower concentrations | RIFM, 2005k |
| 2-Methoxy-4-methylphenol | 93-51-6 | - | - | - | 5.8 | 1450 | 0.42 | 5.8 | | Roberts et al., 2007 |
| 1-Octen-3-yl acetate | 2442-10-6 | 1:3 EtOH:DEP | 7.5, 15.0, 30.0 | 5 | > 30 | > 7500 | >1.76 | > 30 | Should have been tested at higher concentrations | RIFM, 2004g |
| Perillaldehyde <i>p</i> -Mentha-1,8-dien-7-al | 2111-75-3 | 1:3 EtOH:DEP | 0.5, 1.0, 2.5, 5.0, 10.0 | 5 | 9.3 | 2325 | 0.62 | | | RIFM, 2008b |
| Perillaldehyde <i>p</i> -Mentha-1,8-dien-7-al | 2111-75-3 | - | - | - | 8.1 | 2025 | 0.54 | 8.1 | | Roberts et al., 2007 |
| Balsam oil, Peru (<i>Myroxylon pereirae</i> Klotzsch) | 8007-00-9 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | 3.95 | 987 | N/a | 3.95 | | RIFM, 2004h |
| Peru balsam absolute | 8007-00-9 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | 2.5 | 625 | N/a | 2.5 | | RIFM, 2004i |
| Peru balsam absolute | 8007-00-9 | 1:3 EtOH:DEP | 0.5, 1.0, 2.5 | 4 | >2.5 | >625 | N/a | | | RIFM, 2004i |
| Phenylacetaldehyde | 122-78-1 | 4:1 AOO | 2.5, 5, 10, 25, 50 | 4 | 3.0 | 750 | 0.25 | 3.0 | | Basketter et al., 2001 |

Opinion on fragrance allergens in cosmetic products

Annex II . Local lymph node assay (LLNA) data on 59 fragrance substances, based on a summary report submitted by the Research Institute for Fragrance Materials, Inc. (RIFM, 2009)

| | | | | | | | | | | |
|--|-------------|--------------|----------------------------|--------|-------|-------|-------|-------|--|------------------------|
| <i>Phenylacetaldehyde</i> | 122-78-1 | - | - | - | - | - | - | - | No data in the ref (poster abstract) | Basketter et al., 2003 |
| <i>3-Propylidenephthalide</i> | 17369-59-4 | 4:1 AOO | 5, 10, 20 | 4 or 5 | 3.7 | 925 | 0.21 | 3.7 | Should also have been tested at lower concentrations | Gerberick et al., 2004 |
| Tetramethyl acetyloctahydronaphthalenes (OTNE) | 54464-57-2 | 1:3 EtOH:DEP | 2.5, 5.0, 10.0, 25.0, 50.0 | 4 | 25.14 | 6285 | 1.07 | 25.14 | | RIFM, 2005i |
| Trimethylbenzenepropanol <i>Majantol</i> | 103694-68-4 | 4:1 AOO | 3.0, 10.0, 30.0 | 4 | ~30 | ~7500 | ~1.68 | 30 | Should have been tested at higher concentrations | RIFM, 2002d |
| Vanillin | 121-33-5 | 4:1 AOO | 2.5, 5, 10, 25, 50 | 4 | >50.0 | >1250 | >3.3 | >50.0 | | Basketter et al., 2001 |

* source of EC3 value value: % given in the RIFM report or references; µg/cm2 given in the RIFM report and RIFM poster; M calculated by SCCS working group

**material with low levels of oxidation according to RIFM, 2009

- = no data given; A216

References

- Aptula, N., Roberts, D.W., Schultz, T.W., Pease, C., 2007. Reactivity assays for non-animal based prediction of skin sensitisation potential. *Toxicology*, 231(2-3), 117-118
- Ashby J, Basketter D.A., Patton, D., Kimber I. 1995. Structure activity relationships in skin sensitization using the murine local lymph node assay. *Toxicology* 103:177-194
- 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
- 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
- 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
- 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
- Estrada, E., Patlewicz, G., Chamberlain, M., Basketter, D., Larbey, S., 2003. Computer aided Knowledge Generation for Understanding Skin Sensitization Mechanisms: The TOPS-MODE Approach. *Chem. Res. Toxicol.*, 16, 1226-1235
- 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 skin-sensitization testing. *Contact Dermatitis* 50, 274-288
- 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
- Nilsson, A.-M., Bergstrom, M.A., Luthman, K., Nilsson, J.L.G., Karlberg, A.-T., 2005. An alpha,beta-unsaturated oxime identified as a strong contact allergen. Indications of antigen formation via several pathways. *Food and Chemical Toxicology*, 43(11), 1627-1636
- RIFM, 2001a. Local Lymph Node Assay on p-t-Butyl- α -methyl-hydrocinnamic aldehyde in EtOH . RIFM report number 37065 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001b. Local Lymph Node Assay on p-t-Butyl- α -methyl-hydrocinnamic aldehyde in DEP. RIFM report number 37066 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001c. Local Lymph Node Assay on p-t-Butyl- α -methyl-hydrocinnamic aldehyde in 1:3 EtOH:DEP. RIFM report number 37067 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001d. Local Lymph Node Assay on p-t-Butyl- α -methyl-hydrocinnamic aldehyde in 1:3 DEP:EtOH. RIFM report number 37068 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001e. Local Lymph Node Assay on p-t-Butyl- α -methyl-hydrocinnamic aldehyde in 4:1 acetone:olive oil. RIFM report number 41235. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001f. Local Lymph Node Assay on eugenol. RIFM report number 37076. (RIFM, Woodcliff Lake, NJ, USA)

- RIFM, 2001g. Local Lymph Node Assay on eugenol. RIFM report number 37075. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001h. Local Lymph Node Assay on eugenol. RIFM report number 37073. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001i. Local Lymph Node Assay on eugenol. RIFM report number 37074. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001j. Local Lymph Node Assay on geraniol in ethanol. RIFM report number 37069 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001k. Local Lymph Node Assay on geraniol in DEP. RIFM report number 37070 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001l. Local Lymph Node Assay on geraniol in 1:3 EtOH:DEP. RIFM report number 37071 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001m. Local Lymph Node Assay on geraniol in 3:1 EtOH:DEP. RIFM report number 37072 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001n. Local Lymph Node Assay on hydroxycitronellal in 1:3 EtOH:DEP. RIFM report number 37079 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001o. Local Lymph Node Assay on hydroxycitronellal in DEP. RIFM report number 37078 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001p. Local Lymph Node Assay on hydroxycitronellal in 3:1 EtOH:DEP. RIFM report number 37080 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001q. Local Lymph Node Assay on hydroxycitronellal in EtOH. RIFM report number 37080 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001r. Local Lymph Node Assay on p-t-Butyl- α -methyl-hydrocinnamic aldehyde in 4:1 acetone:olive oil. RIFM report number 41235. Unpublished report from Unilever. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001s. Local Lymph Node Assay on isoeugenol in 4:1 acetone:olive oil. RIFM report number 59516. Unpublished report from Firmenich. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001t. Local Lymph Node Assay on isoeugenol in 4:1 acetone:olive oil. RIFM report number 42122. Unpublished report from Firmenich. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001u. Local Lymph Node Assay on isoeugenol in 4:1 acetone:olive oil. RIFM report number 40676. Unpublished report from Firmenich. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001v. Local Lymph Node Assay on isoeugenol in 4:1 acetone:olive oil. RIFM report number 42120. Unpublished report from Firmenich. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2001w. Local Lymph Node Assay on p-isobutyl- α -methyl hydrocinnamaldehyde in 70% Ethanol. RIFM report number 41055. Unpublished report from Givaudan. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2002a. Local Lymph Node Assay on isoeugenol in 4:1 acetone:olive oil. RIFM report number 42139. Unpublished report from Firmenich. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2002b. Local Lymph Node Assay on isoeugenol in 4:1 acetone:olive oil. RIFM report number 42145. Unpublished report from Firmenich. (RIFM, Woodcliff Lake, NJ, USA)

- RIFM, 2002c. Local Lymph Node Assay on isoeugenol in 4:1 acetone:olive oil. RIFM report number 42123. Unpublished report from Firmenich. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2002d. Local Lymph Node Assay on majantol in 4:1 acetone:olive oil. RIFM report number 58693. Unpublished report from Symrise. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003a. Local Lymph Node Assay on cinnamic aldehyde in 3:1 EtOH:DEP. RIFM report number 42032 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003b. Local Lymph Node Assay on cinnamic aldehyde in 3:1 EtOH:DEP with 0.1% tocopherol. RIFM report number 42033 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003c. Local Lymph Node Assay on cinnamic aldehyde in 3:1 EtOH:DEP with 2.0% tocopherol. RIFM report number 42040 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003d. Local Lymph Node Assay on cinnamic aldehyde in 3:1 EtOH:DEP with antioxidant mix. RIFM report number 42034 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003e. Local Lymph Node Assay on cinnamic aldehyde in 3:1 EtOH:DEP with 0.1% Trolox C. RIFM report number 42036 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003f. Local Lymph Node Assay on cinnamic aldehyde in 3:1 EtOH:DEP with 2.0% tocopherol. RIFM report number 42035 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003g. Local Lymph Node Assay on cinnamic aldehyde in 3:1 EtOH:DEP. RIFM report number 42037 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003h. Local Lymph Node Assay on cinnamic aldehyde in 3:1 EtOH:DEP with 0.1% tocopherol. RIFM report number 42038 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003i. Local Lymph Node Assay on cinnamic aldehyde in 3:1 EtOH:DEP with antioxidant mix. RIFM report number 42039 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003j. Local Lymph Node Assay on cinnamic aldehyde in 3:1 EtOH:DEP with 0.1% Trolox C. RIFM report number 42041 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003k. Local Lymph Node Assay on citral in 3:1 EtOH:DEP with 0.1% tocopherol. RIFM report number 42028 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003l. Local Lymph Node Assay on citral in 3:1 EtOH:DEP with antioxidant mix. RIFM report number 42025 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003m. Local Lymph Node Assay on citral in 3:1 EtOH:DEP with antioxidant mix. RIFM report number 42026 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003n. Local Lymph Node Assay on citral in 3:1 EtOH:DEP. RIFM report number 42023 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003o. Local Lymph Node Assay on citral in 3:1 EtOH:DEP with antioxidant mix. RIFM report number 42029 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003p. Local Lymph Node Assay on citral in 3:1 EtOH:DEP with antioxidant mix. RIFM report number 42027 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003q. Local Lymph Node Assay on citral in 3:1 EtOH:DEP with 0.1% Trolox C. RIFM report number 42030 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003r. Local Lymph Node Assay on citral. RIFM report number 43822 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003s. Local Lymph Node Assay on citral. RIFM report number 42024 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2003t. Local Lymph Node Assay on geraniol in 3:1 EtOH:DEP. RIFM report number 43812 (RIFM, Woodcliff Lake, NJ, USA)

- RIFM, 2004a. Local Lymph Node Assay on α -amylcinnamyl alcohol. RIFM report number 45128 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004b. Local Lymph Node Assay on Citral. RIFM report number 45126 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004c. Local Lymph Node Assay on d,l-Citronellol. RIFM report number 48752 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004d. Local Lymph Node Assay on farnesol RIFM report number 47136. Unpublished report from Symrise (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004e. Local Lymph Node Assay on isoeugenol RIFM report number 47326. Unpublished report from Firmenich (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004f. Local Lymph Node Assay on 4-methoxy- α -methyl benzenpropanal. RIFM report number 47809. Unpublished report from IFF (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004g. Local Lymph Node Assay on 1-Octen-3-yl acetate. RIFM report number 47809. Unpublished report from IFF (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004h. Local Lymph Node Assay on Peru Balsam Oil. RIFM report number 44372. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004i. Local Lymph Node Assay on Peru Balsam Absolute. RIFM report number 44371. (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004j. Oakmoss absolute: Local lymph node assay. RIFM report number 43861 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004k. Treemoss absolute: Local lymph node assay. RIFM report number 44368 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004l. d-limonene: Local lymph node assay. RIFM report number 45756 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004m. d-limonene: Local lymph node assay. RIFM report number 45753 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004n. d-limonene: Local lymph node assay. RIFM report number 45755 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2004o. d-limonene: Local lymph node assay. RIFM report number 45754 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2005a. Local Lymph Node Assay on anisyl alcohol. RIFM report number 45755 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2005b. Local Lymph Node Assay on benzyl alcohol. RIFM report number 47376 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2005c. Local Lymph Node Assay on benzyl benzoate. RIFM report number 47377 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2005d. Local Lymph Node Assay on benzyl cinnamate. RIFM report number 48751 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2005e. Local Lymph Node Assay on benzyl salicylate. RIFM report number 47378 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2005f. Local Lymph Node Assay on cinnamyl nitrile. RIFM report number 51626 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2005g. Local Lymph Node Assay on trans-2-hexenal in 1:3 EtOH:DEP. RIFM report number 48756 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2005h. Local Lymph Node Assay on isocyclogeraniol in 1:3 EtOH:DEP. RIFM report number 48755 (RIFM, Woodcliff Lake, NJ, USA)

- RIFM, 2005i. Local Lymph Node Assay on α -Methyl-1,3-benzodioxole- 5-propionaldehyde in 1:3 EtOH:DEP. RIFM report number 50886 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2005j. Local Lymph Node Assay on α -iso-Methylionone in 1:3 EtOH:DEP. RIFM report number 48749 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2005k. Local Lymph Node Assay on Methyl 2-octynoate in 1:3 EtOH:DEP. RIFM report number 48753 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2005l. Local Lymph Node Assay on OTNE in 1:3 EtOH:DEP. RIFM report number 51630 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2005m. Local Lymph Node Assay on tea leaf absolute. RIFM report number 47597. Unpublished report from Robertet (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2006a. Local Lymph Node Assay on α -amylcinnamaldehyde. RIFM report number 52888 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2006b. Local Lymph Node Assay on hexyl salicylate. RIFM report number 51636 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2006c. Local Lymph Node Assay on isocyclocitral. RIFM report number 52892 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2006d. Local Lymph Node Assay on Jasmine Absolute (Grandiflorum). RIFM report number 53024 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2006e. Local Lymph Node Assay on Jasmine Absolute (Sambac). RIFM report number 52885 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2007b. Local Lymph Node Assay on p-t-Butyl-dihydrocinnamaldehyde. RIFM report number 52900 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2007c. Local Lymph Node Assay on carvone. RIFM report number 52902 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2007d. Local Lymph Node Assay on carvone. RIFM report number 52907 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2007e. Local Lymph Node Assay on dibenzyl ether. RIFM report number 52901 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2007f. Local Lymph Node Assay on Ylang Ylang Extra. RIFM report number 52903 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2008a. Local Lymph Node Assay on 2-hexylidene cyclopentanone. RIFM report number 55548 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2008b. Local Lymph Node Assay on p-mentha-1,8-dien-7-al. RIFM report number 54428 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2008c. Local Lymph Node Assay on menthadiene-7-methyl formate. RIFM report number 54429 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM, 2008d. Local Lymph Node Assay on 6-methyl-3,5-heptadien-2-one. RIFM report number 55564 (RIFM, Woodcliff Lake, NJ, USA)
- RIFM. 2009. Research Institute for Fragrance Materials, Inc. Local lymph node assay (LLNA) protocol summaries: Data presented at the 46th Congress of the European Societies of Toxicology
- 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(7), 1019-1030

Smith, C.K., Hotchkiss, S.A.M., 2001. Allergic Contact Dermatitis. Taylor & Francis, New York

Vocanson, M., Goujon, C., Chabeau, G., Castelain, M., Valeyrie, M., Floch, F, Maliverney, C., Gard A., Nicolas, J.F 2006. The skin allergenic properties of chemicals may depend on contaminants - evidence from studies on coumarin. *International Archives of Allergy and Immunology*, 140, 231-238

Warbrick, E.V.R., Dearman J., Ashby J., Schmezer P. and Kimber I. 2001. Preliminary assessment of the skin sensitizing activity of selected rodent carcinogens using the local lymph node assay. *Toxicology*, 163(1), 63-69

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

Annex III - Tabular summary of dose-elicitation studies in sensitised patients**Contents**

| | |
|---|-----|
| Chloroatranol..... | 316 |
| Cinnamal | 318 |
| Hydroxycitronellal | 321 |
| Hydroxyisohexyl 3-cyclohexenecarboxaldehyde (HICC)..... | 323 |
| Isoeugenol | 329 |
| References | 333 |

Chloroatranol

| Chloroatranol (allergen in oak moss absolute: <i>Evernia prunastri</i>) (1) | |
|---|--|
| Design | blinded, randomised with regard to doses and controlled |
| Test subjects | 13 patients previously identified as sensitized to chloroatranol and oak moss absolute |
| Controls | 10 healthy controls |
| Substance | Purity: >99% |
| Patch test | 15 µl solution applied in an 8 mm Finn Chamber occlusion 48 h |
| -dilution steps | 200 ppm to 0.0063 ppm (10 steps) |
| -control/vehicle | ethanol |
| -definition of threshold | lowest concentration giving a visible skin reaction |
| ROAT | volar aspect of forearms |
| area | 3 x 3 cm ² |
| applications/day | two |
| dose | chloroatranol in ethanol: Step 1: 5 ppm Step 2: 25 ppm |
| dose/application/cm ² | step 1: 0.025 µg step2: 0.125 µg |
| control substance | ethanol |
| definition of positive | erythema in at least 25% and at least one papule |
| period | two weeks for each step |
| Results | |
| PT ED10% (95% CI) | 0.013 (0.002-0.03) ppm =0.0004 µg/cm ² |
| PT ED50% (95% CI) | 0.15 (0.077-0.295) ppm =0.0045 µg/cm ² |
| PT no effect level (observed) | / |
| ROAT | Cumulative responses |
| Step 1 (5 ppm) | 12/13 (92%) |
| Step 2 (25 ppm) | 13/13 (100%) |
| Controls | Negative |
| Other information | None relevant |

In a subsequent study chloroatranol and atranol, both ingredients in *Evernia prunastri*, were tested in equimolar concentrations in serial dilution in 10 eczema patients with known sensitization to chloroatranol and oak moss. A positive response was defined as any degree of reaction. Ethanol was included as the control and gave no response. No use tests were done and no control subjects included.

Results: All patients reacted to the highest concentrations of the two substances. For both substances there was a significant dose-dependence and the estimated difference in elicitation potency of chloroatranol relative to atranol was 217%. The dose-response curve is seen in figure 1 below (2).

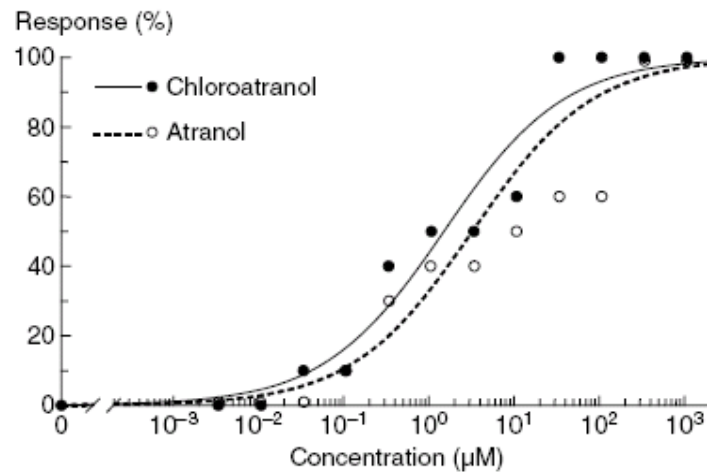


Fig. 1. Observed response rates and fitted parallel logistic dose-response curves for atranol and chloroatranol in equimolar concentrations at patch testing. The response was dichotomized and any reaction other than zero was classified as positive.

Cinnamal

| Cinnamal (3) | |
|----------------------------------|--|
| Design | blinded, randomised and controlled |
| Test subjects | 18 patients with a positive patch test to cinnamal and additional 4 with a doubtful response |
| Controls | 20 healthy controls |
| Substance | Purity: >98% |
| Patch test | 20 mg solution applied in an 8 mm Finn Chamber occlusion 48 h |
| -dilution steps | 2% to 0.01% (7 steps) |
| -control/vehicle | petrolatum |
| -definition of threshold | lowest concentration giving a visible skin reaction in a continuous line of responses |
| ROAT | outer aspect of upper arm |
| area | 5 x 5 cm ² |
| applications/day | two with atomizer pump |
| dose | Step 1: 0.02% Step 2: 0.1% Step 3: 0.8% |
| dose/application/cm ² | Not given |
| control substance | ethanol |
| definition of positive | The response was classified as positive no matter the degree of reaction. |
| period | two weeks for each step; total maximum 6 weeks |
| Results | |
| PT ED10% (95% CI) | / |
| PT ED50% (95% CI) | 0.24% = 96 µg/cm ² (calculated from the data in the paper) |
| PT no effect level(observed) | 0.01 % in pet. = 0.4 µg/cm ² |
| ROAT | Cumulative responses |
| Step 1 (0.02%) | 0/18 |
| Step 2 (0.1%) | 8/18 (44 %) |
| Step 3 (0.8%) | 13/18 (72 %) |
| Controls | No eczema reactions were seen |
| Other information | 2 patients and 2 controls developed immediate reactions to the cinnamal solution |

| | |
|----------------------------------|---|
| Cinnamal (4) | |
| Design | blinded, randomised doses and controlled |
| Test subjects | 17 patients with a positive patch test to cinnamal (8 patients in part 1 and 9 in part two) |
| Controls | 20 controls (non-sensitised dermatitis patients) |
| Substance | purity: / |
| Patch test | 15 µl solution applied in an 8 mm Finn Chamber occlusion 48 h |
| -dilution steps | 2 % to 0.00006 % (17 steps) |
| -control/vehicle | ethanol |
| -definition of threshold | lowest concentration eliciting a + reaction |
| ROAT | Axilla |
| area | 10 x 10 cm ² (estimated) |
| applications/day | two with roll on deodorant (89-700 mg per application of solution) average cases: 263 mg/application controls: only range given |
| dose | Part one: Step 1: 0.032% Step 2: 0.1% Step: 0.32% Part two: Step 1: 0.01% Step 2: 0.032% Step 3: 0.1% |
| dose/application/cm ² | Part two estimated: step one: 0.26 µg; step two: 0.84 µg; 2.63 µg |
| control substance | Deodorant matrix |
| definition of positive | eczematous reaction covering at least 25% of test area |
| period | Part one: one week with each concentration: maximum three weeks Part two: two weeks with each concentration: maximum six weeks |
| Results | |
| PT ED10% (95% CI) | / |
| PT ED50% (95% CI) | / |
| PT no effect level(observed) | 0.002% |
| ROAT | Cumulative responses |
| Step 1 (0.01) | 2/9 (22%) |
| Step 2 (0.032) | 6/9 (67%) |
| Step 3 (0.1) | 8/9 (88%) |
| Controls | No reactions were seen |
| Other information | Only reactions seen to the cinnamal-containing deodorants at ROAT, difference to matrix axilla ($p < 0.001$) and all control |

Opinion on fragrance allergens in cosmetic products

| | |
|--|----------------------------------|
| | persons negative ($p < 0.001$) |
|--|----------------------------------|

Hydroxycitronellal

| Hydroxycitronellal (5) | |
|----------------------------------|---|
| Design | blinded, randomised doses and controlled |
| Test subjects | 7 patients with a positive patch test to hydroxycitronellal |
| Controls | 7 controls (non-sensitised dermatitis patients) |
| Substance | purity: / |
| Patch test | 15 µl solution applied in an 8 mm Finn Chamber occlusion 48 h |
| -dilution steps | 4% to 0.00006% (17 steps) |
| -control/vehicle | ethanol |
| -definition of threshold | lowest concentration eliciting + reaction |
| ROAT | Axilla |
| area | 10 x 10 cm ² (estimated) |
| applications/day | two with roll on deodorant (172-591 per application of solution) average cases: 294 mg/application controls: only range given |
| dose | Step 1: 0.032% Step 2: 0.1% Step: 0.32% |
| dose/application/cm ² | Estimated: step 1: 0.94 µg; step 2: 2.94 µg; step 3: 9.40 µg |
| control substance | Deodorant matrix |
| definition of positive | eczematous reaction covering at least 25% of test area |
| period | two weeks with each concentration: maximum six weeks |
| Results | |
| PT ED10% (95% CI) | / |
| PT ED50% (95% CI) | / |
| PT no effect level(observed) | <0.00012 % |
| ROAT | Cumulative responses |
| Step 1 (0.032) | 4/7 (57%) |
| Step 2 (0.1) | 5/7 (71%) |
| Step 3 (0.32) | 7/7 (100%) |
| Controls | No reactions were seen |
| Other information | Reactions were only seen to the hydroxycitronellal-containing deodorant at ROAT, difference to matrix treated axilla ($p<0.001$) and all control persons negative ($p<0.001$) |

| Hydroxycitronellal (6) | |
|----------------------------------|--|
| Design | double blinded, randomised |
| Test subjects | 13 patients with a positive patch test to hydroxycitronellal |
| Controls | / |
| Substance | purity: unknown |
| Patch test | confirmatory |
| -dilution steps | |
| -control/vehicle | |
| -definition of threshold | |
| ROAT | finger immersion in fragrance solution in 10% ethanol |
| area | / |
| applications/day | Once per day for 10 min |
| dose | Step 1: 10 ppm Step 2: 250 ppm |
| dose/application/cm ² | Not applicable |
| control substance | 10% alcohol |
| definition of positive | clinical grading scale and laser doppler comparison between active and control |
| period | two weeks with each concentration: maximum four weeks |
| Results | |
| PT ED10% (95% CI) | Not relevant |
| PT ED50% (95% CI) | Not relevant |
| PT no effect level(observed) | Not relevant |
| ROAT | Cumulative responses |
| Step 1 (10 ppm) | 1/13 |
| Step 2 (250 ppm) | 5/13 |
| | |
| Vehicle control | 4/13 |
| Other information | No difference between active substance and control application was found. |

Hydroxyisohexyl 3-cyclohexenecarboxaldehyde (HICC)

| Hydroxyisohexyl 3-cyclohexenecarboxaldehyde (HICC) (7) | |
|---|---|
| Design | blinded, randomised and controlled |
| Test subjects | 18 patients with a positive patch test to HICC |
| Controls | 7 healthy controls |
| Substance | Purity: >99% |
| Patch test | 15 µl solution applied in an 8 mm Finn Chamber occlusion 48 h |
| -dilution steps | 6% to 0.0006% |
| -control/vehicle | ethanol |
| -definition of threshold | lowest concentration giving a visible skin reaction in a continuous line of reactions |
| ROAT | volar aspect of lower arm |
| area | 3 x 3 cm ² |
| applications/day | two with droplet bottle (theoretical:30 mg per application of solution) |
| dose | Step 1: 0.5% Step 2: 3% |
| µg/application/cm ² | Step 1: 15.3 (3.4-22.2) Step 2: 126.2 (40.5-226.2) |
| control substance | ethanol |
| definition of positive | erythema in at least 25% and at least one papule |
| period | two weeks for each step; total maximum 4 weeks |
| Results | |
| PT ED10% (95% CI) | 0.9 µg/cm ² 29 (7-69) ppm |
| PT ED50% (95% CI) | 20 µg/cm ² 662 (350-1250)ppm |
| PT no effect level (observed) | / |
| ROAT | Cumulative responses |
| Step 1 (0.5%) | 11/18 (61%) |
| Step 2 (3%) | 16/18 (89%) |
| | |
| Controls | No reactions were seen |
| Other information | Difference between test and control group statistically significant |

| Hydroxyisohexyl 3-cyclohexenecarboxaldehyde (HICC) (8) | |
|---|--|
| Design | blinded, randomised and controlled |
| Test subjects | 15 patients with a positive patch test to HICC |
| Controls | 10 healthy controls |
| Substance | Purity: > 98.8% |
| Patch test | 15 µl solution applied in an 8 mm Finn Chamber occlusion 48 h |
| -dilution steps | 6% to 0.0006% (5 steps) |
| -control/vehicle | ethanol |
| -definition of threshold | lowest concentration giving a visible skin reaction in a continuous line of reactions |
| ROAT | Axilla |
| area | 76 cm ² (template) |
| applications/day | two with roll on deodorant |
| dose | Step 1: 200 ppm Step 2: 600 ppm Step 3: 1800 ppm |
| dose/application/cm ² | median 0.79 µg HICC |
| control substance | deodorant matrix |
| definition of positive | spotty erythema involving at least 25% of the exposed area and infiltration represented by at least one papule. |
| period | two weeks for each step; total maximum 6 weeks |
| Results | |
| PT ED10% (95% CI) | 0.75 µg/cm ² 25 ppm (0.69-120) |
| PT ED50% (95% CI) | 18.3 µg/cm ² 610 ppm (120-2800) |
| PT no effect level (observed) | < 0.0006% |
| ROAT | Cumulative responses |
| Step 1 (200 ppm) | 9/14* (64%) |
| Step 2 (600 ppm) | 12/14* (86%) |
| Step 3 (1800 ppm) | 14/14* (100%) |
| Controls | No reactions were seen |
| Other information | *14 patients completed the use test study Difference between HICC deodorant and matrix deodorant in cases ($p=0.0001$). Difference between controls and patients ($p=0.004$). |

| Hydroxyisohexyl 3-cyclohexenecarboxaldehyde (HICC) (9) | |
|---|--|
| Design | blinded, randomised and controlled |
| Test subjects | 17 patients with a positive patch test to HICC |
| Controls | 15 healthy controls |
| Substance | IFF lot SM/8059062 |
| Patch test | 15 µl solution applied in an 8 mm Finn Chamber occlusion 48 h |
| -dilution steps | 1500 to 0.0022 µg/cm ² HICC (19 steps) |
| -control/vehicle | ethanol |
| -definition of threshold | lowest concentration giving a visible skin reaction in a continuous line of reactions to higher concentrations |
| ROAT | volar aspect of forearms |
| area | 3 x 3 cm (5 areas) |
| applications/day | two with micropipette (20 µl per application) |
| dose | Simultaneous application to 5 areas, four doses each and vehicle |
| µg /application/cm ² | Dose 1:0.0357 Dose 2: 0.357 Dose 3: 3.57 Dose 4: 35.7 |
| control substance | ethanol |
| definition of positive | at least 5 points on a clinical scale, corresponding to erythema in 25% of test area and at least 1 papule |
| period | Three weeks. All concentrations applied simultaneously (randomised) |
| Results | |
| PT ED10% (95% CI) | 0.662 µg/ cm ² (0.052-2.35) |
| PT ED50% (95% CI) | 11.1 µg/ cm ² (3.41- 33.1) |
| PT no effect level(observed) | <0.0022 µg/ cm ² |
| ROAT | Cumulative responses |
| Dose 1 (0.0357) | 0/16* |
| Dose 2 (0.357) | 3/16 (19%) |
| Dose 3 (3.57) | 12/16 (75%) |
| Dose 4 (35.7) | 15/16 (94%) |
| Controls | No reactions were seen |
| Other information | *16 patients completed the use test study The evaporation rate of HICC was calculated to 72% over a 24-h period. ED10% ROAT: 0.064 µg/cm ² (more info see below) |

Opinion on fragrance allergens in cosmetic products

Table 2 The dose per application and accumulated dose after 1, 2 and 3 weeks in the ROAT

| ROAT, dose per application ($\mu\text{g HICC cm}^{-2}$) | Number of applications after 1 week | Total accumulated dose after 1 week ($\mu\text{g HICC cm}^{-2}$) | Number of applications after 2 weeks | Total accumulated dose after 2 weeks ($\mu\text{g HICC cm}^{-2}$) | Number of applications after 3 weeks | Total accumulated dose after 3 weeks ($\mu\text{g HICC cm}^{-2}$) |
|---|-------------------------------------|--|--------------------------------------|---|--------------------------------------|---|
| 35.7 | 14 | 500 | 28 | 1000 | 42 | 1500 |
| 3.57 | 14 | 50 | 28 | 100 | 42 | 150 |
| 0.357 | 14 | 5 | 28 | 10 | 42 | 15 |
| 0.0357 | 14 | 0.5 | 28 | 1 | 42 | 1.5 |

ROAT, repeated open application test; HICC, hydroxyisohexyl-3-cyclohexene carboxaldehyde.

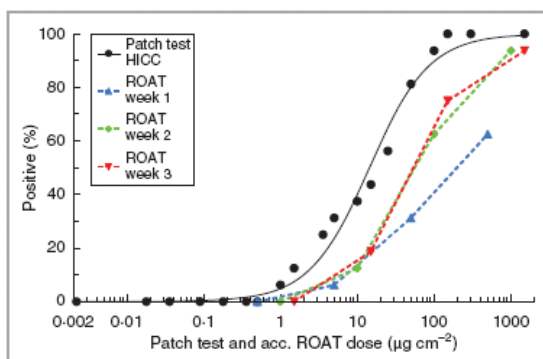


Fig 3. The fitted dose–response curve for the patch test ($n = 16$) and the 1-week, the 2-week 3-week accumulated ROAT doses.

Hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC)

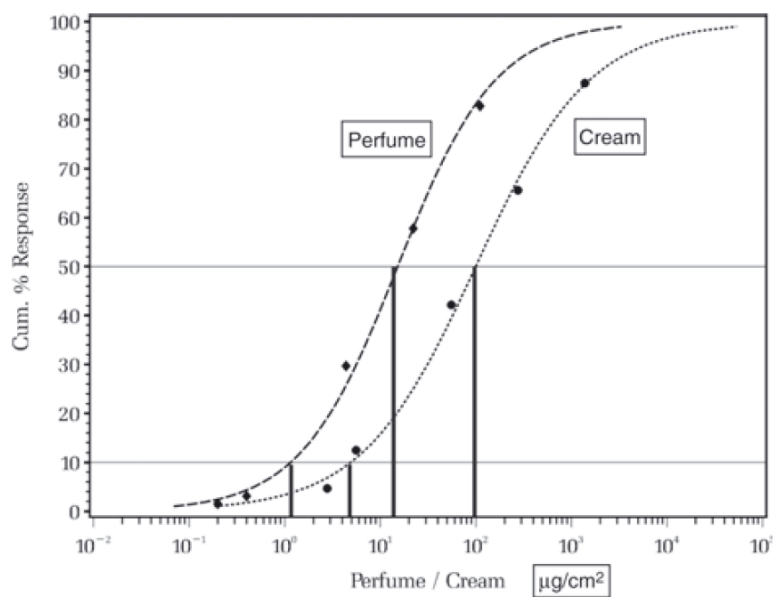
In a study by the German Contact Dermatitis Group, 64 persons previously diagnosed with HICC contact allergy were exposed to increasing doses of HICC in 2 different formulations, a hydrophilic cream and an ethanol solution, to mimic everyday exposures, following a standardised ROAT protocol (10). The concentration of HICC tolerated by 90% of the sensitised was estimated as 1.2 µg/cm² for perfume and 4.9 µg/cm² for cream. The dose-response curve is shown in Fig. 4.3 – 1 below.

| Hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC) (10) | | | | | | | | | | | |
|---|---|---------------------------------|-----------------------------------|---------------------------------|-----------------------------------|----------------------------------|-----------------------------------|-----------------------------------|------------------------------------|------------------------------------|-------------------------------------|
| Design | randomised and vehicle controlled | | | | | | | | | | |
| Test subjects | 67 patients with a previous positive patch test to HICC | | | | | | | | | | |
| Controls | None | | | | | | | | | | |
| Substance | Provided by International Flavor & Fragrances Inc, Hilversum, NL | | | | | | | | | | |
| Patch test | | | | | | | | | | | |
| -dilution steps | 2.5% and 5% | | | | | | | | | | |
| -control/vehicle | petrolatum | | | | | | | | | | |
| -definition of threshold | lowest concentration giving a positive skin reaction in a continuous line to next higher concentration. | | | | | | | | | | |
| ROAT | | | | | | | | | | | |
| area | 3 x 3 cm (4 areas: one test and one control each for alcoholic solution and cream, respectively) | | | | | | | | | | |
| applications/day | two | | | | | | | | | | |
| dose | <table border="0"> <tr> <td>2.8 µg/cm² in cream</td> <td>0.2 µg/cm² in ethanol</td> </tr> <tr> <td>5.6 µg/cm² in cream</td> <td>0.4 µg/cm² in ethanol</td> </tr> <tr> <td>55.6 µg/cm² in cream</td> <td>4.4 µg/cm² in ethanol</td> </tr> <tr> <td>277.8 µg/cm² in cream</td> <td>22.2 µg/cm² in ethanol</td> </tr> <tr> <td>1388.9 µg/cm² in cream</td> <td>111.1 µg/cm² in ethanol</td> </tr> </table> | 2.8 µg/cm ² in cream | 0.2 µg/cm ² in ethanol | 5.6 µg/cm ² in cream | 0.4 µg/cm ² in ethanol | 55.6 µg/cm ² in cream | 4.4 µg/cm ² in ethanol | 277.8 µg/cm ² in cream | 22.2 µg/cm ² in ethanol | 1388.9 µg/cm ² in cream | 111.1 µg/cm ² in ethanol |
| 2.8 µg/cm ² in cream | 0.2 µg/cm ² in ethanol | | | | | | | | | | |
| 5.6 µg/cm ² in cream | 0.4 µg/cm ² in ethanol | | | | | | | | | | |
| 55.6 µg/cm ² in cream | 4.4 µg/cm ² in ethanol | | | | | | | | | | |
| 277.8 µg/cm ² in cream | 22.2 µg/cm ² in ethanol | | | | | | | | | | |
| 1388.9 µg/cm ² in cream | 111.1 µg/cm ² in ethanol | | | | | | | | | | |
| µg /application/cm ² | See above | | | | | | | | | | |
| control substance | Ethanol 96% and glyceryl stearate 15% in water, resp. | | | | | | | | | | |
| definition of positive | (spotty) erythema of at least 25% of the test area along with homogeneous infiltration or papules regardless of the number | | | | | | | | | | |
| period | Two weeks for each step until positive reaction or end of study, whichever occurred first | | | | | | | | | | |
| Results | | | | | | | | | | | |
| PT ED10% (95% CI) | Not calculable; 52 of 60 Patients patch tested positive to 2.5% HICC, 57 / 60 to 5% HICC | | | | | | | | | | |
| PT ED50% (95% CI) | Not calculable | | | | | | | | | | |
| PT no effect level (observed) | Not calculable | | | | | | | | | | |
| ROAT | Cumulative responses: | | | | | | | | | | |
| | <table border="0"> <tr> <td>Cream preparation:</td> <td>Ethanol preparation:</td> </tr> <tr> <td>2.8 µg/cm²: 4.7%</td> <td>0.2 µg/cm²:1.6%</td> </tr> </table> | Cream preparation: | Ethanol preparation: | 2.8 µg/cm ² : 4.7% | 0.2 µg/cm ² :1.6% | | | | | | |
| Cream preparation: | Ethanol preparation: | | | | | | | | | | |
| 2.8 µg/cm ² : 4.7% | 0.2 µg/cm ² :1.6% | | | | | | | | | | |

Opinion on fragrance allergens in cosmetic products

| | | |
|-------------------|---|---|
| | 5.6 $\mu\text{g}/\text{cm}^2$: 12.5% | 0.4 $\mu\text{g}/\text{cm}^2$: 3.1% |
| | 55.6 $\mu\text{g}/\text{cm}^2$: 42.2% | 4.4 $\mu\text{g}/\text{cm}^2$: 29.7% |
| | 277.8 $\mu\text{g}/\text{cm}^2$: 65.6% | 22.2 $\mu\text{g}/\text{cm}^2$: 57.8% |
| | 1388.9 $\mu\text{g}/\text{cm}^2$: 87.5% | 111.1 $\mu\text{g}/\text{cm}^2$: 82.8% |
| Controls | No reactions to vehicle in the patients included into analysis | |
| Other information | See figure below. Three patients were excluded from the study, so results are based on 64 patients. | |

Figure 4.3 – 1: Dose-response curve of 64 patients sensitised to HICC, according to a previous PT, regarding two preparations: perfume and cream, the rhomboid and dot symbol, respectively, indicating the observed response. The curve was fitted by a logistic function (10).



Isoeugenol

| Isoeugenol (11) | |
|--|--|
| Design | blinded, randomised doses and controlled |
| Test subjects | 20 patients with a positive patch test to isoeugenol |
| Controls | 20 healthy controls |
| Substance | purity: 98% |
| Patch test | 20 mg solution applied in an 8 mm Finn Chamber occlusion 48 h |
| -dilution steps | 2% to 0.01% (8 steps) |
| -control/vehicle | petrolatum |
| -definition of threshold | lowest concentration giving a visible skin reaction in a continuous line |
| ROAT | outer aspect of upper arms |
| area | 5 x 5 cm (2 areas: one test and one control) |
| applications/day | two with roll-on |
| dose | 0.2% in ethanol |
| μg /application/cm ² | Doses measured to 0.14 -0.13 mg/application the first 14 days = 5.6 $\mu\text{g}/\text{cm}^2$ |
| control substance | ethanol |
| definition of positive | any degree of reaction |
| period | Two weeks at upper arm and if negative another two weeks including application to base of neck |
| Results | |
| PT ED10% (95% CI) | / |
| PT ED50% (95% CI) | 0.08% 32 $\mu\text{g}/\text{cm}^2$ |
| PT no effect level (observed) | < 0.01% = 0.4 $\mu\text{g}/\text{cm}^2$ |
| ROAT | |
| Dose: 0.2% | 12/19 (63%) |
| Controls | No reactions were seen |
| Other information | |

| Isoeugenol (12) | |
|---------------------------------|--|
| Design | blinded, randomised |
| Test subjects | 27 patients with a positive patch test to isoeugenol |
| Controls | 20 healthy controls |
| Substance | purity: 98% |
| Patch test | 15 µl solution applied in an 8 mm Finn Chamber occlusion 48 h |
| -dilution steps | 2% to 0.00006% (17 steps) |
| -control/vehicle | ethanol |
| -definition of threshold | lowest concentration giving a visible skin reaction in a continuous line of reactions to higher concentrations |
| ROAT | volar aspect of lower arm |
| area | 3 x 3 cm (2 areas) |
| applications/day | two with droplet bottle (30 mg per application) |
| dose | 0.05% in ethanol and 0.2% |
| µg /application/cm ² | Doses were calculated as mean 2.2 µg/cm ² (low conc.) and 9 µg/cm ² (high conc.) |
| control substance | ethanol |
| definition of positive | clear visible erythema |
| period | 28 days |
| Results | |
| PT ED10% (95% CI) | / |
| PT ED50% (95% CI) | / |
| PT no effect level (observed) | < 0.0005% (5 ppm) |
| ROAT | Cumulative responses |
| Dose 1: 0.05% | 10/24 (42%) |
| Dose 2: 0.2% | 16/24 (67%) |
| Controls | No reactions were seen |
| Other information | Response to the low concentration in the ROAT appeared after median 15 days and to the high concentration after median 7 days. |

| Isoeugenol (13) | |
|----------------------------------|--|
| Design | blinded, randomised and controlled |
| Test subjects | 13 patients with a positive patch test to isoeugenol and 4 in part 1 (pre-test) |
| Controls | 10 healthy controls (dermatitis patients) |
| Substance | purity: / |
| Patch test | 15 µl solution applied in an 8 mm Finn Chamber occlusion 48 h |
| -dilution steps | 2% to 0.00006% (w/v) (16 steps) |
| -control/vehicle | ethanol |
| -definition of threshold | lowest concentration eliciting at least + reaction |
| ROAT | Axilla |
| area | 10 x 10 cm ² (estimated) |
| applications/day | two with roll-on deodorant (117-586 mg per application of solution) average cases: 266 mg/application controls: only range given |
| dose | Part 1: Step 1:0.02% Step 2: 0.063% Step 3:0.2% Part 2: Step1:0.0063% Step 2:0.02% Step 3: 0.063% |
| dose/application/cm ² | Part 2: Step 1: 0.167 Step 2: 0.53 Step 3: 1.67 µg/application/cm ² (calculated based on data) |
| control substance | deodorant matrix |
| definition of positive | eczematous response covering 25% of test area |
| period | Part one: one week with each concentration: maximum three weeks Part two: two weeks with each concentration: maximum six weeks |
| Results | |
| PT ED10% (95% CI) | / |
| PT ED50% (95% CI) | / |
| PT no effect level (observed) | <0.0005% (0.15 µg/cm ²) |
| ROAT | |
| Step 1 (0.0063%) | 3/13 (23%) |
| Step 2 (0.02%) | 9/13 (69%) |
| Step 3 (0.063%) | 10/13 (77%) |
| Controls | No reactions were seen |
| Other information | Deodorants containing cinnamal were responsible for all reactions in cinnamal sensitized individuals ($p<0.001$) and all control persons were negative ($p<0.001$) |

References

- 1 Johansen J D, Andersen K E, Svedman C, Bruze M, Bernard G, Gimenez-Arnau E, Rastogi S C, Lepoittevin J P, Menne T. Chloroatranol, an extremely potent allergen hidden in perfumes: a dose-response elicitation study. *Contact Dermatitis* 2003; **49**: 180-4.
- 2 Johansen J D, Bernard G, Gimenez-Arnau E, Lepoittevin J P, Bruze M, Andersen K E. Comparison of elicitation potential of chloroatranol and atranol--2 allergens in oak moss absolute. *Contact Dermatitis* 2006; **54**: 192-5.
- 3 Johansen J D, Andersen K E, Rastogi S C, Menne T. Threshold responses in cinnamic-aldehyde-sensitive subjects: results and methodological aspects. *Contact Dermatitis* 1996; **34**: 165-71.
- 4 Bruze M, Johansen J D, Andersen K E, Frosch P, Lepoittevin J P, Rastogi S, Wakelin S, White I, Menne T. Deodorants: an experimental provocation study with cinnamic aldehyde. *J Am Acad Dermatol* 2003; **48**: 194-200.
- 5 Svedman C, Bruze M, Johansen J D, Andersen K E, Goossens A, Frosch P J, Lepoittevin J P, Rastogi S, White I R, Menne T. Deodorants: an experimental provocation study with hydroxycitronellal. *Contact Dermatitis* 2003; **48**: 217-23.
- 6 Heydorn S, Menne T, Andersen K E, Bruze M, Svedman C, Basketter D, Johansen J D. The fragrance hand immersion study - an experimental model simulating real-life exposure for allergic contact dermatitis on the hands. *Contact Dermatitis* 2003; **48**: 324-30.
- 7 Johansen J D, Frosch P J, Svedman C, Andersen K E, Bruze M, Pirker C, Menne T. Hydroxyisohexyl 3-cyclohexene carboxaldehyde- known as Lylal: quantitative aspects and risk assessment of an important fragrance allergen. *Contact Dermatitis* 2003; **48**: 310-6.
- 8 Jorgensen P H, Jensen C D, Rastogi S, Andersen K E, Johansen J D. Experimental elicitation with hydroxyisohexyl-3-cyclohexene carboxaldehyde-containing deodorants. *Contact Dermatitis* 2007; **56**: 146-50.
- 9 Fischer L A, Menné T, Avnstorp C, Kasting G B, Johansen J D. Hydroxyisohexyl 3-cyclohexene carboxaldehyde allergy: relationship between patch test and repeated open application test thresholds. *Br J Dermatol* 2009; **161**: 560-7.
- 10 Schnuch A, Uter W, Dickel H, Szliska C, Schliemann S, Eben R, Rueff F, Gimenez-Arnau A, Loffler H, Aberer W, Frambach Y, Worm M, Niebuhr M, Hillen U, Martin V, Jappe U, Frosch P J, Mahler V. Quantitative patch and repeated open application testing in hydroxyisohexyl 3-cyclohexene carboxaldehyde sensitive-patients. *Contact Dermatitis* 2009; **61**: 152-62.
- 11 Johansen J D, Andersen K E, Menné T. Quantitative aspects of isoeugenol contact allergy assessed by use and patch tests. *Contact Dermatitis* 1996; **34**: 414-8.
- 12 Andersen K E, Johansen J D, Bruze M, Frosch P J, Goossens A, Lepoittevin J P, Rastogi S, White I, Menne T. The time-dose-response relationship for elicitation of contact dermatitis in isoeugenol allergic individuals. *Toxicol Appl Pharmacol* 2001; **170**: 166-71.
- 13 Bruze M, Johansen J D, Andersen K E, Frosch P, Goossens A, Lepoittevin J P, Rastogi S C, White I, Menne T. Deodorants: an experimental provocation study with isoeugenol. *Contact Dermatitis* 2005; **52**: 260-7.

