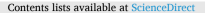
ELSEVIER



# Food and Chemical Toxicology



journal homepage: www.elsevier.com/locate/foodchemtox

## RIFM fragrance ingredient safety assessment, guaiacol, CAS Registry Number 90-05-1

A.M. Api<sup>a</sup>, D. Belsito<sup>b</sup>, D. Botelho<sup>a</sup>, M. Bruze<sup>c</sup>, G.A. Burton Jr.<sup>d</sup>, M.A. Cancellieri<sup>a</sup>, H. Chon<sup>a</sup>, M.L. Dagli<sup>e</sup>, M. Date<sup>a</sup>, W. Dekant<sup>f</sup>, C. Deodhar<sup>a</sup>, A.D. Fryer<sup>g</sup>, L. Jones<sup>a</sup>, K. Joshi<sup>a</sup>, M. Kumar<sup>a</sup>, A. Lapczynski<sup>a</sup>, M. Lavelle<sup>a</sup>, I. Lee<sup>a</sup>, D.C. Liebler<sup>h</sup>, H. Moustakas<sup>a</sup>, M. Na<sup>a</sup>, T.M. Penning<sup>i</sup>, G. Ritacco<sup>a</sup>, J. Romine<sup>a</sup>, N. Sadekar<sup>a</sup>, T.W. Schultz<sup>j</sup>, D. Selechnik<sup>a</sup>, F. Siddiqi<sup>a</sup>, I.G. Sipes<sup>k</sup>, G. Sullivan<sup>a,\*</sup>, Y. Thakkar<sup>a</sup>, Y. Tokura<sup>1</sup>

<sup>b</sup> Member Expert Panel for Fragrance Safety, Columbia University Medical Center, Department of Dermatology, 161 Fort Washington Ave., New York, NY, 10032, USA <sup>c</sup> Member Expert Panel for Fragrance Safety, Malmo University Hospital, Department of Occupational & Environmental Dermatology, Sodra Forstadsgatan 101, Entrance 47. Malmo. SE-20502. Sweden

<sup>d</sup> Member Expert Panel for Fragrance Safety, School of Natural Resources & Environment, University of Michigan, Dana Building G110, 440 Church St., Ann Arbor, MI, 58109. USA

<sup>e</sup> Member Expert Panel for Fragrance Safety, University of Sao Paulo, School of Veterinary Medicine and Animal Science, Department of Pathology, Av. Prof. dr. Orlando Marques de Paiva, 87, Sao Paulo, CEP 05508-900, Brazil

<sup>f</sup> Member Expert Panel for Fragrance Safety, University of Wuerzburg, Department of Toxicology, Versbacher Str. 9, 97078, Würzburg, Germany

<sup>g</sup> Member Expert Panel for Fragrance Safety, Oregon Health & Science University, 3181 SW Sam Jackson Park Rd., Portland, OR, 97239, USA

<sup>h</sup> Member Expert Panel for Fragrance Safety, Vanderbilt University School of Medicine, Department of Biochemistry, Center in Molecular Toxicology, 638 Robinson Research Building, 2200 Pierce Avenue, Nashville, TN, 37232-0146, USA

<sup>i</sup> Member of Expert Panel for Fragrance Safety, University of Pennsylvania, Perelman School of Medicine, Center of Excellence in Environmental Toxicology, 1316 Biomedical Research Building (BRB) II/III, 421 Curie Boulevard, Philadelphia, PA, 19104-3083, USA

<sup>j</sup> Member Expert Panel for Fragrance Safety, The University of Tennessee, College of Veterinary Medicine, Department of Comparative Medicine, 2407 River Dr., Knoxville, TN, 37996- 4500, USA

<sup>k</sup> Member Expert Panel for Fragrance Safety, Department of Pharmacology, University of Arizona, College of Medicine, 1501 North Campbell Avenue, P.O. Box 245050, Tucson, AZ, 85724-5050, USA

<sup>1</sup> Member Expert Panel for Fragrance Safety, The Journal of Dermatological Science (JDS), Editor-in-Chief, Professor and Chairman, Department of Dermatology, Hamamatsu University School of Medicine, 1-20-1 Handayama, Higashi-ku, Hamamatsu, 431-3192, Japan

ARTICLE INFO

Handling editor. Dr. Jose Luis Domingo

Version: 022522. Initial publication. All fragrance materials are evaluated on a five-year rotating basis. Revised safety assessments are published if new relevant data become available. Open access to

(continued on next column)

(continued)

all RIFM Fragrance Ingredient Safety Assessments is here: fragr ancematerialsafetyresource.elsevier. com.

Name: Guaiacol

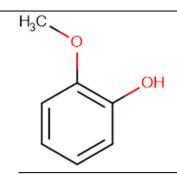
(continued on next page)

\* Corresponding author. E-mail address: gsullivan@rifm.org (G. Sullivan).

https://doi.org/10.1016/j.fct.2022.113168

Received 25 February 2022; Accepted 17 May 2022 Available online 21 May 2022 0278-6915/© 2022 Elsevier Ltd. All rights reserved.

<sup>&</sup>lt;sup>a</sup> Research Institute for Fragrance Materials Inc., 50 Tice Boulevard, Woodcliff Lake, NJ, 07677, USA



CAS Registry Number: 90-05-1

#### Abbreviation/Definition List:

2-Box Model - A RIFM, Inc. proprietary in silico tool used to calculate fragrance air exposure concentration

- AF Assessment Factor
- BCF Bioconcentration Factor
- CNIH Confirmation of No Induction in Humans test. A human repeat insult patch test that is performed to confirm an already determined safe use level for fragrance ingredients (Na et al., 2021)
- Creme RIFM Model The Creme RIFM Model uses probabilistic (Monte Carlo) simulations to allow full distributions of data sets, providing a more realistic estimate of aggregate exposure to individuals across a population (Comiskey et al., 2015, 2017; Safford et al., 2015a, 2017) compared to a deterministic aggregate approach
- DEREK Derek Nexus is an in silico tool used to identify structural alerts
- DRF Dose Range Finding
- DST Dermal Sensitization Threshold
- ECHA European Chemicals Agency
- ECOSAR Ecological Structure-Activity Relationships Predictive Model
- EU Europe/European Union
- GLP Good Laboratory Practice
- IFRA The International Fragrance Association
- LOEL Lowest Observed Effect Level
- MOE Margin of Exposure
- **MPPD** Multiple-Path Particle Dosimetry. An *in silico* model for inhaled vapors used to simulate fragrance lung deposition
- NA North America
- NESIL No Expected Sensitization Induction Level
- NOAEC No Observed Adverse Effect Concentration
- NOAEL No Observed Adverse Effect Level
- NOEC No Observed Effect Concentration
- NOEL No Observed Effect Level
- OECD Organisation for Economic Co-operation and Development
- **OECD TG** Organisation for Economic Co-operation and Development Testing Guidelines
- PBT Persistent, Bioaccumulative, and Toxic
- **PEC/PNEC** Predicted Environmental Concentration/Predicted No Effect Concentration
- **Perfumery** In this safety assessment, perfumery refers to fragrances made by a perfumer used in consumer products only. The exposures reported in the safety assessment include consumer product use but do not include occupational exposures.
- QRA Quantitative Risk Assessment
- QSAR Quantitative Structure-Activity Relationship
- REACH Registration, Evaluation, Authorisation, and Restriction of Chemicals
- RfD Reference Dose
- RIFM Research Institute for Fragrance Materials
- RQ Risk Quotient
- $\label{eq:statistically significant} Statistically significant difference in reported results as compared to controls with a p < 0.05 using appropriate statistical test$
- TTC Threshold of Toxicological Concern

UV/Vis spectra - Ultraviolet/Visible spectra

- VCF Volatile Compounds in Food
- VoU Volume of Use
- $\mathbf{vPvB}$  (very) Persistent, (very) Bioaccumulative
- WoE Weight of Evidence
- The Expert Panel for Fragrance Safety\* concludes that this material is safe as described in this safety assessment.
- This safety assessment is based on the RIFM Criteria Document (Api et al., 2015), which should be referred to for clarifications.
- Each endpoint discussed in this safety assessment includes the relevant data that were available at the time of writing (version number in the top box is indicative of the (continued on next column)

#### (continued)

date of approval based on a 2-digit month/day/year), both in the RIFM Database (consisting of publicly available and proprietary data) and through publicly available information sources (e.g., SciFinder and PubMed). Studies selected for this safety assessment were based on appropriate test criteria, such as acceptable guidelines, sample size, study duration, route of exposure, relevant animal species, most relevant testing endpoints, etc. A key study for each endpoint was selected based on the most conservative endpoint value (e.g., PNEC, NOAEL, LOEL, and NESIL).

\*The Expert Panel for Fragrance Safety is an independent body that selects its own members and establishes its own operating procedures. The Expert Panel is comprised of internationally known scientists that provide RIFM with guidance relevant to human health and environmental protection.

#### Summary: The existing information supports the use of this material as described in this safety assessment.

Guaiacol was evaluated for genotoxicity, repeated dose toxicity, reproductive toxicity, local respiratory toxicity, phototoxicity/photoallergenicity, skin sensitization, and environmental safety. Data show that guaiacol is not genotoxic. Data on read-across analog catechol (CAS # 120-80-9) provide a calculated Margin of Exposure (MOE) > 100 for the repeated dose toxicity and reproductive toxicity endpoints. The skin sensitization endpoint was completed using the Dermal Sensitization Threshold (DST) for reactive materials ( $64 \mu g/cm^2$ ); exposure is below the DST. The phototoxicity/photoallergenicity endpoints were evaluated based on data and ultraviolet/visible (UV/Vis) spectra; guaiacol is not expected to be phototoxic/ photoallergenic. The local respiratory toxicity endpoint was evaluated using the Threshold of Toxicological Concern (TTC) for a Cramer Class I material, and the exposure to guaiacol is below the TTC (1.4 mg/day). The environmental endpoints were evaluated; guaiacol was found not to be Persistent, Bioaccumulative, and Toxic (PBT) as per the International Fragrance Association (IFRA) Environmental Standards, and its risk quotients, based on its current volume of use in Europe and North America (i.e., Predicted Environmental Concentration/Predicted No Effect Concentration [PEC/PNEC]), are <1.

| concentration [1 E0/1 (E0)], are <1.                            |  |  |  |  |
|---|--|--|--|--|
| Human Health Safety Assessment                                  |  |  |  |  |
| Genotoxicity: Not genotoxic.                                    | (ECHA REACH Dossier: Guaiacol; ECHA,         |  |  |  |
|   | 2011)  |  |  |  |
| Repeated Dose Toxicity: $NOAEL =$                               | OECD (2003)                                  |  |  |  |
| 3.3 mg/kg/day.  |  |  |  |  |
| <b>Reproductive Toxicity:</b> NOAEL =                           | (ECHA REACH Dossier: Pyrocatechol;           |  |  |  |
| 160 mg/kg/day.  | ECHA, 2013)                                  |  |  |  |
| Skin Sensitization: Not a concern for skin                      | sensitization under the declared use levels; |  |  |  |
| exposure is below the DST.                                      |  |  |  |  |
| Phototoxicity/Photoallergenicity:                               | (UV/Vis Spectra, RIFM Database; RIFM,        |  |  |  |
| Not phototoxic/not expected to be                               | 2015)  |  |  |  |
| photoallergenic.  |  |  |  |  |
| Local Respiratory Toxicity: No NOAEC                            | available. Exposure is below the TTC.        |  |  |  |
| Environmental Safety Assessment                                 |  |  |  |  |
| Hazard Assessment:  |  |  |  |  |
| Persistence: Critical Measured                                  | (ECHA REACH Dossier: Guaiacol; ECHA,         |  |  |  |
| Value: 97% (OECD 301C)  | 2011)  |  |  |  |
| Bioaccumulation:Screening-level:                                | (EPI Suite v4.11; US EPA, 2012a)             |  |  |  |
| 3.451 L/kg  |  |  |  |  |
| Ecotoxicity:Screening-level: Fish                               | (RIFM Framework; Salvito et al., 2002)       |  |  |  |
| LC50: 653.9 mg/L  |  |  |  |  |
| Conclusion: Not PBT or vPvB as per IFRA Environmental Standards |  |  |  |  |
| Risk Assessment:  |  |  |  |  |
| Screening-level: PEC/PNEC (North                                | (RIFM Framework; Salvito et al., 2002)       |  |  |  |
| America and Europe) $< 1$                                       |  |  |  |  |
| Critical Ecotoxicity Endpoint: Fish                             | (RIFM Framework; Salvito et al., 2002)       |  |  |  |
| LC50: 653.9 mg/L  |  |  |  |  |
| RIFM PNEC is: 0.6539282 μg/L                                    |  |  |  |  |
| • Revised PEC/PNECs (2015 IFRA VoU                              | ): North America and Europe: Not             |  |  |  |
|   |  |  |  |  |

## 1. Identification

- 1. Chemical Name: Guaiacol
- 2. CAS Registry Number: 90-05-1

applicable; cleared at screening-level

- Synonyms: o-Hydroxyanisole; 1-Hydroxy-2-methoxybenzene; o-Methoxyphenol; o-Methylcatechol; Methylcatechol; Phenol, 2methoxy-; Pyroguaiac acid; メトキシフェノール; 2-Methoxyphenol; Guaiacol
- 4. Molecular Formula: C7H8O2
- 5. Molecular Weight: 124.13 g/mol
- 6. RIFM Number: 9

7. Stereochemistry: No stereoisomer possible.

## 2. Physical data

- 1. **Boiling Point:** 205 °C (Fragrance Materials Association), 211.43 °C (EPI Suite)
- 2. Flash Point: 90 °C (Globally Harmonized System)
- 3. Log K<sub>OW</sub>: 1.32 (Smith et al., 2002), 1.34 (Huang et al., 2003), 1.32 ((Abraham and Rafols, 1995)), 1.32 (Smith et al., 2002), 1.34 (EPI Suite)
- 4. Melting Point: 25.15 °C (EPI Suite)
- 5. Water Solubility: 7226 mg/L (EPI Suite)
- 6. Specific Gravity: Not Available
- 7. Vapor Pressure: 0.0664 mm Hg at 20  $^\circ C$  (EPI Suite v4.0), 0.113 mm Hg at 25  $^\circ C$  (EPI Suite)
- 8. UV Spectra: No absorbance between 290 and 700 nm under neutral and acidic conditions. Molar absorption coefficients under the biologically relevant neutral condition (0 L mol<sup>-1</sup> cm<sup>-1</sup>) and acidic conditions (0 L mol<sup>-1</sup> cm<sup>-1</sup>) are below the benchmark (1000 L mol<sup>-1</sup> cm<sup>-1</sup>). The molar absorption coefficient under basic conditions (1140 L mol<sup>-1</sup> cm<sup>-1</sup>) is above the benchmark.
- Appearance/Organoleptic: Colorless, prismatic crystals, or hexagonal prisms with a powerful, smoke-like, somewhat medicinal odor

## 3. Volume of use (Worldwide band)

## 1. 0.1-1 metric ton per year (IFRA, 2015)

# 4. Exposure to fragrance ingredient (Creme RIFM aggregate exposure model v3.1.4)

- 1. 95th Percentile Concentration in Fine Fragrance: 0.00050% (RIFM, 2021)
- 2. Inhalation Exposure\*: 0.000032 mg/kg/day or 0.0023 mg/day (RIFM, 2021)
- 3. Total Systemic Exposure\*\*: 0.00018 mg/kg/day (RIFM, 2021)

\*95th percentile calculated exposure derived from concentration survey data in the Creme RIFM Aggregate Exposure Model (Comiskey et al., 2015; Safford et al., 2015a; Safford et al., 2017; Comiskey et al., 2017).

\*\*95th percentile calculated exposure; assumes 100% absorption unless modified by dermal absorption data as reported in Section V. It is derived from concentration survey data in the Creme RIFM Aggregate Exposure Model and includes exposure via dermal, oral, and inhalation routes whenever the fragrance ingredient is used in products that include these routes of exposure (Comiskey et al., 2015; Safford et al., 2015a; Safford et al., 2017; Comiskey et al., 2017).

## 5. Derivation of systemic absorption

- 1. Dermal: Assumed 100%
- 2. Oral: Assumed 100%
- 3. Inhalation: Assumed 100%

## 6. Computational toxicology evaluation

#### 1. Cramer Classification: Class I, Low

| Expert Judgment | Toxtree v3.1 | OECD QSAR Toolbox v4.2 |
|-----------------|--------------|------------------------|
| Ι               | Ι            | Ι                      |

2. Analogs Selected:

a. Genotoxicity: None

- b. Repeated Dose Toxicity: Catechol (CAS # 120-80-9)
- c. Reproductive Toxicity:: Catechol (CAS # 120-80-9)
- d. Skin Sensitization: None
- e. Phototoxicity/Photoallergenicity: None
- f. Local Respiratory Toxicity: None
- g. Environmental Toxicity: None
- 3. Read-across Justification: See Appendix below

## 7. Metabolism

No relevant data available for inclusion in this safety assessment. Additional References: None.

## 8. Natural occurrence

Guaiacol is reported to occur in the following foods by the VCF\*:

| Beer           | Licorice |
|----------------|----------|
| Cocoa category | Rum      |
| Coffee         | Tomato   |
| Fish           | Whisky   |
| Grape brandy   | Wine     |
|                |          |

\*VCF (Volatile Compounds in Food): Database/Nijssen, L.M.; Ingen-Visscher, C.A. van; Donders, J.J.H. (eds). – Version 15.1 – Zeist (The Netherlands): TNO Triskelion, 1963–2014. A continually updated database containing information on published volatile compounds that have been found in natural (processed) food products. Includes FEMA GRAS and EU-Flavis data.

## 9. REACH dossier

Available; accessed on 10/22/21 (ECHA, 2011).

## 10. Conclusion

The existing information supports the use of this material as described in this safety assessment.

## 11. Summary

## 11.1. Human health endpoint summaries

## 11.1.1. Genotoxicity

Based on the current existing data, guaiacol does not present a concern for genotoxicity.

11.1.1.1. Risk assessment. The mutagenic activity of guaiacol has been evaluated in a bacterial reverse mutation assay conducted in an equivalent manner to OECD TG 471 using the standard plate incorporation method. *Salmonella typhimurium* strains TA98, TA100, TA1535, TA1537, and TA1538 were treated with guaiacol in dimethyl sulfoxide (DMSO) at concentrations up to 5000  $\mu$ g/plate. No increases in the mean number of revertant colonies were observed at any tested concentration in the presence or absence of S9 (ECHA, 2011). Under the conditions of the study, guaiacol was not mutagenic in the Ames test.

The clastogenic activity of guaiacol was evaluated in an *in vivo* micronucleus test conducted in compliance with GLP regulations and in accordance with OECD TG 474. The test material was administered in corn oil via oral gavage to groups of male and female NMRI mice. Doses of 125, 250, or 500 mg/kg body weight were administered. Mice from each dose level were euthanized at 24 and 48 h, and the bone marrow was extracted and examined for polychromatic erythrocytes. The test material did not induce a statistically significant increase in the incidence of micronucleated polychromatic erythrocytes in the bone marrow (ECHA, 2011). Under the conditions of the study, guaiacol was

considered to be not clastogenic in the in vivo micronucleus test.

Based on the data available, guaiacol does not present a concern for genotoxic potential.

Additional References: Florin et al., 1980; Nestmann et al., 1980; Stich et al., 1981; Pool and Lin, 1982; Nestmann and Lee, 1983; Rapson et al., 1980; Haworth et al., 1983; Douglas et al., 1980; Jansson et al., 1986; Ferretti et al., 1977; Tsutsui et al., 1987; Ohshima et al., 1989; Aeschbacher et al., 1989; Rosin (1984); Levan and Tjio, 1948; Hikiba et al., 2005; Miyachi and Tsutsui, 2005; Hamaguchi and Tsutsui, 2000; Someya et al., 2008.

Literature Search and Risk Assessment Completed On: 10/15/21.

#### 11.1.2. Repeated dose toxicity

The MOE for guaiacol is adequate for the repeated dose toxicity endpoint at the current level of use.

11.1.2.1. Risk assessment. There are no repeated dose toxicity data on guaiacol. Read-across material catechol (CAS # 120-80-9; see Section VI) has sufficient data that can be used to support the repeated dose toxicity endpoint. An OECD 422/GLP combined repeated dose toxicity study with reproduction/developmental toxicity screening test was conducted in Wistar rats. Groups of 10 rats/sex/dose were exposed to the test material, catechol, at doses of 30, 80, or 160 mg/kg/day via oral gavage in water once daily, 7 days per week. Males were treated for a minimum of 4 weeks and females for approximately 7 weeks (prior to mating for 2 weeks, through the pairing and gestation periods, until the F1 generation reached day 4 postpartum). Mortality was observed at 160 mg/kg/day; during the pre-pairing period, 1 male and 1 female were found dead on days 3 and 14, respectively, and during the mating period, 1 male was found dead on day 10. Body weight and bodyweight gains were not affected by the treatment for males and females. In males, the level of total bilirubin was statistically significantly increased in the high-dose group as compared to the control. At the high dose, liver weights were statistically significantly increased in males and females. This was considered to be of metabolic nature since only hepatocellular hypertrophy and no liver injury was observed during the histopathological examination. In addition, a statistically significant increase in the absolute weight of kidneys was also observed in males and females. In males, this was caused by the slightly increased severity of hyaline droplets, which were considered to be an incidental increase of a spontaneous lesion. The relative kidney weights for males and females were comparable to the control group. At the high dose, diffuse hepatocellular hypertrophy was recorded at minimal severity in 3 males and 2 females. This was correlated with the significantly increased absolute weight of the liver, as well as with macroscopical findings consisting of enlargement. However, there were no further indicators of liver injury; hence, this lesion was considered to be metabolic and adaptive in nature. In the mid- and high-dose groups, the incidence and severity of squamous hyperplasia in the stomach were increased in both males and females. Thus, the NOAEL for repeated dose toxicity was considered to be 30 mg/kg/day, based on the squamous hyperplasia in the stomach (ECHA, 2013).

In addition, there are carcinogenicity studies available for catechol. Groups of 30 male F344 rats were fed catechol at 0%, 0.1%, 0.2%, 0.4%, and 0.8% (equivalent to 33, 65, 141, and 318 mg/kg/day) for up to 104 weeks. Five rats in each group were euthanized at 34 weeks, and the remaining were euthanized at the end of the treatment period. There were no clinical abnormalities or mortalities related to catechol observed during the treatment. Bodyweight gain was delayed (15%) for the male rats only in the high-dose group. Slight thickening of the pyloric region was observed at 0.4% and 0.8% at week 34. At the end of the study, marked to moderate thickening was also found in rats fed 0.2% of test material and above. Statistically significant adenomas and submucosal hyperplasias of the pyloric glands were developed in rats fed with 0.4% and 0.8% catechol by 34 weeks. Further significant adenomas were developed in rats fed with 0.2% catechol and above by 104 weeks, and significant submucosal hyperplasias were developed in rats at all dose groups. The NOAEL was considered to be 0.1% (33 mg/kg/day) at 34 weeks, and a LOAEL of 0.1% (33 mg/kg/day) was considered at 104 weeks. For 2 year study a NOAEL of 3.3 mg/kg/day was derived by dividing the LOAEL by 10 (33/10 = 3.3 mg/kg/day) (OECD, 2003; Hagiwara et al., 2001).

Therefore, the guaiacol MOE for the reproductive toxicity endpoint can be calculated by dividing the catechol NOAEL in mg/kg/day by the total systemic exposure to guaiacol, 3.3/0.00018, or 18333.

In addition, the total systemic exposure to guaiacol ( $0.18 \mu g/kg/day$ ) is below the TTC ( $30 \mu g/kg/day$ ; Kroes et al., 2007; Laufersweiler et al., 2012) for the repeated dose toxicity endpoint of a Cramer Class I material at the current level of use.

Additional References: None.

Literature Search and Risk Assessment Completed On:  $10/14/\ 21.$ 

## 11.1.3. Reproductive toxicity

The MOE for guaiacol is adequate for the reproductive toxicity endpoint at the current level of use.

11.1.3.1. Risk assessment. There are no reproductive toxicity data on guaiacol. Read-across material catechol (CAS # 120-80-9; see Section VI) has sufficient reproductive toxicity data that can be used to support the reproductive toxicity endpoint. An OECD 422/GLP combined repeated dose toxicity study with reproduction/developmental toxicity screening test was conducted in Wistar rats. Groups of 10 rats/sex/dose were exposed to the test material catechol at doses of 30, 80, or 160 mg/ kg/day via oral gavage in water once daily, 7 days per week. Males were treated for a minimum of 4 weeks and females for approximately 7 weeks (prior to mating for 2 weeks, through the pairing, and gestation periods until the F1 generation reached day 4 postpartum). Mortality was observed at 160 mg/kg/day; during the pre-pairing period, 1 male and 1 female were found dead on days 3 and 14, respectively, and during the mating period, 1 male was found dead on day 10. No treatmentrelated effects were seen with respect to reproductive parameters in males and females. Mean precoital time, conception rate, fertility, and gestation indices were not affected by the treatment. In addition, implantation rate and post-implantation loss were also not affected by the treatment. In the F1 generation, the mean number of pups at birth and on day 4 postpartum was not affected by the treatment at any dose groups. No treatment-related effects were seen in the sex ratio and weight development. At necropsy of pups, no treatment-related findings were noted. Thus, the NOAEL for developmental toxicity and fertility was considered to be 160 mg/kg/day, the highest dose tested (ECHA, 2013).

Therefore, the guaiacol MOE for the developmental toxicity and fertility endpoint can be calculated by dividing the catechol NOAEL in mg/kg/day by the total systemic exposure to guaiacol, 160/0.00018 or 888889.

In addition, the total systemic exposure to guaiacol ( $0.18 \mu g/kg/day$ ) is below the TTC ( $30 \mu g/kg/day$ ; Kroes et al., 2007; Laufersweiler et al., 2012) for the reproductive toxicity endpoint of a Cramer Class I material at the current level of use.

Additional References: None.

Literature Search and Risk Assessment Completed On: 10/14/21.

## 11.1.4. Skin sensitization

Based on existing data and the application of DST, guaiacol does not present a safety concern for skin sensitization under the current, declared levels of use.

#### Table 1

Maximum acceptable concentrations for guaiacol that present no appreciable risk for skin sensitization based on reactive DST.

| IFRA<br>Categorya | Description of<br>Product Type  | Maximum Acceptable<br>Concentrations in<br>Finished Products<br>Based on Reactive<br>DST | Reported 95th<br>Percentile Use<br>Concentrations in<br>Finished Products |
|-------------------|---|--|---|
| 1                 | Products applied to the lips  | 0.0049%  | $4.0\times10^{-4} \%$   |
| 2                 | Products applied to the axillae   | 0.0015%  | $2.0\times10^{-4} \%$   |
| 3                 | Products applied to<br>the face using<br>fingertips   | 0.029%   | $2.0\times10^{-4}\%$  |
| 4                 | Fine fragrance<br>products  | 0.027%   | $5.0\times10^{-4}\%$  |
| 5                 | Products applied to<br>the face and body<br>using the hands<br>(palms), primarily<br>leave-on                     | 0.0070%  | $8.0\times10^{-4}\%$  |
| 6                 | Products with oral<br>and lip exposure  | 0.016%   | 0.0010%   |
| 7                 | Products applied to<br>the hair with some<br>hand contact   | 0.056%   | $8.0\times10^{-4} \%$   |
| 8                 | Products with<br>significant ano-<br>genital exposure   | 0.0029%  | No Datac  |
| 9                 | Products with body<br>and hand exposure,<br>primarily rinse-off   | 0.054%   | 0.0039%   |
| 10                | Household care<br>products with<br>mostly hand contact  | 0.19%  | 0.0039%   |
| 11                | Products with<br>intended skin<br>contact but minimal<br>transfer of fragrance<br>to skin from inert<br>substrate | 0.11%  | No Datac  |
| 12                | Products not<br>intended for direct<br>skin contact,<br>minimal or<br>insignificant<br>transfer to skin           | Not restricted   | 0.5%  |

Note: <sup>a</sup>For a description of the categories, refer to the IFRA/RIFM Information Booklet.

<sup>b</sup>No reported use.

<sup>c</sup>Fragrance exposure from these products is very low. These products are not currently in the Creme RIFM Aggregate Exposure Model.

11.1.4.1. Risk assessment. Limited skin sensitization studies are available for guaiacol. The chemical structure of this material indicates that it would be expected to react with skin proteins directly (Roberts et al., 2007; Toxtree v3.1.0). In a human maximization test, no skin sensitization reactions were observed with 2% or 1380  $\mu$ g/cm<sup>2</sup> guaiacol (RIFM, 1978). Acting conservatively due to the limited data, the reported exposure was benchmarked utilizing the reactive DST of 64  $\mu$ g/cm<sup>2</sup> (Safford, 2008; Safford et al., 2011; Roberts et al., 2015; Safford et al., 2015b). The current exposure from the 95th percentile concentration is below the DST for reactive materials when evaluated in all QRA categories. Table 1 provides the maximum acceptable concentrations for guaiacol that present no appreciable risk for skin sensitization based on the DST approach. However, additional studies may show it could be used at higher levels.

Additional References: (Golberg, 1963); ECHA, 2011.

Literature Search and Risk Assessment Completed On: 10/08/21.

## 11.1.5. Phototoxicity/photoallergenicity

Based on UV/Vis absorbance and *in vitro* study data, guaiacol does not present a concern for phototoxicity. Based on UV/Vis absorbance, guaiacol does not present a concern for photoallergenicity.

11.1.5.1. Risk assessment. UV/Vis absorption spectra indicate no absorption between 290 and 700 nm under the biologically relevant neutral condition, as well as acidic conditions; the corresponding molar absorption coefficients are below the benchmark of concern for phototoxicity and photoallergenicity (Henry et al., 2009). Absorbance was observed under basic conditions, and the molar absorption coefficient was above the benchmark. However, basic conditions for the assay are defined as pH > 10 and may not be biologically relevant for our purposes, where the route of exposure is topical. Furthermore, per the ICH S10 guidance of photosafety Evaluation of Pharmaceuticals, some chromophores, including those with phenolic structures, are considered pH sensitive. In an in vitro 3T3 Neutral Red uptake assay, guaiacol was not predicted to be phototoxic based on mean photo-effect (RIFM, 2015). Based on the in vitro study data and the lack of absorbance under a biologically relevant, neutral pH, guaiacol does not present a concern for phototoxicity. Based on the lack of absorbance under a biologically relevant, neutral pH, guaiacol does not present a concern for photoallergenicity.

11.1.5.2. UV spectra analysis. UV/Vis absorption spectra (OECD TG 101) were obtained. The spectra indicate no absorbance in the range of 290–700 nm under neutral and acidic conditions. The molar absorption coefficients under neutral and acidic conditions (0 L mol<sup>-1</sup> • cm<sup>-1</sup>) are below the benchmark of concern for phototoxic effects, 1000 L mol<sup>-1</sup> • cm<sup>-1</sup> (Henry et al., 2009). Absorbance under the basic condition was greater, and the corresponding molar absorption coefficient (1140 L mol<sup>-1</sup> • cm<sup>-1</sup>) was above the benchmark of concern. However, basic conditions for the assay are defined as a pH of 10 or greater and thus do not represent a biologically relevant condition.

Additional References: None.

Literature Search and Risk Assessment Completed On: 10/15/21.

#### 11.1.6. Local Respiratory Toxicity

The MOE could not be calculated due to a lack of appropriate data. The exposure level for guaiacol is below the Cramer Class I TTC value for inhalation exposure local effects.

11.1.6.1. Risk assessment. There are insufficient inhalation data available on guaiacol. Based on the Creme RIFM Model, the inhalation exposure is 0.0023 mg/day. This exposure is 608.7 times lower than the Cramer Class I TTC value of 1.4 mg/day (based on human lung weight of 650 g; Carthew et al., 2009); therefore, the exposure at the current level of use is deemed safe.

Additional References: Ostrovsky (1964).

Literature Search and Risk Assessment Completed On: 10/15/21.

## 11.2. Environmental endpoint summary

#### 11.2.1. Screening-level assessment

A screening-level risk assessment of guaiacol was performed following the RIFM Environmental Framework (Salvito et al., 2002), which provides 3 tiered levels of screening for aquatic risk. In Tier 1, only the material's regional VoU, its log K<sub>OW</sub>, and its molecular weight are needed to estimate a conservative risk quotient (RQ), expressed as the ratio Predicted Environmental Concentration/Predicted No Effect Concentration (PEC/PNEC). A general QSAR with a high uncertainty factor applied is used to predict fish toxicity, as discussed in Salvito et al. (2002). In Tier 2, the RQ is refined by applying a lower uncertainty factor to the PNEC using the ECOSAR model (US EPA, 2012b), which provides chemical class-specific ecotoxicity estimates. Finally, if necessary, Tier 3 is conducted using measured biodegradation and ecotoxicity data to refine the RQ, thus allowing for lower PNEC uncertainty factors. The data for calculating the PEC and PNEC for this safety assessment are provided in the table below. For the PEC, the range from the most recent IFRA Volume of Use Survey is reviewed. The PEC is then calculated using the actual regional tonnage, not the extremes of the range. Following the RIFM Environmental Framework, guaiacol was identified as a fragrance material with no potential to present a possible risk to the aquatic environment (i.e., its screening-level PEC/PNEC <1).

A screening-level hazard assessment using EPI Suite v4.11 (US EPA, 2012a) did not identify guaiacol as possibly persistent or bioaccumulative based on its structure and physical-chemical properties. This screening-level hazard assessment considers the potential for a material to be persistent and bioaccumulative and toxic, or very persistent and very bioaccumulative as defined in the Criteria Document (Api et al., 2015). As noted in the Criteria Document, the screening criteria applied are the same as those used in the EU for REACH (ECHA, 2012). For persistence, if the EPI Suite model BIOWIN 3 predicts a value < 2.2 and either BIOWIN 2 or BIOWIN 6 predicts a value < 0.5, then the material is considered potentially persistent. A material would be considered potentially bioaccumulative if the EPI Suite model BCFBAF predicts a fish BCF  $\geq$  2000 L/kg. Ecotoxicity is determined in the above screening-level risk assessment. If, based on these model outputs (Step 1), additional assessment is required, a WoE-based review is then performed (Step 2). This review considers available data on the material's physical-chemical properties, environmental fate (e.g., OECD Guideline biodegradation studies or die-away studies), fish bioaccumulation, and higher-tier model outputs (e.g., US EPA's BIOWIN and BCFBAF found in EPI Suite v4.11).

## 11.2.2. Risk assessment

Based on the current Volume of Use (2015), guaiacol does not present a risk to the aquatic compartment in the screening-level assessment.

#### 11.2.2.1. Key studies

- 11.2.2.1.1. Biodegradation. Not available.
- 11.2.2.1.2. Ecotoxicity. Not available.

11.2.1.1.3. Other available data. Guaiacol has been registered for REACH, and the following additional data is available (ECHA, 2011):

The biodegradation of guaiacol was evaluated for 28 days at an initial concentration of 100 mg/L following the OECD 301C method. After 28 days, the measured percentage of biodegradation was 90% (based on Biological Oxygen Demand) and 97% (based on Total Organic Carbon).

The 48-h acute toxicity of guaiacol to *Daphnia magna* was studied under static conditions. The 48-h LC50 was determined graphically by the method of log probit and was found to be 25.9 mg/L.

The 24-h acute toxicity of guaiacol to *Daphnia magna* was studied under static conditions following the AFNOR T90301 of April 1974 modified according to AFNOR T95B DOC 19. The 24-h IC50 was 63 mg/ L.

A 72-h algae acute toxicity study under static conditions was conducted according to the OECD 201 guideline. The growth rate NOEC and EC50 values based on cell density were 10 and > 100 mg/L, respectively.

#### 11.2.3. Risk assessment refinement

Since guaiacol has passed the screening criteria, measured data are included for completeness and have not been used in PNEC derivation.

Ecotoxicological data and PNEC derivation (all endpoints reported in mg/L; PNECs in  $\mu g/L$ ).

Endpoints used to calculate PNEC are underlined.

Exposure information and PEC calculation (following RIFM Framework: Salvito et al., 2002).

| Exposure                            | Europe (EU) | North America (NA) |
|-------------------------------------|-------------|--------------------|
| Log K <sub>ow</sub> Used            | 1.32        | 1.32               |
| Biodegradation Factor Used          | 0           | 0                  |
| Dilution Factor                     | 3           | 3                  |
| Regional Volume of Use Tonnage Band | <1          | <1                 |
| Risk Characterization: PEC/PNEC     | <1          | <1                 |

The RIFM PNEC is  $0.6539282 \ \mu g/L$ . The revised PEC/PNECs for EU and NA are not applicable; therefore, the material does not present a risk to the aquatic environment at the current reported VoU.

Literature Search and Risk Assessment Completed On: 09/28/21.

## Literature Search\*

- **RIFM Database:** Target, Fragrance Structure-Activity Group materials, other references, JECFA, CIR, SIDS
- ECHA: https://echa.europa.eu/
- NTP: https://ntp.niehs.nih.gov/
- OECD Toolbox: https://www.oecd.org/chemicalsafety/risk-assess
   ment/oecd-qsar-toolbox.htm
- SciFinder: https://scifinder.cas.org/scifinder/view/scifinder/scifin derExplore.jsf
- PubMed: https://www.ncbi.nlm.nih.gov/pubmed
- National Library of Medicine's Toxicology Information Services: https://toxnet.nlm.nih.gov/
- IARC: https://monographs.iarc.fr
- OECD SIDS: https://hpvchemicals.oecd.org/ui/Default.aspx
- EPA ACToR: https://actor.epa.gov/actor/home.xhtml
- US EPA HPVIS: https://ofmpub.epa.gov/oppthpv/public\_search. publicdetails?submission\_id=24959241&ShowComments=Yes &sqlstr=null&recordcount=0&User\_title=DetailQuery%20Results &EndPointRpt=Y#submission
- Japanese NITE: https://www.nite.go.jp/en/chem/chrip/chrip\_sear ch/systemTop
- Japan Existing Chemical Data Base (JECDB): http://dra4.nihs.go. jp/mhlw\_data/jsp/SearchPageENG.jsp

|                              | LC50 (Fish)  | EC50          | EC50 (A      | Algae)          | AF      | PNEC (µg/L) | Chemical Class |
|------------------------------|--------------|---------------|--------------|-----------------|---------|-------------|----------------|
|                              | (mg/L)       | (Daphnia)     | (mg/L)       |                 |         |             |                |
|                              |              | (mg/L)        |              |                 |         |             |                |
| RIFM Framework               |              | $\setminus$   |              | /               |         |             |                |
| Screening-level <b>(Tier</b> | <u>653.9</u> | $\mathbf{X}$  | $\mathbf{X}$ |                 | 1000000 | 0.6539      |                |
| 1)                           |              | $/ \setminus$ |              | $\overline{\ }$ |         |             | /              |

#### A.M. Api et al.

- Google: https://www.google.com
- ChemIDplus: https://chem.nlm.nih.gov/chemidplus/

Search keywords: CAS number and/or material names.

\*Information sources outside of RIFM's database are noted as appropriate in the safety assessment. This is not an exhaustive list. The links listed above were active as of 02/25/22.

## Declaration of competing interest

The authors declare that they have no known competing financial

## Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.fct.2022.113168.

## Appendix

Read-across Justification

## Methods

The read-across analog was identified using RIFM fragrance chemicals inventory clustering and read-across search criteria (RIFM, 2020). These criteria are in compliance with the strategy for structuring and reporting a read-across prediction of toxicity as described in Schultz et al. (2015) and are consistent with the guidance provided by OECD within Integrated Approaches for Testing and Assessment (OECD, 2015) and the European Chemical Agency read-across assessment framework (ECHA, 2017).

- First, materials were clustered based on their structural similarity. Second, data availability and data quality on the selected cluster were examined. Third, appropriate read-across analogs from the cluster were confirmed by expert judgment.
- Tanimoto structure similarity scores were calculated using FCFC4 fingerprints (Rogers and Hahn, 2010).
- The physical-chemical properties of the target material and the read-across analogs were calculated using EPI Suite (US EPA, 2012a).
- J<sub>max</sub> values were calculated using RIFM's skin absorption model (SAM). The parameters were calculated using the consensus model (Shen et al., 2014).
- DNA binding, mutagenicity, genotoxicity alerts, and oncologic classification predictions were generated using OECD QSAR Toolbox v4.2 (OECD, 2018).
- ER binding and repeat dose categorization were generated using OECD QSAR Toolbox v4.2 (OECD, 2018).
- Developmental toxicity was predicted using CAESAR v2.1.7 (Cassano et al., 2010), and skin sensitization was predicted using Toxtree v2.6.13.
- Protein binding was predicted using OECD QSAR Toolbox v4.2 (OECD, 2018).
- The major metabolites for the target material and read-across analogs were determined and evaluated using OECD QSAR Toolbox v4.2 (OECD, 2018).
- To keep continuity and compatibility with in silico alerts, OECD QSAR Toolbox v4.2 was selected as the alert system.

|  | Target Material                | Read-across Material  |
|--|--------------------------------|---|
| Principal Name<br>CAS No.  | Guaiacol<br>90-05-1            | Catechol<br>120-80-9  |
| Structure  | HOHO                           | HO  |
| Similarity (Tanimoto Score)<br>Endpoint  |                                | 0.47<br>• Repeated dose<br>toxicity<br>• Reproductive<br>toxicity |
| Molecular Formula<br>Molecular Weight (g/mol)<br>Melting Point (°C, EPI Suite) | $C_7H_8O_2$<br>124.14<br>32.00 | $C_6H_{10}O_2$<br>110.11<br>105.00                                |
| • • • •  |                                | (continued on next page)  |

interests or personal relationships that could have appeared to influence the work reported in this paper. We wish to confirm that there are no known conflicts of interest associated with this publication and there has been no significant financial support for this work that could have influenced its outcome. RIFM staff are employees of the Research Institute for Fragrance Materials, Inc. (RIFM). The Expert Panel receives a small honorarium for time spent reviewing the subject work.

#### (continued)

|  | Target Material  | Read-across Material  |
|--|--|---|
| Boiling Point (°C, EPI Suite)  | 205.00   | 245.50  |
| Vapor Pressure (Pa @ 25°C, EPI Suite)  | 13.73  | 0.49  |
| Water Solubility (mg/L, @ 25°C, WSKOW v1.42 in EPI Suite)  | 18700.00   | 461000.00   |
| Log K <sub>OW</sub>  | 1.32   | 0.88  |
| $J_{max}$ (µg/cm <sup>2</sup> /h, SAM)   | 266.39   | 4556.38   |
| Henry's Law (Pa·m <sup>3</sup> /mol, Bond Method, EPI Suite)                                     | 0.12   | 0.00  |
| Repeated Dose Toxicity   |  |   |
| Repeated Dose (HESS)   | Coumarin<br>(Hepatotoxicity)<br>Alert Toluene<br>(Renal toxicity)<br>Alert | Methyldopa<br>(Hepatotoxicity)<br>Alert Methyldopa<br>(Renal toxicity)<br>Alert Phenols<br>(Mucous membrane<br>irritation) Rank C <br>Styrene (Renal<br>Toxicity) Alert <br>Toluene (Renal<br>toxicity) Alert |
| Reproductive Toxicity  |  | •   |
| ER Binding (OECD QSAR Toolbox V4.2)  | Weak binder, OH<br>group   | Weak binder, OH<br>group  |
| Developmental Toxicity (CAESAR V2.1.6)   | Non-toxicant<br>(moderate<br>reliability)                                  | Non-toxicant (good<br>reliability)  |
| Metabolism   |  |   |
| Rat Liver S9 Metabolism Simulator and Structural Alerts for Metabolites (OECD QSAR Toolbox v4.2) | See<br>Supplemental<br>Data 1  | See Supplemental<br>Data 2  |

## Summary

There are insufficient toxicity data on guaiacol (CAS # 90-05-1). Hence, *in silico* evaluation was conducted to determine read-across analogs for this material. Based on structural similarity, reactivity, physical–chemical properties, and expert judgment, catechol (CAS # 120-80-9) was identified as a read-across analog with sufficient data for toxicological evaluation.

## Conclusions

- Catechol (CAS # 120-80-9) was used as a read-across analog for the target material, guaiacol (CAS # 90-05-1), for the repeated dose toxicity and reproductive toxicity endpoints.
  - o The target material and the read-across analog belong to a class of phenols.
  - o The key difference between the target material and the read-across analog is that the target material has an ortho methoxy substituent, whereas the read-across analog has an ortho hydroxyl substituent. This structural difference makes the read-across analog more reactive than the target material.
  - o The similarity between the target material and the read-across analog is indicated by the Tanimoto score. Differences between the structures that affect the Tanimoto score are toxicologically insignificant.
  - o The physical-chemical properties of the target material and the read-across analog are sufficiently similar to enable comparison of their toxicological properties.
  - o According to the OECD QSAR Toolbox v4.2, structural alerts for toxicological endpoints are consistent between the target material and the readacross analog.
  - o Both the target material and the read-across analog present a renal toxicity alert for the repeated dose (HESS) classification scheme. The data described in the repeated dose toxicity section show that the MOE is adequate at the current level of use. The predictions are superseded by the data.
  - o The target material and the read-across analog are expected to be metabolized similarly, as shown by the metabolism simulator.
  - o The structural alerts for the endpoints evaluated are consistent between the metabolites of the read-across analog and the target material.

## References

- Abraham, M.H., Rafols, C., 1995. Factors that influence tadpole narcosis. An LFER analysis. J. Chem. Soc. Perkin Transact. 2 (10), 1843–1851.
- Aeschbacher, H.U., Wolleb, U., Loliger, J., Spadone, J.C., Liardon, R., 1989. Contribution of coffee aroma constituents to the mutagenicity of coffee. Food Chem. Toxicol. 27 (4), 227–232.
- Api, A.M., Belsito, D., Bruze, M., Cadby, P., Calow, P., Dagli, M.L., Dekant, W., Ellis, G., Fryer, A.D., Fukayama, M., Griem, P., Hickey, C., Kromidas, L., Lalko, J.F., Liebler, D.C., Miyachi, Y., Politano, V.T., Renskers, K., Ritacco, G., Salvito, D., Schultz, T.W., Sipes, I.G., Smith, B., Vitale, D., Wilcox, D.K., 2015. Criteria for the Research Institute for fragrance materials, Inc. (RIFM) safety evaluation process for fragrance ingredients. Food Chem. Toxicol. 82, S1–S19.
- Carthew, P., Clapp, C., Gutsell, S., 2009. Exposure based waiving: the application of the toxicological threshold of concern (TTC) to inhalation exposure for aerosol ingredients in consumer products. Food Chem. Toxicol. 47 (6), 1287–1295.
- Cassano, A., Manganaro, A., Martin, T., Young, D., Piclin, N., Pintore, M., Bigoni, D., Benfenati, E., 2010. CAESAR models for developmental toxicity. Chem. Cent. J. (4 Suppl. 1), S4.
- Comiskey, D., Api, A.M., Barratt, C., Daly, E.J., Ellis, G., McNamara, C., O'Mahony, C., Robison, S.H., Safford, B., Smith, B., Tozer, S., 2015. Novel database for exposure to fragrance ingredients in cosmetics and personal care products. Regul. Toxicol. Pharmacol. 72 (3), 660–672.
- Comiskey, D., Api, A.M., Barrett, C., Ellis, G., McNamara, C., O'Mahony, C., Robison, S. H., Rose, J., Safford, B., Smith, B., Tozer, S., 2017. Integrating habits and practices data for soaps, cosmetics and air care products into an existing aggregate exposure model. Regul. Toxicol. Pharmacol. 88, 144–156.

#### A.M. Api et al.

Douglas, G.R., Nestmann, E.R., Betts, J.L., Mueller, J.C., Lee, E.G.-.H., Stich, H.F., San, R. H.C., Brouzes, R.J.P., Chmelauskas, A., Paavila, H.D., Walden, C.C., 1980. Mutagenic activity in pulp mill effluents. Water Chlorinat. Env. Impact Health Effects 3, 865–880. Ch. 76.

ECHA, 2011. Guaiacol registration dossier. Retrieved from. https://echa.europa.eu/re gistration-dossier/-/registered-dossier/9979/1/2.

ECHA, 2012. Guidance on information requirements and chemical safety assessment. November 2012 v2.1. http://echa.europa.eu/.

- ECHA, 2013. Pyrocatechol registration dossier. Retrieved from. https://echa.europa.eu/ registration-dossier/-/registered-dossier/10516/1/2.
- ECHA, 2017. Read-across assessment framework (RAAF). Retrieved from. https://echa.europa.eu/documents/10162/13628/raaf\_en.pdf/614e5d61-891d-4154-8a47-87efe bd1851a.
- Ferretti, J.J., Lu, W., Liu, M.-B., 1977. Mutagenicity of benzidine and related compounds employed in the detection of hemoglobin. Am. J. Clin. Pathol. 67, 526–527.Florin, I., Rutberg, L., Curvall, M., Enzell, C.R., 1980. Screening of tobacco smoke

FIORIN, I., RUDERS, L., CUTVAI, M., EIIZEH, C.K., 1980. Screening of tobacco sinoke constituents for mutagenicity using the Ames Test. Toxicology 18 (3), 219–232. Golberg, L., 1963. Sensitization to nordihydroguaiaretic acid [Editorial]. Food Chem.

- Toxicol. 1, 114. Hagiwara, A., Takesada, Y., Tanaka, H., Tamano, S., Hirose, M., Ito, N., Shirai, T., 2001. Dose-dependent induction of glandular stomach preneoplastic and neoplastic lesions in male F344 rats treated with catechol chronically. Toxicol. Pathol. 29 (2), 180–186.
- Hamaguchi, F., Tsutsui, T., 2000. Assessment of genotoxicity of dental antiseptics: ability of phenol, guaiacol, p-phenolsulfonic acid, sodium hypochlorite, p-chlorophenol, mcresol or formaldehyde to induce unscheduled DNA synthesis in cultured Syrian hamster embryo cells. Jpn. J. Pharmacol. 83 (3), 273–276.

Haworth, S., Lawlor, T., Mortelmans, K., Speck, W., Zeiger, E., 1983. Salmonella

- mutagenicity test results for 250 chemicals. Environ. Mutagen. 5 (Suppl. 1), 3–142. Henry, B., Foti, C., Alsante, K., 2009. Can light absorption and photostability data be used to assess the photosafety risks in patients for a new drug molecule?
- J. Photochem. Photobiol. B Biol. 96 (1), 57–62.
  Hikiba, H., Watanabe, E., Barrett, J.C., Tsutsui, T., 2005. Ability of fourteen chemical agents used in dental practice to induce chromosome aberrations in Syrian hamster embryo cells. J. Pharmacol. Sci. 97 (1), 146–152.
- Huang, Y.-I., Wang, X., Shao, Y., Chen, D., Dai, X., Wang, L., 2003. QSAR for prediction of joint toxicity of substituted phenols to tadpoles (Rana japonica). Bull. Environ. Contam. Toxicol. 71 (6), 1124–1130.
- IFRA (International Fragrance Association), 2015. Volume of Use Survey, February 2015. Jansson, T., Curvall, M., Hedin, A., Enzell, C.R., 1986. In vitro studies of biological effects of cigarette smoke condensate. II. Induction of sister-chromatid exchanges in human lymphocytes by weakly acidic, semivolatile constituents. Mutat. Res. Genet. Toxicol. 169 (3), 129–139.
- Kroes, R., Renwick, A.G., Feron, V., Galli, C.L., Gibney, M., Greim, H., Guy, R.H., Lhuguenot, J.C., van de Sandt, J.J.M., 2007. Application of the threshold of toxicological concern (TTC) to the safety evaluation of cosmetic ingredients. Food Chem. Toxicol. 45 (12), 2533–2562.

Laufersweiler, M.C., Gadagbui, B., Baskerville-Abraham, I.M., Maier, A., Willis, A., et al., 2012. Correlation of chemical structure with reproductive and developmental toxicity as it relates to the use of the threshold of toxicological concern. Regul. Toxicol. Pharmacol. 62 (1), 160–182.

- Levan, A., Tjio, J.H., 1948. Induction of chromosome fragmentation by phenols. Hereditas (Lund) 34, 453–484.
- Miyachi, T., Tsutsui, T., 2005. Ability of 13 chemical agents used in dental practice to induce sister-chromatid exchanges in Syrian hamster embryo cells. Odontology 93 (1), 24–29.
- Na, M., Ritacco, G., O'Brien, D., Lavelle, M., Api, A., Basketter, D., 2021. Fragrance skin sensitization evaluation and human testing: 30-year experience. Dermatitis 32 (5), 339–352, 2021 Sep-Oct 01.
- Nestmann, E.R., Lee, E.G.-H., 1983. Mutagenicity of constituents of pulp and paper mill effluent in growing cells of saccharomyces cerevisiae. Mutat. Res. Lett. 119 (3–4), 273–280.

Nestmann, E.R., Lee, E.G.-H., Matula, T.I., Douglas, G.R., Mueller, J.C., 1980. Mutagenicity of constituents identified in pulp and paper mill effluents using the salmonella/mammalian-microsome assay. Mutat. Res. Genet. Toxicol. 79 (3), 203–212.

OECD, 2003. SIDS initial assessment profile 1,2-dihydroxybenzene (pyrocatechol, catechol). Retrieved from. https://hpvchemicals.oecd.org/UI/handler.axd?id=0ec bfe38-1c21-4562-aefd-220c345516a3.

- OECD, 2015. Guidance document on the reporting of integrated Approaches to testing and assessment (IATA). ENV/JM/HA(2015)7. Retrieved from. http://www.oecd. org/.
- OECD, 2018. The OECD QSAR Toolbox, v3.2–4.2. Retrieved from. http://www.qsartoo lbox.org/.

- Ohshima, H., Friesen, M., Malaveille, C., Brouet, I., Hautefeuille, A., Bartsch, H., 1989. Formation of direct-acting genotoxic substances in nitrosated smoked fish and meat products: identification of simple phenolic precursors and phenyldiazonium ions as reactive products. Food Chem. Toxicol. 27 (3), 193–203.
- Ostrovsky, M.M., 1964. Toxicology of guaiacol vapors and of its resin. Gigiena i Sanitariia 29 (3), 85-88.
- Pool, B.L., Lin, P.Z., 1982. Mutagenicity testing in the Salmonella typhimurium assay of phenolic compounds and phenolic fractions obtained from smokehouse smoke condensates. Food Chem. Toxicol. 20 (4), 383–391.
- Rapson, W.H., Nazar, M.A., Butsky, V.V., 1980. Mutagenicity produced by aqueous chlorination of organic compounds. Bull. Environ. Contam. Toxicol. 24 (4), 590–596.
- RIFM (Research Institute for Fragrance Materials, Inc.), 1978. Report on Human Maximization Studies. RIFM, Woodcliff Lake, NJ, USA. Report to RIFM. RIFM report number 1787.
- RIFM (Research Institute for Fragrance Materials, Inc.), 2015. Guaiacol: Neutral Red Uptake Phototoxicity Assay in Balb/c 3T3 Mouse Fibroblasts. RIFM, Woodcliff Lake, NJ, USA. RIFM report number 68653.
- RIFM (Research Institute for Fragrance Materials, Inc.), 2020. Clustering a Chemical Inventory for Safety Assessment of Fragrance Ingredients: Identifying Read-Across Analogs to Address Data Gaps. RIFM, Woodcliff Lake, NJ, USA. RIFM report number 76272.
- RIFM, 2021. (Research Institute for fragrance materials, Inc.). Exposure Surv. 31. March 2021.
- Roberts, D.W., Api, A.M., Safford, R.J., Lalko, J.F., 2015. Principles for identification of high potency category chemicals for which the dermal sensitization threshold (DST) approach should not be applied. Regul. Toxicol. Pharmacol. 72 (3), 683–693.
- 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. Chem. Res. Toxicol. 20 (7), 1019–1030.
- Rogers, D., Hahn, M., 2010. Extended-connectivity fingerprints. J. Chem. Inf. Model. 50 (5), 742–754.
- Rosin, M.P., 1984. The influence of pH on the convertogenic activity of plant phenolics. Mutat. Res. Genet. Toxicol. 135 (1), 109–113.

Safford, B., Api, A.M., Barratt, C., Comiskey, D., Daly, E.J., Ellis, G., McNamara, C., O'Mahony, C., Robison, S., Smith, B., Thomas, R., Tozer, S., 2015b. Use of an aggregate exposure model to estimate consumer exposure to fragrance ingredients in personal care and cosmetic products. Regul. Toxicol. Pharmacol. 72, 673–682.

Safford, B., Api, A.M., Barratt, C., Comiskey, D., Ellis, G., McNamara, C., O'Mahony, C., Robison, S., Rose, J., Smith, B., Tozer, S., 2017. Application of the expanded Creme RIFM consumer exposure model to fragrance ingredients in cosmetic, personal care and air care products. Regul. Toxicol. Pharmacol. 86, 148–156.

Safford, R.J., 2008. The dermal sensitisation threshold–A TTC approach for allergic contact dermatitis. Regul. Toxicol. Pharmacol. 51 (2), 195–200.

Safford, R.J., Api, A.M., Roberts, D.W., Lalko, J.F., 2015a. Extension of the dermal sensitization threshold (DST) approach to incorporate chemicals classified as reactive. Regul. Toxicol. Pharmacol. 72 (3), 694–701.

Safford, R.J., Aptula, A.O., Gilmour, N., 2011. Refinement of the dermal sensitisation threshold (DST) approach using a larger dataset and incorporating mechanistic chemistry domains. Regul. Toxicol. Pharmacol. 60 (2), 218–224.

Salvito, D.T., Senna, R.J., Federle, T.W., 2002. A Framework for prioritizing fragrance materials for aquatic risk assessment. Environ. Toxicol. Chem. 21 (6), 1301–1308.

- Schultz, T.W., Amcoff, P., Berggren, E., Gautier, F., Klaric, M., Knight, D.J., Mahony, C., Schwarz, M., White, A., Cronin, M.T., 2015. A strategy for structuring and reporting a read-across prediction of toxicity. Regul. Toxicol. Pharmacol. 72 (3), 586–601.
- Shen, J., Kromidas, L., Schultz, T., Bhatia, S., 2014. An *in silico* skin absorption model for fragrance materials. Food Chem. Toxicol. 74, 164–176.
- Smith, C.J., Perfetti, T.A., Morton, M.J., Rodgman, A., Garg, R., Selassie, C.D., Hansch, C., 2002. The relative toxicity of substituted phenols reported in cigarette mainstream smoke. Toxicol. Sci. 69 (1), 265–278.
- Someya, H., Higo, Y., Ohno, M., Tsutsui, T.W., Tsutsui, T., 2008. Clastogenic activity of seven endodontic medications used in dental practice in human dental pulp cells. Mutat. Res. Genet. Toxicol. Environ. Mutagen 650 (1), 39–47.
- Stich, H.F., Rosin, M.P., Wu, C.H., Powrie, W.D., 1981. The action of transition metals on the genotoxicity of simple phenols, phenolic acids and cinnamic acids. Cancer Lett. 14 (3), 251–260.
- Tsutsui, T., Suzuki, N., Kobayashi, Y., Suzuki, H., Fukuda, S., Maizumi, H., 1987. Assessment of the carcinogenic hazard of 27 substances used in dental practices. Jpn. J. Pharmacol. 43, 132P.
- US EPA, 2012a. Estimation Programs Interface Suite for Microsoft Windows, v4.0–v4.11. United States Environmental Protection Agency, Washington, DC, USA.
- US EPA, 2012b. The ECOSAR (ECOlogical Structure Activity Relationship) Class Program for Microsoft Windows, v2.0. United States Environmental Protection Agency, Washington, DC, USA.