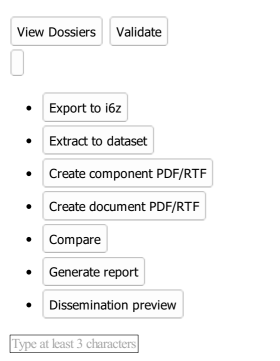
- Dossier preparation manuals
- <u>Q&A</u>
- <u>Create support request</u>
- <u>IUCLID user community</u>
- Additional information
- <u>Video tutorials</u>

SuperUser EPA/ORD/CCTE/SCDCD

- User Settings
- Logout
- Dashboard
- Naphtha (petroleum), steam-cracked ...

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REACH Complete

Naphtha (petroleum), steam-cracked middle arom. [A complex combination of hydrocarbons produced by the distillation of products from a steam-cracking process. It consists predominantly of aromatic hydrocarbons having carbon numbers predominantly in the range of C7 through C12 and boiling in the range of approximately 130°C to 220°C(266°F to 428°F).]

- 1 General information
- 8
- 2 Classification & Labelling and PBT assessment
- 3 Manufacture, use and exposure
- 4 Physical and chemical properties 65
- 5 Environmental fate and pathways 13
- 6 Ecotoxicological information 35

• 7 Toxicological information

181

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- $\circ \ \ eb9c6296‐88f5‐45ac‐a68e‐85172a296bd2$
- 7.1 Toxicokinetics, metabolism and distribution 18
- 7.2 Acute Toxicity
 - 21
- 7.3 Irritation / corrosion
 - 17
- 7.4 Sensitisation
 - 6
- 7.5 Repeated dose toxicity
 - 23
 - 7.5.1 Repeated dose toxicity: oral
 - 12
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- 63e125a0-382a-49b9-895df2a0c3e9e1da
- 7.5.2 Repeated dose toxicity: inhalation
 9
- 7.5.3 Repeated dose toxicity: dermal
 2
- 7.5.4 Repeated dose toxicity: other routes
- 7.6 Genetic toxicity
 - 21
- 7.7 Carcinogenicity
- 7
- 7.8 Toxicity to reproduction 28
- 7.9 Specific investigations
 - 14
- 7.10 Exposure related observations in humans 22
- 7.11 Toxic effects on livestock and pets
- 7.12 Additional toxicological information
- 8 Analytical methods
 - 9 Residues in food and feedingstuffs
- 10 Effectiveness against target organisms
- 11 Guidance on safe use
- 12 Literature search

- 13 Assessment reports
- 14 Information requirements
- Inherited templates

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Compare Docume	ent			
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U				
Administrative data	Data source	Materials and methods	Results and discussion	Overall remarks, attachments
Applicant's summary a	nd conclusion			
Administrative data				
0				
Endpoint sub-chronic toxicity: oral				
Type of information experimental study				
Adequacy of study key study				
Robust study summary				
Used for classification				
Used for SDS				
Study period				
Reliability 1 (reliable without restriction))			
Rationale for reliability incl. d	leficiencies			
Data waiving				
Justification for data waiving				
Justification for type of inform	nation			
Attached justification				
# Attached justification Reas	on / purpose Act	ions		

Cross-reference

Reference

• study report | Unnamed | 1990

Data access

Data protection claimed

Materials and methods

Test guideline

Qualifier Guideline Version / remarks Deviations Actions 1
Qualifier
according to guideline
Guideline
EU Method B.26 (Sub-Chronic Oral Toxicity Test: Repeated Dose 90-Day Oral Toxicity Study in Rodents)
Version / remarks
Deviations

Principles of method if other than guideline

GLP compliance yes Limit test no

Test material

Test material information

• Unnamed | Unnamed | toluene | EC 203-625-9 | 108-88-3

Additional test material information

Specific details on test material used for the study

Specific details on test material used for the study (confidential)

Test animals Species mouse Strain B6C3F1 Details on species / strain selection Sex male/female Details on test animals or test system and environmental conditions

Route of administration oral: gavage

Details on route of administration
Vehicle
Details on oral exposure
Analytical verification of doses or concentrations
Details on analytical verification of doses or concentrations
Duration of treatment / exposure 13 weeks
Frequency of treatment
Doses / concentrations
Dose / conc. Remarks Actions
No. of animals per sex per dose
Control animals
Details on study design
Positive control
Examinations
Observations and examinations performed and frequency
Sacrifice and pathology
Optional endpoint(s)
Other examinations
Statistics
Any other information on materials and methods incl. tables
Results and discussion
Results of examinations
Clinical signs
Description (incidence and severity)
Mortality
Description (incidence)
Body weight and weight changes
Description (incidence and severity)
Food consumption and compound intake (if feeding study)
Description (incidence and severity)

Food efficiency			
Description (incidence and severity)			
Water consumption and compound intake (if drinking water study)			
Description (incidence and severity)			
Ophthalmological findings			
Description (incidence and severity)			
Haematological findings			
Description (incidence and severity)			
Clinical biochemistry findings			
Description (incidence and severity)			
Endocrine findings			
Description (incidence and severity)			
Urinalysis findings			
Description (incidence and severity)			
Behaviour (functional findings)			
Description (incidence and severity)			
Immunological findings			
Description (incidence and severity)			
Organ weight findings including organ / body weight ratios			
Description (incidence and severity)			
Gross pathological findings			
Description (incidence and severity)			
Neuropathological findings			
Description (incidence and severity)			
Histopathological findings: non-neoplastic			
Description (incidence and severity)			
Histopathological findings: neoplastic			
Description (incidence and severity)			
Other effects			
Description (incidence and severity)			
Details on results			
Effect levels			

Key result Dose descriptor Effect level Based on Sex Basis for effect level Remarks on result Actions 1

NOAEL Effect level 625 mg/kg bw/day (nominal) Based on Sex male/female Basis for effect level

• other:

Remarks on result 2 Key result Dose descriptor LOAEL Effect level 1250 mg/kg bw/day (nominal) Based on Sex male/female Basis for effect level

• other:

Remarks on result

Target system / organ toxicity

Key result Critical effects observed Lowest effective dose / conc. System Organ Treatment related Dose response relationship Relevant for humans Actions 1
Key result
Critical effects observed
not specified
Lowest effective dose / conc.

System Organ Treatment related Dose response relationship

Relevant for humans

Any other information on results incl. tables

Overall remarks, attachments

Overall remarks

Attachments

Type Attached (confidential) document Attached (sanitised) documents for publication Remarks Actions

Illustration (picture/graph)

Applicant's summary and conclusion

Conclusions

Executive summary

- Dashboard
- •
- Substances
- Mixtures / Products
- Articles
- •

Categories

Toolbox

- Template
- Manage Reports

Inventory manager

- Contact
- Legal entity
- Sites
- Reference substance
- Test material
- Literature reference

User management

- User Settings
- Users
- Roles

About IUCLID

- About
- Help