



Name: 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol / 118-82-1 / 4,4'-methylenebis(2,6-di-tert-butylphenol) / 118-82-1

Legal entity owner: National Institute of Health Sciences / Kawasaki / Japan

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2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol / 118-82-1

OECD

Health Effects

Repeated dose toxicity: oral

ENDPOINT_STUDY_RECORD: Repeated dose toxicity: oral.001

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Dossier UUID:

Author: SuperUser

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Remarks:

Administrative data

Endpoint

short-term repeated dose toxicity: oral

Type of information

experimental study

Adequacy of study

key study

Robust study summary

false

Used for classification

false

Used for SDS

false

Reliability

1 (reliable without restriction)

Rationale for reliability incl. deficiencies

guideline study

Reliability 1

Data source

Reference

[A 28-day repeat dose oral toxicity study of 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol in rat / Ministry of Health, Labour and Welfare \(MHLW\), Japan / publication](#)

Data access

data published

Materials and methods

Test guideline

Qualifier

according to

Guideline

OECD Guideline 407 (Repeated Dose 28-Day Oral Toxicity in Rodents)

Qualifier

according to

Guideline

other: Study Methods on New Chemical Substances (Chemical Substances Control Law of Japan)

GLP compliance

yes

Test material

Specific details on test material used for the study

2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol / 118-82-1

Test animals

Species

rat

common rodent species

Strain

other: SD [cr1 :CD (SD)]

Sex

male/female

Details on test animals and environmental conditions**TEST ANIMALS**

- Source: Charles River Laboratories Japan, Inc. Atsugi
- Age at study initiation: 5 weeks
- Weight at study initiation: Males: 149-168 g ; Females: 135-155 g
- Housing: bracket-type metallic wire-mesh cages (W 260 × D 380 × H 180 mm)
- Diet: ad libitum
- Water: ad libitum
- Acclimation period: 7 days(male), 8 days(female)

ENVIRONMENTAL CONDITIONS

- Temperature (°C): 22.8-24.7°C (acceptable range: 22±3°C)
- Humidity (%): 51-60 % (acceptable range: 55±10 %)
- Air changes: >10 times / hr
- Photoperiod: 12 hrs dark / 12 hrs light

Administration / exposure

Route of administration

oral: gavage

Vehicle

olive oil

Details on oral exposure**PREPARATION OF DOSING SOLUTIONS:**

Test substance was dissolved in olive oil for injection.

Vehicle

- Name: olive oil (Japanese Pharmacopoeia)
- Lot Number: MI-13
- Manufacturer: Kozakai pharmaceutical Co., Ltd.
- Storage Conditions: Room temperature

Analytical verification of doses or concentrations

yes

Details on analytical verification of doses or concentrations

Test solutions were prepared at least once in 8 days and used within 7 days of preparation.

The test solutions were analyzed concentration after 0days, 4days and 8days preparation.

The results showed that the concentrations were all within the acceptable range.

Duration of treatment / exposure

28 days

Frequency of treatment

once a day

Doses / concentrations

Dose / conc.	
0	mg/kg bw/day (actual dose received)
Dose / conc.	
8	mg/kg bw/day (actual dose received)
Dose / conc.	
40	mg/kg bw/day (actual dose received)
Dose / conc.	
200	mg/kg bw/day (actual dose received)
Dose / conc.	
1000	mg/kg bw/day (actual dose received)

No. of animals per sex per dose

5 or 10/sex/dose

Control animals

yes, concurrent vehicle

Details on study design

Dose selection rationale: Doses in this test were set based on the results of the following dose setting study: a 14-day repeated dose oral toxicity test (doses: 0 (vehicle), 50, 100, 200, 500, and 1000 mg/kg bw/day).

At 100 mg/kg bw/day, significantly lower total cholesterol levels were observed in males and higher relative liver weights in females.

At 200 mg/kg bw/day, significantly higher absolute and relative liver weights were observed in females.

At 500 mg/kg bw/day, significant increases in relative kidney weights were observed in males and relative liver weights in females.

At 1000 mg/kg bw/day, significantly lower total cholesterol levels and higher relative liver weights were observed in males.

At 1000 mg/kg bw/day, significantly higher absolute and relative liver and kidney weights were observed in females.

In this study, the highest dose was 1000 mg/kg bw/day, which is expected to cause toxic effects after repeated administration for 28 days, and the lowest dose was 8 mg/kg bw/day, which is expected to cause no toxic effects.

Therefore, the high dose in this study was set at 1000 mg/kg bw/day, and a low dose of 8 mg/kg bw/day.

- Rationale for animal assignment (if not random): Body weight-balanced randomization

- Post-exposure recovery period in satellite groups: 14 days

Examinations

Observations and examinations performed and frequency

All animals were subjected to detailed clinical observations once before the start of administration. Thereafter, detailed clinical observations were made once a week in dosing and recovery periods.

CAGE SIDE OBSERVATIONS: Yes

Males and females: 4 times/day during the administration period (before and after dosing, 4 hours after dosing), once a day during the recovery period.

DETAILED CLINICAL OBSERVATIONS: Yes

BODY WEIGHT: Yes

Body weights were determined on days 1, 7, 14, 21 and 28 of administration.

Body weights were determined on the day of 7 and 14 of recovery.

Body weights were determined on the day of necropsy.

FOOD CONSUMPTION AND COMPOUND INTAKE (if feeding study): yes

Food consumption was determined on days 5, 12, 19 and 26 of administration for males.

Food consumption was determined on days 5 and 12 of recovery for males.

Food consumption was determined on days 4, 11, 18 and 25 of administration for females.

Food consumption was determined on days 4 and 11 of recovery for females.

OPHTHALMOSCOPIC EXAMINATION: yes

HAEMATOLOGY: Yes

CLINICAL CHEMISTRY: Yes

URINALYSIS: Yes

Animals fasted: no

Sacrifice and pathology

NECROPSY: yes

[brain, pituitary, eye ball, thyroid, spinal cord, heart, trachea, lung, liver, kidney, thymus, spleen, adrenal gland, stomach, ovary, uterus, bladder, sciatic nerve, lymph node, bone marrow]

HISTOPATHOLOGY: yes

Statistics

The data were analyzed for homogeneity of variance by the Bartlett test. If variances were homogeneous, data was analyzed by the Dunnett test, whereas heterogeneous data was analyzed by the steel test.

Comparisons between the two groups were performed by F-test, followed by Student's t-test if variance was uniform, and Aspin-Welch's t-test if variance was not uniform.

Comparisons between multiple groups were performed by Kruskal-Wallis's rank test. When the results showed significant differences, the groups were compared with the control group by the Dunnett type test. Comparisons between the two groups were performed by Mann-Whitney's U-test.

Results and discussion

Results of examinations

Clinical signs

no effects observed

Mortality

no mortality observed

Body weight and weight changes

no effects observed

Food consumption and compound intake (if feeding study)

no effects observed

Haematological findings

no effects observed

Description (incidence and severity)

At 8 mg/kg bw/day in males and females and at 40 mg/kg bw/day in males, low activated partial thromboplastin times were significantly observed at the end of the dosing period.

At 200 mg/kg bw/day, low values of hemoglobin and hematocrit in males and prothrombin time and activated partial thromboplastin time in females were significantly observed.

At 200 mg/kg bw/day, a significant increase in reticulocyte count was observed in females.

However, no dose relationship was observed for these changes.

Clinical biochemistry findings

effects observed, treatment-related

Description (incidence and severity)

At 40 mg/kg bw/day in females, significantly lower values of cholinesterase were observed at the end of the administration period.

At 200 mg/kg bw/day, high levels of LDH and γ -GTP and low total cholesterols and urea-nitrogen were significantly observed in females. However, none of them was dose-related.

At 200 mg/kg bw/day, low creatinine was significantly observed in males. However, the values were within the reference values in the background data and no dose relationship was evident. A significant increase in AST was observed in males at the end of the recovery period. Low total cholesterol levels were significantly observed in males. In females, high levels of γ -GTP and sodium were significantly observed. These changes were considered to be spontaneous, because there were not observed and changes were not dose dependent. and recovered by withdraw.

Urinalysis findings

no effects observed

Description (incidence and severity)

At 200 mg/kg bw/day, a significantly higher number of voids per week of dosing were observed in males. However, these changes were not dose-related, and no changes were observed after Week 2. Therefore, these changes were considered incidental.

Behaviour (functional findings)

no effects observed

Organ weight findings including organ / body weight ratios

effects observed, treatment-related

Description (incidence and severity)

At 40 mg/kg bw/day, higher kidneys relative weights were significantly observed in males in animals sacrificed at the end of the administration period.

At 200 mg/kg bw/day, high absolute and relative liver weights were significantly observed in females. Absolute and relative ovarian and thyroid weights were significantly higher in females than 200 mg/kg bw/day.

At 1000 mg/kg bw/day, a significant increase in relative liver weights was observed in males.

At 1000 mg/kg bw/day, females had significantly lower relative thymic weights.

In the animals sacrificed at the end of the recovery period, high absolute thyroid weights were significantly observed in males, but no changes in relative weights were observed.

Higher relative thymus weights and lower relative pituitary weights were significantly observed in females.

These changes were observed in the organs/tissues, but they were thought to be incidental changes based on the incidence of their occurrence and their histopathological profiles.

Gross pathological findings

no effects observed

Histopathological findings: non-neoplastic

effects observed, treatment-related

Description (incidence and severity)

At the end of the treatment period, in sacrificed animals:

At 200 mg/kg bw/day, diffuse hyperplasia of mild follicular epithelial cells in the thyroid gland was observed in two females.

At 1000 mg/kg bw/day, diffuse hyperplasia of mild follicular epithelial cells in the thyroid gland was significantly observed in 4 females.

Effect levels

Key result

false

Dose descriptor

NOAEL

Effect level

8

mg/kg bw/day (actual dose received)

Based on
test mat.**Sex**
male/female**Basis for effect level**
clinical biochemistry
organ weights and organ / body weight ratios**Target system / organ toxicity**

Key result
false**Critical effects observed**
yes**Lowest effective dose / conc.**

40

mg/kg bw/day (actual dose received)

System
cardiovascular**Organ**
kidney**Treatment related**
yes**Dose response relationship**
yes**Any other information on results incl. tables**

Figures and Tables (in English) are available in the following full report of the study.

http://dra4.nihs.go.jp/mhlw_data/home/pdf/PDF118-82-1b.pdf

Applicant's summary and conclusion

Conclusions

Based on changes in relative kidney weight and blood cholinesterase levels, the NOAEL for repeated-dose toxicity of 4,4'-methylenebis(2,6-di-tert-butylphenol) was determined to be 8 mg/kg bw/day in rats.

Executive summary

The repeated-dose toxicity of 4,4'-methylenebis(2,6-di-tert-butylphenol) was investigated in rats according to the OECD TG 407. Male and female rats (5 or 10 animals/sex/dose) were treated with 4,4'-methylenebis(2,6-di-tert-butylphenol) at doses of 0 (vehicle: olive oil), 8, 40, 200, and 1,000 mg/kg bw/day for 28 days. Five out of the 10 animals/sex treated with 0 and 1,000 mg/kg bw/day were assigned as a recovery group.

There were no deaths in either sex. At doses of ≥ 40 mg/kg bw/day, the relative kidney weight was increased in males, and blood cholinesterase levels were decreased in females. At doses of ≥ 200 mg/kg bw/day, mean corpuscular volume and mean corpuscular hemoglobin levels were decreased in males, and mean corpuscular hemoglobin concentration was decreased in females. Absolute and relative weights of the thyroid gland were increased, and histopathological analysis showed slight diffuse hyperplasia of follicular cells of the thyroid gland was increased with doses of 200 and 1000 mg/kg bw/day. Absolute and relative weights of the ovary were increased without histopathological changes with doses of ≥ 200 mg/kg bw/day. These changes were no longer found after the recovery period. Based on changes in relative kidney weight and blood cholinesterase levels, the NOAEL for repeated-dose toxicity of 4,4'-methylenebis(2,6-di-tert-butylphenol) was determined to be 8 mg/kg bw/day in rats.

ENDPOINT_STUDY_RECORD: Repeated dose toxicity: oral.002

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Author: SuperUser

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Remarks:

Administrative data

Endpoint

short-term repeated dose toxicity: oral

Type of information

experimental study

Adequacy of study

key study

Robust study summary

false

Used for classification

false

Used for SDS

false

Reliability

1 (reliable without restriction)

Rationale for reliability incl. deficiencies

guideline study

Reliability 1

Cross-reference

Reason / purpose

reference to same study

Related information

[OECD / Toxicity to reproduction / Toxicity to reproduction.001 / 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol / 118-82-1 / 4,4'-methylenebis\(2,6-di-tert-butylph...](#)

Remarks

Toxicity to reproduction.001

Data source

Reference

[A reproduction/developmental toxicity screening test in rats treated orally with 2,2',6,6'-tetra-ter... / Ministry of Health, Labour and Welfare \(MHLW\), Japan / publication](#)

Data access

data published

Materials and methods

Test guideline

Qualifier

according to

Guideline

other: OECD Guideline for Testing of Chemicals 421

Qualifier

according to

Guideline

other: Study Methods on New Chemical Substances (Chemical Substances Control Law of Japan)

GLP compliance

yes

Test material

Specific details on test material used for the study

2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol / 118-82-1

Test animals

Species

rat

common rodent species

Strain

other: CrI: CD(SD)

Sex

male/female

Details on test animals and environmental conditions**TEST ANIMALS**

- Source: Charles River Laboratories Japan, Inc. Atsugi
- Age at study initiation: 10 weeks
- Weight at study initiation: Males: 392 - 449g; Females: 228 - 274g
- Housing: bracket-type metallic wire-mesh cages (W 254 × D 350 × H 170 mm)
- Diet: ad libitum
- Water: ad libitum
- Acclimation period: 18 days

ENVIRONMENTAL CONDITIONS

- Temperature (°C): 22-25 (acceptable range:23±3 °C)
- Humidity (%): 38-64 (acceptable range:50±20 %)
- Air changes: 10-15 times / hr
- Photoperiod: 12 hrs dark / 12 hrs light

Administration / exposure

Route of administration

oral: gavage

Vehicle

methylcellulose 0.5 w/v% methylcellulose solution

Details on oral exposure

PREPARATION OF DOSING SOLUTIONS: Test substance was dissolved in 0.5 w/v% methylcellulose solution for injection.

Vehicle

- Name: 0.5 w/v% methylcellulose solution (METOLOSE SM-400, Japanese Pharmacopoeia)
- Lot Number: 8105606
- Manufacturer: Shin-Etsu Chemical Co., Ltd.
- Storage Conditions: In well-closed containers, in a cold and dark place [stored in a refrigerator as the cold and dark place (acceptable values: 1-10°C, measured values: 2-9°C)]

Analytical verification of doses or concentrations

yes

Details on analytical verification of doses or concentrations

Test solutions were prepared at least once in 8 days and used within 7 days of preparation. The test suspensions at all dose concentrations to be used for administration in week 1 or week 6 of administration, dose concentrations and homogeneity were verified by the HPLC method at Gotemba Laboratory, Bozo Research Center Inc. before use for administration. The results showed that the concentrations were 97.5 to 103.5% and the coefficient of variation in the range from 0.5 to 4.6%, both of which were within the acceptable range (concentration: within 100.0 ± 10.0% of the nominal value, homogeneity: CV within 10.0%)

Duration of treatment / exposure

Males: 42 days (14 days prior to mating, 14 days during the mating period and 14 days after the end of the mating period),

Females: 41 to 50 days (14 days prior to mating and throughout the mating and gestation periods until day 4 of lactation)

Non-pregnant females: 53 days

Frequency of treatment

once a day (7 times/week)

Doses / concentrations

Dose / conc.	
0	mg/kg bw/day (actual dose received)
Dose / conc.	
100	mg/kg bw/day (actual dose received)
Dose / conc.	
300	mg/kg bw/day (actual dose received)
Dose / conc.	
1000	mg/kg bw/day (actual dose received)

No. of animals per sex per dose

12 animals/sex/dose

Control animals

yes, concurrent vehicle

Details on study design

The dose levels of this study were selected based on the results of the previously conducted study, "A 14-day oral gavage toxicity study of 2,2',6,6'-tetra- tert-butyl-4,4'-methylenediphenol in rats (dose-finding study)". In that study, there were no toxic changes at 1000 mg/kg, which is the limit test dose prescribed in the OECD Guidelines for Testing of Chemicals 421. Based on these results, the high dose in this reproduction/developmental toxicity screening test was set at 1000 mg/kg, and the lower doses were set at 300 and 100 mg/kg using the common ratio of approximately 3.

- Rationale for animal assignment (if not random): Body weight-balanced randomization
- Post-exposure recovery period in satellite groups: 14 days

Examinations

Observations and examinations performed and frequency

CAGE SIDE OBSERVATIONS: Yes

Males and females: 3 times/day during the administration period (before and after dosing)

DETAILED CLINICAL OBSERVATIONS: Yes

BODY WEIGHT: Yes

males : days 1, 4, 8, 11, 15, 18, 22, 25, 29, 32, 36, 39 and 42 of administration and the day of necropsy,

females : days 1, 4, 8, 11, 15 of administration, days 0, 4, 7, 11, 14, 17 and 20 of gestation, days 0 and 4 of lactation and the day of necropsy

uncopulated females : days 18, 22 and 25 of administration during the mating period),

FOOD CONSUMPTION AND COMPOUND INTAKE (if feeding study):yes

males : days 1, 4, 8, 11, 15, 32, 36, 39 and 42 of administration

females : days 1, 4, 8, 11 and 15 of administration, days 1, 4, 7, 11, 14, 17 and 20 of gestation and days 2 and 4 of lactation.

OPHTHALMOSCOPIC EXAMINATION:yes

HAEMATOLOGY: no

CLINICAL CHEMISTRY: no

URINALYSIS: no

Sacrifice and pathology

Necropsy:yes

[Detailed macroscopic examination: including the external appearance, head, thorax and abdomen]

histopathology: yes

[testes, epididymides, ovaries, uterus, prostate, seminal vesicle, vagina, macroscopic lesions]

Statistics

Group mean with standard deviation was calculated and subjected to a Dunnett-type mean rank test for the difference in mean rank (levels of significance: 0.05 and 0.01, two-tailed).

For the copulation index, insemination index, fertility index and delivery index, the number of copulated animals, number of males impregnated females, number of pregnant females and the number of females that delivered liveborn pups were calculated for each group and analyzed by the chi-square test with Yates' continuity correction (levels of significance: 0.05 and 0.01, two-tailed). If there were cells whose expected frequency was 5 or below, data were analyzed by Fisher's exact test (levels of significance: 0.05 and 0.01, two-tailed) .

Results and discussion

Results of examinations

Clinical signs

no effects observed

Mortality

no mortality observed

Body weight and weight changes

no effects observed

Food consumption and compound intake (if feeding study)

no effects observed

Description (incidence and severity)

In females in the 1000 mg/kg group, significantly low values compared to that of the control group were observed on day 15 of administration and on days 1 and 14 of gestation, but since there were no abnormalities in the body weight development in any period, the changes were thought to be of no toxicological significance.

Organ weight findings including organ / body weight ratios

no effects observed

Gross pathological findings

no effects observed

Description (incidence and severity)

The following changes were observed in the organs/tissues listed below, but they were thought to be incidental changes based on the incidence of their occurrence and their pathological profiles: small testis and epididymis (unilateral) in 1/12 males in the 1000 mg/kg group, white focus in the epididymis in 1/12 males in the 300 mg/kg group, large testis (unilateral) in 1/12 males in the 100 mg/kg group, and diverticulum in the ileum in 1/12 females in the 300 mg/kg group

Histopathological findings: non-neoplastic

no effects observed

Description (incidence and severity)

The some changes were observed in the organs/tissues, but they were thought to be incidental changes based on the incidence of their occurrence and their histopathological profiles.

Any other information on results incl. tables

Figures and Tables (in English) are available in the following full report of the study.

http://dra4.nihs.go.jp/mhlw_data/home/pdf/PDF118-82-1c.pdf

Applicant's summary and conclusion

Executive summary

A reproduction/developmental toxicity screening test (OECD TG 421) was conducted to clarify the effects of 4,4'-methylenebis(2,6-di-tert-butylphenol) on reproductive and developmental toxicity. Male rats were treated with 4,4'-methylenebis(2,6-di-tert-butylphenol) at doses of at 0 [vehicle: 0.5% (w/v) methylcellulose solution], 100, 300, and 1,000 mg/kg bw/day for 14 days prior to mating and throughout the mating period until the day before necropsy (42 days), and female rats were treated for 14 days prior to mating and throughout the mating and gestation periods until day 4 of lactation (41–50 days). There were no mortalities with any dose during the treatment period. There were no adverse effects at 1000 mg/kg/day.

Genetic toxicity in vitro

ENDPOINT_STUDY_RECORD: Genetic toxicity in vitro.001

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Dossier UUID:

Author: SuperUser

Date: 2020-10-09T14:52:02.067+09:00

Remarks:

Administrative data

Endpoint

in vitro gene mutation study in bacteria

Type of information

experimental study

Adequacy of study

key study

Robust study summary

false

Used for classification

false

Used for SDS

false

Reliability

1 (reliable without restriction)

Rationale for reliability incl. deficiencies

guideline study

Reliability 1

Data source

Reference

[A reverse mutation test of 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol using bacteria / Ministry of Health, Labour and Welfare\(MHLW\), Japan / publication](#)

Data access

data published

Materials and methods

Test guideline

Qualifier

according to

Guideline

other: Study Methods on New Chemical Substances (Chemical Substances Control Law of Japan)

Deviations

no

GLP compliance

yes

Type of assay

bacterial reverse mutation assay
in vitro gene mutation study in bacteria

Test material**Specific details on test material used for the study**

2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol / 118-82-1

Method**Species / strain****Species / strain / cell type**

S. typhimurium TA 1535, TA 1537, TA 98 and TA 100
bacteria

Species / strain / cell type

E. coli WP2 uvr A
bacteria

Metabolic activation

with and without

Metabolic activation system

S9 mix, SD male rat liver, induced by phenobarbital (PB) and 5,6-benzoflavone (BF)

Test concentrations with justification for top dose

To set the dose levels for the main tests, a total of 8 dose levels were selected (20, 50, 100, 200, 500, 1000, 2000 and 5000 µg/plate) in the dose-selection test.

In the dose-selection test, growth inhibition by the test substance was not observed all dose levels.

Precipitation by the test substance on the plate was observed at 200 µg/plate and above.

Test were set as the highest dose levels 5000 µg/plate for all tested strains with or without metabolic activation, and a total of 6 dose levels were selected using a common ratio of 2.

Vehicle / solvent

acetone

Controls**Untreated negative controls**

no

Negative solvent / vehicle controls

yes

True negative controls

no

Positive controls

yes

Positive control substance

9-aminoacridine

sodium azide

other: 2-(2-Furyl)-3-(5-nitro-2-furyl)acrylamide (AF-2) 2-Aminoanthracene (2AA)

Details on test system and experimental conditions

METHOD OF APPLICATION: Preincubation

DURATION

- Preincubation period: 20 min

Exposure duration: ca.48 hours

NUMBER OF REPLICATIONS: 3

DETERMINATION OF CYTOTOXICITY

- Method: Cell growth

Evaluation criteria

If two-fold increase in the number of revertant colonies on the test plates or more was observed in comparison with the number of natural revertant colonies (the negative control) and dose response and reproducibility were noted, or if no clear dose response was observed but there was at least two -fold increase in comparison with the number of natural revertant colonies and reproducibility was observed in the two main tests, the test substance was judged to be positive.

Statistics

No statistic method was used for judging of results.

Results and discussion

Test results**Key result**

false

Species / strain

S. typhimurium TA 1535

bacteria

Metabolic activation

with and without

Genotoxicity

negative tested up to max concentration

Cytotoxicity / choice of top concentrations

no cytotoxicity

Vehicle controls validity

valid

Untreated negative controls validity

not examined

True negative controls validity

not examined

Positive controls validity

valid

Key result

false

Species / strain

S. typhimurium TA 1537
bacteria

Metabolic activation

with and without

Genotoxicity

negative tested up to max concentration

Cytotoxicity / choice of top concentrations

cytotoxicity

Vehicle controls validity

valid

Untreated negative controls validity

not examined

True negative controls validity

not examined

Positive controls validity

valid

Key result

false

Species / strain

S. typhimurium TA 98
bacteria

Metabolic activation

with and without

Genotoxicity

negative tested up to max concentration

Cytotoxicity / choice of top concentrations

no cytotoxicity

Vehicle controls validity

valid

Untreated negative controls validity

not examined

True negative controls validity

not examined

Positive controls validity

valid

Key result

false

Species / strain

S. typhimurium TA 100
bacteria

Metabolic activation

with and without

Genotoxicity

negative tested up to max concentration

Cytotoxicity / choice of top concentrations

no cytotoxicity

Vehicle controls validity

valid

Untreated negative controls validity

not examined

True negative controls validity

not examined

Positive controls validity

valid

Additional information on results**TEST-SPECIFIC CONFOUNDING FACTORS**

- Precipitation: Precipitation was observed on plates with concentration of 200 µg/plate or more with and without metabolic activation.

RANGE-FINDING/SCREENING STUDIES:

In range-finding studies, growth inhibition by the test substance was no observed at all dose 20 µg/plate and above for all tested strains.

COMPARISON WITH HISTORICAL CONTROL DATA:

In all test conditions and in all tested strains, the number of revertant colonies of solvent controls and positive controls were within the range of historical control data.

Any other information on results incl. tables _____

Tables in English are attached.

Applicant's summary and conclusion _____**Executive summary**

In a bacterial reverse mutation assay using *S. typhimurium* TA100, TA1535, TA98, and TA1537 and *E. coli* WP2uvrA, 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol was negative with or without metabolic activation.

ENDPOINT_STUDY_RECORD: Genetic toxicity in vitro.002

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Dossier UUID:

Author: SuperUser

Date: 2020-10-09T15:01:44.204+09:00

Remarks:

Administrative data

Endpoint

in vitro cytogenicity / chromosome aberration study in mammalian cells

Type of information

experimental study

Adequacy of study

key study

Robust study summary

false

Used for classification

false

Used for SDS

false

Reliability

1 (reliable without restriction)

Rationale for reliability incl. deficiencies

guideline study

Reliability 1

Data source

Reference

[Chromosome aberration test in cultured chinese hamster cells treated with 2,2',6,6'-tetra-tert-butyl... / Ministry of Health, Labour and Welfare\(MHLW\), Japan / publication](#)

Data access

data published

Materials and methods

Test guideline

Qualifier

according to

Guideline

JAPAN: Guidelines for Screening Mutagenicity Testing Of Chemicals
genetic toxicity in vitro, other

GLP compliance

yes

Type of assay

in vitro mammalian chromosome aberration test

in vitro cytogenicity / chromosome aberration study in mammalian cells

Test material

Specific details on test material used for the study

2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol / 118-82-1

Method

Species / strain**Species / strain / cell type**

Chinese hamster lung (CHL/IU)

mammalian cell line

Metabolic activation

with and without

Metabolic activation system

S9 mix, SD male rat liver, induced by phenobarbital (PB) and 5,6-benzoflavone (BF)

Test concentrations with justification for top dose

+S9 mix(short-term treatment): 0, 250, 500, 1000 µg/mL

-S9 mix(short-term treatment): 0, 250, 500, 1000µg/mL

-S9 mix(24hr-continuous treatment): 0, 250, 500, 1000 µg/mL

-S9 mix(48hr-continuous treatment): 0, 250, 500, 1000µg/ µg/mL

Cell-growth inhibition test was conducted up to the limited concentration of 1000 µg/mL (10 mM)

-Short term treatment and continuous treatment, 50% cell-growth inhibition was not observed any dose(0, 250, 500, 1000 µg/mL)

Vehicle / solvent

acetone

Controls**Untreated negative controls**

no

Negative solvent / vehicle controls

yes

True negative controls

no

Positive controls

yes

Positive control substance

N-ethyl-N-nitro-N-nitrosoguanidine

benzo(a)pyrene

Details on test system and experimental conditions

METHOD OF APPLICATION:

Exposure duration: [continuous treatment]: 24, 48 hrs [short-term treatment]: 6 hrs + 18 hr
SPINDLE INHIBITOR: Colcemid
NUMBER OF REPLICATIONS: 2
NUMBER OF CELLS EVALUATED: 200 cells / dose
DETERMINATION OF CYTOTOXICITY
- Method: relative total growth

Evaluation criteria

For the evaluation of the frequencies of structural aberrations and of polyploidy induced, the following criteria were employed.

Appearance incidence of cells with chromosomal aberrations: Negative (-): < 5%; equivocal (±): 5-10%; positive (+): > 10%.

Finally, the substance is positive when the incidence is considered to be dose-related and reproducible.

Statistics

no

Results and discussion

Test results

Key result

false

Species / strain

Chinese hamster lung (CHL/IU)
mammalian cell line

Metabolic activation

with and without

Genotoxicity

negative :tested up to max concentration

Cytotoxicity / choice of top concentrations

no cytotoxicity

Vehicle controls validity

valid

Untreated negative controls validity

not examined

True negative controls validity

not examined

Positive controls validity

valid

Any other information on results incl. tables

Tables in English are attached.

Applicant's summary and conclusion

Executive summary

It was concluded that 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol is negative for chromosome numerical aberration and chromosome structural aberration under the conditions of this study.

Toxicity to reproduction

ENDPOINT_STUDY_RECORD: Toxicity to reproduction.001

UUID: 937c2fdf-81ea-4c17-90e5-a51010a56e45

Dossier UUID:

Author: SuperUser

Date: 2020-10-09T11:16:03.432+09:00

Remarks:

Administrative data

Endpoint

screening for reproductive / developmental toxicity

Type of information

experimental study

Adequacy of study

key study

Robust study summary

false

Used for classification

false

Used for SDS

false

Reliability

1 (reliable without restriction)

Rationale for reliability incl. deficiencies

guideline study

Reliability 1

Cross-reference

Reason / purpose

reference to same study

Related information

[OECD / Repeated dose toxicity: oral / Repeated dose toxicity: oral.002 / 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol / 118-82-1 / 4,4'-methylenebis\(2,6-di-tert-butylph...](#)

Remarks

Repeated dose toxicity: oral.002

Data source

Reference

[A reproduction/developmental toxicity screening test in rats treated orally with 2,2',6,6'-tetra-ter... / Ministry of Health, Labour and Welfare \(MHLW\), Japan / publication](#)

Data access
data published

Materials and methods

Test guideline

Qualifier

according to

Guideline

OECD Guideline 421 (Reproduction / Developmental Toxicity Screening Test)

Deviations

no

GLP compliance

yes

Test material

Specific details on test material used for the study

2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol / 118-82-1

Test animals

Species

rat

Strain

other: Crl: CD(SD)

Sex

male/female

Details on test animals and environmental conditions**TEST ANIMALS**

- Source: Charles River Laboratories Japan, Inc. Atsugi
- Age at study initiation: 10 weeks
- Weight at study initiation: Males: 392 - 449g; Females: 228 - 274g
- Housing: bracket-type metallic wire-mesh cages (W 254 × D 350 × H 170 mm)
- Diet: ad libitum
- Water: ad libitum
- Acclimation period: 18 days

ENVIRONMENTAL CONDITIONS

- Temperature (°C): 22-25 (acceptable range:23±3 °C)
- Humidity (%): 38-64 (acceptable range:50±20 %)
- Air changes: 10-15 times / hr
- Photoperiod: 12 hrs dark / 12 hrs light

Administration / exposure

Route of administration

oral: gavage

Vehicle

other: 0.5 w/v% methylcellulose solution

Details on mating procedure

M/F ratio per cage:1:1

- Length of cohabitation:up to 14 days

- Proof of pregnancy: [vaginal plug / sperm in vaginal smear] referred to as [day 0] of pregnancy

Analytical verification of doses or concentrations

yes

Details on analytical verification of doses or concentrations

Test solutions were prepared at least once in 8 days and used within 7 days of preparation.

The test suspensions at all dose concentrations to be used for administration in week 1 or week 6 of administration, dose concentrations and homogeneity were verified by the HPLC method at Gotemba Laboratory, Bozo Research Center Inc. before use for administration.

The results showed that the concentrations were 97.5 to 103.5% and the coefficient of variation in the range from 0.5 to 4.6%, both of which were within the acceptable range (concentration: within 100.0 ± 10.0% of the nominal value, homogeneity: CV within 10.0%)

Duration of treatment / exposure

Males: 42 days (14 days prior to mating, 14 days during the mating period and 14 days after the end of the mating period),

Females: 41 to 50 days (14 days prior to mating and throughout the mating and gestation periods until day 4 of lactation)

Non-pregnant females: 53 days

Frequency of treatment

once a day (7 times/week)

Doses / concentrations

Dose / conc.	
0	mg/kg bw/day (actual dose received)
Dose / conc.	
100	mg/kg bw/day (actual dose received)
Dose / conc.	
300	mg/kg bw/day (actual dose received)
Dose / conc.	
1000	mg/kg bw/day (actual dose received)

No. of animals per sex per dose

12 animals/sex/dose

Control animals

yes, concurrent vehicle

Details on study design

The dose levels of this study were selected based on the results of the previously conducted study, "A 14-day oral gavage toxicity study of 2,2',6,6'-tetra- tert-butyl-4,4'-methylenediphenol in rats (dose-finding study)". In that study, there were no toxic changes at 1000 mg/kg, which is the limit test dose prescribed in the OECD Guidelines for Testing of Chemicals 421. Based on these results, the high dose in this reproduction/developmental toxicity screening test was set at 1000 mg/kg, and the lower doses were set at 300 and 100 mg/kg using the common ratio of approximately 3.

- Rationale for animal assignment (if not random): Body weight-balanced randomization
- Post-exposure recovery period in satellite groups: 14 days

Examinations

Parental animals: Observations and examinations

Males and females: 3 times/day during the administration period (before and after dosing)

Observation for completion of delivery (measurement and observation on day 0 of lactation) was conducted twice daily from day 21 of gestation to day 25 of gestation.

Oestrous cyclicity (parental animals)

yes

Sperm parameters (parental animals)

no

Litter observations

no

Postmortem examinations (parental animals)

Necropsy:yes

[Detailed macroscopic examination: including the external appearance, head, thorax and abdomen]

histopathology: yes

[testes, epididymides, ovaries, uterus, prostate, seminal vesicle, vagina, macroscopic lesions]

Postmortem examinations (offspring)

yes

examine for abnormalities in organs/tissues in the external, thoracic and abdominal regions.

Statistics

mean value with standard deviation : [Body weight, body weight gain (from day 1 to day 42 of administration for males, from day 1 to day 15 of administration, from day 0 to day 20 of gestation, and from day 0 to day 4 of lactation for females, and from day 0 to day 4 after birth for liveborn pups), food consumption, number of estruses, estrous cycle, number of elapsed days until copulation, length of gestation period, number of corpora lutea, number of implantation sites, number of pups alive, sex ratio (on day 0 and day 4 after birth) and organ weight (including body weight at necropsy)]

group mean with standard deviation was calculated and subjected to a Dunnett-type mean rank test for the difference in mean rank (levels of significance: 0.05 and 0.01, two-tailed): [stillbirth index, index of external abnormalities, live birth index and viability index on day 4 after birth]

chi-square test with Yates' continuity correction (levels of significance: 0.05 and 0.01, two-tailed): [the copulation index, insemination index, fertility index and delivery index, the number of copulated animals, number of males impregnated females, number of pregnant females and the number of females that delivered liveborn pups were calculated for each group]

If there were cells whose expected frequency was 5 or below, data were analyzed by Fisher's exact test (levels of significance: 0.05 and 0.01, two-tailed)

Reproductive indices

Copulation index (%) = (No. of copulated animals / No. of animals housed together) × 100

Insemination index (%) = (No. of males impregnated females / No. of copulated males) × 100

Fertility index (%) = (No. of pregnant females / No. of copulated females) × 100

Gestation period (days) = Number of days from day 0 of gestation to the day of delivery

Delivery index (%) = (No. of females delivered liveborn pups / No. of pregnant females) × 100

Implantation index (%) = (No. of implantation sites / No. of corpora lutea) × 100

Stillbirth index (%) = (No. of stillborn pups / Total No. of liveborn and stillborn pups) × 100

Index of external abnormalities (%) = (No. of pups with external abnormalities / No. of liveborn pups) × 100

Live birth index (%) = (No. of liveborn pups / No. of implantation sites) × 100

Sex ratio on day 0 after birth = No. of liveborn males / No. of liveborn pups

Sex ratio on day 4 after birth = No. of males alive on day 4 after birth / No. of pups alive on day 4 after birth

Offspring viability indices

Viability index on day 4 after birth (%) = (No. of surviving pups on day 4 after birth / No. of liveborn pups) × 100

Results and discussion

Results: P0 (first parental generation)

General toxicity (P0)

Clinical signs

no effects observed

Mortality

no mortality observed

Body weight and weight changes

no effects observed

Food consumption and compound intake (if feeding study)

no effects observed

Description (incidence and severity)

In females in the 1000 mg/kg group, significantly low values compared to that of the control group were observed on day 15 of administration and on days 1 and 14 of gestation, but since there were no abnormalities in the body weight development in any period, the changes were thought to be of no toxicological significance.

Haematological findings

not examined

Clinical biochemistry findings

not examined

Urinalysis findings

not examined

Behaviour (functional findings)

not examined

Organ weight findings including organ / body weight ratios

no effects observed

Gross pathological findings

no effects observed

Description (incidence and severity)

The following changes were observed in the organs/tissues listed below, but they were thought to be incidental changes based on the incidence of their occurrence and their pathological profiles: small testis and epididymis (unilateral) in 1/12 males in the 1000 mg/kg group, white focus in the epididymis in 1/12 males in the 300 mg/kg group, large testis (unilateral) in 1/12 males in the 100 mg/kg group, and diverticulum in the ileum in 1/12 females in the 300 mg/kg group

Histopathological findings: non-neoplastic

no effects observed

Description (incidence and severity)

The some changes were observed in the organs/tissues, but they were thought to be incidental changes based on the incidence of their occurrence and their histopathological profiles.

Reproductive function / performance (P0)**Reproductive function: oestrous cycle**

no effects observed

Reproductive function: sperm measures

not examined

Reproductive performance

no effects observed

Effect levels (P0)

Key result	false
Dose descriptor	NOAEL
Effect level	1000 mg/kg bw/day (actual dose received)
Based on	test mat.
Sex	male/female
Remarks on result	other: no effects on reproduction

Results: F1 generation

General toxicity (F1)**Clinical signs**

no effects observed

Mortality / viability

mortality observed, non-treatment-related

Description (incidence and severity)

The number of pups that died during the lactation period was 7 in the control group, and 1, 2 and 9 in the 100, 300 and 1000 mg/kg groups, respectively, and there was no significant difference in the viability index on day 4 after birth between the control group and any test article administration group.

Body weight and weight changes

no effects observed

Effect levels (F1)

Key result

false

Dose descriptor

NOAEL

Generation

F1

Effect level

1000

mg/kg bw/day (actual dose received)

Based on

test mat.

Sex

male/female

Remarks on result

other: no effect on development

Any other information on results incl. tables

Figures and Tables (in English) are available in the following full report of the study.

http://dra4.nihs.go.jp/mhlw_data/home/pdf/PDF118-82-1c.pdf

Applicant's summary and conclusion

Conclusions

The NOAEL of 4,4'-methylenebis(2,6-di-tert-butylphenol) for reproductive and developmental toxicity was determined to be 1,000 mg/kg/day (the highest dose tested).

Executive summary

A reproduction/developmental toxicity screening test (OECD TG 421) was conducted to clarify the effects of 4,4'-methylenebis(2,6-di-tert-butylphenol) on reproductive and developmental toxicity. Male

rats were treated with 4,4'-methylenebis(2,6-di-tert-butylphenol) at doses of at 0 [vehicle: 0.5% (w/v) methylcellulose solution], 100, 300, and 1,000 mg/kg bw/day for 14 days prior to mating and throughout the mating period until the day before necropsy (42 days), and female rats were treated for 14 days prior to mating and throughout the mating and gestation periods until day 4 of lactation (41–50 days). There were no mortalities with any dose during the treatment period. There were no effects on reproductive toxicity (fertility and reproductive organs) and developmental toxicity up to the highest dose. The NOAEL of 4,4'-methylenebis(2,6-di-tert-butylphenol) for reproductive and developmental toxicity was determined to be 1,000 mg/kg/day (the highest dose tested).

DOMAIN

SUBSTANCE: 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol / 118-82-1

UUID: c4a3c724-6cba-4624-ae5f-b2bd2dc4cfc9

Dossier UUID:

Author: SuperUser

Date: 2020-10-09T15:01:44.204+09:00

Remarks:

Substance name

2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol / 118-82-1

Legal entity

[National Institute of Health Sciences / Kawasaki / Japan](#)

Identification of substance

Reference substance

[2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol / 4,4'-methylenebis\(2,6-di-tert-butylphenol\) / 118-82-1 / 204-279-1](#)

EC number

204-279-1

EC name

EC Inventory

CAS number

118-82-1

CAS name

IUPAC name

4,4'-methylenebis(2,6-di-tert-butylphenol)

Role in the supply chain

Manufacturer

false

Importer

false

Only representative

false

Downstream user

false

References

REFERENCE_SUBSTANCE: 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol

UUID: ECB5-40091664-5cd2-4b47-95b6-198f2e35e7cc

Dossier UUID:

Author: SuperUser

Date: 2007-05-10T18:00:00.000+09:00

Remarks:

General information

Reference substance name

2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol

Inventory

Inventory name

2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol

Inventory

EC

Inventory number

204-279-1

CAS number

118-82-1

Molecular formula

C₂₉H₄₄O₂

Description

Reference substance information

IUPAC name

4,4'-methylenebis(2,6-di-tert-butylphenol)

Synonyms

Identity
Phenol, 4,4'-methylenebis 2,6-bis(1,1-dimethylethyl)-
Identity
Phenol, 4,4'-methylenebis[2,6-bis(1,1-dimethylethyl)-
Identity
Phenol, 4,4'-methylenebis[2,6-bis(1,1-dimethylethyl)-

CAS information

CAS number
118-82-1

Related substances

Group / category information
DSL Category: Organics
USEPA Category: Neutral Organics;Phenols

Molecular and structural information

Molecular formula
C₂₉H₄₄O₂

Molecular weight

424.6585

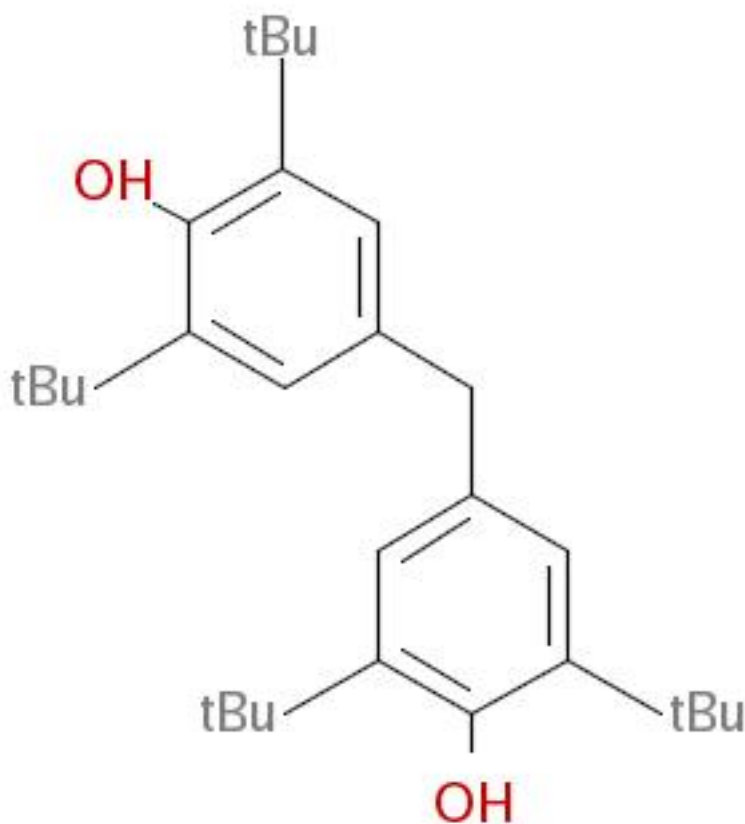
SMILES notation

CC(C)(C)c1cc(Cc2cc(c(O)c2)C(C)(C)C(C)(C)C(C)(C)C)cc(c1O)C(C)(C)C

InChI

InChI=1/C₂₉H₄₄O₂/c1-26(2,3)20-14-18(15-21(24(20)30)27(4,5)6)13-19-16-22(28(7,8)9)25(31)23(17-19)29(10,11)12/h14-17,30-31H,13H2,1-12H3

Structural formula



LITERATURE: A 28-day repeat dose oral toxicity study of 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol in rat

UUID: f37fd311-479b-4fca-b7b9-0ac8548efa1e

Dossier UUID:

Author: SuperUser

Date: 2019-05-22T11:25:03.000+09:00

Remarks:

General information

Reference Type

publication

Title

A 28-day repeat dose oral toxicity study of 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol in rat

Author

Ministry of Health, Labour and Welfare (MHLW), Japan

Year

2010

LITERATURE: A reproduction/developmental toxicity screening test in rats treated orally with 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol

UUID: 94cbeb6e-5aa1-4b9f-b4d8-92b4f7e5e123

Dossier UUID:

Author: SuperUser

Date: 2019-03-27T06:49:02.000+09:00

Remarks:

General information

Reference Type

publication

Title

A reproduction/developmental toxicity screening test in rats treated orally with 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol

Author

Ministry of Health, Labour and Welfare (MHLW), Japan

Year

2012

LITERATURE: A reverse mutation test of 2,2',6,6'-tetra-tert-butyl-4,4'-methylenedi phenol using bacteria

UUID: 7620b1e7-84d8-4e14-b39e-a28aca544fbe

Dossier UUID:

Author: SuperUser

Date: 2019-03-27T05:44:24.000+09:00

Remarks:

General information

Reference Type

publication

Title

A reverse mutation test of 2,2',6,6'-tetra-tert-butyl-4,4'-methylenedi phenol using bacteria

Author

Ministry of Health, Labour and Welfare(MHLW), Japan

Year

2010

LITERATURE: Chromosome aberration test in cultured chinese hamster cells treated with 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol

UUID: f94b33a7-1c0b-4d80-a883-4d98ff599dd4

Dossier UUID:

Author: SuperUser

Date: 2019-05-22T11:37:43.000+09:00

Remarks:

General information

Reference Type

publication

Title

Chromosome aberration test in cultured chinese hamster cells treated with 2,2',6,6'-tetra-tert-butyl-4,4'-methylenediphenol

Author

Ministry of Health, Labour and Welfare(MHLW), Japan

Year

2010

LEGAL_ENTITY: National Institute of Health Sciences

UUID: IUC4-b036ff75-0f3c-323b-b200-ed5f46cf5101

Dossier UUID:

Author: SuperUser

Date: 2019-09-03T10:05:28.000+09:00

Remarks: Disclaimer: The contents in this document were created based on the MHLW (Ministry of Health, Labour and Welfare) peer reviewed study reports (in Japanese) in JECDB (Japan Existing Chemical Database) at http://dra4.nihs.go.jp/mhlw_data/jsp/SearchPageENG.jsp. Authorship is in the Division of Risk Assessment, the National Institute of Health Sciences, and the contents do not reflect any official MHLW opinions or any other regulatory policies.

General information

Legal entity name

National Institute of Health Sciences

Remarks

Disclaimer: The contents in this document were created based on the MHLW (Ministry of Health, Labour and Welfare) peer reviewed study reports (in Japanese) in JECDB (Japan Existing Chemical Database) at http://dra4.nihs.go.jp/mhlw_data/jsp/SearchPageENG.jsp. Authorship is in the Division of Risk Assessment, the National Institute of Health Sciences, and the contents do not reflect any official MHLW opinions or any other regulatory policies.

Identifiers

Other IT system identifiers

IT system LEO
ID 10767
IT system IUCLID4
ID 16558402024DIV750

Contact information

Contact persons

Person Hirose, Akihiko; National Institute of Health Sciences, Japan
Last name Hirose

First name

Akihiko

Organisation

National Institute of Health Sciences, Japan

Department

Division of Risk Assessment

Title

Dr

Country

Japan

Contact address**Address 1**

Tonomachi 3-25-26

Address 2

Kawasaki-ku

Postal code

210-9501

Town

Kawasaki

Region / State

Kanagawa

Country

Japan