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**Name:** 4-Chlorobenzaldehyde / 4-chlorobenzaldehyde / 104-88-1

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

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**Printing date:** 2019-09-03T11:18:16.017+09:00

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# 4-Chlorobenzaldehyde

## CORE

### General information

#### Identification

**SUBSTANCE:** 4-Chlorobenzaldehyde

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**UUID:** IUC5-88396134-556c-42ba-b624-833db483f5ce

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2016-12-21T15:12:44.000+09:00

**Remarks:**

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#### Substance name

4-Chlorobenzaldehyde

#### Legal entity

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

### Identification of substance

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#### Reference substance

[4-chlorobenzaldehyde / 4-chlorobenzaldehyde / 104-88-1 / 203-247-4](#)

**EC number**

203-247-4

**EC name**

EC Inventory

**CAS number**

104-88-1

**CAS name**

**IUPAC name**

4-chlorobenzaldehyde

### Role in the supply chain

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#### Manufacturer

false

#### Importer

false

#### Only representative

false

#### Downstream user

false

# OECD

## Health Effects

Repeated dose toxicity: oral

ENDPOINT\_STUDY\_RECORD: Repeated dose toxicity: oral.001

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**UUID:** IUC5-5c3897a9-5e38-496d-977f-60a8fb98a9cb

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2017-01-17T11:42:25.000+09:00

**Remarks:**

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## Administrative data

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**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**

other: The study was conducted in accordance with Test Guidelines and under GLP

**Cross-reference**

**Reason / purpose**

reference to other study

**Remarks**

7.5.1 Repeated dose toxicity: oral.002

## Data source

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**Reference**

[Twenty-eight-day Repeat Dose Oral Toxicity Test of 4-Chlorobenzaldehyde in Rats / MHLW / publication](#)

**Data access**

data published

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## Materials and methods

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### Test guideline

**Qualifier**

according to

**Guideline**

other: other guideline: Guideline for 28-Day Repeated Dose Toxicity Test in Mammalian Species (Chemical Substances Control Law of Japan)

**Qualifier**

equivalent or similar to

**Guideline**

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

### GLP compliance

yes

## Test material

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### Test material information

[4-chlorobenzaldehyde](#) / [104-88-1](#) / [203-247-4](#)

## Test animals

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**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 Japan, Inc.
- Age at study initiation: 5 weeks old
- Weight at study initiation: male (150-177 g), female (114-131 g)
- Housing: Animals were individually housed in a metallic cage with wire mesh bottoms
- Diet: Solid feed (CRF-1: Oriental Yeast Co., Ltd.) was given ad libitum.
- Water: Tap water was given ad libitum.
- Acclimation and quarantine period: 6 days

**ENVIRONMENTAL CONDITIONS**

- Temperature (°C): 22±3 (actual temperature: 21-25 °C)
- Humidity (%): 50±20% (actual humidity: 39-56%)
- Air changes (per hr): 10-15
- Photoperiod (hrs dark / hrs light): 12 hr dark/12 hr light (light: 8:00~20:00)

## Administration / exposure

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**Route of administration**

oral: gavage

**Vehicle**

corn oil

**Analytical verification of doses or concentrations**

yes

**Duration of treatment / exposure**

28 days

**Frequency of treatment**

once a day

**Doses / concentrations**

**Remarks**

Doses / Concentrations:

0, 8, 40, 200, 1000 mg/kg bw/day

Basis:

actual ingested

**No. of animals per sex per dose**

12/sex (0, 1000 mg/kg bw/day)

6/sex (8, 40, 200 mg/kg bw/day)

**Control animals**

yes, concurrent vehicle

**Details on study design**

- Dose selection rationale:

The dosage levels were determined based on the finding in a 14-day dose-finding study.

In a dose finding study for a 28 day study, CrI:CD(SD) rats were given 4-chlorobenzaldehyde at 0 (corn oil), 8, 40, 200 or 1000 mg/kg/day for 14 days. At 40 mg/kg/day and higher, the urine specific gravity in males was showed a decreasing trend, at 200 mg/kg/day and higher, urine protein in males and females was decreased or showed a decreasing trend, and at 1000 mg/kg/day, an increase in urine volume in males and females, and increases in AST, ALT and ALP in females were observed.

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

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

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## Examinations

**Observations and examinations performed and frequency**

CLINICAL OBSERVATIONS: Yes

- Time schedule: every day during the administration (twice a day: am and pm) and recovery periods (twice a day: am and pm).

DETAILED CLINICAL OBSERVATIONS: Yes

The functional observational battery testing (FOB) was performed on all animals. Among the measures in the FOB, detailed clinical observations were made before the initiation of dosing. Thereafter, detailed clinical observations were made once a week in dosing and recovery periods.

Sensory motor reflexes, forelimb and hindlimb grip strengths, and motor activity were measured on week 4 of administration period (main/recovery group animals) and week 2 of recovery period (recovery group animals).

BODY WEIGHT: Yes

- Time schedule for examinations: Before administration (on days 1, 7, 14, 21 and 28 of the administration period, days 7 and 14 of the recovery period) and the necropsy days after completion of recovery period.

FOOD CONSUMPTION AND COMPOUND INTAKE (if feeding study):

- Food consumption: Yes. Before administration (on days 1, 7, 14, 21 and 28 of the administration period and days 7 and 14 of the recovery period)

OPHTHALMOSCOPIC EXAMINATION: No

HAEMATOLOGY: Yes

- Time schedule for collection of blood: the after completion of the administration and recovery periods

- Anaesthetic used for blood collection: ether

- Animals fasted: Yes (16-22 hours)

- How many animals: all animals

CLINICAL CHEMISTRY: Yes

- Time schedule for collection of blood: the day after completion of the administration and recovery periods

- Animals fasted: Yes

- How many animals: all animals

URINALYSIS: Yes

- Time schedule for collection of urine: on weeks 4 of the administration period and weeks 2 of the recovery period.

- Metabolism cages used for collection of urine: Yes

- Animals fasted: No

- How many animals: all animals

NEUROBEHAVIOURAL EXAMINATION: No

### **Sacrifice and pathology**

GROSS PATHOLOGY: Yes

ORGAN WEIBHT: Yes [brain, pituitary gland, thyroid, adrenal, spleen, heart, liver, kidney, thymus, testis, epididymis, prostate, seminal vesicles (including coagulation gland), ovary, uterus]

HISTOPATHOLOGY: Yes [brain (cerebrum, cerebellum and medulla oblongata), pituitary gland, spinal cord, thymus, thyroid, parathyroid, adrenal glands, spleen, heart, tongue, esophagus, stomach, liver, pancreas, duodenum, jejunum, ileum (including Peyer's patches), cecum, colon, rectal, mesenteric lymph nodes, submandibular lymph nodes, trachea, lung, kidney, bladder, testis, epididymis, prostate, seminal vesicles (including coagulation glands), ovary, uterus, vagina, eye, Harder gland, femur (including bone marrow, right) and the sciatic nerve. (see tables in the study report.)

### **Statistics**

The homoscedasticity was analyzed by Bartlett's test for data of grip strength, motor activity, body weight, body weight gain, food consumption, quantitative items of urinary findings (except for the urine specific gravity), hematological test, biochemical test, organ weight and organ weight ratio. When homogeneity was recognized, one-way analysis of variance (homogeneous data) or Kruskal-Wallis (non-homogeneous data) was conducted. If a significant difference was detected, as the result of one-way analysis of variance, Dunnett's method was applied for comparisons between control and individual treatment groups. And in the case of a significant difference was detected on Kruskal-Wallis, Mann-Whitney's U-test was applied for the same purpose. The trend by the group was analyzed by Kruskal-Wallis for general appearance, detailed clinical observation, qualitative items of urinary findings, and urine specific gravity. If a significant difference was detected as the result of Kruskal-Wallis, Mann-Whitney's U-test as applied for comparisons between control and individual treatment groups.

## **Results and discussion**

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### **Results of examinations**

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#### **Clinical signs**

effects observed, treatment-related

#### **Description (incidence and severity)**

(see Details on results)

#### **Mortality**

mortality observed, treatment-related

**Description (incidence)**

(see Details on results)

**Body weight and weight changes**

effects observed, treatment-related

**Description (incidence and severity)**

(see Details on results)

**Food consumption and compound intake (if feeding study)**

effects observed, treatment-related

**Description (incidence and severity)**

(see Details on results)

**Haematological findings**

effects observed, treatment-related

**Description (incidence and severity)**

(see Details on results)

**Clinical biochemistry findings**

effects observed, treatment-related

**Description (incidence and severity)**

(see Details on results)

**Urinalysis findings**

effects observed, treatment-related

**Description (incidence and severity)**

(see Details on results)

**Behaviour (functional findings)**

effects observed, treatment-related

**Description (incidence and severity)**

(see Details on results)

**Organ weight findings including organ / body weight ratios**

effects observed, treatment-related

**Description (incidence and severity)**

(see Details on results)

**Gross pathological findings**

effects observed, treatment-related

**Description (incidence and severity)**

(see Details on results)

**Histopathological findings: non-neoplastic**

effects observed, treatment-related

**Description (incidence and severity)**

(see Details on results)

**Details on results**

**CLINICAL SIGNS AND MORTALITY**

Transient salivation and tremors were observed in both sexes at 1,000 mg/kg bw/day. Soiled fur was observed in females at 1,000 mg/kg bw/day.

**BODY WEIGHT AND WEIGHT GAIN**



At 1,000 mg/kg bw/day, a decrease in body weight gain was observed in males.

**FOOD CONSUMPTION**

At 1,000 mg/kg bw/day, an increase in food consumption was observed in females.

**HAEMATOLOGY**

At 1,000 mg/kg bw/day, a decrease in platelet was observed in males.

**CLINICAL CHEMISTRY**

At 1,000 mg/kg bw/day, increases in serum aspartate aminotransferase and alanine aminotransferase levels were observed in both sexes, whereas decreases in total protein and  $\beta$ -globulin fraction were observed in males, and increases in alkaline phosphatase and triglyceride levels were observed in females.

**URINALYSIS**

At 1,000 mg/kg bw/day, an increase in urine volume in males and females and decreases in urine pH and protein and specific gravity levels in males were observed.

**NEUROBEHAVIOUR**

At 1,000 mg/kg bw/day, a decrease in grip strength of forearms was observed in males, and a decrease in the locomotor activity was observed in females.

**ORGAN WEIGHTS**

Increased liver weight and decreased ovary and heart weights were observed at 1,000 mg/kg bw/day in females.

**GROSS PATHOLOGY**

At 1,000 mg/kg bw/day, atrophy of lateral vastus muscle in one male and multifocal mucosal black patch in glandular stomach in one female were observed.

**HISTOPATHOLOGY: NON-NEOPLASTIC**

Histopathological examinations revealed hyperostosis metaphysis of the femur in females at 200 mg/kg bw/day. Furthermore, at 1,000 mg/kg bw/day, hyperkeratosis of the mucosal epithelium in the forestomach, grade enhancement of regeneration of the tubular epithelium, dilatation of the renal tubules in the cortex, and cyst-like extension of the collecting duct in the kidney, and hyperostosis metaphysis of the femur were observed in both sexes. Additionally in males at 1,000 mg/kg bw/day, degeneration of nerve fibers in the sciatic nerve and atrophy of muscle fibers in the skeletal muscle were observed. And in females at 1,000 mg/kg bw/day, squamous cell hyperplasia of the boundary edge in the stomach was observed. These changes tended to resolve after the recovery period.

**Effect levels**

<b>Dose descriptor</b>	
NOAEL	
<b>Effect level</b>	
200	mg/kg bw/day (actual dose received)
<b>Based on test mat.</b>	
<b>Sex</b>	
male	
<b>Basis for effect level</b>	
other: see 'Remark'	

At 1,000 mg/kg bw/day in males: Transient salivation and tremors were observed, and decreases in body weight gain and grip strength of forearms were observed, increases in serum aspartate aminotransferase and alanine aminotransferase levels, decreases in total protein and  $\beta$ -globulin fraction were observed, and hyperkeratosis of the mucosal epithelium in the forestomach, grade enhancement of regeneration of the tubular epithelium, dilatation of the renal tubules in the cortex, and cyst-like extension of the collecting duct in the kidney, and hyperostosis metaphysis of the femur were observed. Additionally, degeneration of nerve fibers in the sciatic nerve and atrophy of muscle fibers in the skeletal muscle were observed.

**Dose descriptor**

NOAEL

**Effect level**

40

mg/kg bw/day (actual dose received)

**Based on**  
test mat.

**Sex**  
female

**Basis for effect level**

other: Histopathological examinations revealed hyperostosis metaphysis of the femur in females at 200 mg/kg bw/day.

**Target system / organ toxicity**

**Key result**  
false

**Critical effects observed**  
not specified

**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/PDF104-88-1b.pdf](http://dra4.nihs.go.jp/mhlw_data/home/pdf/PDF104-88-1b.pdf)

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

In a 28-day repeated-dose toxicity test performed according to OECD TG 407, male and female rats (6 animals/sex/dose) were administered 4-chlorobenzaldehyde at 0, 8, 40, 200, and 1,000 mg/kg bw/day. In addition, both sexes (6 animals/sex/dose) were administered 0 and 1,000 mg/kg bw/day of this substance for 28 days and examined after a 14-day recovery period. At 1,000 mg/kg bw/day, transient salivation and tremors were observed in both sexes, decreases in body weight gain and grip strength of forearms were observed in males, and a decrease in the locomotor activity was observed in females. At this dose, increases in serum aspartate aminotransferase and alanine aminotransferase levels were observed in both sexes, whereas decreases in total protein and  $\beta$ -globulin fraction were observed in males, and increases in alkaline phosphatase and triglyceride levels were observed in

females. Increased liver weight and decreased ovary weight were also observed at 1,000 mg/kg bw/day in females. Histopathological examinations revealed hyperostosis metaphysis of the femur in females at 200 mg/kg bw/day. Furthermore, at 1,000 mg/kg bw/day, hyperkeratosis of the mucosal epithelium in the forestomach, grade enhancement of regeneration of the tubular epithelium, dilatation of the renal tubules in the cortex, and cyst-like extension of the collecting duct in the kidney, and hyperostosis metaphysis of the femur were observed in both sexes. Additionally in males at 1,000 mg/kg bw/day, degeneration of nerve fibers in the sciatic nerve and atrophy of muscle fibers in the skeletal muscle were observed. And in females at 1,000 mg/kg bw/day, squamous cell hyperplasia of the boundary edge in the stomach was observed. These changes tended to resolve after the recovery period. On the basis of these effects, NOAELs for repeated-dose toxicity were determined to be 200 mg/kg bw/day and 40 mg/kg bw/day in male and female rats, respectively.

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**ENDPOINT\_STUDY\_RECORD: Repeated dose toxicity: oral.002**

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**UUID:** IUC5-520d8d8b-7f48-4e05-b483-817f6ebe17ce

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2017-01-17T11:45:20.000+09:00

**Remarks:**

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## Administrative data

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**Endpoint**

repeated dose toxicity: oral combined repeated dose and reproduction / developmental screening  
deactivated phrase

**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**

other: GLP guideline study

**Cross-reference**

**Reason / purpose**

reference to same study

**Remarks**

7.8.1 Toxicity to reproduction.001

**Reason / purpose**

reference to other study

**Remarks**

7.5.1 Repeated dose toxicity: oral.001

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## Data source

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**Reference**

[A reproduction/developmental toxicity screening test in rats treated orally with 4-chlorobenzaldehyd... / MHLW Japan / study report](#)

**Data access**

data published

## Materials and methods

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### Test guideline

**Qualifier**

according to

**Guideline**

other: OECD TG 421: Reproduction/developmental toxicity screening test

**Deviations**

no

**GLP compliance**

yes

**Limit test**

no

## Test material

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**Test material information**

[4-chlorobenzaldehyde / 104-88-1 / 203-247-4](#)

## Test animals

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**Species**

rat

common rodent species

**Strain**

other: Crl: CD(SD)

**Sex**

male/female

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

- Source: Charles River Laboratories Japan, Inc. Tsukuba
- Age at study initiation: 10 weeks
- Weight at study initiation: Males: 389-449 g; Females: 234-271 g
- Housing: Steel wire-mesh cage (250 mm x 350 mm x 200 mm )
- Diet: ad libitum
- Water: ad libitum
- Acclimation period: 14 days

**ENVIRONMENTAL CONDITIONS**

- Temperature (°C): 20-25
- Humidity (%): 41-69
- Air changes: 10-15 times / hr
- Photoperiod: 12 hrs dark / 12 hrs light (07:00-19:00)

## Administration / exposure

---

**Route of administration**

oral: gavage

**Vehicle**

corn oil

**Analytical verification of doses or concentrations**

yes

**Duration of treatment / exposure**

(P) Males: 42 days including 14 days pre-mating and mating periods, and thereafter 14 days (P)

Females: 42-45 days including 14 days pre-mating, mating and gestation periods, and the days until day 4 of lactation

**Frequency of treatment**

Once/day, 7days/week

**Doses / concentrations**

**Remarks**

Doses / Concentrations:

0 (vehicle), 40, 200, and 1000 mg/kg bw/day

Basis:

actual ingested

**No. of animals per sex per dose**

12 animals/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 study: 28-day repeated dose oral toxicity test (doses: 0, 8, 40, 200, and 1000 mg/kg bw/day). At 1,000 mg/kg bw/day, transient salivation and tremors were observed in both sexes, decreases in body weight gain and grip strength of forearms were observed in males, and a decrease in the locomotor activity was observed in females. At this dose, increases in serum aspartate aminotransferase and alanine aminotransferase levels were observed in both sexes, whereas decreases in total protein and  $\beta$ -globulin fraction were observed in males, and increases in alkaline phosphatase and triglyceride levels were observed in females. Increased liver weight and decreased ovary weight were also observed at 1,000 mg/kg bw/day in females. Histopathological examinations revealed hyperostosis metaphysis of the femur in females at 200 mg/kg bw/day. Furthermore, at 1,000 mg/kg bw/day, hyperkeratosis of the mucosal epithelium in the forestomach, grade enhancement of regeneration of the tubular epithelium, dilatation of the renal tubules in the cortex, and cyst-like extension of the collecting duct in the kidney, and hyperostosis metaphysis of the femur were observed in both sexes. Additionally in males at 1,000 mg/kg bw/day, degeneration of nerve fibers in the sciatic nerve and atrophy of muscle fibers in the skeletal muscle were observed. And in females at 1,000 mg/kg bw/day, squamous cell hyperplasia of the boundary edge in the stomach was observed. These changes tended to resolve after the recovery period.

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

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**Examinations**

**Observations and examinations performed and frequency**

CAGE SIDE OBSERVATIONS: Yes

- Time schedule:

Males and females: 3 times/day

BODY WEIGHT: Yes

- Time schedule for examinations:

Males: Days 1, 4, 8, 11, 15, 22, 25, 29, 32, 36, 39, 42, and the day of necropsy

Females: Twice a week during the precopulation period (days 1, 4, 8, 11, and 15); gestation days 0, 4, 7, 11, 14, 17, and 20; lactation days 0 and 4; and the day of necropsy

FOOD CONSUMPTION: Yes

Males: Days 1, 4, 8, 11, 15, 32, 36, 39, and 42 in dosing period

Females: Days 1, 4, 8, 11, and 15; gestation days 1, 4, 7, 11, 14, 17, and 20; lactation days 2 and 4

HAEMATOLOGY: No

CLINICAL CHEMISTRY: No

URINALYSIS: No

#### **Sacrifice and pathology**

GROSS PATHOLOGY: Yes (see tables)

HISTOPATHOLOGY: Yes (epididymis, prostate, seminal vesicle, testis, ovary, uterus, vagina, and gross abnormal sites)

#### **Other examinations**

Organ weight: Testes and epididymides

#### **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 ( $p < 0.05$ , two-sided).

## **Results and discussion**

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### **Results of examinations**

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#### **Clinical signs**

effects observed, treatment-related

#### **Description (incidence and severity)**

Nine males and seven females died with tremors and decreased locomotor activity at 1,000 mg/kg bw/day by day 9 of administration.

#### **Mortality**

mortality observed, treatment-related

#### **Description (incidence)**

Nine males and seven females died with tremors and decreased locomotor activity at 1,000 mg/kg bw/day by day 9 of administration.

#### **Body weight and weight changes**

effects observed, treatment-related

#### **Description (incidence and severity)**

Low value of body weight was observed in both sexes at 1,000 mg/kg bw/day.

#### **Food consumption and compound intake (if feeding study)**

effects observed, treatment-related

#### **Description (incidence and severity)**

Low value of food consumption was observed in both sexes at 1,000 mg/kg bw/day.

**Food efficiency**

not examined

**Water consumption and compound intake (if drinking water study)**

not examined

**Ophthalmological findings**

not examined

**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**

effects observed, treatment-related

**Description (incidence and severity)**

see tables.

**Histopathological findings: non-neoplastic**

effects observed, treatment-related

**Description (incidence and severity)**

see tables.

**Histopathological findings: neoplastic**

not examined

**Effect levels**

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**Dose descriptor**

NOAEL

**Effect level**

200

mg/kg bw/day (actual dose received)

**Based on**

test mat.

**Sex**

male/female

**Basis for effect level**

other: Nine males and seven females died with tremors and decreased locomotor activity at 1,000 mg/kg bw/day by day 9 of administration.

**Target system / organ toxicity**

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**Key result**

false

**Critical effects observed**

not specified

**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/PDF104-88-1c.pdf](http://dra4.nihs.go.jp/mhlw_data/home/pdf/PDF104-88-1c.pdf)

**Applicant's summary and conclusion** \_\_\_\_\_

**Conclusions**

In this study, NOAEL for repeated-dose toxicity was determined to be 200 mg/kg bw/day in male and female rats.

**Executive summary**

A reproduction/developmental toxicity screening test was performed according to OECD TG 421. Male and female rats (12 animals/sex/dose) were administered 4-chlorobenzaldehyde at 0, 40, 200, and 1,000 mg/kg bw/day. Males were dosed for 42 days, including a 14 day pre-mating and mating periods. Females were dosed for 42–45 days, including a 14-day pre-mating, mating, and gestation periods and the time until day 4 of lactation. Nine males and seven females died with tremors and decreased locomotor activity at 1,000 mg/kg bw/day by day 9 of administration. In this study, NOAEL for repeated-dose toxicity was determined to be 200 mg/kg bw/day in male and female rats.

## Genetic toxicity in vitro

### ENDPOINT\_STUDY\_RECORD: Genetic toxicity in vitro.001

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**UUID:** IUC5-6657ee02-5cbb-4ca8-9b60-b74d46361195

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2019-09-03T11:17:12.578+09:00

**Remarks:**

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## Administrative data

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### Endpoint

in vitro gene mutation study in bacteria Type of genotoxicity: gene mutation

### 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

other: OECD Test Guideline study under GLP condition

## Data source

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### Reference

[Reverse Mutation Test of 4-chlorobenzaldehyde on Bacteria. / MHLW, Japan / study report](#)

### Data access

data published

## Materials and methods

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### Test guideline

**Qualifier**

according to

**Guideline**

JAPAN: Guidelines for Screening Mutagenicity Testing Of Chemicals

**Qualifier**

according to

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**Guideline**

OECD Guideline 471 (Bacterial Reverse Mutation Assay)  
in vitro gene mutation study in bacteria

**GLP compliance**

yes

**Type of assay**

bacterial reverse mutation assay  
in vitro gene mutation study in bacteria

**Test material**

**Test material information**

[4-chlorobenzaldehyde / 104-88-1 / 203-247-4](#)

**Method**

**Species / strain**

**Species / strain**

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

**Metabolic activation**

with and without

**Metabolic activation system**

rat liver, induced by phenobarbital and 5,6-benzoflavone

**Species / strain**

E. coli WP2 uvr A  
bacteria

**Metabolic activation**

with and without

**Metabolic activation system**

rat liver, induced by phenobarbital and 5,6-benzoflavone

**Test concentrations with justification for top dose**

-S9 mix and + S9 mix: 15.6, 31.3, 62.5, 125, 250, 500 µg/plate (all strains)

**Vehicle**

- Vehicle(s)/solvent(s) used: DMSO

**Controls**

**Negative controls**

no

**Solvent controls**

yes

**True negative controls**

other: tests without all strains, and with vehicle, S9 mix or the highest dose

**Positive controls**

yes

**Positive control substance**

other: -S9 mix: 2-(2-Furyl)-3-(5-nitro-2-furyl) acrylamide (AF2:TA 100, TA98 & WP2 uvrA), sodium azide (SA:TA1535) and 9-aminoacridine hydrochloride (9AA:TA1537). +S9 mix: 2-aminoanthracene (2AA:all strains).

**Details on test system and conditions**

RANGE-FINDING/SCREENING STUDIES:

Concentration: 20-5000 µg/plate

Cytotoxic conc.: Yes, >500 µg/plate

Precipitate: No.

METHOD OF APPLICATION: Preincubation

DURATION

- Preincubation period: 20 min at 37 °C

- Exposure duration:48-49 hrs

NUMBER OF PLATES: 3

NUMBER OF REPLICATIONS: 2

DETERMINATION OF CYTOTOXICITY

- Method: other: growth inhibition

**Evaluation criteria**

In any strain(s) tested with or without S9 mix, when the mean number of revertant colonies per plate increased twice more than that of the negative control and when the increase was shown to be dose-related and reproducible, the chemical was judged mutagenic.

**Statistics**

No.

## Results and discussion

**Test results**

**Key result**

false

**Species / strain**

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

**Metabolic activation**

with and without

**Genotoxicity**

negative

**Cytotoxicity**

yes -S9 mix: 500 µg/plate, +S9 mix: >250 µg/plate

**Vehicle controls valid**

yes

**Negative controls valid**

yes

**Positive controls valid**

yes

**Remarks on result**

other: all strains/cell types tested Migrated from field 'Test system'.

**Key result**

false

**Species / strain**

E. coli WP2 uvr A  
bacteria

**Metabolic activation**

with and without

**Genotoxicity**

negative

**Cytotoxicity**

yes -S9 mix: 500 µg/plate, +S9 mix: >250 µg/plate

**Vehicle controls valid**

yes

**Negative controls valid**

yes

**Positive controls valid**

yes

**Remarks on result**

other: all strains/cell types tested Migrated from field 'Test system'.

**Additional information on results**

Contamination with any other bacterias was not found.

**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/PDF104-88-1e.pdf](http://dra4.nihs.go.jp/mhlw_data/home/pdf/PDF104-88-1e.pdf)

**Applicant's summary and conclusion**

**Conclusions**

Interpretation of results (migrated information):  
negative

**Executive summary**

In a bacterial reverse mutation assay using *S. typhimurium* TA100, TA1535, TA98, and TA1537 and *E. coli* WP2uvrA (OECD TG 471), 4-chlorobenzaldehyde was negative with or without metabolic activation.

## ENDPOINT\_STUDY\_RECORD: Genetic toxicity in vitro.002

---

**UUID:** IUC5-df6301ac-67fc-4017-b622-6ee93cc170fe

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2019-09-03T11:17:36.408+09:00

**Remarks:**

---

## Administrative data

---

### Endpoint

in vitro cytogenicity / chromosome aberration study in mammalian cells Type of genotoxicity: chromosome aberration

### 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

other: OECD Test Guideline study under GLP condition

## Data source

---

### Reference

[In Vitro Chromosomal Aberration Test of 4-chlorobenzaldehyde on Cultured Chinese Hamster Cells. / MHLW, Japan / study report](#)

### Data access

data published

## Materials and methods

---

### Test guideline

#### Qualifier

according to

#### Guideline

OECD Guideline 473 (In Vitro Mammalian Chromosome Aberration Test)  
in vitro cytogenicity / chromosome aberration study in mammalian cells

---

**Deviations**

no

**Qualifier**

according to

**Guideline**

JAPAN: Guidelines for Screening Mutagenicity Testing Of Chemicals

**Deviations**

no

**GLP compliance**

yes

**Type of assay**in vitro mammalian chromosome aberration test  
chromosome aberration**Test material****Test material information**[4-chlorobenzaldehyde / 104-88-1 / 203-247-4](#)**Method****Target gene**

Chromosome

**Species / strain****Species / strain**

other: Chinese hamster lung(CHL/IU) cells

**Metabolic activation**

with and without

**Metabolic activation system**

rat liver, induced by phenobarbital and 5,6-benzoflavone

**Test concentrations with justification for top dose**

-S9 mix (short-term treatment): 0, 21.9, 43.8, 87.5, 175, 263, 350 ug/mL

-S9 mix (short-term treatment, confirmatory test 1): 0, 43.8, 87.5, 175, 219, 263, 307, 350 ug/mL

-S9 mix (short-term treatment, confirmatory test 2): 0, 87.5, 175, 219, 263, 307, 350 ug/mL

+S9 mix (short-term treatment): 0, 87.5, 175, 350, 700, 1050, 1400 ug/mL

+S9 mix (short-term treatment, confirmatory test 1): 0, 87.5, 175, 350, 467, 583, 700 ug/mL

-S9 mix (continuous treatment, 24 h): 0, 5.47, 10.9, 21.9, 43.8, 87.5, 131, 175 ug/mL

-S9 mix (continuous treatment, 24 h, confirmatory test 1): 0, 21.9, 43.8, 87.5, 109, 131 ug/mL

-S9 mix (continuous treatment, 24 h, confirmatory test 2): 0, 21.9, 43.8, 65.7, 87.5, 109, 131 ug/mL

**Vehicle**

- Vehicle(s)/solvent(s) used: DMSO

**Controls****Negative controls**

no

**Solvent controls**

yes

**True negative controls**

no

**Positive controls**

yes

**Positive control substance**

benzo(a)pyrene  
cyclophosphamide  
mitomycin C

**Remarks**

mitomycin C (without S9 mix), benzo[a]pyrene or 3,4-benzopyrene (with S9 mix)

**Details on test system and conditions**

METHOD OF APPLICATION: Exposure duration: [continuous treatment]: 24 hrs [short-term treatment]: 6 hrs + 18 hr

SPINDLE INHIBITOR: Colcemid

NUMBER OF REPLICATIONS: 2-3

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**

not used.

## Results and discussion

**Test results**

**Key result**

false

**Species / strain**

other: Chinese hamster lung (CHL/IU) cells

**Metabolic activation**

with

**Genotoxicity**

positive D20: 0.65 mg/mL (main test), 0.57 mg/mL (confirmatory test)

**Cytotoxicity**

yes >50% cell growth inhibition: >700 ug/mL (short, main), >700 ug/mL (short, confirmatory test)

**Vehicle controls valid**

yes



**Negative controls valid**

not examined

**Positive controls valid**

yes

**Key result**

false

**Species / strain**

other: Chinese hamster lung (CHL/IU) cells

**Metabolic activation**

without

**Genotoxicity**

positive D20: 0.24 mg/mL (main test), 0.26 mg/mL (confirmatory-2)

**Cytotoxicity**

yes >50% cell growth inhibition: >350 ug/mL (short, main), >131 ug/mL (24h, main), >263 ug/mL (short, confirmatory-1), >131 ug/mL (24h, confirmatory-1), >307 ug/mL (short, confirmatory-2), >109 ug/mL (24h, confirmatory-2)

**Vehicle controls valid**

yes

**Negative controls valid**

not examined

**Positive controls valid**

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/PDF104-88-1f.pdf](http://dra4.nihs.go.jp/mhlw_data/home/pdf/PDF104-88-1f.pdf)

**Applicant's summary and conclusion** \_\_\_\_\_

**Executive summary**

An in vitro chromosomal aberration test using CHL/IU cells (OECD TG 473) showed positive.

## Toxicity to reproduction

**ENDPOINT\_STUDY\_RECORD: Toxicity to reproduction.001**

---

**UUID:** IUC5-24c0bbaa-17a7-4f00-b6e7-24491d5a34b4

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2017-01-17T12:13:20.000+09:00

**Remarks:**

---

## Administrative data

---

### Endpoint

screening for reproductive / developmental toxicity based on test type (migrated information)

### 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

other: OECD Test Guideline study under GLP condition

### Cross-reference

#### Reason / purpose

reference to same study

#### Remarks

7.5.1 Repeated dose toxicity: oral.002

## Data source

---

### Reference

[A reproduction/developmental toxicity screening test in rats treated orally with 4-chlorobenzaldehyd... / MHLW, Japan / study report](#)

### Data access

data published

---

## Materials and methods

---

### Test guideline

**Qualifier**

according to

**Guideline**

other: OECD TG 421: Reproduction/developmental toxicity screening test

**Deviations**

no

**GLP compliance**

yes

**Limit test**

no

## Test material

---

**Test material information**

[4-chlorobenzaldehyde / 104-88-1 / 203-247-4](#)

## Test animals

---

**Species**

rat

**Strain**

other: CrI:CD(SD)

**Sex**

male/female

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

- Source: Charles River Laboratories Japan, Inc. Tsukuba
- Age at study initiation: 10 weeks
- Weight at study initiation: Males: 389-449 g; Females: 234-271 g
- Housing: Steel wire-mesh cage (250 mm x 350 mm x 200 mm )
- Diet: ad libitum
- Water: ad libitum
- Acclimation period: 14 days

**ENVIRONMENTAL CONDITIONS**

- Temperature (°C): 20-25
- Humidity (%): 41-69
- Air changes: 10-15 times / hr
- Photoperiod: 12 hrs dark / 12 hrs light (07:00-19:00)

## Administration / exposure

---

**Route of administration**

oral: gavage

**Vehicle**

corn oil

**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

**Duration of treatment / exposure**

(P) Males: 42 days including 14 days pre-mating and mating periods, and thereafter 14 days (P)

Females: 42-45 days including 14 days pre-mating, mating and gestation periods, and the days until day 4 of lactation

**Frequency of treatment**

Once/day, 7days/week

**Doses / concentrations**

**Remarks**

Doses / Concentrations:

0 (vehicle), 40, 200, and 1000 mg/kg bw/day

Basis:

actual ingested

**No. of animals per sex per dose**

12 animals/sex/dose

**Control animals**

yes, concurrent vehicle

---

**Examinations**

**Parental animals: Observations and examinations**

CAGE SIDE OBSERVATIONS: Yes

- Time schedule:

Males and females: 3 times/day

BODY WEIGHT: Yes

- Time schedule for examinations:

Males: Days 1, 4, 8, 11, 15, 22, 25, 29, 32, 36, 39, 42, and the day of necropsy

Females: Twice a week during the precopulation period (days 1, 4, 8, 11, and 15); gestation days 0, 4, 7, 11, 14, 17, and 20; lactation days 0 and 4; and the day of necropsy

FOOD CONSUMPTION: Yes

Males: Days 1, 4, 8, 11, 15, 32, 36, 39, and 42 in dosing period

Females: Days 1, 4, 8, 11, and 15; gestation days 1, 4, 7, 11, 14, 17, and 20; lactation days 2 and 4

OTHER: Females: Numbers of corpus luteum and implantation site on the day of necropsy

**Estrous cyclicity (parental animals)**

Vaginal smears were collected from all females in the main groups and microscopically examined every day from the day after the start of administration until the day copulation was confirmed.

During the pre-mating administration period, vaginal smear pictures were classified as proestrus,

estrus, metestrus or diestrus and examined for the frequency of estrus and interval between estruses (estrous cycle). During the mating period, vaginal smears were examined for the presence of sperm.

#### **Sperm parameters (parental animals)**

Parameters examined in P male parental generations: testes weight, epididymides weight

#### **Litter observations**

PARAMETERS EXAMINED: The following parameters were examined in F1 offspring [number and sex of pups, stillbirths, live births, postnatal mortality, presence of gross anomalies, and weight].

GROSS EXAMINATION OF DEAD PUPS: Yes, for external and internal abnormalities.

#### **Postmortem examinations (parental animals)**

SACRIFICE:

Male animals: Rats were euthanized by exsanguination under ether anesthesia on the day after the last administration.

Maternal animals: Rats were euthanized by exsanguination under ether anesthesia on day 4 of lactation.

GROSS PATHOLOGY: Yes (see tables)

HISTOPATHOLOGY: Yes (epididymis, prostate, seminal vesicle, testis, ovary, uterus, vagina, and gross abnormal sites)

ORGAN WEIGHTS, Yes: Testes and epididymis

#### **Postmortem examinations (offspring)**

GROSS NECROPSY

- Gross necropsy consisted of external and internal examinations including the cervical, thoracic, and abdominal viscera.

#### **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 ( $p < 0.05$ , two-sided).

Especially,

Implantation index, Stillborn index, Liveborn index, External abnormalities, Viability index: the Steel test ( $p < 0.05$  and  $< 0.01$ , two-sided)

Copulation index, Fertility index, Insemination index, Delivery index: Fisher's exact test ( $p < 0.05$  and  $< 0.01$ , two-sided)

#### **Reproductive indices**

Each parameter was determined by the following equations:

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

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

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

Duration of gestation (days) = day 0 of lactation – day 0 of gestation

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

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

Stillborn index (%) = (No. of stillborn pups/Total No. of pups born) × 100

Liveborn index (%) = (No. of liveborn pups/Total No. of pups born) × 100

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

Sex ratio = No. of liveborn male pups/(No. of liveborn male pups + No. of liveborn female pups)

#### **Offspring viability indices**

Viability index (%) = (No. of surviving pup on day 4 after birth/No. of liveborn pups on day 0 after birth) × 100

## **Results and discussion**

---

## Results: P0 (first parental animals)

---

### General toxicity (P0)

---

#### Clinical signs

effects observed, treatment-related

#### Description (incidence and severity)

see 7.5.1 Repeated dose toxicity: oral.002

#### Body weight and weight changes

effects observed, treatment-related

#### Description (incidence and severity)

see 7.5.1 Repeated dose toxicity: oral.002

#### Food consumption and compound intake (if feeding study)

effects observed, treatment-related

#### Description (incidence and severity)

see 7.5.1 Repeated dose toxicity: oral.002

#### Organ weight findings including organ / body weight ratios

no effects observed

#### Description (incidence and severity)

on reproductive organs

#### Gross pathological findings

no effects observed

#### Description (incidence and severity)

on reproductive organs

#### Histopathological findings: non-neoplastic

no effects observed

#### Description (incidence and severity)

on reproductive organs

## Reproductive function / performance (P0)

---

#### Reproductive function: estrous cycle

no effects observed

#### Reproductive function: sperm measures

not examined

#### Reproductive performance

no effects observed

#### Description (incidence and severity)

on reproductive organs

## Effect levels (P0)

---

<b>Dose descriptor</b> NOAEL
---------------------------------

**Effect level**

200

mg/kg bw/day (actual dose received)

**Sex**

male/female

**Basis for effect level**

other: Nine males and seven females died with tremors and decreased locomotor activity at 1,000 mg/kg bw/day by day 9 of administration.

**Results: F1 generation****General toxicity (F1)****Clinical signs**

no effects observed

**Mortality / viability**

no mortality observed

**Body weight and weight changes**

no effects observed

**Sexual maturation**

not examined

**Organ weight findings including organ / body weight ratios**

not examined

**Gross pathological findings**

no effects observed

**Histopathological findings**

not examined

**Effect levels (F1)****Dose descriptor**

NOAEL

**Generation**

F1

**Effect level**

200

mg/kg bw/day (actual dose received)

**Sex**

male/female

**Basis for effect level**

other: Administration of 4-chlorobenzaldehyde at 1,000 mg/kg bw/day was halted because of the frequent deaths in male and female rats.

**Overall reproductive toxicity**

**Key result**

false

**Reproductive effects observed**

not specified

**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/PDF104-88-1c.pdf](http://dra4.nihs.go.jp/mhlw_data/home/pdf/PDF104-88-1c.pdf)

**Applicant's summary and conclusion** \_\_\_\_\_

**Conclusions**

NOAEL for the rat reproductive/developmental toxicity of 4-chlorobenzaldehyde was determined to be 200 mg/kg bw/day.

**Executive summary**

In the reproduction/developmental toxicity screening test (0, 40, 200, and 1,000 mg/kg bw/day) (OECD TG 421), administration of 4-chlorobenzaldehyde at 1,000 mg/kg bw/day was halted because of the frequent deaths in male and female rats. No effects of this substance on reproductive and developmental parameters were observed at 200 mg/kg bw/day. NOAEL for the rat reproductive/developmental toxicity of 4-chlorobenzaldehyde was determined to be 200 mg/kg bw/day.



---

# References

## REFERENCE\_SUBSTANCE: 4-chlorobenzaldehyde

---

**UUID:** ECB5-0baf6234-8c2b-4e38-9592-5f8028824cc8

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2016-12-21T15:12:20.000+09:00

**Remarks:**

---

### General information

---

**Reference substance name**  
4-chlorobenzaldehyde

### Inventory

---

**Inventory name**  
4-chlorobenzaldehyde

**Inventory**  
EC

**Inventory number**  
203-247-4

**CAS number**  
104-88-1

**Molecular formula**  
C<sub>7</sub>H<sub>5</sub>ClO

**Description**

### Reference substance information

---

**IUPAC name**  
4-chlorobenzaldehyde

#### Synonyms

<b>Identity</b> 4-chlorobenzaldehyde
---

<b>Identity</b> Benzaldehyde, 4-chloro-
--

### CAS information

---

**CAS number**  
104-88-1

## Related substances

---

### Group / category information

USEPA Category: Aldehydes;Neutral Organics

## Molecular and structural information

---

### Molecular formula

C<sub>7</sub>H<sub>5</sub>ClO

### Molecular weight

140.567

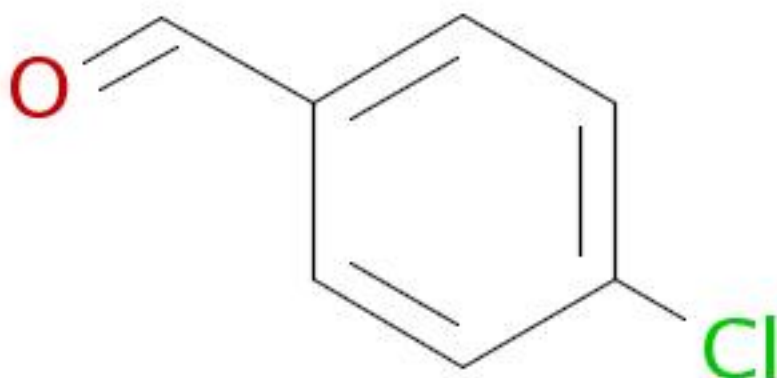
### SMILES notation

Clc1ccc(C=O)cc1

### InChI

InChI=1/C<sub>7</sub>H<sub>5</sub>ClO/c8-7-3-1-6(5-9)2-4-7/h1-5H

### Structural formula



---

# TEST\_MATERIAL\_INFORMATION: 4-chlorobenzaldehyde / 104-88-1 / 203-247-4

---

**UUID:** 3d2fbea2-6dde-3021-a76a-0ed177181828

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2017-01-17T11:42:25.000+09:00

**Remarks:**

---

## Name

4-chlorobenzaldehyde / 104-88-1 / 203-247-4

## Composition

---

### Type

Constituent

### Reference substance

4-chlorobenzaldehyde / 4-chlorobenzaldehyde / 104-88-1 / 203-247-4

### EC number

203-247-4

### EC name

EC Inventory

### CAS number

104-88-1

### CAS name

### IUPAC name

4-chlorobenzaldehyde

## Other characteristics

---

### Details on test material

- Name of test material (as cited in study report): 4-Chlorobenzaldehyde
- Analytical purity: 99.06%
- Lot No.: F7049
- Storage condition of test material: at a cold (temperature 1-10 °C) and dark place, with airtight stopper.
- Stability under test conditions: The stability of test material was identified by analysis of the remainder.

---

# TEST\_MATERIAL\_INFORMATION: 4-chlorobenzaldehyde / 104-88-1 / 203-247-4

---

**UUID:** 6f79d44d-3e27-35b9-9b7c-eae8986a812e

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2017-01-17T12:13:20.000+09:00

**Remarks:**

---

## Name

4-chlorobenzaldehyde / 104-88-1 / 203-247-4

## Composition

---

### Type

Constituent

### Reference substance

4-chlorobenzaldehyde / 4-chlorobenzaldehyde / 104-88-1 / 203-247-4

### EC number

203-247-4

### EC name

EC Inventory

### CAS number

104-88-1

### CAS name

### IUPAC name

4-chlorobenzaldehyde

## Other characteristics

---

### Details on test material

- Name of test material (as cited in study report): 4-chlorobenzaldehyde
- Purity: 99.8%
- Lot/batch No.: QJ5LK
- Stability under test conditions: Stable
- Storage condition of test material: a cool (2-8 °C) and dark place (in a refrigerator), with an airtight stopper
- Dosing solution storage condition: under room temperature (19-23 °C) in a brown glass bottle
- Other: The dosing solution was used within 7 days of preparation.

---

# TEST\_MATERIAL\_INFORMATION: 4-chlorobenzaldehyde / 104-88-1 / 203-247-4

---

**UUID:** 45ad02fc-c5d2-3583-9271-63c80c9ae2e2

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2017-01-17T12:03:06.000+09:00

**Remarks:**

---

## Name

4-chlorobenzaldehyde / 104-88-1 / 203-247-4

## Composition

---

### Type

Constituent

### Reference substance

4-chlorobenzaldehyde / 4-chlorobenzaldehyde / 104-88-1 / 203-247-4

### EC number

203-247-4

### EC name

EC Inventory

### CAS number

104-88-1

### CAS name

### IUPAC name

4-chlorobenzaldehyde

## Other characteristics

---

### Details on test material

- Name of test material (as cited in study report): 4-chlorobenzaldehyde
- Purity: 99.06%
- Lot/batch No.: F7049
- Storage condition of test material: in a hermetically sealed and light-resistant container at cool (2-9 °C) place
- Stability under test conditions: The stability of test material was identified by analysis of the remainder.

---

# TEST\_MATERIAL\_INFORMATION: 4-chlorobenzaldehyde / 104-88-1 / 203-247-4

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**UUID:** 73144eba-5d92-3a50-bce8-9ff650585cd4

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2017-01-17T12:09:01.000+09:00

**Remarks:**

---

## Name

4-chlorobenzaldehyde / 104-88-1 / 203-247-4

## Composition

---

### Type

Constituent

### Reference substance

4-chlorobenzaldehyde / 4-chlorobenzaldehyde / 104-88-1 / 203-247-4

### EC number

203-247-4

### EC name

EC Inventory

### CAS number

104-88-1

### CAS name

### IUPAC name

4-chlorobenzaldehyde

## Other characteristics

---

### Details on test material

- Name of test material (as cited in study report): 4-chlorobenzaldehyde
- Analytical purity: 99.06%
- Lot/batch No.: F7049
- Storage condition of test material: in a hermetically sealed and light-resistant container at cool (2-9 °C) place
- Stability under test conditions: The stability of test material was identified by analysis of the remainder.

# LITERATURE: A reproduction/developmental toxicity screening test in rats treated orally with 4-chlorobenzaldehyde

---

**UUID:** 7a906d06-e2b2-33e0-8940-38ed3aee35a3

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2017-01-17T11:45:20.000+09:00

**Remarks:**

---

## General information

---

**Reference Type**

study report

**Title**

A reproduction/developmental toxicity screening test in rats treated orally with 4-chlorobenzaldehyde

**Author**

MHLW Japan

**Year**

2011

**Bibliographic source**

available in the web of Japan Existing Chemical Data Base (JECDB) at [http://dra4.nihs.go.jp/mhlw\\_data/jsp/SearchPageENG.jsp](http://dra4.nihs.go.jp/mhlw_data/jsp/SearchPageENG.jsp)

**Testing facility**

BoZo Research Center

# LITERATURE: A reproduction/developmental toxicity screening test in rats treated orally with 4-chlorobenzaldehyde

---

**UUID:** 22272062-d1a6-3a5d-b062-26dd76552a05

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2017-01-17T12:13:20.000+09:00

**Remarks:**

---

## General information

---

**Reference Type**

study report

**Title**

A reproduction/developmental toxicity screening test in rats treated orally with 4-chlorobenzaldehyde

**Author**

MHLW, Japan

**Year**

2009

**Bibliographic source**

available in the web of Japan Existing Chemical Data Base (JECDB) at [http://dra4.nihs.go.jp/mhlw\\_data/jsp/SearchPageENG.jsp](http://dra4.nihs.go.jp/mhlw_data/jsp/SearchPageENG.jsp)

**Testing facility**

BoZo Research Center



# LITERATURE: In Vitro Chromosomal Aberration Test of 4-chlorobenzaldehyde on Cultured Chinese Hamster Cells.

---

**UUID:** 6b56bfc3-62e7-3148-942a-7fe72e1f0f90

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2017-01-17T12:09:01.000+09:00

**Remarks:**

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## General information

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**Reference Type**

study report

**Title**

In Vitro Chromosomal Aberration Test of 4-chlorobenzaldehyde on Cultured Chinese Hamster Cells.

**Author**

MHLW, Japan

**Year**

2011

**Bibliographic source**

Japan Existing Chemical Data Base (JECDB)

**Testing facility**

Safety Research Institute for Chemical Compounds Co., Ltd.

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# LEGAL\_ENTITY: National Institute of Health Sciences

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

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2019-09-03T10:05:28.255+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](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.

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## General information

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### 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](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

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### Other IT system identifiers

<b>IT system</b> LEO
<b>ID</b> 10767
<b>IT system</b> IUCLID4
<b>ID</b> 16558402024DIV750

## Contact information

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### Contact address

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**Region / State**

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**Country**

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**Person**

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Akihiko

**Organisation**

National Institute of Health Sciences, Japan

**Department**

Division of Risk Assessment

**Title**

Dr

**Country**

Japan

# LITERATURE: Reverse Mutation Test of 4-chlorobenzaldehyde on Bacteria.

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**UUID:** 73ff8829-35c8-3d27-9841-753cf862a37e

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2017-01-17T12:03:06.000+09:00

**Remarks:**

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## General information

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**Reference Type**

study report

**Title**

Reverse Mutation Test of 4-chlorobenzaldehyde on Bacteria.

**Author**

MHLW, Japan

**Year**

2012

**Bibliographic source**

Japan Existing Chemical Data Base (JECDB)

**Testing facility**

Safety Research Institute for Chemical Compounds Co., Ltd.

# LITERATURE: Twenty-eight-day Repeat Dose Oral Toxicity Test of 4-Chlorobenzaldehyde in Rats

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**UUID:** a17c23ea-ac60-3e55-a207-d084befee301

**Dossier UUID:**

**Author:** SuperUser

**Date:** 2017-01-17T11:42:25.000+09:00

**Remarks:**

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## General information

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**Reference Type**  
publication

**Title**  
Twenty-eight-day Repeat Dose Oral Toxicity Test of 4-Chlorobenzaldehyde in Rats

**Author**  
MHLW

**Year**  
2011

**Bibliographic source**  
available in the web of Japan Existing Chemical Data Base (JECDB) at [http://dra4.nihs.go.jp/mhlw\\_data/jsp/SearchPageENG.jsp](http://dra4.nihs.go.jp/mhlw_data/jsp/SearchPageENG.jsp)

**Testing facility**  
Safety Research Institute for Chemical Compounds Co., Ltd.