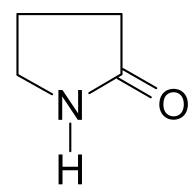
2-Pyrrolidone



CAS Number 616-45-5

Existing Chemical : ID: 616-45-5
CAS No. : 616-45-5
EINECS Name : 2-pyrrolidone
EC No. : 210-483-1
TSCA Name : 2-Pyrrolidinone
Molecular Formula : C4H7NO

Producer related part

Company : Toxicology and Regulatory Affairs

Creation date : 06.10.2002

Substance related part

Company : Toxicology and Regulatory Affairs

Creation date : 06.10.2002

Status : Memo :

Printing date : 13.08.2003

Revision date :

Date of last update : 13.08.2003

Chapter (profile) : Chapter: 1.0.1, 1.2, 2.1, 2.2, 2.3, 2.4, 2.5, 2.6.1, 3.1.1, 3.1.2, 3.3.1, 3.3.2,

3.5, 4.1, 4.2, 4.3, 4.4, 5.1.1, 5.1.2, 5.1.3, 5.1.4, 5.4, 5.5, 5.6, 5.7, 5.8.1, 5.8.2

Reliability (profile) : Reliability: without reliability, 1, 2, 3, 4

Flags (profile)

1. General Information

ld 616-45-5 **Date** 13.08.2003

1.0.1 APPLICANT AND COMPANY INFORMATION

Type : lead organisation

Name : Toxicology and Regulatory Affairs
Contact person : Elmer Rauckman PhD DABT

Date

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Remark: Participating Members of Consortium

BASF Corporation

International Specialty Products

31.12.2002

1.2 SYNONYMS AND TRADENAMES

2-Ketopyrrolidine

08.12.2002

2-Oxopyrrolidine

08.12.2002

2-Pyrol

08.12.2002

4-Aminobutyric acid lactam

08.12.2002

Apha-pyrrolidinone

08.12.2002

Butanoic acid, 4-amino-, lactam

08.12.2002

Butyrolactam

08.12.2002

Gamma-aminobutyric lactam

08.12.2002

Gamma-butyrolactam

08.12.2002

2. Physico-Chemical Data

ld 616-45-5 **Date** 13.08.2003

2.1 MELTING POINT

Value : = 25 °C

Test substance :

2-Pyrrolidone CAS No. 616-45-5

Reliability : (2) valid with restrictions

2 Handbook Value

Flag : Critical study for SIDS endpoint

06.10.2002 (21)

2.2 BOILING POINT

Value : = 245 °C at 1010 hPa

Decomposition

Method

Year

GLP : no data

Test substance

Test substance : CAS No. 616-45-5 2-Pyrrolidone

Reliability : (2) valid with restrictions

Handbook values are assigned 2

Flag : Critical study for SIDS endpoint

06.10.2002 (16)

2.3 DENSITY

Type : density

Value : = 1.116 g/cm³ at 25 °C

Method

Year :

GLP : no data

Test substance

Test substance : CAS No. 616-45-5 2-Pyrrolidone

Reliability : (2) valid with restrictions

2 Handbook Value

Flag : Critical study for SIDS endpoint

06.10.2002 (16)

2. Physico-Chemical Data

ld 616-45-5 **Date** 13.08.2003

2.4 VAPOUR PRESSURE

Value : = .013 hPa at 25 °C

Decomposition

Method

Year

GLP : no data

Test substance

Remark: Given in reference as 0.00949 mm. Converted to hPa by multiplying by

1.33 hPa/mm

Supported by IUCLID 2000 value of 0.04 hPa at 20 C as referenced in

BASF AG, Sicherheitsdatenblatt Pyrrolidon dest. (28.06.1993)

Reliability : (2) valid with restrictions

2 Handbook Value

Flag : Critical study for SIDS endpoint

31.12.2002 (18)

2.5 PARTITION COEFFICIENT

Partition coefficient : octanol-water Log pow : = -.71 at 25 °C

pH value

Method : OECD Guide-line 107 "Partition Coefficient (n-octanol/water), Flask-

shaking Method"

Year

GLP : no data

Test substance

Method :

Approximately 25 ml each of water and 1-octanol were mixed in a shake flask with 0.063, 0.137 or 0.166 grams of test substance in three separate trials at 25 deg C. After separation of the layers, the test substance was determined in quadruplicate in each phase with using gas chromatography. The mean P(OW) values for each of the three trials were 0.193, 0.193 and 0.206. These values were averaged and the log was determined to give a

mean Low K0/w of -0.71

Remark : SRC Physical Properties Data Base lists result 0r -0.85 as published by

Sasaki, H et al. (1991).

EPIWIN, Log Kow (KOWWIN v1.66 estimate) = -0.32 based on smiles

structure.

Test substance

2-Pyrrolidone CAS No. 616-45-5

Reliability : (1) valid without restriction

1, Modern guideline study

Flag : Critical study for SIDS endpoint

31.12.2002 (6)

2. Physico-Chemical Data

Id 616-45-5 **Date** 13.08.2003

2.6.1 SOLUBILITY IN DIFFERENT MEDIA

Solubility in : Water : = at °CValue pH value : = 10 - 11 concentration : 100 g/l at 20 °C Temperature effects :

Examine different pol. :

at 25 °C pKa

Description

Stable

Deg. product

Method

Year

GLP : no data

Test substance

: pH of solution is from: BASF AG, Sicherheitsdatenblatt Pyrrolidon dest. Remark

(28.06.1993)

Result : Miscible

Test substance : CAS No. 616-45-5 2-Pyrrolidone

Reliability : (2) valid with restrictions

2 Handbook value

Flag : Critical study for SIDS endpoint

06.10.2002 (26)

ld 616-45-5 **Date** 13.08.2003

(19)

3.1.1 PHOTODEGRADATION

Type air

Light source

Light spectrum

Relative intensity based on intensity of sunlight

INDIRECT PHOTOLYSIS

Sensitizer

Conc. of sensitizer : 1500000 molecule/cm³

Rate constant .000000000012 cm³/(molecule*sec)

Degradation ca. 50 % after 10.8 hour(s)

Deg. product

Method

Year 2002

GLP

Test substance

Result SMILES: C1CCC(=O)N1

> CHEM: 2-Pyrrolidone MOL FOR: C4 H7 N1 O1

MOL WT: 85.11

- SUMMARY (AOP v1.90): HYDROXYL RADICALS ------Hydrogen Abstraction = 6.4334 E-12 cm3/molecule-sec Reaction with N, S and -OH = 5.5000 E-12 cm3/molecule-sec Addition to Triple Bonds = 0.0000 E-12 cm3/molecule-sec Addition to Olefinic Bonds = 0.0000 E-12 cm3/molecule-sec Addition to Aromatic Rings = 0.0000 E-12 cm3/molecule-sec Addition to Fused Rings = 0.0000 E-12 cm3/molecule-sec

OVERALL OH Rate Constant = 11.9334 E-12 cm3/molecule-sec

HALF-LIFE = 0.896 Days (12-hr day; 1.5E6 OH/cm3)

HALF-LIFE = 10.756 Hrs

: Toxicology and Regulatory Affairs Source Test substance : CAS No. 616-45-5 2-Pyrrolidone

: (2) valid with restrictions Reliability

Calculated by acceptable method : Critical study for SIDS endpoint

Flag 08.12.2002

3.1.2 STABILITY IN WATER

abiotic Type t1/2 pH4 at °C

t1/2 pH7 > 1 year at 25 °C

t1/2 pH9 at °C

Deg. product

Method

Year 2002 **GLP**

Test substance

Method : Estimation using HYDROWIN 1.67.

no

Input was SMILES notation: C1CCC(=O)N1

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Remark : Furthuer supports comes from the "Handbook of Chemical Property

Estimation Methods" (2) in which is it is indicated that the mean hydrolytic

half-life for a series of amides is in the range of 300 years

(2) J.C. Harris in Lyman W, Reehl, W and Rosenblat, D. Handbook of Chemical Property Estimation Methods. American Chemical Society,

Washingotn D.C. 1990, page 7-6

This estimated is supported by the known properties of amides.

For example in the textbook "Organic Chemistry" (1), Vollhardt states that "Amides are the least reactive of the carboxylic derivatives, mainly because of the extra resonance capacity of the nitrogen lone electron pair. As a consequence, their nucleophilic addition-eliminations require relatively harsh conditions. For example, hydrolysis occurs only on prolonged heating in strongly acidic or basic water"

(1) Vollhardt, K. "Organic Chemistry" WH Freeman and Co, New York,

1987, p 815.

Result: HYDROWIN Program (v1.67) Results:

SMILES: C1CCC(=0)N1 CHEM: 2-Pyrrolidone MOL FOR: C4 H7 N1 O1

MOL WT: 85.11

---- HYDROWIN v1.67 Results -----

AMIDE: -N-C(=O)-C-

Compound has an amide group; C=O located at SMILES atom #4

Hydrolysis Rate Extremely Slow or t1/2 > 1 Year

Source : Toxicology and Regulatory Affairs **Test substance** : 2-Pyrrolidone CAS No. 616-45-5

Reliability : (2) valid with restrictions

Estimated using an acceptable method with comfirmation from both chemical principles and experimental data on surrogate compounds.

Flag : Critical study for SIDS endpoint

30.11.2002 (20)

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

Type : fugacity model level III

Media : other: all

Air : % (Fugacity Model Level I)

Water : % (Fugacity Model Level I)

Soil : % (Fugacity Model Level I)

Biota : % (Fugacity Model Level II/III)

Soil : % (Fugacity Model Level II/III)

Method : other Year : 2002

Method : Determined using the Level 3 EQC Model found in EPIWIN 3.05. Actual

values were used for measured physicochemical parameters. The

ld 616-45-5 **Date** 13.08.2003

degredation times applied using the BIOWIN were validated by experimental data on the test substance and/or surrogate compounds

Result :

Level III Fugacity Model (Full-Output):

Chem Name : 2-Pyrrolidone

Molecular Wt: 85.11

Henry's LC: 1.44e-008 atm-m3/mole (Henrywin program)

Vapor Press: 0.00949 mm Hg (user-entered)

Log Kow : -0.71 (user-entered) Soil Koc : 0.0799 (calc by model)

Concentration		Half-Life	Emissions	
()	percent)	(hr)	(kg/hr)	
Air	0.403	21.5	1000	
Water	46.5	360	1000	
Soil	53	360	1000	
Sedimet	0.0776	1440	0	

	Fugacity	Reaction	Advect	Reaction	Advection
	(atm)	(kg/hr)	(kg/hr)	(percent)	(percent)
Air	1.36e-011	153	47.4	5.09	1.58
Water	4.62e-013	3 1050	547	35.1	18.2
Soil	1.94e-011	1200	0	40	0
Sed	3.85e-013	0.439	0.018	0.0146	0.00061

Persistence Time: 392 hr Reaction Time: 489 hr Advection Time: 1.98e+003 hr

Percent Reacted: 80.2 Percent Advected: 19.8

Half-Lives (hr), (based upon Biowin (Ultimate) and Aopwin):

Air: 21.51 Water: 360 Soil: 360 Sediment: 1440

Biowin estimate: 2.957 (weeks)

Advection Times (hr):
Air: 100
Water: 1000
Sediment: 50000

Source

Calculated by Toxicology and Regulatory Affairs, 2002

Test substance

CAS No. 616-45-5 2-Pyrrolidone

Reliability : (1) valid without restriction

Calculated by an acceptable method using measured physicochemical

parameters.

31.12.2002 (19)

3.3.2 DISTRIBUTION

ld 616-45-5 **Date** 13.08.2003

3.5 BIODEGRADATION

Type : aerobic

Inoculum : activated sludge, domestic

Contact time : 28 day(s)

Degradation : = 73 (±) % after 28 day(s) **Result** : readily biodegradable

Deg. product

Method

Year

GLP

Test substance: other TS

Method : Japanese MITI test

Remark :

Surrogate material

Test substance

1-Methyl-2-pyrrolidinone CASNO 872-50-4

Surrogate material

Reliability : (2) valid with restrictions

Published study result

Flag : Critical study for SIDS endpoint

03.08.2003 (17)

Type : aerobic

Inoculum

Contact time

Degradation : (±) % after

Result : readily biodegradable

Deg. product

Method : other: estimation

Year :

GLP :

Test substance :

Method : The structure was run through BIOWIN 4.00, as found in EPIWIN 3.05.

This software predicts, with excellent accuracy, the ease and relative rate of aerobic biodegredation. Estimates are primarily based on a fragment

approach.

Remark

This estimate is supported by the high rate of biodegredation observed in

the Zahn Wellens procedure (BASF AG, Labor Oekologie;

unveroeffentlichte Untersuchung (Pyrrolidon dest., 1977)) and the ready biodegredability of the N-methyl derivitive (NMP, see HSDB) which, based on judgement and BIOWIN modeling, is expected to be slightly more

difficult to biodegrade than 2-Pyrrolidone.

Result :

SMILES: C1CCC(=O)N1 CHEM: 2-Pyrrolidone MOL FOR: C4 H7 N1 O1

MOL WT: 85.11

BIOWIN v4.00 Results

Linear Model Prediction : Biodegrades Fast Non-Linear Model Prediction: Biodegrades Fast

9 / 48

ld 616-45-5 **Date** 13.08.2003

Ultimate Biodegradation Timeframe: Weeks Primary Biodegradation Timeframe: Days

MITI Linear Model Prediction : Biodegrades Fast MITI Non-Linear Model Prediction: Biodegrades Fast

LINEAR BIODEGRADATION PROBABILITY 0.9172 NON-LINEAR BIODEGRADATION PROBABILITY 0.9889

MITI LINEAR BIODEGRADATION PROBABILITY 0.6448
MITI NON-LINEAR BIODEGRADATION PROBABILITY 0.8408

A Probability Greater Than or Equal to 0.5 indicates --> Readily Degradable

A Probability Less Than 0.5 indicates --> NOT Readily Degradable

SURVEY MODEL - ULTIMATE BIODEGRADATION 2.9569 SURVEY MODEL - PRIMARY BIODEGRADATION 3.9304

Interpretation, Primary & Ultimate:

Result Classification:

5.00 -> hours 4.00 -> days 3.00 -> weeks 2.00 -> months

1.00 -> longer

Test substance

2-Pyrrolidone CAS No. 616-45-5

Reliability : (2) valid with restrictions

Estimated using an acceptable method.

31.12.2002

Type : aerobio

Inoculum : other: activated sludge, non-adapted

Contact time

Kinetic of testsubst.

Degradation : > 90 (±) % after 9 day(s)

Result :

: 1 day(s) = 5 % 5 day(s) = 80 % 7 day(s) = 89 % 9 day(s) = 99 %

%

Method: This Inherent Biodegradation test followed the Zahn-Wellens procedure.

Triplicate determinations were made using the test substance at a final concentration of about 500 mg/L and in 2 L of culture containg 100 ml of non-adapted sludge.

Elimination was determined by measuring total organic carbon (TOC) at 0 and 3 hours; and at 1, 5, 7, and 9 days after start of the test.

The methodology follows the Zahn Wellens test procedure.

ld 616-45-5 **Date** 13.08.2003

Remark

Although the conditions do not meet the OECD 301 series, the results clearly demonstrate that non-adapted sludge flora are capable of fully

degrading the test material in a short time.

Technically, this test only indicates inherent biodegradation; however, the rapidity of the biodegradation is consistent with a "readily biodegradable"

material.

Test substance

2-Pyrrolidone, Distilled

Conclusion

The test material is considered "inherently biodegradable" showing rapid

biodegredation.

Reliability : (2) valid with restrictions

The raw data for this triplicate determination was available for review;

although some details were missing the method is scientifically defensible.

03.08.2003 (7)

4.1 ACUTE/PROLONGED TOXICITY TO FISH

Type : static

Species: Brachydanio rerio (Fish, fresh water)

Exposure period : 96 hour(s)
Unit : mg/l

 NOEC
 : = 4600 measured/nominal

 LC0
 : = 4600 measured/nominal

 LC50
 : = 6800 measured/nominal

 LC100
 : = 10000 measured/nominal

Limit test

Analytical monitoring : ye

Method : OECD Guide-line 203 "Fish, Acute Toxicity Test"

Year

GLP : yes Test substance :

Method : METHOD: Followed standard laboratory protocol for OECD 203 (April

1984).

DETAILS OF TEST: Static

DILUTION WATER SOURCE: Municipal water, carbon treated

DILUTION WATER CHEMISTRY: pH 8.0-8.6, total hardness about 2.5 mmol/L, acid capacity about 5.5 mmol/L, TOC not given, TSS not given.

STOCK AND TEST SOLUTION PREPARATION: Test substance added

neat to test water 20 minutes before placing fish in aquaria.

VEHICLE/SOLVENT AND CONCENTRATIONS: Dilution water, concentrations 0, 50, 100, 1000, 2150, 4640, 10000 mg/L

STABILITY OF THE TEST CHEMICAL SOLUTIONS: Assured by analytical

determination

EXPOSURE VESSEL: All-glass aquaria, 30 x22 x 24 cm, containing 10 L $\,$

water and filled to a depth of about 17 cm.

REPLICATES, FISH PER REPLICATE: One replicate, 10 fish per replicate

TEMP PHOTOPERIOD FOOD: Test temperature 22-23 °C, photoperiod 16 hours light and 8 hours dark, food withdrawn one day before exposure,

ANALYTICAL CHEMISTRY DETERMINATIONS: TS measured at one and

96 hours.

Result

Nominal concentrations were: 50, 100, 1000, 2150, 4640 or 10000 mg/L for

test.

Analytical concentrations were: 53, 95, 959, 2146, 4580 or 10221 mg/L at

one-hour

Analytical concentrations were: 38, 98, 947, 2084, 4600 or 9935 mg/L at

96-hours

Id 616-45-5 Date 13.08.2003

> pH measurements at one hour were control to high concentration: 8.6, 8.5, 8.4, 8.5, 8.6, 8.6, 8.6; at 96 hours 8.3, 7.0, 9.8, 8.2, 8.2, nd.

> Oxygen levels were above 7 mg/L in most instances at 1, 24, 48, 73, or 96 hours.

Temperature remained at 22° throughout the study.

Mortality: There was no mortality except at the high concentration (10,000 mg/L) where the cumulative mortality at 24 hours was 6/10, at 48 hours was 8/10 at 72 and 96 hours was 10/10.

Clinical signs: The only reported effects were for the 10,000 mg/L group at 24 hours where apathy and tumbling were reported in surviving fish.

Test substance

2-Pyrrolidone CAS No. 616-45-5 Purity 99.7%

Conclusion

Reliability

The 96-hour LC50 is between 4,600 and 10,000 mg/L (based on nominal concentrations). According to the OECD 203 guideline the geometric mean (6,783) of these concentrations may be used to approximate the

LC50.

or LC50 = 6,800 mg/L: (1) valid without restriction

Guideline study under GLP with no significant problems noted.

Flag : Critical study for SIDS endpoint

13.08.2003 (12)

4.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES

Type static

Species Daphnia magna (Crustacea)

Exposure period : 48 hour(s) Unit : mg/l

EC0 : = 500 measured/nominal **EC50** : > 500 measured/nominal

Limit Test

Method Directive 84/449/EEC, C.2 "Acute toxicity for Daphnia"

Year

GLP no data

Test substance

Method Daphnia magna (2-24 hours old) were exposed to the test substance in

> four replicates of five animals (20/group) at nominal concentrations of 0, 31.25, 62.5, 125, 250, or 500 mg/L for 48 hours. The dilution water was prepared from tapwater by dilution with distilled water to reduce the hardness, addition of sulfuric acid to reduce the alkalinity, filtration to remove particulates and passing the water through activated carbon to remove chlorine. Final dilution water had a total hardness of 2.44 mmol/L, an alkalinity of 0.80 mmol/L (to pH 4.3), a calcium:magnesium ratio (molar) of 4:1, a sodium:potassium (molar) ratio of 10:1 and a pH range of 7.7 to

8.3.

Loading of daphnids was 2 ml/daphnid using 10 ml centrifuge tubes. The temperature was maintained at 293 deg K. Diffuse light was on 16 hours/day at an intensity of 570 microSiemens/cm. The dilution water was

ld 616-45-5 4. Ecotoxicity **Date** 13.08.2003

> bubbled with oil-free air initially to saturate it with oxygen. The test substance dilutions were prepared from a stock at 500mg/l (also the high

concentration) by dilution.

Daphnids were examined at 3, 6, 24 and 48 hours after initiation.

Result

The initial pH did not differ between concentrations and was in the range of 8.11-8.27. The final pH was not concentration dependent and ranged from 7.59 to 8.14. Oxygen concentrations, measure at 0 and 48 hours of the test, were higher at the beginning (9.30-9.42 mg/L) than at the end of the 48 hour exposure period (5.54-8.55) and there was no apparent

relationship of DO levels to test-substance concentration.

No daphnids was found immobilized by the treatment and no adverse

effects were reported at any concentration.

Test substance

2-Pyrrolidone CAS No. 616-45-5, distilled, purity > 99.5%

Conclusion

The NOEC and EC-0 were found to be 500 mg/L

The EC-50 was found to be > 500 mg/L (These are based on nominal concentrations)

Reliability (1) valid without restriction

> Guideline study, with good documentation including copies of raw data. Although the test did not use analytical measurements of test substance

concentration, it is known to be stable in water.

Flag Critical study for SIDS endpoint

03.08.2003 (8)

Type static

Species Daphnia magna (Crustacea)

Exposure period 96 hour(s) Unit ma/l

EC₀ = 1000 measured/nominal **EC50** : > 1000 measured/nominal

Analytical monitoring

Method Groups of 20 Daphnia magna were exposed to the test substance at either

10, 100, or 1000 mg/L. Groups were made up of four replicates of five

daphnids in 300 ml of dilution water containing test substance.

Observations were made at least at 24 hours, 96 hours, 7 days, 14 days

and 21 days.

Remark

The stability of the test substance in water was not established. Other information support the test substance being stable in water for at least the initial 48 hour period. Stability at the 3-week time was likely compromised

by biodegradation of the test substance.

Result

No mortality occurred in the first 96 hours of exposure in any group. At the end of the three-week exposure period the number of surviving daphnids was 17/20, 18/20 and 12/20 for the 10, 100 and 1000 mg/L groups,

respectively.

Test substance

2-Pyrrolidone

Conclusion

The 96-hour EC50 for Daphnia magna is > 1000 mg/L under these

conditions.

Reliability : (2) valid with restrictions

Although this study is old and details are limited, the conduct was similar to

modern guidelines and the study was conducted according to a

scientifically defensible method. The availability of the original data sheets

add to the reliability of the work.

31.12.2002 (27)

Type : static

Species : Daphnia pulex (Crustacea)

Exposure period : 48 hour(s)
Unit : mg/l

EC50 : = 13.21 calculated

Analytical monitoring : no

Method : Daphnia pulex were cultured in 2-L jars of reconstituted hard water (20OC;

pH,7.6-8.0; dissolved oxygen, 60-100% saturation; hardness 160-180 mg/L as CaCO; alkalinity 110-120 mg/L as CaCO). To minimize leaching, dissolution and sorption of toxicants from the water only glassware and tubing made from perfluorocarbon plastic was used for culturing and testing. The daphnid food was a mixture of the four algal species plus cerophyl at a ratio of 1:1:1:1:4. The daphnids were fed five times a week

with 3 mL of food per liter of culture water.

The 48-h tests were conducted with 10 neonates (<24 h old) in five concentrations of each toxicant and the control. Toxicant concentrations (in 150 mL of reconstituted hard water) were at least 50% of the next concentration. The six test beakers, covered with parafilm, were placed in a constant temperature water bath at 20 deg C with a photoperiod of 16 h light, 8 h dark. Test animals were not fed during the experiment. After 48 h the daphnids were pipetted into a watch glass and examined for immobilization.

Mean effective concentration (EC50) and standard error were calculated from the immobilization data for valid toxicity tests (American Society for Testing and Materials 1980). A mean was taken from three valid tests. To calculate EC10, EC50, and EC90 values, we used a computer modification (Peltier et al. 1985) of Finney's (1952) probit analysis. Statistical comparisons were made on logarithmically transformed EC50's using analysis of variance (ANOVA) and Tukey's HSD test (Steel and Torrie 1960).

(Finney DH (1952) Statistical methods in biological assay. C. Griffin and Co Ltd., London, 661 pp)

(Peltier WH, Weber Cl(eds) (1985) Methods for measuring the acute toxicity of effluents to freshwater and marine organisms, 3rd ed Environ Monitor Support Lab, US Environ Protect Agency, Cincinnati, Rep no 600/4-85-013)

(Steel RGD, Torrie JH (1960) Principles and Procedures of Statistics, McGraw Hill, New York)

Result :

The results from all studies in ther report are presented in the table below:

Compound EC50 (mg/L) Mean SE

DDT (D. magna) 0.0011 0.0001

DDT (17 C) 0.0019 0.0001 Chlordane (D. magna) 0.097 0.005 Nicotine 0.242 0.02

Nicotine (170C) 0.326 0.074

Pentachlorophenol (D. magna) 2.00 0.0
Pentachlorophenol 2.5 0.1
1-methylpyrrolidine 2.08 0.20
Isoxanthopterin 2.97 0.47
2-amino-4,6-dimethylpyridine 9.19 1.85

2-pyrrolidinone 13.21 4.02 2-(2-hydroxyethyl)pyridine 13.82 3.60

Mortality as a function of concentration was not given in the article.

The range of toxicity and the reported SE indicate that studies were conducted in the appropriate concentration range for each test material.

Test substance

2-Pyrrolidone CAS No. 616-45-5 Purity >= 97%

Reliability : (2) valid with restrictions

Good, this is a published study by a National Laboratory in a peer reviewed journal conducted using a scientifically defensible method. Stability data

on the test compound are lacking.

13.08.2003 (25)

Type : static

Species : other aquatic mollusc: Planorbella trivolvis

Exposure period : 96 hour(s)
Unit : mg/l

NOEC : = 112 measured/nominal EC0 : = 112 measured/nominal EC50 : > 112 measured/nominal

Limit Test : yes Analytical monitoring : no

Method : One group of 10 snails was exposed to a solution of 100 microliters/L test

substance at a temperature of 18 C for a period of 96 hours. The initial dissolved oxygen level was 9.2 mg/L and the initial pH was 7.7. The final dissolved oxygen level was 2.6 mg/L with a final pH of 7.2. The snails were identified as Helisoma trivolvis, which are currently known as

Planorbella trivolvis.

Result

All snails survived the 96-hour exposure period.

Test substance

2-Pyrrolidone

Conclusion

The 96-hour EC50 for Planorbella trivolvis is > 112 mg/L under these

conditions.

Reliability : (2) valid with restrictions

31.12.2002 (27)

Type : static

Species: other aquatic worm:

Exposure period : 96 hour(s)

Unit : mg/l

NOEC : = 112 measured/nominal EC0 : = 112 measured/nominal EC50 : > 112 measured/nominal

Limit Test : yes **Analytical monitoring** : no

Method

Year :

GLP : no data

Test substance

Method : One group of 10 worms was exposed to a solution of 100 microliters/L test

substance at a temperature of 18 C for a period of 96 hours. The initial dissolved oxygen level was 9.2 mg/L and the initial pH was 7.7. The final dissolved oxygen level was 2.6 mg/L with a final pH of 7.2. The aquatic worms were identified as Dugesia tigrine, which is a common freshwater

platyhelminth.

Test substance :

2-Pyrrolidone

Conclusion

The 96-hour EC50 for Dugesia tigrine is > 112 mg/L under these

conditions.

Reliability : (2) valid with restrictions

31.12.2002 (1) (27)

4.3 TOXICITY TO AQUATIC PLANTS E.G. ALGAE

Species : Scenedesmus subspicatus (Algae)

Endpoint : growth rate
Exposure period : 96 hour(s)
Unit : mg/l

EC10 : = 8 calculated EC50 : = 84 calculated

Limit test

Analytical monitoring : no

Method : other: DIN 38412 L9

Year

GLP : no Test substance :

Method : Cells were placed in quadruplicate cultures of growth medium according to

the method of DIN 38412 L9 containing 0, 25, 50, 100, 250 or 500 mg/L test substance. These concentrations were selected on the basis of a preliminary test at concentrations of 0, 5, 50 or 500 mg/L. Cell counts were determined by counting six replicates from each quadruplicate culture at 0, 24, 48, 72 and 96 hours of incubation. Fluorescence was also determined at these same time-points. pH was measured at the beginning and end of the 90-hour incubation period. The temperature of incubation was a

constant 24.8 deg. C.

Statistical Method: Tallerida and Jacob, The Dose-Response Relation in

Pharmacology Pages 98-103 pub. Springer Verlag 1979

Remark: the ECOSAR (v0.99f) program using the neutral organics model predicts a

96-hour EC50 of 4777

Date 15.00.200

Result

The following results are listed in the order

0, 25, 50, 100, 250 or 500 mg/L: The beginning and end pH values were Start: 7.84, 7.87, 7.89, 7.86, 7.89, 7.88 End: 7.92, 7.99, 8.04, 8.07, 8.12, 8.13

Mean cells counts (X 1000) were:

t= 0: 34, 38, 32, 34, 33, 35 t=24: 106, 94, 88, 62, 51, 51 t=48: 235, 191, 165, 150, 149, 136 t=72: 618, 514, 405, 239, 311, 230 t=96: 1866, 1408, 1042, 334, 279, 407

The changes in fluorescence did not correlate with the cell growth.

From these data the EC10 and EC50 for growth rate at 96 hours were determined to be 20 and 353 mg/L and the EC10 and EC50 for biomass

were determined to be 8 and 84 mg/L.

The 72-hour EC10 and EC50 for biomass were 4 and 253 mg/L

Test substance

2-Pyrrolidone CAS No. 616-45-5, distilled, purity > 99.5%

Reliability : (1) valid without restriction

Guideline study, with good documentation.

Flag : Critical study for SIDS endpoint

31.12.2002 (9)

4.4 TOXICITY TO MICROORGANISMS E.G. BACTERIA

Type : aquatic

Species: Pseudomonas putida (Bacteria)

Exposure period : 17 hour(s)
Unit : mg/l

EC10 : = 9268 calculated

Analytical monitoring : no

Method : other: Bringmann-Kuehn Test

Year : 1988 GLP : no Test substance :

Method : Bacteria were added to flasks containing salts, dilute growth substrate and

test material at 0, 156.25, 312.5, 625, 1250, 2500, 5000, 7500, or 10000 mg/L test material. Flasks were incubated for 17 hours at 297 deg K and

bacterial growth was estimated by absorption of light at 436 nm.

Remark: At concentrations below 10,000 mg/L, the test substance appears to have

stimulated bacterial growth under these conditions.

Result : Bacterial growth, expressed as percent of control after 17 hours incubation

was:

4. Ecotoxicity

ld 616-45-5 **Date** 13.08.2003

TS Conc	Bacterial growth
mg/L	% of control
0	100
156.25	159
312.5	160
625	162
1250	159
2500	150
5000	151
7500	129
10000	73

Test substance : 2-Pyrrolidone, Distilled

Conclusion : The EC10 was calculated to be 9268 mg/L

Reliability : (2) valid with restrictions

Guideline-type study using a scientifically defensible method.

Documentation good.

08.12.2002 (14)

5.1.1 ACUTE ORAL TOXICITY

Type : other: Limit Test Value : > 5000 mg/kg bw

Species : rat

Strain : Sprague-Dawley
Sex : male/female

Number of animals: 10Vehicle: waterDoses: 5000 mg/kg

Method

Year : 1979 GLP : no data

Test substance

Method : Five rats of each sex were given a single oral dose of test material by oral

gavage at a limit dose of 5000 mg/kg-bw. The test material was dissolved in distilled water and administered as a 50% wt/vol solution to Sprague-

Dawley rats that had been fasted overnight. Male rats weighed

approximately 250 grams and females approximately 200 grams at the time of dosing. Animals were observed regularly for mortality and adverse

clinical signs and were weighed on days 4, 7 and 13.

The total observation period before sacrifice was 14 days. Necropsy

findings were not given in the report.

Result: No animal died during the study. Average body weights of males were 250,

236, 269 and 297 g on days 0,4, 7 and 13, respectively. Average body weights of females were 200, 201, 211 and 216 g on days 0,4, 7 and 13,

respectively. No adverse clinical findings were reported.

Test substance : 2-Pyrrolidone, Pure

Conclusion: The acute oral LD50 of the test substance is greater than 5000 mg/kg

bodyweight for both male and female rats.

Reliability : (2) valid with restrictions

Reliability is good as a standard procedure was followed; however, the

study lacks details concerning observations and necropsy.

Flag : Critical study for SIDS endpoint

03.08.2003 (5)

Type : LD50

Value : ca. 8000 mg/kg bw

:

Species : rat
Strain : no data
Sex : no data

Number of animals

Vehicle : water

Doses

Method

Year : 1961

ld 616-45-5 5. Toxicity Date 13.08.2003

GLP : no

Test substance

Method The study was conducted as part of the "toxicological pre-testing" for this

> material. The pre-testing consisted of acute oral dosing of rats, inhalation risk-test in rats, i.p. ALD determination in mice, skin and eye irritation.

Details of each procedure are not given in the report.

In this study, the ALD50 (Approximate Median Lethal Dose) was stated as Result

about 8.0 g/kg at both 24 hours and 8 days. It is presumed that the observation time was 8 days. Clinical signs were given as convulsions, dyspnea and lying on side: however, it cannot be determined from the report if these signs refer to mice administered TS i.p. or the rats administered TS orally. Likewise, there is no indication of the dose

corresponding to these signs or the time of their occurrence.

2-Pyrrolidone, Distilled, solid Test substance

21.11.2002 (13)

5.1.2 ACUTE INHALATION TOXICITY

Type other: Inhalation Risk Test

Value

Species rat

Strain

Sex

Number of animals 6 Vehicle

Doses

Exposure time 8 hour(s)

Method other: BASF Inhalation Risk Test

Year 1961 GLP no Test substance

Method The study was conducted as part of the "toxicological pre-testing" for this

material. The pre-testing consisted of acute oral dosing of rats, inhalation risk-test in rats, i.p. ALD determination in mice, skin and eye irritation.

Details of each procedure are not given in the report.

Result : Under the conditions of this study no animal died as a result of the

> exposure to saturated vapor for 8 hours. It is noted in the report that no abnormalities were detected at necropsy; however, the length of the post-

exposure observation period is not specified in the report.

Test substance 2-Pyrrolidone, Distilled, solid

Conclusion : It can be concluded that the 8-hour inhalation LD50 for 2-Pyrrolidone is

greater than the air saturation concentration of the test substance in air at

30 deg C. Which is approximately 80 ppm.

Reliability : (2) valid with restrictions

A reliability of 2 is assigned. Although some important details are lacking

this study was conducted according to a standard procedure that is

scientifically defensible.

21.11.2002 (13)

5.1.3 ACUTE DERMAL TOXICITY

Type : LD50

Value : > 2000 mg/kg bw

Species : rabbit

Strain : New Zealand white

Sex : male/female

Number of animals : 10

Vehicle

Doses : 2000

Method : OECD Guide-line 402 "Acute dermal Toxicity"

Year : 1992 **GLP** : yes

Test substance :

Method

Following a quarantine period of at least one week, five healthy male and five healthy female New Zealand Albino rabbits were randomly assigned to the treatment group. The pretest weight range was 2.3 - 2.6 kg for males and 2.1 - 2.5 kg for females. The animals were housed 1/cage in suspended wire mesh cages. Bedding was placed beneath the cages and changed twice/week. Fresh Purina Rabbit Chow (Diet #5321) was provided daily. Water was available ad libitum. The animal room, reserved exclusively for rabbits on acute tests, was temperature controlled, had a 12 hour dark/light cycle.

The test article was used as received and the dose was based on the sample weight as calculated from the specific gravity. The test article was applied to the prepared dermal site, one time, by syringe type applicator at a dose level of 2.0 g/kg. The test site was covered with a gauze patch, secured with non-irritating tape and gentle pressure was applied to the gauze to aid the distribution of the test article over the area. The torso was wrapped with plastic that was secured with non-irritating tape. At 24-hours after initiation, the patches were removed and residual test article was removed with distilled water.

.

The animals were observed 1, 2 and 4 hours post dose and once daily for 14 days for toxicity and pharmacological effects. Animals were observed twice daily for 14 days for mortality. The test sites were scored for dermal irritation at 24 hours post dose and on days 7 and 14 using the numerical Draize scale

Body weights were recorded pretest, weekly and at death or termination. All animals were examined for gross pathology. Abnormal tissues were preserved in 10% buffered formalin and saved for possible future microscopic examination.

Result

All animals survived the 2000 mg/kg dermal application. There were no abnormal systemic signs noted in 9/10 animals. One male exhibited red staining of the nose/mouth area and an apparent cataract in the right eye on day 5, with the ocular abnormality persisting through day 14 but this was considered to result from a slef-inflicted injury unrelated to test material administration. Body weight gains were normal at all weighing periods. Dermal reactions were slight to well-defined on day 1 but were absent on days 7 and 14. Necropsy did not reveal any treatment related changes.

Test substance : 2-Pyrol, no further information

Conclusion: The dermal LD50 was found to be > 2000 mg/kg-bw

Reliability : (1) valid without restriction

Guideline study under GLP with no significant problems noted.

Flag : Critical study for SIDS endpoint

30.11.2002 (24)

5.4 REPEATED DOSE TOXICITY

Type : Sub-chronic

Species : rat

Sex : male/female
Strain : Wistar
Route of admin. : drinking water
Exposure period : 90 days
Frequency of treatm. : daily
Post exposure period : none

Doses : 600, 2400, 7200 or 15000 ppm in drinking water

Control group : yes, concurrent vehicle

NOAEL : = 2400 ppm **LOAEL** : = 7200 ppm

Method : OECD Guide-line 408 "Subchronic Oral Toxicity - Rodent: 90-day Study"

Year : 1981 GLP : yes Test substance :

rest substance

Method : 2-Pyrrolidone was administered to groups of 10 male and 10 female Wistar rats at doses of 0; 600; 2,400; 7,200 and 15,000 ppm in the drinking water

over a period of 3 months.

Wistar rats (Chbb: THOM (SPF)) were obtained from Dr. Karl Thomae GmbH, Biberach/Riss, FRG. Rats were identified unambiguously by ear tattoo. Animals were individually housed in type DK III stainless steel wire cages Becker & Co., Castrop-Rauxel). Animal rooms were air-conditioned with temperatures in the range 20 - 24°C and relative humidity in the range 30 - 70%. The day/night cycle was 12 hours (light from 06.00 a.m. - 06.00 p.m.).

Test solutions were analysed at the start and end of the study to assure that the concentrations were correct and the 4-day stability was assessed as 97%. The mixtures were prepared at no less than 4-day intervals. Water consumption was determined once/week over a period of 4-days. Animals were weighed weekly and given a thorough physical examination at each weighing. Food consumption was determined weekly. Urine samples were taken on day 85, blood was sampled and analyzed on study day 88, the final bodyweight was recorded on day 91 and necropsies were conducted over days 92 to day 95,

Food consumption, water consumption and body weight were determined each week. The animals' state of health was checked each day. When the animals were weighed they were subjected to an additional comprehensive clinical examination

Clinincal chemistry parameters were: alanine aminotransferase, aspartate aminotransferase, alkaline phosphatase - serum-gamma-glutamyltransferase

Blood chemistry parameters were: sodium, potassium, chloride, inorganic

5. Toxicity

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phosphate, calcium, urea, creatinine, glucose, total bilirubin, total protein, albumin, globulins, triglycerides, cholesterol, magnesium.

In addition complete hematology and urinalysis were conducted.

At necropsy, major organs were weighed and sections were fixed for histopathology. All animals were subjected to gross-pathological assessment, followed by histopathological examination using a complete tissue list.

Statistical methods: Means and standard deviations for the variables food consumption, body weight, body weight change, water consumption and test substance intake (except control group) were calculated for the animals of each test group. They were printed out in the summary and individual value tables, with the exception that for test substance intake and body weight change only summary tables were prepared. For the parameters food consumption, water consumption, body weight and body weight change a parametric one-way analysis of variance was done via the F-test (ANOVA). If the resulting p-values were equal to or less than 0.05, a comparison of each dose group with the control group was carried out. These comparisons were performed simultaneously via Dinnett's test for the hypothesis of equal means. If the results of this test were significant, labels (* for,p < 0.05, ** for p < 0.01) were printed together with the group means in the tables. Both tests were performed two-sided. Statistical analysis of histopathology was conduced with a proprietary computer program.

The following tissues were examined and preserved at necropsy:

- brain
- pituitary gland
- thyroid and parathyroid glands
- thymus
- trachea
- lungs
- heart
- aorta
- salivary glands (mandibular and sublingual)
- liver
- spleen
- kidneys
- adrenal glands
- pancreas
- testes/ovaries
- uterus/vagina
- epididymides, prostate, seminal vesicles
- skin
- esophagus
- stomach (forestomach and glandular stomach)
- duodenum
- jejunum
- ileum
- cecum
- colon
- rectum
- urinary bladder
- lymph nodes (mesenteric, mandibular)

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- female mammary gland
- skeletal muscle
- sciatic nerve
- bone marrow (femur)
- eyes
- femur with knee joint
- sternum with marrow
- spinal cord (cervical, thoracic and lumbar)
- extraorbital lacrimal gland
- all gross lesions

The Following Tissues Were examined microscopically in high-dose and control animals (and other groups as indicated)

- brain
- pituitary gland
- thyroid
- parathyroid
- thymus (and all females in all groups)
- trachea
- lungs (and all animals in all groups)
- heart
- aorta
- salivary glands (mandibular and sublingual)
- liver (and all animals in all groups)
- spleen
- kidneys (and all animals in all groups)
- adrenal glands (and all animals in all groups)
- pancreas
- testes/ovaries
- uterus/vagina
- epididymides, prostate, seminal vesicles
- skin
- esophagus
- forestomach (and all animals in all groups)
- glandular stomach (and all animals in all groups)
- duodenum
- jejunum
- ileum
- cecum
- colon
- rectum
- urinary bladder
- lymph nodes (mesenteric, mandibular)
- female mammary gland
- skeletal muscle
- sciatic nerve
- sternum with marrow
- femur with joint
- bone marrow (femur)
- eyes
- femur with knee joint
- sternum with marrow
- spinal cord (cervical, thoracic and lumbar)
- All gross lesions were examined in all groups

For a full description of the procedures used to examine reproductive organs see the robust summary for fertility.

Remark

The study was carried out according to following guidelines:

- EC Commission Directive 87/302/EEC of 18 November, 1987; Part B: Methods for the determination of Toxicity; Sub-chronic Oral Toxicity Test; 90-day repeated oral dose using rodent species; Official Journal of the European Communities No. L 133, p. 8-11, 1988
- OECD Guidelines for Testing of Chemicals; Method No. 408: Subchronic Oral Toxicity Rodent: 90-day study; May 12, 1981

Result

Substance intake::

Mean test material consumption in mg/kg- day were:

- + males: 33, 184, 529 and 1062 mg/kg
- + females 42, 230, 643 and 1189 mg/kg

No animal died during the study and no adverse clinical signs were noted.

Other effects by dose group:

- *** Test group 4 (15,000 ppm; about 1,125 mg/kg body weight)
- -decreased food and water consumption in both sexes
- decreased body weight gains, male's BW were 9% lower than controls and female's were 8% lower than controls on day 91
- prolonged prothrombin times in rats of each sex
- decrease in total protein, globulins, triglycerides and creatinine in both sexes
- increased urinary specific gravity in the males reduced urinary volume in the males
- dark yellow discoloration of urine specimens in the males
- increase in the mean relative kidney weights in males and females
- *** Test group 3 (7,200 ppm; about 586 mg/kg body weight)
- slight decrease of food consumption in female animals
- slight decrease of water consumption in both sexes
- slightly decreased body weights in females, 6% less than controls on day 91
- decreased body weight gains of 7% (males) and 16% (females) on day 91
- decrease in creatinine in both sexes
- decrease in total protein in the females
- increased urinary specific gravity in the males reduced urinary volume in the males
- dark yellow discoloration of urine specimens in the males
- increase in the mean relative kidney weights in males
- *** Test group 2 (2,400 ppm; about 207 mg/kg body weight) and
- no substance-related effects
- *** Test group 1 (600 ppm; about 37 mg/kg body weight)
- no substance-related effects

Mean Terminal Body and Kidney Weights (Absolute and Relative)

MALES (grams)

	(3.5	- /		
Group Body		Kidney	Kidney	
		(absolute)	(relative)	
0	471	2.97	0.68	
600	460	2.93	0.69	
2400	458	3.05	0.72	
7200	452	3.11	0.73*	
15000	428*	3.13	0.77**	

FEMALES (grams)

Group	Body	Kidney	Kidney	
	•	(absolute)	(relative)	
0	265	1.92	0.79	
600	263	2.00	0.83	
2400	269	1.99	0.80	
7200	248	1.93	0.84	
15000	242*	2.03	0.89**	

Note: A finding of "altered cellular composition of the thymic cortex" was reported in all dosed groups of females. A second 90-day study was conducted at 0, 50 and 15,000 ppm in drinking water using groups of five female rats to investigate the significance of this finding. It this second study the identical finding was present; however, it also occurred in controls. In addition, retrieval and examination of thymus slides from controls animals in other studies were examined and were also found to have the same "pathology". Therefore, this was considered incidental and not compound related.

Test substance

2-Pyrrolidone CAS No. 616-45-5 Purity 99.7%

Conclusion

The kidney appears to be a target organ at dose levels of 7,200 ppm (about 586 mg/kg) in the drinking water and above. The NOAEL is 2,400

ppm in drinking water or about 207 mg/kg-bw-day

Reliability : (1) valid without restriction
Flag : Critical study for SIDS endpoint

03.08.2003 (10)

5.5 GENETIC TOXICITY 'IN VITRO'

Type : Salmonella typhimurium reverse mutation assay

System of testing

Test concentration : 0, 0.1, 1.0, 5.0, 10, 25, 50, 100 and 150 microliters per plate

Cycotoxic concentr. : 150 microliters per plate

Metabolic activation: with and without

Result : negative
Method : other
Year : 1987
GLP : yes
Test substance :

27 / 48

Method

: S. typhimurium strains TA1535, TA1538, TA100, TA1537, TA98 were tested using a plate incorporation technique both with and without metabolic activation. Aroclor 1254 induced rat liver S-9 was used for metabolic activation at a rate of 0.5 ml S-9 per plate when used with the overlay procedure. Test and control materials were incorporated directly into the overlay agar with the bacteria.

Plates were prepared and read in triplicate and the entire assay was repeated a second time (independent repeat). Colonies were counted using an automated Biotran II colony counter except when accurate counts could not be obtained (e.g. precipitate formation).

Concentrations of test substance were selected based on a preliminary toxicity assay at 14 concentration levels using two-fold dilutions from a high concentration of 150 microliter per plate (for liquids). 150 microliters per plate was used as the top concentration in the studies because this is the limit dose for the test and because this concentration reduced the number of TA-100 revertant colonies by approximately 50% in a preliminary doserangefinding test.

Concentrations tested were 0, 0.1, 1.0, 5.0, 10, 25, 50, 100 and 150 microliters per plate for all strains in both of the two independent repeats.

The solvent and negative control substance was distilled water. Positive controls were:

Without metabolic activation Sodium azide at 10 mcg/ plate for strain TA-1535 and TA-100 Quinacrine mustard at 5 mcg/ plate for strain TA-1537 2-Nitrofluorene at 10 mcg/ plate for strains TA-1538 and TA-98

With metabolic activation, 2-Anthramine at 2.5 mcg/ plate for all strains

Statistical Methods

Formal statistical methods were not used to evaluate the data. Evaluations considered if a dose-response was observed and strain-specific evaluation criteria.

For strains TA-1535, TA-1537 and TA-1538, the data set is evaluated as positive if a dose-response is observed over a minimum of three test concentrations and the increase in revertants is equal to or greater than three times the solvent control value at the peak of the dose-response. The solvent control value should be within the normal range for evaluating the results.

For strains TA-98 and TA-l00, the data set is evaluated as positive if a dose-response is observed over a minimum of three test concentrations and the increase in revertants achieves a doubling of the solvent control value at the peak of the dose-response. The solvent control value should be within the normal range for evaluating the results.

In the preliminary study on TA-100, the test material was toxic to the indicator only at 150 microliters per plate as evidenced by the reduced number of revertants on the minimal media plates (about a 50% reduction).

The results of the initial and independent assays conducted on the test material at dose levels ranging from 0.1 to 150 microliters per plate in the

Result

absence and presence of metabolic activation did not exhibit increased numbers of his+ revertant colonies.

The positive control treatments in both the nonactivation and S9 activation assays induced large increases in the revertant numbers with all the indicator strains, which demonstrated the effectiveness of the S9 activation system and the ability of the test system to detect known mutagens.

Test substance : 2-Pyrrolidone CAS No. 616-45-5 Purity by GLC 99.9 Area % source

BASF

Conclusion: The test material, 2-Pyrrolidone, did not exhibit genetic activity in any of

the assays conducted in this evaluation and was not mutagenic to the Salmonella tvphimurium indicator organisms under the test conditions

according to the established evaluation criteria.

Reliability : (1) valid without restriction

Guideline-like study under GLP

Flag : Critical study for SIDS endpoint

06.08.2003 (22)

Type : other: Aneuploidy Induction in Yeast

System of testing : Saccharomyces cerevisiae

Test concentration : 0, 289.6, 321.0, 352.2, 383.3, 414.2, or 445.0 mM

Cycotoxic concentr. : 321 and above

Metabolic activation : without Result : positive

Method

Year : 1987 GLP : no data

Test substance

Method

Diploid strain D61.M of Saccharomyces cerevisiae, developed by F.K. Zimmermann, was used for the detection of aneuploidy and other genetic events. Its genetic constitution and the detailed procedures for its use in detecting aneuploidy have been previously described in detail. In brief: recessive alleles (cyh2, cycloheximide resistance; ade6, white-adenine requirement; leul, leucine requirement) of three genes are arranged on both sides of the centromere on one copy of chromosome VII. Simultaneous expression of all three recessive alleles in the same clone can result either from loss of the homologous chromosome VII carrying the wild-type alleles or from simultaneous multiple events of recombination or mutation, which are expected to be extremely rare.

Ten parallel 5-ml cultures were grown in YEPD medium until they attained a titer of approximately 5-7 x 10exp7 cells/ml. A 0.1-ml aliquot was removed from each culture and plated onto the cycloheximide-YEPD medium to select cultures with low spontaneous rates of cycloheximide resistance. The 5-ml cultures were stored at 4°C until use. A culture that was determined to have a low spontaneous frequency of cycloheximide resistance (typically < 1 x 10exp6) was diluted 1:10 into fresh YEPD medium and incubated at 28C for 4 hr to bring the cells into exponential growth phase before addition of the test chemical.

The exponential phase culture was adjusted to 5 x 10exp6 cells/ml in YEPD medium. Treatments were carried out in 2-nil aliquots in glass test tubes by adding microliter quantities of the test chemical either directly or from a stock solution of the chemical in water prepared just before use. The concentration of the stock solutions was dictated by the level of toxicity, which had been determined in preliminary experiments. The growing yeast

cells were treated in a shaker water bath at 28°C for 4 hr; then the cultures were refrigerated at 4°C in a water bath for 16 hr. The cold holding period was followed by a second 4-hr incubation at 28°C before the cultures were diluted and plated on the appropriate media. (The interruption of growth by cold temperature storage greatly enhances the induction of aneuploidy by a number of solvent chemicals). When necessary, cultures were diluted to approximately 1-2 x 10exp7 cells/ml, and 0.1-ml aliquots were plated directly onto the selective cycloheximideYEPD medium to determine the resistant population. Appropriate dilutions were plated onto YLPD medium to determine the surviving population. Plates were incubated for 5-7 days, and colonies were enumerated. On selective cycloheximide-YEPD medium the resistant colonies were either red or white. The red colonies resulted from the occurrence of genetic events such as gene conversion or mutation affecting the CYH2 locus only and not from chromosome malsegregation. The cycloheximide-resistant white colonies are presumably due to chromosome loss because the recessive cyh2 and the recessive ade6 alleles are being simultaneously expressed. To confirm that the white resistant colonies are really monosomic for chromosome VII, each colony to be tested was streaked onto YEPD master plates, which were incubated overnight at 28C, and then replicas were plated onto both a synthetic complete medium and onto the same medium lacking leucine. White (ade6) and cycloheximide-resistant (cyh2) colonies must also require leucine (leul) to be considered monosomic.

Remark

: In a subsequent paper, these same authors found no aneuploidy potentiation of 2-Pyrrolidinone with nocodazole. They discussed the potential mechanism of solvent-induced aneuploidy in terms of the fact that microtubles dissociate in the cold to their tubulin subunits and polymerize again as the temperature is raised. The solvents were speculated to inhibit or accelerate the rate of repolymerization (Mayer and Goin, Mut Rech. 201:413-421, 1988).

Several factors indicate that this result is not relevant to hazard assessment to man.

Solvent-induced aneuploidy appears to be a special case.

Solvent-induced aneuploidy is enhanced by cold incubation, which was part of the protocol in this investigation.

The concentration range where effects are reported is narrow range and coincides with toxicity.

The concentrations where effects are reported are extremely high and impossible to achieve under normal industrial conditions in man.

Common non-genotoxic solvents such as acetone are known to induce this effect under the special conditions employed in this study.

Result

Positive results on the induction of aneuploidy by 1-methyl-2-pyrrolidirone and 2-pyrrolidinone were recorded as the number of cycloheximide-resistant white colonies observed and the fraction of these colonies that were Leu-. Aneuploidy frequencies were calculated by using these numbers as the numerator and the population screened as the denominator. In cases in which only a few white colonies were found, all were tested for their leucine requirement. When many white colonies were observed, all were counted, and a representative number (usually 25) was tested. The number of red cycloheximide-resistant colonies was

determined and was found not to increase with test material concentration. As red-resistant colonies arise as a result of other genetic events, they served as a control showing that other genetic effects such as mutation or recombination were not induced by the test chemical.

The frequency of aneuploidy increased with the dosage of each test chemical. 1-Methyl-2-pyrrolidinone was active between 150 and 230 mM, while 2-pyrrolidinone was active between 350 and 450 mM, and appeared to be slightly less toxic in comparable ranges. As there was no increase with concentration for either chemical in the frequency of the red cycloheximide-resistant colonies. Therefore, aneuploidy rather than other nuclear genetic effects were being induced by these chemicals.

Data are shown in the table.

Test substance Attached document 2-Pyrrolidone CAS No. 616-45-5 from Aldrich Chemical Co

Y-table-HP600.bmp

Test M Conc (mM)	Percent Survival	Pop Screened X 10 ⁶	Total White Colonies	Aneuploidy Frequency x10 ⁶ CFU
0	100	4.73	10	1.27
289.6	98	5.20	42	6.79
321.0	61	4.35	48	9.71
352.2	42	4.43	65	10.56
383.3	23	3.28	98	17.93
414.2	8	1.75	120	21.94
445.0	7	1.50	120	19.20

Reliability : (2) valid with restrictions

The method was well described and sufficient details and data were

presented to indicate that this study has good reliability.

28.11.2002 (23)

Type : Cytogenetic assay

System of testing

Test concentration

Cycotoxic concentr. : High doses minimally cytotoxic.

Metabolic activation: with and without

Result : negative

Method : OECD Guide-line 473

Year : 1987 **GLP** : yes

Test substance :

Method : 2-Pyrrolidone was tested for its ability to induce chromosomal aberrations in human lymphocytes following in vitro exposure in the presence and absence of a metabolizing system.

Based on a pretest to determine the highest experimental dose and in consideration of the cytotoxicity actually found in the present cytogenetic investigations, 3500 mcg/ml, 2500 mcg/ml and 1250 mcg/ml culture medium in the experiment without S-9 mix. or 6000 mcg/ml, 5000 mcg/ml and 2500 mcg/ml culture medium in the experiment with metabolic activation, were selected. This selection was based on the quality of the metaphases and not on the mitotic index because the test substance

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concentrations causing reduction in the mitotic index are at dose levels that severely affect chromosomes; thus, no longer allowing evaluation.

Duplicate cultures were used for all experimental points. The solvent was distilled water.

Negative controls (untreated and solvent) and positive controls both without S-9 mix (0.2 mcg mitomycin C/ml culture medium) and with metabolic activation (6 mcg cyclophosphamide/ml culture medium) were also tested.

Heparinized human venous blood was added to the culture medium (chromosome medium 1A with PHA). After mitogen stimulation of the lymphocytes using PHA and incubation at 37°C for 48 hours. The cultures were treated with test substance without S-9 mix for 24 hours; in the experiment with S-9 mix (from Aroclor-induced rats) test substance treatment lasted 2 hours followed by a reincubation for 22 hours using fresh culture medium without test substance. About 2 - 3 hours prior to harvesting the cells, Colcemid was added to arrest cells in a metaphase-like stage of mitosis (C-metaphase). After preparation of the lymphocyte chromosomes and staining with Giemsa, 100 metaphases of each culture in the case of the test substance, untreated control and solvent control, or 50 cells of each culture in the case of positive controls, were analyzed for chromosomal aberrations.

Statistical Procedure:

The Fisher exact test was applied to determine significant differences between the relative frequencies of a characteristic of two groups, and it was used to answer the questions of whether there are significant differences between control groups (untreated controls and solvent controls) and dose groups with regard to the rate of structural aberrant metaphases.

Result

** Assay without metabolic activation:::

Untreated controls

10 (5.0%) aberrant cells including gaps and 2 (1,0%) aberrant cells excluding gaps were found

Solvent controls:

12 (6.0%) aberrant metaphases including gaps and 5 (2.5%)aberrant metaphases excluding gaps were found

3500 mcg/ml:

8 (4.0%) chromosomally damaged cells including gaps and 2 (1.0%) aberrant cells excluding gaps were detected.

2500 mcg/L:

14 (7.0%) aberrant metaphases including gaps and 6 (3.0%) chromosomally damaged cells excluding gaps were observed.

1250 mcg/ml:

17 (8.5%) aberrant cells including gaps and 2 (1.0%) aberrant metaphases excluding gaps were found.

0.2 mcg mitomycin C/ml:

With 44 (44%) aberrant cells including gaps and 37 (37%) aberrant mitosis excluding gaps including 2 multiple aberrant metaphases and 5 cells with

exchanges, the positive control substance led to the expected increase in the number of chromosomally damaged cells.

No differences regarding aneuploidies (hyperploid metaphases) and polyploidies between the various dose groups and the negative controls were observed.

Assay with metabolic activation:::

Untreated control:

4 (2.0%) aberrant mitosis including gaps only were found.

Solvent contro1:

15 (7.5%) aberrant metaphases including gaps and 4 (2.0%) chromosomally damaged cells excluding gaps were found.

6000 mcg/ml:

17 (8.5%) chromosomally damaged cells including gaps and 2 (1.0%) aberrant cells excluding gaps were observed.

5000 mcg/ml:

16 (8.0%) chromosomally damaged cells including gaps and 1 (0.5%) aberrant cells excluding gaps were observed.

2500 mcg/ml:

13 (6.5%) chromosomally damaged cells including gaps and 1 (0.5%) aberrant cells excluding gaps were observed.

6 mcg cyclophosphamide/ml:

27 (27%) chromosomally damaged cells including gaps and 20 (20%) aberrant cells excluding gaps were observed, which was the expected increase for positive controls.

No differences regarding aneuploidies (hyperploid metaphases) and polyploidies between the various dose groups and the negative controls were observed.

Test substance Conclusion

: 2-Pyrrolidone CAS No. 616-45-5 Purity 99.9%

: According to the results of the present study, the test substance 2-pyrrolidone did not lead to any increase in the number of aberrant metaphases including and excluding gaps when compared to the solvent controls either without S-9 mix or after adding a metabolizing system. 2-Pyrrolidone is evaluated not to be a chromosome-damaging (clastogenic) agent under in vitro conditions using human lymphocytes, under these experimental conditions.

Reliability : (1) valid without restriction

Guideline study under GLP with no significant problems noted.

Flag : Critical study for SIDS endpoint

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5.6 GENETIC TOXICITY 'IN VIVO'

Type : Micronucleus assay

Species: mouseSex: male/femaleStrain: NMRIRoute of admin.: i.p.

Exposure period : 16, 24 and 48 hours

Doses : 2000, 1000, and 500 mg/kg-bw

Result : negative

Method : OECD Guide-line 474 "Genetic Toxicology: Micronucleus Test"

Year : 1993 GLP : yes Test substance :

Method :

Male and female animals (NMRI mice, Charles River GmbH, WIGA) were assigned to the test groups using a randomization plan prepared with an appropriate computer program. Animals were housed in Makrolon cages, in groups of 5 according to sex in fully air-conditioned rooms with a range of 20 - 24°C for temperature and a range of 30 - 70% for relative humidity. Before treatment, animals were transferred to Makrolon cages and housed individually under the same conditions until the end of the test. The day/night rhythm was 12 hours (light from 6.00 - 18.00 hours). Standardized pelleted feed (Kliba Haltungsdidt, Klingentalmühle AG) and drinking water from bottles were available ad libitum.

Doses selected were 2000, 1000 and 500 mg/kg-bw and were selected on the basis of a preliminary toxicity study. In this study, the highest recommended dose of 2000 mg/kg was administered and survived by all animals but led to signs of toxicity such as irregular respiration, piloerection, abdominal position, apathy and squatting posture; the general state of the animals was poor.

Five Male and female animals per sacrifice interval and dose group were given test substance dissolved in distilled water 2000 mg/kg, 1000 mg/kg and 500 mg/kg body weight. Treatment consisted of a single intraperitoneal administration with a volume of 10 ml/kg body weight. As a positive control, 20 mg of cyclophosphamide/kg body weight or 0.15 mg of vincristine/kg body weight, both dissolved in distilled water, were administered to groups (five animals total, either 2 or 3 of each sex) of male and female animals once intraperitoneally each in a volume of 10 ml/kg body weight. All test substance formulations were prepared immediately before administration.

Sacrifice intervals per dose-group were:

2000 mg/kg; 16, 24 and 48 hours

1000 mg/kg; 24 hours 500 mg/kg 24 hours Controls 24 hours 5. Toxicity

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Preparation of bone marrow: After cutting off the epiphyses, the bone marrow was flushed out of the diaphysis into a centrifuge tube using a cannula filled with fetal calf serum which was at 37°C (about 2 ml/femur). The suspension was mixed thoroughly with a pipette, centrifuged at 1500 rpm for 5 minutes, the supernatant removed the cells were resuspended. One drop of this suspension was dropped onto clean microscopic slides. Smears were prepared using slides with ground edges, the preparations were dried in the air and subsequently stained in eosin and methylene blue solution for 5 minutes, rinsed, placed in fresh distilled water for 2 or 3 minutes and finally stained in Giemsa solution for 12 minutes. After being rinsed twice and clarified with xylene, the preparations were embedded in Corbit-Balsam. Slides were coded before microscopic analysis.

Evlauations: In general, 1000 polychromatic erythrocytes from each male and female animal of every test group was evaluated and investigated for micronuclei. The normochromatic erythrocytes which occur were also scored. The following parameters were recorded:

Number of polychromatic erythrocytes

Number of polychromatic erythrocytes containing micronuclei Number of normochromatic erythrocytes

Number of normochromatic erythrocytes containing micronuclei Ratio of polychromatic to normochromatic erythrocytes Number of small micronuclei (d < D/4) and of large micronuclei (d > D/4)

No statistical methods were employed in data analysis.

Result

Clinical examinations: The single intraperitoneal administration of the solvent in a volume of 10 ml/kg body weight was tolerated by all animals without any signs or symptoms. A dose of 2000 mg/kg body weight of test substance, led to irregular respiration, piloerection, abdominal position and apathy about 30 minutes after administration; the general state of some animals was poor. After treatment of the animals with 1000 or 500 mg/kg, only irregular respiration and piloerection were observed after about 30 minutes. After about 1 - 2 hours clinical signs were no longer observed. Neither the single administration of the positive control substance cyclophosphamide in a dose of 20 mg/kg-bw nor that of vincristine at 0.15 mg/kg-bw caused any evident signs of toxicity.

Micronulei: Mean polychromatic erythrocytes containing micronuclei were:

Negative control (24 hrs)	1.5%
2000 mg/kg (16 hrs)	1.2%
2000 mg/kg (24 hrs)	1.7%
2000 mg/kg (48 hrs)	1.6%
1000 mg/kg (24 hrs)	2.4%
2000 mg/kg (16 hrs)	1.2%
Cyclophosphamide (24 hrs)	13.6%
Vincristine (24 hrs)	83.2%

Administration of test substance did not lead to any increase in the rate of micronuclei. The number of normochromatic or polychromatic erythrocytes containing small micronuclei (d < D/4) or large micronuclei (d > D/4) did not deviate from the solvent control value at anv sacrifice interval. No inhibition of erythropoiesis induced by the treatment of mice with Pyrrolidon-2 was detected; the ratio of polychromatic to normochromatic erythrocytes was always in the same range as that of the control values in all dose groups.

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> The number of normochromatic erythrocytes containing micronuclei did not differ to any appreciable extent in the negative control or in the various

dose groups at any of the sacrifice intervals.

Test substance 2-Pyrrolidone CAS No. 616-45-5 Purity > 99.5%

Conclusion

The number of normochromatic erythrocytes containing micronuclei did not differ to any appreciable extent in the negative control or in the various

dose groups at any of the sacrifice intervals.

Reliability (1) valid without restriction

Guideline study under GLP with no significant problems noted.

Flag Critical study for SIDS endpoint

29.11.2002 (4)

CARCINOGENICITY

5.8.1 TOXICITY TO FERTILITY

Type other: Reproductive Organ Examination from 90-Day Study

Species rat

Sex male/female Strain Wistar

Route of admin. drinking water Exposure period 90-days Frequency of treatm. daily

Premating exposure period

Male Female

Duration of test

No. of generation

studies

Doses 600, 2400, 7200 or 15000 ppm in drinking water

Control group yes, concurrent vehicle

Method 2-Pyrrolidone was tested for subchronic toxicity in a 90-day study. The test

> substance was administered in drinking water to groups of 10 Wistar rats (Strain Chbb:THOM (SPF)) 10 of each sex - in dose groups of 15000,

7200, 2400, 600 and 0 ppm.

Methods followed the European and international guidelines:

EC Commission Directive 87/302/EEC of November 18, 1987; Part B: Methods for the determination of Toxicity Sub-chronic Oral Toxicity Test, 90-day repeated oral dose using rodent species; Official Journal of the European Communities No. L 133, pages 8-11, 1988; and OECD Guideline for Testing of Chemicals; Method No. 408: Subchronic Oral Toxicity -

Rodent: 90-day study; May 12, 1981.

For a more detailed description of the 90-day study conduct and results please see the robust summary for the study in the repeated-dose section.

Briefly: Test solutions were analysis at the start and end of the study to assure that the concentrations were correct and the 4-day stability was assessed as 97%. The mixtures were prepared at no less than 4-day intervals. Water consumption was determined once/week over a period of 4-days. Animals were weighed weekly and given a thorough physical

ood consumption was determined weekly.

examination at each weighing. Food consumption was determined weekly. Urine samples were taken on day 85, blood was sampled and analyzed on study day 88, the final bodyweight was recovered on day 91 and necropsies were conducted over days 92 to day 95.

This robust summary will describe the methods and results of reproductive organ evaluation.

Organ Weights

The testes were weighed in all male rats, and the ovaries were weighed in all female rats. Absolute weights as well as relative weights (related to the terminal body weight) were determined and assessed statistically, using Dunnett's test (two-sided).

Gross lesions

During necropsy, specific attention was given to gross lesions of male reproductive organs (testes, epididymides, prostate gland, seminal vesicles and coagulating glands) and female reproductive organs (ovaries, including oviducts, uterus, including cervix uteri and vagina). In addition, the adrenal glands of all animals were inspected grossly during necropsy and after appropriate fixation; the pituitary glands of all animals were assessed grossly during necropsy after removal of the brain as well as during removal from the skull after appropriate fixation. Further, in females, special attention was given to the gross appearance of the mammary gland and the external genitalia (males: penis, preputium, scrotum, processus vaginalis; females: vulva) were also inspected carefully during necropsy.

Histopathology

The reproductive organs of male rats (testes, epididymides, prostate gland, seminal vesicles and coagulating glands) and the reproductive organs of female rats (uterus, including cervix uteri, ovaries, including oviducts and vagina) were fixed routinely in a 4% aqueous solution of formaldehyde for at least 48 hours. In addition, the adrenal glands and the pituitary gland (both sexes) and parts of the mammary gland (female rats) of all animals were fixed in formaldehyde solution. Any gross lesions noted during necropsy in the external or internal sex organs of male or female rats were also fixed in 4% aqueous formaldehyde solution.

After fixation, the reproductive organs of male (both testes, both epididymides - comprising caput, corpus and cauda epididymidis, prostate gland - comprising dorsolateral and ventral parts, seminal vesicles with attached coagulating glands) and of female rats (both ovaries, uterus, including cervix uteri and vagina) as well as the pituitary gland and the female mammary gland were trimmed, processed to paraplast blocks, cut at a thickness of approximately 3 microns and stained with hematoxylin and eosin (H.& E.). The slides of all animals of the control and of the high dose group were assessed using a light microscope with primary magnifications between 25-400 x. Adrenal glands of all animals were processed, stained with H.& E. and assessed histopathologically.

During histopathological evaluation of reproductive organs, the following were specifically considered:

Testes: histopathology was performed on mid cross sections through both testes. Besides gross lesions such as atrophy or tumors, testicular

histopathological examination looked for treatment-related effects such as focal or diffuse atrophy of the seminal epithelium, retained spermatids, missing germ cell layers or types, multinucleated giant cells or sloughing of spermatogenic cells into the tubular lumen. In addition, attention was given to the morphology of the Sertoli cells (vacuolization) and to the interstitial cells of Leydig (number and morphology).

Epididymides: the examination was performed on a mid longitudinal section through both epididymides, comprising caput, corpus and cauda epididymidis. Besides gross lesions such as atrophy, special attention was given to the presence of sperm granulomas, leukocytic infiltration (inflammation), aberrant cell types within the lumen, and oligospermia or aspermia.

Prostate Gland: histopathology was performed on cross sections through the dorso-lateral and ventral parts of the gland. Special attention was given to looking for inflammatory reactions (acute or chronic, purulent, mixed cellular or lymphocytic, in the glandular acini or in the interstitium). Moreover, the morphology of the acinar cells was assessed carefully (hypertrophy, hyperplasia, atrophy) as was the functional status of the gland (amount of colloid in the acini and its staining properties).

Seminal vesicles: both glands were investigated using cross sections through the mid part of the gland. The attached coagulating glands were also examined (although they were not protocol organs and were, hence, not separately mentioned in the tables of the report). Special attention was given to looking for findings of inflammatory reactions (acute or chronic, purulent, mixed cellular or lymphocytic, in the glandular acini or in the interstitium). Moreover, the morphology of the acinar cells was assessed carefully (hypertrophy, hyperplasia, atrophy) as well as the functional status of both glands (amount of fluid in the acini and its staining properties). Ovaries: histopathological examination was performed on mid cross sections through both organs. Assessment was focused on possible detection of qualitative depletion of the primordial and growing follicle populations, as well as the presence/absence of antral follicles (Graafian follicles) and corpora lutea. Special attention was given to the ovarian interstitium and its cell populations with regard to, atrophy, hypertrophy and/or hyperplasia. A differential ovarian follicle count (DOFC, to detect a quantitative depletion of primordial and/or growing follicles) was not performed.

Oviducts: Not investigated histopathologically.

Uterus (including cervix uteri, which was not listed separately): histopathology was performed on cross sections through the mid part of each uterus horn and on a mid longitudinal section through the cervix uteri with the portio on one side and the base of the uterine horns on the other side. Special attention was give to looking for findings of inflammatory reactions (acute or chronic, purulent, mixed cellular or lymphocytic, in the mucosa or in the glands). Moreover, the morphology of the epithelium, the glands and the musculature were assessed carefully (e.g. for hypertrophy, hyperplasia, atrophy). No specific consideration was given to the status of the sexual cycle according to the cellularity in uterus and cervix uteri.

Vagina: a longitudinal section was performed. Major possible findings which were investigated were inflammatory reactions (acute or chronic, purulent, mixed cellular or lymphocytic in the lumen and/or the wall).

Moreover, the morphology of the epithelium and the underlying musculature was assessed carefully (e.g. for hypertrophy, hyperplasia, atrophy). No specific consideration was given to the status of the sexual cycle according to the cellularity in the vagina.

Remark

Comparison of methodology to pathology performed in a reproductive study:

With the exception of the oviducts, all organs of the male and female genital tract that are examined in a "modern" reproduction toxicity study (e.g. US-EPA OPPTS 870.3800) were investigated both grossly and histopathologically. Further, all other organs required for histopathology in the OPPTS 870.3800 guideline - namely the pituitary and the adrenal glands - were investigated histopathologically. The only significant deviation from OPPTS 870.3800 was that uterus (with oviducts and cervix), epididymides (total weights for both and cauda weight for either one or both), seminal vesicles (with coagulating glands and their fluids), prostate gland, pituitary gland and spleen were not weighed. These organs, however, were grossly inspected and histopathologically assessed. If treatment-related weight changes had occurred, they would likely have been identified by the detailed histopathological examination.

Methods followed the European and international guidelines:

EC Commission Directive 87/302/EEC of November 18, 1987; Part B: Methods for the determination of Toxicity Sub-chronic Oral Toxicity Test, 90-day repeated oral dose using rodent species; Official Journal of the European Communities No. L 133, pages 8-11, 1988; and OECD Guideline for Testing of Chemicals; Method No. 408: Subchronic Oral Toxicity - Rodent: 90-day study; May 12, 1981.

Result

- : Results of Reproductive Organ Weight Determinations
 - 1. MALES: There were no statistically significant deviations of the mean absolute or relative testes weights between treated and control animals.
 - 2. FEMALES: the mean absolute weight of the ovaries was statistically significantly decreased (- 17.0%) in the 7200-ppm dose group. The mean relative ovary weight was the lowest in the 7200-ppm dose group (0.035 mg% = 12.5%); however, this was not statistically significant.

Terminal Body and Reproductive Organ Weights

MALES	3	
Group	Body W	Testes
0	471	3.59
600	460	3.47
2400	458	3.60
7200	452	3.50
15000	428*	3.51

FEMALES Group Body W Ovaries (mg) 265 97.5 600 263 89.6 269 93.8 2400 80.9** 7200 248 15000 242* 89.5

Results of Gross Examination of Reproductive Organs:

- 1. MALES: No gross lesions were noted in the reproductive organs of male rats of any group.
- 2. FEMALES: One female iin the control group, the low dose group, the low mid and the high mid dose groups revealed slight or moderate dilation of the lumen of one or both horns of the uterus. The dilated areas contained a clear (water-like) fluid. No similar or other gross lesions were recorded in animals of the high dose group.

Results of Histopathologic Examination

In the epididymides of one high dose animal, the only microscopic finding recorded was a minimal, unilateral mononuclear cell infiltration.

In the pituitary gland of one control male and one control female animal, small cysts were noted (location was not specified in the report, however, most likely in the distal/glandular part).

Cystic dilation of the uterus (i.e. one or both horns) was observed in each one female rat of the control, low, low mid and high mid dose groups, whereas two high dose females displayed this finding. The severity was graded as moderate in animals that showed this finding on gross examination (one animal each in the control, low, low mid and high mid dose groups). The two high dose females affected only showed slight dilation, which was not seen on gross examination at necropsy.

No microscopic findings were noted in the adrenal cortex, adrenal medulla, female mammary gland, ovaries, prostate gland, seminal vesicles (including the attached coagulating glands), testes and vagina.

Test substance

Conclusion

2-Pyrrolidone CAS No. 616-45-5 Purity 99.7%

: All organs of the male and female genital tract examined in a "modern" reproduction toxicity study, with the exception of oviducts, were investigated grossly and histopathologically. All other organs required for histopathology in the OPPTS 870.3800 guideline were investigated histopathologically. No gross lesions and no microscopic findings were detected that were indicative of an alteration of male or female reproductive performance. The few gross lesions and microscopic findings reported in these organs were all interpreted as incidental lesions, with respect to both incidence and severity.

Although organ weights for uterus (with oviducts and cervix), epididymides (total weights for both and cauda weight for either one or both), seminal vesicles (with coagulating glands and their fluids), prostate gland, pituitary gland and spleen were not taken, these organs were grossly inspected and

histopathologically assessed. If treatment-related adverse effects had occurred they would have been identified histopathologically or grossly.

In summary, the results of the 90-day subchronic toxicity with 2-Pyrrolidone in male and female Wistar rats are regarded as valid to interpret the potential reproductive performance of the animals as being un-altered by

administration of the test article via drinking water.

Reliability : (1) valid without restriction

Guideline study, with good documentation.

Flag : Critical study for SIDS endpoint

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5.8.2 DEVELOPMENTAL TOXICITY/TERATOGENICITY

Species : rat Sex : female

Strain : Sprague-Dawley

Route of admin. : gavage

Exposure period: days 6-15 of gestation

Frequency of treatm. : Daily

Duration of test

Doses : 190, 600, 1900

Control group : yes, concurrent vehicle

NOAEL maternal tox. : = 190 mg/kg bw

NOAEL teratogen. : = 600 mg/kg bw

Result : Not Specific Developmental Toxin
Method : OECD Guide-line 414 "Teratogenicity"

Year

GLP : yes

Test substance :

Method :

Groups of 25 pregnant rats were exposed to the test substance by oral gavage using distilled water as vehicle at dose levels of 0, 190, 600 or 1900 mg/kg-bw. On day 20 of gestation, each female was killed and given a gross pathological examination. The gravid uterus was weighed, its contents were examined and all the fetuses were weighed and examined externally. Of these fetuses, approximately half were given a fresh internal examination, their heads removed and examined by the technique of Wilson. The remaining fetuses were eviscerated. All fetuses were stained with Alizarin Red S and their skeletons examined.

Female Sprague-Dawley rats [Crl:CD (SD) BR] were obtained from Charles River Breeding Laboratories, Kingston, New York. After arrival, animals were examined by a veterinary aide; any animals found in poor condition were rejected from the study. After an acclimation period of 14 days, each female was placed in a cage with a proven male breeder of the same strain and source. On the day of mating (Day 0 of gestation), the females were 80-93 days of age and weighed between 231 and 320 g. Pregnancy was assumed when there was positive identification of spermatozoa in the daily vaginal lavage and this was termed day 0 of gestation. Animals were individually housed except during mating.

MATERNAL IN-LIFE DATA: Animals were checked twice daily for mortality and clinical signs. Pregnant females were examined prior to and following dosing for reactions to treatment, indications of poor health and abnormal

behavior from day 6 to day 15 of gestation. Animals were weighed once each week during the acclimatization period and on days 0, 6, 9, 12, 15, 18 and 20 of gestation. Food intake was assessed for all animals on days 0 to 6, 6 to 9, 9 to 12, 12 to 15, 15 to 18 and 18 to 20 of gestation. On day 20 of gestation, female rats were killed by carbon dioxide asphyxiation followed by exsanguination from the abdominal aorta, each was given a complete gross pathological examination.

MATERNAL EXAMINATION: The reproductive tract of each female was dissected out, the ovaries removed and the corpora lutea counted. The uterus was weighed. The uterine contents were examined and the number and position of live fetuses, dead fetuses, early (endometrial gland with or without some placental tissue), middle (discernible placental and fetal tissue present) and late (fetal structure apparent) resorptions were recorded. The fetuses were then removed from the uterus for examination. The uterus of any animal judged to be nonpregnant was stained with 10% aqueous (v/v) ammonium sulphide solution and was then examined for implantation sites.

FETAL EXAMINATION: Each fetus was weighed, given a detailed external examination with external sex being recorded and then killed. A detailed internal examination using a dissecting microscope was performed on approximately one half of the fetuses, selected randomly from each litter, which were then eviscerated. The heads of these fetuses were removed and placed in Bouin's fluid for examination by the technique of Wilson. The remaining fetuses in each litter were eviscerated; these and the bodies of those fetuses examined internally were placed in 85% ethanol/15% methanol for subsequent staining with Alizarin Red S using a modified Dawson technique for skeletal examination.

Abnormalities were classified as major malformations, minor visceral or skeletal anomalies or common skeletal variants.

STATISTICAL METHODS: The group mean body weights and body weight gains of animals with live fetuses were calculated. The group mean corrected body weights for day 20 of gestation (body weight on day 20 minus gravid uterus weight) and the corrected body weight gains from day 6 to 20 (corrected body weight day 20 minus body weight day 6) were calculated (Data for non-pregnant animals were not included). These parameters were analyzed using one-way analysis of variance, and where the F value was found to be of significance (P < 0.05), intergroup differences between control and treated groups were examined using Dunnett's "t" test.

The group mean live litter size, corpora lutea count, number of implants and number of resorptions were calculated. The individual and group litter mean for the sex ratio and pre- and post-implantation losses were calculated. Statistical analyses were performed using the Kruskal-Wallis test and where the "H" value was significant (P < 0.05) the Mann-Whitney "U" test was used to analyze for differences between control and test groups.

The litter mean fetal weights and group mean fetal weights were calculated and statistical analysis was performed using an analysis of variance (one-way classification) and Dunnett's "t" test.

The incidences of major malformations and minor anomalies were reported as the number of litters with abnormalities in each group and the number of

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fetuses affected. Statistical analyses comparing the number of litters (containing major malformations) in each test group with the control values were performed using either the chi-square test (with Yate's correction factor) or Fisher's exact probability test. The incidence of minor anomalies was analyzed in the same manner. The incidence of common skeletal variants was reported as the litter mean percentage of fetuses affected. Statistical analyses were performed by comparing the litter mean percentage incidences of each test group with the control group using the Kruskal-Wallis and Mann-Whitney "U" tests.

Result

No animals died during the study and no treatment-related clinical signs were reported.

BODY WEIGHT: Between day 6 and day 9 of gestation, the 1,900 mg/kg-day group lost weight while the body weight gains were significantly reduced in the 600-mg/kg-day group. There were significantly reduced body weight gains over the day 9 to 12 interval in the 1,900-mg/kg-day group. These reduced body weight gains resulted in significantly reduced body weights from day 9 to 20 of gestation in both the 600 and 1,900 mg/kg-day groups. The corrected body weights were significantly decreased in the 600 and 1,900 mg/kg-day groups and the corrected body weight gain was decreased significantly in the 1,900-mg/kg-day group. FOOD CONSUMPTION: (Table 5, Appendix 3)

Over days 6 to 9 and 9 to 12 of gestation, food consumption in both the 600 and 1,900-mg/kg-day groups was significantly reduced. Food consumption continued to be significantly reduced over days 12 to 15 of gestation in the 1,900-mg/kg-day group only.

GROSS PATHOLOGICAL FINDINGS: (Table 1, Appendix 6)
Gross pathological examinations revealed no abnormalities related to treatment other than a few incidental findings among mid and low-dose animals on the study.

UTERINE FINDINGS: (Tables 1 and 8, Appendix 7)

The pregnancy rate was at least 88.0% in all groups. Ammonium sulphide staining revealed no other pregnancies.

Gravid uterus weights were significantly reduced in the high-dose group. There were no significant differences between control and treated groups for the following ovarian and uterine parameters: total corpora lutea, total implantation sites, numbers of male and female fetuses, sex, ratio, number of live fetuses, number of dead fetuses, early, middle and late resorptions, total resorptions and pre- and post-implantation losses.

FETAL FINDINGS:

FETAL WEIGHTS were significantly reduced for males, females and totals only in the high-dose group.

MAJOR MALFORMATIONS, In the high-dose group there was a significant increase in the incidence of litters and fetuses with major malformations with 5 fetuses affected. All had acaudia or microcaudia and anal atresia. In addition, one of these fetuses had absence of some thoracic and all lumbar, sacral and caudal vertebrae and absence of 9 pairs of ribs. The incidence of major malformations in the mid and low-dose groups was not different from controls.

MINOR VISCERAL ANOMALIES: There was no effect upon the overall incidence of litters with minor visceral anomalies, but the incidence of fetuses affected was significantly increased in the high-dose group.

MINOR SKELETAL ANOMALIES: The overall incidence of fetuses with minor skeletal anomalies was significantly increased at the high dose. This increase was primarily the result of significantly increased incidences of

several findings which included reduced ossification of frontal bones, irregular ossification of supraoccipital bones, reduced number of pre-sacral vertebrae and ossification centers on the seventh cervical vertebra. In the mid and low-dose groups, statistically significant differences in the incidences of reduced ossification of the interparietal bone, ossification centers on the first lumbar vertebra, reduced ossification of the pubic bones, reduced ossification of the ischial bones or absent ribs were attributed to intergroup variation.

COMMON SKELETAL VARIANTS: The percentage of fetuses with thoracic centrum variants was significantly decreased in the 1900 mg/kg-day group. There was a statistically significant reduction in the percentage of fetuses with sternebral (5 or xiphisternum) variants in the 190-mg/kg-day group that was attributed to intergroup variation.

The accompanying table presents most of the fetal results in tabular form.

2-Pyrrolidone CAS No. 616-45-5, Purity 99.6% Tab-Dev-01.bmp

Test substance Attached document

Dose(mg/kg)	0	190	600	1900
Dams Pregnant	22	25	23	24
Corpora lutea:	17.5	18.3	17.4	17.5
Implantations:	16.3	16.4	16.4	15.5
Postimplantation Loss:	0.7	0.8	1.0	0.8
Live Fetuses/Litter	15.5	15.5	15.4	14.8
Total # Dead Fetuses	0	0	0	0
Total # Live Fetuses:	341	388	355	354
Mean Fetal Weight (g):	3.45	3.54	3.40	3.12
Sex Ratio (male):	0.43	0.46	0.46	0.51
Major Malformations	0	1	1	5*
Litters with Maj Malf	0	1	1	5*
Minor Visceral Malf.	1	2	1	7
Litters with MVM	1	2	1	5
Minor Skeletal Anoml	82	98	60	140**
Litters with MSA	19	23	19	23

Conclusion

Treatment of pregnant rats with 2-pyrrolidone, by gavage, at dosages of up to 1,900 mg/kg-day, throughout major organogenesis, resulted in significant maternal toxicity at the 600 and 1,900 mg/kg-day levels, as evidenced by decreased body weights and food consumption. At the 1,900 mg/kg-day level there were increased incidences of major malformations, minor visceral and skeletal anomalies and decreased fetal weights. No effect upon postimplantation loss was observed.

Therefore, 2-pyrrolidone at a dose of 1,900 mg/kg-day was considered embryo- and fetotoxic but not embryolethal. No effect upon embryonic development was seen at the 600 mg/kg/day level where a significant level of maternal toxicity occurred. The 190 mg/kg/day group was considered the no effect level for maternal toxicity. Based upon these data, the A/D (adult/developmental) ratio was calculated to be <1, indicating 2-pyrrolidone did not show selective toxicity to the rat fetus.

Reliability : (1) valid without restriction

Modern Guideline study under GLP

Flag : Critical study for SIDS endpoint

31.12.2002 (15)

Species : rat

Sex

Strain : Sprague-Dawley

Route of admin. : gavage

Exposure period: days 6-15 of gestation

Frequency of treatm. : daily

Duration of test : 10 days

Doses : 1700 microliters/kg-bw Control group : yes, concurrent no treatment

Result : Not teratogenic in the rat by oral gavage

Method : other: FDA 1966

Year : 1971 GLP : no Test substance :

Method

Test substance was administered in distilled water to 25 presumedpregnant dams on days 6-15 of gestation. Dosing solution was prepared fresh daily. Controls (26 dams) were untreated. Animals were checked daily for adverse clinical signs and mortality. Animals were weighed three times a week during the dosing period. The dose of the test substance was based on the weight of the rat on day 0. The concentration of the solutions was adjusted in such a way that the amount of test substance to be administered for 100 g body weight was contained in a volume of 0.5 ml. On the 20th day of post coitum all the animals were sacrificed, the uteri were removed, the implantation and resorption sites were recorded, the number of live and deed fetuses, their body length, their weight and sex, and the weight of the placentas were determined. The fetuses were examined macroscopically for any malformations. A third of the fetuses of each dam were fixed in Bouin's solution and transversal sections were prepared and assessed according to Wilson's method (Wilson, Warkany: Teratology, Principles und Techniques, 1965). For the assessment of the skeletal system, the remaining fetuses were fixed in 96% strength alcohol, clarified with potassium hydroxide solution and stained with Alizarin red-S using a modified Dawson method. The uteri of the apparently nonpregnant animals or the empty uterine horns in the case of single-horn pregnancy were stained in 10% strength ammonium sulfide solution and then assessed again in order to determine early resorptions.

Remark

The dose level was 1700 microliters/kg-bw. Based on the specific gravity of 1.103, this is approximately 1875 mg/kg-bw.

Without the maternal body-weight gain data the maternal toxicity cannot be adequately assessed. This dose was approximately the same a as that used in the three-dose level 1990 developmental toxicity study and the results are similar in that there was not a major teratogenic effect.

Result

All the pregnant rats tolerated the 10 oral administrations of test material without visible signs of toxicity. One dam died on the 17th day post coitum. The animal proved to be not pregnant. No substance-induced changes could be observed macroscopically. The mean number of implantations and the percentage of resorptions did not differ between the test and control groups. Maternal weights, although recorded, were not included the report.

MACROSCOPIC FETAL EFFECTS: The mean weight and length of the fetuses in the test group did not differ from the values in the control group.

The mean weights of the placentas in the test group and untreated control group were also comparable. The percentage of malformed live fetuses was 2.8 in both groups; similarly, the percentage of runts was the same in the test and control groups.

SKELETAL ASSESSMANT: In treated animals, one fetus (dam No. 6) had a bipartite 12th thoracic vertebral centrum. One fetus (dam No. 10) was observed to have anasarca and two other fetuses of this dam had a cleavage of the eleventh thoracic vertebral centrum. Dam No. 22 had one malformed fetus. The tail of this fetus was missing and atresia was also reported. One fetus of dam No. 24 had a bipartite eleventh thoracic vertebral centrum.

In Untreated animals: One fetus (dam No. 30) had a bipartite eleventh thoracic vertebral centrum. One fetus (dam No. 33) had a bipartite twelfth thoracic vertebral centrum. One fetus of each of dams Nom. 34 and 35 had a bipartite eleventh thoracic vertebral centrum. The presphenoid was missing in one fetus of dam No. 44. One fetus of dam No. 47 had a bipartite 12th thoracic vertebral centrum.

TRANSVERSE SECTIONS: No malformations were found in the fetuses of test or control animals.

Test substance

: 2-Pyrrolidone CAS No. 616-45-5

Conclusion

: The pregnant dams tolerated the 10 oral administrations of test material without any visible symptoms of toxicity or any macroscopically evident pathological changes. The malformations or anomalies found in the fetuses of the test group corresponded in type and number to those of the controls and historical controls. The test material does not have teratogenic effects in Sprague-Dawley rats.

Reliability

: (2) valid with restrictions

A reliability of 2 is assigned. Although some important details are lacking this study was conducted according to a standard procedure that is scientifically defensible. It has value as a supporting study.

08.12.2002

(3)

9. References Id 616-45-5

Date 13.08.2003

(1) All flatworms survived the 96-hour exposure period. (2) BASF AG, Abt. Toxikologie, unpublished study report (86/286), 26.11.1987 BASF AG, Abt. Toxikologie, unveroeffentlichte Untersuchung (XIX/421), 04.08.1971 (3) BASF AG. Abteilung Toxikologie; unpublished report. Cytogenetic Study In Vivo of (4) Pyrrolidon-2 in Mice, Micronucleus test. (92/1491), 28.06.93 (5)BASF AG, Abteilung Toxikologie; unveroeffentliche Untersuchungen (79/409), 09.04.1981 BASF AG, Analytisches Labor; Unpublished Stiudy (J.Nr.129300/04 vom 14.06.88) (6) (7) BASF AG, Labor Oekologie; unveroeffentlichte Untersuchung (Pyrrolidon dest., 1977) BASF AG, Labor Oekologie; unveroeffentlichte Untersuchung, (0701/88) (8) (9)BASF AG, Labor Oekologie; unveroeffentlichte Untersuchung, (0701/88, Fa.Noack) BASF AG, Report of the Subchronic oral toxicity with 2-Pyrrolidone in Wistar rats, 3-month (10)drinking water, Project No. 52S0014/92038 June 4, 1998 BASF AG, Report of the Subchronic oral toxicity with 2-Pyrrolidone in Wistar rats, 3-month (11)drinking water, Project No. 52S0014/92038 June 4, 1998. Dr. med. vet. C. Gembardt, Special Report: "Assessment of the reproductive performance of 2-Pyrrolidone in male and female Wistar rats from data obtained in a subchronic toxicity study" 15 July 2003 (12)BASF AG: Abt. Toxikologie, unpublished report, (92/14), 01.08.1995 BASF AG: Abt. Toxikologie, unveroeffentlichte Untersuchung,(XI/407), 07.11.1961 (13)(14)BASF Labor Okologie, unpublished study, 28.06.88 Bio-Research Laboratories Inc, An Oral Teratoloty Study of 2-Pyrrolidone in the Rat. (15)Project # 83880, Dec. 19, 1990 Sponsored by GAF Chemicals and BASF AG (16)Budavari, S. (ed.). The Merck Index - An Encyclopedia of Chemicals, Drugs, and Biologicals. Whitehouse Station, NJ: Merck and Co., Inc., 1996. 1378 Chem Inspect Test Inst; Biodegradation and Bioaccumulation Data of Existing Chemicals (17)Based on the CSCL Japan; Published by Japan Chemical Industry Ecology-Toxicology & Information Center. ISBN 4-89074-101-1 p. 5-5 (1992) (18)Daubert, T.E. and Danner, R.P. Physical and Thermodynamic Properties of Pure Chemicals: Data Compilation. Design Institute For Physical Property Data, American Institute Of Chemical Engineers. Hemisphere Pub. Corp., New York, NY., 5 Vol, 1997 (19)EPIWIN 3.05 caluclation SRC Syracuse NY (20)Estimated using HYDROWIN 1.67 as found in EPIWIN 3.05, SRC Syracuse NY

ld 616-45-5 9. References

Date 13.08.2003

- (21)Flick, E.W. (ed.). Industrial Solvents Handbook 4 th ed. Noyes Data Corporation., Park Ridge, NJ., 1991. 918, as cited in Hazardous Substance Data Base, NLM, Revison of 8-6-2002
- (22)Jagannath, D.R., Mutagenicity Test on 2-Pyrrolidone in the Ames Salmonella/Microsome Reverse Mutation Assay, Final Report, Hazleton Labs, GAF Sponsor April 24, 1987.
- Mayer, V.W. Goin, C. J. and Taylor-Mayer, R. E. Aneuploidy Induction in Saccharomyces (23)cerevisiae by Two Solvent Compounds, 1-Methyl-2-pyrrolidinone and 2-Pyrrolidinone. Environmental and Molecular Mutagenesis 11:31-40, 1988
- (24)MB Research Laboratories Inc project number MB-92-1432 Sponsored by International Specialty Products, 4/29/1992.
- Perry, C.M., Smith, S.B. Toxicity of Six Heterocyclic Nitrogen Compounds to Daphnia pulex. (25)Bull. Environ. Contam. Toxicol.41, 604-608, (1988)
- Riddick, J.A.; Bunger, W.B.; and Sakano, T.K. Organic Solvents: Physical Properties (26)And Methods Of Purification. Techniques Of Chemistry. 4th Ed. New York, NY: Wiley-Interscience. 2: Pp.1325, 1986 (as cited in CIS 4-2002)
- (27)Submission to U.S. EPA: Raw data for ecotoxicity information on 2-Pyrrolidinone (CAS Reg. No 616-45-5), with cover letter dated 01/29/86 Source: EPA/OTS; Doc #FYI-OTS-0794-1152 Submitted by Eastman Kodak Company