CLH report

Proposal for Harmonised Classification and Labelling

Based on Regulation (EC) No 1272/2008 (CLP Regulation), Annex VI, Part 2

Substance Name: Metaldehyde

EC Number: 203-600-2

CAS Number: 108-62-3

Index Number: 605-005-00-7

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Part A.

1 PROPOSAL FOR HARMONISED CLASSIFICATION AND LABELLING

1.1 Substance

Table 1: Substance identity

Substance name:	Metaldehyde
EC number:	203-600-2
CAS number:	108-62-3
Annex VI Index number:	605-005-00-7
Degree of purity:	985 g/kg
Impurities:	Relevant Impurity: Acetaldehyde max. 1.5 g/kg

1.2 Harmonised classification and labelling proposal

Table 2: The current Annex VI entry and the proposed harmonised classification

	CLP Regulation	Directive 67/548/EEC (Dangerous Substances Directive; DSD)
Current entry in Annex VI, CLP	Flam. Sol. 2, H228	F; R11
Regulation	Acute Tox. 4, H302	Xn; R22
Current proposal for consideration	Flam. Sol. 2, H228	F; R11
by RAC	Acute Tox. 3, H301 STOT RE 2, H373	Xn; R22 - R48/22
	Aquatic Chronic 2, H411	R51/53
Resulting harmonised classification		
(future entry in Annex VI, CLP		
Regulation)		

1.3 Proposed harmonised classification and labelling based on CLP Regulation and/or DSD criteria

 Table 3:
 Proposed classification according to the CLP Regulation

CLP Annex I	Hazard class	Proposed classification	Proposed SCLs and/or M-	Current classification 1)	Reason for no classification 2)
ref		0.44 55222 0.02 52	factors	C10 052110001011	CAU DD 222CW01 042
2.1.		The MSDS submitted indicates under Dust explosion class: St(H)2: strong dust explosion, indicator 2.	-	-	Conclusive, but not sufficient for classification
	Explosives	Classification for dust explosiveness is neither foreseen according to CLP Regulation nor according to Directive 67/548/EEC			
2.2.	Flammable gases	1	ı	1	Conclusive, but not sufficient for classification
2.3.	Flammable aerosols	-	-	-	Conclusive, but not sufficient for classification
2.4.	Oxidising gases	-	-	-	Conclusive, but not sufficient for classification
2.5.	Gases under pressure	-	-	-	Conclusive, but not sufficient for classification
2.6.	Flammable liquids	1	-	-	Conclusive, but not sufficient for classification
2.7.	Flammable solids	Flam. Sol. 2, H228	-	F; R11	
2.8.	Self-reactive substances and mixtures	-	-	-	Conclusive, but not sufficient for classification
2.9.	Pyrophoric liquids	-	-	-	Conclusive, but not sufficient for classification
2.10.	Pyrophoric solids	1	ı	-	Conclusive, but not sufficient for classification
2.11.	Self-heating substances and	-	-	-	Data inconclusive

	mixtures				
	Substances and mixtures which in contact with water emit flammable gases				Conclusive, but not sufficient for classification
	Oxidising liquids				Conclusive, but not sufficient for classification
2.14.	Oxidising solids				Conclusive, but not sufficient for classification
2.15.	Organic peroxides				Conclusive, but not sufficient for classification
2.16.	Substance and mixtures corrosive to metals				Conclusive, but not sufficient for classification
3.1.	Acute toxicity - oral	H301	-	H302	-
	Acute toxicity - dermal	-	-	-	Conclusive but not sufficient for classification
	Acute toxicity - inhalation	-	-	-	Conclusive but not sufficient for classification
3.2.	Skin corrosion / irritation	-	-	-	Conclusive but not sufficient for classification
3.3.	Serious eye damage / eye irritation	-	-	-	Conclusive but not sufficient for classification
3.4.	Respiratory sensitisation	-	-	-	Data lacking
3.4.	Skin sensitisation	-	-	-	Conclusive but not sufficient for classification
3.5.	Germ cell mutagenicity	-	-	-	Conclusive but not sufficient for classification
3.6.	Carcinogenicity	-	-	-	Conclusive but not sufficient for classification
3.7.	Reproductive toxicity	-	-	-	Conclusive but not sufficient for classification
3.8.	Specific target organ toxicity –single exposure	-	-	-	Conclusive but not sufficient for classification
3.9.	Specific target organ toxicity – repeated exposure	H373	-	-	-
3.10.	Aspiration hazard	-	-	-	Data lacking
4.1.	Hazardous to the aquatic environment	H411	-	-	

5.1.	Hazardous to the ozone layer				Data lacking
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¹⁾ Including specific concentration limits (SCLs) and M-factors

Labelling: Signal word:

Danger

Hazard statements:

Flam. Sol. 2, **H228**: Flammable Solid Acute Tox. 3, **H301**: Toxic if swallowed

STOT RE 2, H373: May cause damage to organs through prolonged or repeated exposure if

swallowed

Aquatic Chronic 2, H411: Toxic to aquatic life with long lasting effects

Precautionary statements: (resulting from hazard statements according to Annex I to

Regulation (EC) No. 1272/2008 without any further selection)

Prevention:

P210: Keep away from heat/sparks/open flames/hot surfaces. – No smoking.

P240: Ground/bond container and receiving equipment.

P241: Use explosion proof electrical/ventilating/lighting/.../equipment.

P280: Wear protective gloves/protective clothing/eye protection/face protection.

P264: Wash ... thoroughly after handling.

P270: Do not eat, drink or smoke when using this product.

P260: Do not breathe dust/fume/gas/mist/vapours/ spray.

Response:

P370+P378: In case of fire: Use water spray, dry powder, foam for extinction.

P301+P310: IF SWALLOWED: Immediately call a POISON CENTER or a

doctor/physician.

P321: Specific treatment (see ... on this label).

P330: Rinse mouth.

P314: Get medical advice/attention if you feel unwell.

Proposed notes assigned to an entry: -

²⁾ Data lacking, inconclusive, or conclusive but not sufficient for classification

Table 4: Proposed classification according to DSD

Hazardous property	Proposed classification	Proposed SCLs	Current classification 1)	Reason for no classification
Explosiveness	No classification	-	No classification	Conclusive, but not sufficient for classification
Oxidising properties	No classification	-	No classification	Conclusive, but not sufficient for classification
Flammability	F; R11	-	F; R11	-
Other physico-chemical properties [Add rows when relevant]	No classification	-	No classification	-
Thermal stability	No classification	-	No classification	Data lacking
Acute toxicity	R22	-	R22	-
Acute toxicity – irreversible damage after single exposure	-	-	-	Conclusive but not sufficient for classification
Repeated dose toxicity	R48/22	-	-	-
Irritation / Corrosion	-	-	-	Conclusive but not sufficient for classification
Sensitisation	-	-	-	Conclusive but not sufficient for classification
Carcinogenicity	-	-	-	Conclusive but not sufficient for classification
Mutagenicity – Genetic toxicity	-	-	-	Conclusive but not sufficient for classification
Toxicity to reproduction – fertility	-	-	-	Conclusive but not sufficient for classification
Toxicity to reproduction – development	-	-	-	Conclusive but not sufficient for classification
Toxicity to reproduction – breastfed babies. Effects on or via lactation	-	-	-	Conclusive but not sufficient for classification
Environment 1) Including SCLs	R51/53			

1) Including SCLs

Labelling: Indication of danger:

F Highly flammable

Xn Harmful

R-phrases:

R11 Highly flammable.

R22 Harmful if swallowed.

R48/22 Harmful: danger of serious damage to health by prolonged exposure if swallowed.

R51/53 Very toxic to aquatic organisms, may cause long term effects in the environment

²⁾ Data lacking, inconclusive, or conclusive but not sufficient for classification

S-phrases:

- S2 Keep out of the reach of children.
- S7 Keep container tightly closed.
- S13 Keep away from food, drink and animal feedingstuffs.
- S16 Keep away from sources of ignition No smoking.
- S20/21 When using do not eat, drink or smoke.
- S36/37 Wear suitable protective clothing and gloves.
- S46 If swallowed, seek medical advice immediately and show this container or label.

2 BACKGROUND TO THE CLH PROPOSAL

2.1 History of the previous classification and labelling

Metaldehyde is a molluscicide for the control of slugs and snails, and was approved in 2008 for Annex I listing as a 3A Review compound under Council Directive 91/414/EEC, with Austria as Rapporteur Member State. In accordance with Article 36(2) of the CLP Regulation, metaldehyde should now be considered for harmonised classification and labelling. Therefore, this proposal considers all physico-chemical, human health and environmental end points. This Annex VI dossier presents a classification and labelling proposal based mainly on the information presented in the assessment of metaldehyde under Directive 91/414/EEC. The assessment made under that Directive is attached to the IUCLID 5 dossier.

Metaldehyde is already listed in Annex VI of the CLP Regulation (it was inserted into Annex I of Directive 67/548/EEC in the 19th ATP and updated with the 31st ATP [Tox data were not discussed for update with 31st ATP]) with the classifications as F; R11 and Xn; R22.

This proposal seeks to update these classification and additionally, to include classification for repeated dose toxicity. During the peer review for Annex I Inclusion of metaldehyde Member States and EFSA agreed that Austria should flag the new proposal for classification and labelling to ECHA, including repeated dose toxicity.

No REACH registration dossiers were available, when resubmitting the CLH-dossier.

2.2 Short summary of the scientific justification for the CLH proposal

Current classification according to Annex VI, Table 3.1 in the CLP Regulation for Metaldehyde is Flam. Sol. 2, H228.

The lowest LD₅₀ value of 283 mg/kg for acute oral toxicity was found in rats. According to Regulation (EC) No. 1272/2008 metaldehyde belongs to acute toxicity category 3 (50 < ATE \leq 300 mg/kg bw) and requires classification and labelling with **H301 "Toxic if swallowed"**.

Metaldehyde requires classification with H373 "May cause damage to organs through prolonged or repeated exposure (if swallowed)" based on the findings of mortality at 30 mg/kg bw/d and testicular findings (moderate to marked diffuse atrophy and/or degeneration of the germinative epithelium) at 90 mg/kg bw/d in a 52-week dog study. Criteria as specified in the Regulation (EC) No. 1272/2008: For the oral route the guidance values to assist in Category 2 classification are $10 < \text{dose} \le 100 \text{ mg/kg}$ bodyweight/day. These guidance values refer to effects seen in a standard 90-day toxicity study conducted in rats.

Regarding environment (considering 2nd ATP criteria) following classification will be proposed:

DSD: N, R51/53 (DSD)

CLP: Aquatic Chronic 2, H411

Aquatic Acute classification is based on:

• LC50 value for *Pomacea canaliculata* = >3.33 mg/L (Scholtz 2004), resulting in N, R51 (DSD) and no classification (CLP)

Aquatic chronic classification is based on:

- Metaldehyde is not considered as ready biodegradable/rapid degradable. Therefore a <u>R53 (DSD)</u> classification is proposed.
- chronic aquatic toxicity studies
 Based on the non rapid degradability and on the toxicity to *Pomacea canaliculata*(Scholtz 2004) with a NOEC= 0.5 mg/L. a classification with <u>Aquatic Chronic 2</u>,
 H411 (CLP) is proposed.

2.3 Current harmonised classification and labelling

2.3.1 Current classification and labelling in Annex VI, Table 3.1 in the CLP Regulation

Classification:

Flam. Sol. 2, H228 Acute Tox. 4, H302

Labelling:

pictograms: GHS02, GHS07

signal word: Dgr

hazard statement codes: H228, H302

2.3.2 Current classification and labelling in Annex VI, Table 3.2 in the CLP Regulation

Classification:

F; R11 Xn; R22

Labelling:

F; Xn

R: 11-22

S: (2-)13-16-25-46

2.4 Current self-classification and labelling

2.4.1 Current self-classification and labelling based on the CLP Regulation criteria

No current self-classification and labelling based on CLP Regulation criteria.

2.4.2 Current self-classification and labelling based on DSD criteria

Classification and labelling as proposed in the Draft Assessment Report for metaldehyde, Volume 1, Level 2, Section 2.5.1 (August 2010):

On the basis of the available data the following classification and labelling is proposed according to Directive 67/548/EEC in combination with Directive 93/21/EEC:

Hazard symbols:		Black St. Andrew's Cross on orange square Flame on orange square
Indication of	Xn	Harmful

danger:	F	Highly flammable		
Risk phrases: R11		Highly flammable		
	R22	Harmful if swallowed.		
	R48/22	Harmful: danger of serious damage to health by prolonged exposure if swallowed.		
	R52/53	Harmful to aquatic organisms, may cause long-term adverse effects in the aquatic environment.		
Safety phrases:	S2	Keep out of the reach of children.		
	S7	Keep container tightly closed.		
	S13	Keep away from food, drink and animal feedingstuffs.		
	S16	Keep away from sources of ignition - No smoking.		
	S20/21	When using do not eat, drink or smoke.		
	S36/37	Wear suitable protective clothing and gloves.		
	S46	If swallowed, seek medical advice immediately and show this container or label.		

Justification for the proposal:

Physical and chemical properties: The classification and labelling with F follows the risk phrase R11. R11 is set as the propagated combustion over the 100 mm is < 45 seconds under test condition. (Tremain, S. P., (2001) Doc. No. 119-002)

Human health effects: The classification and labelling with Xn follows from risk phrases R22 and R48/22. R22 was assigned due to the acute oral toxicity observed in rats ($LD_{50} = 283 \text{ mg/kg}$, Jones J. and Collier T, 1987; $LD_{50} = 654 \text{ mg/kg}$, Durando J., 2009) and mice ($LD_{50} = 411-443 \text{ mg/kg}$; Coles R., 1990a). R 48/22 is required as mortality as well as atrophy of the germinal epithelium of the testes occurred at the dose of 30 mg/kg in a 52-week dog study (Leuschner J., 2003; Leuschner J. and Drommer W., 2006). S2, S7, S13, S16, S20/21 follow from a general precautionary approach. S36/37 follows from the risk assessment of operator exposure, while S46 follows from the acute oral toxicity of metaldehyde.

Effects on aquatic organisms: R52/53 is proposed because the lowest acute LC_{50} for fish is 75 mg/L (Bogers, 1990a) and metaldehyde is not readily biodegradable (Wuethrich, 1990a). S56, S57, S60 and S61 follow from a general precautionary approach.

3 JUSTIFICATION THAT ACTION IS NEEDED AT COMMUNITY LEVEL

Metaldehyde is used as a pesticide. For pesticides there is no need for justification (cf. Article 36(3) CLP Regulation).

Part B.

SCIENTIFIC EVALUATION OF THE DATA

1 IDENTITY OF THE SUBSTANCE

1.1 Name and other identifiers of the substance

Table 5: Substance identity

EC number:	203-600-2
EC name:	Metaldehyde 2,4,6,8-tetramethyl-1,3,5,7- tetraoxacyclooctane
CAS number (EC inventory):	203-600-2
CAS number:	108-62-3
CAS name:	2,4,6,8-tetramethyl-1,3,5,7-tetraoxacyclooctane 1,3,5,7-Tetroxocane, 2,4,6,8-tetramethyl-
IUPAC name:	r-2, c-4, c-6, c-8-tetramethyl-1,3,5,7-tetroxane
CLP Annex VI Index number:	605-005-00-7
Molecular formula:	$C_8H_{16}O_4$
Molecular weight range:	176.2 g/mol

Structural formula:

1.2 <u>Composition of the substance</u>

Table 6: Constituents (non-confidential information)

Con	stituent	Typical concentration	Concentration range	Remarks
Meta	aldehyde	Min. purity: 985 g/kg	983 – 998 g/kg	-

Current Annex VI entry:

Table 3.1

List of harmonised classification and labelling of hazardous substances

Index No	International Chemical Identification	EC No CAS No					Labelling		Specific	Notes
			No	Hazard Class and Category Code(s)	Hazard stateme nt Code(s)	Pictogram, Signal Word Code(s)	Hazard statement Code(s)	Suppl. Hazard statement Code(s)	Conc. Limits, M-factors	
605-005-00-	2,4,6,8-tetramethyl- 1,3,5,7- tetraoxacyclooctane; metaldehyde	203- 600-2	108- 62-3	Flam. Sol. 2 Acute Tox. 4 *	H228 H302	GHS02 GHS07 Wng	H228 H302			

Table 7: Impurities (non-confidential information)

Impurity	Typical concentration	Concentration range	Remarks
Acetaldehyde (relevant impurity)	Max. content: 1.5 g/kg	-	-

Current Annex VI entry:

Table 3.1

List of harmonised classification and labelling of hazardous substances

Index No	International Chemical Identification	EC No CAS No		Classification		Labelling		Specific	Notes	
			Hazard Class and Category Code(s)	Hazard stateme nt Code(s)	Pictogram, Signal Word Code(s)	Hazard statement Code(s)	Suppl. Hazard statement Code(s)	Conc. Limits, M-factors		
605-003-00-	acetaldehyde; ethanal	200- 836-8	75-07- 0	Flam. Liq. 1 Carc. 2 Eye Irrit. 2 STOT SE 3	H224 H351 H319 H335	GHS02 GHS08 GHS07 Dgr	H224 H351 H319 H335			

Table 8: Additives (non-confidential information)

Additive	Function	Typical concentration	Concentration range	Remarks
Confidential	Stabilizer	confidential	-	-

Current Annex VI entry: no entry

1.2.1 Composition of test material

Physico-chemical properties: purity of tested technical material in the range from 99.1% to 99.5%)

<u>Human health hazard assessment:</u> purity of tested technical material in the range from 90.14% to 94.8%

 $\underline{\text{Environmental hazard assessment:}}$ purity of tested technical material in the range from 90.2% to 98.7%

1.3 <u>Physico-chemical properties</u>

Table 9: Summary of physico - chemical properties

Study	Method	Results	Conclusion/Comment	Reference
B.2.1.1 Melting point, freezing point or solidification point (IIA 2.1.1)	EEC/A1 and OECD 102 Metal block and Differential scanning calorimetry method (DSC)) GLP	TGAI purity: 99.5 % (w/w) Two methods according OECD 102 were used in this testing: Metal block method and DSC. Metal block method determination: Heating rate: 1 °C/min -sealed tube In the metal block method the samples (in duplicate) was placed in a sealed tube. The samples went straight to stage B (i.e. a clearance between the sample and the wall due to shrinkage of the melt) as described in method OECD 102 (figure 2) at a temperature of 191 °C. The remaining stages of melting were not observed. Complete sublimation was achieved at 201 °C and at 197 °C respectively. Differential scanning calorimetry determination: Heating rate: 10 °C/min range: 25 °C to 400 °C in atmosphere of air. Tests were performed in open and in sealed crucibles. In each case a sharp endotherm peak with a maximum between 200 and 210 °C was detected. The extrapolated onset temperatures for the sharp endotherm peak were 182 °C for the open and 185 °C for the sealed crucible tests. The endotherms started with broad features with onsets at approx. 100 °C which indicates the initiation of changes in the test substance at this temperature. After each test the crucible was empty. Metaldehyde starts to sublime at 191 °C.	Acceptable The use of technical material is acceptable since purity is > 98 %(w/w)	Comb, A. L. (2007) (Doc. No. 112-002)

Study	Method	Results	Conclusion/Comment	Reference
B.2.1.2 Boiling point (IIA 2.1.2)	Statement Tier 2	Not applicable, as the test item is a solid	Not applicable as sublimation is more realistic	
B.2.1.3 Temperature of decomposition or sublimation (IIA 2.1.3)	EEC/A1 and OECD 102 Metal block and Differential scanning calorimetry method (DSC)) GLP	TGAI purity: 99.5 % (w/w) Metaldehyde starts to sublime at 191 °C.	Acceptable The use of technical material is acceptable since purity is > 98 %(w/w)	Comb, A. L. (2007) (Doc. No. 112-002)
B.2.1.4 Relative density (IIA 2.2)	OECD 109 (Gas comparison pycnometer) GLP	Purified product purity: 99.1 % (w/w) Density: $1.27 \times 10^3 \text{ kg/m}^3 (20 \pm 0.5 \text{ °C})$	Acceptable Method is equivalent to EEC/A3 Relative density is not reported	Hogg, A.S.; (1998) (Doc. No. 112-001)
B.2.1.5 Vapour pressure (IIA 2.3.1)	OECD 104 (Static method) not GLP*	Purified product purity: 99.3 % (w/w) Vapour pressure: 6.6 ± 0.3 Pa (25 °C) 4.4 ± 0.2 Pa (20 °C)	Acceptable	Cardinaals, J.M. (1988) (Doc. No.115-001)
B.2.1.6 Volatility, Henry's law constant (IIA 2.3.2)	Calculation	3.5 Pa.m³.mol ⁻¹ (20 °C) <u>values used for calculation:</u> water solubility: 0.222 g/L at 20 °C vapour pressure: 4.4 Pa at 20 °C	Acceptable	Cardinaals, J.M. (1988) (Doc. No.115-002)

Study	Method	Results	Conclusion/Comment	Reference
B.2.1.7 B.2.1.8 Appearance: physical state (IIA 2.4.1)	Visual examination	TGAI purity: 99.5 % (w/w) White crystalline powder	Acceptable The use of technical material is acceptable since the purity is > 98%.	Comb, A. L. (2007) (Doc. No. 112-002)
		Technical product purity: 99.3% (w/w) White powder	Acceptable	O'Connor, B. J., Mullee, D. M. (2000) (Doc. No. 172-001)
B.2.1.10 Spectra of the active substance (IIA 2.5.1)	UV/VIS - Spectroscopy OECD guideline No.101 GLP	Purified product purity: 99.1 % (w/w) $c = 1.02 \times 10^{-3} \text{mol/L} (0.18 \text{g/L})$ Solvent MeOH/HCl no significant absorption occurs at any wavelength. v/v]	Acceptable	O'Connor, B.J., Mullee, D.M., (2001) (Doc. No. 119-001)
		MeOH no significant absorption occurs at any wavelength. MeOH/NaOH no significant absorption occurs at any [90/10 (0.1 N) wavelength. v/v]		

Study	Method	Results	Conclusion/Comment	Reference
	¹ H and ¹³ C-NMR FTIR (KBr, 4000 - 600 cm ⁻¹) MS (EI 70eV)	UV/VIS, IR, NMR and MS spectra including interpretation data were submitted. The spectra confirm the molecular structure. Optical purity: not relevant as Metaldehyde has no optical isomers.	Acceptable	
B.2.1.11 Spectra of relevant impurities (IIA 2.5.2)	OECD 101 GLP except NMR spectrum	Purified product purity: 99.5% (w/w) Acetaldehyde is considered to be a relevant impurity. Ultraviolet/visible, infrared, nuclear magnetic resonance and mass spectra were recorded and found to be consistent with the assigned structure of the molecule.	Acceptable Although the NMR spectrum was not performed according to GLP because the laboratory (London Metropolitan University) is not a member of the UK GLP compliance programme the structure of acetaldehyde is confirmed by IR, MS and UV.	Comb, A. L. (2009) (Doc. No. 157-001)
B.2.1.12 Solubility in water (IIA 2.6)	EEC A.6 OECD 105 (Flask shaking method) not GLP*	$\begin{array}{llllllllllllllllllllllllllllllllllll$	Acceptable	Bohle, J.F., (1989) (Doc. No. 114-001)
	OECD 105 (Flask shaking method) not GLP*	Purified product purity: 99.3 % (w/w) 19.9 - 23.0 °C 0.222 g/L; pH 6.5; Milli-Q-water (Millipore)	Acceptable OECD 105 is comparable to EEC/A6	Cardinaals, J.M. (1988b) (Doc. No. 114-003)

Study	Method	Results			Conclusion/Comment	Reference
B.2.1.13 Solubility in	EEC A.6 OECD 105	Purified product purity:	>99.5% (w/w)		Acceptable	Bohle, J.F., (1989)
organic	(Flask shaking method)	solvent	solubility at 20.3-22.4 °C [g/L]			(Doc. No.
solvents (IIA 2.7)	not GLP*	hexane methanol	1.73	10 ⁻³		114-004) Comb, A. L.
		toluene tetrahydrofurane 1,2-dichloroethane	0.53 1.56 3.08			(2007) (Doc. No.
		acetone	1.46			114-006)
	EEC A.6 (Flask shaking method) GLP	Purified product purity:	99.5% (w/w)		Acceptable	O'Connor, B.J., Mullee,
		ethyl aceto acetate	solubility at 20.0 ± 0.5 °C [g/L] 0.754			D.M., (2001a) (Doc. No. 114-005)
B.2.1.14 Partition coefficient n-	OECD 107 (Shake flask method) not GLP*	$\log P_{\text{ow}} = 0.12 \ P_{\text{ow}} = 1.33 \pm 0$	•	6.7	Acceptable The method is comparable to the EEC/A8 shake flask	Cardinaals, J.M. (1988b) (Doc. No.
octanol/water (IIA 2.8)		Effect of pH (4 to 10) is relative an acid nor a base.	not required, because Metaldehydo	e is	method	114-002)
		Because metaldehyde is no phase is not buffered	ot an ionisable compound the w	ater		

Study	Method	Results	Conclusion/Comment	Reference
B.2.1.15 Hydrolysis rate (IIA 2.9.1)	US EPA-FIFRA N- 161-1 40 CFR 158.130 not GLP*	 14C-Metaldehyde radiochemical purity: > 99% Study performed at 25 ± 1 °C for 30 days in the dark (in pH 5, 7 and 9 buffers) Metaldehyde is hydrolytically stable, no accurate half-life could be determined due to insignificant degradation 	Acceptable For details see B 8.4 Fate and behaviour in water	Carpenter, M., (1989) (Doc. No. 711-001)
B.2.1.16 Direct phototrans- formation (IIA 2.9.2)	US EPA-FIFRA N- 161-1 40 CFR 158.130 not GLP*	 14C-Metaldehyde radiochemical purity: > 97.7% Study performed at 25 ± 1°C for 30 days (in pH 7 buffer) On day 30, 97.5% of the initially administered parent compound was still found. Metaldehyde is photolytically stable, no accurate half-life could be determined due to insignificant degradation 	Acceptable For details see B 8.4 Fate and behaviour in water	Carpenter, M., (1989a) (Doc. No. 712-001)
B.2.1.17 Quantum yield (IIA 2.9.3)	FAO revised guidelines on Environmental Criteria for the Registra-tion of Pesticides SETAC 1995	$^{14}\text{C-Metaldehyde radiochemical purity:} > 97.7\%$ Quantum yield $\Phi \approx 0$ Test substance is stable in the absence of a photo-sensitiser, therefore no separate determination is necessary	Acceptable For details see B 8.4 Fate and behaviour in water	Carpenter, M., (1989a) (Doc. No. 712-001)
B.2.1.18 Dissociation constant (pKa) (IIA 2.9.4)			Not relevant as Metaldehyde does not dissociate in water	

Study	Method	Results	Conclusion/Comment	Reference
B.2.1.19 Stability in air, photochemical oxidative degradation (IIA 2.10)	Atkinson calculation	$K_{OH} = 73.8 \times 10^{-12} \text{ cm}^3 \text{ x molecule}^{-1} \text{ x sec}^{-1}$ half-life in troposphere: $t\frac{1}{2}$: 5.3 hours	Acceptable For details see B 8.7.1 Fate and behaviour in air	Voget, M., (1994) (Doc. No. 782-001)
B.2.1.20 Flammability (IIA 2.11)	EEC/A10 GLP	Technical product purity: 99.5% (w/w) As the preliminary test was positive, the main test according to EEC/A10 was performed with the result, that Metaldehyde is highly flammable (the propagated combustion over the 100 mm was < 45 seconds)	Acceptable The study was performed with the TGAI of high purity. R11 for classification is required	Tremain, S.P., (2001) (Doc. No. 119-002)
B.2.1.21 Auto- flammability (IIA 2.11.2)	EEC/A16 GLP	Technical product purity: 99.5% (w/w) No self ignition up to 400 °C	Acceptable The study was performed with the TGAI of high purity.	Tremain, S.P., (2001) (Doc. No. 119-002)
B.2.1.22 Flash point (IIA 2.12)			Not applicable because material is a solid with a melting point > 40°C	

Study	Method	Results	Conclusion/Comment	Reference
B.2.1.23 Explosive properties (IIA 2.13)	EEC/A14 GLP	Technical product purity: 99.5% (w/w) Thermal sensitivity test: no explosion after 5 minutes (nozzle diameter: 2.0 mm and 6.0 mm) Shock test: no explosion occurred within 6 tests using a mass of 10 kg from a height of 0.4 m Friction test: no explosion occurred within 6 tests using a 36 kgf (≈360 N) loading. A black mark on the porcelain plate and peg indicates decomposition	Acceptable The study was performed with the TGAI of high purity. Although dust explosion does not cover this annex point, the MSDS indicates that metaldehyde is classified as St(H)2: strong dust explosion, indicator 2	Tremain, S.P., (2001) (Doc. No. 119-002) Anonymous, (2001) (Doc. No. 955-004)
B.2.1.24 Surface tension (IIA 2.14)	EEC/A5 Ring method GLP	Purified product purity: 99.5% (w/w) $\sigma = 71.9 \text{ mN/m at } 19.5 \text{ °C} \pm 0.5 \text{ °C}$ (0.204 g/L aqueous solution)	Acceptable The compound is not considered as surface active	O'Connor, B.J., Mullee, D.M., (2001) (Doc. No. 119-001)
B.2.1.25 Oxidising properties (IIA 2.15)	Statement	Oxidizing properties are not expected, considering the overall chemical structure and the oxygen balance of Metaldehyde.	Acceptable	Weiss, A. (2009) (Doc. No. 143-001)

^{*} not GLP: as no certificate of the relevant GLP authority is attached, but due to the date of study, not necessary.

2 MANUFACTURE AND USES

2.1 Manufacture

Not relevant for Classification and Labelling.

2.2 Identified uses

Metaldehyde is a molluscicide for the control of slugs and snails.

3 CLASSIFICATION FOR PHYSICO-CHEMICAL PROPERTIES

Table 10: Summary table for relevant physico-chemical studies (DSD)

Method	Results	Remarks	Reference
EEC/A10	highly flammable (the propagated combustion over the 100 mm was < 45 seconds)	-	Tremain, S.P., (2001) (Doc. No. 119-002)

3.1

3.1.1 Summary and discussion of physico-chemical properties

Current classification according to Annex VI, Table 3.1 in the CLP Regulation for Metaldehyde is Flam. Sol. 2, H228.

3.1.2 Comparison with criteria

Not relevant since no test/study is reported in Annex VI, Table 3.1 to be compared with method EEC/A10.

3.1.3 Conclusions on classification and labelling

Current classification according to Annex VI, Table 3.1 in the CLP Regulation for Metaldehyde is Flam. Sol. 2, H228.

4 HUMAN HEALTH HAZARD ASSESSMENT

4.1 Toxicokinetics (absorption, metabolism, distribution and elimination)

4.1.1 Non-human information

One single study on ADME of metaldehyde was conducted in Sprague-Dawley CD rats and included single oral dosing with 10 and 100 mg/kg and repeated oral dosing with 10 mg/kg. In this study, absorption, distribution, excretion as well as metabolism were investigated.

<u>Absorption</u>: Metaldehyde was absorbed rapidly and practically completely in male and female rats after single low and high dose (10 and 100 mg/kg) and repeated oral low doses (10 mg/kg). Maximum concentrations in blood were found within 1-2 h in males and 2-4 h in females, with calculated half-lives of 3.4 h for males and 8.8 h for females indicating a faster metabolism and excretion in males than in females. Oral absorption was essentially complete as the majority of radioactivity (approximately 80% after 48 h and 85% after 7 days) was expired as CO₂. 10% of the administered radioactivity was still found in the body after 7 days, and 2-5% of the dose was found in urine.

<u>Distribution</u>: Following single or repeated oral dosing, radioactivity was widely distributed into the whole body. The majority of radioactivity found in the body was detected in the carcass (approximately 80%) and the rest was distributed more or less evenly in tissues and organs investigated. Small amount of metaldehyde (0.5% males and 0.59% females) were found in the brain, indicating that metaldehyde can pass the blood-brain barrier. There was no indication of bioaccumulation as there was no increase of tissue residue levels after repeated administration compared with tissue levels after single application. The amount of radioactivity found in the whole body markedly decreased over time as demonstrated by the amounts found at peak blood level, first half-life, second half-life and 7 days after dosing.

Excretion: Most of the administered radioactivity was found in the expired air, regardless of single or repeated dosing: approximately 80% after 48 h and 85% after 7 days. The amounts found in urine (2-5%) and faeces (2-3%) were comparably low and independent from the dosing scheme. Most of the radioactivity was excreted during the first 24- 48 h after application, however, males expired the radioactivity more rapidly than females during this time period.

Metabolism: Analysis of blood samples collected at different time points showed that metaldehyde is metabolised to acetaldehyde. No other metabolites were found in the blood. When urine samples were analysed, ¹⁴C residues appeared as multiple peaks with retention times associated with very polar compounds. No intact metaldehyde was detected in urine. The analysis of trapping solutions of the expired air showed that almost all of the radioactivity was expired as CO₂. Taken together it can be assumed that metaldehyde is extensively metabolised to acetaldehyde. The metabolic profile of acetaldehyde is well established and involves conversion of acetaldehyde to acetyl-CoA. The physiological molecule acetyl-CoA is oxidised through the Krebs cycle to CO₂ or utilised in the various anabolic reactions involved in the synthesis of cholesterol, fatty acids or other tissues constituents. The oxidation reactions account for the large amount of ¹⁴C-labelled CO₂ found in the expired air while the anabolic reactions account for the slow decline of ¹⁴C residues observed in the carcass.

4.1.2 Human information

Not available.

4.1.3 Summary and discussion on toxicokinetics

Absorption, distribution, excretion and metabolism (toxicokinetics)

Rapid and essentially complete (>95%) based on Rate and extent of oral absorption excretion via air (80% within 48h; 85% within 7 days) and urinary excretion (2-5% within 7 days). Further 8-10% was still present in the body after 7 days. Distribution Widely distributed; most of radioactivity found in carcass and not in specific organs Potential for accumulation No evidence of accumulation Rate and extent of excretion After 7 days: expired air (85%); urine (2-5%); faeces (2-Metabolism in animals 85% metabolised to acetaldehyde and expired as CO₂; 2-5% metabolised and excreted via polar degradates in the urine (no parent compound in urine); 2-3% in faeces not identified.

4.2 Acute toxicity

Table 11: Summary table of relevant acute toxicity studies

Method	Results	Remarks	Reference
Acute oral toxicity in the rat (OECD guideline 401)	$LD_{50} = 283 \text{ mg/kg bw (M+F)}$ R22	-	Jones, J., Collier, T.; 1987
Acute oral toxicity in the rat, up- and-down procedure (OECD guideline 425)	LD ₅₀ = 654 mg/kg bw (F) R22	This study was not requested by the RMS or any other MS.	Durando, J.; 2009
Acute oral toxicity in the mouse (OECD guideline 401)	LD ₅₀ = 411 mg/kg bw (M) LD ₅₀ = 443 mg/kg bw (F) R22	-	Coles, R.; 1990
Acute percutaneous toxicity in rats	LD ₅₀ > 5000 mg/kg bw	Limited validity	Davies, R., Collins, C.; 1974

Several information regarding acute toxicity can be found in public literature e.g. http://toxnet.nlm.nih.gov/cgi-bin/sis/search/a?dbs+hsdb:@term+@DOCNO+1735 or http://www.inchem.org/documents/pims/chemical/pim332.htm. However the dossier submitter does not have access to the original studies. Therefore no evaluation on reliability of the respective information could be performed and this information has not been included into the CLH-report of metaldehyde.

4.2.1 Non-human information

4.2.1.1 Acute toxicity: oral

Reference: P0071: OECD 401 Acute oral toxicity in the rat, Project Nun		
	102/9A	
Author(s), year:	Jones J., Collier T., 1987	
Report/Doc. number:	Doc.No. 521-002, Lonza Report No. 1354, Conducting laboratory:	
	Safepharm Laboratories Limited, Derby, UK	
Guideline(s):	OECD Guideline 401 (1981)	
GLP:	Yes	
Deviations:	No	
Validity:	Valid	

Material and Methods:

Groups of 5 rats/sex received single doses of 0 (vehicle control), 100, 200, 400 and 800 mg/kg bw metaldehyde (sponsor's identification: P0071; batch no. 3157; purity 99.3 %) suspended in arachis oil by oral gavage. This dose selection was based upon a range-finding study where one male and one female rat per test group were dosed with 100, 250, 500, 1000, 2000 and 5000 mg/kg bw. At the beginning of the study the rats (strain: Sprague-Dawley CFY; source: Interfauna Ltd., UK) were approximately five to eight weeks old and weighed 126-148 g (males) and 114-148 g (females).

Animals were observed immediately after dosing and at 6, 12 and 18 h after dosing and subsequently once daily for 14 days. Death and evidence of clinical signs of toxicity were recorded

at each observation. Bodyweights were determined on the day of treatment (day 0), days 7 and 14, and at death. All animals were subjected to gross necropsy examination for any macroscopic abnormalities. No tissues were retained.

Findings:

Range finding study: No deaths were observed at 100 and 250 mg/kg, while 2 deaths were noted at doses of 500 mg/kg and above. Therefore mortality data indicated an oral LD $_{50}$ between 250 and 500 mg/kg.

Main study: The observed death rates were 0/5 for males and females at 100 mg/kg, 2/5 for males and 1/5 for females at 200 mg/kg, 3/5 for males and 4/5 for females at 400 mg/kg, and 5/5 for both males and females at 800 mg/kg. Two males treated with 800 mg/kg and one males treated with 400 mg/kg were found dead immediately after dosing. All other deaths were noted six or twelve hours after dosing. The acute oral median lethal dose (LD₅₀) was calculated to be 283 mg/kg bodyweight. Clinical signs were observed at all dose levels immediately after dosing. Principal signs of toxicity noted in both decedents and surviving animals were hunched posture, pilo-erection, lethargy and decreased respiratory rate. Occasional or isolated signs were increased salivation, ptosis, (occasional) body tremors, red/brown staining around the eyes, snout and mouth with diuresis, diarrhoea, tonic convulsions, ataxia and coma. Surviving animals showed no signs 2-8 days after dosing. All surviving animals had expected gains in bodyweight over the study period. Common findings noted in decedents were red or haemorrhaged lungs, dark or patchy pallor of the liver and congestion of the small intestine. Sloughing of the gastric mucosa was also noted. No abnormalities were noted at necropsy of animals killed at the end of the study period.

Conclusion:

Metaldehyde (suspended in arachis oil) is of moderate toxicity to rats after oral administration. The acute oral LD_{50} was calculated to be 283 (210 – 382) mg/kg bw.

Reference: Acute Oral Toxicity Study with META Metaldehyde techn. CAS No.

108-62-3: Up-And-Down Procedure in Rats

Author(s), year: Durando J., 2009

Report/Doc. number: Eurofins/Product Safety Laboratories, USA; Laboratory Identification No.

26776

Lonza Report No. 4377, Doc. No.: 521-003

Guideline(s): US EPA, OPPTS 870.1100

OECD Guidelines for the Testing of Chemicals, Test No. 425

GLP: Yes
Deviations: No
Validity: Valid

Material and methods:

Test material META Metaldehyde techn.

Lot/Batch 38596 Purity 99.10%

Vehicle Distilled water; Satellite group: corn oil

Species Rat, females

Strain Sprague-Dawley derived, Albino

Age 9-11 weeks

Weight at dosing 164-213 g

Source Ace Animals, Inc., Boyertown, PA, USA

Main Test: Based on the Sponsor's estimation of an LD_{50} of 350 mg/kg, the test was conducted using a default starting dose level of 111 mg/kg (as a 30% w/w mixture in distilled water), which was administered to one healthy female rat by oral gavage. Following the Up and Down test procedure, additional females (three at each dose level) were tested at levels of 350 and 1110 mg/kg.

Confirmatory Test: Following completion of the main test using distilled water as a vehicle (the preferred vehicle according to the guideline), a confirmatory limit test at 350 mg/kg using corn oil was performed. A number of previous gavage toxicity studies with the test substance have been conducted with corn oil as the vehicle, and it was considered useful to confirm that the vehicle would not make a significant difference to toxicity. The test substance diluted in corn oil at 30 % w/w was administered to one female rat by oral gavage at 350 mg/kg. Due to the absence of mortality in this animal, four additional females were sequentially treated at the same dose level.

Females were selected for the test because they are frequently more sensitive to the toxicity of test compounds than males. All animals were observed for mortality, signs of gross toxicity, and behavioural changes for a 14-day period post dose administration or until death occurred. Body weights were recorded prior to administration and again on Study days 7 and 14 or after death. Necropsies were performed on all animals.

Findings:

Table 12: Main test (vehicle: distilled water)

Dosing Sequence	Animal No.	Sex dosed	Dose level (mg/kg)	Dose volume (ml)	Outcome*
1	3101	F	111	0.077	S
2	3102	F	350	0.21	S
3	3103	F	1110	0.59	D
4	3104	F	350	0.20	S
5	3105	F	1110	0.65	D
6	3106	F	350	0.20	S
7	3107	F	1110	0.63	D

^{*} S – Survival, D – Death

Main test:

111 mg/kg Dose Level (1 animal): no clinical signs, no effect on body weight, no gross abnormalities

350 mg/kg Dose Level (3 animals): All animals survived and gained body weight during the study. Following administration, two animals were hypoactive and/or exhibited reduced fecal volume and the third animal was observed with piloerection. However, all animals recovered by Day 2 and appeared active and healthy for the remainder of the 14-day observation period. No gross abnormalities were noted for these animals.

1110 mg/kg Dose Level (3 animals): All animals died within one day of test substance administration. Prior to death these animals were hypoactive and/or exhibited hunched posture and tremors. Gross necropsy of the decedents revealed red discoloration of the lungs, red intestines, red oral discharge and/or ano-genital staining.

Dosing Sequence	Animal No.	Sex dosed	Dose level (mg/kg)	Dose volume (ml)	Outcome*
1	3108	F		0.22	S
2	3109	F		0.24	S
3	3110	F	350	0.23	D
4	3111	F		0.24	S
5	3112	F		0.24	S

^{*} S – Survival, D – Death

Confirmatory test:

One animal died within one day of test substance administration. Prior to death this animal was hypoactive and exhibited tremors. Following administration, the surviving animals were hypoactive, however, they recovered by Day 1, gained body weight and appeared active and healthy for the remainder of the 14-day observation period. Gross necropsy of the decedent revealed discoloration of the intestines and lungs. No gross abnormalities were noted for any of the euthanized animals when necropsied at the conclusion of the 14-day observation period.

Conclusion:

Based on the results of the study, the acute oral LD_{50} of the test substance is estimated to be 654 mg/kg in female rats with approximate 95% Confidence Limits of 1110 mg/kg (upper) and 350 mg/kg (lower). A confirmatory limit test indicated that animals are not substantially more sensitive to the test substance when administered in corn oil than when administered in water.

Reference: P0071: Acute oral toxicity test in the mouse, Project Number 102/50
Author(s), year: Coles R., 1990
Report/Doc. number: Doc.No. 521-001, Lonza Report No. 1325, Conducting laboratory:

Report Boe. named.

Safepharm Laboratories Limited, Derby, UK

Guideline(s): OECD Guideline 401 (1981)); US EPA Pesticide Assessment Guidelines

Subdivision F, No 81-1; Annex V method B1 of EEC Commission

Directive 84/449/EEC

GLP: Yes Deviations: No Validity: Valid

Material and Methods:

Groups of 5 mice/sex received single doses of 0 (vehicle control), 400, 526, 693, 912 and 1200 mg/kg bw metaldehyde (sponsor's identification: P0071; batch no. 5448; purity 99.3 %) suspended in arachis oil by oral gavage. As the mortality data derived from these dose groups did not permit calculation of the acute oral LD₅₀ value, an additional group of 5 mice/sex were treated with 304 mg/kg. The dose selection was based upon a range-finding study when one male and one female mouse per test group were dosed with 500, 1000, 3000 and 5000 mg/kg bw. At the beginning of the study the mice (strain: BKW; source: Bantin & Kingman Ltd., Hull, UK) weighed 21-30 g (males) and 20-25 g (females), and were approximately six to eight weeks old. Animals were observed 1 and 4 hours after dosing and subsequently once daily for 14 days. Deaths and evidence of overt toxicity were recorded at each observation. Individual bodyweights were recorded on the day of treatment (day 0), days 7 and 14, or at death. All animals were subjected to gross necropsy. No tissues were retained.

Findings:

Range finding study: No deaths were observed at 500 mg/kg, while 2 deaths were noted at doses of 1000 mg/kg and above. The mortality data indicated an oral LD₅₀ between 500 and 1000 mg/kg.

Main study: The observed death rates were 0/5 for both sexes at 304 mg/kg, 4/5 for males and 3/5 for females at 400 mg/kg, 3/5 for both sexes at 526 mg/kg, and 5/5 for males and 4/5 for females at 693 mg/kg, 5/5 for both sexes at 912 mg/kg, and 5/5 for males and 4/5 for females at 1200 mg/kg. Deaths were noted 1-4 hours after dosing and on day 1 (one female at 400 mg/kg) and day 7 (one male and one female at 526 mg/kg) after treatment. Common signs of toxicity included hunched posture, lethargy and piloerection. Additional or isolated signs of toxicity noted were ataxia, ptosis, pallor of the extremities, decreased respiratory rate, occasional body tremors and tonic convulsions. Surviving animals appeared normal 1, 2 or 7 days after treatment. All animals treated with 304 mg/kg appeared normal throughout the study. Incidents of reduced bodyweight gain and bodyweight loss were noted in all treatment groups during the study period. Macroscopic findings found at necropsy of animals that died during the study were red lungs, dark liver or patchy pallor of the liver, pale spleen, dark kidneys, haemorrhage of the glandular gastric epithelium and large intestine. No findings were noted at necropsy of surviving animals.

Conclusion:

Metaldehyde (suspended in arachis oil) is of moderate toxicity to mice after oral administration. The acute oral LD_{50} was calculated to be 411 (346-489) mg/kg bw for males and 443 (333-591) mg/kg bw for females.

4.2.1.2 Acute toxicity: inhalation

The available study on acute inhalation toxicity shows major limitations and was considered not to be of sufficient validity for assessing the acute inhalation toxicity of metaldehyde.

Reference: Acute inhalation toxicity to the rat of metaldehyde dust

Author(s), year: Berczy Z., Cobb L., Cherry C., 1973

Report/Doc. number: Eurofins/Product Safety Laboratories, USA; Laboratory Identification No.

26776

Lonza Report No. 4377, Doc. No.: 521-003

Guideline(s): No test guideline is mentioned in the study report.

GLP: No. When the study was performed (1973), GLP was not compulsory Deviations: Several deviations with respect to current guidelines were noted: reduced number of animals/group; two dose levels: no information on substance

number of animals/group; two dose levels; no information on substance identification; major reporting deficiencies like air flow, oxygen content,

actual concentration, temperature, humidity and MMAD

Validity: limited scientific validity

Material and Methods:

Groups of 4 male and 4 female rats (strain: Sprague Dawley; source: Charles River Ltd., UK) weighing 205-245 g (males) and 173-204 g (females) were exposed for a 4 hour period (whole body exposure) to dust aerosols of metaldehyde. Neither the purity of the test substance nor the batch no. is presented in the study report. The nominal concentrations were 0 (dry air), 1 and 15 mg a.i./L air. These concentrations were produced by the selection of suitable powder feed and air flow rates in the dust generator. The test athmosphere dust was passed through a central air inlet into a 100 L

exposure chamber containing a group of 8 rats separated into individual compartments by radial grills. Particle size distribution was monitored by collecting dust on a glass slide and examination of the slide by light microscopy. After the 4 hour exposure period the animals were kept under observation for 14 days. Body weights were measured on the day of exposure (day 1) and then on days 2, 5, 8, 12 and 14. At the end of the study macroscopy was performed on all animals.

Findings:

Regarding particle size it was reported that 83-90 % of the particles ranged in a size of 1-5 μ m and therefore were considered sufficiently small to reach the respiratory surfaces of the lung. Another 4-7 % of the particles had a reported size of 5-15 μ m and were regarded as inhalable. The remaining 6-10 % had a particle size larger than 15 μ m.

One female rat died overnight following the exposure with the high dose of 15 mg/L air. Slight temporary dyspnea and occasional sneezing was observed during exposure of the low dose animals. Eye irritation, dyspnea, sneezing, discomfort and increased nasal and oral secretion were seen in animals exposed to the high dust concentration. After exposure, these signs of irritation to the respiratory system disappeared within one hour, but the animals remained lethargic for two days (low concentration and one week (high concentration), respectively, post-exposure. Slight reductions of bodyweight were recorded on the day following the exposure procedure in both test groups, but the rate of growth returned to normal for the rest of the observation period. In the rat that died following dust exposure, marked congestion of the lung and pleural fluid in the thorax was noted; however, these findings could possibly be due to autolytic changes. In surviving rats, no macroscopic findings were noted at termination of the study.

Conclusion:

The results of the study showed irritating effects to the respiratory system, however, it is not clear if this irritation was an intrinsic characteristic of the test compound or a secondary result from the dust distributed in the chamber in this study of limited validity. Only nominal but not actual concentrations are presented in the study report. No LC50 value was calculated. Regarding the observed death rate, the results indicate that the LC50 value (4 h whole body exposure) for male and female rats is in excess of 15 mg/L air (nominal concentration). However, considering the limitations of the study, no exact statement on the actually inhaled concentration of metaldehyde is possible.

An attempt was made to initiate a new inhalation toxicity study, but the particle size could not be reduced to meet the requirements of OECD 403 (Griffiths, 2009; summarized in the additional report to the DAR). Based on this study, it is not feasible to perform a valid inhalation toxicity study due to the physico-chemical properties of metaldehyde.

Reference: Outcome of technical pre-trials for an acute inhalation toxicity study

with Metaldehyde

Author(s), year: Griffiths D.R., 2009

Report/Doc. number: Harlan Laboratories Ltd., Derbyshire, UK; Ref: L260109-01/vm

Lonza Report No. 4336, Doc. No.: 581-004

Guideline(s): Not applicable GLP: Not applicable Deviations: Not applicable

Validity: Valid

Material and Methods:

A report in the form of a letter (2 pages) was submitted. In this letter, the work which was carried out to generate appropriate test atmospheres for the purposes of performing an acute inhalation toxicity study as recommended in OECD Test Guideline 403 is described. With respect to the particle size it is referred to the Draft OECD Test Guideline 403 in the current version of 1996 which requests a Mean Mass Aerodynamic Diameter (MMAD) of $< 4 \,\mu m$. This reflects the state-of-the-art requirements for the particle size of a test item to be tested in an acute inhalation toxicity study.

Due to the crystalline nature of the test material it was considered unsuitable for the purposes of direct atmosphere generation. Therefore attempts were made to grind the test material.

Findings:

With different grinding techniques like a Braun mini grinder, a Retsch Centrifugal Ball Mill with a Wrights Dust Feeder or a SAG 410 Solid Aerosol Generator, the smallest particle size which could be achieved was larger than 4 μ m (MMAD = 11.07 μ m).

With a particle separator which was introduced before the aerosol entered the exposure chamber in order to remove large particles and thereby increase the inhalable portion of the generated aerosol, the MMAD was still $9.58 \, \mu m$.

Conclusion:

It was considered that all techniques available were exhausted to reduce the MMAD of the test material. Thus, the MMAD of approximately 10 µm was the best that could be practically achieved in these series of experiments. It was concluded that the nature of the test material has been demonstrated to be unsuitable for the performance of an acute inhalation toxicity study according to the state-of-the-art requirements for particle sizes reflected in the Draft OECD 403 of 1996.

4.2.1.3 Acute toxicity: dermal

An acute percutaneous toxicity study was performed in rats. However, this study shows some deviations and is therefore considered to be of limited scientific validity: No test guideline is mentioned in the study report. It complies only partly with OECD guideline 402. Following deviations were noted: no specification of the test material; several reporting deficiences concerning housing and feeding conditions; no detailed clinical observation findings and individual necropsy findings. The study is not conform with GLP. When the study was performed (1974), GLP was not compulsory.

Despite of the mentioned deviations, a conclusion can be drawn from this study regarding dermal toxicity. Groups of 5 rats/sex received a topical application of metaldehyde (neither purity nor batch no. is presented) suspended in water at dose levels of 0 (vehicle control) and 5000 mg/kg bw. The test substance was prepared as a 50% suspension in water and applied evenly to the intact clipped skin on the dorso-lumbar region, equivalent to 10% of the total body surface. The treated area was then covered with aluminium foil which was held in contact with waterproof plaster. After the 24 hour exposure period, the dressings were carefully removed and the treated skin washed with warm dilute soap solution and afterwards rinsed with clean warm water. The animals were observed during a period of 14 days. No mortality was noted at 5000 mg/kg. The only signs of clinical toxicity were slight lethargy and piloerection on the day of treatment (number of animals concerned not given). There was no local reaction like erythema or oedema. All treated animals and controls gained normal bodyweight during the study. Terminal autopsy revealed darkening of the liver and spleen together with pale or mottled kidneys (no number of incidences reported).

Conclusion: Several limitations of the study design were noted, however, the results demonstrated adequate evidence that the acute dermal toxicity of metaldehyde is low. The acute dermal toxicity LD_{50} of metaldehyde (suspended in water) is considered to be greater than the maximum dose tested (5000 mg/kg bw).

4.2.1.4 Acute toxicity: other routes

No data available.

4.2.2 Human information

Many poisonings are reported after accidental or suicidal intake of metaldehyde with intoxications ranging from mild to lethal outcome. Clinical signs include gastrointestinal symptoms which may be followed by convulsions, somnolence, apnoe, cyanosis, coma and death.

Several cases of metaldehyde poisoning are reported and summarized in a dissertation from Borbely A. (1970, Doc.No. 592-001). They were derived from a total of 223 cases of metaldehyde intoxication which were reported to the Swiss Toxicological Information Centre between march 1966 and June 1969. The intoxications were all caused by metaldehyde tablets or snail pellets containing metaldehyde. The data were derived mainly from report forms filled in by the telephone information service and completed by the treating physician. 122 intoxications were due to metaldehyde tablets (pure metaldehyde), and 101 cases to snail pellets (5 - 7 %) metaldehyde. 189 children, 24 adults and 10 animals were involved. Most of the children affected were aged 2 – 4 years. Apart from 20 attempted suicides by adults, all the intoxications were accidental. The course of the poisoning is known in 128 cases: 87 cases were mild, 25 moderately severe, 14 severe and 2 fatal. 22 case histories illustrating particularly severe and typical intoxications are presented in the dissertation and reported here as follows:

Case 1

A 50 year old woman had taken approximately 10-15 metaldehyde tablets about two hours before hospitalisation. She complained about intense nausea and vomited white matter. Some degree of psychological contact was achieved with the patient but she was unable to give exact details about the number of tablets ingested or when. Gastric lavage with sodium bicarbonate, activated charcoal and Carlsbad salt was adopted. During day 1, mental confusion increased. Epileptiform seizures with tonic-clonic spasms and subsequent respiratory failure and cyanosis were observed. Deep respiration emerged in the evening. Calcium was administered because of tetany signs. In the morning of day 2, the patient developed respiratory acidosis and coma. During days 2 and 3, 47 seizures were counted. There was a gradual reduction in frequency of seizures terminally, but increased occurrence of respiratory failure. On day 4, there was a final respiratory failure of the patient in coma.

Case 2

While playing, a 2 year old boy discovered a portion of a metaldehyde bar which was not completely burnt and swallowed it. Shortly afterwards the boy began to vomit spontaneously. In the evening, he had mild diarrhea and spent a restless night, awakening frequently. When taken out of bed the next morning, the child could neither walk nor stand. The boy showed somewhat slow reactions; contact was possible. The neurological status was as follows: intact bilateral reflexes, no pathological reflexes, Chvostek's sign and muscles slightly hypertonic. No other pathological findings were noted. The child was treated with Valium for immediate sedation and relevant measures were undertaken to promote diuresis. No further symptoms were observed. The child was discharged as cured the next day.

Case 3

A nearly 3 year old child ingested snail pellets containing metaldehyde five hours prior to hospitalization. On admission, the child was shivering from head to foot, with heavily flushed face. Tendon reflexes were markedly enhanced, but not diffuse. Chvostek's sign were initially negative and later positive. Positive signs of ankle clonus were noted. As therapy gastric lavage with Glauber's salts, charcoal, and Luminal by intravenous drip were chosen. On day 2, muscle rigidity and trembling were no longer observed but low-grade hyperreflexia was persistent. Blood alkalosis with otherwise normal Astrup values, presumably due to the vomiting, was noted. The child was discharged the following day.

Case 4

A 33 year old male patient swallowed 6-8 metaldehyde tablets four hours before hospitalisation. Vomiting of white mucus occurred shortly afterwards, followed by violent trembling and shaking over the whole body and intense motor restlessness. Hyperventilation with tetany spasms, loud groaning and clouding of consciousness, choreatic movements, Trousseau's sign, bilateral dorsal flexion of toes, intermittent convulsive choking and white-coated tongue were observed. No enlargement of liver was noted and pulmonary and cardiac findings were normal. Generalized seizures with tonic-clonic spasms and transient loss of consciousness occurred three times. Treatment was performed with morphine-scopolamine and Valium; no signs of renal or hepatic damage were noted. After regression of acute symptoms, the patient remained free of seizures under Luminal.

Case 5

One hour before hospitalisation, the patient (male, 24 years old) had swallowed half a metaldehyde tablet. The status on admission was: intense sensation of burning in the stomach and intense gastric pain, nausea, mild cyanosis, spasms of all extremities, restlessness, anxiety, slight sensory dulling, hyperpnea, slurring of speech, pure cardiac sounds, tachycardia and occasional extrasystoles. No information on progress.

Case 6

A 2 year old girl had been playing with metaldehyde snail pellets. After an unspecified period it no longer responded on being spoken to, and began to twitch and convulse. Slight bouts of vomiting occurred. Findings on admission were: stiff, unconscious child, extension spasms, clonic spasms, pink skin, pupils moderately dilated with little reaction to light, deep, accelerated breathing, tachycardia, increased muscle tone, reacting to touch with spasms and enhanced tendon reflexes. The child was given Somnifen and Taractan and was intubated for gastric lavage which revealed snail pellets. On day 2 the girl was again bright and responsive. The neurological status on discharge was normal (five days after admission).

Case 7

While on military service, the patient (adult male) distributed metaldehyde tablets and then had a meal without first washing his hands. An hour and a half later, he complained of nausea and abdominal pain (data on the further course are lacking).

Case 8

At an unspecified time, the patient (adult male, 30 years old) had ingested metaldehyde tablets. His condition on being found was characterised by vomiting, salivation, epileptiform convulsions and risus sardonicus. He was unresponsive when addressed but did react to mild stimuli. The patient became completely responsive within three days.

Case 9

About two hours previously the patient (male, 18 years old) had swallowed four metaldehyde tablets. Facial twitching, particularly at the side of the mouth occurred. Chvostek and Trousseau signs were positive, increasing restlessness, spasm of the hands and then of the arms and legs were noticed. Debris of metaldehyde tablets were obtained after several gastric lavages and administration of large amounts of laxatives. No acidosis was detected. The spasm-like subsided under Valium and the patient was able to sleep. He was discharged as cured one week later.

Case 10

Two hours previously, the patient (male, 24 years old) had swallowed some metaldehyde powder (an amount approximately equivalent to one-tenth of a metaldehyde tablet) by mistake and complained about a burning sensation in the stomach. After gastric lavage with sodium bicarbonate and liquid paraffin, no further symptoms were observed.

Case 11

Sixteen hours prior to hospitalisation a 2 ½ year old child had swallowed half a metaldehyde tablet and had vomited three times. 24 hours after admission slight trembling of the upper limbs was noted which subsided after several hours.

Case 12

The mother of the patient (male, 17 years old) had prepared rat poison with metaldehyde in the following manner: she had crushed half a metaldehyde tablet, mixed it with some chocolate and made a chocolate truffle out of it. The son ate half of this by mistake. Gastric lavage was adopted. No symptoms of poisoning were detectable.

Case 13

On blowing out a lamp, the patient (adult male) had aspirated metaldehyde dust. Three and half hours later, he complained about nausea, stomach pain and headache. He was treated with Antrenyl and Torecan suppositories and was free of symptoms ten hours after the incident.

Case 14

A two year old girl had been sucking a piece of metaldehyde and had possibly swallowed a small amount. After 25 minutes gastric lavage with sodium bicarbonate and prophylactic treatment with penicillin were adopted. Apart from acetone in urine, no symptoms of poisoning occurred.

Case 15

A 2 ½ year old girl had eaten pieces of metaldehyde scattered in the garden as snail pellets. Some hours later, first spasms, which lasted about sixteen hours were observed despite administration of Luminal. No subsequent spasms occurred. The patient was discharged as cured after seven days.

<u>Case 16</u>

About two and a half hours after ingestion of some snail pellets, the parents noticed that the child's hand was unsteady while drinking from a cup, often spilling the drink. The child (male, four years old) also walked unsteadily and exhibited muscle twitching. Findings on admission were red face, slight tachypnea and swollen abdominal wall. Neurological status was normal, apart from signs of spasticity of the lower limbs and muscle twitching. Within 24 hours after gastric lavage and administration of adsorbent charcoal, the symptoms had regressed. Two days after hospitalisation, however, rather extensive paravertebral hematomas on the back and over the left spina iliaca

anterior superior occurred. Platelets, coagulation time and prothrombin time were normal. The patient was discharged as cured after four days.

<u>Case 17</u>

A 15 year old boy had swallowed an unknown amount of metaldehyde snail pellets 30 minutes previously. During the period of hospitalisation, no symptoms of poisoning occurred, apart from acidosis demonstrable only biochemically.

Case 18

One hour before gastric lavage, the a 2 year old boy has swallowed one metaldehyde tablet or less. No symptoms of poisoning apart transient exanthema after 24 hours were observed.

Case 19

An hour and a quarter prior to hospitalisation and gastric lavage, a 1 year old boy had swallowed no more than one half tablet of metaldehyde. Status and blood count were normal. Gastric lavage with water and then sodium bicarbonate as well as administration of charcoal were adopted. No symptoms of intoxication occurred. Metaldehyde in lavage returns was chemically demonstrated.

Case 20

A 4 year old child had swallowed an unknown amount of metaldehyde tablets. On admission, no symptoms of poisoning except for a reddened face were noted. No subsequent symptoms occurred.

Case 21

The patient (adult male) had been boiling water with metaldehyde for some time when he began to suffer from nausea, dizziness and vomiting (intoxication by aspiration). The dizziness subsided only 24 hours later, after he had vomited another eight times. The patient was treated with oral sodium bicarbonate.

Case 22

Several days previously the patient (no information on age and sex) had swallowed 10 tablets of metaldehyde. He then began to suffer from disturbed vision, seeing triangles and red, green and black images. The disturbances of vision were said to have occurred only on the day of ingestion; no subsequent symptoms were noted. The patient had undergone gastrectomy shortly before. Acid was nonetheless demonstrated in the residual stomach.

In a publication by Moody J. and Inglis F. (1992, Doc.No. 592-029), a single case of metaldehyde intoxication is presented. A 37 year old Asian male was admitted to hospital suffering from a suspected drug overdose and in a comatose condition. He developed pyrexia (temperature up to 38.5°C) and multiple violent seizures which were treated with diazepam and phenytoin. On admission, blood urea, glucose and electrolytes were normal and gastric lavage returned clear fluid. He was treated with intravenous fluids (5 % dextrose and normal saline) but developed hypokalaemia and mild renal impairment. All indices of liver function were normal apart from the AST which was slightly raised on admission (43 U L-1), increasing to 660 U L-1 after 3 days and declining to 95 U L-1 on the fifth day. On the sixth day the patient had recovered sufficiently to be transferred to a psychiatric hospital. Relatives found two empty containers of Slugit liquid in the garage. Slugit liquid contained approximately 20 % metaldehyde and 9 % ethylene glycol. A minimum intake of 35-50 ml was estimated after losses from nausea and vomiting. Subsequent examination of serum and urine specimens by gas chromatography confirmed the presence of metaldehyde. The serum levels remained elevated for 35 hours. Ethylene glycol did not appear to have contributed significantly to toxicity.

Thompson J., Casey P. and Vale J. (1995; Doc.No. 592-030) reported a case of suicidal intake of two 250 ml bottles of Murphy Slugit Liquid (20 % Metaldehyde, 9 % ethylene glycol) and 1 ½ packets of Ratak (difenacoum). The 45 year old male patient vomited and subsequently became comatose and developed convulsions. The patient was ventilated mechanically but died after one week. He was treated with vitamin K and his prothrombin time remained normal. The conclusion of the authors was that the clinical and post mortem findings were in keeping with metaldehyde poisoning, though ethylene glycol (not confirmed by analysis) may also have contributed to the fatal outcome.

A case report of massive metaldehyde poisioning is presented by Longstreth W. and Pierson D. (1982; Doc.No. 592-033). A 32 year old woman swallowed approximately 470 ml of a commercial slug bait that contained 4 % metaldehyde. In total, 28.9 g metaldehyde or approximately 330 mg/kg bw was ingested. All other ingredients were reported to be inert without known toxicity. The patient soon had nausea and vomiting. Two hours after ingestions she had the first of many generalised convulsions. Initially she was treated at a local hospital with gastric lavage, activated charcoal and diazepam, but later that day was transferred to a Medical Center because of continuing convulsions, decreased mental status and muscle spasms. By the time transfer was completed the patient was comatose. On neurologic examination she was noted to be unresponsive to voice or painful stimuli. Pupils were reactive, eye movements were full on horizontal oculocephalic reflexes and corneal reflexes were present. Tone and tendon reflexes were diffusely increased though there were flexor plantar responses. Chvostek's sign was present. The arterial blood gas determinations showed respiratory alkalosis with pH 7.57. Urine pH was 5.5 with ketones present. Serum calcium and magnesium levels were normal. Toxicologic studies on blood and gastric contents were negative except for an unidentified substance. The hospital course was complicated by severe muscle spasms and repeated generalised convulsions despite therapeutic concentrations of phenytoin and phenobarbital in the blood. Diazepam administered intravenously only briefly controlled the spasms and convulsions. Although the serum creatine kinase level was elevated to four times normal, significant myoglobinuria did not develop. She also had pneumonia, increased oral and tracheobronchial secretions and elevated serum transaminase levels, which peaked during the second week and returned to normal by the time of discharge. Convulsions continued for three days, during which time an interictal electroencephalogram showed diffuse slowing with scattered epileptiform discharges. The coma lasted for seven days but tracheal intubation was prolonged to nine days due to general weakness and excessive secretions. When communication became possible the patient was noted to have pronounced memory deficits, an exaggerated glabellar reflex and prominent snout and palmomental reflexes. Three months before this admission, the findings of a detailed neurologic examination (done during a hospital stay for a suicide attempt) had been entirely normal. The patient's strength improved and administration of anticonvulsant drugs was discontinued. Findings on lumbar puncture and contrast-enhanced cranial computerised tomography were normal. Although her mental status improved slowly, at best she had a poor recent and remote memory, flat affect and pronounced latency of response. She was subsequently transferred to the psychiatric service. A repeat electroencephalogram was normal. Primarily she had an adaptive problem-solving impairment and severe impairment of memory in both verbal and visual-spatial areas. The patient was discharged 51 days after the attempted suicide. One year after the poisoning, the patient reported by telephone that her memory had returned almost to normal.

In a review publication by Booze T. and Oehme F., (1985; Doc.No. 592-025), 2 cases of metaldehyde poisoning are cited, of which one case is identical with the case above (32 year old woman) presented by Longstreth W. and Pierson D. (1982; Doc.No. 592-033). The other cited case was derived from EPA database and concerned a 30 year old female which ingested 16 – 19 g of a liquid slug bait. The clinical signs included convulsions for 3 days, fever, coma, memory loss, respiratory depression, frontal lobe damage, regression to infantile reflexes and general apathy. It was not reported if the patient recovered.

No detrimental effects were reported on health in manufacturing personnel.

Conclusion:

Data regarding human intoxication are primarily available from exposures to single doses after accidental or suicidal oral intake. Chronic poisonings at low doses are not known and are not likely to occur because of the rapid elimination of metaldehyde. The course of the intoxication is characterised by a first phase involving gastrointestinal signs such as nausea, salivation, vomiting and later abdominal pain and diarrhoea. This phase may be followed by convulsions, somnolence, coma, apnoea, cyanosis, memory loss and decreased blood pressure.

4.2.3 Summary and discussion of acute toxicity

Two acute oral toxicity tests in rats showed that metaldehyde is of <u>moderate acute oral toxicity</u>. Also in mice moderate acute oral toxicity was demonstrated. At high dose levels, unspecific symptoms like reduced activity and lethargy, hunched posture and decreased respiratory rate but also convulsions and ataxia were observed in both species tested. Target organs in these studies were the lungs, liver and gastrointestinal tract.

Despite some limitations of the study with <u>acute dermal</u> exposure to rats, the results obtained demonstrated low acute toxicity of metaldehyde by this route of administration. The available study on <u>acute inhalation toxicity</u> (Berczy et al., 1973) shows major limitations and was considered not to be of sufficient validity for assessing the acute inhalation toxicity of metaldehyde. An attempt was made to initiate a new inhalation toxicity study, but the particle size could not be reduced to meet the requirements of OECD 403 (Griffiths, 2009). Based on the new experience (Griffiths, 2009) it is not feasible to perform a valid inhalation toxicity study due to the physico-chemical properties of metaldehyde.

Based on the experimental results and according to the classification criteria of Directive 67/548/EEC, metaldehyde requires classification and labelling for acute oral toxicity with $\underline{Xn, R22}$ (Harmful if swallowed).

4.2.4 Comparison with criteria

The lowest LD₅₀ value of 283 mg/kg for acute oral toxicity was found in rats. Under Directive 67/548/EEC these data trigger classification and labelling with Xn; **R22** "Harmful if swallowed" (LD₅₀ oral, rat: $200 < \text{LD}_{50} \le 2000$ mg/kg bw). According to Regulation (EC) No. 1272/2008 metaldehyde belongs to acute toxicity category 3 ($50 < \text{ATE} \le 300$ mg/kg bw) and requires classification and labelling with **H301** "Toxic if swallowed".

For acute dermal toxicity the LD_{50} value was greater than 5000 mg/kg in rabbits. According to the Commission Directive 67/548/EEC (as amended) and to Regulation (EC) No. 1272/2008, metaldehyde does not require any classification and labelling for acute dermal toxicity.

4.2.5 Conclusions on classification and labelling

Directive 67/548/EEC: Xn; R22

Regulation (EC) No. 1272/2008: Acute Tox. 3, H301

development

al (Neeper-

4.3 Specific target organ toxicity – single exposure (STOT SE)

4.3.1 Summary and discussion of Specific target organ toxicity – single exposure

Table 14: Summary of effects observed in rats, mice dogs and rabbits in comparison to cut off vales

Species- Route (Reference)	Maximum applications	Cut off value Cat 1 STOT SE (1272/2008) [mg/kg bw]	Cut off value Cat 2 STOT SE (1272/2008) [mg/kg bw]	Effects below cut off value	Significance of toxicologic effect (1272/2008) below c off value
Rat- acute oral (gavage) (Jones J. et al., 1987)	1	300	2000	-≥ 200 mg/kg bw: death, hunched posture, pilo-erection, lethargy, decreased respiratory rate	Lethal effect, already covered by acute toxicity classification
Rat- acute oral (gavage) (Durando J., 2009)	1	300	2000	-≥ 350 mg/kg bw: hypoactive and/or reduced fecal volume, piloerection -≥ 1100 mg/kg bw: death, hypoactive, hunched posture, tremors	Absence of significant toxicity at 350 mg/kg bw Lethal effect, already covere by acute toxicity classification
Mouse- acute oral (gavage) (Coles R., 1990)	1	300	2000	-≥ 400 mg/kg bw: death, hunched posture, lethargy, piloerection, ataxia, ptosis, pallor of the extremities, decreased respiratory rate, tremors, tonic convulsions	Lethal effect, already covered by acute toxicity classification
Mouse- 90 days oral (Gill M. et al., 1990)	1-2	300	2000	- ≥ 743 (F) mg/kg bw: death - 1919 (M)- 2996 (F) mg/kg bw: death	Lethal effect, already covered by acute toxicity classification
Dog- 28 days oral (Leuschner J., 2002)	1	300	2000	-≥ 60 mg/kg bw/d: reduced motility, clonic convulsions, increased respiratory rate, emesis -≥ 75 mg/kg bw/d: tonoclonic convulsions, mydriasis, inflated stomach, slight tremor - 90 mg/kg bw/d: ataxia, salivation, abdominal/lateral position, pale gingivial; moribund condition of 1/3 females: shaking of the head, lateral position, difficulty in breathing- no pathological findings (macroscopic) after necropsy	The intensity/severity of the symptoms declined with time and had almost disappeared towards the end of the 4-week treatment Moribund condition, already covered by acute toxicity classification
Dog- 52 weeks oral (Leuscher J., 2003)	1-2	300	2000	- 90 mg/kg bw: ataxia, reduced motility, emesis, tremor, twitching, salivation- no changes in histopathology	Incidence and severity declined from study week 19 onwards. No changes in histopathology.
Rat-	1-2	300	2000	Dams:	Lethal effect, already covere

by acute toxicity

- 150 mg/kg bw: 6/25 death, 6/25

Species- Route (Reference)	Maximum applications	Cut off value Cat 1 STOT SE (1272/2008) [mg/kg bw]	Cut off value Cat 2 STOT SE (1272/2008) [mg/kg bw]	Effects below cut off value	Significance of toxicological effect (1272/2008) below cut off value
Bradley T. et al., 1990)				ataxia, 3/25 tremor, 3/25 twitching, 1/25 hyperactive, 1/25 prostration, 1/25 paresis	classification
Rabbit- development al (Neeper- Bradley T., 1990a)	1-2	300	2000	Dams: -≥ 100 mg/kg bw: 1/5 tremor -≥ 200 mg/kg bw: 2/5 death, 3/5 tremor, 2/5 ataxia, 1/5 paresis, 1/5 broken vertebrae -≥ 350 mg/kg bw: 2/5 death, 2/5 tremor, 2/5 ataxia, 1/5 convulsions, 1/5 prostration, 1/5 broken vertebrae - 500 mg/kg bw: 4/5 death, 3/5 tremor, 4/5 ataxia, 3/5 twitch, 2/5 broken vertebrae	Absence of significant toxicity at 100 mg/kg bw. Lethal effect ≥ 200 mg/kg bw, already covered by acut toxicity classification
Rat- oral (gavage) acute neurotoxicity (Haferkorn J., 2009)	1	300	2000	-≥ 150 mg/kg bw: slight tremor (F), piloerection, diarrhea, impaired ability for wired manoeuvre, impaired gait (F), ↓ resistance during limb rotation (F) - 250 mg/kg bw: 5/10 death (F), reduced motility (F), ataxia (F), tremor (M+F), reduced muscle tone (F), tonic convulsions (F), ↑ body temperature, ↓ hindleg splay (F), ↓ righting reflex (F), ↓ toe/tail pinch response (F)	Reversible changes in clinic signs Transient findings in neurological screening No macroscopic or microscopic findings in the nervous tissue or in the vasculature of the nervous tissue Lethal effect ≥ 250 mg/kg bw, already covered by acut toxicity classification
Rat- oral 90 day repeated dose neurotoxicity (Jones L. et al. 2003)	10	300	2000	- 240 mg/kg bw: 1 female: loss of limb function with no sign of recovery (premature sacrifice on day 22) considered to result from spinal cord injury	No macroscopic or microscopic findings Premature sacrifice at 240 mg/kg bw, already covered by acute toxicity classification

4.3.2 Comparison with criteria

There was no evidence of any specific, non-lethal target organ toxicity arising from a single exposure to metaldehyde. (Reversible) clinical signs of toxicity were observed after single exposures to metaldehyde but were considered to be non-specific signs of general acute toxicity. According to the Guidance on the Application of the CLP Criteria (ECHA 2009): "Acute toxicity refers to lethality and STOT-SE to non lethal effects. However, care should be taken not to assign both classes for the same toxic effect, essentially giving a "double classification", even where the criteria for both classes are fulfilled". The dose-response curve for metaldehyde seems to be rather

steep, meaning that significant acute toxicity occurs only at doses which already lead to mortality. Therefore no classification as STOT SE is proposed.

4.3.3 Conclusions on classification and labelling

Directive 67/548/EEC: no classification proposed

Regulation (EC) No. 1272/2008: no classification proposed

4.4 Irritation

4.4.1 Skin irritation

Table 15: Summary table of relevant skin irritation studies

Method	Results	Remarks	Reference
Skin irritation and corrosivity study in rabbits	Not irritating	-	Jones, J.; 1983

4.4.1.1 Non-human information

A primary skin irritation and corrosivity study was performed in female New Zealand White rabbits. No guideline is mentioned in the study report, but the design complies to a large extent with the requirements of the OECD guideline 404. The study is conform with GLP and is considered to be scientifically valid and acceptable.

A portion of metaldehyde (0.5 g moistened with water) was applied to the clipped dorsal skin. The patch was occluded (no further details given) and secured with a strip of impermeable adhesive tape for a 4-hour exposure period. Following the exposure period, the patches were removed and the skin wiped with a paper towel moistened with water to remove any remaining test material. The skin sites were examined 60 minutes after treatment and then daily for 3 days. Skin reactions were evaluated according to Draize. No skin reactions were noted on any treated site during the observation period.

4.4.1.2 Human information

The irritation and sensitisation potential of 36 substances including metaldehyde was tested (*Lisi P.*, *Carafini S.*, *Assalve D.*; 1987; *Doc.No.* 592-002). The number of persons tested for metaldehyde was 442, of whom 89 were agricultural workers, 30 ex-agricultural workers, and 323 others. Patch tests were performed on the upper back and were read after 48 and 72 h. Irritant and allergic reactions were evaluated according to Cronin's criteria. The concentration tested was 1%. Neither irritant nor allergic reactions were observed in the test persons.

4.4.1.3 Summary and discussion of skin irritation

Metaldehyde (moistened with water) is not irritating to the skin.

4.4.1.4 Comparison with criteria

Estimated skin irritation scores are below the criteria for triggering classification and labelling (according to both DSD and CLP).

4.4.1.5 Conclusions on classification and labelling

Directive 67/548/EEC: no classification proposed

Regulation (EC) No. 1272/2008: no classification proposed

4.4.2 Eye irritation

Table 16: Summary table of relevant eye irritation studies

Method	Results	Remarks	Reference
Acute eye irritation testing rabbits	Very slightly irritang	-	Coles, R.; 1990

4.4.2.1 Non-human information

An acute eye irritation test was performed in three adult female New Zealand White rabbits. The study was conducted according to GLP and to OECD guideline 405. The study is considered scientific valid and acceptable.

The animals received single applications of 0.1 mL (equivalent to approximately 82 mg) of the undiluted test substance (batch no. 5448, purity not given) into the conjunctival sac of the right eye. The eyelids were held together for one second following application. The left eyes remained untreated and served for control purposes. The eyes were examined for ocular reactions according to Draize (1 hour after the instillation and then 1, 2, and 3 days thereafter).

Metaldehyde (undiluted) was found to be slightly irritating to the eyes of rabbits. Iridial inflammation (grade 1) was noted in all treated animals 1 hour after treatment but no longer at 24, 48 and 72 h. Minimal conjunctival irritation (grade 1) was noted in all treated eyes 1 and 24 hours after treatment. Conjunctival chemosis and discharge were found only 1 h after treatment. All treated eyes appeared normal after 48 h. No corneal effects were noted during the study.

4.4.2.2 Human information

Not available.

4.4.2.3 Summary and discussion of eye irritation

Metaldehyde (undiluted) was slightly irritating to the eyes of rabbits.

4.4.2.4 Comparison with criteria

Estimated eye irritation scores (24 – 72 hours; 0 (conjunctival chemosis), 0.33 (conjunctival redness) and 0 (iritis) and 0 (corneal opacity) are below the criteria for triggering classification and labelling (according to both DSD and CLP).

4.4.2.5 Conclusions on classification and labelling

Directive 67/548/EEC: no classification proposed

Regulation (EC) No. 1272/2008: no classification proposed

4.4.3 Respiratory tract irritation

4.4.3.1 Non-human information

One acute inhalation toxicity study of limited validity (Berczy, Z.; 1973) showed some irritating effects to the respiratory system: Eye irritation, dyspnea, sneezing, discomfort and increased nasal and oral secretion were seen in animals exposed to the high dust concentration. After exposure, these signs of irritation to the respiratory system disappeared within one hour.

4.4.3.2 Human information

Not available.

4.4.3.3 Summary and discussion of respiratory tract irritation

Mild and transient signs of respiratory tract irritation were observed in one non-valid acute inhalation toxicity study.

4.4.3.4 Comparison with criteria

Mild and transient signs of respiratory tract irritation seen in a non-valid acute inhalation toxicity study are not sufficient for classifying as respiratory tract irritant.

4.4.3.5 Conclusions on classification and labelling

Directive 67/548/EEC: no classification proposed

Regulation (EC) No. 1272/2008: no classification proposed

4.5 Corrosivity

See chapter 4.4.1 Skin irritation.

4.6 Sensitisation

4.6.1 Skin sensititsation

Table 17: Summary table of relevant skin sensitisation studies

Method	Results	Remarks	Reference
Buehler test (guinea pig)	Not sensitising	Limited validity, some major deviations	Nitka, S.; 1984
Local Lymph Node Assay (mouse)	Not sensitising	-	Bull, A. D.; 2007

4.6.1.1 Non-human information

Skin sensitization in the guinea pig was investigated in a study according the methods of Buehler (1965) but not according to current guidelines. Although no sensitizing effects were observed, the study was considered of limited validity and not adequate to draw conclusions on the sensitizing properties of metaldehyde. Therefore, a new Local Lymph Node Assay was performed. The application of metaldehyde at concentrations of 5%, 10% or 25% w/v in acetone: olive oil (4:1) resulted in ³H-methylthymidine incorporation which was less than 3-fold at all three concentrations (0.4, 0.9 and 1.0, respectively).

4.6.1.2 Human information

The irritation and sensitisation potential of 36 substances including metaldehyde was tested (*Lisi P.*, *Carafini S.*, *Assalve D.*; *1987*; *Doc.No. 592-002*). The number of persons tested for metaldehyde was 442, of whom 89 were agricultural workers, 30 ex-agricultural workers, and 323 others. Patch tests were performed on the upper back and were read after 48 and 72 h. Irritant and allergic reactions were evaluated according to Cronin's criteria. The concentration tested was 1%. Neither irritant nor allergic reactions were observed in the test persons.

4.6.1.3 Summary and discussion of skin sensitisation

Metaldehyde was not sensitising to the skin in a Local Lymph Node Assay, in a Buehler test of limited validity and in a patch test performed in human volunteers.

4.6.1.4 Comparison with criteria

Metaldehyde was not sensitising to the skin in a Local Lymph Node Assay.

4.6.1.5 Conclusions on classification and labelling

Directive 67/548/EEC: no classification proposed

Regulation (EC) No. 1272/2008: no classification proposed

4.6.2 Respiratory sensitisation

No data available.

4.7 Repeated dose toxicity

 Table 18:
 Summary table of relevant repeated dose toxicity studies

Method	Results	Remarks	Reference
Sprague-Dawley rat 4 weeks oral (dietary)	0, 2500, 5000, 10000 and 20000 ppm/diet (equivalent to 0, 197, 382, 761 and 1547 mg/kg bw/d for males; 0, 233, 454 and 875 mg/kg bw/d for females) NOAEL could not be determined LOAEL = 2500 ppm based on: - increased liver weights - histopathological findings in the liver (hepatocellular hypertrophy)	Dose finding study, some deviations, supplementary information only	Van Miller, J.; 1989
Sprague-Dawley rat 90 days oral (dietary)	0, 250, 750 and 2500 ppm/diet (equivalent to 0, 21, 65 and 215 mg/kg bw/d for males and females) NOAEL = 250 ppm Effects at LOAEL: - histopathological findings in the liver (hepatocellular hypertrophy)	-	Thomas, O., Bartlett, A., Brooks, P.; 1998
CD-1 mouse 90 day oral (dietary)	0, 100, 300, 1000, 3000 and 10000 ppm/diet (equivalent to 0, 19, 54, 178, 560 and 1919 mg/kg bw/d for males; 0, 24, 70, 235, 743 and 2996 mg/kg bw/d for females) NOAEL could not be determined LOAEL = 100 ppm based on: - increased liver weights - histopathological findings in the liver (hepatocellular hypertrophy, necrosis, acute inflammation, anisokaryosis)	Dose finding study, limited investigations, supplementary information only	Gill, M., Wagner, C.; 1990
Beagle dog 4 weeks oral (dietary)	Escalating doses: 30, 60, 75 and 90 mg/kg bw/d Fixed doses: 75 and 90 mg/kg bw/d NOAEL could not be determined Observed effects: - mortality at 90 mg/kg bw/day - clinical signs: reduced motility, tremor, convulsions, ataxia, emesis, increased respiratory rate, lateral/ abdominal position, moderate salivation, pale gingival, mydriasis	Dose finding study	Leuschner, J.; 2002
Beagle dog 26 weeks oral (dietary)	0, 20, 60 and 90 mg/kg bw/d NOAEL = 20 mg/kg bw/d Effects at LOAEL: - histopathological findings in the testes: diffuse atrophy of the germinative epithelium - histopathological findings in the prostate: diffuse atrophy	-	Neumann, W.; 1980, 1991 + Re-evaluation of histopathological findings in the testes Leuschner, J.; 2009
Beagle dog 52 weeks oral (dietary)	0, 10, 30 and 90 mg/kg bw/d via diet NOAEL = 10 mg/kg bw/d Effects at LOAEL:	-	Leuschner, J.; 2003 + Re-evaluation of histopathological

	histopathological findings in the testes (atrophy of the germinal epithelium) mortality		findings in the testes Leuschner, J.,
NZW rabbit 21 day dermal	0, 100, 300 and 1000 mg/kg bw/d NOAEL = 1000 mg/kg bw/d	-	Drommer, W.; 2009 Hermansky, S., Wagner, D.; 1991

4.7.1 Non-human information

4.7.1.1 Repeated dose toxicity: oral

28-day toxicity (rat)

Reference: Twenty-Eight day Dietary Oral Toxicity Study with Metaldehyde in

rats

Author(s), year: Van Miller J., 1989

Report/Doc. number: Lonza Report No. 1380, Doc.No. 532-001,

Conducting laboratory: Bushy Run Research Center, Pennsylvania, USA

Guideline(s): No guideline is mentioned in the study report but the design is similar to

OECD guideline 407.

GLP: Yes

Deviations: limited histopathology, no NOAEL could be determined

Validity: This dose finding study is scientific valid but due to the deviations

mentioned considered as supplementary information only.

Material and Methods:

Groups of 10 male and 10 female rats of approximately 8 weeks of age (strain: Sprague-Dawley CD; source: Charles River Breeding Laboratories, MI) received diets containing 0, 2500, 5000, 10000 or 20000 ppm metaldehyde (batch no. 5448; purity 99.0 %), equivalent to 0, 197, 382, 761 and 1547 mg/kg bw/d in males and 0, 233, 454 and 875 mg/kg bw/d in females, resp., for 4 weeks. Due to 100 % mortality in females at the top dose, no daily intakes could be calculated for this group. Diets were prepared weekly; concentrations of metaldehyde in the diet, and stability and homogenicity of the test substance were confirmed by analysis.

Animals were observed twice daily for clinical signs or reaction to treatment. Detailed clinical observations were performed each week. Body weight and food consumption data were collected weekly. Prior to sacrifice, blood samples were taken from fasted animals for haematological investigations (haematocrit, haemoglobin, RBC, erythrocyte indices, reticulocyte count, total leucocyte count, WBC differential, platelets count) and clinical chemistry investigations (aspartate aminotransferase (ASAT), alanine aminotransferase (ALAT), alkaline phosphatase (AP), gamma glutamyl transpeptidase, total protein, albumin, globulin, A/G ratio, blood urea nitrogen, creatinine, glucose, total, direct and indirect bilirubin, sodium, potassium, chloride, calcium, inorganic phosphorus). No urinalysis was performed.

At necropsy, suriving animals were subjected to a detailed gross pathological examination, and weights of selected organs (adrenals, brain with stem, heart, kidneys, liver, ovaries, spleen, testes) were recorded for each animal. Histopathological examinations were limited to adrenals, brain, kidneys, liver, spinal cord and peripheral nerve (sciatic) from all control animals (both sexes), from male rats of the 20000 ppm group and from females of the 10000 ppm group.

Findings:

General observations: At the highest dose level of 20000 ppm, mortality occurred in 100 % of females and 40 % of males. Death occurred in the females during the first week of treatment (1 animal on day 2, 1 on day 3, 1 on day 4, 1 on day 5, 4 on day 6, 2 on day 7) while no males died prior to day 6. Also, 60 % of the females receiving 10000 ppm died (between day 4 and 21) or were sacrificed in a moribund condition. Animals were sacrificed when found with hind limb paralysis/paresis. In all cases the paralysis/paresis was associated with spinal fracture/luxation which resulted in pinching of the spinal cord (see pathology). According to the study report it was assumed that convulsions, although observed in only one high dose female (on day 2), occurred prior to and resulted directly in the fractures/luxations. No other clinical signs of toxicity were considered related to metaldehyde toxicity.

Concerning food consumption, there was some dose-related decrease in both sexes from all treatment groups during the first week of treatment and food intake tended to be slightly lower throughout the remainder of the study. However, these differences were considered to be the result of moderate aversion to the test diet, particularly in the first week. There were also dose-related decreases in body weight gain for males from the 5000, 10000 and 20000 ppm groups, and for females from the 5000 and 10000 ppm groups during the first week of the study. Mean weight and the total weight gain for all groups, however, were similar to controls at the end of the study.

Table 19: 28 day feeding study in rats Food consumption and body weight / body weight gain (mean values)

	0 ppm	2500 ppm	5000 ppm	10000 ppm	20000 ppm
Males					
body weight (g)					
week 0	255.9	255.5	255.1	254.7	254.7
week 1	298.1	293.8	286.1*	280.8**	270.4**
week 2	332.2	329.5	319.7	314.5*	307.1**
week 3	356.3	357.2	347.8	342.0	342.9
week 4	376.8	380.2	370.9	365.1	369.1
body weight gain (g)					
week 0 - 1	42.1	38.2	31.0*	26.1**	13.8**
week 0 - 4	121.3	124.6	115.7	110.4	111.0
food consumption					
(g/animal/day)					
week 0 - 1	25.4	24.1*	22.6**	21.6**	20.1**
week 3 - 4	26.3	26.4	25.0	24.3*	25.4
Females					
body weight (g)					
week 0	164.4	163.5	162.9	164.0	162.8
week 1	186.5	181.1	177.9	172.7	-
week 2	202.7	198.4	196.4	193.7	-
week 3	214.3	210.3	210.1	211.1	-
week 4	226.8	219.5	217.7	218.6	-
body weight gain (g)					
week 0 - 1	22.1	17.6	15.0*	6.8**	-
week 0 - 4	62.4	56.0	54.9	53.2	-
food consumption					
(g/animal/day)					
week 0 - 1	19.0	16.9**	15.3**	13.3**	-
week 3 - 4	18.9	18.2	18.5	18.2	-

^{**} $(p \le 0.01)$, * $(p \le 0.05)$; significantly different from controls

Haematological investigations at study termination revealed a decrease in erythrocyte count for males from the 10000 and 20000 ppm groups. In addition, a trend (not statistically significant) towards decreased values of hemoglobin and hematocrit was observed in all treatment groups of males and in the 2500 and 10000 ppm groups of females. Investigations on clinical chemistry parameters demonstrated a significant increase in urea nitrogen for females from the 2500 and

10000 ppm groups. It was stated in the report that the biological significance of all these findings mentioned is unclear. No other differences between treatment groups and controls were considered treatment-related.

Table 20: 28 day feeding study in rats Haematological and clinical chemistry findings (mean values)

	0 ppm	2500 ppm	5000 ppm	10000 ppm	20000 ppm
Males					
Erythrocyte (106/μL)	7.8	7.7	7.7	7.3*	7.2*
Hematocrit (%)	43.6	42.8	42.8	41.8	41.0
Hemoglobin (g/dL)	16.1	15.7	15.9	15.8	14.9
Urea nitrogen (mg/L)	144	153	148	153	135
Females					
Erythrocyte (106/µL)	7.1	6.9	7.0	6.7	-
Hematocrit (%)	40.1	39.4	40.2	38.9	-
Hemoglobin (g/dL)	15.0	14.8	15.0	14.3	-
Urea nitrogen (mg/L)	141	162*	155	180**	-

^{** (}p \leq 0.01), * (p \leq 0.05); significantly different from controls

<u>Pathology</u>: Gross lesions related to treatment were restricted to the animals that died or were sacrificed in a moribund condition. These lesions were consistent with those to be expected in animals that suffered convulsions and with the observations of hind limb paralysis/paresis. The lesions included color change (congestion and/or haemorrhage) of various organs, traumatic injuries to the back and spinal cord (hemorrhages and fractures/luxations) and lesions of the urinary tract (hydronephrosis, distended bladders and blood ingested urine). The urinary tract lesions were presumably secondary to urine stasis resulting from paralysis.

<u>Organ weights</u>: There was a dose-dependent increase in absolute as well as relative liver weights in all dose groups as compared to controls. There were also increased relative kidney weights in males at 5000 and 20000 ppm. Due to lack of dose response in the 10000 ppm group and lack of histological changes in kidneys, these findings were considered of questionable toxicological significance.

<u>Histopathology</u>: The microscopic lesions found, particularly those in rats which died or were sacrificed moribund, confirmed the lesions observed grossly. They included hemorrhage and/or congestion of various organs, vertebral luxations or fractures, paravertebral and spinal cord hemorrhage and cord compression, vacuolization and malacia. Hydronephrosis, nephritis and urinary bladder hemorrhage occurred more frequently in rats with spinal trauma than in controls. In addition to the above lesions which were mainly the secondary results of convulsive episodes, most of the surviving animals showed dose-related hepatocellular hypertrophy which was rated mild to moderate. Similar findings were observed in animals that died or were sacrificed prior to termination of the study, although the lesions were more evident and severe in rats that survived longer. Furthermore, sporadic foci of hepatocellular degeneration were noted in animals that survived to termination of the study.

Table 21: 28 day feeding study in rats Organ weights and histopathological findings

	0 ppm	2500 ppm	5000 ppm	10000 ppm	20000 ppm
Males					
Liver weight:					
absolute (g)	9.696	12.335**	13.017**	12.751**	16.205**
relative (% of bw)	2.755	3.458**	3.738**	3.700**	4.665*
Hepatocellular	0/10	8/10	9/10	10/10	6/6
hypertrophy					
Kidney weight:	2.716	2.895	3.001	2.826	3.076

	0 ppm	2500 ppm	5000 ppm	10000 ppm	20000 ppm
absolute (g)	0.774	0.813	0.863**	0.820	0.886**
relative (% of bw)					
Females					
Liver weight:					
absolute (g)	6.652	7.053	7.328	8.077	-
relative (% of bw)	3.144	3.389	3.557**	3.922**	-
Hepatocellular	0/10	10/10	10/10	4/4	-
hypertrophy					

^{** (}p \leq 0.01), * (p \leq 0.05); significantly different from controls

Conclusion:

Continuous treatment of rats with metaldehyde at dietary concentrations of 2500 ppm or more over 28 days caused systemic toxicity as shown by increased liver weight and hepatocellular hypertrophy. At higher dose levels (10000 ppm for females and 20000 ppm for both sexes), mortality and spinal injuries, presumably as a result of convulsions, occurred. Changes in haematology (decrease in erythrocytes) were found in males at 10000 ppm and above. In addition, transient reduction of body weight gain and food consumption was noted at higher dose levels. A NOAEL could not be determined in this study. The LOAEL was 2500 ppm (equivalent to 197 mg/kg bw/d for males and 233 mg/kg bw/d for females).

90-day toxicity (rat)

Reference:	P0071: Ninety day sub-chronic oral (dietary) toxicity study in the rat
Author(s), year:	Thomas O., Bartlett A., Brooks P., 1998
Report/Doc. number:	Lonza Report No. 2974, Doc.No. 533-003,
	Conducting laboratory: Safepharm Laboratories Limited, Derby, UK
Guideline(s):	Japanese MAFF Guidelines for Toxicological Studies 59 No 4200, 1985
GLP:	Yes
Deviations:	No
Validity:	Yes

Material and Methods:

10 rats/sex/group (strain: Sprague-Dawley Crl:CD BR; source: Charles River Ltd., Kent, UK) received dietary concentrations of 0, 250, 750 or 2500 ppm metaldehyde (P0071; batch no. 22654; purity 99.1 %) over a period of 90 days. The mean achieved dose levels were 0, 21, 65 or 215 mg/kg bw/day for both sexes. At the start of the treatment the animals were approximately 5-8 weeks old and weighed 156-212 g (males) and 132-194 g (females). Diets were prepared at monthly intervals and stability and homogeneity of the test material in the diet were determined and confirmed by analysis.

All animals were examined for clinical signs once daily. Bodyweights were recorded at weekly intervals and at terminal kill. Food consumption was recorded for each cage group weekly throughout the study. Water intake was observed daily for each cage group by visual inspection of the water bottles for any overt change. The eyes of all control and high dose animals were examined at the beginning and at the end of the study. Prior to sacrifice, blood samples were taken from nonfasted animals from the lateral tail vein. Hematology included haemoglobin, erythrocyte count, hematocrit, MCH, MCV, MCHC, total and differential leucocyte count, platelet count, reticulocyte count, prothrombin time and partial thromboplastin time. Clinical chemistry parameters assessed were urea, glucose, total protein, albumin, albumin/globulin ratio, sodium, potassium, chloride, calcium, inorganic phosphorus, aspartate aminotransferase (ASAT), alanine aminotransferase

(ALAT), alkaline phosphatase (AP), creatinine and total bilirubin. The following parameters were measured in freshly collected urine: volume, specific gravity, pH, protein, glucose, ketones, bilirubin, urobilinogen, reducing substances, blood and microscopic examination of the sediment. All animals were subjected to a full external and internal macroscopic examination. Organ weights were determined from adrenals, brain, heart, kidneys, liver, ovaries, spleen and testes. Histopathological examinations included samples of the following tissues and organs of all animals: adrenals, aorta (thoracic), bone & bone marrow (femur including stifle joint, sternum), brain (three levels), caecum, colon, duodenum, exorbital lachrymal gland, eyes, gross lesions, heart, ileum, jejunum, kidneys, liver, lungs with bronchi, lymph nodes (cervical and mesenteric), mammary gland, skeletal muscle, oesophagus, ovaries, pancreas, pituitary, prostate, rectum, salivary glands (submaxillary), sciatic nerve, seminal vesicles, skin (hind limb), spinal cord (cervical, mid-thoracic and lumbar), spleen, stomach, testes with epididymides, thymus, thyroid/parathyroid, tongue, trachea, urinary bladder and uterus.

Preliminary study: The dose levels were chosen after the conduction of a 14-day preliminary study with 3 rats/sex/group receiving 0, 800, 2500 or 7500 ppm (estimated to be equivalent to 0, 40, 125 or 375 mg/kg bw/d). One female of the highest dose group developed clinical signs on study day 10 including dehydration, pallor of the extremities, hindlimb paralysis, piloerection, ptosis and body tremors and was sacrificed in extremis immediately. Bodyweight gain and also body weight was reduced during week 1 in females receiving 7500 ppm. One female from either the remaining treatment groups showed a bodyweight loss at day 4. A reduction in bodyweight gain was also apparent amongst males receiving 7500 ppm but only slightly in lower dose groups. Animals receiving 7500 ppm and females receiving 2500 ppm showed a reduced dietary intake during the first week of the study. Absolute and relative liver weights were elevated in both sexes at 7500 ppm. Microscopic examination revealed centrilobular hepatocyte enlargement in both sexes at 2500 and 7500 ppm and in two males at 800 ppm.

Findings:

<u>General observations</u>: One female from the highest dose group was found dead on study day 11 without any clinical precursors that might indicate a deterioration in health. Also histopathological examination failed to determine a precise cause of this death. There were no clinical signs of toxicity detected during the study.

<u>Body weight</u>: Females treated with 2500 ppm showed a statistically significant reduced bodyweight gain during the first week of treatment when compared with controls. Bodyweight development returned to normal thereafter although these animals completed the treatment period with a lower terminal bodyweight than controls.

<u>Food consumption</u>: There was no adverse effect on dietary intake during the study. Females treated with 2500 ppm did, however, show a reduction in food efficiency (ratio of bodyweight gain to dietary intake) during week 1 which recovered thereafter. No overt intergroup differences in water consumption were found during the study.

Table 22: 90 day feeding study in rats: Body weight / body weight gain

	Dose group level (ppm)								
		Ma	les		Females				
	0	250	750	2500	0	250	750	2500	
Body weight gain (g)									
- week 1	55	56	57	53	27	24	24	17**	
- week 2	51	45	48	47	21	18	20	19	
Body weight (g)									
- day 0	189	188	190	182	166	163	158	156	
- day 90	501	502	493	493	310	299	294	269	

** (p< 0.01); significantly different from controls

Ophthalmoscopic examination: There were no treatment-related effects observed.

<u>Hematology</u>, <u>clinical chemistry and urinalysis</u>: No treatment-related effects or any significant intergroup differences were observed.

Organ weights: Individual males and females of the 2500 ppm group showed a slight increase in relative liver weight compared to controls, however, statistical significance was not achieved. Females receiving 2500 ppm showed a statistically significant reduction in absolute heart and spleen weight. In the absence of a reduction of relative organ weights and histopathological findings, these differences were considered to be incidental or as a result of lower terminal bodyweight of these animals. Females of this group showed also a significant increase in relative brain weight which is often reported when bodyweight was reduced during treatment. A reduction of relative adrenal weights in females receiving 750 ppm was clearly not dose related and considered to be of no toxicological significance.

Table 23: 90 day feeding study in rats: Absolute and relative (% of bodyweight) organ weights

	Dose group level (ppm)								
		Ma	ales		Females				
	0	250	750	2500	0	250	750	2500	
Liver									
- absolute	16.07	15.43	15.46	17.05	10.39	10.03	9.27	9.61	
- relative (% bw)	3.21	3.08	3.14	3.45	3.36	3.37	3.19	3.66	
Heart									
- absolute	1.63	1.55	1.62	1.70	1.07	0.99	0.98	0.89**	
- relative (% bw)	0.33	0.31	0.33	0.35	0.35	0.34	0.34	0.34	
Spleen									
- absolute	0.88	0.82	0.80	0.81	0.59	0.52	0.52	0.46**	
- relative (% bw)	0.18	0.16	0.16	0.17	0.19	0.18	0.18	0.18	
Brain									
- absolute	2.21	2.15	2.09	2.30	1.94	1.95	1.87	1.97	
- relative (% bw)	0.45	0.43	0.43	0.48	0.64	0.67	0.65	0.76**	
Adrenals									
- absolute	0.062	0.070	0.065	0.068	0.085	0.077	0.072**	0.075	
- relative (% bw)	0.012	0.014	0.013	0.014	0.028	0.026	0.025	0.029	

^{** (}p< 0.01); significantly different from control group

<u>Necropsy</u>: Enlarged liver was reported for 5 males receiving 2500 ppm although this was only partly reflected in the liver weights. The female found dead (2500 ppm) showed pale kidneys and spleen together with normally expected post-mortem changes. No other treatment-related findings were noted.

<u>Histopathology</u>: Centrilobular hepatocyte enlargement was noted in both sexes at doses of 750 ppm and above. No further treatment-related effects were found.

Table 24: 90 day feeding study in rats: Histopathological findings in the liver

	Dose group level (ppm)								
		Ma	les		Females				
	0	250	750	2500	0	250	750	2500	
Centrilobular hepatocyte									
enlargement									
- minimal	0/10	0/10	5/10	1/10	0/10	0/10	5/10	4/10	
- slight	0/10	0/10	0/10	8/10	0/10	0/10	0/10	5/10	

Conclusion:

The predominant treatment-related effects were found in the liver. Centrilobular hepatocyte enlargement was observed at 750 ppm and above while liver enlargement and increased relative liver weight were noted in individual animals only in the high dose group (2500 ppm). Females receiving 2500 ppm showed a reduction of body weight gain and food efficiency during the first week of treatment which resulted in a reduced (not statistically significant) terminal body weight. In conclusion, the NOAEL is considered to be 250 ppm (equivalent to 21 mg/kg bw/d for both sexes).

90-day toxicity (mouse)

Reference: Ninety-Day Dietary Dose Range Finding Study with Metaldehyde in

Mice

Author(s), year: Gill M. and Wagner C., 1990

Report/Doc. number: Lonza Report No. 1546, Doc.No. 533-002,

Conducting laboratory: Bushy Run Research Center, Pennsylvania, USA

Guideline(s): Dose finding study, no guideline is mentioned in the study report

GLP: Yes

Deviations: No haematology and clinical/urine chemistry were performed.

Validity: This dose finding study is scientific valid but due to the limited

investigations considered as supplementary information only

Material and Methods:

In this dose finding study for an oncogenicity study 15 male and 15 female mice per dose group received diets containing 0, 100, 300, 1000, 3000 or 10000 ppm metaldehyde (batch no. 5448, purity: 99.0 %) over 90 days. The actual intake of metaldehyde was 0, 19, 54, 178, 560 and 1919 mg/kg bw/d for males and 0, 24, 70, 235, 743 and 2996 mg/kg bw/d for females. The animals (strain: CD-1, source: Charles River Breeding Laboratories, MI, USA) were approximately 7 weeks of age at first dosing. Stability and homogeneity of the test material in the diet were determined and confirmed by analysis.

During the treatment period, observations for mortality were made twice daily. Detailed clinical observations were performed once each a week, and observations for overt clinical signs were made on all other days. Body weight and food consumption data were collected weekly. No haematology, clinical chemistry or urinalysis were performed. All animals were subjected to complete necropsy. Organ weights (absolute, relative to body weight, relative to brain weight) were determined for liver, kidneys, brain with stem and testes. Histopathological examinations were performed on the following tissues and organs from 10 animals/sex/group randomly selected from the surviving animals of the control and 10000 ppm group. In addition, these tissues were examined for all animals that died during the study: gross lesions, brain (cerebral cortex, cerebellar cortex, medulla/pons), pituitary, thyroid-parathyroid complex, thymic region, lungs with mainstream bronchi, heart, liver (three lobes), spleen, esophagus, stomach, duodenum, jejunum, ileum, cecum, colon, representative lymph nodes (mesenteric and submandibular), kidneys, adrenals, pancreas, testes, epididymis and ovaries. From the other dose groups, only selected organs were examined for 10 animals/sex/group: liver, kidneys, testes, stomach, lungs, duodenum and gross lesions.

Findings:

General observations: There were no treatment-related clinical signs observed during the study. However, mortality occurred in 5 males and 1 female of the 10000 ppm group: 3 males on study day 1, one female on study day 2, one male on study day 4 and one male on study day 8. A female from the 3000 ppm treatment group was found dead on study day 24. The cause of death for these

animals was not apparent based on gross or microscopic examination. In addition, one male from the controls died of disseminated lymphosarcoma on study day 80. Finally, a female receiving 3000 ppm died as a result of a cage accident on study day 36.

<u>Food consumption</u>: There was no treatment-related effect on food consumption.

<u>Body weight</u>: Body weight gain was lower in males receiving 10000 ppm during the first week of the study. Body weight for females in the 3000 and 10000 ppm groups tended to be higher than for controls beginning in the 4th week of the study and occasional statistically significant differences from control were observed thereafter for body weight and/or body weight gain. Body weight findings probably should be considered together with the increases in liver weight observed during the study. For example, body weight reduction in males receiving 10000 ppm could be masked by the significant increase in liver weight. In addition, the increase in body weight for females may reflect increases observed in liver weight.

Table 25: 90 day feeding study in mice: Body weight / body weight gain

	Dose group level (ppm)								
	0	100	300	1000	3000	10000			
Males									
body weight gain (g)									
-week 0-1	1.2	1.1	1.1	0.8	1.5	0.4**			
-week 0-13	6.6	6.7	6.5	7.6	7.4	6.7			
Females									
body weight gain (g)									
-week 0-1	1.1	0.8	0.9	1.2	0.9	0.8			
-week 0-13	5.5	5.0	6.1	6.1	7.0**	6.7**			

^{**} $(p \le 0.01)$, significantly different from controls

Organ weights: Dose-related increases (mostly statistically significant) in absolute and relative liver weights were observed in both sexes at dose levels of 300 ppm and above. The percent increase for absolute liver weights compared to controls ranged from 6-181 % for males and 12-104 % for females. 15-17 % decreases in absolute and relative kidney weights were observed in males in the 10000 ppm treatment group.

Table 26: 90 day feeding study in mice: Absolute and relative (% of bodyweight, % of brain weight) organ weights

			Dose group	level (ppm)		
	0	100	300	1000	3000	10000
Males:						
Liver weight						
- absolute	1.902	1.930	2.018	2.288**	2.926**	5.347**
- relative (% body weight)	5.525	5.645	5.895**	6.520**	8.340**	15.558**
- relative (% brain weight)	398	409	431*	468**	602**	1100**
Kidney weight						
- absolute	0.650	0.648	0.611	0.642	0.627	0.549
- relative (% body weight)	1.890	1.909	1.787	1.827	1.795	1.607**
- relative (% brain weight)	136	138	130	131	129	113**
Females:						
Liver weight						
- absolute	1.495	1.552	1.677**	1.840**	2.168**	3.049**
- relative (% body weight)	5.461	5.589	5.856*	6.536**	7.556**	10.607**
- relative (% brain weight)	311	310	333	371**	432**	615**
Kidney weight						
- absolute	0.408	0.404	0.430	0.389	0.404	0.407
- relative (% body weight)	1.496	1.454	1.502	1.395	1.434	1.419
- relative (% brain weight)	85	80	85	79	81	82

^{*} $(p \le 0.05)$; ** $(p \le 0.01)$; significantly different from controls

<u>Necropsy</u>: Treatment-related gross lesions were limited to the livers of males in the 10000 ppm treatment group (7 swollen and/or size increased) and 300 ppm treatment group (2 swollen).

<u>Histopathology</u>: Hepatic lesions were observed in all treatment groups and included hepatocellular necrosis, hypertrophy and/or hyperplasia, inflammation, anisokaryosis, hepatocellular vacuolization, cholestasis and biliary hyperplasia. The number of the lesions observed for individual animals, the number of animals affected within a treatment group, and the severity of individual lesions increased in a dose-related manner for animals in all treatment groups. The severity of lesions were minimal for the 100 and 300 ppm groups, minimal and/or moderate for the 1000 and 3000 ppm groups, and in the range of minimal to severe for animals in the 10000 ppm group.

Table 27: 90 day feeding study in mice: Histopathological findings in the liver

	Dose group level (ppm)								
	0	100	300	1000	3000	10000			
Males:									
Liver, total number examined	11	10	10	10	10	15			
hepatocellular hypertrophy	-	1	2	10	10	11			
hepatocellular hyperplasia	-	-	-	-	-	3			
hepatocellular necrosis	-	4	3	8	10	10			
inflammation, chronic	2	1	-	2	2	4			
inflammation, acute	-	4	4	7	10	11			
vacuolization	-	-	1	0	1	7			
anisokaryosis	-	1	5	10	10	11			
cholestasis	-	-	-	-	-	4			
biliary hyperplasia	-	ı	1	1	3	3			
Females:									
Liver, total number examined	10	10	10	10	11	11			
hepatocellular hypertrophy	-	1	4	10	11	10			
hepatocellular hyperplasia	-	-	-	-	-	1			
hepatocellular necrosis	2	2	3	6	7	8			
inflammation, chronic	2	4	2	5	3	8			
inflammation, acute	3	3	3	5	7	8			
vacuolization	-	-	1	-	-	7			
anisokaryosis	-	3	8	9	11	11			
cholestasis	-	-	-	-	-	-			
biliary hyperplasia	-	-	-	-	-	2			

Conclusion:

Treatment with 10000 ppm metaldehyde in the diet resulted in the death of 5 males and 1 female within the first eight days of treatment. Effects on the liver were observed in all treatment groups (100 ppm and above): increased liver weight, swelling of the liver, hepatocellular hypertrophy, hyperplasia, necrosis, inflammation, anisokaryosis, vacuolization, cholestasis and biliary hyperplasia. Increases in body weight in females at 3000 and 10000 ppm were possibly related to the increases in liver weights. The decreases in absolute and relative kidney weights in males of the 10000 ppm treatment group were not associated with microscopic lesions in the kidney. Based on the results of this study, dosage levels of 20, 100 and 300 ppm were selected for a subsequent 18-month oncogenicity study. No NOAEL could be derived from this study. The LOAEL was 100 ppm (equivalent to 19 mg/kg bw/d for males and 24 mg/kg bw/d for females).

4-weeks toxicity (dog)

Reference: 4-week dose-range-finding study for a 52-week chronic toxicity study

of metaldehyde by oral administration via the diet to Beagle dogs

Author(s), year: Leuschner J., 2002

Report/Doc. number: LPT Laboratory of Pharmacology and Toxicology KG, Hamburg,

Germany; LPT Report No. 14543/01

Lonza Report No. 3506, Doc. No.: 532-003

Guideline(s): Dose-range-finding study based on OECD Guideline 452

GLP: Yes

Deviations: Not applicable

Validity: Valid

Material and methods:

Test material Metaldehyde

Lot/Batch 30202
Purity 98.3%
Vehicle Diet
Species Dog
Strain Beagle
Age 7 months

Weight at dosing 6.5 - 10.5 kg (males), 6.4 - 9.6 kg (females)

Source Stefano Morini, S. Polo D'Enza, Italy

A dose-range-finding study was conducted in order to select the dose levels for a 52-week chronic toxicity study of metaldehyde in dogs. The study was divided into two experiments. The first experiment was called "Escalating dose levels" and lasted 22 days in total. In this experiment, metaldehyde was administered orally via the diet to 4 animals (2 males, 3 females as 1 female was sacrificed prematurely) initially at a dose level of 90 mg/kg bw/day and subsequently by escalating dose levels of 30, 60, 75 and 90 mg/kg bw/day. Each of the doses was administered for 3 days followed by a 2-day wash-out period. Due to sings of toxicity at the initial dose of 90 mg/kg bw/day, the dosing was stopped after the first administration and after a regular wash-out period the dosing was continued as described above. The second experiment called "Fixed dose levels" was a four week treatment. One group of animals (2 males, 2 females) received 75 mg/kg bw/day and a second group (2 males, 2 females) received 90 mg/kg bw/day orally via the diet. Parameters evaluated in all animals included body weight, food and drinking water consumption, clinical signs and mortality. Blood samples for haematology and clinical biochemistry were taken from all phase 2 animals (fixed dose levels) fasted overnight before the first administration and at the end of test week 4. Additionally, recording of heart rate (ECG) for all animals of the fixed dose level experiment was carried out on test day 1 (before feeding, and 2h and 4h after start of feeding) and at the end of test week 4 (before feeding and 2h after start of feeding). At the end of the study all surviving animals were allowed to recover. Autopsy and macroscopic inspection of the animal which died prematurely was carried out as soon as possible after exitus.

The substance-food mixture was freshly prepared daily. The metaldehyde concentration was adjusted for each dosing interval or for each week to the weekly mean food consumption per group employing the food consumption data of the previous week. Daily intakes were not calculated.

Due to the low number of animals, a statistical evaluation of the results was not possible.

	v	0 0	
Group	Duration of administration (days)	Dose in mg/kg bw/day via diet	Number and sex of animals
	Phase 1:	Escalating dose levels	
1	1*	90	2 males
	3-day wash	n-out period	
	3	30	2 females
	2-day wash	n-out period	
	3	60	
	2-day wash	i-out period	
	3	75	
	2-day wash	i-out period	
	3	90	
	Phase	2: Fixed dose level	
2	28	75	2 males, 2 females
3	28	90	2 males, 2 females

Table 28: Study design of the dose-range-finding study

Findings:

Clinical signs and mortality: Due to signs of toxicity at the intended lowest dose level of 90 mg/kg bw/day the dosing was stopped after the first administration. The following clinical symptoms were observed: Ataxia and clonic convulsions were noted in both male and female animals. Additionaly, both female animals exhibited emesis. In female animal no. 3, lateral position, difficulty in breathing and shaking of the head were noted 4 hours after application. The animal had to be sacrificed prematurely in moribund condition approximately 6 hours after application. Macroscopic inspection at necropsy revealed no substance-related pathological findings.

During the <u>escalating dose period</u> no behavioural changes were noted following the 3-day administration of 30 mg/kg bw/day. Reduced motility was observed in both male and female animals and clonic convulsions in one male and two females on the first day of administering 60 mg/kg bw/day. In addition, increased respiratory rate and emesis were noted in one female animal. The 3-day treatment with 75 mg/kg bw/day led to reduced motility in one male and one female animal on two or all three administration days and to tonoclonic convulsions in one male and one female animal on two or all three administration days. Emesis was also observed in one female on all three administration days. A further increase in the dose to 90 mg/kg bw/day led to ataxia in all animals and to occasional or regular slight tremor in one male and the two females during the treatment days. In addition, tonoclonic convulsion and emesis were observed in all female animals and salivation in one female animal during the application period. Abdominal position was noted in both females on the first administration day. None of the animals of the escalating dose phase starting at 30 mg/kg bw/day died.

In the <u>fixed dose level experiment</u>, treatment with 75 mg/kg bw/day led to slight tremor, inflated stomach, emesis, mydriasis and increased respiratory rate of both male and female dogs. All the effects were observed starting 1-2 hours after administration and lasted up to 6 hours. The intensity of the symptoms subsided with time. No signs of toxicity were observed from day 7 onwards.

Treatment with 90 mg/kg bw/day led to slight tremor, clonic and tonoclonic convulsions, inflated stomach, emesis, increased respiratory rate, slight ataxia, lateral position, abdominal position, moderate salivation, pale gingival and mydriasis of both male and females starting on day 1 onwards. All the effects were observed starting from 20-60 min after administration and lasted up to

^{*} Due to signs of toxicity at the intended lowest dose level of 90 mg/kg bw/day, the dosing was stopped after the first administration. One female animal had to be sacrificed prematurely in a moribund condition approximately 6 hours after application. The study was continued starting with a new lowest dose level of 30 mg/kg bw/day and another female animal to keep a group size of 2 males and 2 females.

6 hours. Overall, the severity of all symptoms declined with time and the signs of toxicity had almost disappeared towards the end of the 4-week treatment. No mortality occurred in the fixed dose level experiment.

<u>Body weight</u>: No substance-related effect was measured in the escalating dose experiment. In the fixed dose level experiment, male dogs treated with 90 mg/kg bw/day revealed a slight reduction of body weight gain approximately -5% relative to the 75 mg/kg bw/day dose group.

<u>Food and water consumption</u>: Both were not influenced by administration of the test substance in both experiments.

<u>Electrocardiography</u>: The visual assessment of the electrical complexes of the ECG and the determination of the heart rate did not show any substance-related changes at 75 or 90 mg/kg bw/day during the 28-day treatment.

<u>Haematology</u>: No substance-related influence was observed for haemoglobin content, number of erythrocytes and leucocytes, differential blood count, haematocrit value, platelet and reticulocyte counts, thromboplastin time, activated partial thromboplastin time, erythrocyte sedimentation rate, MCV, MCH and MCHC.

<u>Clinical biochemistry</u>: In test week 4, increased cholesterol, creatinine and glucose levels compared to pre-dose levels were found in all male and female dogs. However, no dose-relationship was observed and no concurrent control group was included in this test. Thus no conclusion on treatment-relationship is possible.

There was no influence on bile acids, bilirubin (total), protein (total), protein electrophoresis, urea (in blood), calcium, chloride, potassium and sodium. The activity of ALAT/GPT, ALP, ASAT/GOT and LDH was not increased compared to pre-dose levels.

Conclusion:

On the basis of the results of this study, dose levels of 10, 30 and 90 mg/kg bw/day were proposed for the 52-week main study by the study director.

26-weeks toxicity (dog)

Reference: 26-weeks-toxicity of metaldehyde 99% - called "Metaldehyd" - in

Beagle dogs after oral administration

and

Supplement No. 1 for 26-weeks-toxicity of metaldehyde 99% - called

"Metaldehyd" - in Beagle dogs after oral administration

Author(s), year: Neumann W., 1980

and

Neumann W., 1991

Report/Doc. number: LPT Laboratory of Pharmacology and Toxicology KG, Hamburg,

Germany; LPT Lonza Report No. 1379 Part 1, Doc. No.: 533-001

and

Lonza Report No. 1379 Part 2, Doc. No.: 533-001

Guideline(s): No, study was performed before adoption of OECD Guideline 452

(adopted May 1981)

GLP: No, study was performed before implementation of GLP

Deviations: Not applicable

Validity: Valid

Reference: Histological re-examination of the testes and re-evaluation of the

findings of the 26-week toxicity of metaldehyde 99% in Beagle dogs after oral administration (LPT Study report dated March 31, 1980)

Author(s), year: Leuschner J., 2009

Report/Doc. number: LPT Laboratory of Pharmacology and Toxicology, Hamburg, Germany;

LPT Report No. 24158, Doc. No.: 581-005

Guideline(s): Not applicable, as the report is a histological re-examination and re-

evaluation of samples generated in the study of Neumann W., 1980

GLP: Not applicable Deviations: Not applicable

Validity: Valid

Material and methods:

Test material Metaldehyde

Lot/Batch No batch number; test material obtained from current production in January

1979

Purity >99%
Vehicle Diet
Species Dog
Strain Beagle
Age 8 months
Weight at dosing Not reported

Source Chr. Fred Leuschner & Co., Laboratory of Toxicological and

Pharmacological Examinations, Loehndorf/Post Wankendorf, Germany

Metaldehyde was administered orally via diet to male and female Beagle dogs (6 dogs/sex/dose) to achieve dose levels of 0, 20, 60 and 90 mg/kg bw/day for a period of 26 weeks. All animals were adapted to laboratory condition for a period of four weeks prior to study initiation. The dose levels were set on the basis of a 14-day preliminary test (1979) using 1 male and 1 female per dose at the dose levels of 90 and 180 mg/kg bw/day. No intolerance was observed at 90 mg/kg bw/day but the

dose level of 180 mg/kg bw/day caused vomiting and was within the lethal range. The test substance was admixed to the food homogeneously every day for each dose group. 50 g/kg bw/day of the food was offered to each dog for one hour between 8 and 9 a.m. In case of animals with poor appetite, the food was served for a longer period up to 8 hours. The residue was removed and weighed.

The following examinations were carried out:

- Behaviour including examination of general reflexes and external appearance, faeces: daily
- Food consumption, drinking water: daily
- <u>Body weight</u>: once a week (for the calculation of the daily dose of the test compound, the weight was determined daily without registration)
- <u>Haematology</u>: before the first administration and after 4, 8, 16, 20 and 26 test weeks in all animals: haemoglobin, erythrocytes, leucocytes, differential blood count, haematocrit value, thromboplastin time, erythrocyte sedimentation rate, blood clotting time, platelets, reticulocytes
- <u>Clinical biochemistry</u>: before the first administration and after 4, 8, 16, 20 and 26 test weeks in all animals: ALAT, ASAT, ALP, blood urea, glucose, sodium, potassium, calcium, chloride, total protein, uric acid, total bilirubin, albumin, globulin, liver function, creatinine, free cholesterol, total cholesterol, non-esterified fatty acids, esterified fatty acids, LDH, direct bilirubin, gamma-GT
- <u>Urinalysis</u>: before the first administration and after 8, 16 and 26 test weeks in all animals: colour, specific weight, protein, glucose, bilirubin, haemoglobin, ketone bodies, pH, urinary sediment
- <u>Electrocardiography</u>: on the first test day and after 13 and 26 weeks in all animals; examination before and 2 hours after administration on the lying animal in the limb leads I-III; limb lead II was evaluated and heart rate was determined
- Examination of circulary functions: after 26 test weeks in the animals of the highest dose group and the control; the dogs were kept under narcosis and after surgical interventions, systolic and diastolic pressure in general and pulmonary circulation were measured before and after administration of norepinephrine.
- Ophthalmological, auditory and dental examinations: before the first administration and after 4, 8, 16, 20 and 26 test weeks in all animals: ophthalmological examination included cornea, anterior chamber, pupil, lens, vitreous body, fundus of the eye; the auditory check was done with a simple noise test
- Macroscopic and microscopic evaluation: after 26 weeks in all animals; macroscopic inspection; organ weights: heart, liver, lungs, spleen, kidneys, adrenals, thymus, pituitary, gonads, thyroid, brain; histopathological examination: heart, lungs, liver, spleen, kidneys, adrenals, thymus, pituitary, gonads, thyroid, brain, prostate/uterus, stomach, duodenum, jejunum, ileum, colon, rectum, salivary glands, eye, urinary bladder, bone marrow, trachea, aorta, oesophagus, pancreas, lymph node, peripheral nerve, skeletal muscle, skin, tongue, spinal cord, gall bladder, bone, mamma
- <u>Statistical evaluation</u>: Analysis of variance and Student's t-test were carried out. Limits of significance were p ≤ 0.01

• <u>Timeframe</u>: the experiments were performed between June 15th 1979 and December 15th 1979

In a supplement to the main report, a re-evaluation of the macroscopic and microscopic pathology data of liver, testes and prostate and reformatted individual animal data were presented.

Findings:

- Behaviour including examination of general reflexes and external appearance, faeces: At all three dose levels, behaviour and external appearance were unchanged during 26 weeks of treatment. The neurological examinations did not lead to pathological findings. Pupillary light reflex, corneal reflex, patellar reflex, flexor, extensor and postular reflexes were not changed.
- <u>Food consumption, drinking water</u>: Food and drinking water consumption were within the normal range during the 26 week test period. The real intake of metaldehyde is shown in the following table:

Table 29: Real intake of metaldehyde in the 26-week dog study (mg)

Group	Males	Females
20 mg/kg bw/day	20.2 ± 0.4	19.7 ± 2.8
60 mg/kg bw/day	61.5 ± 4.1	62.2 ± 5.6
90 mg/kg bw/day	91.8 ± 7.8	86.7 ± 3.8

- Body weight: no effect on body weight or body weight gain
- <u>Haematology, clinical biochemistry and urinalysis</u>: no effects of test substance were found
- <u>Electrocardiography and examination of circulatory functions</u>: No effect attributable to the test substance was observed in the ECG. The minimal changes in the functional picture were within the normal range and explained by the fact that the ECG in the dogs could not be recorded in a state of actual rest. At all tested dose levels systolic and diastolic pressure was unchanged in both general and pulmonary circulation.
- Ophthalmological, auditory and dental examinations: no effects were observed during the test period
- Macroscopic and microscopic evaluation:

Results of macroscopic evaluation, organ weighing and histopathology were presented in the original report. However, due to the request from US EPA, a Supplement of the study (Neumann W., 1991) was provided including detailed individual tables, redesigned summary tables and information on the severity and extent of all lesions described. A review on liver, testes and prostate findings was undertaken at request from the sponsor, which basically confirmed the findings of the original report. No effects were noted at macroscopic evaluation and organ weighing.

Histopathology:

<u>Liver</u>: The overall evaluation of the histopathological findings in the liver did not show any treatment-related effect. The few very slight lesions recorded are considered to be spontaneous findings occurring in dogs.

<u>Testes</u>: Diffuse atrophy of the testes was found only in the mid (2/6 males) and high dose group (4/6 males). The atrophy seen in one control animal was a consequence of

cryptorchism. One animal from the low dose group had very slight focal atrophy of the tubules. This was considered a spontaneous finding because it occurred also in one untreated control animal. The dose level of 20 mg/kg bw/d was therefore considered the NOAEL for the testes effects.

<u>Prostate</u>: Diffuse atrophy of the prostate was found in the mid dose group (4/6 males) and the high dose group (2/6 males). One male from the high dose group also showed focal atrophy. No such effects were seen in the control or low dose group.

Mesenteric lymph nodes: Examination of the lymph nodes revealed a considerable infestation with parasites, especially in the higher dose groups, which could have possibly contributed to the higher incidences of findings at these dose levels. For precautionary reasons a clear NOAEL of 20 mg/kg bw/day was set.

Table 30: Histopathological findings in the 26-week dog study (according to Supplement 1 of the original study, Neumann W., 1980, 1991)

		ntrol	bw/	ng/kg /day	bw/	ng/kg /day	bw/	ng/kg /day
	3	2	3	2	3	2	3	9
LIVER								
examined	6	6	6	6	6	6	6	6
no abnormalities detected	5	4	5	2	4	5	4	2
very slight single cells necrosis with	0	2	1	3	2	1	1	2
predominantly mononuclear cell								
proliferation								
very slight periportal infiltration of	0	0	0	1	0	0	2	2
lymphocytes, histiocytes and eosinophilic								
granulocytes								
very slight hydropic swelling in focal areas	1	0	0	0	0	1	2	2
(H.E. staining)								
PROSTATE								
examined	6	-	6	=.	6	-	6	-
no abnormalities detected	6	-	6	-	2	-	3	-
atrophy, diffuse	0	-	0	-	4	-	2	-
atrophy, focal	0	-	0	-	0	-	1	-
TESTICLES								
examined	7#	-	6	-	7#	-	6	-
no abnormalities detected	5	-	5	-	2	-	2	-
diffuse atrophy	1	-	0	-	2	-	4	-
focal atrophic tubules	1	-	1	-	3	-	0	-
LYMPH NODE, MES.								
examined	6	6	6	6	6	6	6	6
no abnormalities detected	4	4	4	5	3	2	1	1
follicular hyperplasia	0	0	0	0	0	0	3	3
inflammation - chronic	1	1	1	1	3	3	1	1
erythrocytes (medullary funicle)	1	1	1	0	0	1	0	0
lymphocytes depletion - moderate	0	0	0	0	0	0	2	0
parasitic granuloma	1	1	2	1	3	4	4	2

[#] one animal of this group was examined in both testicles

In the original 26-week-toxicity study of metaldehyde in Beagle dogs (Neumann W., 1980, Doc. No. 533-001), the findings in the testes were not graded according to severity. Thus, a microscopical re-examination of H. & E. stained paraffin and PAS-stained sections of the testes of all 24 male dogs of the 26-week toxicity study was performed in 2009 with the objective to score findings using a current severity grading system. This up-grade was considered necessary to gain direct comparability of findings in the different dog reports (Leuschner J., 2009, Doc. No. 581-005).

Findings of the histological re-examination of the testes and re-evaluation:

Lesions related to the administration of the test substance: Cryptorchism and a moderate atrophy of the testes (1 of 6 animals) and a mild to moderate focal atrophy of the germinative epithelium (3 of 6 animals) were noted for the animals of the 60 mg/kg bw/d dose group. The changes in the 90 mg/kg bw/d dose group consisted of a mild to moderate diffuse atrophy/degeneration of the germinative epithelium. Four of 6 animals were affected, the mean severity grade of the high dose animals for diffuse atrophy of the left and right testes was 1.67 and 1.83 compared to 0.00 and 0.00 in the control. The changes are morphologically clearly distinct from the spontaneous findings in the 20 mg/kg bw/d dose group and in the control group and therefore attributable to the treatment with the test item.

Lesions unrelated to the administration of the test item: A focal minimal to mild atrophy of the germinative epithelium of the testes was noted in control animals and animals from the 20 mg/kg bw/d dose group. Incidence and severity indicate an incidental distribution of this finding for the control and for the low dose treated animals.

Table 31: Histological re-examination of testes (Leuschner J., 2009)

Dose group	Contr	ol	20 mg	/kg bw/day	60 mg	/kg bw/day	90 mg	/kg bw/day
No. animals	6		6		6		6	
	#	SEV	#		#	SEV	#	SEV
Testis (I)			•	•		•		
Atrophy of germin. epith., focal	4	0.67	2	0.50	3	1.33	1	0.17
Atrophy of germin. epith., diff.	0	-	0	-	0	-	4	1.67
Atrophy of testis	1	0.33	0	-	1	0.50	0	-
Cryptorchism	1	-	0	-	1	-	0	-
Testis (II)			•	•		•		
Atrophy of germin. epith., focal	2	0.33	2	0.50	4	1.50	1	0.17
Atrophy of germin. epith., diff.	0	-	0	-	0	-	4	1.83
Testis (I) PAS-stain	•	•		•	•	•	•	
Atrophy of germin. epith., focal	4	0.67	2	0.50	3	1.33	1	0.17
Atrophy of germin. epith., diff.	0	-	0	-	0	-	4	1.67
Atrophy of testis	1	0.33	0	-	1	0.50	0	-
Cryptorchism	1	-	0	-	1	-	0	-
Testis (II) PAS-stain			•				•	
Atrophy of germin. epith., focal	2	0.33	2	0.50	4	1.50	1	0.17
Atrophy of germin. epith., diff.	0	-	0	-	0	-	4	1.67

Severity grading (SEV): 1= minimal, 2= mild, 3= moderate, 4= marked, mean values are presented

Conclusion:

A NOAEL of 20 mg/kg bw/day is set for this 26-week-toxicity study in Beagle dogs based on the histopathological findings in testes (mild to moderate atrophy/degeneration of the germinative epithelium) and prostate (diffuse atrophy). The histopathological re-examination of the testes confirmed the pattern of testicular findings: the severity score of focal findings was increased at 60 mg/kg bw/day while at the high dose level of 90 mg/kg bw/day a clear increase of diffuse atrophic changes was found. The NOAEL for this study was therefore confirmed to be 20 mg/kg bw/day.

52-weeks toxicity (dog)

Reference: 52-week chronic toxicity study of metaldehyde by repeated oral

administration via the diet to Beagle dogs

Author(s), year: Leuschner J., 2003

Report/Doc. number: Lonza Report No. 3657, Doc.No. 537-003, Conducting laboratory: LPT

Laboratory of Pharmacology and Toxicology KG, Hamburg, Germany

Guideline(s): OECD Guideline 452 (1981); EC Guideline L133 Part B, Chronic Toxicity

Test (1988); Japanese MAFF, 12 NohSan No. 8147 (2000)

GLP: Yes
Deviations: No
Validity: Yes

Material and Methods:

Metaldehyde (batch no. 30202, purity: 98.3 %) was administered with the diet to 4 male and 4 female Beagle dogs per treatment group at dose levels of 0, 10, 30 and 90 mg/kg bw/d over a period of 52 weeks. The Beagle dogs (source: Stefano Morini, Reggio Emilia, Italy) were approximately 6 months of age at the beginning of the study and weighed 5.4-9.2 kg (males) and 5.8-9.2 kg (females). The amount of the test substance given was adjusted to each animal's actual body weight and mean food consumption weekly. The test substance-diet mixture was freshly prepared daily. The diet was checked for stability, concentration and homogeneity at the beginning of the study and thereafter every 3 months. The dose levels for this study were selected based on the results of a 4-week dose finding study in dogs.

All animals were checked at least once daily for clinical signs. These observations included skin/fur, eyes, mucous membranes, respiratory and circulatory systems, somatomotor activity and behaviour patterns. Mortality was checked twice daily, and as soon as possible after exitus, postmortem examination was performed. Body weights were recorded at study initiation and thereafter in weekly intervals. Food consumption was recorded on a daily basis throughout the experimental period. The report included weekly mean values. Daily monitoring by visual appraisal of the drinking water consumption was maintained throughout the study. In case of food not consumed during the first part of the daily feeding, the quantity of remaining food was recorded and the amount of test substance intake of the animal was corrected accordingly. The report includes weekly mean values of uptake of the test substance. Blood samples were taken from animals fasted overnight at the beginning of the study and at the end of test weeks 13, 26 and 52. Haematology included erythrocytes, hematocrit, haemoglobin, MCV, MCH, MCHC, leucocytes, differential blood count, reticulocytes, platelets, thromboplastin time, activated partial thromboplastin time and erythrocyte sedimentation rate. The following clinical chemistry parameters were determined: albumin, albumin/globulin ratio, total bilirubin, total cholesterol, creatinine, glucose, total protein, urea in blood, calcium, chloride, potassium, sodium, alanine amino transferase (ALAT), aspartate amino transferase (ASAT), alkaline phosphatase (ALP), gamma-glutamyl-transferase (GGT), triglycerides and inorganic phosphorus. Urine was collected from all animals in a metabolism cage in the morning of the appropriate test day 3 hours prior to the administration after the dogs received 50 mL tap water/kg bw orally by gavage. Urinalysis was performed at the beginning of the study and at the end of test weeks 13, 26 and 52 and included volume, pH, specific gravity, protein, glucose, bilirubin, urobilinogen, ketones, haemoglobin, nitrite, color and microscopic examination. Ophthalmological and auditory examinations were performed prior to the first administration and during test weeks 13, 26 and 52. All animals were examined at necropsy. The weights of the following organs were determined: adrenals, brain, heart, kidneys, liver, lungs, ovaries, pituitary,

prostate, spleen, testes and thyroids including parathyroids. The following organs and tissues were examined histopathologically: adrenals, aorta abdominalis, bone (sternum and os femoris with joint), bone marrow (sternum and femur), brain (transverse section through optic chiasma, infundibulum, midbrain, brain stem and cerebellum), caecum, epididymis, eye with optic nerve, gall bladder, gross lesions, heart (3 levels: left and right ventricle, septum), small intestine (duodenum, jejunum, ileum), large intestine (colon, rectum), kidney and ureter, lacrimal gland, liver, lungs (with mainstream bronchi and bronichioles), lymph node (cervical, mesenteric), mammary gland, muscle (skeletal, thigh and tibia), nerve (sciatic and tibial), esophagus, ovaries, pancreas, pituitary, prostate, salivary glands (mandibular, parotid, sublingual), seminal duct, skin (left flank), spinal cord (cervical, thoracic, lubar) incl. spinal ganglion and spinal root, spleen, stomach, testes, thymus, thyroids incl. parathyroids, tissues masses or tumours, trachea incl. larynx, urinary bladder, uterus (incl. cervix and oviducts) and vagina.

Findings:

General observations: Clinical signs were only observed in the highest dose group from study week 1 onwards. The following symptoms were noted: ataxia, reduced motility, emesis, tremor, twitching and salivation. Individual to all animals were affected. Incidence and severity appeared to decline from study week 19 onwards with none to two animals only affected.

<u>Mortalilty</u>: 1 male and 1 female from the 30 mg/kg dose group and 1 female of the 90 mg/kg dose group were found dead between study days 260 and 322. The deaths are regarded to be test-substance related. No premortal symptoms were recorded.

<u>Body weight</u>: The body weight of the treated animals was not influenced compared to controls. Body weight gain appeared to be reduced in the high dose group, however, no statistical significance was noted.

<u>Food consumption</u>: During some weeks of the study there was a slight statistically significant increase of food intake noted for male and female animals of the high dose group. This effect might be due to the slight decrease in body weight gain noted in this dose group. No overt intergroup differences in water consumption were found during the study.

be due to the	slight decrease in	body weight gain noted in this dose group. No overt intergroup							
lifferences in water consumption were found during the study.									
Table 32:	Table 32: 52 week feeding study in dogs Body weight / body weight gain								
		Dose group level (mg/kg bw/d)							

	Dose gr	Dose group level (mg/kg bw/d)								
	Males	Males				Females				
	0	10	30	90	0	10	30	90		
Body weight (kg)										
- week 0	7.70	7.55	7.55	8.00	7.33	6.85	6.83	6.70		
- week 52	13.33	11.43	11.77	11.00	10.83	10.68	10.07	9.13		
Body weight gain										
- absolute (kg)	5.63	3.88	4.22	3.00	3.5	3.83	3.24	2.43		
- relative (%)	73	51	56	38	48	56	47	36		

Ophthalmoscopic examination: There were no treatment-related effects observed.

<u>Auditory examination</u>: In test week 13 one male and one female dog treated with 90 mg/kg did not react to the noise test. None of the animals of this dose group was affected in test weeks 26 or 52. There was no indication of any impairment in animals of lower dose groups.

<u>Haematology</u>: Test substance-related changes were seen as an increase in haemoglobin, erythrocytes and haematocrit of animals of the highest dose groups. This effect was observed in both sexes in study week 13 (statistically significant only for females) and in females only in study week 26 (not longer statistically significant). At the end of the study, no changes were noted for any

of the haematological parameters. The decrease of activated thromboplastin time in males of the highest dose group in week 13 is regarded as spontaneous and within the normal variability.

Table 33: 52 week feeding study in dogs Relevant haematological findings

	Dose group level (mg/kg bw/d)							
	Males			Female	Females			
	0	10	30	90	0	10	30	90
Erythrocytes (1012/L)								
- Week 0	5.38	5.45	5.23	5.33	5.35	5.18	5.40	5.60
- Week 13	5.45	5.85	5.45	6.23	5.73	5.70	6.08	7.15**
- Week 26	5.70	6.13	5.65	5.55	5.48	5.58	5.70	6.35
- Week 52	6.43	6.78	6.60	6.25	6.45	6.18	7.13	6.27
Hemoglobin (mmol/L)								
- Week 0	7.33	7.48	7.13	7.10	7.23	6.95	7.28	7.45
- Week 13	7.50	8.08	7.58	8.88	7.95	7.80	8.28	9.90**
- Week 26	7.95	8.70	8.00	8.28	7.93	7.78	7.90	8.95
- Week 52	9.00	9.30	9.33	9.18	9.18	8.58	9.80	8.47
Hematocrit (%)								
- Week 0	38.3	38.5	36.8	36.8	37.0	35.3	37.3	38.3
- Week 13	36.3	39.3	37.3	43.0	39.0	38.0	40.8	49.3**
- Week 26	38.5	42.0	38.8	39.3	38.0	37.5	37.5	43.0
- Week 52	43.0	45.3	45.0	43.8	43.8	41.0	47.0	40.7

^{** (}p< 0.01); significantly different from control group

<u>Clinical chemistry</u>: In the highest dose group, test substance-related changes were noted for bilirubin (females, week 13), triglycerides (females, weeks 13 and 26) and AP levels (both sexes, weeks 13, 26 and 52).

Table 34: 52 week feeding study in dogs Relevant clinical chemistry findings

	Dose group level (mg/kg bw/d)							
	Males			Females				
	0	10	30	90	0	10	30	90
Bilirubin (µmol/L)								
- Week 0	3.73	2.83	3.68	3.23	2.78	3.30	2.95	3.28
- Week 13	2.48	2.30	2.95	2.85	2.53	2.33	2.95	3.98**
- Week 26	3.30	3.38	3.00	2.60	3.00	3.05	3.33	3.35
- Week 52	3.88	3.65	3.53	3.73	3.43	3.38	3.27	3.93
Triglycerides								
(mmol/L)	0.243	0.215	0.218	0.200	0.195	0.190	0.180	0.200
- Week 0	0.265	0.220	0.285	0.258	0.238	0.260	0.195	0.490**
- Week 13	0.305	0.280	0.258	0.283	0.235	0.340	0.245	0.553**
- Week 26	0.230	0.215	0.223	0.278	0.208	0.235	0.217	0.280
- Week 52								
AP (U/L)								
- Week 0	250	258	287	302	259	235	240	220
- Week 13	143	149	174	296**	147	135	138	236
- Week 26	85	102	108	221**	79	97	79	156
- Week 52	71	94	94	216**	66	79	70	151**

^{** (}p< 0.01); significantly different from control group

<u>Urinalysis</u>: No substance-related changes were noted at urinalysis. The slightly decreased specific gravity of the urine noted for the intermediate males in week 52 is regarded as fortuitous as no dose-relationship was noted. The slightly decreased pH-value observed for high dose males in week 52 is regarded to be within the normal variability.

<u>Necropsy</u>: The macroscopic necropsy examination during dissection revealed no findings in the organs and tissues in any treatment group. The gross findings such as indurated, emphysematous

and/or discoloured lungs were observed in 2 mid dose animals and 1 high dose animal and were regarded as spontaneous.

<u>Organ weights</u>: A treatment-related increase of absolute and relative liver weights was found in males and females of the mid and high dose groups whith statistical significance achieved only at the high dose group.

Table 35: 52 week feeding study in dogs Absolute and relative (% of bodyweight) organ weights

	Dose g	Dose group level (mg/kg bw/d)						
	Males	Males			Females			
	0	10	30	90	0	10	30	90
Liver								
- absolute	430	360	457	541	351	350	362	418
- relative (% bw)	34.2	33.9	41.2	51.6**	33.6	34.5	38.1	48.8**

^{** (}p< 0.01); significantly different from control group

<u>Histopathology</u>: The histopathological findings in testes were re-examined and discussed in the "Expert statement on the histological findings (giant cells, atrophy and degeneration of the germinative epithelium) in the 52-week toxicity study in Beagle dogs with metaldehyde" (Leuschner J., Drommer W., 2006), see below.

Mild atrophy of the prostate was noted in 3 of 4 male animals of the high dose group and 1 of 4 animals from the mid dose group.

No treatment-related findings were noted in any other organ examined. In particular, no changes were noted in the spinal cord with ganglion/root (cervical, thoracal and lumbar region). A minimal fatty infiltration of hepatocytes without degeneration was observed in the liver of control and test animals. None of the histopathological findings in the male of the mid dose group which died prematurely could be seen as a cause of death. Both females which died prematurely (1 from the mid dose and 1 from the high dose group) revealed microscopic findings in the lungs: moderate interstitial pneumonia and moderate bronchopneumonia were found which might have contributed to the deaths of the animals, although they seemed to be not directly related to the test substance. The study authors regarded the premature deaths of these animals as substance-related.

Table 36: 52 week feeding study in dogs Relevant histopathological findings (testes and prostate)

	0 mg/kg	10 mg/kg	30 mg/kg	90 mg/kg
Testis I (not reported if left or				
right testis)				
no. examined	4	4	41)	4
Atrophy of the germinal	-	-	2	2
epithelium			(no.18	(no.25 mild
(animal no., grade)			moderate	no.27 marked)
			no.20 mild)	
Degeneration of the germinal	-	1	-	3
epithelium		(no.10 mild)		(no.26 marked
(animal no., grade)				no.27 mild
				no.28 mild)
Giant cells	-	2	1	2
(animal no., grade)		(no.10 mild,	(no.18 mild)	(no.25
		no.12 minimal		minimal no.28
)		mild)
Juvenile testis	-	1	-	-
(animal no.)		(no.12)		
Testis II (not reported if left or				
right testis)				

	0 mg/kg	10 mg/kg	30 mg/kg	90 mg/kg
no. examined	4	4	41)	4
Atrophy of the germinal epithelium (animal no., grade)	-	-	(no.18 minimal no.20 minimal)	3 (no.25 mild no.27 marked no.28 moderate)
Degeneration of the germinal epithelium (animal no., grade)	-	1 (no.10 mild)	-	3 (no.26 marked no.27 mild no.28 mild)
Giant cells (animal no., grade)	-	1 (no.10 mild)	-	2 (no.25 mild no.28 minmal)
Juvenile testis (animal no.)	-	1 (no.12)	-	-
Prostate				
no. examined	4	4	41)	4
Atrophy (animal no., grade)	-	-	1 (no.19 mild)	3 (no.25 mild no.26 mild no.28 mild)
Lymphocytic infiltration (animal no., grade)	(no.3 minimal)	1 (no.9 moderate)	-	-
Suppurative prostatitis (animal no., grade)	-	1 (no.9 minimal)	-	-
Autolysis (animal no., grade)	-	-	1 (no18 moderate)	-
Cysts (animal no., grade)	-	-	1 (no.19 mild)	1 (no.27 minimal)

Severity grading: minimal, mild, moderate, marked

1) including animal no. 18 which died prematurely. In animal no. 18, atrophy of the germinal epithelium in both testes (minimal to moderate), giant cells in testis I (mild), and autolysis in the prostate (moderate) were recorded.

Conclusion:

Clinical signs of toxicity (ataxia, reduced motility, emesis, tremor, twitching, salivation) were observed at the high dose group (90 mg/kg) from study week 1 onwards with incidence and severity declining over the time. 1 male and 1 female animal from the mid dose group (30 mg/kg) and 1 female from the high dose group were found dead between study day 260 and 322. As no obvious cause of death could be determined, the deaths were considered to be related to treatment. A reduction of body weight gain was noted in high dose animals but without statistical significance. Also in the high dose group, 1 male and 1 female animal did not react to the noise test in week 13 but were not affected in week 26 and 52. Red blood cell parameters (increased erythrocytes, haemoglobin and hematocrit) were affected predominantely in females at the highest dose group. The effects were strongest at week 13 but declined until week 26 and were no longer seen at week 52. A similar situation was observed for bilirubin and triglyceride values in females of the high dose group, which were increased at week 13 but returned to normal until the end of the study. In contrast, elevated levels of alkaline phosphatase were observed in males and females at weeks 13, 26 and 52. Relative and absolute liver weights were increased in both sexes of the high dose group. In histopathology, in one animal of the mid dose group and 3 animals of the high dose group mild atrophy of the prostate was observed. The histopathological findings in testes and setting of the NOAEL is reported below.

Reference: Expert statement on the histological findings (giant cells, atrophy and

degeneration of the germinative epithelium) in the 52-week toxicity

study in Beagle dogs with metaldehyde

Author(s), year: Leuschner J., Drommer W., 2006

Report/Doc. number: LPT Laboratory of Pharmacology and Toxicology, Hamburg, Germany;

LPT Report No. 15050/01, Doc. No.: 581-001

Guideline(s): Not applicable, as the report is a histological re-examination and re-

evaluation of samples generated in the 52-week study in dogs (Leuschner

J., 2003)

GLP: Not applicable Deviations: Not applicable

Validity: Valid

Findings:

A complete picture of the results of the original 52-week toxicity study in dogs is presented in the DAR. In this Report, an additional table is included describing the clinical signs observed at the beginning of this 52-week toxicity study in detail. Furthermore, the histopathological findings in testes are discussed for the setting of the NOAEL in this 52-week toxicity study.

General observations/clinical signs

Clinical signs were only observed in the highest dose group (90 mg/kg bw/d) from study week 1 onwards. The following symptoms were noted: ataxia, reduced motility, emesis, tremor, twitching and salivation. Individual to all animals were affected. Incidence and severity appeared to decline from study week 19 onwards with none to two animals only affected.

Table 37: Clinical signs observed during Week 1 in the 52-week dog study in the highest dose group of 90 mg/kg bw/d

Animal number, sex	Observed on test day	Total number of days	Observation
25 m	1, 3-7	6	Ataxia
	2-5, 7	5	Tremor
	1-7	7	Emesis
	1-4	4	Salivation
	1-5, 7	6	Twitching
26 m	1, 3-5, 7	5	Ataxia
	1-7	7	Tremor
	1-5	5	Emesis
	1-2	2	Salivation
	2-7	6	Twitching
27 m	1-7	7	Ataxia
	1-5, 7	6	Tremor
	1-5	5	Emesis
	2	1	Salivation
	2-7	6	Twitching
	2-3	2	Lateral position
28 m	1, 3-7	6	Ataxia
	2-7	6	Tremor
	1-2, 6-7	4	Emesis
	1-2	2	Salivation
	1-5, 7	6	Twitching
	1	1	Lateral position
29 f	1-7	7	Ataxia
	1-7	7	Tremor

	1-3, 5-7	6	Emesis
	1	1	Salivation
	1-4, 6-7	6	Twitching
30 f	1-4, 6-7	6	Ataxia
	1-7	7	Tremor
	1-7	7	Emesis
	1	1	Salivation
	1-7	7	Twitching
31 f	1-4, 6-7	6	Ataxia
	1-7	7	Tremor
	7	1	Emesis
	1-7	7	Twitching
32 f	6-7	2	Ataxia
	1-7	7	Tremor
	4-6	3	Emesis
	1-7	7	Twitching

Histopathological findings in testes

Table 38: Atrophy and/or degeneration of the germinative epithelium in the testes (Leuschner J., 2003- see table 35)

	0 mg/kg	10 mg/kg	30 mg/kg	90 mg/kg
Testis I (not reported if left or right testis)				
no. examined	4	4	41)	4
Atrophy of the germinal epithelium (animal no., grade)	-	-	2 (no.18 moderate no.20 mild)	2 (no.25 mild no.27 marked)
Degeneration of the germinal epithelium (animal no., grade)	-	1 (no.10 mild)	-	3 (no.26 marked no.27 mild no.28 mild)
Giant cells (animal no., grade)	-	(no.10 mild, no.12 minimal)	1 (no.18 mild)	2 (no.25 minimal no.28 mild)
Juvenile testis (animal no.)	-	1 (no.12)	-	-
Testis II (not reported if left or right testis) no. examined	4	4	41)	4
Atrophy of the germinal epithelium (animal no., grade)	-	-	2 (no.18 minimal no.20 minimal)	3 (no.25 mild no.27 marked no.28 moderate)
Degeneration of the germinal epithelium (animal no., grade)	-	1 (no.10 mild)	-	3 (no.26 marked no.27 mild no.28 mild)
Giant cells (animal no., grade)	-	1 (no.10 mild)	-	2 (no.25 mild no.28 minmal)
Juvenile testis (animal no.)	-	1 (no.12)	-	-

Severity grading: minimal, mild, moderate, marked

Upon request of the notifier the histopathological slides of the testes were re-examined by the study pathologist in order to verify the original diagnosis. The results of the re-examination were

¹⁾ including animal no. 18 which died prematurely.

presented in an expert statement. In this paper it was explained that the histological changes of testes seen at the highest dose (90 mg/kg), where all animals were affected, consisted of mainly moderate to marked diffuse atrophy and/or degeneration of the germinative epithelium. At the two lower dose levels (10 and 30 mg/kg), findings were more <u>focal</u> in nature and <u>of minimal to mild severity</u>.

Concerning the "moderate severity" finding in one animal (No. 18), which died premature, of the mid dose (30 mg/kg), it was explained that the finding was limited to only one testis (the other testis showed only minimal focal atrophy) and was therefore regarded to be a spontaneous change.

Furthermore, it is stated in the position paper that the histopathological findings seen in testes at 10 and 30 mg/kg bw/day are of spontaneous nature and morphologically distinct from the treatment related findings seen at 90 mg/kg bw/day. Historical background data for atrophy and degeneration of the germinative epithelium performed at LPT Laboratory of Pharmacology and Toxicology KG, Hamburg, Germany during the years 2003-2006 were made available, which should demonstrate that incidences observed at the low and mid dose level were comparable to historical control data.

In conclusion, it was proposed in the position paper to set the NOAEL at 10 mg/kg bw/d based on mortality occurring at 30 and 90 mg/kg bw/d.

Table 39: Historical background data of atrophy and/or degeneration of the germinative epithelium in the testes of control animals (Leuschner J., Drommer W., 2006)

Finding	Study 1	Study 2	Study 3	Study 4	Study 5	Study 6
Testis I (not reported if left or right testis) no. examined	4	5	5	5	5	4
Atrophy of the germinal epithelium	0	0	0	1	1	0
Degeneration of the germinal epithelium	0	0	0	0	0	0
Giant cells	1	0	0	1	1	1
Testis II (not reported if left or right testis) no. examined	4	5	5	5	5	4
Atrophy of the germinal epithelium	0	0	0	3	3	0
Degeneration of the germinal epithelium	0	0	0	0	0	1
Giant cells	2	2	0	1	1	1

Conclusion:

Considering the historical control data on testes findings, the NOAEL of the 52-week study is set at 10 mg/kg bw/d based on testes findings (atrophy of the germinal epithelium) and mortality observed at 30 mg/kg.

4.7.1.2 Repeated dose toxicity: inhalation

No data available.

4.7.1.3 Repeated dose toxicity: dermal

In a 21-day dermal toxicity study in New Zealand White rabbits, no substance-related effects were observed up to the highest dose tested of 1000 mg/kg bw/d. Signs of local irritation occurred in control and treatment groups and were attributed to the dosing and occlusion procedures. The NOAEL for repeated dermal toxicity in rabbits was 1000 mg/kg bw/d.

4.7.1.4 Repeated dose toxicity: other routes

No data available.

4.7.1.5 Human information

Not available.

4.7.1.6 Other relevant information

Not available.

4.7.1.7 Summary and discussion of repeated dose toxicity

Rat, oral

28 days:

Sprague-Dawley CD rats received diets containing 0, 2500, 5000, 10000 and 20000 ppm metaldehyde (equivalent to 0, 197, 382, 761 and 1547 mg/kg bw/d for males; 0, 233, 454 and 875 mg/kg bw/d for females). Continuous treatment of rats with metaldehyde at dietary concentrations of 2500 ppm or more over 28 days caused systemic toxicity as shown by increased liver weight and hepatocellular hypertrophy. At the highest dose level of 20000 ppm, mortality occurred in 100% of females and 40% of males. Death occurred in the females during the first week of treatment while no males died prior to day 6. Also, 60% of the females receiving 10000 ppm died (between day 4 and 21) or were sacrificed in a moribund condition. In most cases, animals were sacrificed when found with hind limb paralysis/paresis presumably resulting from convulsions since these findings were associated with spinal fracture/luxation and pinching of the spinal cord. Changes in haematology (decrease in erythrocytes) were found in males at 10000 ppm and above. In addition, transient reduction of body weight gain and food consumption was noted at higher dose levels. A NOAEL could not be determined in this study. The LOAEL was 2500 ppm (equivalent to 197 mg/kg bw/d for males and 233 mg/kg bw/d for females).

90 days:

Sprague-Dawley rats received dietary concentrations of 0, 250, 750 and 2500 ppm metaldehyde (equivalent to 0, 21, 65 and 215 mg/kg bw/d for males and females) for 90 days. The predominant treatment-related effects were found in the liver. Centrilobular hepatocyte enlargement was observed at 750 ppm and above while liver enlargement and increased relative liver weight were noted in individual animals only in the high dose group (2500 ppm, not statistically significant). Females receiving 2500 ppm showed a reduction of body weight gain and food efficiency during the first week of treatment which resulted in a reduced (not statistically significant) terminal body weight. In conclusion, the NOAEL is considered to be 250 ppm (equivalent to 21 mg/kg bw/d for both sexes).

Mouse, oral

90 days:

CD-1 mice received 0, 100, 300, 1000, 3000 and 10000 ppm per diet (equivalent to 0, 19, 54, 178, 560 and 1919 mg/kg bw/d for males; 0, 24, 70, 235, 743 and 2996 mg/kg bw/d for females). Treatment with 10000 ppm metaldehyde in the diet resulted in the death of 5 males and 1 female within the first eight days of treatment. Effects on the liver were observed in all treatment groups (100 ppm and above): increased liver weight, swelling of the liver, hepatocellular hypertrophy, hyperplasia, necrosis, inflammation, anisokaryosis, vacuolization, cholestasis and biliary hyperplasia. Increases in body weight in females at 3000 and 10000 ppm were possibly related to the increases in liver weights. The decreases in absolute and relative kidney weights in males of the 10000 ppm treatment group were not associated with microscopic lesions in the kidney. Based on the results of this study, dosage levels of 20, 100 and 300 ppm were selected for a subsequent 18-month oncogenicity study. No NOAEL could be derived from this study. The LOAEL was 100 ppm (equivalent to 19 mg/kg bw/d for males and 24 mg/kg bw/d for females). For details see DAR.

Dog, oral

Originally, only the 52-week study in Beagle dogs was submitted. For the resubmission of metaldehyde, also an oral 28-day study and an oral 26-week study in Beagle dogs were submitted.

In the <u>28-day dose-range-finding study</u>, severe clinical symptoms occurred at 90 mg/kg in the escalating dose experiment so that one female dog was sacrificed in moribund condition. In the fixed dose experiment, 75 and 90 mg/kg led to clinical symptoms but no mortality occurred. No histopathology was performed.

In the <u>26-week toxicity study</u>, no clinical symptoms were observed at all three dose levels of 20, 60 and 90 mg/kg. No effects were noted for body weights, haematology, clinical biochemistry, urinalysis, ophthalmology, auditory examination, macroscopic evaluation and organ weights. At histopathological examination, diffuse atrophy of the testes was found in the mid (2/6 males) and high dose group (4/6 males). A histopathological re-examination was performed in 2009, when also severity scores were investigated. The re-examination confirmed the pattern of testicular findings: the severity score of focal findings was increased at 60 mg/kg bw/day while at the high dose level of 90 mg/kg a clear increase of diffuse atrophic changes was found. The NOAEL for the 26-week study was therefore confirmed to be 20 mg/kg.

In the 52-week study in Beagle dogs (10, 30 and 90 mg/kg), mortality occurred in the mid and high dose group which was considered treatment-related. Clinical signs of toxicity (ataxia, reduced motility, emesis, tremor, twitching, salivation) were observed at the high dose group (90 mg/kg) from study week 1 onwards with incidence and severity declining over the time. Body weight gain was only slightly but not statistically significant reduced in the high dose group. There was indication of hearing impairment at the high dose group. Also in the high dose group, increases of erythrocytes, haemoglobin, hematocrit, bilirubin and triglycerides were observed during the conduct of the study but had largely resolved until the end of the study. In contrast, increased levels of alkaline phosphatase were observed in this group until the end of the study. The increase in liver weight in both sexes of the high dose group was not correlated with any histopathological finding. Male reproductive organs (testes, prostate) were target organs of metaldehyde as incidence and severity of microscopic lesions increased with dose. Prostate atrophy was observed in 1/4 males (25%) of the mid dose group and 3/4 (75%) males from the high dose group. In testes, atrophy and/or degeneration of the germinal epithelium was observed which mostly correlated with the occurrence of giant cells. The incidences for atrophy and/or degeneration of the germinal epithelium were 1/4 (25%) in the low dose, 2/4 (50%) in the mid dose and 4/4 (100%) in the high dose group. The single finding of juvenile testes in one animal of the low dose group appeared together with

giant cells. Upon request of the notifier, the histopathological slides of the testes were re-examined by the study pathologist in 2006 and the results presented in an expert statement. In this statement the effects occurring at 10 and 30 mg/kg were judged to be of spontaneous nature and morphologically distinct from the findings at 90 mg/kg. Historical background data were also submitted. In conclusion, it was proposed in the expert statement to set the NOAEL at 10 mg/kg bw/day based on mortality occurring at 30 and 90 mg/kg.

Considering the historical control data on testes findings, the RMS followed the majority of comments received from the Member States, EFSA and the notifier to set the NOAEL of the 52-week study at 10 mg/kg bw/d based on testes findings (atrophy of the germinal epithelium) and mortality observed at 30 mg/kg.

Rabbit, dermal

In a 21-day dermal toxicity study in rabbits, no substance-related effects were observed. Signs of local irritation occurred in control and treatment groups and were attributed to the dosing and occlusion procedures. The NOAEL for repeated dermal toxicity in rabbits was 1000 mg/kg bw/d (highest dose tested).

4.7.1.8 Summary and discussion of repeated dose toxicity findings relevant for classification according to DSD

Table 40: Summary of effects observed in rats, mice, dogs and rabbits in comparison to cut off vales

Species- Route (Reference)	Study duration	Cut off value R 48/22 (67/548/EC) [mg/kg bw/d]	Effects below cut off value	Significance of toxicological effect (67/548/EC) below cut off value
Mouse- oral (Gill M. et al., 1990)	90 days	50	≥ 19 (M)-24 (F) mg/kg bw/d: hepatocellular hypertrophy, necrosis, inflammation, anisokaryosis	Liver lesions of minimal severity- no evidence of organ dysfunction
Dog- oral (Leuschner J., 2002)	28 days	? *	-≥ 60 mg/kg bw/d: reduced motility, clonic convulsions, increased respiratory rate, emesis -≥ 75 mg/kg bw/d: tonoclonic convulsions, mydriasis, inflated stomach, slight tremor - 90 mg/kg bw/d: ataxia, salivation, abdominal/lateral position, pale gingivial; moribund condition of 1/3 females: shaking of the head, lateral position, difficulty in breathing- no pathological findings (macroscopic) after necropsy	The intensity/severity of the symptoms declined with time and had almost disappeared towards the end of the 4-week treatment Moribund condition, already covered by acute toxicity classification
Dog- oral (Neumann W., 1980, 1991, Leuschner J., 2009)	26 weeks	? *	-≥ 60 mg/kg bw/d: diffuse atrophy pf the prostate, moderate atrophy of the testes (1/6), mild to moderate focal atrophy of the germinative epithelium (3/6), parasitic granuloma in mes. lymph nodes - 90 mg/kg bw/d: mild to moderate diffuse atrophy of the germinative epithelium, follicular hyperplasia mes. lymph nodes	Severe organ damage

Species- Route (Reference)	Study duration	Cut off value R 48/22 (67/548/EC) [mg/kg bw/d]	Effects below cut off value	Significance of toxicological effect (67/548/EC) below cut off value
Dog- oral (Leuscher J., 2003)	51 weeks	?*	- ≥ 30 mg/kg bw/d: mortality - 90 mg/kg bw: ataxia, reduced motility, emesis, tremor, twitching, salivation- no changes in histopathology, transient hearing loss, ↑ AP, moderate to marked diffuse atrophy and/or degeneration of the germinative epithelium above the historical control data	Incidence and severity of clinical signs declined from study week 19 onwards. No changes in histopathology. Mortality Severe organ damage
Rabbit- oral (gavage) (Neeper- Bradley T., 1990a)	Developmental (12 days of dosing)	150?	Dams: -≥ 100 mg/kg bw: 1/5 tremor, 1/5 hypoactive and death	No gross lesions observed. No dose response regarding mortality after administration of repeated doses

^{*} For cut off values in dog studies, the only available document is ECBI/64/06 "Dose limits for classification with R48 based on dogs studies", 2006. In this document it is proposed that the cut off values for dog studies should be below the limit dose for the rat.

At histopathological examination in the <u>26-week dog toxicity study</u> diffuse atrophy of the testes was found in the mid (2/6 males) and high dose group (4/6 males). A histopathological re-examination was performed in 2009, when also severity scores were investigated. The re-examination confirmed the pattern of testicular findings: the severity score of focal findings was increased at 60 mg/kg bw/day while at the high dose level of 90 mg/kg a clear increase of diffuse atrophic changes was found.

In the 52-week dog study, all animals showed histological changes of testes at the highest dose (90 mg/kg), which consisted of mainly moderate to marked diffuse atrophy and/or degeneration of the germinative epithelium. At the two lower dose levels (10 and 30 mg/kg), findings were more focal and of minimal to mild severity and were thus considered to be of spontaneous nature and morphologically distinct from the treatment related findings seen at 90 mg/kg bw/day. Historical background data for atrophy and degeneration of the germinative epithelium demonstrated that incidences observed at the low and mid dose level were comparable to historical control data.

The notifier argued that a classification and labelling of metaldehyde with R48/22 cannot be derived from testicular changes seen in the 52-week dog study. It was questioned if the severity of effects and the dose where they appear are sufficient for classification. It was agreed that the testicular effects may be borderline for triggering classification and labelling with Xn; R48/22, as only the effects at 90 mg/kg bw/day seem to be severe enough to justify R48/22 in the case of testicular findings.

However, mortality observed in dogs at 30 mg/kg bw/day triggers classification with R48/22. One male and one female dog from the 30 mg/kg bw/d group and one female dog from the 90 mg/kg bw/d group were found dead between study days 260 and 322. It was clearly stated by the study authors that these deaths are considered treatment-related. The dose of 30 mg/kg bw/d is below the guidance value of 50 mg/kg bw/d (rat) listed in Directive 2001/59/EC, Annex VI, Point 3.2.3.

In conclusion, the RMS is of the opinion that metaldehyde should be classified with **R48/22** "Harmful: danger of serious damage to health by prolonged exposure if swallowed" based on the findings of mortality at 30 mg/kg bw/d and supported by the testicular findings in dogs.

4.7.1.9 Comparison with criteria of repeated dose toxicity findings relevant for classification according to DSD

Metaldehyde requires classification with R48/22 "Harmful: danger of serious damage to health by prolonged exposure if swallowed" based on the findings of mortality at 30 mg/kg bw/d and supported by the testicular findings in dogs.

Criteria as specified in the Directive 67/548/EEC: "Evidence indicating that R48 should be applied: a) substance related deaths [...] d) severe organ damage noted on microscopic examination following autopsy [...]". Substances should be classified "when these effects are observed at levels of the order of oral, rat \leq 50 mg/kg (bodyweight)/day".

4.7.1.10 Conclusions on classification and labelling of repeated dose toxicity findings relevant for classification according to DSD

Directive 67/548/EEC: Xn; R48/22

4.8 Specific target organ toxicity (CLP Regulation) – repeated exposure (STOT RE)

4.8.1 Summary and discussion of repeated dose toxicity findings relevant for classification as STOT RE according to CLP Regulation

For description of findings see chapter 4.7.1.8 Summary and discussion of repeated dose toxicity findings relevant for classification according to DSD.

Table 41: Summary of effects observed in rats, mice, dogs and rabbits in comparison to cut off values

Species- Route (Reference)	Study duration	Cut off value Cat 1 STOT RE (1272/2008) [mg/kg bw/d]	Cut off value Cat 2 STOT RE (1272/2008) [mg/kg bw/d]	Effects below cut off value	Significance of toxicological effect (1272/2008) below cut off value
Rat- oral (van Miller J.,1989)	28 days	30	300	-≥ 197 (M)- 233 (F) mg/kg bw/d: ↑ liver weights, hepatocelluar hypertrophy	Changes in liver weight with no evidence of organ dysfunction
Rat- oral (Thomas O. et al., 1998)	90 days	10	100	- 65 mg/kg bw/d only: centrilobular hepatocyte enlargement	No evidence of organ dysfunction
Mouse- oral (Gill M. et al., 1990)	90 days	10	100	≥ 19 (M)-24 (F) mg/kg bw/d: hepatocellular hypertrophy, necrosis, inflammation, anisokaryosis - ≥ 54 (M)-70 (F) mg/kg bw/d: ↑ liver weight, vacuolisation, bilary hyperplasia	Liver lesions of minimal severity- no evidence of organ dysfunction
Dog- oral (Leuschner	28 days	? *	? *	- ≥ 60 mg/kg bw/d: reduced motility, clonic convulsions,	The intensity/severity of the symptoms declined with time and had almost disappeared

Species- Route (Reference)	Study duration	Cut off value Cat 1 STOT RE (1272/2008) [mg/kg bw/d]	Cut off value Cat 2 STOT RE (1272/2008) [mg/kg bw/d]	Effects below cut off value	Significance of toxicological effect (1272/2008) below cut off value
J., 2002)				increased respiratory rate, emesis -≥75 mg/kg bw/d: tonoclonic convulsions, mydriasis, inflated stomach, slight tremor - 90 mg/kg bw/d: ataxia, salivation, abdominal/lateral position, pale gingivial; moribund condition of 1/3 females: shaking of the head, lateral position, difficulty in breathing- no pathological findings (macroscopic) after necropsy	towards the end of the 4-week treatment Moribund condition, already covered by acute toxicity classification
Dog- oral (Neumann W., 1980, 1991, Leuschner J., 2009)	26 weeks	? *	? *	-≥ 60 mg/kg bw/d: diffuse atrophy pf the prostate, moderate atrophy of the testes (1/6), mild to moderate focal atrophy of the germinative epithelium (3/6), parasitic granuloma in mes. lymph nodes - 90 mg/kg bw/d: mild to moderate diffuse atrophy of the germinative epithelium, follicular hyperplasia mes. lymph nodes	Severe organ damage
Dog- oral (Leuscher J., 2003)	51 weeks	? *	?*	-≥ 30 mg/kg bw/d: mortality - 90 mg/kg bw: ataxia, reduced motility, emesis, tremor, twitching, salivation- no changes in histopathology, transient hearing loss, ↑ AP, moderate to marked diffuse atrophy and/or degeneration of the germinative epithelium above the historical control data	Incidence and severity of clinical signs declined from study week 19 onwards. No changes in histopathology. Mortality Severe organ damage
Rabbit- oral (gavage) (Neeper- Bradley T., 1990a)	Developm ental (12 days of dosing)	30?	300?	Dams: -≥ 100 mg/kg bw: 1/5 tremor, 1/5 hypoactive and death	No gross lesions observed. No dose response regarding mortality after administration of repeated doses

^{*} For cut off values in dog studies, the only available document is ECBI/64/06 "Dose limits for classification with R48 based on dogs studies", 2006. In this document it is proposed that the cut off values for dog studies should be below the limit dose for the rat.

According to Regulation (EC) No. 1272/2008, mortality occurring at 30 mg/kg bw/d and histopathological testes findings observed at 90 mg/kg bw/d in a 52-week dog study trigger classification with Category 2 for Specific target organ toxicity – repeated exposure, **H373** "May cause damage to organs through prolonged or repeated exposure (if swallowed)".

4.8.2 Comparison with criteria of repeated dose toxicity findings relevant for classification as STOT RE

Metaldehyde requires classification with **H373** "May cause damage to organs through prolonged or repeated exposure (if swallowed)" based on the findings of mortality at 30 mg/kg bw/d and testicular findings at 90 mg/kg bw/d in dogs.

Criteria as specified in the Regulation (EC) No. 1272/2008: For the oral route the guidance values to assist in Category 2 classification are $10 < dose \le 100$ mg/kg bodyweight/day. These guidance values refer to effects seen in a standard 90-day toxicity study conducted in rats.

4.8.3 Conclusions on classification and labelling of repeated dose toxicity findings relevant for classification as STOT RE

Regulation (EC) No. 1272/2008: STOT RE 2, H373

4.9 Germ cell mutagenicity (Mutagenicity)

4.9.1 Non-human information

Table 42: Summary table of relevant in vitro and in vivo mutagenicity studies

Method	Results	Remarks	Reference
In vitro studies			
Reverse mutation assay (S. typhimurium TA 98, TA 100, TA 1535, TA 1537 and E. coli WP2uvrA ⁻)	0, 50, 150, 500, 1500 and 5000 μg/plate suspended in DMSO Negative (+/- S-9 mix)	-	Thompson, P.; 1998
Reverse mutation assay (S. typhimurium TA 98, TA 100, TA 1535, TA 1537 and TA 1538)	0, 0.26, 1.28, 6.4, 32 and 160 µg/plate (1 st experiment) 0, 4, 6, 16 and 32 µg/plate (2 nd experiment) dissolved in DMSO Negative (+/- S-9 mix)	Supplementary information only	Friederich, U., Wuergler, F.; 1981
Gene mutation assay in L5178Y mouse lymphoma cells	0, 20, 50, 100 and 200 μ g/ml (- S-9 mix) 0, 20, 50, 100 and 167 μ g/ml (+ S-9 mix) dissolved in HEPES-buffered cell culture medium Negative (+/- S-9 mix)	-	Debets, F., Enninga, I.; 1986
Chromosome aberration test in CHO cells	0, 20, 50, 100 and 200 μ g/ml (- S-9 mix) 0, 20, 50, 100 and 167 μ g/ml (+ S-9 mix) dissolved in HEPES-buffered cell culture medium Negative (+/- S-9 mix)	-	Debets, F.; 1986
Lethal DNA damage in Escherichia coli (WP2, WP67, CM871)	0, 100, 316, 1000, 3160 and 10000 μg/ml suspended in 0.15 % aqueous agar Negative (+/- S-9 mix)	-	May, K.; 1992
In vivo studies			
Oral micronucleus test in BKW mice	0, 25, 50 and 100 mg/kg bw suspended in arachis oil Negative	-	Jenkinson, P.; 1990

4.9.2 Human information

Not available.

4.9.3 Other relevant information

Not available.

4.9.4 Summary and discussion of mutagenicity

Metaldehyde was tested in a battery of *in vitro* genotoxicity testings including gene mutation in bacterial strains and L5178Y mouse lymphoma cells, chromosomal aberration in CHO cells and lethal DNA damage in Escherichia coli. None of these *in vitro* tests indicated genotoxicity of metaldehyde. In addition, an *in vivo* micronucleus assay in mice showed no genotoxic potential of metaldehyde. In conclusion, there was no indication that metaldehyde was genotoxic *in vitro* or *in vivo*.

4.9.5 Comparison with criteria

Metaldehyde was tested negative for genotoxicity in a battery of in vitro and one in vivo test.

4.9.6 Conclusions on classification and labelling

Directive 67/548/EEC: no classification proposed

Regulation (EC) No. 1272/2008: no classification proposed

4.10 Carcinogenicity

Table 43: Summary table of relevant carcinogenicity studies

Method	Results	Remarks	Reference
Chronic toxicity / Oncogenicity study in Sprague Dawley CD rats	0, 50, 1000 and 5000 ppm/diet (equivalent to 0, 2, 44 and 224 mg/kg bw/d for males; 0, 3, 60 and 314 mg/kg bw/d for females) NOAEL = 50 ppm Effects at LOAEL: -decreased body weight and body weight gain -increased serum cholesterol -hepatocellular hypertrophy Increase of hepatocellular adenomas in females of the 5000 ppm group was within the historical control range	-	Gill, M., Wagner, C.; 1992
Chronic toxicity study in Wistar rats	0, 200, 1000 and 5000 ppm/diet (no information on actual test substance intake is presented in the publication) NOAEL Could not be determined LOAEL = 200 ppm based on: - posterior paralysis - lordosis	This study is of limited validity.	Verschuuren, H. et al.; 1975
Oncogenicity study in CD-1 mice	0, 25, 100 and 300 ppm/diet (equivalent to 0, 4, 16 and 49 mg/kg bw/d for males; 0, 5, 20 and 60 mg/kg bw/d for females) NOAEL = 100 ppm Effects at LOAEL: - hepatocellular hypertrophy	-	Chun, J., Wagner, C.; 1993
Oncogenicity study in CD-1 mice	0 and 1000 ppm/diet (equivalent to 0 and 135 mg/kg bw/d for males; 0 and 163 mg/kg bw/d for females) LOAEL = 1000 ppm based on: - increased liver weight - hepatocellular toxicity - benign hepatocellular adenoma	Follow-up study to the oncogenicity study of Chun and Wagner, 1993.	Beyrouty, P.; 1998

4.10.1 Non-human information

4.10.1.1 Carcinogenicity: oral

Rat

Reference: Chronic dietary toxicity / oncogenicity study with metaldehyde in rat

Author(s), year: Gill M. and Wagner C., 1992

Report/Doc. number: Doc.No. 537-002, Lonza Report No. 1550

Conducting laboratory: Bushy Run Research Center, Pennsylvania, USA

Guideline(s): US EPA Guideline 83-5 (1984), OECD Guideline 453 (1981)

GLP: Yes
Deviations: No
Validity: Yes

Material and Methods:

60 male and 60 female Sprague Dawley CD rats per group received 0, 50, 1000 or 5000 ppm metaldehyde (batch no. 5448, purity 99 %) via the diet. The animals (source: Charles River Breeding Laboratories, Portage, MI, US) were approximately 8 weeks of age at the first dose. Two untreated control goups were included in this study. These groups were treated as independent entities for all activities performed during the study. The purpose was to collect data that would provide some information regarding the range of normal or control values for the parameters evaluated in this study. It was not considered appropriate to combine the data from the two control groups for the purposes of comparing the combined control data to those from the treated groups.

Table 44: Combined chronic toxicity / carcinogenicity study in Sprague Dawley CD rats; Experimental design and test substance intake

Group	Number of animals per group	Concentration in the diet	Test substanc (mg/kg bw/d)	e intake
		(ppm)	males	females
Control 1	60 m. / 60 f.	0	0	0
Control 2	60 m. / 60 f.	0	0	0
Low	60 m. / 60 f.	50	2	3
Mid	60 m. / 60 f.	1000	44	60
High	60 m. / 60 f.	5000	224	314

A 28-day dose range finding study was performed with rats being exposed to 0, 2500, 5000, 10000 and 20000 ppm in the diet. Mortality and traumatic injury to the back and spinal cord were observed for male and female rats at the 20000 ppm level and for female rats also at the 10000 ppm level. Hepatocellular hypertrophy was observed in most animals that survived to sacrifice. The severity of the hepatocellular hypertrophy was dose-related. In addition, sporadic foci of individual hepatocellular degeneration were noted in some animals from all treatment groups. Associated with hepatocellular hypertrophy there was a dose-related increase in liver weight for all treated male rats, and for female rats from the 5000 and 10000 ppm groups. Based on this dose finding study, dose levels of 50, 1000 and 5000 ppm were selected for the long term study.

Test diets were prepared weekly from a concentrated premix by appropriate dilutions. Homogeneity, concentration and stability were checked by gas chromatography analysis.

<u>In-life observations</u>: During the treatment period, observations for mortality were made twice daily. Detailed clinical observations including palpations were performed once a week. Observations of overt clinical signs were made once daily during the treatment period except on days with detailed clinical observations. Body weight and food consumption data were collected for all animals weekly for the first 14 weeks of the study and every second week thereafter. Ophthalmoscopic

examinations were performed for all animals prior to the start of the study and prior to final sacrifice.

<u>Clinical Pathology</u>: Hematology and clinical chemistry evaluations were conducted on 15 fasted animals/sex/group at 26, 52, 78 and 104 weeks of the study. Blood was collected from the retroorbital sinus. Whenever possible, urinalysis was conducted on the same 15 animals/sex/group during study weeks 25, 51, 77 and 103. For urine collection the rats were placed in metabolism cages for 24 hours.

<u>Hematology</u>: erythrocytes, haemoglobin, hematocrit, MCV, MCH, MCHC, platelet count, total leukocyte count, differential leukocyte count, reticulocyte count

<u>Clinical chemistry</u>: glucose, urea nitrogen, creatinine, aspartate aminotransferase (ASAT), alanine aminotransferase (ALAT), creatine kinase (CK), gamma glutamyl transpeptidase (GGT), alkaline phosphatase (AP), total protein, total cholesterol, albumin, globulin (calculated), A/G ratio (calculated), total bilirubin, direct bilirubin, indirect bilirubin (calculated), calcium, phosphorus, sodium, potassium and chloride

<u>Urinalysis</u>: color, appearance, specific gravity, total volume, pH, protein, glucose, ketone, bilirubin, blood, urobilinogen and microscopic elements

Pathology: Following the 104-week treatment period, terminal necropsy of all animals was undertaken. A complete necropsy was performed on all animals sacrificed at study termination, found dead or sacrificed moribund. Organ weights were determined for all animals sacrificed at termination: liver, kidneys, spleen, heart, brain with stem, adrenal glands, testes and ovaries. Complete histopathology was performed for all animals in both control groups and the high dose group and included gross lesions, spinal cord (cervical, midthoracic, lumbar), brain (cerebral cortex, cerebellar cortex, medulla/pons), pituitary, thyroid (with parathyroid), thymic region, trachea, lungs (with mainstem bronchi), heart, salivary gland (mandibular), liver, spleen, kidneys, adrenals, pancreas, testes, epididymis, prostate, seminal vesicles, ovaries, uterus (corpus and cervix), vagina, mammary gland, esophagus, stomach, duodenum, jejunum, ileum, cecum, colon, rectum, urinary bladder, skin, representative lymph nodes (mesenteric, submandibular), peripheral nerve (sciatic), sternum (including marrow), femur (including articular surface), thigh musculature, eyes and aorta. In the low and mid dose groups, only gross lesions, liver, lungs and kidneys were examined microscopically.

Findings:

Mortality and clinical signs: There were no treatment-related effects on the incidence of mortality observed. The mortality rates for male rats (including those sacrificed moribund / sacrificed moribund due to enlarged mass, but excluding procedural and accidental deaths) were 65 %, 58 %, 68 %, 60 % and 65 % for control 1, control 2, low, mid and high dose groups, respectively. For the females, mortality rates were 50 %, 45 %, 50 %, 60 % and 58 %, respectively. The tendency for increased mortality for females of the mid and high dose groups resulted from an increase in the number of animals in these groups that died during the last weeks of the study and was not considered to reflect a direct response to treatment. This conclusion was supported by the fact that the mean survival times for animals in the mid and high dose groups were slightly greater than those for the control and low dose groups. No significant alterations in clinical signs were noted throughout the study. However, with respect to the results of a published study described later in this section (Verschuuren H., 1975), the clinical observations regarding behaviour/CNS are described in the table below.

Table 45: Combined chronic toxicity / carcinogenicity study in Sprague Dawley CD rats; Mortality rates

	Dose g	Dose group level (ppm)								
	Males					Females				
	01	02	50	1000	5000	01	02	50	1000	5000
Total number of animals	60	60	60	60	60	60	60	60	60	60
Number sacrificed	20	24	17	22	20	30	31	30	24	25
Number found dead	24	22	32	26	24	10	9	12	17	19
Number sacrificed moribund	14	11	8	6	13	8	9	16	15	12
Number sacrificed due to enlarged mass	1	2	1	4	2	9	6	2	1	1
Number sacrificed due to ulcerated mass	0	0	1	2	1	3	3	0	3	3
Number cage accidents	1	0	0	0	0	0	2	0	0	0
Number procedural deaths	0	1	1	0	0	0	0	0	0	0
Mean survival time (days)	532	536	548	525	520	530	527	531	541	546

¹ Control group 1; ² Control group 2

Table 46: Combined chronic toxicity / carcinogenicity study in Sprague Dawley CD rats; Clinical signs on behaviour / CNS (number of animals affected; earliest to latest day a finding was observed)

	Dose group level (ppm)						
	01	02	50	1000	5000		
Males							
Hyperactive	2 (477-696)	1 (407-408)	-	1 (624)	1 (400)		
Hypoactive	16 295-728)	4 (477-722)	8 (393-710)	13 (557-713)	11 (283-718)		
Aggressive	-	1 (646)	-	1 (631-634)	1 (630)		
Paresis					, , ,		
- leg-hind both	4 (666-728)	1 (407-408)	3 (575-728)	3 (609-722)	3 (565-728)		
- leg-hind left	(708-722)	-	(666-673)	(636)	-		
- leg-hind right	(666-673)	1 (715-722)	-	-	-		
Paralysis							
- leg-hind both	-	-	1 (708-722)	-	-		
- leg-hind right	-	1 (708)	-	-	-		
Ataxia	4 (516-691)	4 (407-722)	3 (554-702)	2 (568-660)	8 (489-715)		
Tremor	3 (646-691)	1 (587)	1 (714)	(635-646)	4 (564-663)		
Clonic convulsions	-	-	-	1 (630)	-		
Tonic convulsions	-	-	-	1 (575)	-		
Helicoptering	1 (604-610)	-	1 (708)	-	2 (393-505)		
Circling	4 (484-728)	-	1 (631)	1 (610-631)	6 (456-718)		

	Dose group	level (ppm)			
	01	02	50	1000	5000
Prostration	6	4	4	2	7
Tiostation	(406-680)	(481-615)	(340-714)	(575-691)	(406-722)
Head tilt	8	-	4	2	6
	(516-728)		(528-702)	(575-660)	(365-718)
Tucked posture	1	-	1	1	-
•	(604-610)		(645)	(568-571)	
Exaggerated hind limb	1	1	3	1	-
placement	(726-727)	(624-722)	(624-687)	(610-631)	
Females	1 .		1		•
Hyperactive	-	1	-	1	1
		(463)		(484-498)	(491-503)
Hypoactive	7	13	11	11	9
	(456-729)	(463-729)	(421-704)	(323-729)	(421-728)
Paresis					
- leg-hind both	-	-	-	1	3
				(428-429)	(274-686)
- paw-hind both	-	-	-	-	1
					(615-621)
Paralysis					
- leg-hind both	-	-	-	-	1
					(484-485)
Ataxia	6	5	9	8	11
	(464-666)	(530-729)	(422-729)	(344-725)	(435-728)
Tremor	2	-	3	1	3
	(358-582)		(574-680)	(565)	(609-678)
Clonic convulsions	-	-	1	-	-
			(565)		
Helicoptering	-	2	1	-	1
		(463-576)	(652-708)		(603-666)
Circling	2	1	5	3	2
	(421-593)	(468)	(446-652)	(477-666)	(435-624)
Prostration	3	5	11	8	3
	(481-582)	(468-723)	(547-728)	(349-716)	(548-609)
Head tilt	6	8	11	10	4
	(400-729)	(435-729)	(441-729)	(505-729)	(421-708)
Tucked posture	3	2	1	2	1
	(505-616)	(533-631)	(446-470)	(505-554)	(611)
Exaggerated hind limb	2	5	5	5	3
Placement Control group 1: ² Control	(624-666)	(631-729)	(589-729)	(582-725)	(603-708)

¹ Control group 1; ² Control group 2

Body weights: Body weights and/or body weight gain were significantly decreased in males of the high dose group during the study. The mean absolute body weight and body weight gain were decreased generally 2-4 % and 6-8 % throughout the first year of the study and generally 4-8 % and 7-12 % by week 78, respectively. After that time this effect was no longer apparent as mean body weights for the groups began to vary as the incidence of mortality increased. The mean body weight and body weight gain for the mid dose group of males was slightly lower than controls during the early part of the study. These differences were, however, not statistically significant except for body weight gain in study week 1. No effect on body weight was observed in low dose males. Also in females, body weights and/or body weight gain were significantly decreased in the high dose group throughout the study until the last several measurement periods. The mean absolute body weight was generally 5-7 % lower than in controls and the mean body weight gain was

^a statistically significant different from control group 1 (p<0.05)

b statistically significant different from control group 2 (p<0.05)

generally 8-14 % lower than controls for most measurement periods. The mean body weight and body weight gain for the females from the mid dose group were lower than controls for approximately the first year of the study. The decrease in weight gain for the mid dose group was statistically significant for the first 10 weeks and intermittently through week 18. Statistically significant reductions in mean absolute body weight occurred for females in the mid dose group for some periods during the first 16 weeks of the study. There was no effect on body weight observed in the low dose group.

Table 47: Combined chronic toxicity / carcinogenicity study in Sprague Dawley CD rats; Body weight and body weight gains

	Dose gr	roup leve	l (ppm)							
	Males	_				Female	S			
	01	02	50	1000	5000	01	02	50	1000	5000
Body weight										
(g)										
-week 0	306.4	303.7	306.6	305.6	305.5	205.1	202.0	203.2	203.9	203.2
-week 14	593.1	589.3	598.1	579.8	574.6	320.3	320.2	315.0	309.7a,	303.7a,
-week 26	669.1	662.3	680.4	652.5	645.6a	353.7	355.3	350.3	b	b
-week 52	765.1	770.7	782.9	751.7	736.0b	405.5	411.4	409.1	344.4	332.5a,
-week 78	823.0	794.9	832.7	801.7	761.2a	452.2	476.3	463.3	404.4	b
-week 104	713.1	699.7	747.2	725.7	742.6	486.3	437.7	479.9	457.9	386.1
									503.7	441.4
										473.6
Body weight										
gain (g)										
-week 0-1	49.5	49.3	49.9	43.0a,b	38.4a,b	23.2	21.3	20.0a	15.9a,b	14.0a,b
-week 0-14	286.7	285.2	291.5	274.2	269.1a,	115.2	118.2	111.8	105.7a,	100.4a,
-week 0-26	362.7	358.2	373.3	346.9	b	148.6	153.8	147.0	b	b
-week 0-52	459.0	466.8	475.3	445.8	340.1a	200.6	209.9	205.9	140.4a,	129.2a,
-week 0-78	517.7	492.0	526.0	495.2	430.2a,	248.1	274.8	259.8	b	b
-week 0-104	405.4	441.6	398.2	441.9	b	281.9	237.2	277.1	200.6	182.8
					454.9b				254.4	237.9
					422.5				299.7	270.3

¹ Control group 1; ² Control group 2

<u>Food consumption</u>: Some statistically significant increases and decreases of food consumption in males and females were considered incidental and not related to treatment.

<u>Ophthalmoscopic examination</u>: Corneal crystals, keratitis and cataracts were the most common ophthalmic abnormalities noted in this study. The distribution across dosage groups indicated no test substance related effects.

<u>Hematology</u>: There were no differences in the mean values which were considered to be treatment-related. Female rats in the 5000 ppm group had a statistically significant decrease in MCV and MCH at week 26 and 78 while no statistical significance was reached at week 78 and 104. As the decreases were not consistent throughout the study and did not demonstrate a pathologic process, they were not considered to be biologically significant.

Table 48: Combined chronic toxicity / carcinogenicity study in Sprague Dawley CD rats; Hematology findings

Dose gr	oup level	(ppm)							
Males					Females	3			
01	02	50	1000	5000	01	02	50	1000	5000

^a statistically significant different from control group 1 (p<0.05)

^b statistically significant different from control group 2 (p<0.05)

	Dose g	roup leve	el (ppm)								
	Males	_				Femal	Females				
	01	02	50	1000	5000	01	02	50	1000	5000	
MCV (µm³)											
- week 26	49.8	51.5	50.7	50.8	50.4	54.6	54.6	55.9	54.3	52.8a,b	
-week 52	50.9	53.1a	51.9	51.6b	51.3b	55.9	56.0	56.5	55.3	54.2	
-week 78	52.4	53.6	51.9	52.9	51.4	56.0	56.2	56.4	55.6	54.0a,b	
-week 104	52.5	53.1	54.2	52.8	53.7	57.5	56.8	56.1	56.0	56.2	
MCH (pg)											
-week 26	18.4	19.2	18.8	18.9a	18.7b	20.5	20.7	20.9	20.4	19.9a,b	
-week 52	19.0	19.9	19.4	19.2	19.2	20.6	20.5	20.9	20.5	20.0	
-week 78	19.3	19.9	19.2	19.6	19.2	21.1	21.2	21.5	20.9	20.4a,b	
-week 104	17.7	18.0	18.3	17.8	18.1	19.5	19.5	19.2	18.9	19.1	

Clinical chemistry: In males, no treatment-related effects were observed. Female rats developed an apparent treatment-related effect on cholesterol (increased levels) at all measurement periods in the 1000 and 5000 ppm groups. Evidence of probable treatment-related increases in total protein, globulin and corresponding decrease in the A/G ratio were also present in the 5000 ppm females at week 26. The increase in globulins and corresponding decrease in the A/G ratio was persistent up to week 52 and 78. A shift in mean values at week 78, relative to week 52, reflect a change in methodology. All other statistically significant differences between mean values for control and treated animals occurred in a random fashion and were not supported by treatment-related trends and therefore not considered to be related to treatment.

Combined chronic toxicity / carcinogenicity study in Sprague Dawley CD rats; **Table 49:** Clinical chemistry findings in females

	Dose gro	Dose group level (ppm)									
	Females	=									
	01	02	50	1000	5000						
Cholesterol (g/L)											
- week 26	1.09	1.13	1.13	1.28a	1.57a,b						
-week 52	1.15	1.23	1.32	1.60a,b	1.76a,b						
-week 78	0.96	1.07	1.05	1.50a,b	1.55a,b						
-week 104	1.25	1.24	1.18	1.48	1.67						
Total protein (g/L)											
- week 26	69	70	66b	69	73a						
-week 52	68	70	70	70	71						
-week 78	74	77	78	77	79						
-week 104	70	72	72	70	71						
Globulin (g/L)											
- week 26	32	33	32	33	36a,b						
-week 52	32	33	34	34	36a,b						
-week 78	34	35	36	38a	38a						
-week 104	31	33	34	32	33						
A/G Ratio											
-week 26	1.14	1.11	1.08	1.08	1.01a,b						
-week 52	1.14	1.13	1.06	1.04	0.98a,b						
-week 78	1.24	1.19	1.19	1.06a	1.07a						
-week 104	1.32	1.26	1.17	1.22	1.20						

¹ Control group 1; ² Control group 2

<u>Urinalysis</u>: There were no statistically significant differences between control and treatment groups for male or female rats at weeks 25, 51 or 103. At week 77, female rats in the 1000 and 5000 ppm groups had statistically significant increases in urine total volume. Since these increases were not

¹ Control group 1; ² Control group 2 ^a statistically significant different from control group 1 (p<0.05)

b statistically significant different from control group 2 (p<0.05)

^a statistically significant different from control group 1 (p<0.05)

b statistically significant different from control group 2 (p<0.05)

noted again at study termination and there was no supporting evidence for kidney functional changes, they were not considered treatment-related.

<u>Organ weights</u>: Mean liver weights (absolute and relative to body weight or brain weight) were increased in males (10-16 %) and females (21-32 %) in the 5000 ppm treatment groups. These increases were statistically significant for females. No other effects on organ weights were noted.

Table 50: Combined chronic toxicity / carcinogenicity study in Sprague Dawley CD rats; Liver weights

	Dose g	group lev	el (ppm)								
	Males	Iales					Females				
	01	02	50	1000	5000	01	02	50	1000	5000	
Liver weights (g)											
-absolute weight	19.6	20.3	20.4	20.9	22.9	13.1	12.1	12.4	13.9	15.9a,b	
-% body weight	2.95	2.82	3.15	3.04	3.25	2.85	2.97	2.79	2.95	3.58a,b	
-% brain weight	847	861	873	895	971	630	599	608	680	772a,b	

¹ Control group 1; ² Control group 2

<u>Gross pathology</u>: Females had more liver masses and nodules in the high dose group but only at the scheduled sacrifice at week 104.

<u>Histopathology</u>: Based upon the preliminary findings for this study by the designated study pathologist that identified the liver as a target organ, the Sponsor requested an internal peer review of the livers by a second pathologist. Following completion of the peer review, both the designated and the second pathologist discussed the discrepancies between their diagnoses. The designated pathologist then made edits he deemed appropriate to his original diagnoses. The results which are described and discussed here reflect the final diagnoses. For reasons of transparency, also histopathological diagnoses from the initial liver microscopy and from the internal peer review of liver microscopy are presented in an additional table.

Non-neoplastic lesions: Dose related increases in the incidence and severity of hepatocellular hypertrophy were observed for male and female rats in the 1000 and 5000 ppm treatment groups. This effect was statistically significant for males at 1000 and 5000 ppm and females at 5000 ppm. The severity generally ranged from minimal to mild. Hepatocellular hypertrophy was usually centrilobular in distribution especially in those animals exposed to metaldehyde. In control animals and a few treated animals it tended to be periportal in distribution. No effect was noted on hepatocellular hyperplasia. Foci of cellular alteration were observed in control and treated animals. No clear relation to dose was observed though values were statistically significant for mid dose males found dead and for high dose males sacrificed at week 104. As part of the peer review, the altered cell foci were tabulated by cell type, i.e. vacuolated, clear, mixed, eosinophilic or basophilic. Again, there were no clear treatment-related differences.

Neoplastic lesions: Hepatocellular adenomas were observed in 6/60 females of the high dose group compared with incidences of 1/60 and 0/60 in the control groups, being statistically significant when compared to the second control group. In males, no hepatocellular adenomas were observed in metaldehyde treated animals. Hepatocellular carcinomas were observed for some males and females in treated and control groups. The incidences were not dose-related. In males of the mid dose group, incidences of carcinomas reached statistical significance in animals sacrificed in study week 104. When the combined incidences of hepatocellular adenomas and carcinomas were

^a statistically significant different from control group 1 (p<0.05)

b statistically significant different from control group 2 (p<0.05)

analysed statistically, females of the high dose group had a significantly higher incidence of tumors when compared to the second control group. Historical control data were supplied from two studies conducted at the same laboratory (Bushy Run Research Center, Report Numbers 53-543 and 53-566), using the same study design and source of animals. In study no. 1 incidences of hepatocellular carcinomas and adenomas were in the same range as in the present study while in study no. 2 the incidences were somewhat lower.

In the present study, hepatocellular adenomas and carcinomas were usually apparent upon gross examination but were never very large or widespread within the organ. They were never identified as cause of death. These observations suggest that hepatocellular neoplasms developed late in the lives of the animals. The earliest to appear was a carcinoma in a male rat in the low dose group at 86 weeks of age.

There were no other neoplasms which occurred with significantly greater incidence in treated animals than in controls. Tumors of the pituitary were by far the most frequently occurring neoplasms, affecting 49 % of the males and 85 % of the females. A variety of tumors was reported in mammary glands of females. Numerous other tumors occurred in control and treated animals with similar frequencies and were expected in a study of this nature. There were no rare or unusual tumors which occurred in such a distribution as to suggest a relationship to treatment.

Table 51: Combined chronic toxicity / carcinogenicity study in Sprague Dawley CD rats; Histopathology findings in the liver

	Dose	group l	level (p	om)						
	Male	S				Fem	ales			
	01	02	50	1000	5000	01	02	50	1000	5000
Non-neoplastic lesions										
Hepatocellular										
hypertrophy										
-sacrificed at week 104	-	2	3	11a,b	20a,b	1	3	-	5	20a,b
-found dead / sacrificed	-	-	3	9a,b	18a,b	4	2	-	6	16a,b
moribund										
-all animals on study	-	2	6a	20a,b	38a,b	5	5	-b	11	36a,b
Hepatocellular										
hyperplasia										
-sacrificed at week 104	1	-	-	-	-	-	-	2	1	2
-found dead / sacrificed	-	1	-	2	1	2	1	2	-	1
moribund										
-all animals on study	1	1	-	2	1	2	1	4	1	3
Focus of cellular										
alteration										
-sacrificed at week 104	10	18	9	10	18a	13	13	17	17	12
-found dead / sacrificed	7	8	6	16a	9	5	8	5	10	8
moribund										
-all animals on study	17	26	15	26	27	18	21	27	20	21
Neoplastic lesions								-		
Hepatocellular										
adenoma										
-sacrificed at week 104	1	-	-	-	-	1	-	-	-	5b
-found dead / sacrificed	-	-	-	-	-	-	_	1	-	1
moribund										
-all animals on study	1	-	-	-	-	1	-	1	-	6b
Hepatocellular										
carcinoma										
-sacrificed at week 104	1	-	1	4a	-	-	-	1	_	1
-found dead / sacrificed	1	-	3	-	2	1	_	-	_	-
moribund										
-all animals on study	2	-	4	4	2	1	-	1	_	1

	Dose	group l	level ((ppm)								
	Male							Females				
	01	02	50	1000	5000	01	02	50	1000	5000		
Hepatocellular												
adenomas + carcinomas	3	-	4	4	2	2	-	2	-	7b		
Historical control data (f	rom co	nductin	ıg lab	oratory)								
	Cont	rol 1		Control 2	,	Cont	trol 1		Control 2			
	(male	es)		(males)		(fem	ales)		(females)			
Study 1												
-adenomas	1			7		6			6			
-carcinomas	-			1		-			-			
-adenomas+carcinomas	1			8		6			6			
Study 2												
-adenomas	3			3		1			-			
-carcinomas	-			_		-			-			
-adenomas+carcinomas	3			3		1			-			

¹ Control group 1; ² Control group 2

Table 52: Combined chronic toxicity / carcinogenicity study in Sprague Dawley CD rats; Initial and peer review liver histopathology for adenomas and carcinomas

	Dose	group	level (pj	pm)						
	Male	es				Fema	Females			
	01	02	50	1000	5000	01	02	50	1000	5000
INITIAL										
liver histopathology										
-adenoma	3	1	4	3	5	1	-	1	2	7
-carcinoma	-	-	-	2	1	1	-	1	-	-
-adenoma + carcinoma	3	1	4	5	6	2	-	2	2	7
PEER REVIEW										
liver histopathology										
-adenoma	2	-	-	-	-	1	-	1	_	5
-carcinoma	1	-	4	4	3	1	-	1	-	3
-adenoma + carcinoma	3	-	4	4	3	2	-	2	_	8

¹ Control group 1; ² Control group 2

Conclusion:

Administration of metaldehyde in the diet for 104 weeks did not result in toxicologically significant alterations in mortality, clinical signs of toxicity, palpable masses or food consumption. Body weight and body weight gain were reduced throughout the study period in males of the 5000 ppm group and females of the 1000 and 5000 ppm groups. Occasionally decreased MCV and MCH values in females of the 5000 ppm dose group remained of questionable toxicological significance. A clear treatment-related effect was observed in clinical chemistry evaluation when increased cholesterol levels were observed in females of the 1000 and 5000 ppm groups. Additionally, some alterations in total protein, globulin and albumin / globulin ratio values were found in females of the 5000 ppm group. Pathology findings revealed the liver to be the target organ of metaldehyde. Increased liver weights (absolute and relative to body and brain weight) were found in the 5000 ppm dose group as well as an increased rate of liver masses and nodules. The incidence and severity of hepatocellular hypertrophy increased with dose. Statistical significance was reached in males already at a dose of 50 ppm, but this finding did not correlate with other liver findings. Hepatocellular tumors were found in males and females in treatment and control groups. No dose relation was observed in male animals. In females, statistical significance was reached for the 5000 ppm group when compared to one of the two concurrent control groups. However, when tumor incidences were compared with historical control data, a relation to treatment with metaldehyde was found to be questionable. In conclusion, a NOAEL of 50 ppm (2 and 3 mg/kg

^a statistically significant different from control group 1 (p<0.05)

b statistically significant different from control group 2 (p<0.05)

bw/d for males and females, respectively) is defined based on slightly lower body weight and body weight gain during the first year of the study for females, increased serum cholesterol for females, and hepatocellular hypertrophy for both sexes. No carcinogenic potential of metaldehyde was assumed based on the results of this study.

In the valid long term toxicity / oncogenicity study in Sprague Dawley CD rats (Gill, M., Wagner, C.; 1992- see above), the main target organ identified was the liver, as it was observed also in the short term toxicity studies. Body weight and body weight gain were decreased in the mid dose (1000 ppm, females only) and high dose groups (5000 ppm, both sexes). Increased absolute and relative liver weights were noted in rats of the high dose group. The incidence and severity of hepatocellular hypertrophy increased with dose in both sexes of the mid and high dose groups. At the same dose levels, blood cholesterol levels were increased in females. Additionally, some alterations in total protein, globulin and albumin/globulin ratio were found in females of the high dose group. Hepatocellular tumors were found in males and females in treatment and control groups. While no dose relation was observed in males, statistical significance against one of the two control groups was reached in females of the high dose group regarding hepatocellular adenomas alone and the sum of hepatocellular adenomas and carcinomas. As the incidences were within the historical background range of the conducting laboratory, a relation to treatment with metaldehyde was not assumed. No effects were noted at the low dose level of 50 ppm.

The second study in Wistar rats (Verschuuren, H. et al.; 1975) was of limited validity due to study design and reporting. The dose levels tested (200, 1000 and 5000 ppm) were comparable to the other study regarding the mid and high dose level, however, the outcome of the study was different. Effects on the liver were only found at the 5000 ppm level in form of increased liver weights reaching statistical significance in males only. No treatment-related changes in liver histopathology were reported. On the other hand, clinical signs of posterior paralysis were observed in 1/25 males from the 200 ppm group (first sign at day 569), 1/25 males (first sign at day 657) and 1/25 females of the 1000 ppm group (first sign at day 652) and 5/25 females from the 5000 ppm group (first sign at day 19, 641, 625, 659 and 559). The clinical signs were reflected in histopathology where transverse lesions of the spinal cord were observed in 3 of the females showing posterior paralysis. In the 3 animals with posterior paralysis receiving 200 and 1000 ppm, only lordosis was detected. The histological investigations showed that the lesions were caused by trauma i.e. a "sudden kink" in the vertebral column leading to a fracture or distortion of the vertebrae and a subsequent compression of the spinal cord. This effect is similar as described in the 28-day study in rats, when animals receiving very high doses (10000 and 20000 ppm) developed hind limb paralysis/paresis. No NOAEL could be derived. The LOAEL of this very limited study is 200 ppm.

Mouse

The original oncogenicity study in CD-1 mice (Chun and Wagner; 1993) showed no changes in mortality, clinical signs, body weight, food consumption, haematology or necropsy findings. Like in rats, the liver was found to be the target organ. Liver weight (absolute and relative) was slightly but not significantly increased in the high dose males (300 ppm). In histopathology, a significant increase in the incidence of hepatocellular hypertrophy was noted in both sexes of the high dose group. An increase of hepatocellular adenomas in high dose males was statistically not significant, however slightly exceeding historical control data (HCD; animals of the same strain, sex and source at a comparable time period). In conclusion, a NOAEL of 100 ppm (16 and 20 mg/kg bw/d for males and females, respectively) was derived based on the increased incidence of hepatocellular hypertrophy in both sexes at 300 ppm (49 and 60 mg/kg bw/d for males and females, respectively).

An additional carcinogenicity study (Beyrouty, 1998) was conducted as a supplement to the previous 18-month mouse study. CD-1 mice were administered doses of 0 and 1000 ppm in order to satisfy the regulatory requirements for a maximum tolerated dose. The treatment with 1000 ppm (equivalent to 135 and 163 mg/kg bw/d for males and females, respectively) metaldehyde resulted in changes in the liver of males and to a lesser extent of females. These changes included increased liver weight and an increase in the incidence of animals with hepatocellular toxicity and benign hepatocellular adenoma. The nature of these findings is consistent with an adaptive hypertrophic response of the liver to an increase in metabolic demand with subsequent development of proliferative changes and hepatocellular toxicity. A summary of hepatocellular tumour incidences observed in both studies is given in the following table which is taken from the resubmission Additional Report (AR, June 2010) to the DAR for metaldehyde.

Table 53: Overview on hepatocellular tumour incidences and historical control values in CD-1 mice

Hepatocellular Neoplasms		Tumour incidence	2
-		Males	Females
Hepatocellular ADENOMA			
Study of Chun and Wagner, 1993			
Concurrent Control Group 1	0 ppm	8/60	0/60
Concurrent Control Group 2	0 ppm	8/60	1/60
Low Dose Group	25 ppm	4/60	1/60
Mid Dose Group	100 ppm	9/60	1/60
High Dose Group	300 ppm	15/60	0/60
Study 1#: Historical Control Group 1	0 ppm	6/60	No information available #
Study 2#: Historical Control Group 2	0 ppm	13/60	No information available #
Study of Beyrouty, 1998			
Concurrent Control Group 1	0 ppm	4/60	1/60
Concurrent Control Group 2	0 ppm	5/60	0/60
Single Dose Group	1000 ppm	14/601,2	5/602
Hepatocelluar CARCINOMA			
Study of Chun and Wagner, 1993			
Concurrent Control Group 1	0 ppm	1/60	0/60
Concurrent Control Group 2	0 ppm	3/60	0/60
Low Dose Group	25 ppm	6/60	0/60
Mid Dose Group	100 ppm	3/60	1/60
High Dose Group	300 ppm	3/60	0/60
Study of Beyrouty, 1998			
Concurrent Control Group 1	0 ppm	2/60	0/60
Concurrent Control Group 2	0 ppm	2/60	0/60
Single Dose Group	1000 ppm	4/60	0/60

statistically significantly different from concurrent control group 1, p<0.05

Historical control data for Studies 1 and 2 were derived from the original study report which was submitted for DAR evaluation (Chun and Wager, 1993, "Chronic Dietary Oncogenicity study with Metaldehyde in Mice", Appendix 3, Anatomic Pathology Report, page 9-10). Only values for male animals are presented there. In the Pathology Report of the carcinogenicity study, Study 1 and 2 are described as two studies conducted with animals of the same strain, sex and source at a comparable time period (BRRC Project Report Numbers 53-515 and 53-528). Historical control data for male mice from both studies (53-515 and 53-528) were cited also in the Position Paper submitted by the notifier in April 2010.

The issue of a possible cancerogenic potential of metaldehyde has been discussed at the **PRAPeR** 79 (Pesticide Risk Assessment Peer Review) expert meeting in July 2010:

² statistically significantly different from concurrent control group 2, p<0.05

For hepatocellular carcinomas in CD-1 mice, there is no dose relation or statistical significance for males. For females, there was only one incidence of carcinoma at mid dose without statistical significance. The experts agreed that the hepatocellular carcinomas are of not concern.

In the first mouse study (Chun and Wagner, 1993), there was a dose related increased incidence of hepatocellular adenomas in males, but no statistical significance. Incidence was above HCD at the high dose of 300 ppm. No HCD is available for females.

In the second mouse study (Beyrouty, 1998), for both males and females there is a statistically significant increase in hepatocellular adenomas at 1000 ppm (single dose study) compared to the concurrent control groups. It was agreed that the HCD presented in Table 6.5.3-5 are only for the first study (Chun and Wagner, 1993). It was commented that these HCD may not be reliable for the second study (Beyrouty, 1998) due to the time interval (5 years) and different laboratories used. Furthermore, it was mentioned that males may have exceeded the tolerable limit in the Beyrouty study since in the 90-day preliminary mouse study (Gill and Wagner, 1990) liver toxicity was observed at all doses \geq 100 ppm (19 mg/kg bw/day and above). This liver toxicity may have contributed to the incidence of adenomas observed in the long term carcinogenicity studies.

The experts considered whether the hepatocellular adenomas in CD-1 mice are enough to trigger classification based on the above data. Finally, the experts voted on whether classification with R40 should be proposed: **The majority of experts voted to not classify with R40.**

This is in accordance with Directive 67/548/EEC which states that "if the only available tumour data are liver tumours in certain sensitive strains of mice, without any further other supplementary evidence, the substance may not be classified in any of the categories".

4.10.1.2 Carcinogenicity: inhalation

No data available.

4.10.1.3 Carcinogenicity: dermal

No data available.

4.10.2 Human information

Not available.

4.10.3 Other relevant information

Not available.

4.10.4 Summary and discussion of carcinogenicity

Metaldehyde showed no cancerogenic potential in rats. In the first mouse study (Chun and Wagner, 1993), there was a dose related increased incidence of hepatocellular adenomas in male CD-1 mice, but no statistical significance. The incidence was slightly exceeding HCD at the high dose of 300 ppm. In the second mouse study (Beyrouty, 1998), for both male and female CD-1 mice there was a statistically significant increase in hepatocellular adenomas at 1000 ppm (single dose study) compared to the concurrent control groups. No HCD is available for the second mouse study. At the PRAPeR 79 (Pesticide Risk Assessment Peer Review) expert meeting in July 2010 the experts decided to not classify metaldehyde with R40 based on the hepatocellular adenomas found in CD-1 mice (majority after vote).

4.10.5 Comparison with criteria

The decision to not classify metaldehyde with R40 based on the hepatocellular adenomas found in CD-1 mice is in accordance with Directive 67/548/EEC which states that "if the only available tumour data are liver tumours in certain sensitive strains of mice, without any further other supplementary evidence, the substance may not be classified in any of the categories".

4.10.6 Conclusions on classification and labelling

Directive 67/548/EEC: no classification proposed

Regulation (EC) No. 1272/2008: no classification proposed

4.11 Toxicity for reproduction

Table 54: Summary table of relevant reproductive toxicity studies

Method	Results	Remarks	Reference
Two generation study in CD rats	0, 50, 1000 and 2000 ppm/diet (equivalent to 0, 3.2, 65 and 134 mg/kg bw/d for males; 0, 4.0, 81 and 160 mg/kg bw/d for females) Reproduction NOAEL: 2000 ppm Parental NOAEL: 50 ppm Offspring NOAEL: 1000 ppm Parental and offspring toxicity effects at LOAEL: - reduction of bw and bw gain	-	Chun, J., Neeper-Bradley, T.; 1993
Two generation study in Wistar rats	0, 200, 1000 and 5000 ppm/diet (no information on actual test substance intake is presented in the publication)	No NOAELs are defined due to the limited reporting and the limited validity of this study.	Verschuuren, H. et al.; 1975
Developmental toxicity study in Sprague Dawley CD rats	0, 25, 75 and 150 mg/kg bw/d oral gavage Maternal NOAEL: 75 mg/kg bw/d Fetal NOAEL: 150 mg/kg bw/d Maternal effects at LOAEL: - mortality and clinical signs - reduced bw gain	-	Neeper-Bradley, T., Chun, J.; 1990
Dose range finding study in NZW rabbits	0, 50, 100, 200, 350 and 500 mg/kg bw/d oral gavage Clinical signs of maternal toxicity leading to premature death were already observed at the dose level of 100 mg/kg	This is a dose finding study. No NOAELs were defined due to limited fetal examinations.	Neeper-Bradley, T.; 1990
Developmental toxicity study in NZW rabbits	0, 10, 40 and 80 mg/kg bw/d oral gavage Maternal NOAEL: 80 mg/kg bw/d Fetal NOAEL: 80 mg/kg bw/d	-	Neeper-Bradley, T.; 1990

4.11.1 Effects on fertility

4.11.1.1 Non-human information

Two two-generation studies were submitted. The newer study (*Chun J. and Neeper-Bradley T.*, 1993) was conducted according to international guidelines and GLP and is considered valid. The older study (*Verschuuren H. et al.*, 1975) was not conducted according to guidelines or GLP and is presented in the form of a publication. It shows several limitations like short reporting, missing individual numbers on several study parameters and limited statistical evaluations which do not allow to set sound and scientifically based NOAELs. Furthermore, this study was conducted with a test material from a different supplier and obviously not from the notifier. Therefore, the results from this study are of limited validity.

Reference: Two generation reproduction study in CD rats with metaldehyde

administered in the diet

Author(s), year: Chun J. and Neeper-Bradley T., 1993

Report/Doc. number: Bushy Run Research Center, Pennsylvania, USA

Lonza Report No. 1544, Doc. No. 543-001

Guideline(s): US EPA Guideline 83-4, OECD Guideline 416 (1983)

GLP: Yes
Deviations: No
Validity: Valid

Material and Methods:

Groups of 28 male and 28 female outbred albino CD rats (source: Charles River Laboratories, Portage, MI, USA) received metaldehyde (batch no. 5448, purity > 99 %) in dietary concentrations of 0, 50, 1000 and 2000 ppm. The mean body weight range of F0 animals on the day of first treatment was 227.0 – 277.7 g for males and 164.3 – 165.3 g for females. The F0 animals were exposed to the diet for a prebreed period of 10 weeks. Following the prebreed period the F0 animals were randomly paired within dose groups for a 21-day period to produce the F1 generation. Exposure to the test diets continued through mating, gestation, parturition and lactation. After the F1 pups were weaned, the F0 females were sacrificed. F0 males were necropsied after delivery of the F1 litters. At weaning, 28 F1 weanlings/sex/group were randomly selected as parents of the next generation. The F1 weanlings were exposed to the same dietary concentrations of metaldehyde as their parents for at least 10 weeks. In addition, 10 F1 pups/sex/group were randomly selected for necropsy. After the prebreed period, the F1 animals were paired as described above to produce the F2 offspring. One week after weaning of F2 pups, 10 pups/sex/group were randomly selected for necropsy and the remaining F2 offspring were examined for gross external abnormalities, euthanized and discarded.

<u>In-life evaluations</u>: Parental animals were examined twice daily for mortality and overt signs of toxicity and once daily for any clinical signs of toxicity. Detailed clinical examination was conducted weekly. Body weights and food consumption were measured weekly. All pups from the F1 and F2 generations were examined as soon as possible on the day of birth (day 0) to determine the number of viable and stillborn pups per litter. Litters were evaluated twice daily for survival. On day 4 after birth, the size of each litter was adjusted by eliminating extra pups to four males and four females in each litter. Culled pups were examined externally, sacrificed and discarded. All pups were examined for abnormalities at birth and throughout the lactational period. All pups dying during lactation were necropsied when possible to investigate the cause of death.

<u>Necropsy</u>: Animals that died or were sacrificed prior to scheduled sacrifice were subjected to full necropsy as soon as possible after they were found. Adult males surviving the treatment period were examined externally and sacrificed after parturition of the litters. All surviving females were similarly examined and sacrificed after weaning of their offspring. Necropsy included examination of the external surfaces, all orifices, cranial cavity, carcass, external surfaces of the brain and spinal cord, the thoracic, abdominal and pelvic cavities and their viscera including reproductive organs, and cervical tissues and organs. In addition, spinal cords of adult animals that died or were sacrificed in a moribund condition were retained.

<u>Histopathology</u>: The following tissues from F0 and F1 parental animals from the control and high dose groups were examined histopathologically: spinal cord, vagina, uterus, ovaries, testes, epididymides, seminal vesicles, prostate, and tissues with gross lesions.

<u>Dose finding study</u>: The dose levels selected for this study were based on the results of a reproductive toxicity dose range finding study in CD rats conducted at dietary concentrations of 0, 625, 1250, 2500 and 5000 ppm metaldehyde. No treatment-related clinical signs were observed in

males. One female from the high dose group was observed with hind limb paresis on day 10 and was sacrificed. Necropsy of this animal indicated luxation of the spine between the 9th and 10th thoracic vertebrae. Body weight and weight gain as well as food consumption tended to be reduced during prebreed period for all treated males and females. During gestation/lactation, six additional females from the high dose group were sacrificed due to hind limb paralysis/paresis. Necropsy of this animals indicated vertebral fractures/luxations and perivertebral hemorrhaging in all animals. In the last week of lactation, five females from the 2500 ppm group were observed with hind limb paralysis/paresis. These animals were also observed with vertebral fractures/luxations and/or perivertebral hemorrhaging. Pup body weights were reduced in the last two weeks of lactation in the 2500 and 5000 ppm groups presumably resulting from direct intake of the test diets.

Findings:

<u>In-life observations</u>: No treatment-related mortality or clinical signs of toxicity were observed in F0 and F1 males during the entire study period. In F0 females, no clinical signs were noted during prebreed, mating and gestation. During lactation on days 16-18, three females from the 2000 ppm group were sacrificed due to bilateral hind limb paralysis. In F1 females, no treatment-related mortality was observed. One F1 female (no. 13923) of the 50 ppm group that showed paresis on day 43 had lymphosarcoma and was sacrificed. One non-gravid female from the 2000 ppm dose group (no. 13973) showed prostration, tremors, abdominal breathing and rapid respiration on day 102 but survived to scheduled sacrifice on day 134.

Table 55: Reproductive toxicity study in CD rats; Clinical signs on behaviour / CNS (number of animals affected; earliest to latest day a finding was observed)

	Dose group level (ppm)							
	0	50	1000	2000				
F0 females								
Paralysis (leg-hind-both)	-	-	-	3 (110-113)				
Twitch (entire body)	-	-	-	1 (113)				
F1 females								
Paresis (leg-hind-left)	-	1 (43)	-	-				
Ataxia	-	-	-	1 (94)				
Tremor, Prostration	-	-	-	1* (102)				

^{*} non-gravid female no. 13973

<u>Reproduction parameters</u>: No reproductive parameters including mating, fertility and gestational indices as well as gestational length were affected by treatment in both the F0 and F1 generation.

Table 56: Reproductive toxicity study in CD rats; Reproductive parameters

	Dose group level (ppm)					
	0	50	1000	2000		
F0 generation						
No. F0 pairs	28	28	28	28		
No. males impregnating females	28	27	26	28		
No. plug / sperm-positive females	28	27	27	28		
No. pregnant	25	26	25	27		
No. males siring litters	25	25	24	26		
No. live litters on postnatal day 0	25	25	24	27		
Gestational length (days)	22.0 ± 0.4	22.1 ± 0.6	22.0 ± 0.6	22.0 ± 0.4		

	Dose group level (ppm)				
	0	50	1000	2000	
Mating index males1)	100	96.4	92.9	100	
Mating index females2)	100	96.4	96.4	100	
Fertility index males3)	89.3	92.6	92.3	92.9	
Fertility index females4)	89.3	96.3	92.6	96.4	
Gestation index5)	100	96.2	96.0	100	
F1 generation					
No. F0 pairs	28	27a)	28	28	
No. males impregnating females	28	26	27	27	
No. plug / sperm-positive females	28	27	28	28	
No. pregnant	27	24	23	24	
No. males siring litters	27	22	23	23	
No. live litters on postnatal day 0	27	22	23	22	
Gestational length	22.0 ± 0.5	22.0 ± 0.7	22.0 ± 0.5	22.0 ± 0.6	
Mating index males1)	100	96.3	96.4	96.4	
Mating index females2)	100	100	100	100	
Fertility index males3)	96.4	84.6	85.2	85.2	
Fertility index females4)	96.4	88.9	82.1	85.7	
Gestation index5)	100	91.7	100	91.7	

- 1) Number of males impregnating females x 100 / Total number of males paired
- 2) Number of plug-/sperm-positive females x 100 / Total number of females paired
- 3) Number of males siring litters x $100\,/$ Number of males impregnating females
- 4) Number of pregnant females x $100\,/$ Number of plug-/sperm-positive females
- 5) Number of females with live litters x 100 / Number of females pregnant
- a) One female was sacrificed on day 43 prior to pairing

F1 and F2 offspring: There were no treatment-related effects on litter size or sex ratio for the F1 and F2 pups. Pup body weight of the treated groups was similar to control values through the first four days of lactation. From lactation day 7-28, mean body weights were slightly reduced in the 2000 ppm groups in F1 and F2 pups. These reductions corresponded to a decreased body weight gain in this period of time. A single statistically significant decrease in pup weight gain (lactational day 7-14) in F2 pups receiving 50 or 1000 ppm was not considered to be biologically significant due to the small magnitude of effects and due to the fact that absolute body weight was not changed. There was no effect on pup viability and survival in F1 and F2 pups. While the number of dead pups appeared to be increased for the 2000 ppm dose group on day 21, all 24 of the dead pups were sacrificed due to concurrent maternal sacrifices on lactational days 16, 17 or 18. At necropsy, there were no gross lesions observed that were attributed to treatment.

Table 57: Reproductive toxicity study in CD rats; Pup body weight and body weight gain (g)

	Dose group level (ppm)								
	Males				Females	Females			
	0	50	1000	2000	0	50	1000	2000	
F1 pups									
Body weight									
Lactation day 1	7.27	7.50	7.39	7.34	6.85	7.00	6.99	6.84	
day 7	17.78	17.88	17.59	17.10	16.90	16.69	16.82	16.03	
day 14	36.51	36.44	37.07	34.52	34.97	34.40	35.70	33.04	
day 21	59.67	59.41	60.37	56.61	56.76	55.56	57.60	53.79	
day 28	104.19	104.40	105.25	99.90	94.89	93.39	95.24	90.56	
Body weight gain									
Lactation day 1-4	3.65	3.75	3.59	3.57	3.50	3.54	3.48	3.39	
day 4-7	6.86	6.62	6.60	6.19	6.54	6.17	6.35	5.80**	
day 7-14	18.73	18.56	19.49	17.42	18.07	17.71	18.88	17.01	
day 14-21	23.16	22.97	23.30	21.88	21.79	21.16	21.90	20.48	
day 21-28	44.51	44.99	44.88	43.29	38.13	37.83	37.63	36.80	
F2 pups								•	

	Dose group level (ppm)							
	Males				Females			
	0	50	1000	2000	0	50	1000	2000
Body weight								
Lactation day 1	7.52	7.75	7.21	7.03	7.15	7.14	6.77	6.74
day 7	17.57	17.98	17.34	16.95	17.08	17.19	16.55	16.21
day 14	37.16	36.62	35.59	34.47	36.61	35.04	34.49	33.23**
day 21	60.54	59.56	58.16	56.24**	58.63	56.49	55.94	53.93**
day 28	100.89	100.51	99.56	95.91	93.50	91.51	90.04	88.25
Body weight gain								
Lactation day 1-4	3.23	3.47	3.81	3.24	6.82	6.76	6.32	6.66
day 4-7	3.22	3.36	3.63	3.11	6.71	6.57	6.16	6.36
day 7-14	19.59	18.63	18.26	17.52	19.23	17.85*	17.94*	17.02**
day 14-21	23.38	22.94	22.56	21.77	22.32	21.45	21.45	20.71
day 21-28	40.35	40.95	41.40	39.29	34.87	35.03	34.10	34.31

^{*} Significantly different from control group (p < 0.05)

Necropsy and histopathology of F0 parental animals: There were no treatment-related necropsy findings for F0 males and females surviving to scheduled sacrifice. Necropsy of F0 females from the 2000 ppm dose group that were sacrificed due to hind limb paralysis indicated vertebral fractures or spinal cord luxations for two of the three females. Additional gross findings associated with spinal cord injury included dilation / distention of the urinary bladder and/or blood in the bladder. Microscopic examination for all three F0 females with hindlimb paralysis from the 2000 ppm group showed hemorrhage and/or necrosis of the spinal cord. In addition, minimal spinal cord hemorrhage was also observed microscopically for one female from the 2000 ppm group that survived to scheduled sacrifice without previous clinical signs of injury. There were no other treatment-related microscopic findings for F0 animals that survived to scheduled sacrifice. Organ weights, necropsy and histopathology of F1 parental animals: In F1 males, relative liver weight was increased at the 2000 ppm dose level. In F1 females, both absolute and relative liver weights were increased for the 2000 ppm dose group. No necropsy or microscopic findings observed in F1 animals of either sex or any dose level were attributed to treatment with metaldehyde.

Two F1 females from the 50 ppm group were sacrificed due to clinical signs or found dead during the study.

Female (13920) showed no clinical signs but was found dead. Histopathology for this animal exhibited among other findings: meningitis, myelitis and hemorrhage of the spinal cord. However, these effects were in all probability a result of infection with septic emboli to various organs.

Female (13923) showed paresis and was sacrificed due to severe clinical signs on day 43. However, histopathology of this animal exhibited lymphosarcoma but no adverse histopathological findings of the spinal cord.

Table 58: Reproductive toxicity study in CD rats; Histopathology findings

	Dose group level (ppm)					
	Females	;				
	0	50	1000	2000		
F0 females						
Necropsy						
Paralysis	-	-	-	3/281)		
Spinal cord: consistency change,	-	-	-	1/281)		
shape/contour change, color change						
Histopathology						
Spinal cord: hemorrhage	-	-	-	31)		
Spinal cord: necrosis	-	-	-	21)		
F1 females		<u>.</u>	<u>.</u>	<u> </u>		

^{**} Significantly different from control group (p < 0.01

	Dose group level (ppm)					
	Females					
	0	50	1000	2000		
Histopathology						
Spinal cord: hemorrhage, meningitis,	-	12)	-	-		
myelitis						
Lymphosarcoma	-	13)	-	-		

- 1) found in the three females that were sacrificed due to clinical signs of paralysis (leg-hind-both)
- 2) died of infection with septic emboli to various organs
- 3) the female was sacrificed due to clinical signs of paresis (leg-hind-left); autopsy revealed lymphosarcoma

Table 59: Reproductive toxicity study in CD rats; Histopathology findings in female adults sacrificed at week 19

	Dose group level (ppm)					
	Females					
	0	50	1000	2000		
F0 females						
Histopathology						
Spinal cord: hemorrhage	1	-	-	1		
Spinal cord: vacuolization	1	-	-	0		
Spinal cord: neuronal vacuolization	1	-	-	2		
F1 females						
Histopathology						
Spinal cord: hemorrhage	1	-	=	-		

Conclusion:

No effects on reproduction were observed in this study. The NOAEL for reproductive toxicity therefore is larger than 2000 ppm (equivalent to 134 mg/kg bw/d in males and 160 mg/kg bw/d in females).

Systemic parental toxicity was evident in the high dose level (2000 ppm) when three females of the F0 generation developed paralysis (both hind legs) and were sacrificed moribund. Histopathology showed hemorrhage and necrosis of the spinal cord. One F1 female of the low dose group (50 ppm) showed signs of paresis of the left hind leg and was sacrificed. Autopsy revealed lymphosarcoma and hemorrhage, meningitis and myelitis. Though a connection to treatment cannot be excluded, this single case of paresis in the F1 females was not considered relevant for setting the NOAEL as no effects were observed at the two higher dose levels and the severity of the effect (paresis) was lower than in F0 females (paralysis). Body weight and body weight gain was affected only in F1 females. Statistically significant changes were observed at 1000 ppm and above during preebreeding, gestation and lactation. Absolute and/or relative liver weights were increased at the high dose level in both sexes. Offspring toxicity was demonstrated at the high dose level as body weight and body weight gain were decreased predominantely in F2 pups. In conclusion, the NOAEL for parental systemic toxicity was 50 ppm (equivalent to 3.2 mg/kg bw/d in males and 4.0 mg/kg bw/d in females) based on reduction of body weight. For offspring toxicity, the NOAEL was 1000 ppm (equivalent to 65 mg/kg bw/d in males and 81 mg/kg bw/d in females), again based on reduction of body weight and body weight gain.

4.11.1.2 Human information

Not available.

4.11.2 Developmental toxicity

4.11.2.1 Non-human information

Developmental toxicity studies according to GLP and international guidelines were conducted in rats and rabbits.

Reference: Developmental toxicity evaluation of metaldehyde administered by

gavage to CD (Sprague Dawley) rats

Author(s), year: Neeper-Bradley T. and Chun J., 1990 Report/Doc. number: Doc.No. 551-003, Lonza Report No. 1545

Conducting laboratory: Bushy Run Research Center, Pennsylvania, USA

Guideline(s): US EPA Guideline 83-3 (1982), OECD Guideline 414 (1981)

GLP: Yes
Deviations: No
Validity: Valid

Material and Methods:

Groups of 25 presumed pregnant rats (strain: CD of Sprague Dawley origin; source: Charles River Breeding Laboratories, Michigan, USA) received metaldehyde (batch no 5448; purity 99.0 %; suspended in corn oil) from day 6 to 15 of pregnancy by oral gavage at a constant dosing volume of 5 ml/kg bw. Dose levels were 0 (vehicle control), 25, 75 and 150 mg/kg bw/d. These dose levels were selected on the results of a preliminary study (reported in summary form only) with dose levels of 0, 50, 100, 150, 200 and 250 mg/kg bw/d given to groups of 6 pregnant rats. In this doserange finding study, maternal toxicity including mortality was observed at dose levels \geq 200 mg/kg bw/d. At 150 mg/kg bw/d, treatment-related clinical signs and reductions in body weight and food consumption were observed. There were no indications of developmental toxicity at any treatment level in this preliminary study.

Observations for mortality and clinical signs were made at least daily. Body weights were recorded on days 0, 6, 9, 12, 15, 18 and 21 of gestation, food consumption was measured at three-day intervals throughout gestation. On gestation day 21, females were subjected to a post-mortem macroscopic examination. The reproductive tract and abdominal and thoracic organs were examined grossly. In addition, maternal liver and uterine weights were determined. Further investigations included the number of corpora lutea/ovary, implantation and resorption sites, number and distribution of live and dead fetuses, weight and sex of each fetus and external malformed fetuses. Half of the fetuses from each litter were processed for the examination of visceral (thoracic and abdominal) abnormalities. These fetuses were then decapitated and their heads examined for abnormalities of craniofacial structures. The remaining half of fetuses were processed for skeletal staining and examined for skeletal malformations and variations. Dosing suspensions were prepared three times during the main study and were analyzed for test substance content prior to use.

Findings:

Maternal toxicity: Following 1 - 2 days of treatment, 6/25 dams at 150 mg/kg were found dead, which was considered related to treatment (all 6 dams were pregnant). In addition, at this dose level, treatment-related clinical signs of toxicity occured in the surviving rats during treatment period and included ataxia, tremor and rapid respiration. One of the six dams also showed paresis of both hind legs. No clinical signs were seen in the other dose groups. Maternal body weight was significantly reduced at 150 mg/kg on gestation day 9. A statistically significant reduction in mean body weight gain at 150 mg/kg was observed for the same period of time on gestation days 6-9 and 6-15. Also

maternal food consumption was reduced in this group for the first three days of treatment (days 6-9), but increased subsequent to treatment (days 15-21) which can be considered a compensatory effect. No dams (at any dose level) aborted, delivered early or were removed from the study. At scheduled laparotomy, one dam each at 0, 25 and 150 mg/kg and three dams at 75 mg/kg were not pregnant. All remaining pregnant dams had one or more live fetuses at scheduled sacrifice (no litters were fully resorbed). 18 litters were available for examination at 150 mg/kg and 22-24 litters were available for examination at each of the other dose groups.

At necropsy, the pathological findings in the 6 dams which died before termination included perioral and perinasal wetness and encrustation, colour changes in the lungs, ulceration in the glandular and nonglandular portions of the stomach, blood and colour changes of the urinary bladder and distended intestines. Pathological findings in females of the high dose group considered to be related to treatment comprised hydronephrosis and dilated renal pelvis (3 dams) and paravertebral hemorrhages in the thoracic (2 dams) and lumbar region (1 dam). There were no significant treatment-related effects on uterine and liver weight at any dose level.

Dam No	Treatment day	Gestation day	Behavirour/ CNS	Time of death (gestation day)	Lesion (probably CNS related)- gross examination
24533	1	6	Twitch, ataxia	Scheduled sacrifice	-
24599	1	6	Ataxia, twitch, tremor	7	Paravertebral hemorrhage thoracic area
24588	1	6	Tremor	Scheduled sacrifice	-
24591	2	7	Ataxia, hyperactive	8	-
24497	2	7	Ataxia	8	-
24466	1	6	Tremor, ataxia	7	-
24524	1	6	Twitch	Scheduled sacrifice	Paravertebral hemorrhage thoracic and lumbar areas
24472	2	7	-	7	-

8

Ataxia, prostration,

paresis

Table 60: Death, behaviour/CNS effects in dams at 150 mg/kg bw

<u>Litter data/fetal parameters</u>: There were no significant effects of treatment on any gestational parameter. The number of corpora lutea, total implantations, pre- and postimplantation losses, sex ratio, and mean fetal and placental weights gave no indication of any response to treatment in any dose group. Examination of the fetuses at necropsy showed no treatment-related increases in individual external, visceral or skeletal malformations. The only finding in fetuses showing statistical significance was an increased incidence of bilobed centrum of the 12th thoracic vertebra-considered a skeletal variation - at 25 mg/kg dose group (39/158) when compared with controls (24/157), but not in the 75 (23/145) and 150 mg/kg dose groups (25/150). The incidence of all other variations (external, visceral or skeletal) were not significantly altered by the administration of metaldehyde.

Significant findings of this study are given in table 61.

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Table 61: Rat oral developmental toxicity (teratogenicity) study; Summary of substantial study findings (group mean values)

Parameter	0 mg/kg	25 mg/kg	75 mg/kg	150 mg/kg
No. on study	25	25	25	25
No. that died	0	0	0	6
No. examined at laparotomy	25	25	25	19
No. pregnant at laparotomy	24	24	22	18

Parameter	0 mg/kg	25 mg/kg	75 mg/kg	150 mg/kg
Maternal body weight gain (g)				
day 0 - 6	25.97	26.98	29.93	26.23
day 6 - 15	34.53	35.88	31.31	25.41**
day 15 - 21	88.27	86.49	88.67	93.82
Food consumption (g/animal/day)				
day 0 - 6	20.94	21.19	21.60	21.02
day 6 - 15	19.40	19.57	19.10	16.91**
day 15 - 21	25.52	25.65	27.40*	27.33*
Total implants	14.5	14.3	15.3	14.0
No. of live fetuses (% males)	13.7 (46.4 %)	13.6 (48.7 %)	13.8 (54.4 %)	13.3 (46.4 %)
No. of early resorptions	0.9	0.7	1.5	0.7
No. of late resorptions	0	0	0	0.1
No. of dead fetuses	0	0	0	0
Fetal body weight per litter (g)	5.130	5.093	5.208	5.295
Total number of fetuses with	10	10	12	10
malformations				

^{**} $(p \le 0.01)$, * $(p \le 0.05)$; significantly different

Conclusion:

The maternal NOAEL in this study can considered to be 75 mg/kg bw/d based on mortality and clinical signs, but also on slight reduction in body weight gain and food consumption during the treatment period at 150 mg/kg bw/d. Fetal survival and growth was not affected in any dose group. In addition, there was no teratogenic effect observed. Therefore the fetal NOAEL can be set at 150 mg/kg bw/d.

Reference:	Developmental toxicity dose range-finding study of metaldehyde administered by gavage to New Zealand White rabbits
Author(s), year:	Neeper-Bradley T., 1990a
Report/Doc. number:	Doc.No. 551-001, Lonza Report No. 1503
	Conducting laboratory: Bushy Run Research Center, Pennsylvania, USA
Guideline(s):	Dose range finding study according to US EPA Guideline 83-3 (1982) and
	OECD Guideline 414 (1981)
GLP:	Yes
Deviations:	low animal number and fetal examinations restricted to external
	malformations and variations
Validity:	Valid

Material and Methods:

The objective of this study was to determine the appropriate dose levels of metaldehyde for a subsequent definitive developmental toxicity study in New Zealand White rabbits. Five mated females per group (source: Hazelton-Dutchland Laboratories Inc., Denver, PA, USA) received metaldehyde (batch no. 5448, purity 99.0 %, suspended in corn oil) from gestation days 6 to 18 by oral gavage at a dosing volume of 2 ml/kg. The dose levels were 0 (vehicle control), 50, 100, 200, 350 and 500 mg/kg bw/d. Dosing suspensions were prepared twice times during the main study and were analyzed for test substance content prior to use.

Observations for mortality and clinical signs were made twice daily during the treatment period and daily throughout the rest of the study. Body weights were recorded on days 0, 6, 13, 19, 24 and 29 of gestation. All surviving females were sacrificed on gestation day 29 and subjected to a macroscopic examination. The reproductive tract and abdominal and thoracic organs were examined grossly. Maternal liver and uterine weights were determined. Further investigations

included the number of corpora lutea, implantation and resorption sites, number and distribution of live and dead fetuses, fetal body weight per litter, external malformations and variations.

Findings:

Maternal toxicity: Following 1 - 2 days of treatment, 4/5 does at 500 mg/kg and 2/5 does at 350 and 200 mg/kg were found dead. 1/5 females receiving 100 mg/kg died on gestation day 12. All of those females were pregnant. The remaining females of the 200, 350 and 500 mg/kg dose groups were sacrificed on gestation day 8 due to the severity of clinical signs. All euthanized females were pregnant with the exception of one at 200 mg/kg. Characteristic clinical signs of these dose groups were ataxia, whole body twitching, tremors and rapid respiration. Rapid respiration was observed during treatment of does receiving 100 mg/kg. One doe of the 50 mg/kg group developed abdominal breathing and rapid respiration on gestation day 16 and aborted on gestation day 26. Of the 13 does surviving to the scheduled sacrifice, 2 does (one each at 0 and 100 mg/kg) were nonpregnant. Four litters were evaluated at 0 and 50 mg/kg and three litters were evaluated at 100 mg/kg.

There were no statistically significant effects on maternal body weight and body weight gain. However, food consumption was reduced from gestation days 6-7 in females receiving 100, 200 and 350 mg/kg. At necropsy, some of the does at 100, 200, 350 and 500 mg/kg which died prior to scheduled sacrifice exhibited broken vertebrae and gastrointestinal tract lesions, including color changes and ulcerations of the glandular stomach and distended intestines with no evidence of technical dosing error. In addition, color changes of the liver and ulcerations and color changes in the nonglandular portion of the stomach were observed at doses 100 mg/kg and above. Does surviving to scheduled sacrifice on gestation day 29 did not exhibit any treatment-related gross lesion. Maternal organ weights at scheduled sacrifice showed no statistically significant differences among groups. There were no apparent differences in corrected body weight or corrected gestational body weight change (corrected for gravid uterine weight). Relative liver weight was non statistically significant increased at 100 mg/kg.

Table 62: Death, behaviour/CNS effects in dams

Dam No	Dose group (mg/kg bw)	Treatment day	Gestation day	Behavirour/ CNS	Time of death (gestation day)	Lesion (probably CNS related)- gross examination
9896	100	3	9	Hypoactive	11	-
9888	100	1/9-10	6/ 15-16	Tremor (6), head tilt (15-16)	Scheduled sacrifice	-
9891	200	2	7	tremor	8 (euthanized, sponsor requested)	-
9892	200	2	7	Ataxia, twitch, paresis, tremor	8	Vertebral column broken lower lumbar region
9902	200	1	6	Tremor, ataxia	7	-
9889	350	2	7	Tremor, tonic convulsions, prostration, apparent broken back	7	Vertebral column broken
9909	350	1	6	Tremor, ataxia,	7	-
9918	500	1	6	Tremor, twitch	7	Vertebral column broken near first lumbar vertebra
9913	500	1	6	Ataxia, tremor, twitch	7	Vertebral column broken near first lumbar vertebra

9887	500	1	6	Ataxia	7	-
9900	500	1/2	6/7	Tremor (6), ataxia	7 (euthanized,	-
				(7)	sponsor	
					requested)	
9890	500	1	6	Twitch, ataxia	7	-

<u>Litter data/fetal parameters</u>: Gestational parameters for does at scheduled sacrifice were approximately equivalent across all groups from 0 to 100 mg/kg, with no statistically significant changes in the number of corpora lutea, live fetuses, early or late resorptions, dead fetuses, implantations, percent live fetuses per litter or sex ratio. While there were greater numbers of live fetuses at 50 and 100 mg/kg, due to the larger number of total implants, the number of non-viable implants was also increased. The apparent increase in the number of dead fetuses was most probably due to the greater number of total implants. Fetal body weights per litter were equivalent across groups. There were no external malformations or variations observed in any fetus of this study.

Significant findings of this study are given in table 63.

Table 63: Rabbit oral developmental dose finding study; Summary of substantial study findings (group mean values)

Parameter	0 mg/kg 5	50 mg/kg	100 mg/kg	200 mg/kg 5	350 mg/kg 5	500 mg/kg 5
No. on study						
No. found dead	-	-	1	2a)	2a)	4a)
Abortions	-	1	-	-	-	-
No. examined at laparotomy	5	4	4	-	-	-
No. pregnant at laparotomy	4	4	3	-	-	-
Total implants	5.3	10.8	9.0	-	-	-
No. of live fetuses	5.0	9.5	7.7	-	-	-
No. of early resorptions	0	0.5	0.7			
No. of late resorptions	0	0	0	-	-	-
No. of dead fetuses	0.2	0.7	0.7			
Fetal body weight per litter (g)	45.00 (n=4)	40.18 (n=4)	40.21 (n=3)	-	-	-

a) the remaining females were sacrificed on gestation day 8

Conclusion:

In this dose range finding study in NZW rabbits, severe maternal toxicity was observed at the high dose levels of 100 mg/kg and above. Clinical signs like ataxia, whole body twitching, tremors and rapid respiration were observed, leading to premature death of 1/5, 2/5, 2/5 and 4/5 females of the 100, 200, 350 and 500 mg/kg dose group. At necropsy, broken vertebrae and gastrointestinal tract lesions were found in some of these animals. The remaining females receiving 200, 350 or 500 mg/kg were euthanized due to clinical signs on gestation day 8. Treatment rapid respiration was also observed in one doe of the 50 mg/kg dose group, which aborted on gestation day 26.

There were no indications of fetal toxicity in this study. Also, no external malformations or variations were observed. No examination for visceral anomalies or detailed skeletal examination were performed.

Based on the effects indicative for maternal toxicity at 100 mg/kg bw/day (mortality and clinical signs) and the slight effect observed at 50 mg/kg (abdominal breathing and rapid respiration in one dam), the doses for the main study were selected as 0, 10, 40 and 80 mg/kg bw/d. No NOAELs are

^{**} $(p \le 0.01)$, * $(p \le 0.05)$; significantly different

set for this dose finding study. Due to low animal number and missing detailed fetal investigations this study is of supplementary information only.

Reference: Developmental toxicity evaluation of metaldehyde administered by

gavage to New Zealand White rabbits

Author(s), year: Neeper-Bradley T., 1990b

Report/Doc. number: Doc.No. 551-002, Lonza Report No. 1504

Conducting laboratory: Bushy Run Research Center, Pennsylvania, USA

Guideline(s): US EPA Guideline 83-3 (1982), OECD Guideline 414 (1981)

GLP: Yes
Deviations: No
Validity: Valid

Material and Methods:

16 mated females per group (source: Hazelton-Dutchland Laboratories Inc., Denver, PA, USA) received metaldehyde (batch no. 5448, purity 99.0 %, suspended in corn oil) from gestation days 6 to 18 by oral gavage at a dosing volume of 2 ml/kg. The dose levels of 0 (vehicle control), 10, 40 and 80 mg/kg bw/d were selected on the basis of a dose finding study (Neeper-Bradley T., 1990a), when maternal mortality was observed at a dose level of 100 mg/kg. Dosing suspensions were prepared three times during the study and were analyzed for test substance content prior to use.

All females were thoroughly examined daily for clinical signs of toxicity (twice daily during dosing period). In addition, the animals were examined twice daily for mortality and morbidity. All females on study were weighed on gestation day 0, 6, 13, 19, 24 and 29. Food consumption was recorded daily. All surviving females were sacrificed on gestation day 29 and subjected to a macroscopic examination. The reproductive tract and abdominal and thoracic organs were examined grossly. Maternal liver and uterine weights were determined. Further investigations included the number of corpora lutea, implantation and resorption sites, number and distribution of live and dead fetuses and fetal body weight per litter. All fetuses were examined for external malformations and variations, visceral anomalies and skeletal malformations and variations. Half of the fetuses were also examined for soft tissue craniofacial structures.

Findings:

Maternal toxicity: Pregancy rate was approximately equivalent across all groups ranging from 75 – 100 %. One doe receiving 40 mg/kg was found dead on gestation day 17, however, no reason for this death is explained in the study report. Necropsy of this female showed color changes of liver and spleen and blood in the urinary bladder, while histopathology revealed no abnormalities. There were no clearly treatment-related clinical signs of toxicity observed in any of the treatment groups, though one female of the 80 mg/kg dose group developed rapid respiration during the treatment period. There were no statistically significant differences among groups for maternal body weight, body weight gain or food consumption. Necropsy findings in does surviving to the scheduled sacrifice were not considered to be treatment-related. There were no apparent differences among groups in gravid uterine weight, corrected body weight or body weight change. While not statistically significant, there appeared to be dose-dependent increases in liver weight and relative liver weight. However, at the 80 mg/kg dose, liver weights were only increased by 6 %. Gestational parameters showed no statistically significant changes in the number of corpora lutea, live fetuses, early or late resorptions, dead fetuses or sex ratio.

<u>Fetal examination</u>: No statistical changes in fetal body weights per litter were observed. There were no treatment-related increases in the incidence of individual fetal malformations by category (external, visceral, skeletal) or of total malformations observed at any dose level in this study. There

were also no treatment-related increases in the incidence of individual fetal variations, by category or total, observed.

Significant findings of this study are given in table 64.

Table 64: Rabbit oral developmental toxicity (teratogenicity) study; Summary of substantial study findings (group mean values)

Parameter	0 mg/kg	10 mg/kg	40 mg/kg	80 mg/kg
No. on study	16	16	16	16
No. that died	0	0	1	0
No. examined at laparotomy	16	16	15	16
No. pregnant at laparotomy	12	13	13	16
Maternal body weight gain (g)				
day 0 - 6	123	113	124	132
day 6 - 19	14	-103	-164	-1
day 19 - 29	279	312	340	326
Food consumption (g/animal/day)				
day 0 - 6	189	169	181	184
day 6 - 19	92	81	70	79
day 19 - 29	155	147	152	182
Liver weight				
-absolute	95	95	98	100
-relative to body weight	2.73	2.75	2.81	2.87
Total implants	10.8	9.8	9.4	10.4
No. of live fetuses	9.3	8.0	8.2	8.7
No. of early resorptions	0.2	1.2	0.3	0.7
No. of late resorptions	0.2	0.1	0.2	0.1
No. of dead fetuses	1.1	0.6	0.7	0.7
Fetal body weight per litter (g)	40.22	41.09	39.25	40.64
Total % fetuses with	2.7	1.9	0.9	4.3
malformations (external, soft				
tissue, skeletal)				
Total % fetuses with variations	99.1	99.0	100	99.3
(external, soft tissue, skeletal)				

^{**} $(p \le 0.01)$, * $(p \le 0.05)$; significantly different

Conclusion:

In this study, no maternal toxicity was observed at the dose levels employed (0, 10, 40 and 80 mg/kg bw/d). The lack of maternal toxicity at least at the high dose level is in contrast to the dose range finding study, when clear toxicity resulting in the death of 1/5 females was observed at the dose level of 100 mg/kg. Rapid respiration, a characteristic sign of toxicity in the dose range finding study, was noted in one female at 80 mg/kg in the actual study, which suggests that the dose of 80 mg/kg was just below the level of clear maternal toxicity in the population of rabbits in the actual study. Obviously, metaldehyde has a very steep dose – effect relationship regarding acute toxic effects including mortality, that is why the study is acceptable for evaluation of the teratogenic potential of this test substance even when no maternal toxicity was observed. In conclusion, the maternal NOAEL of this study was set at 80 mg/kg bw/d. Fetal survival and growth were not affected in any dose group. In addition, there was no teratogenic effect observed. Therefore the fetal NOAEL is greater than 80 mg/kg bw/d.

4.11.2.2 Human information

Not available.

4.11.3 Other relevant information

Not available.

4.11.4 Summary and discussion of reproductive toxicity

Two two-generation studies were submitted. The newer study (*Chun J. and Neeper-Bradley T.*, 1993) was conducted according to international guidelines and GLP and is considered valid. The older study (*Verschuuren H. et al.*, 1975) was not conducted according to guidelines or GLP and is presented in the form of a publication. It shows several limitations like short reporting, missing individual numbers on several study parameters and limited statistical evaluations which do not allow to set sound and scientifically based NOAELs. Furthermore, this study was conducted with a test material from a different supplier and obviously not from the notifier. Therefore, the results from this study are of limited validity.

In the newer two generation study conducted in CD rats, no effects on reproductive performance were noted throughout the study. The NOAEL for reproductive toxicity therefore is higher than the highest dose tested: > 2000 ppm (equivalent to 134 mg/kg bw/d in males and 160 mg/kg bw/d in females). Systemic parental toxicity was evident at the high dose level (2000 ppm) when three females of the F0 generation developed paralysis (both hind legs) and were sacrificed moribund. Histopathology showed hemorrhage and necrosis of the spinal cord. Such findings were also observed in the older two generation study, where 50-75% of the females of the 5000 ppm dose group and up to three females of the 1000 ppm dose group developed posterior paralysis with transverse lesions of the spinal cord. None of the males was affected in any generation in both studies. The finding of hind limb paralysis was also observed in repeated dose toxicity studies at high dose levels, however, the increased body weight of the pregnant females may have promoted this effect. Further signs of systemic toxicity were reduced body weight in F1 females receiving 1000 or 2000 ppm and increased liver weight in both sexes receiving 2000 ppm. Offspring toxicity was evident at 2000 ppm and resulted in decreased body weight and body weight gain. In conclusion, the NOAEL for parental systemic toxicity was 50 ppm (equivalent to 3.2 mg/kg bw/d in males and 4.0 mg/kg bw/d in females) based on reduction of body weight. For offspring toxicity, the NOAEL was 1000 ppm (equivalent to 65 mg/kg bw/d in males and 81 mg/kg bw/d in females), again based on reduction of body weight and body weight gain.

In the <u>rat</u> study, severe maternal toxicity was observed at the highest dose level of 150 mg/kg, including ataxia, tremor, rapid respiration, paresis of hind legs, and finally mortality. In these animals, necropsy revealed hydronephrosis, dilated renal pelvis and paravertebral hemorrhages. Also body weight and body weight gain as well as food consumption were slightly reduced in the dams of the high dose group. No effects regarding maternal toxicity were observed at the low and mid dose of 25 and 75 mg/kg. No effects on gestational parameters or on fetal toxicity were observed at any dose. Also, no evidence of teratogenicity was observed in this study. In conclusion, the NOAEL for maternal toxicity in rats was 75 mg/kg bw/d and the NOAEL for fetal toxicity and teratogenicity was greater than 150 mg/kg bw/d.

The teratogenicity of metaldehyde in <u>rabbits</u> was investigated in one dose finding study and in the main study. In the dose finding study, severe maternal toxicity was observed at dose levels of 100 mg/kg and above, leading to ataxia, tremors, whole body twitching, rapid respiration and finally to death following 1-2 days of treatment. The remaining females of the high dose groups (200, 350 and 500 mg/kg) were sacrificed on gestation day 8 due to the severity of clinical signs. At necropsy, some of the high dose females showed broken vertebrae and gastrointestinal lesions, including color changes and ulcerations of the glandular stomach and distended intestines. All animals surviving to the scheduled sacrifice did not exhibit any treatment-related gross lesion. One female of the 50 mg/kg dose group of the dose finding study developed rapid respiration during dosing period,

which was regarded to be related to treatment. There was no effect on body weight of the dams and the fetuses. Limited fetal examinations showed no external malformations or variations. Based on the findings of this study, dose levels of 10, 40 and 80 mg/kg bw/d were set for the main study. However, in the main study no maternal toxicity occurred. Rapid respiration, a characteristic sign of toxicity in the dose range finding study, was noted in one female at 80 mg/kg in the main study, which suggests that the dose of 80 mg/kg was just below the level of clear maternal toxicity in the population of rabbits in the main study. Obviously, metaldehyde has a very steep dose-effect-relationship regarding acute toxic effects including mortality. Therefore the study is acceptable for evaluation of the teratogenic potential of this test substance even if no maternal toxicity was observed. No effects on gestational parameters, fetal survival and growth were observed. There was no evidence of teratogenicity. In conclusion, the maternal and fetal NOAEL for developmental toxicity in rabbits was 80 mg/kg bw/d.

4.11.5 Comparison with criteria

In a valid two generation study conducted in CD rats, no effects on reproductive performance were noted throughout the study. Systemic parental toxicity was evident at the high dose level (2000 ppm) when three females of the F0 generation developed paralysis (both hind legs) due to transverse lesions of the spinal cord. Further signs of systemic toxicity were reduced body weight in F1 females in the mid and high dose group (receiving 1000 or 2000 ppm, respectively) and increased liver weight in both sexes in the high dose group. Offspring toxicity was evident only in the high dose group and resulted in decreased body weight and body weight gain. In conclusion, the NOAEL for reproductive toxicity is > 134 mg/kg bw/d, the NOAEL for parental systemic toxicity is 3.2 mg/kg bw/d (based on reduction of body weight) and the NOAEL for offspring toxicity is 65 mg/kg bw/d (based on reduction of body weight and body weight gain).

In the 26 and 52 week dog toxicity studies diffuse atrophy of the testes and/or degeneration of the germinative epithelium were observed from 60 mg/kg bw/d onwards. However as fertility was not affected in the two generation study this effect was not considered relevant for reproductive toxicity.

In the rat developmental toxicity study severe maternal toxicity was observed at the highest dose level of 150 mg/kg, including mortality, clinical signs and reduced body weight and body weight gain. No effects regarding maternal toxicity were observed at the low and mid dose. No effects on gestational parameters or on fetal toxicity were observed at any dose. In conclusion, the NOAEL for maternal toxicity in rats was 75 mg/kg bw/d and the NOAEL for fetal toxicity and teratogenicity was greater than 150 mg/kg bw/d. In the main developmental toxicity study in rabbits no maternal toxicity as well as no effects on gestational parameters and development were observed up to the highest dose tested of 80 mg/kg bw/d.

Taken together, no specific effects on fertility and no developmental toxicity was observed following metaldehyde administration to test animals. No classification is proposed.

4.11.6 Conclusions on classification and labelling

Directive 67/548/EEC: no classification proposed

Regulation (EC) No. 1272/2008: no classification proposed

4.12 Other effects

4.12.1 Non-human information

4.12.1.1 Neurotoxicity

Table 65: Summary table of relevant neurotoxicity studies

Method	Results	Remarks	Reference
Acute neurotoxicity study in rats	0, 75, 150 and 250 mg/kg bw by gavage NOAEL = 75 mg/kg bw Effects at LOAEL: transient clinical signs and findings in neurological screening	-	Haferkorn, J.; 2009
90-day repeated dose oral neurotoxicity study in rats	0, 100, 500 and 2500 ppm/diet (equivalent to 0, 8, 39 and 185 mg/kg bw/d)	-	Jones, L., Finn, J., Mullee, D.; 2003
Relevant aspects of neurological effects associated with exposure to Metaldehyde	Neurofunctional effects of metaldehyde are consequences of acute toxicity; no classical neurotoxin	Position paper	Harder, V., Roth, T., Hofer, M.; 2010

Reference: Acute neurotoxicity study in rats by oral administration of

metaldehyde

Author(s), year: Haferkorn J., 2009

Report/Doc. number: LPT Laboratory of Pharmacology and Toxicology, Hamburg, Germany;

LPT Report No. 23443, Doc. No.: 541-002

Guideline(s): EC method B.43; OECD Guideline 424; OPPTS 870.6200

GLP: Yes Deviations: No Validity: Yes

Material and methods:

Test material META Metaldehyde techn.

Lot/Batch 38337 Purity 99.6% Vehicle Corn oil

Species Rat, males and females Strain CD, Crl:CD(SD)

Age Males: 44-47 days, Females: 50-53 days Weight at dosing Males: 178-228 g; Females: 159-198 g

Source Charles River Laboratories, Sulzfeld, Germany Number of animals 40 males, 40 females (10 animals/sex/group)

Metaldehyde was administered orally by gavage as single administration. Three dose levels and one control group were tested. Dose level groups of 20 (10 male and 10 female) animals each were used for functional testing and detailed clinical observations. 10 (5 male and 5 female) animals from each dose level group were selected for perfusion in situ and neurohistopathology. In case animals showed neurotoxic effects those animals were included in the subgroup selected for perfusion in situ and neurohistopathology. At the start of the experiment the animals were weighted and assigned to the groups by randomization. The animals were kept singly in Makrolon cages.

Table 66:	Acute neurotoxicity study: Study design	1

Group	Metaldehyde dose (mg/kg bw, p.o.)	Number and sex of animals
1	0	10 m
	Control	10 f
2	75	10 m
	(low dose)	10 f
3	150	10 m
	(intermediate dose)	10 f
4	250	10 m
	(high dose)	10 f

Test item stability, its concentration and homogeneity in the vehicle was shown to be adequate in a separate study (Flügge, 2009, Doc. No. 411-002).

Observations:

General health condition: All animals were checked at least once daily at the same time for clinical signs. Daily cage-side observations included skin/fur, eyes, mucous membranes, respiratory and circulatory systems, somatomor activity and behaviour patterns.

Mortality/Morbidity: All animals were checked at least twice daily for signs of morbidity or mortality. Premortal symptoms were recorded in detail; as soon as possible after exitus, a post mortem examination was performed.

Body weight: The weight of each rat was recorded on the day of treatment and weekly thereafter throughout the experimental period.

Food and water consumption: The quantity of food left by individual animals was recorded on a weekly basis. Drinking water consumption was monitored daily by visual appraisal throughout the study.

Detailed clinical observations were made before the administration, 2 and 8 hours after administration and during the 14 day recovery period at days 7 and 14 after dosing. These observations were made outside the home cage in a standard arena and at the same time. Signs noted included changes in skin, fur, eyes, mucous membranes, occurrence of secretions and excretions and autonomic activity (e.g. lacrimation, pilo-erection, pupil size, unusual respiratory pattern). Changes in gait, posture and response to handling as well as the presence of clonic or tonic movements, stereotypies (e.g. excessive grooming, repetitive circling) or bizarre behaviour (e.g. self-mutilation, walking backwards) would have been recorded using the scoring systems that include criteria or scoring scales for each measurement in the observations.

Functional tests:

The functional tests were performed before dosing, at 2 and 8 hours after administration and 7 and 14 days after dosing in all animals.

Righting reflex

The animal was grasped by its tail and flipped in the air (approx. 60 cm) above the cart surface so that it turned head over heels. The normal animal should land squarely on its feet in which case the score sheet space was given as '0'. If it landed on its side, 1 point was scored; if it landed on its back, 2 points were scored. This test was repeated five times and the total scores were recorded.

Body temperature

An electronic probe thermometer (with a blunt probe) was used to take a rectal temperature, being allowed to equilibrate for 30 seconds before the reading was recorded.

Salivation

Discharge of clear fluid from mouth, most frequently seen as beads of moisture on lips in rats. Normal state was to see none, in which case the score sheet space was given as '0'. If present, a plus sign was recorded.

Startle response

With the animal on the cart, the metal cage was struck with the blunt probe. The normal animal should exhibit a marked but short-lasting response, in which case the space on the scoring sheet was given as '0'. If present, a plus sign was entered.

Respiration

While at rest on the cart, the animal's respiration cycle was observed and evaluated in terms of a scale from 1 (reduced) to 5 (increased), with 3 being normal.

Mouth breathing

Rats are normally obligatory nose-breathers. Each animal was observed, whether it was breathing through its mouth or not. In the normal case a '0' was given. If present, a plus sign was recorded.

Urination

When an animal was removed from its cage, the pan beneath the animal's cage was examined while returning the animal to its cage. The signs of urination were evaluated on a scale of 0 (lacking) to 5 (polyurea).

Convulsions

If clonic or tonic convulsions were observed to occur, they were graded on intensity (from 1, minor, to 5, marked) and the type and intensity were recorded. In the normal case a '0' was given.

Pilo-erection

The fur of the animal's back was observed, whether it was raised or elevated. In the normal case (no pilo-erection) a '0' was given. If pilo-erection was present, a plus sign was entered.

Diarrhea

In examining the pan beneath an animal's cage, it was noted if there were any signs of loose or liquid stools. Normal state was for there to be none ('0'), in case of diarrhea the intensity was recorded on a scale of 1 (slight) to 5 (much increased).

Pupil size

The pupils were determined if they were constricted or dilated and were graded on a scale of 1 (constricted) to 5 (dilated), with 3 being normal.

Pupil response

The beam of light from the pen light was played across the eyes of the animal and the changes in pupil size were noted. In the normal animal, the pupil is constricted when the beam is on it and then dilate back to normal when the light is removed ('+' = normal). It was noted if there was no response (in which case a minus sign was recorded in the blank space).

Lacrimation

The animal was observed for the secretion and discharge of tears. In rats the tears contain a reddish pigment. No discharge is normal and in this case the box was given as '0'. If the discharge was present, a plus sign was entered.

Impaired gait

The occurrence of abnormal gait was evaluated. The most frequent impairments were waddling (W), hunched gait (H), or ataxia (A, the inability of all the muscles to act in unison). The extent of any impairment was recorded on a scale of 1 (slight) to 5 (marked). The normal reaction was given as '0'.

Stereotypy

Each animal was evaluated for stereotypic behaviour (isolated motor acts or partial sequences of more complex behavioural patterns, occurring out of context and with an abnormal high frequency). These were graded on a scale of 0 (= normal) to 5 if such signs were present.

Toe pinch

The blunt probe was used to bring pressure to bear on one of the digits of the hindlimb. This should evoke a response from the normal animal, graded on a scale from 1 (absent) to 5 (exaggerated); 3 = normal.

Tail pinch

The procedure detailed above was utilised with the animal's tail instead of its hindlimb and was graded on the same scale with 1 (absent) to 5 (exaggerated); 3 = normal.

Wire maneuver

The animal was placed on the metal rod suspended parallel to the cart approx. 60 cm above it. Its ability to move along the rod was evaluated. If impaired, a score of 1 (slightly impaired) to 5 (unable to stay on wire) was recorded. The normal reaction was given as '0'.

Hind leg splay

Using an ink pad, the hind paws were marked with ink. The rat was then held 30 cm above a sheet of blotting paper on the cart. The animal was dropped and the distance between the prints of the two hind paws was measured.

Positional passivity

When placed in an awkward position (such as on the edge of the top of the wire bottomed cage) on the cart surface, it was examined whether the animal immediately moved into a more normal position ('0'). If not, a score was recorded on a scale of 1 (slightly impaired) to 5 (cataleptic).

Tremors

Periods of continued fine movements, usually starting in the limbs (and perhaps limited to them). The normal case is to have none, in which case a '0' was recorded. If present, they were graded on a scale of 1 (slight and infrequent) to 5 (continuous and marked).

Positive geotropism

The animal was placed on the inclined (at an angle of approx. 30°) top surface of the wire cage with its head facing downward. It should turn 180° and face 'uphill', in which case a '0' was given. If this did not occur, a negative sign was recorded.

Limb rotation

One of the animal's hindlimbs was taken and moved through its normal plane of rotation. In the normal state, it should rotate readily but there should be some resistance. The variations from normal were from no resistance (1) to markedly increased resistance or rigidity (5), with 3 being normal.

Auditory function

Each animal was placed into a container and observed for Preyer's reflex (twitching of the pinna) in response to a high frequency sound stimulus. The stimulus was repeated, if necessary, up to 3 times. A normal reaction was given as '+' in the blank, a lacking reaction was indicated by '0'.

Grip strength

Prior to testing, the gauge (Chatillon, Model DPP - 1.0 kg) was calibrated with a set of known weights and the apparatus adjusted for the size of the animal (about 1 cm clearance on both sides of the animal). After the strain gauge was zeroed and set in the record mode, the animal was placed into the trough with the forepaws inside the triangular grasping ring. Using one hand, the animal was grasped about 2.5 cm of the way up toward the base of the tail and steadily pulled (approx. 2.5 cm/sec) away from the ring until the grip was broken. The animal continued to be pulled along the trough until the hindlimbs grasped the T-bar. The trial was completed when grip of the hindlimbs was broken. Three successive readings were taken for each animal with an intertrial interval long enough to record the data and zero both meters for the next trial.

Locomotor activity

The rats were individually placed into a motility system with microprocessor control and automatic statistical evaluation. Each unit had 2 individually adjustable channels so that slight static movements and active moving could be differentiated and separately scored (TSE Systems, Bad Homburg). The motility meter created a magnetic field, movement of the animals interfered with the magnetic field. Two separate magnetic fields allowed two sensitivity levels to be chosen to distinguish between two types of movements:

- -High sensitivity: Stereotype, static movement, (static movements) (e.g. grooming, a stationary movement of the animal without leaving its own position)
- -Low sensitivity: Active locomotion (active moving)

Necropsy and histopathology:

The nervous system from 5 animals/sex/group was fixed in situ according to standard perfusion procedure for neurohistopathological examination. The following organs or parts of organs of all animals were fixed in 7% buffered formalin except the eyes which were preserved in Davidson's solution for optimum fixation: center of cerebrum (incl. hippocampus, midbrain, cerebellum, pons, medulla oblongata), cauda equina, dorsal root ganglia, dorsal and ventral root fibres, eye with optic nerve and retina (2), forebrain, muscle, skeletal, calf, sciatic nerve (proximal), spinal cord (cervical and lumbal swellings), tibial nerve (proximal, knee) and tibial nerve (calf muscle branches).

The afore-listed organs of 5 animals/sex of groups 1 and 4 were examined neurohistologically after preparation of sufficient paraffin sections with haematoxylin-eosin staining. The spinal cord and peripheral nerve sections included both cross or transverse and longitudinal sections. Attention was

paid to the vasculature of the nervous system. In addition, to identify the dorsal root ganglia that were not present in the original slide, several H&E stained step sections seperated by approximately 100 µm of the cervical and/or lumbar spinal cord (transverse and longitudinal section) were prepared in a number of animals (nos. 1, 2, 3, 6, 7, 11, 12, 16, 17, 18, 61, 62, 63, 69, 70, 73, 75, 78 and 80) and also examined microscopically. The frequency and severity of each lesion were recorded.

Findings:

Mortality: 5 of 10 female rats treated with 250 mg/kg died prematurely within 24 hours after the administration.

<u>Clinical signs</u>: None of the male and female rats treated with 75 mg/kg revealed any changes of behaviour or external appearance. Oral treatment with 150 mg/kg caused slight tremor in females. The high dose of 250 mg/kg caused death (5/10), reduced motility, ataxia, tremor, reduced muscle tone, tonic convulsions, piloerection and diarrhea in all females on test day 1. All high dosed males revealed tremor and diarrhea on test day 1. All casualities and clinical symptoms started 1 to 6 hours after administration and were limited to the day of administration.

<u>Food and drinking water consumption</u>: Food consumption was slightly decreased in the 5 surviving females of the high dose group during the experimental phase. Drinking water consumption was not changed.

Body weights and body weight gain: There was no treatment-related effect observed at any dose level 8 and 14 days after administration.

Table 67: Mortality, clinical signs, body weight and food consumption

Symmtoma!	Con	ntrol	75 mg	/kg bw	150 m	g/kg bw	250 m	g/kg bw
Symptoms/ Criteria	m	f	m	f	m	f	m	f
Criteria	(n=10)	(n=10)	(n=10)	(n=10)	(n=10)	(n=10)	(n=10)	(n=10)
Clinical signs								
Reduced motility	none	none	none	none	none	none	none	+ to ++ 3 to 6h (10)
Ataxia	none	none	none	none	none	none	none	+ 6h (10)
Tremor	none	none	none	none	none	+ 6h (10)	+ 3 to 6h (10)	+ to ++ 3 to 6h (10)
Reduced muscle tone	none	none	none	none	none	none	none	++ 6h (10)
Tonic convulsions	none	none	none	none	none	none	none	+ 6h (10)
Pilo-erection	none	none	none	none	none	none	none	+ 3 to 6h (8)
Diarrhea	none	none	none	none	none	none	+ 1 to 3h (10)	+ 1 to 3h (10)
Mortality								
Within 6h	0	0	0	0	0	0	0	0

Symptoma!	Control		75 mg	/kg bw	150 mg	g/kg bw	250 mg/kg bw	
Symptoms/ Criteria	m (n=10)	f (n=10)	m (n=10)	f (n=10)	m (n=10)	f (n=10)	m (n=10)	f (n=10)
Within 24h	0	0	0	0	0	0	0	5
Within 7d	0	0	0	0	0	0	0	5
Within 14d	0	0	0	0	0	0	0	5
Mean body weight [g] / (Mean l	ody weight	gain [%])					
Before admin.	196.2	173.7	204.5	177.7	203.3	180.2	193.7	170.7
Test day 8	265.7 (+35.5)	203.2 (+17.1)	279.8 (+37.2)	208.3 (+17.3)	277.5 (+36.9)	213.9 (+18.8)	262.9 (35.9)	198.6 (+15.1)
Test day 15	309.3 (+57.9)	219.6 (+26.5)	314.3 (+54.2)	214.8 (+20.9)	310.7 (+53.3)	224.6 (+24.7)	306.3 (+58.3)	207.8 (+20.4)
Food consumption [Food consumption [g/week/rat]							
Testweek 1	173.6	134.3	189.5	135.9	184.7	139.6	172.8	110.0
Testweek 2	149.5	118.5	170.0	139.5	159.3	131.7	153.5	104.4

h: hours p.a.; d: days p.a..; +: slight/observed; ++: moderate

<u>Neurological screening</u>: No treatment-related effects were observed after administration of 75 mg/kg. Oral treatment with 150 and 250 mg/kg caused changes in the appearance, behaviour and reactivity of male and female rats 2 and 8 hours after administration. No changes were observed in control animals except for the extreme low values for "hind-leg splay" 8 hours and 7 days after treatment compared to the other observation time points.

Pilo-erection, diarrhea and impaired ability for wire maneuver were noted in all male rats treated orally with 150 mg/kg. The female rats revealed the same symptoms and, in addition, impaired gait, tremor and decreased resistance during limb rotation. These findings were observed 8 hours after treatment and subsided 7 days after administration.

The high dose of 250 mg/kg metaldehyde revealed further signs of toxicity in form of tremor in the male rats and decreased hindleg splay in the female rats 2 hours after administration. Furthermore, significantly increased body temperature was observed in both sexes and a reduced righting reflex, convulsions, impaired gait, a reduced toe/tail pinch response and a decreased resistance during limb rotation were noted in the female rats 8 hours after administration. A significantly increased hindleg splay was noted for the females 8 hours after administration but as the values of the controls were extremely low compared to other scoring time points, it is considered to be spontaneous.

All clinical changes had sibsided on test day 2 and all neurological changes on the next examination time point for neurological screening on test day 7.

The study authors considered all symptoms observed, especially in the 250 mg/kg dose group with a mortality rate of 50 % in females, to be related to the overall toxicity of metaldehyde at lethal or close-to-lethal doses and not signs of neurological toxicity.

Table 68: Grip strength and locomotor activity

Symptoms/ Time point	Con	Control		/kg bw	150 mg	150 mg/kg bw		250 mg/kg bw	
	m	f	m	f	m	f	m	f	
	(n=10)	(n=10)	(n=10)	(n=10)	(n=10)	(n=10)	(n=10)	(n=10)	
Grip strength forelimb [g]									
pre-dose	306.0	297.6	320.1	297.0	328.4	278.9	322.7	300.8	
2 hours p.a.	303.6	298.3	320.4	295.7	333.7	298.2	317.4	290.8	
8 hours p.a.	334.2	293.0	323.9	288.9	318.2	277.3	244.6	nd	
7 days <i>p.a.</i>	390.2	386.4	374.9	342.5	389.1	348.0	404.2	393.7	
14 days <i>p.a.</i>	399.2	348.9	356.5	327.8	383.6	326.7	387.4	298.8	

Samuel	Cor	ntrol	75 mg	/kg bw	150 mg	g/kg bw	250 mg	g/kg bw
Symptoms/ Time point	m	f	m	f	m	f	m	f
	(n=10)	(n=10)	(n=10)	(n=10)	(n=10)	(n=10)	(n=10)	(n=10)
Grip strength hindl	imb [g]							
pre-dose	138.7	137.6	134.9	129.6	139.8	132.3	141.1	144.2
2 hours p.a.	148.9	138.3	139.7	123.0	148.0	134.5	132.1	137.4
8 hours <i>p.a.</i>	148.0	134.9	140.0	131.4	133.1	113.3	104.3	nd
7 days <i>p.a</i> .	274.2	250.9	136.4	128.2	145.0	130.7	307.2	263.7
14 days <i>p.a</i> .	158.1	139.3	142.1	138.8	147.7	134.3	145.5	131.1
Spontaneous motilit	ty – high sen	sitivity (slig	ht moveme	nts) [numbe	r of moveme	ents/10 min]		
pre-dose	685.2	687.0	697.3	860.8	894.4	800.5	815.3	832.8
2 hours p.a.	320.4	281.2	526.7	706.0	349.7	612.2	549.8	617.1
8 hours p.a.	398.4	307.5	453.0	545.0	590.9	652.4	531.4	555.8
7 days <i>p.a</i> .	503.7	570.2	570.0	811.7	561.3	998.6	630.0	930.6
14 days <i>p.a</i> .	340.7	597.0	516.7	778.1	706.4	858.6	591.2	763.2
Spontaneous motilit	ty – low sens	sitivity (activ	ve moving) [number of 1	movements/	10 min]		
pre-dose	140.5	106.4	119.2	145.8	178.9	141.8	154.0	131.9
2 hours p.a.	36.1	29.5	68.1	82.5	45.4	69.6	60.3	70.9
8 hours p.a.	53.1	33.5	69.6	82.3	87.9	79.4	71.5	73.3
7 days <i>p.a</i> .	103.2	93.5	109.1	146.3	97.0	216.5	117.4	163.0
14 days <i>p.a</i> .	63.5	111.9	92.7	133.1	139.0	155.0	108.6	104.6

nd: not determined; **: stat. sign. with $p \le 0.01$

Table 69: Neurological screening, functional observations

	Cor	ntrol	75 mg	/kg bw	150 mg	g/kg bw	250 mg	g/kg bw
Parameter	m	f	m	f	m	f	m	f
	(n=10)	(n=10)						
		Predose / 2h a	fter dosing / 8h af	ter dosing / 7d afte	er dosing / 14d afte	r dosing		
Rightning reflex	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/ 1.0 /-/-
Body temperature [°C]	36.7/37.0/36.9/	37.0/37.1/37.0/	37.0/36.9/37.1/	36.8/37.0/37.0/	37.0/37.1/37.3/	36.9/37.1/37.3/	37.1/37.2/ 37.5 */	37.0/37.2/ 38.0* /
	37.3/37.6	37.4/37.7	37.5/37.8	37.5/37.8	37.5/37.8	37.6/37.8	37.3/37.8	37.2/37.7
Salivation	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-
Startle response	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-
Respiration	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-
Mouth breathing	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-
Urination	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-
Convulsion	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/2.0/-/-
Pilo-erection	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/+/-/-	-/-/+/-/-	-/-/+/-/-	-/+/+/-/-
Diarrhea	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/1.0/-/-	-/-/1.0/-/-	-/-/1.0/-/-	-/-/1.0/-/-
Lacrimation	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-
Impaired gait	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/1.0/-/-	-/-/-/-	-/2.0/3.0/-/-
Stereotypy	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-
Toe pinch	-/-/2.9/3.0	-/-/3.0/3.2/-	-/-/-3.0/3.0	-/-/3.0/3.0/-	-/-/-3.0/3.0	-/-/3.0/3.0/-	-/-/-3.1/3.0	-/-/ 2.0 /3.4/-
Tail pinch	-/-/2.9/3.0	-/-/3.0/2.9/-	-/-/-3.0/3.0	-/-/3.0/3.0/-	-/-/-3.0/3.0	-/-/3.0/3.0/-	-/-/2.9/3.0	-/-/ 2.0 /3.0/-
Wire maneuver	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/1.0/-/-	-/-/2.0/-/-	-/-/1.0/-/-	-/-/-/-
Hind leg splay	6.5/6.9/6.1/6.5/ 6.9	5.2/5.2/4.9/4.9/ 5.4	6.4/6.7/6.7/6.8/ 7.0	5.2/5.2/5.1/6.1/ 5.4	6.7/6.7/6.3/6.9/ 7.0	5.3/5.5/5.4/6.0/ 5.5	6.6/6.6/6.7/6.6/ 6.9	5.6/4.9/ 6.7* ¹ /5. 1/ 5.3
Positional passivity	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-
Tremors	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/1.0/-/-	-/1.0/1.0/-/-	-/1.0/3.0/-/-
Positive geotropism	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/nd/-/-	-/-/nd/-/-	-/-/nd/-/-	-/nd/nd/-/-
Limb rotation	-/-/-/-	-/-/3.0/-/-	-/-/-/-	-/-/3.0/-/-	-/-/-/-	-/-/2.0/-/-	-/-/-/-	-/-/1.0/-/-
Auditory function	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-

^{-:} no influence; +: slight/observed; ++:moderate; nd: not determined; *: stat. sign. with $p \le 0.01$

¹⁾ considered to be spontaneous due to low control values at 8h after dosing

<u>Necropsy and histopathology</u>: Administration of 75, 150 and 250 mg/kg metaldehyde did not produce any macroscopic or microscopic findings considered to be related to treatment. There were no microscopic alterations recorded in the nervous tissues or in the vasculature of the nervous tissue.

Conclusion:

CD-1 rats were treated with a single dose of 75, 150 or 250 mg/kg metaldehyde orally by gavage. 5 of 10 female rats treated with 250 mg/kg died prematurely within 24 hours after administration. Oral treatment with 150 mg/kg and 250 mg/kg caused transient clinical signs and findings in the neurological screening with were recorded only on the administration day with females being more affected than males. The neurological screening revealed several changes like pilo-erection, tremor, diarrhea, increased body temperature, convulsions, impaired ability for wire maneuver, impaired gait, decreased resistance during limb rotation, increased hindleg splay, a reduced righting reflex and/or a reduced toe/tail pinch response. The histomorphological examination of the nervous system did not reveal any pathological evidence. The study author concluded that all neurological changes should be regarded related to the overall toxicity of metaldehyde at lethal or close-to-lethal doses and not to signs of neurological toxicity, as no histopathological changes were found. However, the RMS feels that the effects observed in the neurological screening at 150 and 250 mg/kg cannot be solely attributed to general acute toxicity, as they already appear at the non-lethal dose level of 150 mg/kg. Furthermore, even if these findings are acute toxic effects, they are also neurotoxic effects. The NOAEL of this study for both the systemic toxicity and the neurotoxicity is therefore set at 75 mg/kg bw metaldehyde, based on the findings in the neurological screening at the dose level of 150 mg/kg.

Reference: LZ1060, metaldehyde: ninety day repeated dose oral (dietary)

neurotoxicity study in the rat

Author(s), year: Jones L., Finn J., Mullee D., 2003

Report/Doc. number: Safepharm Laboratories Limited, Derbyshire, UK

SPL Project Number: 102/420; Lonza Report No.3644; Doc. No. 533-004

Guideline(s): OECD Guideline 424 (1997); JMAFF Guideline 12 Nohsan No.8147

(2000)

GLP: Yes Deviations: No Validity: Yes

Material and Methods:

10 male and 10 female Sprague-Dawley Crl:CD rats (source: Charles River Ltd., Kent, UK) were fed diets containing concentrations of 0, 100, 500 and 2500 ppm metaldehyde (LZ1060, batch no. 31509, purity 98.3 %) over a period of ninety days. At the start of treatment the animals were six weeks old and weighed 179-243 g (males) and 143-186 g (females). The mean achieved dose levels were equivalent to 0, 8, 39 and 185 mg/kg bw/d. Dietary admixtures were prepared prior to treatment and then twice during the three month study period at approximately monthly intervals. They were shown to be stable for 6 weeks. Homogeneity and concentration of the samples were confirmed by analysis.

<u>Clinical observations</u>: All animals were examined for overt signs of toxicity or behavioural changes once daily.

<u>Functional observations</u>: Prior to the start of treatment and during weeks 2, 4, 8 and 12, all animals were observed for signs of functional/behavioural toxicity together with functional performance

tests and an assessment of sensory reactivity to different stimuli. Behavioural assessment was performed for each animal using a purpose built arena, including the following parameters: gait, tremors, twitches, convulsions, bizarre/abnormal/stereotypic behaviour, salivation, piloerection, exophthalmia, lachrymation, hyperthermia, hypothermia, skin color, respiration, palpebral closure, urination, defecation, transfer arousal and tail elevation. Functional performance tests included motor activity and forelimb/ hindlimb grip strength. Motor activity was assessed with purpose built automated activity monitors. Animals of one sex were tested at each occasion and were randomly allocated to the activity monitors. The tests were performed at approximately the same time each day, under similar laboratory conditions. The percentage of time each animal was active and mobile was recorded for a one hour period. Forelimb/ hindlimb grip strength was tested using an automated grip strength meter. Each animal was allowed to grip the proximal metal bar of the meter with its forepaws. The animal was pulled by the base of the tail until its grip was broken. The animal was drawn along the trough of the meter by the tail until ist hind paws gripped the distal metal bar. The animal was pulled by the base of the tail until its grip was broken. A record of the force required to break the grip for each animal was made. Three consecutive trials were performed for each animal. Sensory reactivity: Each animal was individually assessed for sensory reactivity to auditory, visual and proprioceptive stimuli. The following parameters were observed: grap response, vocalisation, toe pinch, tail pinch, finger approach, touch escape, pupil reflex, startle reflex and blink reflex.

Assessment of motor activity:

Twenty purpose built 44 infra-red beam automated activity monitors were used to assess motor activity. Animals of one sex were tested at each occasion and were randomly allocated to the activity monitors. The tests were performed at approximately the same time each day, under similar laboratory conditions. The evaluation period was one hour for each animal. The percentage of time each animal was active and mobile was recorded for a one hour period and also during the final 20% of the period (considered to be the asymptotic period).

Bodyweight: Individual body weights were recorded on day 0 and at weekly intervals thereafter.

<u>Food and water consumption</u>: Food consumption was recorded for each cage group at weekly intervals throughout the study. Water intake was observed daily, for each cage group, by visual inspection of the water bottles for any overt changes.

<u>Ophthalmoscopic examination</u>: The eyes of all control and high dose animals were examined pretreatment and during the final week of the study. Examinations included observation of the anterior structures of the eye, pupillary and corneal blink reflexes, and following pupil dilation with tropicamide solution, detailed examination of the internal structure of the eye.

<u>Pathology</u>: All surviving animals were killed by intravenous overdose of sodium pentobarbitone. Five males and five females from each dose group were then perfused with glutaraldehyde:paraformaldehyde via the heart, following initial perfusion with heparinised saline. The remaining animals were subjected to gross examination only, and any macroscopic abnormalities were recorded.

Organ weights: The brain from all perfused animals was weighed following immersion in buffered 10 % formalin.

<u>Histopathology</u>: Microscopic evaluation was performed on tissues from the perfused control and 2500 ppm dose groups. Samples of the following tissues from all animals were immersed in buffered 10 % formalin (spinal cord dissection was performed to reveal dorsal and ventral root ganglia/fibres following fixation): brain (olfactory bulb, forebrain centre of cerebrum including hippocampus, midbrain, cerebellum, pons and medulla oblongata), dorsal root ganglia (cervical and

lumbar regions), dorsal and ventral root fibres (longitudinal cervical and lumbar sections), eyes (longitudinal sections), optic nerve (longitudinal sections), sciatic nerve (proximal-longitudinal and transverse sections), tibial nerve (proximal at knee and calf muscle branches – longitudinal and transverse sections), skeletal calf muscle (transverse sections), spinal cord (longitudinal and transverse cervical and lumbar sections).

Findings:

Clinical observations: A female receiving 2500 ppm showed loss of limb function on day 10 together with increased respiratory rate. The animal was housed individually and breathing recovered by day 13. Hind limb function, however, remained absent. The condition of the animal remained stable with full forelimb movement, but grooming appeared to regress over the following week. In the study report it is stated that loss of the hind limb function in this animal was considered to result from spinal cord injury, even though the lesion was not histopathologically examined. According to the study author in the absence of any similar effects in the other animals throughout the treatment period, this neurotoxic finding is considered to be consistent with an episode of acute toxicity associated with the large of metaldehyde administered at the beginning of the study. The doses received by the concerned animal can be estimated to be in excess of 240 mg/kg, which clearly lays in the acute toxic range of metaldehyde. The condition of the animal failed to improve and it was humanely killed on day 22. There was no evidence of neurotoxicity in any of the 2500 ppm animals. Incidents of hunched posture (and one incident of tiptoe gait on day 14) were noted in one male and three females from day 28 to day 31 but these were isolated and transient. The remaining clinical signs were confined to generalised fur loss and scab formation in treated and control animals. One control animal developed a limp on the left hind limb from day 77 onwards. All these findings were regarded as commonly seen, low incidence findings in laboratory rats.

<u>Functional observations</u>: Behavioural assessments: Detailed open-field observations during week 2, 4, 8 and 12 showed no evidence of neurotoxic effects. Functional performance tests: There were no treatment-related changes in the functional performance parameters measured. The statistically significant intergroup differences detected in motor activity parameters were considered to be isolated changes. A slight but statistically significant increase in hind limb grip strength was detected for 2500 ppm males during week 4. The increase was minimal and, in isolation was considered to be fortuitous. Sensory reactivity assessments: A statistically significant increase in startle reflex parameters was detected for 500 ppm females during week 12, but in the absence of a dose-response relationship, the differences were considered to be incidental and of no toxicological importance.

Some statistically significant intergroup differences were detected in motor activity parameters. As there were no clear dose-response-relationships and also large standard deviations occurred, these findings were considered to be incidental.

All group mean motor activity values for males and females are presented in the following table:

Table 70: Group mean motor activity values and standard deviations

Dietary	Number	MALES				FEMALES			
concentration	of	Motor ac	tivity			Motor ac	tivity		
(ppm)	animals	Overall		Final 20%	6 of trial	Overall		Final 20%	% of trial
		%	%	%	%	%	%	%	%
		activity	mobile	activity	mobile	activity	mobile	activity	mobile
			activity		activity		activity		activity
PRETEST									
0 (control)	10	19.6	5.0	9.2	0.8	8.2	0.1	26.9	7.6
		Sd 4.3	Sd 1.2	Sd 7.6	Sd 1.5	Sd 5.7	Sd 0.2	Sd 10.2	Sd 3.7
100	10	29.9	4.6	25.0	0.9	11.5	0.6	30.3	9.7

	Dietary	Number	MALES				FEMALI	ES			
No. No.	concentration	of	Motor ac	tivity			Motor activity				
	(ppm)	animals	Overall		Final 20%	% of trial	Overall	<u> </u>	Final 20%	% of trial	
School S			%	%			%	%	%	%	
Sd 19.0 Sd 1.5 Sd 28.6 Sd 1.2 Sd 9.6 Sd 1.0 Sd 6.7 Sd 2.6			activity		activity		activity	mobile	activity	mobile	
500				activity		activity		activity		activity	
Sel 15.6 Sel 13.8 Sel 29.5 Sel 0.4 Sel 15.8 Sel 8.2 Sel 10.8 Sel 4.6 Sel 13.8 Sel 4.6 Sel 13.8 Sel 13.6 Sel 1			Sd 19.0	Sd 1.5	Sd 28.6	Sd 1.2	Sd 9.6	Sd 1.0	Sd 6.7	Sd 2.0	
Description	500	10	30.2	6.3	19.8	0.4	13.4	4.3	38.6	13.9	
WEEK 2 WEEK 2 0 (control) 10 8.4 0.4 18.8 5.1 29.7 4.7 41.4 8.9 100 10 8.8 0.9 18.3 4.6 15.0 3.8 44.3 13.8 100 10 5.8 0.9 18.3 4.6 15.0 3.8 44.3 13.8 500 10 *18.2 1.6 35.9 7.6 41.2 5.2 56.5 15.5 500 10 **18.2 1.6 35.9 7.6 41.2 5.2 56.5 15.5 2500 10 **12.2 3.4 29.6 8.2 35.6 8.3 40.6 11.2 56.7 56.5 56.5 56.5 56.5 56.5 56.5 56.5 15.5 44.2 14.2 12.2 31.1 74 44.2 44.2 14.2 14.2 31.1 74 44.2 14.2 14.2 14.1 36.6 77.7 <td< td=""><td></td><td></td><td>Sd 15.6</td><td>Sd 1.3</td><td>Sd 29.5</td><td>Sd 0.4</td><td>Sd 15.8</td><td>Sd 8.2</td><td>Sd 10.8</td><td>Sd 4.6</td></td<>			Sd 15.6	Sd 1.3	Sd 29.5	Sd 0.4	Sd 15.8	Sd 8.2	Sd 10.8	Sd 4.6	
WEEK 2	2500	10	23.4								
O (control)			Sd 6.4	Sd 1.7	Sd 3.9	Sd 1.1	Sd 4.6	Sd 1.3	Sd 13.6	Sd 3.6	
Sd 9.8 Sd 0.5 Sd 6.2 Sd 2.2 Sd 26.7 Sd 5.6 Sd 20.0 Sd 3.2	WEEK 2										
100	0 (control)	10	8.4	0.4	18.8	5.1	29.7	4.7	41.4	8.9	
Sd 7.0 Sd 1.0 Sd 6.7 Sd 1.8 Sd 19.9 Sd 6.4 Sd 11.0 Sd 3.5			Sd 9.8	Sd 0.5	Sd 6.2	Sd 2.2	Sd 26.7	Sd 5.6	Sd 20.0	Sd 3.3	
Signature Sign	100	10	5.8	0.9	18.3	4.6		3.8	44.3	13.8	
Sd 32.5 Sd 3.4 Sd 24.5 Sd 3.4 Sd 34.6 Sd 4.9 Sd 12.9 Sd 4.1			Sd 7.0	Sd 1.0	Sd 6.7	Sd 1.8	Sd 19.9	Sd 6.4	Sd 11.0	Sd 3.9	
The image of the	500	10	*18.2		35.9	7.6	41.2	5.2	56.5	15.5	
WEEK 4 Sd 21.9 Sd 7.8 Sd 10.6 Sd 3.7 Sd 21.6 Sd 6.7 Sd 3.0 WEEK 4 0 (control) 10 10.8 0.4 20.4 4.2 14.2 1.2 31.1 7.4 100 10 14.4 0.6 27.8 5.6 27.7 6.7 37.1 8.7 500 10 15.5 1.2 27.8 5.6 27.7 6.7 37.1 8.7 500 10 15.5 1.2 27.8 5.9 40.2 7.5 46.9 *11.7 500 10 4.9 1.1 23.5 6.6 40.5 50.6 80.1.8 80.2 7.5 46.9 *11.7 2500 10 4.9 1.1 23.5 6.6 40.5 50.7 80.9 41.2 \$2.7 2500 10 21.2 2.7 29.5 8.2 17.3 1.6 36.0 7.0 WEEK 8 0			Sd 32.5	Sd 3.4	Sd 24.5	Sd 3.4	Sd 34.6	Sd 4.9	Sd 12.9	Sd 4.1	
WEEK 4 0 (control) 10 10.8 0.4 20.4 4.2 14.2 1.2 31.1 7.4 100 10 14.4 0.6 27.8 Sd 2.6 27.7 6.7 37.1 8.7 500 10 15.5 1.2 27.8 5.9 40.2 7.5 46.9 *11.7 500 10 15.5 1.2 27.8 5.9 40.2 7.5 46.9 *11.7 2500 10 4.9 1.1 23.5 6.6 40.5 10.1 43.4 **11.7 2500 10 4.9 1.1 23.5 6.6 40.5 10.1 43.4 **11.7 2500 10 21.2 2.7 8d 15.1 8d 3.3 8d 22.1 8d 8.3 8d 14.1 8d 6.6 WEEK 8 0 (control) 10 21.2 2.7 29.5 8.2 17.3 1.6 36.0 7.0 Sd 25.3 8d 3.7	2500	10	**12.2	3.4	29.6				40.6	11.2	
0 (control) 10 10.8 Sd 13,2 Sd 0.5 0.4 Sd 9.2 Sd 1.3 14.2 Sd 27.9 Sd 3.5 31.1 Sd 9.9 Sd 1.2 100 10 14.4 0.6 27.8 Sd 9.9 Sd 19.5 Sd 2.6 5.6 27.7 6.7 Sd 13.1 Sd 18.6 37.1 8.7 Sd 12.6 500 10 15.5 1.2 27.8 Sd 19.5 Sd 2.6 Sd 38.3 Sd 9.2 Sd 18.6 Sd 1.8 Sd 1.8 Sd 22.9 Sd 1.9 Sd 25.7 Sd 7.6 Sd 14.3 Sd 3.9 34.9 250.0 Sd 19.5 Sd 22.9 Sd 1.9 Sd 25.7 Sd 7.6 Sd 14.3 Sd 3.9 2500 10 4.9 1.1 23.5 6.6 40.5 Sd 22.9 Sd 1.9 Sd 22.1 Sd 8.3 Sd 14.1 Sd 6.6 WEEK 8 0 (control) 10 21.2 2.7 Sd 25.3 Sd 3.7 Sd 11.5 Sd 4.1 Sd 29.5 Sd 3.7 Sd 15.3 Sd 2.4 100 10 20.7 2.1 35.1 8.6 17.0 3.9 40.0 10.8 Sd 12.3 Sd 3.6 500 10 20.7 2.1 35.1 8.6 17.0 3.9 40.0 10.8 Sd 12.3 Sd 3.6 Sd 12.8 Sd 13.1 Sd 2.9			Sd 21.9	Sd 7.8	Sd 10.6	Sd 3.7	Sd 21.6	Sd 6.7	Sd 6.7	Sd 3.0	
Sd 13,2 Sd 0.5 Sd 9.2 Sd 1.3 Sd 27.9 Sd 3.5 Sd 9.9 Sd 1.2	WEEK 4										
100 10 14.4 0.6 27.8 5.6 27.7 6.7 37.1 8.7 500 10 15.5 1.2 27.8 5.9 40.2 7.5 46.9 *11.7 2500 10 4.9 1.1 23.5 6.6 40.5 10.1 43.4 **11.7 2500 10 4.9 1.1 23.5 6.6 40.5 10.1 43.4 **11.7 2500 8 30.7 8d 15.1 8d 3.3 8d 22.1 8d 25.7 8d 14.3 8d 3.9 40 4.9 1.1 23.5 6.6 40.5 10.1 43.4 **11.7 8d 7.8 8d 2.7 8d 15.1 8d 3.3 8d 22.1 8d 8.3 8d 14.1 8d 6.6 WEEK 8 80 2.7 29.5 8.2 17.3 1.6 36.0 7.0 9 60 20.7 2.1 35.1 8.6 17.0 3.9 40.0 10.8	0 (control)	10	10.8	0.4	20.4	4.2	14.2	1.2	31.1	7.4	
Sd 26.0 Sd 0.9 Sd 19.5 Sd 2.6 Sd 38.3 Sd 9.2 Sd 18.6 Sd 1.8			Sd 13,2	Sd 0.5	Sd 9.2	Sd 1.3	Sd 27.9	Sd 3.5	Sd 9.9	Sd 1.2	
500 10 15.5 1.2 27.8 5.9 40.2 7.5 46.9 *11.7 Sd 26.6 Sd 1.5 Sd 22.9 Sd 1.9 Sd 25.7 Sd 7.6 Sd 14.3 Sd 3.9 2500 10 4.9 1.1 23.5 6.6 40.5 10.1 43.4 **11.7 Sd 7.8 Sd 2.7 Sd 15.1 Sd 3.3 Sd 22.1 Sd 8.3 Sd 14.1 Sd 6.6 WEEK 8 0 (control) 10 21.2 2.7 29.5 8.2 17.3 1.6 36.0 7.0 Sd 25.3 Sd 3.7 Sd 11.5 Sd 4.1 Sd 29.5 Sd 3.7 Sd 15.3 Sd 2.4 100 10 20.7 2.1 35.1 8.6 17.0 3.9 40.0 10.8 500 10 15.8 3.0 31.0 8.4 20.7 3.3 47.7 12.6 500 10 30.4 3.7 44.6 9.4 16.5 <td< td=""><td>100</td><td>10</td><td>14.4</td><td>0.6</td><td>27.8</td><td>5.6</td><td></td><td>6.7</td><td>37.1</td><td>8.7</td></td<>	100	10	14.4	0.6	27.8	5.6		6.7	37.1	8.7	
Sd 26.6 Sd 1.5 Sd 22.9 Sd 1.9 Sd 25.7 Sd 7.6 Sd 14.3 Sd 3.9 2500 10 4.9 1.1 23.5 6.6 40.5 10.1 43.4 **11. Sd 7.8 Sd 2.7 Sd 15.1 Sd 3.3 Sd 22.1 Sd 8.3 Sd 14.1 Sd 6.6 WEEK 8 0 (control) 10 21.2 2.7 29.5 8.2 17.3 1.6 36.0 7.0 Sd 25.3 Sd 3.7 Sd 11.5 Sd 4.1 Sd 29.5 Sd 3.7 Sd 15.3 Sd 2.4 100 10 20.7 2.1 35.1 8.6 17.0 3.9 40.0 10.8 500 10 15.8 3.0 31.0 8.4 20.7 3.3 47.7 12.6 2500 10 30.4 3.7 44.6 9.4 16.5 4.2 37.9 10.1 8d 21.8 8d 4.8 8d 21.6 8d 4.2 8d 18.1 8d 5.0 8d			Sd 26.0	Sd 0.9	Sd 19.5	Sd 2.6	Sd 38.3	Sd 9.2	Sd 18.6	Sd 1.8	
2500	500	10	15.5	1.2	27.8	5.9	40.2	7.5	46.9	*11.7	
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WEEK 8 0 (control) 10 21.2 2.7 Sd 29.5 Sd 4.1 Sd 29.5 Sd 3.7 Sd 15.3 Sd 2.4 8.2 17.3 Sd 29.5 Sd 3.7 Sd 15.3 Sd 2.4 10.6 Sd 29.5 Sd 3.7 Sd 15.3 Sd 2.4 10.6 Sd 29.5 Sd 3.7 Sd 15.3 Sd 2.4 10.6 Sd 15.3 Sd 2.4 10.8 Sd 12.3 Sd 2.4 10.8 Sd 12.3 Sd 2.4 10.8 Sd 12.3 Sd 3.6 10.8 Sd 24.1 Sd 24.1 Sd 7.8 Sd 12.3 Sd 3.6 10.8 Sd 12.3 Sd 12.3 Sd 12.3 Sd 12.3 10.8 Sd 12.3 Sd 12.3 Sd 12.3 Sd 12.3 Sd 14.0 Sd 15.3 10.8 Sd 12.3 Sd 12.3 Sd 12.3 Sd 14.0 Sd 15.3 10.8 Sd 14.0 Sd 15.3 Sd 15.1 Sd 15.3 Sd 15.1 Sd 15.3 Sd 15.1 Sd 15.3 Sd 15.1 Sd 15.4 Sd 15.5 Sd 15.5 Sd 15.6 Sd 15.4 Sd 15.5 Sd 15.5 Sd 15.6 Sd 15.4 Sd 15.5 Sd	2500	10	4.9	1.1	23.5	6.6	40.5	10.1	43.4	**11.7	
0 (control) 10 21.2 2.7 29.5 8.2 17.3 1.6 36.0 7.0 100 10 20.7 2.1 35.1 8.6 17.0 3.9 40.0 10.8 500 10 15.8 3.0 31.0 8.4 20.7 3.3 47.7 12.6 500 10 15.8 3.0 31.0 8.4 20.7 3.3 47.7 12.6 Sd 19.8 Sd 4.5 Sd 13.6 Sd 3.8 Sd 20.6 Sd 4.3 Sd 14.0 Sd 5.3 2500 10 30.4 3.7 44.6 9.4 16.5 4.2 37.9 10.1 WEEK 12 5d (control) 10 5.4 0.4 24.4 4.3 12.7 1.3 29.6 6.2 WEEK 12 5d (control) 10 5.4 0.4 24.4 4.3 12.7 1.3 29.6 6.2 100 10 12.9 1.6 27.7 <td< td=""><td></td><td></td><td>Sd 7.8</td><td>Sd 2.7</td><td>Sd 15.1</td><td>Sd 3.3</td><td>Sd 22.1</td><td>Sd 8.3</td><td>Sd 14.1</td><td>Sd 6.6</td></td<>			Sd 7.8	Sd 2.7	Sd 15.1	Sd 3.3	Sd 22.1	Sd 8.3	Sd 14.1	Sd 6.6	
Sd 25.3 Sd 3.7 Sd 11.5 Sd 4.1 Sd 29.5 Sd 3.7 Sd 15.3 Sd 2.4 100 10 20.7 2.1 35.1 8.6 17.0 3.9 40.0 10.8 500 10 15.8 3.0 31.0 8.4 20.7 3.3 47.7 12.6 Sd 19.8 Sd 4.5 Sd 13.6 Sd 3.8 Sd 20.6 Sd 4.3 Sd 14.0 Sd 5.3 2500 10 30.4 3.7 44.6 9.4 16.5 4.2 37.9 10.1 WEEK 12 30.4 3.7 44.6 9.4 16.5 4.2 37.9 10.1 WEEK 12 0 (control) 10 5.4 0.4 24.4 4.3 12.7 1.3 29.6 6.2 WEEK 12 5d 3.9 Sd 0.4 Sd 16.6 Sd 2.4 Sd 14.4 Sd 2.1 Sd 13.3 Sd 2.9 100 10 5.4 0.4 24.4 4.3 12.7 13.8 8.4	WEEK 8										
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Sd 17.3 Sd 1.7 Sd 14.5 Sd 4.2 Sd 24.1 Sd 7.8 Sd 12.3 Sd 3.6 500 10 15.8 3.0 31.0 8.4 20.7 3.3 47.7 12.6 Sd 19.8 Sd 4.5 Sd 13.6 Sd 3.8 Sd 20.6 Sd 4.3 Sd 14.0 Sd 5.3 2500 10 30.4 3.7 44.6 9.4 16.5 4.2 37.9 10.1 WEEK 12 0 (control) 10 5.4 0.4 24.4 4.3 12.7 1.3 29.6 6.2 Sd 3.9 Sd 0.4 Sd 16.6 Sd 2.4 Sd 14.4 Sd 2.1 Sd 13.3 Sd 2.9 100 12.9 1.6 27.7 5.5 18.8 *4.1 44.4 12.0 Sd 15.7 Sd 3.8 Sd 15.1 Sd 3.1 Sd 24.9 Sd 8.9 Sd 13.1 4.3 500 10 6.5 0.9 22.3 5.2 *32.9 *5.6 51.8 <t< td=""><td></td><td></td><td>Sd 25.3</td><td>Sd 3.7</td><td>Sd 11.5</td><td>Sd 4.1</td><td>Sd 29.5</td><td>Sd 3.7</td><td>Sd 15.3</td><td>Sd 2.4</td></t<>			Sd 25.3	Sd 3.7	Sd 11.5	Sd 4.1	Sd 29.5	Sd 3.7	Sd 15.3	Sd 2.4	
500 10 15.8 Sd 19.8 Sd 4.5 Sd 13.6 Sd 3.8 Sd 20.6 Sd 4.3 Sd 14.0 Sd 5.3 2500 10 30.4 Sd 21.8 Sd 4.8 Sd 21.6 Sd 4.2 Sd 18.1 Sd 5.0 Sd 16.1 Sd 6.3 WEEK 12 0 (control) 10 5.4 Sd 3.9 Sd 0.4 Sd 16.6 Sd 2.4 Sd 14.4 Sd 2.1 Sd 13.3 Sd 2.9 100 10 12.9 Sd 15.7 Sd 3.8 Sd 15.1 Sd 3.1 Sd 24.9 Sd 8.9 Sd 13.1 4.3 500 10 6.5 Sd 9.4 Sd 2.2 Sd 10.6 Sd 2.8 Sd 24.6 Sd 5.4 Sd 15.5 Sd 3.2 2500 10 6.8 O.5 Z4.1 Sd 10.6 Sd 2.8 Sd 24.6 Sd 5.4 Sd 15.5 Sd 3.2 2500 10 6.8 O.5 Z4.1 Sd 10.6 Sd 2.8 Sd 24.6 Sd 5.4 Sd 15.5 Sd 3.2	100	10	20.7	2.1	35.1	8.6	17.0	3.9	40.0	10.8	
Sd 19.8 Sd 4.5 Sd 13.6 Sd 3.8 Sd 20.6 Sd 4.3 Sd 14.0 Sd 5.3 2500 10 30.4 3.7 44.6 9.4 16.5 4.2 37.9 10.1 10.1 5.4 5d 21.8 5d 21.6 5d 4.2 5d 18.1 5d 5.0 5d 16.1 5d 6.3 WEEK 12 0 (control) 10 5.4 0.4 24.4 4.3 12.7 1.3 29.6 6.2 6.2 5d 13.3 5d 2.9 5d 13.1 44.4 12.0				Sd 1.7	Sd 14.5	Sd 4.2				Sd 3.6	
2500 10 30.4 Sd 21.8 3.7 Sd 44.6 9.4 Sd 21.6 16.5 Sd 4.2 4.2 Sd 5.0 37.9 Sd 16.1 10.1 Sd 6.3 WEEK 12 0 (control) 10 5.4 Sd 3.9 Sd 0.4 Sd 16.6 Sd 2.4 Sd 14.4 Sd 2.1 Sd 13.3 Sd 2.9 100 10 12.9 1.6 Sd 3.8 Sd 15.1 Sd 3.1 Sd 24.9 Sd 8.9 Sd 13.1 4.3 500 10 6.5 Sd 9.4 Sd 2.2 Sd 10.6 Sd 2.8 Sd 24.6 Sd 5.4 Sd 5.4 Sd 15.5 Sd 3.2 2500 10 6.8 0.5 24.1 5.4 *36.7 *7.5 *45.9 12.1	500	10		3.0	31.0	8.4	20.7	3.3	47.7	12.6	
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WEEK 12 0 (control) 10 5.4 Sd 3.9 Sd 0.4 Sd 16.6 Sd 2.4 Sd 14.4 Sd 2.1 Sd 13.3 Sd 2.9 100 10 12.9 Sd 15.7 Sd 3.8 Sd 15.1 Sd 3.1 Sd 24.9 Sd 8.9 Sd 13.1 4.3 12.7 Sd 3.8 Sd 15.1 Sd 3.1 Sd 24.9 Sd 8.9 Sd 13.1 4.3 500 10 6.5 Sd 9.4 Sd 2.2 Sd 10.6 Sd 2.8 Sd 24.6 Sd 5.4 Sd 5.4 Sd 15.5 Sd 3.2 2500 10 6.8 0.5 24.1 5.4 *36.7 *7.5 *45.9 12.1	2500	10	30.4	3.7						10.1	
0 (control) 10 5.4 Sd 3.9 0.4 Sd 0.4 Sd 16.6 24.4 Sd 2.4 Sd 14.4 Sd 2.1 Sd 13.3 Sd 2.9 100 10 12.9 Sd 15.7 Sd 3.8 Sd 15.1 Sd 3.1 Sd 24.9 Sd 8.9 Sd 13.1 4.3 500 10 6.5 Sd 9.4 Sd 2.2 Sd 10.6 Sd 2.8 Sd 24.6 Sd 5.4 Sd 15.5 Sd 3.2 2500 10 6.8 0.5 24.1 5.4 *36.7 *7.5 *45.9 12.1			Sd 21.8	Sd 4.8	Sd 21.6	Sd 4.2	Sd 18.1	Sd 5.0	Sd 16.1	Sd 6.3	
Sd 3.9 Sd 0.4 Sd 16.6 Sd 2.4 Sd 14.4 Sd 2.1 Sd 13.3 Sd 2.9 100 12.9 1.6 27.7 5.5 18.8 *4.1 44.4 12.0 Sd 15.7 Sd 3.8 Sd 15.1 Sd 3.1 Sd 24.9 Sd 8.9 Sd 13.1 4.3 500 10 6.5 0.9 22.3 5.2 **32.9 **5.6 51.8 12.7 Sd 9.4 Sd 2.2 Sd 10.6 Sd 2.8 Sd 24.6 Sd 5.4 Sd 15.5 Sd 3.2 2500 10 6.8 0.5 24.1 5.4 *36.7 *7.5 *45.9 12.1	WEEK 12										
100 10 12.9 1.6 27.7 5.5 18.8 *4.1 44.4 12.0 Sd 15.7 Sd 3.8 Sd 15.1 Sd 3.1 Sd 24.9 Sd 8.9 Sd 13.1 4.3 500 10 6.5 0.9 22.3 5.2 **32.9 **5.6 51.8 12.7 Sd 9.4 Sd 2.2 Sd 10.6 Sd 2.8 Sd 24.6 Sd 5.4 Sd 15.5 Sd 3.2 2500 10 6.8 0.5 24.1 5.4 *36.7 *7.5 *45.9 12.1	0 (control)	10									
Sd 15.7 Sd 3.8 Sd 15.1 Sd 3.1 Sd 24.9 Sd 8.9 Sd 13.1 4.3 500 10 6.5 0.9 22.3 5.2 **32.9 **5.6 51.8 12.7 Sd 9.4 Sd 2.2 Sd 10.6 Sd 2.8 Sd 24.6 Sd 5.4 Sd 15.5 Sd 3.2 2500 10 6.8 0.5 24.1 5.4 *36.7 *7.5 *45.9 12.1			Sd 3.9	Sd 0.4					Sd 13.3	Sd 2.9	
500 10 6.5 0.9 22.3 5.2 **32.9 **5.6 51.8 12.7 Sd 9.4 Sd 2.2 Sd 10.6 Sd 2.8 Sd 24.6 Sd 5.4 Sd 15.5 Sd 3.2 2500 10 6.8 0.5 24.1 5.4 *36.7 *7.5 *45.9 12.1	100	10									
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2500 10 6.8 0.5 24.1 5.4 *36.7 *7.5 *45.9 12.1	500	10									
										Sd 3.2	
	2500	10									
Sd = Standard deviation			Sd 2.7	Sd 0.5	Sd 6.9	Sd 1.6	Sd 16.2	Sd 6.0	Sd 15.2	Sd 6.3	

Sd = Standard deviation

<u>Body weight</u>: Females receiving 2500 ppm showed a slight but statistically significant reduction in body weight gain during the first week of treatment when compared to controls. Thereafter body weight development was normal. No such effect was observed for 2500 ppm males or for animals of either sex treated with 500 or 100 ppm.

<u>Food consumption</u>: A slight reduction in dietary intake was evident for 2500 ppm females during weeks 3 and 4 but this was considered attributable to the physical condition of the solitary housed

^{*} significantly different from control group p<0.05

^{**} significantly different from control group p<0.01

female which had suffered from spinal injury, and was unrelated to test material toxicity. Food efficiency was unaffected throughout the study period.

Table 71: 90 day neurotoxicity study in Sprague-Dawley Crl:CD rats Body weight / body weight gain

	Dose group level (ppm)									
	Males	_			Femal	es				
	0	100	500	2500	0	100	500	2500		
Body weight (g)										
- day 0	219	220	221	222	164	164	162	164		
- day 90	501	501	516	508	288	294	271	272		
Body weight gain (g)										
- week 1	42	42	45	40	20	21	16	14*		
- week 5	24	20	20	20	9	14	10	9		
- week 9	14	9	8	13	5	6	5	6		
- week 13	11	14	9	10	0	4	5	4		

^{* (}p< 0.05); significantly different from controls

Ophthalmoscopic examination: There were no treatment-related effects observed.

<u>Brain weights</u>: There were no treatment-related changes in brain weights, absolute and relative to body weight.

<u>Necropsy</u>: No macroscopic abnormalities were detected in test or control animals at terminal kill. The 2500 ppm female which developed loss of hind limb function and which was sacrificed prematurely showed no macroscopic abnormalities.

Histopathology: There were no treatment-related changes in the tissues examined.

Conclusion:

No evidence of neurotoxicity was revealed from the subchronic administration of metaldehyde at any dose level tested (100, 500 and 2500 ppm). The NOAEL for neurotoxic effects is therefore considered to be 2500 ppm (equivalent to 185 mg/kg bw/d). At the high dose level, one female showed loss of hind limb function together with an increased respiratory rate. Loss of the hind limb function was considered to result from spinal cord injury, even though the lesion was not histopathologically examined. In the absence of any similar effects in the other animals throughout the treatment period, this finding was considered to be an acute toxic effect. In the females receiving 2500 ppm, a slight and transient reduction of bodyweight gain was detected during the first week of treatment. Based on the finding of reduced body weight gain and the loss of hind limb function in one female at the high dose, the NOAEL for systemic toxicity was considered to be 500 ppm (equivalent to 39 mg/kg bw/d).

In the acute neurotoxicity study, CD rats were treated with a single oral dose of 75, 150 or 250 mg/kg metaldehyde by gavage. 250 mg/kg led to mortality in 5 of 10 female rats within 24 hours after administration. Treatment with 150 mg/kg and 250 mg/kg caused transient clinical signs and findings in the neurological screening with were recorded only on the administration day with females being more affected than males. The neurological screening revealed several changes like pilo-erection, tremor, diarrhea, increased body temperature, convulsions, impaired ability for wire maneuver, impaired gait, decreased resistance during limb rotation, increased hindleg splay, a reduced righting reflex and/or a reduced toe/tail pinch response. The histomorphological examination of the nervous system did not reveal any pathological evidence. The NOAEL of this

study for both systemic toxicity and neurotoxicity is therefore set at 75 mg/kg metaldehyde, based on the findings in the neurological screening at the dose level of 150 mg/kg.

A 90 day neurotoxicity study was conducted in Sprague-Dawley Crl:CD rats. No evidence of neurotoxicity was observed following subchronic administration of metaldehyde at any dose level tested (100, 500 and 2500 ppm). At the high dose level of 2500 ppm (185 mg/kg bw/d), one female showed loss of hind limb function together with an increased respiratory rate. Loss of the hind limb function was considered to result from spinal cord injury, even though the lesion was not histopathologically examined. In the females receiving 2500 ppm, a slight and transient reduction of bodyweight gain was detected during the first week of treatment. The NOAEL for both systemic toxicity and neurotoxicity is considered to be 500 ppm (equivalent to 39 mg/kg bw/d) based on the finding of reduced body weight gain and the loss of hind limb function in one female at 2500 ppm.

In order to clarify the neurotoxic potential of metaldehyde, a position paper was requested from the notifier, as convulsions (or other signs of neurotoxicity) were observed in most of the studies and species (acute oral, 28-d rats, 1-year dogs, chronic toxicity rats, 2 generation rats, developmental rat and rabbit). The **notifier** considers the central nervous system not a specific target organ of short term toxicity of metaldehyde. The following argumentation was provided by the notifier:

- Metaldehyde is rapidly absorbed by rats after oral exposure with no evidence of accumulation as metaldehyde is rapidly and effectively metabolised to acetaldehyde, then further oxidised through the citric acid cycle and the major part expired as carbon dioxide (Selim, 1990, Doc. No. 512-001, see DAR).
- This kinetic profile determines the toxic profile which is characterized by acute and transient effects occurring immediately after dosing in the period of the most elevated plasma concentrations. Studies with repeated daily administration caused repeatedly daily clinical symptoms of acute intoxication which were transient.
- Both single and repeated treatments induced acute toxic effects including partly pronounced neurological symptoms.
- Metaldehyde induced neurofunctional effects like ataxia, twitching, tremors and/or convulsions following repeated exposure. As these effects did not persist, they were not in line with sustained dysfunctions normally induced by classical neurotoxins.
- Because of the distinct differences between Metaldehyde and classical neurotoxins, the term "neurotoxic effect" was not considered adequate to characterize the toxicity profile of metaldehyde.
- In part of the metaldehyde studies, there were individual cases of mortality. These cases were not seen frequently and did not follow a dose and time related incidence pattern that would point towards the presence of cumulative effects or other typical modes of actions related to subchronic dosing. With respect to the above kinetic and toxic profile these cases are interpreted as the combination of a rapid increase of significant metaldehyde plasma concentration with individual cardiovascular susceptibility of the animal in question at the day of administration.
- Hind limb paralysis/paresis due to mechanic vertebral lesions was not very frequently observed. They were considered to occur secondary to pronounced acute neurological symptoms causing mechanical impact on the lumbar region of the vertebral column. A genetic predisposition of certain rat in- and outbred strains for mechanical liability of the lumbar vertebrae could be a contributing factor for the observed lesions. This finding is clearly not caused by a neurotoxic mechanism leading to degeneration or other toxic damage to the central or peripheral nerve tissue. These single cases are relevant for the acute toxicity profile of metaldehyde, but not relevant for classification and labelling as neurotoxicant.

The **RMS** (i.e. dossier submitter) can follow the notifier's arguments that the neurofunctional effects (e.g. tremor, convulsions, ataxia, paresis) following exposure to metaldehyde only occur at doses which are clearly acute toxic and are therefore covered by acute toxicity classification.

4.12.1.2 Immunotoxicity

No data available.

4.12.1.3 Specific investigations: other studies

Not relevant for classification and labelling of metaldehyde.

4.12.1.4 Human information

See also section 4.2 Acute Toxicity

Signs and symptoms of acute poisoning: Some information is presented in the Toxicology Update published in the Journal of Applied Toxicology (*Von Burg R., 1991; Doc.No. 592-028*), which is a brief review on toxicology data found in literature data bases.

However, more detailed information is found in the dissertation of *Borbely A.* (1970, *Doc.No.* 592-001). The neurological symptoms of metaldehyde are described as follows:

Confusion, restlessness, haziness, drowsiness, coma, spasms, tremor, muscle twitching, chorea, abnormal reflexes, ataxia, elevated muscle tone, hypersensitivity, Chvostek's sign, Trousseau's sign, disturbed vision, amnesia, respiratory arrest, risus sardonicus.

4.12.2 Summary and discussion

Neurofunctional effects (e.g. tremor, convulsions, ataxia, paresis) following exposure to metaldehyde are considered to occur only at doses which are clearly acute toxic. These effects do not persist (with exception of hind limb paresis following spinal cord injury) and are not in line with sustained dysfunctions normally induced by classical neurotoxins. Because of the distinct differences between metaldehyde and classical neurotoxins, the term "neurotoxic effect" was not considered adequate to characterize the toxicity profile of metaldehyde. Therefore in the EFSA peer review it has been agreed that metaldehyde does not require classification as neurotoxicant. The accurate conclusion drawn by EFSA is "Acute toxic effects following metaldehyde administration include partly pronounced neurological symptoms, without specific neurotoxic mechanism leading to degeneration or other toxic damage to the central or peripheral nerve tissue. Therefore these reversible effects at high doses are not relevant for classification and labelling as neurotoxicant." (see peer review report, evaluation table, 2.2. point of clarification, http://registerofquestions.efsa.europa.eu/roqFrontend/outputLoader?output=ON-1856)

4.12.3 Comparison with criteria

Not relevant, not required.

4.12.4 Conclusions on classification and labelling

Directive 67/548/EEC: no classification proposed

Regulation (EC) No. 1272/2008: no classification proposed

5 ENVIRONMENTAL HAZARD ASSESSMENT

5.1 Degradation

Table 72: Summary of relevant information on degradation

Method	~ 	Results			Remarks	Reference
Guideline	Type of study	Matrix	Temp.	Result/Half-life		
EPA Subdiv. N, Section 161-1	Hydrolysis	pH 5, 7 and 9, sterile buffer solutions	25 °C	hydrolytically stable		Carpenter M. (1989a)
EPA Subdiv. N, Section 161-2	Photolysis	pH 7, sterile buffer solution	25 °C	photolytically stable		Carpenter M. (1989b)
OECD 301E	Biological degradation (ready)	test medium inoculated with active sewage sludge	24 – 26.5 °C	not readily biodegradable		Wüthrich V. (1990a)
OECD 301F	Biological degradation (ready)	test medium inoculated with active sewage sludge	22 °C	not readily biodegradable		Lebertz, H. (2008)
OECD 302B	Biological degradation (inherent)	test medium inoculated with active sewage sludge	19 – 23.5 °C	not inherently biodegradable		Wüthrich V. (1990b)
BBA part IV-5-1	Water/Sediment study	water: pH 8.4 - 9 sediment: pH 7.8 - 7.9	20 ± 2 °C	Water DT ₅₀ : 11.35 d (S1), 10.25 d (S2) DT ₉₀ : 37.71 d (S1), 34.07 d (S2) Whole system: Metaldehyde DT ₅₀ : 4.10 d (S1), 4.42 d (S2) DT ₉₀ : 13.61 d (S1), 14.97 d (S2) Acetaldehyde DT ₅₀ : 30.98 d (S1), 19.01 d (S2) DT ₉₀ : 102.90 d (S1), 63.14 d (S2)		Möllerfeld J., Römbke J. & Heller M. (1993) – Calculations reviewed June 2009
OECD 308	Water/Sedi- ment study	water: pH 5.1 – 7.8 sediment: pH 7.4 – 8.0	20 ± 2 °C	Water DT ₅₀ : > 1000 d (Silt loam), 473 d (Sand) DT ₉₀ : > 1000 d (Silt loam), > 1000 d (Sand) Whole system: Metaldehyde DT ₅₀ : >1000 d (Silt loam), >1000 d (Sand) DT ₉₀ : > 1000 d (Silt loam), >1000 d (Sand)		Kane, T. (2009)
OECD 307	Aerobic degradation in soil	pH: 6.1 – 7.3 OC 1 % - 4.2 %	20 °C	DT ₅₀ Hockey-stick 6.6 d – 19.4 d		Juozenaite, A. (2009)

5.1.1 Stability

Reference: Carpenter M. (1989a): Hydrolysis of metaldehyde as a function of pH at 25 $^{\circ}$ C.; LONZA Report No. 1410; Document No. 711-001

<u>Guideline:</u> EPA, Pesticide Assessment Guidelines, Subdivision N, Section 161-1, Hydrolysis studies, October 1982

GLP: Yes

<u>Test item</u>: [U-¹⁴C] metaldehyde, radiochemical purity >99 %, batch no. 8232

Material and methods:

The hydrolytic stability of [U- 14 C] metaldehyde in an aqueous solution of a nominal concentration of 25 µg/L was studied at pH 5, 7 and 9. Two different pH 7 buffers were applied to evaluate buffer catalysis of the degradation process: TRIS buffer (tris(hydroxymethyl)aminomethane/HCL) and HEPES buffer (N-2-Hydroxyethylpiperazine-N $^{-2}$ -ethane-sulfonic acid). The pH 5 and pH 9 buffer solutions consisted of acetic acid / sodium acetate and boric acid / borax. The test vessels were incubated in the dark under sterile conditions at 25 $^{\circ}$ C up to 32 days. Duplicate samples were taken at days 0, 1, 3, 6, 9, 14, 22 and 32 and analysed by LSC, TLC and reverse phase HPLC. The DT₅₀ values were calculated with linear regression assuming first order kinetics.

Findings:

The pH values measured on each sampling day showed deviations of only 0.01 units for the pH values showing sufficient stability of the buffer solutions. The concentration of metaldehyde remained constant until the end of the study. The measured amounts of applied radioactivity in all samples were within the range of 98.6 - 100 %. No hydrolytic degradation products were observed except for one sample at pH 5 at day 22 where small amounts of paraldehyde and acetaldehyde were detected. Since this was the only sample at which degradation was observed the sample was considered to be an outlier (rejected based on the "Q" test) and was not further taken into account.

Conclusion:

Metaldehyde was hydrolytically stable at a temperature of 25 °C and pH values of 5-9. Comment (RMS):

The study was considered to be acceptable.

Photochemical Degradation

Reference: Carpenter M. (1989b): Photodegradation of metaldehyde in pH 7 buffered solution. LONZA Report No. 1412; Document No. 712-001

Guideline: EPA Pesticide Assessment Guidelines, Subdivision N, Section 161-2,

Photodegradation studies in water, October 1982

GLP: Yes

<u>Test item</u>: [14C]-metaldehyde, radiochemical purity 97.7 %, batch no. 8232

Material and methods:

 14 C-metaldehyde was applied to a pH 7 buffer solution at an initial concentration of 30 μg/L. Additionally photosensitized samples were prepared containing 1 % by volume acetone. The test vessels were incubated at 25 ± 1 °C under sterile conditions and continuous irradiation with a xenon arc lamp (wavelength 300 – 750 nm, 0.02937 – 0.8649 W/m²) up to 30 days. Duplicate samples were taken on days 0, 1, 3, 7, 14 and 30 and analysed by LSC and HPLC. Findings:

No degradation of metaldehyde was observed in the dark controls, in irradiated sensitized samples and in irradiated non-sensitized samples. The measured concentration of metaldehyde

remained stable during the test within a range of 95.9 - 98.1 % of applied radioactivity. Since no photolytical degradation was observed (regardless of the presence of acetone as a photosensitizer) no estimation of the photolytical half-life was conducted.

Conclusion:

Metaldehyde is photolytically stable at pH 7 and a temperature of 25 °C.

Comment (RMS):

The study was considered to be acceptable.

5.1.2 Biodegradation

5.1.2.1 Biodegradation estimation

5.1.2.2 Screening tests

Reference: Wüthrich V. (1990a): Ready biodegradability: Modified OECD Screening Test for

P0071. LONZA Report No. 1490; Document No. 713-002

Guideline: OECD 301 E: Ready Biodegradability: Modified OECD Screening Test (1981)

GLP: Yes

Test item: Metaldehyde, purity 99.3 %, batch no. 5448

Material and methods:

Metaldehyde was dissolved at a nominal concentration of 100 mg/L in a test medium containing 0.5 mL active sewage sludge per 1000 mL. The flasks were loosely covered with an aluminium foil. Duplicate test vessels with metaldehyde, aniline and blank controls were incubated for 28 days in the dark at $24 - 26.5^{\circ}\text{C}$. Samples were taken for DOC analysis on days 0, 7, 21 and 28. Findings:

Aniline reached a level of biodegradation of 97 % within 7 days showing sufficient biological activity of the sewage sludge. 18 % degradation of metaldehyde was observed within the 28-day incubation period. Metaldehyde was slightly degraded but it failed clearly the trigger value of 70 % removal of DOC. Therefore it has to be classified as not readily biodegradable.

Conclusion:

Metaldehyde is not readily biodegradable.

Comment (RMS):

The pH value was not measured, but since the test medium was prepared according to OECD guideline 301, the study was considered to be acceptable.

Reference: Wüthrich V. (1990b): Inherent biodegradability: "Modified Zahn-Wellens Test".

LONZA Report No. 1488; Document No. 713-001

Guideline: OECD 302 B: Inherent Biodegradability: Modified Zahn-Wellens Test (1981)

GLP: Yes

Test item: Metaldehyde, purity 99.3 %, batch no. 5448

Material and methods:

Metaldehyde was dissolved in a test medium at a nominal concentration of 100 mg/L. The test medium was prepared according to OECD guideline 302. An amount of sludge corresponding to 0.2 g dry material was added per litre test medium. Two replicates of the sample treated with metaldehyde one sample treated with aniline and two blank controls were incubated in the dark for 28 days at 19 - 23.5 °C. The flasks were aerated with a flow rate of 0.5 - 0.7 L/minute resulting in an oxygen concentration of $8.4 - 9.7 \text{ mg } O_2$ per litre. The pH values were in the range of 7.5 - 7.8. Samples were taken for DOC analysis immediately after application and at $3 - 9.7 \text{ mg} O_2$

hours, 7, 14, 21 and 28 days after treatment.

<u>Findings:</u> Aniline degraded by 98 % within 14 days showing sufficient biological activity of the sewage sludge. After 28 days only 6 % degradation of metaldehyde was observed. The level of 20 % reduction in DOC within 28 days was not reached and therefore metaldehyde has to be classified as not inherently biodegradable.

Conclusion:

Metaldehyde is not inherently biodegradable.

Comment (RMS):

The study was considered to be acceptable.

Reference: Study on ready biodegradability of Metaldehyde

Author(s), year: Lebertz, H., 2008

Report/Doc. number: Lonza Report No. 4317, Doc. No. 713-003

Guideline(s): OECD 301 F

GLP: Yes
Deviations: None
Validity: Yes

Material and methods:

A test medium was prepared according to the OECD Guideline 301 F and inoculated with active sewage sludge. Metaldehyde was dissolved in the test medium at a nominal concentration of 100 mg/L. The incubation flasks were closed and incubated for 28 days at 22°C. The oxygen uptake was continuously measured by using a manometer. Sodium benzoate was used as a positive control.

Findings:

Metaldehyde showed a degradation of 2.8 % within the 28-day incubation period and is classified as not readily biodegradable in terms of this test (Table 73). The positive controle sodium benzoate reached a level of 91 % biodegradation within 28 days showing an adequate activity of the used inoculum.

Table 73: Ready biodegradability test with Metaldehyde and the positive control Sodium benzoate (results expressed as % degradation)

Time after application [days]	application Degradation of Metaldehyde		on Degradation of Metaldehyde		Degradation of Sodium Benzoate
	Set 1	Set 2			
0	0.0	0.0	0		
7	1.0	0.3	79		
14	2.1	0.6	88		
21	2.8	1.6	90		
27	3.3	2.1	91		
28	3.3	2.3	91		

Conclusion:

Due to the fact that only 2.8 % of the Metaldehyde was degraded during the 28 day incubation period it has to be classified as not readily biodegradable in terms of this test.

Comments (RMS):

The study is considered acceptable

5.1.2.3 Simulation tests

Reference: Determination of the degradation and persistence of ¹⁴C-Metaldehyde in

the water/sediment-system

Author(s), year: Möllerfeld, J., Römbke, J., Heller, M., 1993 Report/Doc. number: Lonza Report No. 2120, Doc. No. 714-001

Guideline(s): BBA Guideline

GLP: Yes
Deviations: None
Validity: Yes

For details on material and methods, please refer to DAR.

The study described under PD June 2009, IIA, 7.1/12 showed that Acetaldehyde is trapped by soda lime but not liberated with HCl. The rate of degradation of Metaldehyde in water/sediment systems was determined using non-linear regression assuming first-order reaction kinetics. On request of the member states and EFSA, the DT₅₀ and DT₉₀ values of Metaldehyde were recalculated during dossier preparation according to the recommendations of the FOCUS Kinetic Working group.

Findings:

The formation of CO₂ was very high, accounting for a maximum of 61.57 % and 68.80 % of the applied radioactivity on day 100 for the sandy and loamy system, respectively. The study described under PD June 2009, IIA, 7.1/12 confirmed that only CO₂ is liberated by adding of HCl. Therefore, the radioactivity presented for the soda lime traps is only related to the volatile product CO₂. Other volatile compounds did not exceed 0.12 % of the applied radioactivity.

The amount of ¹⁴C-Metaldehyde in the water phases decreased in the sandy and loamy system continuously to a minimum of 0.33 and 0.38 % of applied radioactivity at the end of the study, respectively (Table B.8.10.3-2). In the sediment extracts of the sandy system, the ¹⁴C-Metaldehyde concentration increased to 10.92 % of applied radioactivity on day 7 and subsequently decreased to 0.16 % of applied radioactivity on day 100. In the sediment of the loamy system, the Metaldehyde concentration increased to a maximum value of 19.49 % of applied radioactivity on day 7 and thereafter continuously decreased to 0.28 % of applied radioactivity on day 100.

The major degradation product both in the extract of the sediment and in the water phase was Acetaldehyde. The amount of Acetaldehyde in the water phases increased continuously to a maximum of 22.32 % and 21.73 % of applied radioactivity at day 30 in the sandy and loamy system, respectively. Thereafter, the amount of Acetaldehyde in the water phase decreased to 5.39 and 1.58 % of applied radioactivity on day 100, respectively. In the sediment, Acetaldehyde occurred with a maximum of 4.70 % and 3.76 % at day 14 in the sandy and loamy system, respectively

The degradation of Metaldehyde in water/sediment systems was described by first order

degradation kinetics. The DT_{50} values in the water phase were calculated to be 11.8 and 11.2 days for the sandy and loamy system, respectively, and the DT_{90} values were determined to be 39.1 days and 37.1 days, respectively. In the total water sediment systems Metaldehyde was degraded with half-lives of 12.4 days (sandy system) and 11.9 days (loamy system). The DT_{90} values for the total systems were calculated to be 41.0 and 39.4 days, respectively (Table B.8.10.3-3).

On request of the member states and EFSA, the DT_{50} and DT_{90} values of Metaldehyde were recalculated during dossier preparation according to the recommendations of the FOCUS Kinetic Working group to separate between degradation and dissipation half-lives with the model KINGUI Version 1.1 (Peter, S. 2009, Doc. No. 782-015, PD June 2009, IIA, 7.2/10). The degradation DT_{50} values in the entire system were calculated to be 4.10 and 4.42 days for the sandy and loamy system, respectively, and the DT_{90} values were determined to be 13.61 days and 14.97 days, respectively (Table B.8.10.3-4). In the water and sediment phase Metaldehyde dissipated with half-lives of 11.35 and 10.78 days (sandy system) and 10.25 and 9.78 days (loamy system). The DT_{90} values for the water and sediment phases ranged from 31.50 to 37.71 days in both systems.

The DT_{50} and DT_{90} values of the main degradation product Acetaldehyde in the total water/sediment systems were estimated during the dossier preparation according to the recommendations of the FOCUS Kinetic Working group. The DT_{50} values were calculated to be 30.98 and 19.01 days in the sandy and loamy system, respectively. The DT_{90} values were calculated to be 102.90 and 63.14 days, respectively (Table B.8.10.3-5).

Table B.8.10.3-1: Percent distribution of applied radioactivity and mass balance in water/sediment systems treated with 14C-Metaldehyde (results expressed as % of applied radioactivity)

	Organia		Water	Sediment E	Extractables	Sediment	Total
Day	Organic Volatiles	olatiles CO ₂ phase DCM Met		Methane Extract	Un- extractables	Recov ered	
			Sand	y system			
0	<0.1	0.06	98.97	1.45	0.51	<0.01	101.0
0.25	<0.1	0.13	96.18	3.31	0.95	0.15	100.7
1	<0.1	0.27	91.75	7.72	1.92	0.65	101.7
2	<0.1	0.27	86.48	7.74	2.17	0.57	102.3
7	<0.1	0.57	85.02	10.92	3.65	0.62	100.8
14	<0.1	6.39	59.47	9.28	4.70	8.58	88.42
30	0.11	27.48	23.10	1.09	3.17	20.54	75.49
62	<0.1	50.57	13.41	0.43	1.82	17.03	83.26
100	<0.1	61.57	5.72	0.16	0.87	19.05	87.37
			Loam	y system			
0	0	-	99.46	0.95	0.09	<0.10	100.5
0.25	0	0.12	96.75	4.66	0.62	0.11	102.3
1	0.12	0.22	88.53	9.85	1.42	0.29	100.2
2	0	0.35	83.61	12.82	1.80	0.48	99.1
7	0	0.68	76.44	19.49	2.73	0.84	100.2
14	0	5.08	57.23	15.01	3.76	6.13	87.2
30	0	36.82	22.99	1.26	2.84	13.33	77.2
62	0.01	64.77	5.56	0.24	1.90	18.88	91.4
100	0.01	68.80	2.02	0.28	1.22	10.63	78.7

¹Dichloromethan

Table B.8.10.3-2: Percent distribution of applied radioactivity and mass balance in water and sediment phase of an aerobic aquatic systems treated with 14C-Metaldehyde

	Metaldo	ehyde	Acetald	lehyde
Day	Water phase	Sediment	Water phase	Sediment
	<u>.</u>	[% of applied	d radioactivity]	
		Sandy syste	m	
0	95.87	1.45	3.11	0.51
0.25	93.28	3.31	2.90	0.95
1	89.10	7.72	2.66	1.92
2	87.14	7.74	2.34	2.17
7	82.61	10.92	2.41	3.65
14	40.05	9.28	19.42	4.70
30	0.78	1.09	22.32	3.17
62	3.20	0.43	10.21	1.82
100	0.33	0.16	5.39	0.87
		Loamy syste	m	
0	96.38	0.95	3.08	<0.10
0.25	93.68	4.66	3.07	0.62
1	85.87	9.85	2.66	1.42
2	81.00	12.82	2.61	1.78
7	74.10	19.49	2.35	2.73
14	36.01	15.01	21.22	3.76
30	1.27	1.26	21.73	2.84
62	0.38	0.24	5.18	1.90
100	0.44	0.28	1.58	1.22

Table B.8.10.3-3: DT50 and DT90 values for Metaldehyde in water/sediment systems

	DT ₅₀ (days)	DT ₉₀ (days)	\mathbb{R}^2	Method of Calculation
		Sandy System		
Water Phase	11.8	39.1	0.749	First order
Total System	12.4	41.0	0.698	
		Loamy System		
Water Phase	11.2	37.1	0.643	First order
Total System	11.9	39.4	0.939	

Table B.8.10.3-4: DT50 and DT90 values for Metaldehyde in water/sediment systems

System	Phase	DT ₅₀ [d]	DT ₉₀ [d]	Model
Sandy System	System	4.10	13.61	Single first order ¹⁾
	Water	11.35	37.71	Single first order
	Sediment	10.78	35.82	Single first order
Loamy System	tem System 4.42		14.97	Single first order ¹⁾
	Water	10.25	34.07	Single first order
	Sediment	9.48	31.50	Single first order

¹⁾Without taking into account the data of the lag phase (first 14 days of incubation)

Table B.8.10.3-5: DT50 and DT90 values for Acetaldehyde in total water/sediment systems (re-calculated during dossier preparation with KINGUI Version 1.1)

System	DT ₅₀ (days)	DT ₉₀ (days)	Chi ²	Model
Sandy System	30.98	102.90	17.0	Single first order
Loamy System	19.01	63.14	16.7	Single first order

Proposed degradation pathways of Metaldehyde in aquatic systems

Conclusions:

Regarding the sequence of the used traps in the water/sediment study, it is very likely that the CO_2 trap was arranged at the last trapping vessel (i.e. any traps collecting metabolites are arranged between the incubation vessel and the CO_2 trap).

During the evaluation process, questions related to the trapping of Metaldehyde and its possible degradation products in laboratory studies have been raised in the reporting tables by the Member States and by EFSA. Therefore, several tests were conducted to identify appropriate media to trap Metaldehyde and any volatile degradation products which could be formed during the degradation of Metaldehyde in the laboratory studies (for details please see Juozenaite, A., 2009, Doc. No. 741-002, PD June 2009, IIA 7.1/12). Based on these results a trapping system was proposed and used in the new conducted water/sediment study (Kane, T. 2009).

Comment RMS: The review is considered acceptable.

Reference: ¹⁴C-Metaldehyde Aerobic Transformation in Aquatic Sediment Systems

Author(s), year: Kane, T., 2009

Report/Doc. number: Lonza Report No. 4392, Doc. No. 714-003

Guideline(s): OECD 308

GLP: Yes
Deviations: None
Validity: Yes
MATERIAL AND METHODS:

Test item: ¹⁴C-labelled Metaldehyde:

Batch 2860DCR021-3

Radiochemical purity >97% (checked before use in this study)

Specific radioactivity: 4.03 MBq/mg

Radiochemical purity >97% (checked before use in this study)

Non-radiolabelled test substance:

Lot number 37801

Purity 99.6% (expiry date 26 February 2011)

Incubation 20 ± 2 °C

temperature:

Application rate: 0.07 mg/L corresponding to 700 g ai/ha assuming a 30 cm water

column

Table B.8.10.3-6: Water/Sediment-systems used to investigate the route and rate of degradation of 14C-Metaldehyde

	Syst	em I	Syste	m II
Parameters		bbey Lake, ire, UK)	(Swiss Lake, Derbyshire, UK)	
		Sedi	ment	
Soil texture (UK classification)	Silt I	oam	Sar	nd
% sand (2000 μm-63 μm)	1	0	90)
% silt (63 μm-2 μm)	7	2	4.0)
% clay (<2 μm)	1	8	6.0)
Organic carbon (%)	4	.1	0.0	6
pH value (H ₂ O)	7	.8	5.1	1
Redox-Potential [mV]				
(beginning of study)	25		75	;
(end of study)	2	.2	26	3
N-total (%)	0.	39	0.2	3
P-total (mg/kg sediment)	946		16	1
CEC (mVal N/100 g dry soil)	15	5.5	2.2	2
Microbial characterisation (colony forming untis/g)	Day 0	Day 97	Day 0	Day 97
Aerobic bacteria	747500	312250	590000	630000
Aerobic bacterial spores	146500	23050	60000	28000
Anaerobic bacteria	151500	61250	11950	11350
Anaerobic bacterial spores	54500	8850	9675	9225
Actinomycetes	3160	14575	1755	103
Fungi	1010	2150	1030	548
		Wa	ter	
N-total (mg/L)	2	.3	1.2	2
P-total (mg/L)	<().1	<0.	1

Organic carbon (mg/L)	(6.7	4.7	7
Total hardness as CaCO₃(mg/L)	1	71	26	S
pH value				
(beginning of study)	3	3.0	7.4	1
(end of study)	7	7.2	6.7	7
Oxygen content [mg/L]				
(beginning of study)	7	1.1	70.5	
(end of study)	7	1.0	63.9	
Redox-Potential [mV]				
(beginning of study)	3	368	36	4
(end of study)	3	882	461	
Microbial characterisation (colony forming untis/g)	Day 0	Day 97	Day 0	Day 97
Aerobic bacteria	46200	11550	2270	4200
Aerobic bacterial spores	155	133	60	<18
Actinomycetes	<13	<223	<10	<10
Fungi	<13	120	<10	<13

The degradation of ¹⁴C-Metaldehyde under aerobic aquatic conditions was studied in a silt loam and in a sand water/sediment system which were collected from two different lakes in Derbyshire, UK. The sediment and water characteristics of both systems are summarised in Table B.8.10.3-6. Incubation flasks for the test system were filled with wet sediment and water in such a way to obtain a sediment layer of 2.5 to 3 cm and supernatant water layer of 7.5 to 9 cm. Afterwards, the water/sediment systems were allowed to acclimatise in the dark at 20 ± 2 °C for 19 days. The equilibrium period resulted in a complete phase separation and stabilisation of the oxygen concentration, pH and redox potential; these parameters were also measured at each sampling point. ¹⁴C-Metaldehyde was dissolved in acetonitrile and applied separately onto the water surface of each test system at a rate of 0.07 mg/L corresponding to a field application rate of approximately 700 g ai/ha assuming an even distribution in a 30 cm water column. Acetonitrile was applied to control flasks at a corresponding rate to determine the microbial activity at the end of the study. Provisions were made for the quantitative trapping of any volatiles by the installation of a polyurethane foam bung (trapping of Metaldehyde), two silica gel cartridges coated with 4-dinitrophenylhydrazine (DNPH, trapping of Acetaldehyde), a vessel containing ethyl digol (any additional organic volatile) and two vessels containing 1 M KOH trapping solutions (trapping of CO₂). Several trapping media were tested before study start to have an appropriate trapping system for this study in order to trap and to quantify volatile Metaldehyde, any possible volatile organic degradation products and CO₂ separately. For details see point B.8.1.1. Samples were taken immediately after application and after 1, 2, 7, 14, 29, 59 and 97 days of incubation. The microbial activity of the water and sediment phase was measured separately for different groups of organisms (i.e aerobic and anaerobic bacteria, aerobic and anaerobic bacterial spores, actinomycetes and fungi) at study initiation and at 97 days after treatment. The results demonstrate that the microbial viability of the study soil at the time of dosing and during the incubation period was representative of microbially active systems.

The water and sediment phase was separated and the radioactivity in the water phase was determined by LSC. Sediment samples were extracted two times with approximately 400 mL and 200 mL methanol. Day 97 samples were additionally extracted with approximately 200 mL acetonitrile. The extracts were combined and the radioactivity in the extracts was quantified by LSC and characterised by HPLC. The results of selected samples were confirmed by TLC

analysis. Extracted sediment samples were combusted to determine levels of un-extractable residues. Polyurethane bungs were extracted with acetonitrile. Each DNPH cartridge was eluted with acetonitrile and the eluates from the two cartridges associated with each sample were combined. Radioactivity in the trapping solutions, extracts and eluates was quantified by LSC. The presence of CO₂ in the KOH trapping solutions was confirmed by precipitation with barium carbonate.

The kinetic analysis of Metaldehyde in the water, sediment phase and entire system was assessed using different kinetic models according to the recommendations of the FOCUS Kinetic Working group.

Findings:

Total mean recoveries obtained during the study ranged from 96.8 % to 101.7 % and from 97.5 % to 105.8 % of the applied radioactivity for the silt loam and sand system, respectively (Table B.8.10.3-7).

Radioactivity in the water phases of both test systems decreased slowly throughout the incubation period reaching 68.2 % (silt loam system) and 73.6 % of applied radioactivity (sand system) after 97 days.

Non-extractable residues in the sediment did not exceed 4 % in the silt loam system and 1.0 % of applied radioactivity in the sand system during the entire incubation period.

Volatilised Metaldehyde, volatilised Acetaldehyde and other volatile organics did not exceed 2.0% of applied radioactivity. The formation of CO_2 was low, accounting for a maximum of 4.8% and 8.1% of the applied radioactivity on day 97 for the silt loam and sand system, respectively.

The amount of ¹⁴C-Metaldehyde in the water phases decreased in the silt loam and sand system to 72.7 % and 78.5 % of applied radioactivity at day 7 (Table B.8.10.3-8). Afterwards, the level of Metaldehyde in the water phase remained stable in both systems ranging from 64.6 % to 67.4 % of applied radioactivity in silt loam system and from 71.1 % to 78.1 % in sand system. In the sediment extracts of the silt loam system, the ¹⁴C-Metaldehyde concentration increased to 21.2 % of applied radioactivity on day 14 and remained at similar levels on day 97 (Table B.8.10.3-8). In the sediment of the sand system, the Metaldehyde concentration increased to a maximum value of 13.5 % of applied radioactivity on day 29 and thereafter the residue level did not change obviously.

In both systems, up to three unknown compounds were detected during the entire incubation period none of them exceeded 2.0 % of applied radioactivity in the water phase as well in the sediment phase. The reaming radioactivity distributed throughout the regions of the chromatogram other than those specified was below 3.0 % of applied radioactivity in both systems and in both phases. The proposed degradation pathway of Metaldehyde in aquatic sediment systems under aerobic conditions is presented under Figure B.8.10.3-1.

The degradation kinetics of Metaldehyde in water/sediment systems was evaluated by different kinetic models according to the recommendations of the FOCUS Group. The best fit model based on statistical values and visual assessment was chosen. The DT_{50} values in the water phase and total system were calculated to be > 1000 days for the silt loam system using DFOP and FOMC model (Table B.8.10.3-10). For the sand system, the DT_{50} values in water and total

system were determined to be 473 days and 714 days using HS and DFOP model, respectively. The corresponding DT_{90} values were determined to be > 1000 days in both phases of both systems. Since no decline of Metaldehyde occurred in the sediment phase of both systems from the maximum level onwards, no kinetic evaluation could be conducted for this phase.

Table B.8.10.3-7: Percent distribution of applied radioactivity and mass balance in water/sediment systems treated with 14C-Metaldehyde (results expressed as % of applied radioactivity)

Time		Volatile	s			Sediment		
after application [d]	Met- aldehyde	Acet- aldehyde	Other organics	CO ₂	Water phase	Extract- ables	Un- extract- ables	Total Recovered
			Silt loan	n syster	n			
0	-	-	-	-	100.3	1.2	0.3	101.7
1	<loq< td=""><td>0.2</td><td>0.2</td><td>0.3</td><td>92.3</td><td>5.6</td><td>1.5</td><td>100.0</td></loq<>	0.2	0.2	0.3	92.3	5.6	1.5	100.0
2	<loq< td=""><td>0.1</td><td>0.4</td><td>0.6</td><td>90.2</td><td>7.5</td><td>1.8</td><td>100.5</td></loq<>	0.1	0.4	0.6	90.2	7.5	1.8	100.5
7	0.1	0.1	0.9	1.6	75.7	18.5	2.6	99.4
14	0.2	0.1	1.3	2.3	69.6	21.7	3.9	99.0
29	0.3	0.3	2.0	3.6	69.2	20.9	2.7	98.8
59	0.4	0.6	1.5	4.6	67.1	20.1	2.8	96.8
97	0.5	0.2	2.3	4.8	68.2	21.3	3.3	100.6
			Sand :	system				
0	-	-	-	-	99.8	2.5	0.2	102.5
1	<loq< td=""><td>0.7</td><td>0.4</td><td>0.8</td><td>92.8</td><td>6.5</td><td>0.3</td><td>101.3</td></loq<>	0.7	0.4	0.8	92.8	6.5	0.3	101.3
2	<loq< td=""><td>0.2</td><td>0.1</td><td>0.5</td><td>96.3</td><td>8.7</td><td>0.2</td><td>105.8</td></loq<>	0.2	0.1	0.5	96.3	8.7	0.2	105.8
7	<loq< td=""><td>1.0</td><td>0.1</td><td>1.2</td><td>82.2</td><td>12.9</td><td>0.5</td><td>97.9</td></loq<>	1.0	0.1	1.2	82.2	12.9	0.5	97.9
14	<loq< td=""><td>0.8</td><td>0.2</td><td>2.0</td><td>80.7</td><td>13.3</td><td>0.6</td><td>97.5</td></loq<>	0.8	0.2	2.0	80.7	13.3	0.6	97.5
29	0.1	1.2	0.9	3.5	77.3	14.1	0.7	97.7
59	0.3	1.2	1.4	5.0	76.9	13.2	0.6	98.4
97	0.5	1.2	1.3	8.1	73.6	13.3	0.6	98.5

Table B.8.10.3-8: Characterisation of radioactivity in the water phase of aerobic aquatic systems treated with 14C-Metaldehyde (expressed in % of applied radioactivity)

Time after application [d]	Metaldehyde	Unknown 1	Unknown 2	Unknown 3	Others ¹⁾
Silt loam syster	n				
0	97.6	1.2	nd	nd	1.5
1	89.9	0.2	0.3	0.1	1.9
2	87.9	nd	0.1	nd	2.3
7	72.7	0.5	0.2	nd	2.4
14	67.4	0.3	0.2	nd	1.7
29	67.0	nd	nd	nd	2.2
59	64.6	0.1	nd	0.3	2.1
97	66.4	nd	nd	0.3	1.6
Sand system					
0	97.0	0.7	0.3	nd	1.9
1	90.0	1.3	nd	nd	1.6
2	92.3	1.6	0.4	nd	2.0
7	78.5	0.6	0.1	nd	3.0

Time after application [d]	Metaldehyde	Unknown 1	Unknown 2	Unknown 3	Others ¹⁾
14	78.1	0.3	0.1	nd	2.3
29	74.8	nd	0.1	nd	2.4
59	75.0	nd	nd	nd	1.9
97	71.1	nd	nd	nd	2.5

¹⁾ radioactivity distributed throughout regions of the chromatogram other than those specified and which did not contain any discrete radioactive components;

nd not detected

Table B.8.10.3-9: Characterisation of radioactivity in the sediment phase of aerobic aquatic systems treated with 14C-Metaldehyde (expressed in % of applied radioactivity)

Time after application [d]	Metaldehyde	Unknown 1	Unknown 2	Unknown 3	Others ¹⁾			
Silt loam system								
0	1.1	nd	nd	nd	0.1			
1	5.1	nd	nd	0.1	0.4			
2	7.0	nd	nd	< LOQ	0.2			
7	17.4	0.1	nd	0.5	0.5			
14	21.2	0.1	nd	nd	0.5			
29	20.6	nd	nd	nd	0.3			
59	19.5	nd	nd	0.1	0.9			
97	20.5	nd	nd	0.1	0.8			
Sand system								
0	2.5	nd	nd	nd	0.1			
1	6.1	0.1	nd	nd	0.3			
2	8.3	0.1	nd	nd	0.4			
7	12.3	0.2	nd	nd	0.5			
14	12.6	0.1	0.1	0.1	0.6			
29	13.5	nd	0.1	0.1	0.4			
59	12.8	nd	nd	0.1	0.3			
97	12.5	nd	nd	0.1	0.7			

¹⁾ radioactivity distributed throughout regions of the chromatogram other than those specified and which did not contain any discrete radioactive components;

nd not detected

Table B.8.10.3-10: DT50 and DT90 values for Metaldehyde in water/sediment systems

Compartment	Kinetic model	DT ₅₀ (days)	DT ₉₀ (days)	Chi ²				
Silt loam system								
Water Phase	SFO	184	610	5.5				
	FOMC	>1000	>1000	1.8				
	DFOP*	>1000	>1000	1.3				
	HS	>1000	>1000	0.8				
Total System	SFO	579	>1000	2.8				
	FOMC*	>1000	>1000	1.1				
	DFOP	>1000	>1000	1.1				
	HS	nc	nc	nc				
Sand system								
Water Phase	SFO	287	953	5.5				
	FOMC	>1000	>1000	1.8				
	DFOP	494	>1000	1.3				
	HS*	473	>1000	0.8				

Compartment	Kinetic model	DT ₅₀ (days)	DT ₉₀ (days)	Chi²				
Silt loam system								
Total System	SFO	486	>1000	2.1				
	FOMC	>1000	>1000	0.8				
	DFOP*	714	>1000	0.6				
	HS	703	>1000	0.6				

nc not calculated

Conclusion:

Metaldehyde dissipated from the water phase with a DT_{50} value of >1000 days in silt loam system and of 473 days in sand systems. DT_{50} values for the dissipation from the total system were >1000 days in both systems. Metaldehyde was degraded within both water and sediment phases and up to four degradation products (including $^{14}CO_2$) were formed. Significant quantities of intermediate products were not produced during Metaldehyde degradation in both systems.

Comments (RMS):

The study is considered acceptable.

^{*} indicates the best fit with respect to Chi²value and visual assessment of the fit to the measured data

5.1.2.4 Degradation in soil under aerobic conditions

The *aerobic metabolism* of metaldehyde is suggested to occur via microbial and/or chemical degradation building mainly CO₂ (80.4% AR after 21 days). No volatilised Metaldehyde or other organic volatiles are observed at relevant concentrations. The aerobic metabolism studies showed that Metaldehyde degraded rapidly after a lag-phase between 6 and 19 days following hockey-stick kinetic. No metabolites are formed over 5 % AR. The non-extractable residues increased during the incubation period from a mean of 0.5% AR to a mean of about 36% AR.

The proposed metabolic pathway in soil is shown in the figure below:

DEGRADATION IN SOIL UNDER STANDARD CONDITIONS

The degradation rate of Metaldehyde under *aerobic* conditions was investigated in four soils, and the degradation occurred rapidly under aerobic conditions after a lag phase of 5.8 to 19 days, which was essentially no degradation of Metaldehyde during this period. The compound degraded almost completely over a very short period of time with extensive mineralisation to carbon dioxide and incorporation into bound residues. CO_2 accounted for up to 80 % of applied radioactivity and the maximum level of bound residues was 26 %. A decline of the non-extractable residues was observed during the entire incubation period. No intermediate degradation products were produced in relevant amounts (i.e. < 4 % of applied radioactivity). Using a modified hockey-stick kinetic model, the DT_{50} and DT_{90} values ranged from 6.6 to 19.5 days and from 8.5 to 22.1 days, respectively. The half-life of Metaldehyde during the rapid decline phase ranged from 0.5 to 4.1 days. The half-lives proposed for further evaluation were determined by back-calculation of the DT_{90} values (i.e. $DT_{90}/3.32$) resulting in pseudo- DT_{50} values ranging from 2.6 to 6.7 days.

The degradation rates are summarised in the tables below:

Table 74: Laboratory studies

Metaldehyde	Aerobic	Aerobic conditions					
Soil type (study Ref.)	ОС	рН	Temperatu re/ Moisture content	DT ₅₀ /DT ₉₀ Hockey-stick (d)	Duration of lag-phase (d)	DT ₅₀ Pseudo single-first- order	Chi ² (%)
Silt Loam, 02-A Juozenaite, A. (2009)	1.2	6.5	20 ℃/pF 2	19.5/20.6	19.0	6.2	2.0
Sandy clay loam, Elmton (294) Juozenaite, A. (2009)	4.2	7.0	20 °C/pF 2	11.6/21.0	7.5	6.3	9.6
Sandy clay loam, Fladbury Juozenaite, A. (2009)	3.1	6.1	20 °C/pF 2	15.9/22.1	13.2	6.7	9.3
Sandy loam, Lanoe Juozenaite, A. (2009)	1.0	7.3	20 °C/pF 2	6.6/8.5	5.8	2.6	7.8
Geometric mean						5.1	

PHOTOLYTIC DEGRADATION ON SOIL SURFACE

It can be concluded that metaldehyde was found to be photolytically stable on soil surface under both tested conditions. Besides the parent compound no further radioactive fractions were observed in the extracts. Since no significant degradation of metaldehyde was detected during the study in either irradiated or non-irradiated systems it was not possible to estimate accurate DT_{50} and DT_{90} values.

5.1.3 Summary and discussion of degradation

Summary: Biotic degradation	Test guideline / design	GLP (y/n)	Reliability
Ready biodegradability			
Due to the fact that only 2.8 % of the Metaldehyde was degraded during the 28 day incubation period it has to be classified as not readily biodegradable in terms of this test.			

Summary: Biotic degradation	Test guideline / design	GLP (y/n)	Reliability
Water/sediment system (simulation test) In laboratory incubations in dark aerobic natural sediment water systems (four systems investigated), metaldehyde exhibited low to very high persistence. In the 2 systems where metaldehyde exhibited low persistence (where conditions were more oxidising, as indicated by the negative sediment redox potentials measured for the pertinent systems), microbial degradation of Metaldehyde was rapid with DT50 values of 4.1 - 4.4 days for the whole systems and the major metabolite acetaldehyde was formed (max. ca. 22 % AR in water and 5% in sediment). In these systems acetaldehyde exhibited moderate persistence. In the two less oxidising systems metaldehyde exhibited very high persistence with DT50 (SFO) values in the range of 579 - 486 days and no major metabolites were formed. The unextractable sediment fraction (not extracted using methanol or dichloromethane) was a sink for the carbon radiolabels (all carbons uniformly labelled), accounting for 0.6 – 19 % AR at study end (97-100 days). Mineralisation of these radiolabels accounted for 5-8 % AR in the lower oxidation state systems and 62-69% AR under the more oxidising systems, at the end of the studies.		у	у
Degradation in soil: The degradation rate of Metaldehyde under aerobic conditions was investigated in four soils, and the degradation occurred rapidly under aerobic conditions after a lag phase of 5.8 to 19 days, which was essentially no degradation of Metaldehyde during this period. The compound degraded almost completely over a very short period of time with extensive mineralisation to carbon dioxide and incorporation into bound residues. Mineralisation of the carbon radiolabels (all carbons uniformly labelled) to carbon dioxide accounted for 50 - 78 % AR after 22-60 days (termination times of the incubations). The formation of unextractable residues (not extracted using methanol) for these radiolabels accounted for 13 – 20 % AR after 60 days.		у	n

Summary: Abiotic degradation	Test guideline / design	GLP (y/n)	Reliability
Hydrolysis: Metaldehyde was hydrolytically stable at a temperature of 25 °C and pH values of 5 – 9.		у	n
Photolysis Metaldehyde was photolytically stable at a temperature of 25 °C and a pH value of 7.		У	n
Soil Photolysis It can be concluded that metaldehyde was found to be photolytically stable on soil surface under both tested conditions. Besides the parent compound no further radioactive fractions were observed in the extracts. Since no significant degradation of metaldehyde was detected during the study in either irradiated or non-irradiated systems it was not possible to estimate accurate DT50 and DT90 values.		у	n

Conclusion: The criteria for rapid degradation are not fulfilled because

Metaldehyde is hydrolytically and photolytically stable at a temperature of 25 $^{\circ}$ C and environmentally relevant pH values.

Metaldehyde is not readily biodegradable under test conditions within 28 days.

In UK simulation study (two less oxidising systems) DT50 whole system is >> 16 d, therefore Metaldehyde is considered not to be ready biodegradable/rapid degradable.

5.2 Environmental distribution

5.2.1 Adsorption/Desorption

The adsorption behaviour of metaldehyde was studied in 8 soils using the batch equilibrium method. The K_F values were in the range of 0.432 to 0.977 L/kg. The K_{FOC} values were calculated to be in the range of 38 to 149 L/kg with a median value of 60.4 L/kg, indicating very high to high mobility. The Freundlich coefficient 1/n was in the range of 0.675 to 1.023 with a median value of 0.96. A dependency between organic content and K_{FOC} values is given. The K_{FOC} values increased with decreasing organic content values.

The results are summarised in the table below:

Table 75: Soil adsorption/desorption

Metaldehyde					
Soil Type (Ref. Study)	OC %	Soil pH	K _F	K _{FOC}	1/n
Sand (Heim & Daly,1999)	0.29	7.4	0.432	173	0.9099
Sandy loam (Heim & Daly,1999)	0.46	6.5	0.644	161	0.9651
Silt loam (Heim & Daly,1999)	1.39	7.1	0.685	57	0.9918
Clay loam (Heim & Daly,1999)	1.51	7.5	0.962	75	0.9953
Humic sand (de Vette & Aalderink, 2002)	1.9	5.3	0.735	38	0.974
Sandy loam (de Vette & Aalderink, 2002)	1.56	7.7	0.633	40	1.023
Loam (de Vette & Aalderink, 2002)	1.45	7.5	0.807	56	0.961
Low humic content sand (de Vette & Aalderink, 2002)	0.76	7.8	0.675	78	0.675
Median	•	•	•	60.4	0.96
pH dependence, Yes or No No				•	

5.2.2 Volatilisation

With a Henry's constant of 3.5 Pa $\rm m^3/mol~(20^{\circ}C)$ and a vapour pressure of 4.4 Pa $\rm (20^{\circ}C)$ metaldehyde is expected to volatilise.

In addition, the volatility of metaldehyde from a liquid formulation applied to a soil surface was experimentally investigated. The experiment showed that metaldehyde was only slowly released from the formulated product. However once released, most of the metaldehyde was found in the trapping solutions for volatile products whereas only small amounts were found as extractable or un-extractable residues in soil. Therefore, it can be concluded that the potential for volatilization of metaldehyde in formulated form seems to be significantly reduced. The photochemical oxidative half-life was estimated by a model calculation according to Atkinson to be 1.7 hours (12 hours day - 1.5 x 10^6 OH-radicals/cm³) indicating a quick degradation of metaldehyde in the troposphere.

Summary: Evironmental D	istribution (not relevant for classification and labelling)
Adsorption/Desorption	$K_{F,OC}$ values were calculated to be in the range of 5.7 to 83.3 L/kg with an arithmetic mean of 36.4 L/kg, indicating high mobility.
Volatilisation	Henry's constant of 1.6 x 10 ⁻⁶ Pa m ³ /mol (20° C) Vapour pressure of 1.3 x 10 ⁻⁵ Pa (20° C)

5.2.3 Distribution modelling

No data/information available

5.3 Aquatic Bioaccumulation

Table 76: Summary of relevant information on aquatic bioaccumulation

Method	Results	Remarks	Reference
Partition coefficient n-octanol/water OECD 107 (Shake flask method)	Purified product purity: 99.3 % (w/w) at 19.9-20.1 °C log $P_{ow} = 0.12$ $P_{ow} = 1.33 \pm 0.04$ at pH 6.7 Effect of pH (4 to 10) is not required, because Metaldehyde is neither an acid nor a base. Because metaldehyde is not an ionisable compound the water phase is not buffered	Acceptable The method is comparable to the EEC/A8 shake flask method	Cardinaals, J.M. (1988b) (Doc. No. 114-002)
Bioconcentration in fish OECD 305 E (1981), EPA Pesticide Assessment Guidelines, Subdivision N Series 165-4 (1989)	BCF (steady-state, whole fish, based on total radioactive residues): 11 Lipid content: Not measured	*	Sved, D. W., Holmes, C. M., Smith, G. J. (1992), Document No. 872-001

^{*} Due to the low and variable residues in fish tissues, negligible depuration, and the lack of detailed information about the kinetic models used for the estimation of uptake rate and depuration rate constants in the study report (k1 and k2) the estimates for these parameters and the kinetic BCF estimates are not considered to be reliable. However, the study is of sufficient quality to demonstrate that metaldehyde does not bioconcentrate in fish and hence the study is acceptable.

5.3.1 Aquatic bioaccumulation

5.3.1.1 Bioaccumulation estimation

5.3.1.2 Measured bioaccumulation data

<u>Reference:</u> Sved, D. W., Holmes, C. M., Smith, G. J. (1992): A bioconcentration study with metaldehyde in the bluegill (*Lepomis macrochirus*). Document No. 872-001

<u>Test guideline</u>: OECD 305 E (1981), EPA Pesticide Assessment Guidelines, Subdivision N Series 165-4 (1989)

GLP: Yes

Material and methods:

Test substance:

Metaldehyde, purity: 99 %, batch: 5448; [U- 14 C]metaldehyde, radiochemical purity: \geq 98.4 % *Test organism*:

Bluegill sunfish (*Lepomis macrochirus*), mean length of fish collected during the test: 55 mm (s.d. 10.4), mean weight of fish collected during the test: 2.4 g (s.d. 1.57). Detailed information on lengths and weights of fish at the start of exposure and the length and weight development during the test are not given in the study protocol. No information on age of fish at test initiation is provided.

Treatments: 0.1 mg/L, methanol was used as a solvent (0.1 mL/L), solvent control: 0.1 mL methanol/L

Number of animals: 2 replicates (A and B) per treatment and control, 70 fish per replicate

Duration:

Replicates A: 28 days uptake phase, 21 days depuration phase, replicates B: 28 days uptake phase plus 1 additional day of exposure at an increased rate of specific radioactivity to increase the concentration of radioactivity in fish tissues for possible metabolite identification *Test medium:*

Dilution water: Fresh water obtained from a well 45 m deep, pumped through a sand filter, aerated, filtered to remove microorganisms and particles, analytical results of the used well water indicate acceptable quality for the purpose of this study, hardness: 132 - 148 mg/L as CaCO₃, dissolved oxygen: 5.4 - 8.3 mg/L (> 60 % saturation throughout the study), pH: 7.7 - 8.2

Test conditions and test design:

Flow-through system, 6.4 volume additions every 24 hours, test chambers: Teflon $^{\$}$ -lined, loading: 0.61 g fish/L test medium that passed through test chambers in 24 hours, at any given time the biomass of fish/L of test water did not exceed 3.9 g fish/L, feeding: flaked fish food, excess food was siphoned from the test chambers daily after feeding, photoperiod: 16 hours light, 8 hours dark, temperature: 20.4 - 22.6 $^{\circ}$ C.

Water and fish data from replicates A were used to determine the BCF, the rate and degree of uptake, and the rate of depuration. Water and fish samples from replicates B were used to characterise the radioactive residues with regard to their polarity, extractability, and possible identification.

Biological observations:

Daily fish were observed for mortality and sublethal signs of toxicity.

Analytical measurements:

Concentration verifications: On days 0, 3, 7, 14, 21 and 28 of the uptake phase and on days 1, 3, 7, 10, 14 and 21 of the depuration phase water samples were collected from the solvent control and treated test chambers and analysed for total radioactivity by LSC. The concentrations of radioactivity in the water samples were converted to mg metaldehyde equivalents/L. Radioactivity in the samples was corrected for background radioactivity and counting efficiency by an external standard method. LOQ: 0.0078 mg/L.

Determination of total radioactivity in fish tissues: On days 0, 3, 7, 14, 21 and 28 of the uptake phase and on days 1, 3, 7, 10, 14, and 21 of the depuration phase four fish were sampled from replicates A of the treated and solvent control groups. Non edible (fins, heads, and viscera) and edible tissue fractions from the four fish were pooled. Analysis of total radioactivity was performed by homogenisation of tissue samples followed by combustion and LSC analysis of the trapped CO₂. The concentrations of radioactivity in the tissue samples were converted to mg metaldehyde equivalents/kg of tissue. Radioactivity in the samples was corrected for background radioactivity, counting efficiency, and combustion efficiency. LOQ: 0.29 mg/kg. Determination of polar, non polar and non extractable residues in fish tissues: On day 28 of the uptake phase 10 fish were collected from replicates B of the treated and solvent control groups. The edible parts were pooled for each group. Proportions of polar, non polar, and non extractable residues were determined by sequential extraction with methanol and hexane and combustion (trapping of CO₂) followed by LSC.

Collection of water and fish tissues for possible metabolite identification: Water samples were collected from chambers A and B of the treated and control groups on days 14 and 28 of the uptake phase and from chamber B of the treated group on day 29 after delivering [\frac{14}{C}]-metaldehyde at an increased specific activity for approximately 21 hours. Fish samples (20 individuals) were taken from chambers B of the treated and control groups on days 14 and 28 of the uptake phase. The 18 fish remaining in chamber B of the treated group were sampled on day 29 after exposure to [\frac{14}{C}]-metaldehyde at an increased specific activity. The edible parts of fish were pooled for each fish sample and frozen for possible metabolite identification. However,

this identification was not conducted, because there was evidence that metaldehyde is metabolised to acetaldehyde which is used in anabolic pathways (see findings described below). *Statistical evaluation:*

The BIOFAC computer program for characterising the rates of uptake and clearance of chemicals in aquatic organisms (The Dow Chemical Company, Midland, MI) was used to estimate uptake rate constants (k2), depuration rate constants (k1) and kinetic BCF values. Findings:

Verification of test concentrations:

Mean concentrations of metaldehyde in replicates A and B of the treated group were 0.10 mg/L (s.d. 0.022) and 0.10 mg/L (s.d. 0.017), respectively. Total radioactivity in test chambers ranged from 0.078 - 0.14 mg metaldehyde equivalents/L. The concentration of metaldehyde in water of the control replicates and in all replicates during the depuration phase was less than the LOQ.

Table 77: 14C-residues in water samples and tissue samples of Lepomis macrochirus during the uptake and depuration phase of a bioconcentraiton test with metaldehyde

	Sample	Residues [metaldehyde equivalents/L or kg]						
	(chamber A)	day 1	day 3	day 7	day 14	day 21	day 28	
Uptake	Water	0.088	0.10	0.078	0.14	0.09	0.11	
phase	Fish edible	< LOQ	< LOQ	< LOQ	0.75	0.30	0.87	
	Fish non edible	< LOQ	< LOQ	0.62	2.0	1.0	2.0	
	Whole fish	< LOQ	< LOQ	0.38	1.4	0.66	1.4	
Depuration	Water	< LOQ	< LOQ	< LOQ	< LOQ	< LOQ	< LOQ	
phase	Fish edible	0.87	1.1	0.48	0.64	0.75	1.2	
	Fish non edible	1.4	2.8	1.2	0.97	0.96	1.7	
	Whole fish	1.1	1.9	0.83	0.79	0.85	1.4	

The amount of radioactivity in edible and non edible tissues did not decrease during the 28 days of depuration. In the study report this finding is explained as follows: In mammals metaldehyde is quantitatively metabolised to acetaldehyde which is readily converted to acetyl-CoA. Acetyl-CoA is utilised in various anabolic pathways. These anabolic reactions should account for the slow depuration and persistence of ¹⁴C-residues in tissues if the metabolic pathways for metaldehyde in fish are similar to those in mammals.

Fractionation of metaldehyde residues in edible tissues after 28 days of uptake into polar, non polar and non extractable residues yielded 28 % polar, 12 % non polar and 59 % non extractable residues expressed as % of total residues. The high portion of non extractable residues supports the view that metaldehyde is metabolised to acetyl-CoA which is transferred via anabolic pathways to body carbon pools.

Table 78: Fish tissue bioconcentration factors for [U-14C]-metaldehyde in Lepomis macrochirus

Days of uptake	Resi		vater and fish equiv./L or kg	Bioc	oncentration	factors	
uptake	Water *	Edible	Non edible	Whole fish	Edible	Non edible	Whole fish
0	0.088	< LOQ	< LOQ	< LOQ	-	-	-
3	0.094	< LOQ	< LOQ	< LOQ	-	-	-
7	0.089	< LOQ	0.62	0.39		-	4.4
14	0.10	0.75	2.0	1.4	7.5	20	14

Days of uptake	Resi		vater and fish equiv./L or kg	Bioc	oncentration	factors	
иртакс	Water *	Edible	Non edible	Whole fish	Edible	Non edible	Whole fish
21	0.099	0.30	1.0	0.61	3	10	6.2
28	0.10 0.87 2.0 1.4				8.5	20	14
mean BCF	mean BCF at steady state (days 14 - 28)					17	11

^{*} Average concentration over time.

Table 79: Estimated uptake and depuration parameters of 14C-residues in Lepomis macrochirus estimated with the BIOFAC computer program

Parameters	Edible tissues ^a	Non edible tissues	Whole fish
Uptake rate constant k1 [mg/g/d]	-	0.94	0,62
Depuration rate constant k2 [1/d]	-	0.024	0.017
Time to reach 90 % steady state [d]	-	96	135
Time to reach 50 % clearance [d]	-	29	41
Estimated BCF		39	36

^a The BIOFAC model could not produce reasonable parameter estimates due to the low and variable tissue residues and the apparent lack of depuration

Lipid content: Not measured Mortalities and clinical signs:

Four fish in the control group and two fish in the treated group died during the uptake phase. No fish died during the depuration phase. Since post-mortem examinations revealed no signs of toxicity, these deaths are not considered to be treatment related.

Conclusions:

The uptake of metaldehyde in fish was slow and low. Depuration of total radioactivity was negligible during 28 days. The low depuration can be explained by the metabolisation of metaldehyde to acetaldehyde which enters anabolic pathways via acetyl-CoA and hence ¹⁴C-residues are incorporated into body carbon pools. This interpretation is supported by the high portion of non extractable residues found in edible fish tissues after 28 days of exposure. The data of total radioactive residues in fish tissues suggest that steady-state is reached after 14 days, however, residues thereafter were variable.

BCF (steady-state, whole fish, based on total radioactive residues) : 11 Comments (RMS):

Due to the low and variable residues in fish tissues, negligible depuration, and the lack of detailed information about the kinetic models used for the estimation of uptake rate and depuration rate constants in the study report (k1 and k2) the estimates for these parameters and the kinetic BCF estimates are not considered to be reliable. However, the study is of sufficient quality to demonstrate that metaldehyde does not bioconcentrate in fish and hence the study is acceptable.

5.3.2 Summary and discussion of aquatic bioaccumulation

The determined BCF (steady-state, whole fish, based on total radioactive residues) 11 and the log $P_{\rm ow}$ (0.12at pH 6.7) demonstrate that metaldehyde does not bioconcentrate in fish. This was

Summary Bioconcentration					
	Active substance Metaldehyde				
$log P_{O/W}$	0.12 (19.9 - 20.1 °C, pH: 6.7)				
Bioconcentration factor (BCF)	11 (whole fish at steady state)				
Conclusion	Metaldehyde is considered to have a low				
	bioaccumulation potential				

5.4 Aquatic toxicity

Table 80: Summary of relevant information on aquatic toxicity

	Results					Refere			
Method	Test organism	Test condition	Duration	Endpoint	Test conc	NOEC [mg ai/L]	EC ₅₀ /LC ₅₀ [mg ai/L]	Remarks	acceptal
EPA Pesticide Assessment Guidelines, Subdivision E, 72-1 (1985), OECD 203	Oncorhynchu s mykiss	semi-static	96 h	mortality	n	56	75		Bogers (199 yes
OECD 203	Cyprinus carpio	semi-static	96 h	mortality	n	100	> 100		Wetton et a
OECD 202 (1984), US EPA 540/9-85-005	Daphnia magna	static	48 h	immobility	mm	90	> 90		Wuethrich yes
No guideline available, OECD 202 (2004) was taken into account	Planorbarius corneus	static	48 h	immobility	n	200	> 200		Egeler et al
OECD 201	Scenedesmus subspicatus	static	96 h	biomass growth rate	n			no reliable toxicity values could be derived	Wuethrich / no
OECD 201 (2006)	Desmodesmu s subspicatus	static	72 h	yield growth rate	n	25 25	> 200 > 200		Egeler et al / yes
OECD 204	Oncorhynchu s mykiss	semi-static	21 d	mortality body weight	n	37.5	-		Bogers (199 yes
OECD 202, Part II	Daphnia magna	semi-static	21 d	immobility reproduc.	n	90	-		Wuethrich yes
No guideline	Pomacea canaliculata	1		by of metaldehyde against as in the Philippines.	P.	Estimated by RMS NOEC = < 0.5 (14 d)	Recalculated by RMS >3.33 (48h)	Recalculation: see comment page 167	Scholtz (20

n=nominal; mm= mean measured;

5.4.1 Fish

Short-term toxicity to fish

<u>Reference:</u> Bogers, M. (1990a): 96-hour acute toxicity study in the rainbow trout with P0071 in a semi-static system. Document no. 821-001

<u>Test guideline:</u> EPA Pesticide Assessment Guidelines, Subdivision E, 72-1 (1985), OECD 203 (1984)

GLP: Yes

Material and methods:

Range finding test

In a preliminary 96-hour fish toxicity study (semi-static system) rainbow trouts were exposed to a range of 0.1 to 1000 mg metaldehyde/L forming a geometric progression with a factor of 10 (1000 mg/L was a supersaturated solution with undissolved substance particles)

Main test:

Test substance: P0071 (metaldehyde), purity: 99.3 %, batch: 5448

Test organism:

Rainbow trout (*Oncorhynchus mykiss*), mean length: 5.3 cm (s.d. 0.64), mean weight: 1.64 g (s.d. 0.389)

Treatments: 0, 32, 56, 100 and 180 mg/L

Number of animals: 2 replicates with 5 fish per concentration and control

Duration: 96 hours Test conditions:

Semi-static conditions, renewal of the test media after 48 hours, test vessel were of glass and had a volume of 12 L, no feeding from 24 hours prior to the test start and during the total test period, 16 hours photoperiod daily, temperature between 11 and 13.5 °C (on one occasion 9.5 – 9.7 °C), loading: 0.68 g fish/L

Test medium:

Analytical results of the used tap-water indicate in general adequate quality for the purpose of this study, however the pH was relatively high, hardness: 2.2 mmol/L ($220 \text{ mg CaCO}_3/L$), pH between 8.4 and 8.8, dissolved oxygen concentration between 10.5 and 12.9 mg/L

Biological observations:

Mortality and sublethal effects were recorded at 2, 24, 48, 72 and 96 hours

Analytical measurements: Stability test: Additionally to the test vessels with fish a vessel without fish and 100 mg metaldehyde/L was set up to confirm the stability of the test substance. From this vessel duplicate samples were taken at 0 and 48 hours.

Concentration verifications: From each replicate of all treatment groups samples were taken at test start and from replicates of the 32 and 0.56 mg/L treatment groups samples of new test media were taken at 48 hours.

Method of analysis: Derivatisation with 2,4-dinitrophenylhydrazin followed by HPLC *Statistical evaluation*:

Since no test concentration resulted in partial mortalities, no statistical evaluation of the data was performed. The LC_{50} was estimated as the geometric mean between the lowest concentration with 100 % mortality and the highest concentration with 0 % mortality (this is in accordance with OECD 203).

Findings:

Range finding study:

1000~mg/L (supersaturated solution): 100~% mortality, 100~mg/L: one out of the four fish exposed was found dead

Main study:

Analytical results:

Stability test: Measured concentrations were 92 % (0 hours) and 87 % (48 hours) of nominal.

Concentration verification: Measured concentrations were between 82 and 100 % of nominal. *Mortality:*

Up to and including 56 mg/L no fish died, at 100 mg/L and 180 mg/L all fish died within 24 hours of exposure.

Sublethal effects:

At 56 mg/L one fish was found discoloured 48 hours after exposure. Since this was the only incidence that a sublethal effect was observed it is not considered to be treatment related. Conclusion:

LC₅₀: 75 mg/L (nominal), NOEC: 56 mg/L (nominal)

Comment (RMS):

Deviations from test guidelines: On one occasion temperatures of both replicates from the 100 mg/L and of one replicate from the 180 mg/L treatment group had decreased to values ranging from $9.5 \text{ to } 9.7 \,^{\circ}\text{C}$, respectively. This deviation from the guideline is not considered to have affected the test results.

Due to the relatively high pH of the used tap water the pH of test media (8.4 - 8.8) exceeded slightly the limit of the optimal range given in the guidelines. However, no effects were observed in the control and measured test concentrations were acceptable. Therefore this deviation from guidelines is also considered to have not affected the results of the study. The study is considered acceptable.

<u>Reference:</u> Wetton, P. M., Mullee, D. M. (2001): Meta[®]-Metaldehyde (Code LZ1060): Acute Toxicity to common carp (*Cyprinus carpio*). Document no. 821-002

Test guideline: OECD 203 (1992)

GLP: Yes

Material and methods:

Range finding study

3 fish per treatment were exposed to nominal concentrations of 0, 1, 10 and 100 mg/L Main study:

Test substance: META^R metaldehyde (tech. metaldehyde), purity: not stated, batch: 29103 Test organism:

Common carp (*Cyprinus carpio*), mean length: 4.2 cm (s.d. 0.1), mean weight: 1.82 g (s.d. 0.21) at the end of the study, loading rate: 0.91 g/L, food: commercial crushed carp pellets. Feeding was discontinued approximately 24 hours prior to the start of exposure.

Treatments: Based on the results of the range-finding study a limit test at 100 mg/L was conducted.

Number of animals: 10 fish for the control and 2 replicates with 10 fish each per treatment group

Duration: 96 hours *Test conditions:*

20 L glass vessels, semi-static test regime, daily renewal of test media, photoperiod: 16 hours light and 8 hours dark with 20 minutes transition periods, temperature was held constant at $21\,^{\circ}\text{C}$.

Test medium:

Laboratory tap water of sufficient quality, dechlorinated by passage through an activated carbon filter and partly softened, total hardness of approximately $100 \text{ mg CaCO}_3/L$, pH: 7.6 - 7.9, dissolved oxygen: 7.4 - 8.7 mg/L

Biological observations:

Mortalities and sublethal effects were recorded at 3, 6, 24, 48, 72 and 96 hours.

Analytical measurements:

Method of analysis: Gas chromatography using an external standard

Test substance verification: Samples from the control and each replicate test vessel were taken at 0, 24 and 96 hours.

Stability test: Pre-study test samples were prepared and analysed initially and after storage for approximately 24 hours (period between media renewal) in sealed vessels at ambient temperature in light and dark conditions.

Findings:

Range finding study

No mortalities or sublethal effects could be observed up to the highest tested concentration of 100 mg/L.

Main study

Analytical results:

Analysis of the test preparations at 0, 24 and 96 hours gave measured values ranging from 84 % to 108 % of nominal concentrations. The test material was stable in the test media for the period between renewals (24 hours).

Mortalities and sublethal effects:

After 48 hours one fish was observed to be moribund and killed in extremis. Since no sublethal effects were observed in the remaining fish after 96 hours, this mortality was considered not to be treatment related. No further mortalities occurred.

Conclusion:

LC₅₀: > 100 mg/L, NOEC: 100 mg/L, based on a nominal limit concentration

Comment (RMS):

The study is considered acceptable.

Long-term toxicity to fish

<u>Reference:</u> Bogers, M. (1990b): Prolonged toxicity study in the rainbow trout with P0071. Document No. 826-001

Test guideline: OECD 204 (1984)

GLP: Yes

Material and methods:

Test substance: P0071 (metaldehyde), purity: 99.3 %, batch: 5448

Test organism:

Rainbow trout (Oncorhynchus mykiss), mean length: 4.8 cm (s.d. 0.12, n=10), mean weight:

1.18 g (s.d. 0.022, n=10)

Treatments: Control, 4.5, 9.5, 19, 37.5 and 75 mg/L

Number of animals: One replicate with ten fish per concentration and control

Duration: 21 days Test medium:

Analytical results of the used tap water indicate acceptable quality for the purpose of this study. Oxygen concentrations were generally in the range of 6.5 and 10.9 mg/L which corresponds to saturation levels above 60 %. However, on days 8 and 16 saturation levels had dropped to about 35 % and 50 % in old test media of the 75 mg/L and 19 mg/L treatments (% saturation levels were estimated by the RMS from measured temperatures and measured oxygen concentration levels). pH: 7.6 to 8.7, values above 8.5 were only found at the test concentration of 75 mg/L; hardness: 220 mg CaCO₃/L

Test conditions:

Semi-static test system with renewal of test media every 48 hours, test vessels: 15 L all-glass, photoperiod: 16 hours light and 8 hours dark, temperature: range of 12.0 to 15.0 °C except on day 1 and day 9 where temperature had dropped to 9.5 °C (75 mg/L) and 10.5 °C (at all test concentrations), loading: 0.91 g fish/L at the start of the test, feeding: daily 0.6 g (first week), 0.9 g (second week) and 1.2 g (third week) Trouvit per vessel

Analytical measurements:

Method of analysis: Derivatisation with 2,4-dinitrophenylhydrazin followed by HPLC Sampling: At the start of the test, after 48 hours (old media) and just before the last renewal

duplicate samples from the 4.5, 19, and 75 mg/L test vessels were taken. Extra samples were taken from all test concentrations and stored at -20 °C for additional analyses if necessary. *Biological observations:*

Fish were daily observed for mortality. Effects on behaviour and appearance were recorded every 48 hours at test media renewals.

Statistical evaluation:

Fish weights and lengths recorded for treated groups at the end of the test were compared to those of the control with the Steel-test.

Findings:

Analytical measurements:

Only samples taken from concentrations that were considered relevant in terms of effects (19, 37.5 and 75 mg/L) were analysed because the analytical method used was of relatively low sensitivity. Measured concentrations were in the range of 86 to 105 % of nominal concentrations.

Mortalities:

Only at 75 mg/L mortalities were observed, one fish was found dead on day 1 and one on day 10 of exposure (total mortality: 20 %).

Sublethal effects: Up to and including the test concentration of 37.5 mg/L no effects on fish weights and body lengths could be observed. At 75 mg/L the mean weight of fish was statistically significantly reduced compared to the control.

Table 81: Mean weights and lengths of rainbow trouts after 21 days of exposure to metaldehyde

Test concentration [mg/L]	Mean weight (s.d.) [g]	Mean length (s.d.) [cm]
0 (control)	2.79 (0.59)	6.0 (0.49)
4.5	2.79 (0.56)	6.2 (0.58)
9.5	2.69 (0.62)	6.1 (0.51)
19	2.66 (0.90)	6.1 (0.64)
37.5	2.46 (0.90)	5.9 (0.68)
75	1.63 (0.43) *	5.3 (0.50)

^{*} statistically significantly different from control (p<0.05)

Conclusion:

NOEC = 37.5 mg/L (nominal) based on mortalities and reduced weights of fish at 75 mg/L Comment (RMS):

At test concentrations of 19 and 75 mg/L on one occasion dissolved oxygen concentrations were found to be below 60 % and at the highest test concentration pH values were repeatedly slightly above 8.5 (max 8.7), the upper limit given in OECD 204. The RMS is of the opinion that the singular cases of lowered oxygen saturation and the slightly elevated pH at the highest test concentration as well as the short periods of reduced temperature did not have an influence on the integrity of the study. Hence the study is considered acceptable.

5.4.2 Aquatic invertebrates

Short-term toxicity to aquatic invertebrates

<u>Reference:</u> Wuethrich V. (1990a): 48-hour acute toxicity of P0071 to *Daphnia magna* (OECD-immobilization test). Document no. 822-001

<u>Test guideline:</u> OECD 202 (1984), US EPA 540/9-85-005 Acute Toxicity Test for Freshwater Invertebrates (1985)

GLP: Yes

Material and methods:

Test substance: P0071 (metaldehyde), purity: 99.3 %, batch: 5448

Test organism: Daphnia magna, less than 24 hours of age

Treatments: Control, solvent control (0.01 % methanol per volume), 1.125, 2.25, 4.5, 9 and 90 mg/L. The concentration of the solvent methanol was 0.01 % per volume at the highest.

Number of animals: 2 replicates with 10 animals each per control and treatment

Duration: 48 hours *Test conditions*:

Test vessels: 50 mL glass beakers, loading: 10 daphnids per 20 mL of test medium, static system, temperature: 21 °C (no values for individual measurements provided in the study protocol), photoperiod: 16 hours light and 8 hours dark, no information about feeding is provided

Test medium:

Reconstituted water prepared according to Directive 84/449 EEC, Part C.2: "Acute Toxicity for Daphnia" was used. Total hardness: 250 mg $CaCO_3/L$, dissolved oxygen concentration: 8.0 - 8.8 mg/L, pH: 8.1 - 8.3. Oxygen and pH were measured at the start and end of the test. *Biological observations*:

Mobility of daphnids was recorded after 24 and 48 hours of exposure. Observations for sublethal effects were not conducted.

Analytical measurements:

Method of analysis: Derivatisation with 2,4-dinitrophenylhydrazin followed by HPLC. Concentration verification: Samples from the solvent control and 1.125, 4.5 and 90 mg/L vessels were taken at the start and end of the test.

Stability test: From an additional test vessel containing 90 mg/L samples were taken at the beginning and after 48 hours of exposure.

Statistical evaluation:

Due to the lack of treatment related effects, no statistical evaluation of the data was performed. Findings:

Analytical results:

The test substance was shown to be stable over the test duration. Measured concentrations (mean of two replicates) were in the range of 75.4 - 81.7 % of nominal concentrations at the beginning of the test and 71.6 - 95.7 % of nominal concentrations at the end of the test. Since measured concentrations were repeatedly below 80 % of nominal, the EC50 was derived from mean measured concentrations.

Immobility:

Table 82: Effect of P0071 (metaldehyde) on immobility of Daphnia magna

Treatment	Replicate	Cumulative imm	obilisation [%]
(nominal concentrations)	Neplicate	24 h	48 h

Treatment	Donlingto	Cumulative imm	obilisation [%]
(nominal concentrations)	Replicate	24 h	48 h
Control without methanol	Α	0	0
Control without methanol	В	0	0
Control with methanol	Α	0	0
(0.01 % per volume)	В	0	0
1.125 mg/L	Α	0	0
1.125 Hig/L	В	0	0
2.25 mg/L	Α	0	0
2.25 Hg/L	В	0	0
4.5 mg/l	Α	0	0
4.5 mg/L	В	10	10
0 ma/l	Α	100	100
9 mg/L	В	0	0
00 mg/l	Α	0	0
90 mg/L	В	0	0

The observed 100 % immobilisation in one replicate at a test concentration of 9 mg/L is not regarded to be treatment related because at the second replicate at this concentration and at a treatment rate of 90 mg/L no immobilisation was recorded.

Conclusion:

 $EC_{50} > 78.4 \text{ mg/L}$ based on mean measured concentrations

Comments (RMS):

The study is acceptable.

Long-term toxicity to aquatic invertebrates

<u>Reference:</u> Wuethrich, V. (1990c): Influence of P0071 on the reproduction of *Daphnia magna*. Document No. 827-001

Test guideline: OECD 202, Part II

GLP: Yes

Material and methods:

Test substance: P0071 (metaldehyde), purity: 99.3 %, batch: 5448

Test organism: Daphnia magna, age: < 24 hours

Treatments:

Control, solvent control, 5.63, 11.25, 22.5, 45 and 90 mg/L, the concentration of the solvent methanol was in all test solutions and in the solvent control 0.01 % v/v

Test duration: 21 days

Test medium:

A mixture of 2/3 of reconstituted water (Directive 84/449 EEC, Part C.2: "Acute Toxicity for Daphnia") and 1/3 of pond water (Anwiler Weiher, CH-4461 Anwil) was used as dilution water. Hardness of the mixture: 256-278 mg CaCO₃/L (measured once per week), pH: 8.0-8.6 in fresh test media and 8.0-9.0 in old test media (measured in the control, solvent control, the highest and lowest test concentrations), dissolved oxygen: 5.9-11.3 mg/L (≥ 60 % saturation) and on one occasion 5.2 mg/L (slightly below 60 % saturation) at the test concentration of 5.63

mg/L.

Test design:

Daphnids were kept in groups of 2 x 10 individuals per treatment in glass beakers of 200 mL until eggs could be observed in the brood pouches (day 5). Then for each treatment ten daphnids with eggs in their brood pouches were separated and kept individually in beakers with 50 mL test medium. The reproduction rate of these individually held daphnids was recorded until the end of the study. The daphnids not selected for individual holding were further on kept in groups (at least 5 daphnids per beaker) and were observed for mortality until the end of the test. In the study protocol no information is provided on the criteria which were applied to select 10 daphnids with eggs in their brood pouches for the reproduction part of the test out of the original 20 individuals per concentration.

Test conditions:

Semi-static test system, test medium renewal: every Monday, Wednesday and Friday, temperature: 20-21 °C (checked at renewal days), photoperiod: 16 hours light and 8 hours dark, food: a mixture of yeast and algae (*Scenedesmus subspicatus*) was fed, feeding was performed with increasing amounts of food, the information provided in the study report is not conclusive on the amounts of food used per test chamber

Analytical measurements:

Method of analysis: Derivatisation with 2,4-dinitrophenylhydrazin followed by HPLC Concentration verification: From the solvent control, the lowest, medium and highest test concentration samples were taken at days 0, 2 and 19 (fresh media) and at day 21 (old media). Stability test: From an additional test vessel containing 90 mg/L test article (without feed and daphnids), samples were taken on days 0, 2 and 21.

Biological observations:

The mortality of adults and the number of young was recorded 3 times per week at test medium renewals.

Statistical evaluation: Steel-test

Findings:

Analytical measurements:

Table 83: Measured concentrations of metaldehyde in test media

Sampling day	Test medium	Test concentrations [% of nominal]				
Sampling day	rest medium	5.63 mg/L	22.5 mg/L	90 mg/L	90 mg/L (stability test)	
0	fresh	83.1	80.0	73.0	90.1	
2	fresh	94.7	90.1	86.2	95.9	
19	fresh	82.2	82.3	67.7	not sampled	
21	old	89.5	83.9	101.4	99.4	

The measured concentrations from day 19 (fresh test media) and day 21 (respective old test media) indicate that the test article was not completely dissolved in fresh test media and that the dissolution process continued throughout the period between medium renewals. This finding was more pronounced at the highest test concentration of nominal 90 mg/L.

Biological observations:

At the second water renewal on day 5 eggs were observed in the brood pouches of daphnids and then 10 daphnids per test concentrations were selected and held individually.

Table 84 Effects of metaldehyde on survival and reproduction of Daphnia magna after 21 days of exposure

Treatment [mg ai/L]	Cumulative no. of young / 10 adults	Mean no. of live young / adult ± s.d.	Immobile out of 20 individuals [%]
Control	1209	121 ± 9	10
Solvent control	1337	134 ± 13	5
5.63	1389	139 ± 7	5
11.25	1390	139 ± 5	5
22.5	1213	135 ± 8	5
45	1263	126 ± 33	5
90	1421	142 ± 18	0

Conclusion:

NOEC = 90 mg/L based on nominal concentrations or 68 mg/L based on mean measured concentrations of fresh test media of the 90 mg/L treatment group, LOEC > 90 / 68 mg/L (nominal / mean measured)

Comment (RMS):

The study was conducted according to a test design which deviates from accepted test guidelines. Daphnids were first held in groups of 2 x 10 individuals per concentration. On day 5 of exposure, when eggs were observed in the brood pouches of daphnids, 10 animals per concentration with eggs in their brood pouches were selected and then held individually. Such a change from group exposure to individual exposure of a subsample of the original group is not foreseen in any accepted test guideline. In the study protocol no information is provided on the criteria which were applied for the selection of the 10 daphnia for individual exposure per treatment level. Also the number of adults with eggs in their brood pouches per concentration on the day of change of exposure regime (from group to individual exposure) is not given. Further deviations from the applied test guideline (OECD 202, part II) are: Daphnids were fed three times a week, they should have been fed daily, the loading was 20 mL of test medium per animal, it should have been 40 mL per animal, the dilution water was a mixture of reconstituted water and pond water, the processing of pond water (e.g. filtration) is not stated, no water quality parameters such as TOC of the pond water are provided in the study protocol. Despite these deviations from accepted test guidelines and shortcomings of the study design, the RMS thinks that the results can be accepted for risk assessment for the following reasons: Up to the highest tested concentration of 90 mg/L no reproductive effects were observed on the 10 selected and individually held daphnids, additionally no treatment related mortalities were observed. The validity criteria of the guidelines OECD 211 and EPA OPPTS 850.1300 are met: mortality of parent animals ≤ 20 % (met for all daphnids of the test), mean number of offspring per surviving control parent at the end of the test is ≥ 60 and no ephippia are produced by control animals (met for the individually held daphnids for which offspring was recorded). Since no effects were observed, it can be assumed, that the quality of the used pond water was acceptable for this test. The mean Koc of metaldehyde is 85 (s.d. 53, n = 8) and hence the adsorption to organic carbon will be rather low. Therefore it can be assumed that the TOC added with the pond water (1/3 of the dilution water) will not have had a major effect on the bioavailability of the test substance.

The study is considered acceptable.

5.4.3 Algae and aquatic plants

<u>Reference:</u> Wuethrich, V. (1990b): Acute toxicity of P0071 to *Scenedesmus subspicatus* (OECD- Algae growth inhibition test). Document no.: 823-001

Test guideline: OECD 201

GLP: Yes

The study is not considered valid for the following reasons:

Concentrations of 0, 1.56, 3.13, 6.25, 12.5, 25, 50 mg metaldehyde/L were tested. Regarding the appearance of the test substance in the test medium the following is stated in the study protocol: "not completely soluble, slightly turbid, particles similar to glass splinters rise on glass – was above liquid level". It was not mentioned at which test concentrations this was observed. Therefore it is not clear whether the test substance was sufficiently solved and hence bioavailable in the test media at the highest test concentrations (25 and 50 mg/L).

Regarding the composition of the nutrient solution for algal cultivation only a reference is provided in the study protocol. From this reference it cannot be judged whether the limits for P, N, chelators and hardness as given in OECD 201 are met.

The pH was adjusted to 7.7 at the beginning of the test, but it was not stated which buffer was used.

After 24 and 48 hours the number of algae was estimated by means of a microscope. After 72 and 96 hours the number of algae was determined spectrophotometrically using a standard curve established with known (counted) cell densities. No information on the validity of the spectrophotometrical biomass estimation (e.g. calibration data) was provided and hence the reliability of the used method could not be evaluated.

No values for growth rates were provided in the study report. Therefore the RMS calculated growth rates from the provided cell density data and found that growth rates were unexpectedly high for the used species *Scenedesmus subspicatus* within the first 24 hours of the study. From 24 to 48 hours growth rates dropped drastically and recovered from 48 to 72 hours (see table B.9.2.1-2).

The ErC_{50} values (72 h and 96 h) as given in the study report are not considered valid by the RMS. The 72 h ErC_{50} was stated to be 21.7 mg/L although even at 50 mg/L (the highest test concentration) the inhibition of the mean growth rate was below 20 %. Also the derivation of the ErC_{50} value for 96 hours could not be comprehended.

In the OECD guideline it is stated that the highest test concentration should inhibit growth at least by 50 % and preferably stop growth completely. As demonstrated in other toxicity tests on fish and *Daphnia magna*, higher test concentrations than 50 mg/L can be achieved and therefore should be tested.

Table B.9.2.1-2: Changes in growth rates of Scenedesmus subspicatus over time at different time at din	ent
treatments with metaldehyde	

Time period	Treatments [mg ai/L]							
Politica	Control	1.56	3.13	6.25	12.5	25	50	
0-24 h	2.44	2.83	2.97	3.05	2.71	1.82	1.36	
24-48 h	0.51	0.39	0.54	0.42	0.22	0.53	0.44	
48-72 h	0.93	0.92	0.32	0.18	0.47	0.76	1.39	
72-96 h	1.65	1.49	1.39	1.65	1.62	1.87	1.91	
Mean 0-72 h	1.30	1.38	1.28	1.22	1.14	1.04	1.06	
Mean 0-96 h	1.39	1.41	1.30	1.33	1.26	1.25	1.28	

Reference: Metaldehyde - A study on the toxicity to algae (Desmodesmus

subspicatus) over 72 hours

Author(s). year: Egeler, P., Junker, T. Knoch, E. (2007)

Report/Doc. number: 823-002

Guideline(s): OECD 201 (2006)

GLP: Yes

Deviations: None of relevance Validity: Acceptable

Test substance: Metaldehyde, batch no.: 36605, purity: 99.5 % (w/w)

MATERIAL AND METHODS:

Test species: Desmodesmus subspicatus

Test concentrations: Nominal: 0 (medium control), 12.5, 25, 50, 100 and 200 mg/L No. of replicates: 3 replicates per test concentration and 6 replicates for the control, 2

replicates for a test substance stability check without algae

Initial loading: 5 x 10³ cells/mL Test type / duration: Static, 72 hours

Nutrient medium: AAP-medium adjusted to a pH of 7.5

Test conditions: Continuous illumination at $76.4 \pm 4.1 \,\mu\text{E m}^{-2}\,\text{s}^{-1}$ (fluorescent tubes,

universal white type) Continuous stirring

Temperature: 22.8 ± 0.24 °C

pH at test start: 7.3 - 7.6. pH at test end: 8.1 - 8.8

Observations: Cell concentrations were evaluated by fluorescent measurements and

conversion of fluorescence units into biomass concentration using a

calibration curve.

Cell measurements after 24, 48 and 72

Microscopic observation of algal cells at the beginning and at the end of

the test

Analytical For chemical analysis of the test substance samples of test solutions from

measurements: all treatment levels were taken at test start and test end. The lowest test

concentration sample (12.5 mg/L) was not analysed.

Analytical method: Gas chromatography with mass selective detection

(GC-MSD)

Statistical evaluation: EC₅₀: no statistics, NOEC: ANOVA followed by Williams' Test

RESULTS

Validity criteria: Biomass increase in the control cultures: 132 (required: \geq 16 within 72 h)

Mean coefficient of variation for section-by-section specific growth rates

in control cultures: 11.6 % (required: \leq 35 %)

Coefficient of variation of average specific growth rates during the test

period in replicate control cultures: 6.7 % (required: ≤ 7 %)

Analytical results: Measured concentrations were found to be 98.2 – 102.8 % of nominal at

test start and 91.3 – 102.2 % at test end

Morphological effects: In the pre-culture cells appeared normal and healthy

0 up to 25 mg/L: Cells appeared healthy at the end of the test \geq 50 mg/L: Cells were partially deformed at the end of the test

Table 85: Effects of metaldehyde on yield and growth rate of Desmodesmus subspicatus after 72 hours of exposure

Metaldehyde	Inhibition after 72 hours [%]				
[mg/L]	Yield	Growth rate			
12.5	6.1	1.5			
25	1.5	0.2			
50	30	7.6			
100	39	9.9			
200	39	10.3			

CONCLUSION:

 $E_y C_{50}$ (72 h): > 200 mg/L $E_r C_{50}$ (72 h): > 200 mg/L

NOEC (72 h): 25 mg/L (based on inhibition of yield and growth rate)

Based on mean nominal concentrations.

5.4.4 Other aquatic organisms (including sediment)

Reference: Metaldehyde- A study on the toxicity to the Great Ramshorn Snail

(Planorbarius corneus) over 48 h

Author(s). year: Egeler, P., Goth, M., Knoch, E. (2007)

Report/Doc. number: 825-001

Guideline(s): No guideline available, OECD 202 (2004) was taken into account

GLP: Yes

Not applicable (no guideline available) **Deviations:**

Acceptable Validity:

Metaldehyde technical, batch no.: 36605, purity: 99.5 % (w/w) Test substance:

Test species: Great Ramshorn Snail (Planorbarius corneus), snails of similar size were used,

body index (ratio of fresh weight to volume): 0.51 - 1.41

Nominal: 0 (control), 9, 19, 41, 91 and 200 mg/L Treatments:

4 replicates with 5 snails each per treatment and control No. of replicates: Static, 48 h followed by a 24 h post exposure period Test type / duration:

Test medium: Elendt Medium M4

Test conditions: Temperature: 20.1 ± 0.5 °C, pH: 7.8

Dissolved oxygen: 4.6 - 9.5 mg/L (51 – 103 % saturation)

Hardness: 268 mg/L as CaCO₃

Photoperiod: 16 light / 8 h dark, No aeration, No feeding during the exposure period

Behaviour: The number of moving snails was recorded before and 1 minute Observations:

after gentle agitation of the vessels.

Immobilisation: After 48 h exposure snails were transferred to vessels

containing clean medium and a piece of cucumber as food source. The location of each animal in the vessel was marked by clearly visible circles on the outer

bottom of the vessel. The location of each snail was also recorded by

photography immediately after transfer. Observations were performed 3, 6 and 24 h after exposure. All animals having changed location were recorded as

mobile.

Analytical At the beginning of the test and at test termination (48 h) samples from all test measurements:

levels were taken, however, only samples from the control, 91 and 200 mg/L

groups were analysed for the test substance.

Analytical method: Gas chromatography with mass selective detection (GC-

MSD)

None (all snails were mobile after 24 hours) Statistical evaluation:

RESULTS

Analytical results: Measured concentrations were in the range of 80 - 98 % of nominal.

After 24 h of exposure in the two highest test concentrations (91 and 200 mg/L) Sublethal effects

95 and 90 % of snails did not move even after gentle agitation of the test (behaviour):

vessels. After 48 h 85 and 100 % of snails did not move at the two highest test concentrations. In the highest test concentration snails were lying on the lateral

side, foot retracted into the shell.

Immobilisation: All animals up to and including the highest test concentration appeared to be

mobile 24 h after termination of exposure.

CONCLUSION: Based on nominal concentrations.

 EC_{50} (48 h): > 200 mg/L (immobility 24 h after termination of exposure) NOEC (48 h): 200 mg/L (immobility 24 h after termination of exposure) Reference: Scholtz, R. (2004): **Toxicity of Metaldehyde-containing slug pellets to Golden**Snails in rice paddy fields. Summary and calculation of acute lethal effect dose using data from efficacy trials conducted by Dupoh Bio Research Centre during 1997 in the

Philippines. Lonza Report No. 3873 Document no.: not assigned

<u>Remark (RMS):</u> The notifier submitted this report to cover the need for information on the toxicity of the molluscicide metaldehyde to aquatic molluscs.

Test guideline: Non

GLP: No

Material and Methods

The aim of the study was to test the efficacy of a metaldehyde-containing pelleted formulation (content of metaldehyde: 10 %) against the Golden Snail (*Pomacea canaliculata*), which is a pest in rice. The study was conducted in the Philippines. The efficacy of 5 different application rates, each used at two different water depths and with 3 replicates per depth was tested. Each replicate consisted of a 20 m² plot in which 50 live adult Golden Snails were set out. The efficacy was measured by counting dead snails up to 14 days after application.

Findings

The notifier used the percentages of dead snails for each treatment at day 5 and 14 to estimate respective LC_{50} values of 1.61 and 1.97 mg a.s./L by probit analyses.

Comment (RMS):

In the submitted study summary percentages of mortalities for 1, 2, 3, 5, and 14 days after application are presented. However, the given mortality data are not conclusive since partly more than 100 % mortality is stated if they were seen as mortalities from one census to the next. On the other hand data cannot be cumulative values since percentages for later time points are partly lower than percentages for earlier censuses (see table 86). Apparently in the study summary numbers of survived snails of each data point were set to 100 %, to calculate the next data point in the timeline. Therefore the RMS asked the notifier for clarification and the submission of the raw data. However, the notifier was not able to submit the raw data and no clarification was given. Thus, the LC_{50} derived from summarised data can not be accepted for classification and labelling. For this reason a recalculation was done by RMS based on the addition of mortalities on each data point which results in a logical series. Results and were presented in table 87.

Table 86: Efficacy of a Metaldehyde-containing pelleted formulation (Metaldehyde 10% granules) against the Golden Snail in rice paddy fields under 2 different levels of water cover. Location: DBRC Farm, Calauan, Laguna, Philippines; data presented in the study summary

Concentration	N	Mean % dead	l snails¹		
per Liter (mg)'	1 DAA ²	2 DAA	3 DAA	5 DAA	14 DAA
0.5	3	0	0	5.33	7.33
0.625	4	2.33	3	4.67	2.33
0.75	0	0	11.33	20	10.67
0.875	3.67	6.67	20	17.67	6.67
1	0	2.67	26	48	19
1.67	7.33	5.33	20.67	40	30
2.08	5.67	21.87	46	61.67	47.67
2.5	7.67	34.33	68.67	79.67	70.33
2.9	14.33	35.33	60	70.67	72
3.33	12.33	36.33	75.67	86	81.67

¹⁾ Mean of 3 replicates

²⁾ Day After Application (DAA)

Table 87: Recalculation of % mortality (by RMS)

	mean % dead snails ¹					
	0 DAA ²	1 DAA	2 DAA	3 DAA	5 DAA	14 DAA
conc mg/L						
0.500	0.00	3.00	3.00	3.00	8.17	14.90
0.625	0.00	4.00	6.24	9.05	13.30	15.32
0.750	0.00	0.00	0.00	11.33	29.06	36.63
0.875	0.00	3.67	10.10	28.08	40.79	44.73
1.000	0.00	0.00	2.67	27.98	62.55	69.66
1.670	0.00	7.33	12.27	30.40	58.24	70.77
2.080	0.00	5.67	26.30	60.20	84.75	92.02
2.500	0.00	7.67	39.37	81.00	96.14	98.85
2.900	0.00	14.33	44.60	77.84	93.50	98.18
3.330	0.00	12.33	44.18	86.42	98.10	99.65

- 1) Mean of 3 replicates
- 2) Day After Application (DAA)

<u>Determined LC₅₀ recalculated Values</u>

14 days after application LC50	> 0.875 and < 1.00 mg/L
5 days after application LC50	> 0.875 and < 1.00 mg/L
3 days after application LC50	> 1,670. and < 2.080 mg/L
calculated LC50 value (with excel logarithmic trend)	1.6 mg/L
2 days after application LC50	> 3.330 mg/L
calculated LC50 value (with excel logarithmic trend)	4.8 mg/L

NOEC Value (14 d) = < 0.5 mg/L

5.4.5 Summary and discussion: Acute (short-term) aquatic toxicity

Data element: Acute (short-	term) aquatic to	xicity Ge	nerally expressed in terms of LC ₅	or EC ₅₀ (r	ng/L)
	L(E)C [mg/I		Test guideline / design	GLP (y/n)	Reliability *
		Fish (96 l	nr LC ₅₀):		
Oncorhynchus mykiss	75		EPA Pesticide Assessment Guidelines, Subdivision E, 72-1 (1985), OECD 203	у	у
Cyprinus carpio	> 10	0	OECD 203	у	
	C	Crustacea (4	8 hr EC ₅₀):		
Daphnia magna	90		OECD 202 (1984), US EPA 540/9-85-005	У	у
	Alg	gae (72 or	96 hr E _r C ₅₀):		
Scenedesmus subspicatus	biomass growth rate		OECD 201	у	n
Desmodesmus subspicatus	yield growth rate	> 200 > 200	OECD 201 (2006)	У	у
		Gastro	ppods		
Planorbarius corneus	> 200 (4	18 h)	No guideline available, OECD 202 (2004) was taken into account	n	у
Pomacea canaliculata	> 3.330 ((48 h)	No guideline	n	y

Conclusion:

For classification and labeling the most sensitive species *Pomacea canaliculata* with a recalculated LC50 = >3.3 mg/L was used.

5.4.6

5.4.7 Summary and discussion: Chronic (long-term) aquatic toxicity

-term) aquatic tox	cicity			
NOEC (mg/L)				
		Test guideline / design	GLP (y/n)	Reliability *
Fi	sh (21 d NOEC):			
mortality body weight	37.5	OECD 204	у	у
Crus	tacea (21 d NOEC):			
immobility reproduction	90	OECD 202, Part II	у	у
Algae/ac	quatic plants (NOEC	2):		
biomass growth rate		OECD 201	у	n
yield growth rate	25 25	OECD 201 (2006)	у	у
Mortality	< 0.5 mg	No guideline	n	y
	Final mortality body weight Crus immobility reproduction Algae/act biomass growth rate yield growth rate	NOEC (mg/L) NOEC [mg/L] Fish (21 d NOEC): mortality body weight Crustacea (21 d NOEC): immobility reproduction Algae/aquatic plants (NOEC) biomass growth rate yield growth rate 25 growth rate 25	NOEC (mg/L)	NOEC (mg/L)

For classification and labeling the most sensitive species *Pomacea canaliculata* with a NOEC = < 0.5 mg/L was

5.5 Comparison with criteria for environmental hazards (sections 5.1 - 5.4)

Endpoint			on Criteria a in bold)	Conclusion for Metaldehyd
	CLP (2 nd ATP)	DSD	
Degradation				
	Metaldehyde is photo pH values.	olytically stable and hyd	Irolytically stable at environmentally relevant	The classification as R53 according to Directive 67/548/EEC is based on the fact that the active substance is not considered
	In UK simulation stu	eadily biodegradable un 1dy (two less oxidising sy 1ese data, the substance is 1 degradable.	as ready biodegradable/rapid degradable.	
Bioaccumulatio	n			
Criteria LogKow		K_{ow} is < 4 $Log K_{\text{ow}} = 0.12$	$\begin{array}{c} \text{Log } K_{ow} \text{ is } < 3 \\ \text{Metaldehyde Log } K_{ow} = 0.12 \end{array}$	The measured BCF is 11 and is below the classification criteria of 100 (DSD) and below the classification criteria of
Criteria BCF	BCF < 500 Metaldehyde BCF is 11		BCF < 100 Metaldehyde BCF is 11	500 (CLP). Therefore depending on classification criteria Metaldehyde is considered to have a low bioaccumulation potential.
Acute aquatic to	xicity			
Criteria			$LC/EC_{50} > 1 \text{ mg/L} \le 10 \text{ mg/L}$	Metaldehyde is toxic to <i>Pomacea canaliculata</i> with an EC50 of >3.33 mg/L and fulfills the criteria for the proposed
			Pomacea canaliculata LC50 > 3.33 mg/L	classification as R51 according to Directive 67/548/EEC. No classification was proposed according to Regulation EC 1272/2008.
Chronic aquatic	toxicity			
Criteria	For non rapidly degradable substances: 0.1 <noec l<="" mg="" td="" ≤1=""><td></td><td>Metaldehyde is chronic toxic to aquatic molluscs with a NOEC of < 0.5 mg/L Therefore Metaldehyd fulfills the criteria for the proposed classification as H411 according to Regulation</td></noec>			Metaldehyde is chronic toxic to aquatic molluscs with a NOEC of < 0.5 mg/L Therefore Metaldehyd fulfills the criteria for the proposed classification as H411 according to Regulation
	Pomacea canaliculata	NOEC(14 d) = <0.5 mg/L		EC 1272/2008.

5.6 Conclusions on classification and labelling for environmental hazards (sections 5.1 – 5.4)

Conclusion of environmental classification according to Directive 67/548/EEC

Metaldehyde should be classified "Dangerous for the Environment" with the following risk and safety phrases:

N Dangerous for the Environment

R51 Very toxic to aquatic organisms

R53 May cause long term effects in the environment

Conclusion of environmental classification according to Regulation EC 286/2011 (2nd ATP to EC 1272/2008)

Based on the CLP Regulation, metaldehyde should be classified as:

Classification categories aquatic environmental hazard chronic category 2

GHS Pictogram



Signal Word Warning

Hazard Statement H411 'Toxic to aquatic life with long lasting effects'

6 OTHER INFORMATION

Environmental fate properties and environmental hazard assessments of this CLH report are based on studies and summaries of the Draft Assessment Report and its addenda.

7 REFERENCES

7.1 Physico-chemical properties

Author(s)	Year	Title Source (where different from company) Company, Report No GLP or GEP status (where relevant), Published or not	Data Protection Claimed Y/N-R/NR	Owner
Bohle, J. F.	1989	Determination of the water solubility of P0071 RCC, 026021 LR 1390 N	N	LON
Bohle, J. F.	1989	Determination of the solubility of P0071 in different solvents RCC, 016457 LR 1391 Yes No	No	LON
Cardinaals, J. M.	1988	Determination of the vapour pressure of P0071 RCC, 00854/C559; LR 1386 No No	N	LON
Cardinaals, J. M.	1988	Calculation of Henry's law constant of P0071 RCC, 00854/C588 LR 1389 No No	N	LON
Cardinaals, J. M.	1988	Determination of the solubility of P0071 in water RCC, 00854/C558 LR 1388 No No	N	LON
Cardinaals, J. M.	1988b	Determination of the partition coefficient of P0071 RCC, 00854/C560 LR 1387 No No	N	LON
Carpenter, M.	1989	Hydrolysis of Metaldehyde as a function of pH at 25℃ ABC, 37146 LR 1410 No No	N	LON
Carpenter, M.	1989a	Photodegradation of Metaldehyde in ph 7 buffered solution ABL, 37766 LR 1412 No No	N	LON
Carpenter, M.	1989a	Photodegradation of Metaldehyde in ph 7 buffered solution ABL, 37766 LR 1412 No No	N	LON

Author(s)	Year	Title Source (where different from company) Company, Report No GLP or GEP status (where relevant), Published or not	Data Protection Claimed Y/N-R/NR	Owner
Comb, A.L.	2007	Metaldehyde - Melting temperature Huntingdon Life Science Report No.: LZA 0287/072574 LR 4077 GLP, unpublished Doc. No.: 112-002	Y	LON
Comb, A.L.	2009	!! Confidential !! - Acetaldehyde Spectra Huntingdon Life Science Lonza Rep. No. 4384 Report No.: LZA0315 GLP, unpublished Doc. No.: 157-001	Y	LON
Comb, A.L.	2007	Metaldehyde - Solvent solubility Huntingdon Life Science Report No.: LZA0288/072599 LR 4078 GLP, unpublished Doc. No.: 114-006	Y	LON
Hogg, A. S.	1998	Meta Metaldehyde: determination of general physico- chemical properties SPL, 102/278 LR 2993 Yes No	N	LON
O'Connor, B. J., Mullee, D. M.	2001	Meta Metaldehyde (CODE LZ1060): determination of general physico-chemical properties and spectra SPL, 102/353 LR 3340 Yes No	Y	LON
O'Connor, B. J., Mullee, D. M.	2001	Meta Metaldehyde (CODE LZ1060): determination of general physico-chemical properties and spectra SPL, 102/353 LR 3340 Yes No	Y	LON
O'Connor, B. J., Mullee, D. M.	2001	Meta Metaldehyde (CODE LZ1060): determination of general physico-chemical properties and spectra SPL, 102/353 LR 3340 Yes No	Y	LON
O'Connor, B. J., Mullee, D. M.	2001a	Meta Metaldehyde (CODE LZ1060): determination of solubility in ethyl aceto acetate SPL, 102/388 LR 3332 Yes No	Y	LON
Tremain, S. P.	2001	Meta Metaldehyde (Code LZ1060): determination of hazardous physico-chemical properties SPL, 102/354 LR 3338 Yes No	Y	LON

Author(s)	Year	Title Source (where different from company) Company, Report No GLP or GEP status (where relevant), Published or not	Data Protection Claimed Y/N-R/NR	Owner
Voget, M.	1994	Calculation of the photochemical oxidative half-life of Meta Metaldehyde ECO, 94-23-11 LR 2293 no No	N	LON
Weiss, A.	2009	Statement related to the oxidising properties of Metaldehyde Scientific Consulting Company, Wendelsheim, Germany Report No.: 262-013 Not GLP, unpublished Doc. No.: 143-001	Y	LON

7.2 Toxicological properties

Author(s)	Year	Title Source (where different from company) Company, Report No GLP or GEP status (where relevant), Published or not	Data Protection Claimed Y/N-R/NR	Owner
Selim S.	1992	Pharmacokinetic study of ¹⁴ C-metaldehyde in the rat following oral administration Biological Test Center; CA, USA Doc.No. 512-001, Lonza Report No. 1899 GLP: yes unpublished	N	LONZA
Jones J., Collier T.	1987	P0071: OECD 401 Acute oral toxicity test in the rat, Project Number 102/9A Safepharm Laboratories Ltd., Derby, UK Doc.No. 521-002, Lonza Report No.1354 GLP: yes unpublished	N	LONZA
Coles R.	1990a	P0071: Acute oral toxicity test in the mouse, Project Number 102/50 Safepharm Laboratories Ltd., Derby, UK Doc.No. 521-001, Lonza Report No. 1325 GLP: yes unpublished	N	LONZA
Durando, J.	2009	Acute Oral Toxicity Study with META Metaldehyde techn. CAS No. 108-62-3: Up-And-Down Procedure in Rats Eurofins/Product Safety Laboratories, USA Laboratory Identification No. 26776 Lonza Report No. 4377 / Doc. No.: 521-003 GLP Unpublished	Y	Lonza
Davies R., Collins C.	1974	Acute percutaneous toxicity to rats of metaldehyde Huntington Research Centre, UK Doc.No. 522-001, Lonza Report No. 1360 GLP: no unpublished	N	LONZA
Berczy Z., Cobb L., Cherry C.	1973	Acute inhalation toxicity to the rat of metaldehyde dust Huntington Research Centre, UK Doc.No. 523-001, Lonza Report No. 1362 GLP: no unpublished	N	LONZA
Griffiths, D.R.	2009	Outcome of technical pre-trials for an acute inhalation toxicity study with Metaldehyde Harlan Laboratories Ltd., Derbyshire, UK; Ref: L260109-01/vm Lonza Report No. 4336, Doc. No.: 581-004 GLP: not applicable Unpublished	Y	Lonza
Jones J.	1983	P0071: A primary skin irritation and corrosivity study in the rabbit Hazleton Laboratories Europe Ltd., UK Doc.No. 565-001, Lonza Report No. 1373 GLP: yes unpublished	N	LONZA

Author(s)	Year	Title Source (where different from company) Company, Report No GLP or GEP status (where relevant), Published or not	Data Protection Claimed Y/N-R/NR	Owner
Coles R.	1990b	P0071: Acute eye irritation test in the rabbit, project number 102/63 Safepharm Laboratories Ltd., Derby, UK Doc.No. 566-001, Lonza Report No. 1351 GLP: yes unpublished	N	LONZA
Nitka S.	1984	Guinea pig sensitization (Buehler) Consumer Product Testing, NJ, USA Doc.No. 567-001, Lonza Report No. 1378 GLP: yes unpublished	N	LONZA
Bull, A.D.	2007	LZ1060 Metaldehyde. Assessment of skin sensitization potential using the Local Lymph Node Assay in the mouse Huntingdon Life Sciences Limited, Cambridgeshire UK; LZA 0292/064237/LN Lonza Report No. 4064, Doc. No.: 567-003 GLP Unpublished	Y	Lonza
Van Miller J.	1989	Twenty-Eight Day Dietary Oral Toxicity Study with Metaldehyde in rats Bushy Run Research Center, PA, USA Doc.No. 532-001, Lonza Report No. 1380 GLP: yes unpublished	N	LONZA
Thomas O., Bartlett A., Brooks P.	1998	P0071: Ninety day sub-chronic oral (dietary) toxicity study in the rat Safepharm Laboratories Ltd., Derby, UK Doc.No. 533-003, Lonza Report No. 2974 GLP: yes unpublished	Y	LONZA
Gill M., Wagner C.	1990	Ninety-Day Dietary Dose Range Finding Study with Metaldehyde in Mice Bushy Run Research Center, PA, USA Doc.No. 533-002, Lonza Report No. 1546 GLP: yes unpublished	N	LONZA
Leuschner, J.	2002	4-week dose-range-finding study for a 52-week chronic toxicity study of metaldehyde by oral administration via the diet to Beagle dogs LPT Laboratory of Pharmacology and Toxicology KG, Hamburg, Germany; LPT Report No. 14543/01 Lonza Report No. 3506, Doc. No.: 532-003 GLP Unpublished	Y	Lonza
Neumann W. + Neumann W.	1980 + 1991	26-weeks-toxicity of metaldehyde 99% - called "Metaldehyd" - in Beagle dogs after oral administration + Supplement No. 1 for 26-weeks-toxicity of metaldehyde 99% - called "Metaldehyd" - in Beagle dogs after oral administration LPT Laboratory of Pharmacology and Toxicology KG, Hamburg, Germany; LPT Lonza Report No. 1379 Part 1+2, Doc. No.: 533-001 GLP: no, study was performed before adoption of OECD Guideline 452 Unpublished	N	Lonza

Author(s)	Year	Title Source (where different from company) Company, Report No GLP or GEP status (where relevant), Published or not	Data Protection Claimed Y/N-R/NR	Owner
Leuschner, J.	2009	Histological re-examination of the testes and re-evaluation of the findings of the 26-week toxicity of metaldehyde 99% in Beagle dogs after oral administration (LPT Study report dated March 31, 1980) LPT Laboratory of Pharmacology and Toxicology, Hamburg, Germany; LPT Report No. 24158, Doc. No.: 581-005 GLP: not applicable Unpublished	Y	Lonza
Leuschner J.	2003	52-week chronic toxicity study of metaldehyde by repeated oral administration via the diet to Beagle dogs LPT Laboratory of Pharmacology and Toxicology KG, Hamburg, Germany Doc.No. 537-003, Lonza Report No. 3657 GLP: yes unpublished	Y	LONZA
Leuschner, J. Drommer W.	2006	Expert statement on the histological findings (giant cells, atrophy and degeneration of the germinative epithelium) in the 52-week toxicity study in Beagle dogs with metaldehyde LPT Laboratory of Pharmacology and Toxicology, Hamburg, Germany; LPT Report No. 15050/01, Doc. No.: 581-001 GLP: not applicable Unpublished	Y	Lonza
Hermansky S., Wagner C.	1991	21-day repeated cutaneous dose toxicity study with metaldehyde in New Zealand White rabbits Bushy Run Research Center, PA, USA Doc.No. 532-002, Lonza Report No. 1800 GLP: yes unpublished	N	LONZA
Thompson P.	1998	P0071: Reverse mutation assay "Ames Test" using Salmonella typhimurium and Escherichia coli Safepharm Laboratories Ltd., Derby, UK Doc.No. 557-006, Lonza Report No. 2972 GLP: yes unpublished	Y	LONZA
Friederich U., Wuergler F.	1981	Salmonella / Microsome assay with metaldehyde Institut für Toxikologie, Universitaet Zuerich, Switzerland Doc.No. 557-001, Lonza Report No. 1381 GLP: no unpublished	N	LONZA
Debets F., Enninga I.	1986	Evaluation of the mutagenic activity of P0071 in an in vitro mammalian cell gene mutation test with L5178Y mouse lymphoma cells NOTOX C.V., `S-Hertogenbosch, Netherlands Doc.No. 557-002, Lonza Report No. 1382 GLP: yes unpublished	N	LONZA

Author(s)	Year	Title Source (where different from company) Company, Report No GLP or GEP status (where relevant), Published or not	Data Protection Claimed Y/N-R/NR	Owner
Debets F.	1986	Evaluation of the ability of P0071 to induce chromosome aberrations in cultured Chinese Hamster Ovary (CHO) cells NOTOX C.V., `S-Hertogenbosch, Netherlands Doc.No. 557-003, Lonza Report No. 1383 GLP: yes unpublished	N	LONZA
Мау К.	1992	P0071: Assessment of its ability to cause lethal DNA damage in strains of <i>Escherichia coli</i> Life Science Research Ltd., Suffolk, England Doc.No. 557-004, Lonza Report No. 1900 GLP: yes unpublished	N	LONZA
Jenkinson P.	1990	P0071: OECD 474 Micronucleus test in the mouse Safepharm Laboratories Ltd., Derby, UK Doc.No. 557-005, Lonza Report No. 1507 GLP: yes unpublished	N	LONZA
Gill M., Wagner C.	1992	Chronic dietary toxicity / oncogenicity study with metaldehyde in rats Bushy Run Research Center, PA, USA Doc.No. 537-002, Lonza Report No. 1550 GLP: yes unpublished	N	LONZA
Chun J., Wagner C.	1993	Chronic dietary oncogenicity study with metaldehyde in mice Bushy Run Research Center, PA, USA Doc.No. 537-001, Lonza Report No. 1549 GLP: yes unpublished	N	LONZA
Beyrouty P.	1998	A chronic dietary oncogenicity study with metaldehyde in mice ClinTrials BioResearch Ltd., Senneville, Quebec H9X 3R3, Canada Laboratory Project I.D. 87013 Lonza Report No. 2976, Doc. No.: 555-002 GLP Unpublished	N	Lonza
Harder V., Roth T., Hofer M.	2010	Review of carcinogenicity studies with Metaldehyde SCC Scientific Consulting Company, Wendelsheim, Germany SCC Project No. 262-004; Report date 2010-04-22, Doc. No.: 581-009 GLP: not applicable Unpublished	Y	Lonza
Chun J., Neeper- Bradley T.	1993	Two generation reproduction study in CD rats with metaldehyde administered in the diet Bushy Run Research Center, PA, USA Doc.No. 543-001, Lonza Report No. 1544 GLP: yes unpublished	N	LONZA

Author(s)	Year	Title Source (where different from company) Company, Report No GLP or GEP status (where relevant), Published or not	Data Protection Claimed Y/N-R/NR	Owner
Neeper-Bradley T., Chun J.	1990	Developmental toxicity evaluation of metaldehyde administered by gavage to CD (Sprague Dawley) rats Bushy Run Research Center, PA, USA Doc.No. 551-003, Lonza Report No. 1545 GLP: yes unpublished	N	LONZA
Neeper-Bradley T.	1990a	Developmental toxicity dose range-finding study of metaldehyde administered by gavage to New Zealand White rabbits Bushy Run Research Center, PA, USA Doc.No. 551-001, Lonza Report No. 1503 GLP: yes unpublished	N	LONZA
Neeper-Bradley T.	1990b	Developmental toxicity evaluation of metaldehyde administered by gavage to New Zealand White rabbits Bushy Run Research Center, PA, USA Doc.No. 551-002, Lonza Report No. 1504 GLP: yes unpublished	N	LONZA
Haferkorn J.	2009	Acute neurotoxicity study in rats by oral administration of metaldehyde LPT Laboratory of Pharmacology and Toxicology, Hamburg, Germany; LPT Report No. 23443, Doc. No.: 541-002 GLP Unpublished	Y	Lonza
Jones L., Finn J., Mullee D.	2003	LZ1060, metaldehyde: ninety day repeated dose oral (dietary) neurotoxicity toxicity study in the rat Safepharm Laboratories Ltd., Derbyshire, UK Doc.No. 533-004, Lonza Report No. 3644 GLP: yes unpublished	Y	LONZA
Harder V., Roth T., Hofer M.	2010	Relevant aspects of neurological effects associated with exposure to Metaldehyde SCC Scientific Consulting Company, Wendelsheim, Germany SCC Project No. 262-004; Report date 2010-04-22, Doc. No.: 581-008 GLP: not applicable Unpublished	Y	Lonza
Flügge C.	2009	Validation of an analytical method for the determination of metaldehyde in test item formulations by GC-FID method LPT Laboratory of Pharmacology and Toxicology, Hamburg, Germany; LPT Report No. 23202, Doc. No.: 411-002 GLP Unpublished	Y	Lonza
Booze T., Oehme F.	1986	An investigation of metaldehyde and acetaldehyde toxicities in dogs Fundamental and Applied Toxicology 6, 440-446 (1986) Doc.No. 592-026 GLP: no Published	N	Published data

Author(s)	Year	Title Source (where different from company) Company, Report No GLP or GEP status (where relevant), Published or not	Data Protection Claimed Y/N-R/NR	Owner
Sparks S.E. et al.	1996	Metaldehyde molluscicide action in mice: distribution, metabolism and possible relation to GABAergic system Pesticide Biochemistry and Physiology 55, 226-236, 1996 Doc. No. 592-005 GLP: no Published	N	Published data
Joos R., Matter S.	2003	"To whom it may concern" (Statement regarding medical information on plant personnel) Lonza AG Doc.No. 574-002 GLP: not applicable unpublished	N	LONZA
Borbely A.	1970	The problem of metaldehyde poisoning Inaugural dissertation for obtaining a doctorate at the Medical Faculty of the University of Zurich Doc.No. 592-001 GLP: no published	N	N.R.
Moody J., Inglis F.	1992	Persistence of metaldehyde during acute molluscicide poisoning Human & Experimental Toxicology 11, 361-362 (1992) Doc.No. 592-029 GLP: no published	N	N.R.
Thompson J., Casey P., Vale J.	1995	Deaths from pesticide poisoning in England and Wales 1990-1991 Human & Experimental Toxicology 14, 437-445 (1995) Doc.No. 592-030 GLP: no published	N	N.R.
Longstreth W., Pierson D.	1982	Metaldehyde poisoning from slug bait ingestion The Western Journal of Medicine 137, 134-137 (1982) Doc.No. 592-033 GLP: no published	N	N.R.
Booze T. and Oehme F.	1985	Metaldehyde toxicity: a review Vet Hum Toxicol 27(1), 11-19 (1985) Doc.No. 592-025 GLP: no published	N	N.R.
Lisi P., Caraffini S., Assalve D.	1987	Irritation and sensitization potential of pesticides Contact Dermatitis 17, 212-218 (1987) Doc.No. 592-002 GLP: no published	N	N.R.
Booze T. and Oehme F.	1985	Metaldehyde toxicity: a review Vet Hum Toxicol 27(1), 11-19 (1985) Doc.No. 592-025 GLP: no published	N	N.R.

Author(s)	Year	Title Source (where different from company) Company, Report No GLP or GEP status (where relevant), Published or not	Data Protection Claimed Y/N-R/NR	Owner
Von Burg R.	1991	Toxicology Update: Metaldehyde Journal of Applied Toxicology 11(5), 377-378 (1991) Doc.No. 592-028 GLP: no published	N	N.R.
Borbely A.	1970	The problem of metaldehyde poisoning Inaugural dissertation for obtaining a doctorate at the Medical Faculty of the University of Zurich Doc.No. 592-001 GLP: no published	N	N.R.
Mayer S.	1991	Poison metaldehyde In Practice, March 1991 Doc.No. 592-010 GLP: no published	N	N.R.
Booze T. and Oehme F.	1986	An investigation of metaldehyde and acetaldehyde toxicities in dogs Fundamental and Applied Toxicology 6, 440-446 (1986) Doc.No. 592-026 GLP: no published	N	N.R.
Anonymous	1991	Safety Data Sheet Meta Metaldehyde techn. Lonza Group, Version 20.08.2001 Doc.No. 955-004 GLP: not applicable unpublished	N	LONZA
Mayer S.	1991	Poison metaldehyde In Practice, March 1991 Doc.No. 592-010 GLP: no published	N	N.R.
Mayer S.	1991	Poison metaldehyde In Practice, March 1991 Doc.No. 592-010 GLP: no published	N	N.R.

7.3 Environmental effects

7.3.1 Environmental fate and behaviour

		Title	Data	Owner
Author(s)	Year	Source (where different from company)	Protection	
		Company, Report No	Claimed	
		GLP or GEP status (where relevant),		
		Published or not	Y/N-R/NR	

Author(s)	Year	Title Source (where different from company) Company, Report No GLP or GEP status (where relevant), Published or not	Data Protection Claimed Y/N-R/NR	Owner
Cardinaals, J. M.	1990	Calculation of the volatility of P0071 Generated by: RCC Document No: 1510 GLP / GEP Yes unpublished	N	LONZA
Carpenter, M.	1989 a	Hydrolysis of Metaldehyde as a function of pH at 25°C Generated by: ABC Document No: 1410 GLP / GEP Yes unpublished	N	LONZA
Carpenter, M.	1989 b	Photodegradation of Metaldehyde in ph 7 buffered solution Generated by: ABL Document No: 1412 GLP / GEP Yes unpublished	N	LONZA
Cranor, W.	1990 a	Aerobic soil metabolism of ¹⁴ C-Metaldehyde Generated by: ABC Document No: 1472 GLP / GEP Yes unpublished	N	LONZA
Cranor, W.	1990 b	Anaerobic soil metabolism of ¹⁴ C-Metaldehyde Generated by: ABC Document No: 1424 GLP / GEP Yes unpublished	N	LONZA
De Vette, H. Q. M. Aalderink, G. H.	2002	A study on the adsorption of [U-14-] Metaldehyde to soil particles in four soil types Generated by: TNO Document No: 3551 GLP / GEP Yes unpublished	Y	LONZA
Groß, R.	2000	Determination of the aerobic soil degradation of ¹⁴ C-Metaldehyde – Recalculation of DT ₅₀ and DT ₉₀ values Generated by: SCC Document No: 1634 GLP / GEP not applicable unpublished	N	LONZA
Heim, D. Daly, D.	1999	Soil/ sediment adsorption-desoprtion of ¹⁴ C-Metaldehyde Generated by: ABC Document No: 3123 GLP / GEP Yes unpublished	N	LONZA
Kabler, K.	1990	Determination of the photolysis rate of Metaldehyde on the surface of soil Generated by: ABC Document No: 1425 GLP / GEP Yes unpublished	N	LONZA
Möllerfeld, J. Römbke, J. eller, M.	1993	Determination of the degradeability and persistence of ¹⁴ C-Metaldehyde in the water / sediment-system Generated by: BAT Document No: 2120 GLP / GEP Yes unpublished	N	LONZA

Author(s)	Year	Title Source (where different from company) Company, Report No GLP or GEP status (where relevant), Published or not	Data Protection Claimed Y/N-R/NR	Owner
Mônego, J.G.	2005	Inherent biodegradability of ¹⁴ C-Metaldehyde in Brazilian Soils. Generated by: Bioensaios Document No: 3949 GLP / GEP Yes unpublished	Y	LONZA
Römbke, J. Möllerfeld, J.	1991	Determination of the aerobic soil degradation of ¹⁴ C-Metaldehyde Generated by: BAT Document No: 1634 GLP / GEP Yes unpublished	N	LONZA
Juozenaite, A.	2009	 14C-Metaldehyde route and rate of degradation in aerobic soil '02-A' Doc. No. 722-004 GLP Yes unpublished 	Y	LONZA
Juozenaite, A.	2009	 ¹⁴C -Metaldehyde Route and Rate of Degradation in Aerobic Soil 'Elmton (294)' Doc. No. 722-005 GLP Yes unpublished 	Y	LONZA
Juozenaite, A.	2009	14C -Metaldehyde route and rate of degradation in aerobic soil 'Fladbury' Doc. No. 722-006 GLP Yes unpublished	Y	LONZA
Juozenaite, A.	2009	14C -Metaldehyde route and rate of degradation in aerobic soil 'Fladbury' Doc. No. 722-007 GLP Yes unpublished	Y	LONZA
Juozenaite, A.	2009	Investigation of Media to Trap Volatile Products of Metaldehyde Degradation Doc. No. 741-002 GLP No unpublished	Y	LONZA
Selim, S.	1993	Laboratory volatility of Metaldehyde from soil Generated by: BTC Document No: 742-001 GLP / GEP Yes unpublished	N	LONZA
Voget, M.	1994	Calculation of the photochemical oxidative half-life of Meta Metaldehyde Generated by: ECO Document No: 2293 GLP / GEP Yes unpublished	N	LONZA
Wuethrich, V.	1990 a	Ready biodegradability: Modified OECD screening test for P0071 Generated by: RCC Document No: 1490 GLP / GEP Yes unpublished	N	LONZA

Author(s)	Year	Title Source (where different from company) Company, Report No GLP or GEP status (where relevant), Published or not	Data Protection Claimed Y/N-R/NR	Owner
Wuethrich, V.	1990 b	P0071: Inherent biodegradability "modified Zahn-Wellens Test" Generated by: RCC Document No: 1488 GLP / GEP Yes unpublished	N	LONZA
Lebertz, H.	2008	Study on ready biodegradability of Metaldehyde Document No: 713-003 GLP / GEP Yes unpublished	Y	LONZA
Kane, T.	2009	14C-Metaldehyde Aerobic Transformation in Aquatic Sediment Systems Document No: 714-003 GLP Yes unpublished	Y	LONZA
Klein, C.	2009	91/414/EEC Review of Metaldehyde – Revised estimation of the degradation rates of Metaldehyde to be used in environmental fate modelling Doc. No. 782-011 Not GLP unpublished	Y	LONZA
Peter, S.	2009	91/414/EEC Review of Metaldehyde - Evaluation of the degradation kinetics of Metaldehyde and its degradation product Acetaldehyde in aquatic systems Document No: 782-015 GLP No unpublished	Y	LONZA
Römbke, J.	2009	Statement about the water/sediment study with ¹⁴ C-Metaldehyde - Arrangement of the traps Doc. No: 714-002 GLP No unpublished	Y	LONZA

7.3.2 Ecotoxicology

Author(s)	Year	Title Source (where different from company) Company name, Report No., GLP status (where relevant) published or not	Data protect. claimed	Owner
Bogers, M.	1990a	96-hour acute toxicity study in the rainbow trout with P0071 in a semi-static system RCC Report No. 000348, LR 1489 Document No.: 821-001 GLP: Yes, not published	No	Lonza
Bogers, M.	1990b	Prolonged toxicity study in the rainbow trout with P0071 RCC Report No. 029295, LR 1517 Document No. 826-001 GLP: Yes, not published	No	Lonza
Brooke, L. T., Call, D. J., Geiger, D. L., Northcott, C. E. (eds.)	1984	Acute toxicities of organic chemicals to fathead minnows (<i>Pimephales promelas</i>), Volume I, Center for Lake Superior Environmental Studies, University of Wisconsin-Superior, pp 1-13 and 29-30. Document No. 892-009 GLP: No, published	No	not reported
Egeler, P. Junker, T. Knoch, E.	2007	Metaldehyde - A study on the toxicity to Algae (Desmodesmus subspicatus) over 72 h SGS Institut Fresenius GmbH Report No.: AR1AO LR 4090 GLP, unpublished Doc. No.: 823-002	Yes	Lonza
Geiger, D. L., Brooke L. T., Call, D. J. (eds.)	1990	Acute toxicities of organic chemicals to fathead minnows (<i>Pimephales promelas</i>), volume V, Center for Lake Superior Environmental Studies, University of Wisconsin-Superior, pp 33-34. Document No. 892-010 GLP: No, published	No	not re- ported
Sved, D. W. Holmes, C. M. Smith, G. J.	1992	A bioconcentration study with Metaldehyde in the bluegill (<i>Lepomis macrochirus</i>) WLD Report No. 289A-102, LR 1959 Document No. 872-001 GLP: Yes, not published	No	Lonza
Wetton, P. M., Mullee, D. M.	2001	Meta Metaldehyde (Code LZ1060): aute toxicity to common carp (<i>Cyprinus carpio</i>) SPL Report No.: 102/355, LR 3345, Document No. 821-002 GLP: Yes, not published	Yes	Lonza
Wuethrich, V.	1990a	48-hour acute toxicity of P0071 to <i>Daphnia magna</i> (OECD-immobilization test) RCC Report No. 267658, LR 1539 Document No. 822-001 GLP: Yes, not published	No	Lonza

Author(s)	Year	Title Source (where different from company) Company name, Report No., GLP status (where relevant) published or not	Data protect. claimed	Owner
Wuetherich, V.	1990b	Acute toxicity of P0071 to Scenedesmus subspicatus (OECD - algae growth inhibition test) RCC Report No. 259953, LR 1540 Document No. 823-001 GLP: Yes, not published	No	Lonza
Wuetherich, V.	1990c	Influence of P0071 on the reproduction of <i>Daphnia magna</i> RCC Report No. 259942, LR 1538 Document No. 827-001 GLP: Yes, not published	No	Lonza
Scholtz, R	2004	Toxicity of Metaldehyde-containing slug pellets to Golden Snails in rice paddy fields. Summary and calculation of acute lethal effect dose using data from efficacy trials conducted by Dupoh Bio Research Centre during 1997 in the Philippines. Lonza Report No. 3873 GLP: No	Yes	Lonza
Egeler, P., Goth, M., Knoch, E.	2007	Metaldehyde- A study on the toxicity to the Great Ramshorn Snail (<i>Planorbarius corneus</i>) over 48 h Report/Doc. number: 825-001 Guideline(s): No guideline available, OECD 202 (2004) was taken into account GLP: Yes	Yes	Lonza

8 ANNEXES