FOREWORD

INTRODUCTION

TRIETHYLENE TETRAMINE
CAS Nº: 112-24-3

SIDS Initial Assessment Report for SIAM 8

(Paris, 28-30 October 1998)

Chemical Name : Triethylenetetramine

CAS No: 112-24-3

Sponsor Country: Germany

National SIDS Contact Point in Sponsor Country: Dr Jan Ahlers

HISTORY:

The SIDS Initial Assessment Report was discussed at SIAM 5 & 6 and adopted at SIAM 8.

COMMENTS:

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SIDS INITIAL ASSESSMENT PROFILE

CAS No.	112-24-3	
Chemical Name	Triethylene tetramine	
Structural Formula	H ₂ N-CH ₂ -CH ₂ -NH-CH ₂ -CH ₂ -NH-CH ₂ -CH ₂ -NH ₂	
CONCLUSIONS AND RECOMMENDATIONS		

Environment

The chemical is toxic to algae, but PEC/PNEC ratios are lower than 1. It is currently considered of low potential risk and low priority for further work.

Human Health

The chemical is genotoxic *in vitro*, a severe irritant to skin and eyes and a skin sensitiser, but exposure is low and well-controlled. Therefore, it is currently considered of low potential risk and low priority for further work. However due to its hazard character appropriate classification and labelling are recommended.

SHORT SUMMARY WHICH SUPPORTS THE REASONS FOR THE CONCLUSIONS AND RECOMMENDATIONS

The production volume of triethylenetetramine (TETA) in 1990 is 1200-1500 t/a in Germany, ca. 6000 t/a in the Netherlands, >11000 t/a in the USA and ca. 1800 t/a in Japan. TETA is mostly used as intermediate in chemical synthesis. Ca. 160 t/a are directly used as curing agent for epoxy resins in Germany. For Sweden, a similar use pattern was described. TETA is stable in neutral solution and is classified as "non biodegradable". The most sensitive environmental species to TETA is the alga *Scenedesmus subspicatus* (72h-EC10 = 0.67 mg/l). A PNEC of 13.4 µg/l is determined.

TETA has a moderate acute toxicity: LD50 (oral, rat) > 2000 mg/kg bw, LD50 (dermal, rabbit) = 550.805 mg/kg bw. The NOAEL for repeated dose toxicity is 600 ppm (92 (male), 99 (female) mg/kg bw) for mice (oral, 90 days). In *in vitro* tests the substance showed genetic toxicity whereas in *in vivo* test negative results were found. There are no animal data on reproductive toxicity available. From experience with humans TETA reveals no effects on reproduction. TETA is a severe irritant to skin and eyes. TETA induces skin sensitisation in guinea pigs, mice and man.

The highest aquatic local PEC during processing as an intermediate was estimated to be 4.5 µg/l.

The estimated human exposure at the workplace is estimated at < 0.143 resp. < 0.0143 mg/kg bw. Data on consumer exposure are not available.

NATURE OF FURTHER WORK RECOMMENDED

Appropriate classification and labelling are recommended.

CAS-N	CAS-NO.: 112-24-3 PROTOCOL RESULTS				
PHYSI	CAL CHEMICAL				
2.1	Melting-Point	1	NA	12 °C	
2.2	Boiling-Point		NA	ca. 280°C (at kPa)	
2.3	Density		NA	ca. 980 kg/m ³	
2.4	Vapour Pressure		NA	1.3 Pa at 20°C	
2.5	Partition Coefficient (Log Pow)		(calc.)	- 1.4	
2.6 A	Water solubility		NA	completely miscible	
В	pH		NA	10.7. at 10 g/l	
	рКа		20 °C	pKa1 = 3.32 pKa2 = 6.67	
				pKa3 =9.2 pKa4 = 9.92	
2.12	Oxidation : Reduction potential		/	mV	
	CONMENTAL FATE / CGRADATION				
3.1.1	Photodegradation		calc. (Atkinson)	In air $T_{1/2} = 1.7$ hour	
3.1.2	Stability in water		NA	no hydrolysis	
3.2	Monitoring data			In air = $/mg/m^3$ In surface water = $/\mu g/l$ In soil / sediment = $/\mu g/g$ In biota = $/\mu g/g$	
3.3	Transport and Distribution		calculated (fugacity level 1 type)	In air/ %In water/ %In sediment/ %In soil/ %In biota/ %	
3.5	Biodegradation		OECD 301 D OECD 302 B	not readily biodegradable not inherently biodegradable	

FULL SIDS SUMMARY

CAS-N	10.:112-24-3	SPECIES	PROTOCOL	RESULTS
ECOT	OXICOLOGY			
4.1	acute/prolonged toxicity to fish	Poecilia reticulata	84/449/EEC, C.1	$LC_{50} (96 \text{ hr}) = 570 \text{mg/l}$
4.2	acute/prolonged toxicity to aquatic invertebrates (daphnia)	Daphnia magna	84/449/EEC, C.2	EC ₅₀ (24hr) =31.1mg/l
4.3	toxicity to aquatic plants e. g. algae	Scenedesmus subspicatus	DIN 38412 part 9	$EC_{50} (72hr) = 2.5 mg/l$ $EC_{10} (72hr) = 0.67 mg/l$
4.4	toxicity to microorganisms	Pseudomonas fluorescens	DEV, L 8	EC_0 (24 hr) =500 mg/l
4.5.2	chronic toxicity to aquatic invertebrates (daphnia)	Daphnia magna	OECD 202 part 2	NOEC (21d) =1mg/l
(4.6.3)	mammalian terrestrial species	Agelaius phoeniceus	NA	$LD_{50} (18hr) => 101mg/kg$
	(including birds)			
TOXIC	COLOGY			
5.1.1	acute oral toxicity	rat mouse rabbit	N A N A N A	LD ₅₀ =2500 mg/kg LD ₅₀ =1600 mg/kg LD ₅₀ =5500 mg/kg
5.1.2	acute inhalation toxicity			$LC_{50} = mg/m^3$
5.1.3	acute dermal toxicity	rabbit	NA	$LD_{50} = 550 \text{ mg/kg}$
5.4	repeated dose toxicity	mouse	NA	NOAEL =92mg/kg bw
5.5	genetic toxicity in vitro			
	bacterial test (gen mutation)	S. typhimurium	Ames test	positive (with and witout metabolic activation)
	non-bacterial in vitro test (chromosomal abberations)	CHO cells		positive (with and witout metabolic activation)
5.6	genetic toxicity in vivo	mouse	Micronucleus assay	negative
5.8	toxicity to reproduction			NOEL =mg/kg (general toxicity) NOEL =mg/Kg (rep. tox. parental) NOEL =mg/Kg (rep. tox. F1)
5.9	developmental toxicity / teratogenicity			NOEL =750mg/kg (general toxicity) NOEL =750mg/Kg (pregnancy/litter) NOEL =750mg/Kg (foetal data)
5.11	experience with human exposure			

SIDS Initial Assessment Report

1.Identity

Name:	Triethylenetetramine (TE	TA)
CAS Nr.:	112-24-3	
Empirical Formula:	$C_6H_{18}N_4$	
Structural Formula:	H ₂ N-CH ₂ -CH ₂ -NH-CH	₂ -CH ₂ -NH-CH ₂ -CH ₂ -NH ₂
Purity of industrial product:	60 - 70 %	
Major impurities: N,N'-Bis-(2-aminoethyl)J N-[1-(2-Piperazin-1-yl-et Tris-(2-aminoethyl)-amin Diethylenetriamine Water	hyl)]-ethane-1,2-diamine	11 - 13 % 10 - 13 % 4 - 6 % <= 3 % <=0.5 %

2. Exposure

2.1 General discussion

Triethylenetetramine is produced by the reaction of aqueous ammonia with 1,2dichloroethane. This process yields the entire family of ethyleneamines: ethylenediamine, piperazine, diethylenetriamine, triethylenetetramine, tetraethylenepentamine, pentaethylenehexamine and aminoethylpiperazine. These polyamines are produced as their hydrochloride salts, and must be neutralized, typically with aqueous caustic soda, to obtain the free amines. The by-product salt produced in the neutralisation step is separated and the individual products are isolated by fractional distillation (8).

TETA can be used as an intermediate in a number of production processes (10):

- The reaction with polyisobutenylsuccinic anhydride yields the corresponding polybutenylsuccinimides, which are ashless, dispersant-detergent additives for motor oil.
- Polyamide -epichlorohydrin resins are produced by the reaction of epichlorohydrin with a polyamide, such as those formed by polymerisation of adipic acid and TETA. These are used in the paper industry as wet-strength additives for liner board, toweling, tissue and sanitary applications.
- The ethoxylated products of TETA are curing agents for epoxy resins. The largest application is surface coatings (35%).
- Imidazolines from the condensation of TETA with two moles of fatty acid are cationic surfactants used as fabric softeners, asphalt emulsifiers, oil field corrosion inhibitors, ore flotation agents and epoxy curing agents.
- Reactive polyamides from the polymerisation of dimer acids with TETA are mostly used as curing agents for epoxy surface coatings.

In 1989 - 1991, 1200 - 1500 t/a were produced in Germany. Production capacities as of 1990 for other countries are available as well (8):

Netherlands	ca. 6000 t/a	(2 sites)
USA	> 11000 t/a	(3 sites)
Japan	ca. 1800 t/a	(1 site)

According to the German producer, ca. 40 to 50% are sold in Germany (> 10 clients) and ca. 40 - 50 % are exported; the rest is further processed by the same producer. Import volumes are estimated by the producer at ca. 1500 t/a. The total consumption in Germany amounts to ca. 2200 t/a.

In Germany, triethylenetetramine (TETA) is mainly used as

- intermediate for curing agents for epoxy resins (ca.1600 t/a)
- direct curing agent for epoxy resins (ca. 160 t/a)
- intermediary for auxiliary agents used in the paper industry, the textile industry and in glues (ca. 330 t/a)
- intermediate for asphalt emulsifiers (ca. 110 t/a)

Ca. 100 t/a are used by the producer as an intermediate. No information is available on the processing at other chemical manufacturers.

In Sweden, the use pattern of TETA is similar to the use pattern described for Germany:

- intermediate for transport, fertilizer and plastics industry (200 533 t/a)
- adhesive, binding agent (4 6 t/a)
- hardener for plastic (1 4 t/a)
- others (max 5 t/a)

The use pattern for other countries is not available.

2.2 Environmental e xposure

2.2.1 General/Environmental fate

TETA is completely miscible with water forming an alkaline solution (pH 10 at 10 g/l). The technical product has a vapour pressure of ca. 1 Pa at 20 °C. The calculated Log Pow (unprotonated form) amounts to ca. -1.4 and indicates a low potential for bioaccumulation. There are no measured Koc-values available. For ethylenediamine (CAS Nr. 107-15-3) and diethylenetriamine (CAS Nr. 111-40-0), Koc-values of 4766 and 19111 were measured respectively (1). The high adsorption is most likely due to electrostatic interaction. A comparable Koc can be expected for TETA, which would suggest a high potential for geoaccumulation.

Based on the physical-chemical properties the target compartment of TETA in the environment is the hydrosphere (the estimation of the distribution with a Fugacity model is not opportune due to the protophile behaviour of TETA).

TETA is not readily biodegradable (0% after 20 days, OECD GL 301 D; same result with adapted inoculum). Also, in a test on inherent biodegradability with industrial sludge, TETA was not degraded (0 % DOC removal after 28 days, OECD GL 302 B). TETA has therefore to be regarded as **non biodegradable**. Adsorption onto sewage sludge was not observed.

In a test on hydrolysis, TETA was not found to have undergone hydrolysis after 36 days.

Direct photolysis of TETA in the hydrosphere is not to be expected (molar extinction coefficient $< 10 \text{ l} / (\text{mol} \cdot \text{cm}) \text{ at} > 240 \text{ nm}$). The half-life due to photooxidative degradation by OH-radicals in the atmosphere is estimated to be 1.7 hours. As TETA does have a low tendency to pass from water to air, this does not represent a significant removal process from the environment.

Based upon the physical-chemical and biodegradation properties of TETA, no elimination in waste water treatment plants is assumed.

2.2.2 Exposure assessment

a) Local concentrations

Considering the above described use pattern, point releases are to be expected during production and processing.

production

According to the German producer, no continuous releases occur during the production process to waste water. During cleaning operations of the production facility and the distillation column, the releases are estimated by the German producer at ca. 1 g/t related to the production capacity (8). For a production capacity of 5000 t/a (worst case assumption) a release of 5000 g TETA during one day (assuming one cleaning operation per year) can be estimated. Assuming no elimination in the WWTP, 5000 g are released into a river with a flow of 60 m³/s, according to the generic release scenario for production in (3). A **PEC**_{local} of 1 µg/l is calculated.

processing

Many processes involving TETA as intermediate with different release rates are to be expected.

Specific data are available only from one German producer, using ca. 100 t TETA per year for processing with fatty acids: a maximum of 2.4 kg/a are released to the waste water (8).

For a generic estimation, the following worst case situation according to the release scenario for intermediates described in (3) is used.

For a processing site using 1000 t/a of TETA, a release factor of 0.7 % is assumed. Considering no elimination in the WWTP, 7 t/a are released into a river with a flow of 60 m³/s. Assuming release over 300 days per year, a concentration of $PEC_{local} = 4.5 \ \mu g/l$ is calculated.

b. <u>Regional concentrations</u>

Diffuse release into the environment would occur through the direct use of TETA as a curing agent. Also, the curing agents produced from TETA contain residual concentrations of TETA (approx. 7.9%).

The final extent of conversion of TETA during curing reactions is not known. On the other hand, the conversion of diethylenetriamine was determined to be 60 to 80 % (2) (related to the total NH-functions). As TETA presents 6 NH-functions, a molecular conversion rate of > 90% can be assumed.

About 160 t/a of TETA are used directly as curing agent. With a conversion factor of 90%, ca. 16 t are available as free molecules in the resins. On the worst case assumption , that 10% are released through migration from the matrix (3), a maximum of 1.6 t/a are released into the environment through this path.

About 1600 t/a are processed to yield curing agents containing an average of 7.9% free TETA. For a rough estimate, it is assumed that TETA reacts with the same amount of chemicals so that 3200 t of curing agents with ca. 250 t of free TETA result. Of these, max. 10% (see above) remain unreacted in the curing process and 10 % of these may be released through migration, i.e. a maximum of 2.5 t/a.

For the calculation of the regional PEC the use of a fugacity model is not opportune due to the ionic nature of TETA. The regional concentration can be estimated in a first approach with the following formula (9):

EMIS

PEC_{regional} = —

 $FLOW + V \cdot k$

- with: EMIS: emission into surface water = 1.6 + 2.5 = 4.1 t/a
 - FLOW: flow through the water compartment
 - V: Volume of water compartment
 - k: first order biodegradation rate constant

The default values described in (3) will be used for the calculation:

- a small but densely populated area is considered: 200x200 km with 20 million inhabitants;
- with an area fraction of water of 0.02 and a mixing depth of 3 m, V = $2.4 \cdot 10^9 \text{ m}^3$
- with an average residence time of the water of 40 days, FLOW = $6 \cdot 10^7 \text{ m}^3/\text{d}$
- TETA being non-biodegradable, k = 0

=> PEC_{regional} = 0.18 µg/l

2.3 Consumer exposure

Where epoxy resins are cured in do-it-yourself applications (e.g. in coatings, adhesives, and epoxy-fiber composites), consumers may come into contact with TETA or TETAderived curing agents, either when mixing the ingredients, or when grinding and polishing the solidified product whereby unreacted TETA may be set free.

2.4 Occupational exposure

The production unit simultaneously produces ethylenediamine, diethylenetriamine, triethylenetetramine and other substances from ammonia and 1,2-dichloroethane.

To date, exposure to triethylenetretamine (TETA) has not been measured directly. Instead, exposure is estimated on the basis of measurements of ethylenediamine (according to TRGS 402) - the end product with the lowest boiling point.

The MAK-value of 25 mg/m³ for ethylenediamine is consistently met. All measurements indicate that exposure is below 1 mg/m^3 .

Substance	Boiling Point	Vapour Pressure
Ethylenediamine	116.5 °C	12.1 hPa
TETA	approx. 280 °C	< 0.1 hPa

Due to ethylenediamine's significantly lower boiling point and its greater vapour pressure (by a factor of 100) it can be concluded with certainty that the concentration of TETA in the air during synthesis and processing does not exceed 0.1 mg/m^3 .

Exposure is, therefore, clearly below the actual occupational exposure limit of 6 mg/m³ in Sweden.

3. Toxicity

3.1 Human Toxicity

a) <u>Acute Toxicity</u>

Triethylene tetramine is of low acute toxicity on oral administration (LD₅₀ rat > 2000 mg/lkg bw) and moderate toxicity on dermal application (LD₅₀ rabbit 550-805 mg/kg bw). Exposition to saturated vapour was tolerated without impairment whereas the exposition to aerosol leads to reversible irritations of the mucous membranes in the respiratory tract. According to EC Directive 67/584/EEC triethylene tetramine is labelled as harmful in contact with skin (R 21).

Conclusion: Moderate acute toxicity Priority setting: low priority or concern

b) <u>Repeated Dose Toxicity</u>

In a subacute study (rat, oral, up to 2980 mg/kg bw) retarded body weight gain and elevated liver and kidney weights were observed in the highest dose groups. From this study, a NOAEL of 500 mg/kg was derived.

In a subacute study, undiluted test substance was rubbed into the skin of pregnant and non-pregnant guinea pigs (4 mg/guinea pig and day = ca. 9 mg/kg bw) daily for 55 days. In the course of the experiment the death of test animals (2/9) as well as of the control animals (6/11) occurred (11). In another study, dermal application to pregnant and non-pregnant guinea pigs (4 mg/animal = ca. 9 mg/kg bw) daily for the first 10 days and every second day for next 45 days resulted in reduced weight gain, and from the 5th day of treatment in inflammatory alterations at the application site with subsequent erosions. In the course of the experiment 7/11 pregnant and 7/11 non-pregnant animals died (12). It is unclear whether the death of the animals is due to the strong irritant and/or the skin sensitization potential of the test substance.

In an additional study F344 rats and B6C3F1 mice received triethylenetetramine dihydrochloride in the drinking water at concentrations of 0, 120, 600, 3000 ppm (target concentration) for up to 92 days. Each dose group were fed either cereal based (NIH-31) or purified (AIN-76A) diet both containing nutritionally adequate levels of copper. An additional control group of rats and mice received a Cu-deficient AIN-76A diet. Sign of triethylenetetramine dihydrochloride toxicity were noted only in B6C3F1 mice fed AIN-76A diet given 3000 ppm triethylenetetramine dihydrochloride. These toxic signs included inflamation of the lung interstitium, hemapoetic cell proliferation of the spleen, liver periportal fatty infiltration, kidney weight reduction, reduced renal cytoplasmatic vacuolization and body weight gain reduction. From this study a NOAEL of 600 ppm for mice was derived. According to the authors, the signs observed in F344 rats appear to be related to copper deficiency (13).

Lifelong dermal application to mice (1.2 mg/mouse and application) caused no skin tumours or any tumours.

In a former inhalation study with rats, mice, guinea pig and rabbit (aerosol: 0.4 ml in 0.5 ml ethanol in a 4001 chamber, 10 d), no irritations or other toxic effects were observed.

Conclusion:

Signs of impairment only in mice following subchronic oral dosing of 3000 ppm triethylenetetramine dihydrochloride. NOAEL: 600 ppm [92 (male), 99 (female) mg/kg bw].

Priority setting low priority or concern

c) Reproductive/Developmental Toxicity

In rabbits, triethylene tetramine does not cause embryotoxic and teratogenic effects, even at maternally toxic dose levels (4).

In rats, there are several studies concerning developmental toxicity. The oral treatment of rats with 75, 375 and 750 mg/kg resulted in no effects on dams and fetuses, except slight increased fetal body weight (5). After oral treatment of rats with 830 or 1670 mg/kg bw only in the highest dose group increased fetal abnormalities in 27/44 fetus (69,2%) were recorded, when simultanously the copper content of the feed was reduced. Copper-supplementation in the feed reduced significant the fetal abnormalities of the highest dose group to 3/51 (6,5% fetus. These findings suggest that the developmental toxicity is produced as a secondary consequence of the chelating properties of triethylene tetramine (6).

In chapter 3.1.b) 2 studies on pregnant guinea pigs dermally treated with 4 mg/animal = ca. 9 mg/kg bw daily for 55 days or daily for 10 days and every second day for the next 45 days, respectively, were described (11, 12). Beside the clear mortality rate and the local effects, necrotic changes of the placenta and miscarriage or mortification of the fetuses and stillbirth of malformed fetuses were observed. Due to the clear maternal toxicity and due to the lack of dose-response relationship the reported studies are not suitable to evaluate developmental toxicity.

There are no data on effects on fertility with triethylene tetramine .In the subchronic toxicity studies with mice and rats, which were described in chapter 3.1.b, the reproductive organs are examined. In mice, there were no treatment related effects on the reproductive organs. According to the authors the only finding which may be attributable to trien-2HCl occured in AIN-76A-fed females rats. There was a significant dose-related trend toward an increased prevalence of uterine dilatation (13). There are no changes of the vagina and the ovaries. Therefore dilatation of uterus in isolation cannot be regarded as hormonal effects. Thus, this finding is not suitable to evaluate any reproductive toxicity. In addition, oral treatment of rats with the analogue diethylene triamine caused no adverse effects respective mating index, fertility index and number of live and dead pups.

Triethylene tetramine is used in the therapy of Wilsons' disease. While taking 400 to 800 mg triethylene tetramine 3 times a day for about 120 months, there have been six pregnancies in four female patients. There were no miscarriages and no fetal abnormalities. All six children developed normally (7).

Conclusion:

From experiences with humans (substance given as a drug) there is no reason to assume that the substance reveals effects on reproduction.

Priority setting: low priority or concern

d) Genetic Toxicity

The results of the genetic toxicity testing are not uniform. In vitro, triethylene tetramine has clear genotoxic activity in the Ames-test and in mammalian cytogenetic tests. Whereas in vivo, triethylene tetramine is not clastogenic in the mouse micronucleus test following intraperitonal injections of 130 to 600 mg/kg bw. The study was conducted in accordance with GLP standards. In addition, there is a further micronucleus test using oral application (14) which yielded a negative result as well. In this study, mice received once 1500, 3000 and 6000 mg/kg bw. These doses are within the range of and/or greater than the LD50 value for mice, which is cited in the basic data set: LD50(mice) = 1600 mg/kg bw (15). The test design and test performance was carried out according to W. Schmid and coworkers who developed the test (see references).

Following 1500 and 3000 mg/kg bw the percentage of erythrocytes containing micronuclei corresponds with the percentage of those in the concurrent solvent control. Following 6000 mg/kg bw a decrease in erythrocytes containing micronuclei was noted and was thus lower than those in the concurrent solvent control.

Triethylene tetramine revealed no mutagenic activity in the SLRL test in Drosophila melanogaster.

Conclusion:

As triethylene tetramine revealed no mutagenic activity in relevant in-vivo tests there is no reason to assume genotoxicity.

Priority setting: low priority or concern

e) Sensitizatio

The sensitization potency of triethylene tetramine was investigated in the Guinea Pig Maximization Test (GPMT) and in the Mouse Ear Swelling Test (MEST).

One of the GPMTs (16) used triethylene tetramine as a commercial product (no further information on purity of the substance). The method used was in accordance with the original description of the GPMT by Magnusson and Kligman (20, 21). Control animals received vehicle only. Induction concentration was 0.5 % in water and challenge concentration was 2 %. 12/15 animals (80 %) showed positive reactions 24 hours after removal af the patch. In the second GPM test, carried out according to OECD Guideline 406, purified TETA (purity: 99.5 %) was used and the applied concentrations were for induction 0.5 % and for challenge 2 % as well. As positive control served dinitrochlorobenzene. 9/10 animals (90 %) showed positive reactions (17). As additional test, the MEST was performed with 10 mice (17). The concentration of the purified TETA (purity: 99.5%) for the induction procedure was 10 % and the challenge concentration was 2.5 %. Oxazolone served as positive control. In 4/10 mice positive reactions were seen.

Cross reactions between triethylene tetramine, ethylenediamine and diethylenetriamine were also observed in guinea pigs (18).

Numerous reports concern the sensitizing potential of triethylene tetramine in humans (18).

In Poland, 20 - 51.2 % out of 20 - 447 examined workers exposed to epoxy resins reacted positive to triethylene tetramine (19). At another factory dermatitis was observed in 126 out of 422 workers. Skin tests were carried out on 99 patients. A positive reaction was observed in 55.1 % of these cases (18). In an examination of 20 workers exposed to casting resins and triethylene tetramine 5 showed positive reaction to triethylene tetramine whereas in another group of 23 epoxy resin-workers, suffering from dermatitis, none

reacted positive on a patch test with triethylene tetramine (18). In a control group of 112 persons 2 persons (1.5 %) gave positive patch test results (18).

Cross reactions between triethylene tetramine, diethylenetriamine and ethylenediamine were also reported (18).

Conclusion:

Triethylene tetramine induces skin sensitization in guinea pigs, mice and man. According to EC Directive 67/584/EEC triethelyene tetramine is labelled: R 43 = may cause sensitization by skin contact.

3.2 Ecotoxicity

3.2.1 Aquatic organisms

a) <u>Toxicity to fish</u>

Other test results with *Leuciscus idus* and *Pimephales promelas*, which could not be validated, are in the same order of magnitude.

b) <u>Toxicity to invertebrates</u>

Daphnia magna	48h-EC ₅₀	31.1 - 33.9 mg/l	
(several tests)			
Effect: immobilisation	21d-EC ₅₀	> 3.2 - < 10 mg/l	
	21d-NOEC	1 mg/l	
(immobilisation of parental organisms was the most sensitive effect parameter)			

Furthermore, concentrations of 293 - 7313 mg/l had no teratogenic effects on sea-urchin (*Paracen trotus lividus*) eggs. The larvae were most sensitive and showed delay of development at 293 mg/l

c) Toxicity to algae

Scenedesmus subspicatus	$72h-E_BC_{50}$	2.5 mg/l
	$72h-E_BC_{10}$	0.67 mg/l
	$72h-E_{\mu}C_{50}$	>= 100 mg/l
	$72h-E_{\mu}C_{10}$	0.95 mg/l
Effects enough inhibition (D	hismesses susveth uses)	

Effect: growth inhibition (B = biomass; μ = growth rate)

Due to the intensive growth of the algae the pH in the control and in the concentrations up to 1 mg/l increased within 72 h to 10.2 - 10.3.

Selenastrum capricornutum	72h-EC ₅₀	20 mg/l
Effect: growth inhibition (biomass)	72h-NOEC	< 2.5 mg/l
Selenastrum capricornutum	96h-EC ₅₀	3,7 mg/l

Effect: growth inhibition (biomass)

A further test with Chlorella pyrenoidosa was considered to be non valid.

d) Toxicity to microorganisms

Pseudomonas fluorescens24h-EC0500 mg/lEffect: growth inhibition (biomass)

e) Derivation of PNEC

Algae are clearly the most sensitive species to TETA. According to the EU-Technical Guidance Document (3), the value of the safety factor is $\mathbf{F} = 50$ (long term tests have been performed for two trophic levels and with the organisms which were the most sensitive in the acute tests).

With the lowest aquatic effect concentration of 0.67 mg/l:

$$PNEC = \frac{670}{50} = 13.4 \ \mu g/l$$

3.2.2 Terrestrial organisms

Acute oral toxicity to the redwinged blackbird (*Agelaius phoeniceus*) was determined to be $18h-LD_{50} > 101 \text{ mg/kg bw}$.

4. Initial Assessment

4.1 Human toxicity

4.1.1 Identification of critical toxic effects

Triethylene tetramine is a severe irritant to skin and eyes and induces skin sensitizations. Triethylene tetramine is of moderate acute toxicity: LD50(oral, rat) > 2000 mg/kg bw, LD50(dermal, rabbit) = 550 - 805 mg/kg bw. Acute exposure to saturated vapour via inhalation was tolerated without impairment.

Following repeated oral dosing via drinking water only in mice but not in rats at concentration of 3000 ppm there were signs of impairment. The NOAEL is 600 ppm [92 mg/kg bw (oral, 90 days)]. Lifelong dermal application to mice (1.2 mg/mouse) did not result in tumour formation.

There are differing results of the genetic toxicity for triethylene tetramine. The positive results of the in vitro tests may be the result of a direct genetic action as well as a result of an interference with essential metal ions. Due to this uncertainty of the in vitro tests, the genetic toxicity of triethylene tetramine has to be assessed on the basis of in vivo tests. The in vivo micronucleus tests (i.p. and oral) and the SLRL test showed negative results.

There are no data on reproductive toxicity (fertility assessment). The analogue diethylene triamine had no effects on reproduction. Triethylene tetramine shows developmental toxicity in animal studies if the chelating property of the substance is effective. The NOEL is 830 mg/kg bw (oral).

Experience with female patients suffering from Wilson's disease demonstrated that no miscarriages and no fetal abnormalities occur during treatment with triethylene tetramine.

4.1.2 Comparison of Exposure and Critical effects

Workplace

There are no measurements of the concentration of triethylene tetramine in the air at the workplace. To estimate the exposition at the workplace adequately the results of the concentration measurements of the product with the lowest boiling point has to be applied: ethylene diamine (see chapter 4.2). All results of these measurements are below 1 mg/m³ (TLV: 25 mg/m³). Because of the higher boiling point and the lower vapour pressure of triethylene tetramine it can be assumed that the concentration in the air at the workplace is below or equal than 0.1 mg/m³.

The EHE (Estimated Human Exposure) can be calculated according to the following equitation:

 $EHE = \frac{\text{respiratory rate (10 m^3) * exposition (mg/m^3)}}{\text{body weight (70 kg)}}$

 $\begin{array}{l} exposition < 1 \ mg/m^3 \\ exposition < 0.1 \ mg/m^3 \end{array}$

EHE < 0.143 mg/kg bwEHE < 0.0143 mg/kg bw

Thus the estimated human exposure is far below the NOAEL described in animal experiments of 92 mg/kg bw for subacute toxicity and a NOAEL of 850 mg/kg bw for teratogenicity. The safety margin based on the lowest NOAEL is between:

 92 mg/kg bw 92 mg/kg bw

 ----- > 643.4

 < 0.143 mg/kg bw < 0.0143 mg/kg bw

and thus does not suggest a particular risk.

Isolated cases of exposure through skin contact cannot be ruled out. However, the risk is to be assumed very low.

Consumer area

Data on consumer exposure are not available. However, it cannot be excluded that products containing triethylene tetramine give off small amounts of the substance. Due to the low toxicity in animal experiments it can be assumed that the probability of acute poisoning is very low. In addition, the application of triethylene tetramine as drug excluded high toxicity to humans. Also multiple administration of TETA to animals did cause neither significant systemic effects nor the formation of tumours.

Exposure via the environment

Data are not available on exposure of the general population. Exposure of the population via the hydrosphere is considered to be minimal, even assuming the concentration in drinking water to be equal to the regional predicted concentration in surface waters (0.18 μ g/l). With 2 l drinking water/person/da y, the daily dose would be 0.005 μ g/kg bw/day. Compared to the exposure at the working place the exposure through the environment is negligible.

4.2 Assessment of environmental hazards

In the following table, the PEC/PNEC ratios for the different exposure scenarios are presented:

Scenario	PEC _{local} + PEC _{regional}	PEC/PNE
	[µg/l]	С
production (site)	1 + 0.18	0.08
processing (site)	4.5 + 0.18	0.35

A PEC/PNEC < 1 in all scenarios, a low potential risk to the aquatic compartment is at present to be expected.

A significant exposure to the **terrestrial** compartment could not be identified. Further work is presently not necessary for an assessment of risks to this compartment.

5. Conclusions and Recommendations

An environmental hazard assessment of triethyle netetramine was possible with the available data and showed that the compound was presently of low concern to the environment. No further work is recommended.

On the basis of the known facts and properties, triethylene tetramine may represent a hazard for human health. The chemical is a severe irritant to skin and eyes and induces skin sensitization. The substance is classified and labelled accordingly within the EU: R 34 =causes burns; R 43 = may cause sensitization by skin contact.

From experience with humans (substance given as a drug) there is no reason to assume that the substance reveals further toxic effects. Besides appropriate classification and labelling no further work is recommended.

References

- 1 OECD SIDS Diethylenetriamine (111-40-0), draft from April 1994
- 2. Kamon, T. and Saito, K., Kobunshi ronbunshu 41 (1984), H. 5, 293-299
- 3. Technical Guidance Documents in support of the Commissions Directive 93/67/EEC on Risk Assessment for New Notified Substances and the Commission Regulation (EC) 1488/94 on Risk Assessment for Existing Substances, 1996
- 4. Union Carbide Corporation, Bushy Run Research Center, Project Report 50-127 (1988) cited in: BG Chemie, Toxikologische Bewertung Nr. 181, 1991
- 5. Ciba Geigy Ltd, Report on TK10458, Report No. 830035 (1984) cited in BG Chemie, Toxikologische Bewertung Nr. 181, 1991
- 6. Cohen N. L. et al., DrugNutr. Interact. 2, 203-210 (1983)
- 7. Walshe J. M., The Lancet 1, 643-647 (1982)
- 8. BUA-Report 89: Triethylentetramin; Juni 1992; ISSN 0179-2601
- 9. van den Meent: Simplebox: a generic multimedia evaluation model; RIVM report no. 672720001; (1993)
- 10. Wm. K. Johnson: CEH Product Review "Ethyleneamines", SRI 1989
- 11. Dobryszycka W. et al., Archivum Immun. Therap. Exp. 23, 867-870 (1975)
- 12. Szacki J. et al., Archivum Immun. Therap. Exp. 22, 123-128 (1974)
- 13. Greenman D.L. et al., Fundam. Appl. Toxicol. 29, 185-193 (1996)
- 14. Heinz N., Schroeder H.F., Drug Res. 31, 950-953 (1981)
- 15.Stavreva M., Khig. Zdraveopaz. 22, 179-182 (1979)
- 16. Thorgeirsson A., Acta Derm. (Stockholm) 58, 332-336 (1978)
- 17. Maisey J. et al., contact Dermatitis 18, 133-137 (1988)
- 18. BG: Berufsgenossenschaft der chemischen Industrie, Toxikologische Bewertung triethylene tetramine No. 18, 1991
- 19. Ruddzki E., contact dermatitis 6, 235-236 (1980)
- 20. Magnusson B., Kligman A.M., J. Invest. Dermatol. 52, 586 (1969)
- 21. Magnusson B., Kligman A.M., cited in: Identification of contact Allergens, Ch.C. thomas Publisher, Springfield, Ill., 1970

IUCLID Data Set

Existing Chemical ID: 112-24-3 112 - 24 - 3CAS No. trientine EINECS Name EC No. 203-950-6 TSCA Name 1,2-Ethanediamine, N,N'-bis(2-aminoethyl)-Molecular Formula C6H18N4 Producer Related Part Bayer AG Company: Creation date: 15-MAR-1993 Substance Related Part Bayer AG Company: Creation date: 15-MAR-1993 Memo: AKTUELL OECD-SIDS Printing date: 24-JUL-2002 Revision date: 17-MAY-1993 Date of last Update: 27-JAN-1998 Number of Pages: 56 Chapter: 1, 2, 3, 4, 5, 6, 7, 8, 10 Chapter (profile): Reliability: without reliability, 1, 2, 3, 4 Reliability (profile): Flags (profile): Flags:without flag, confidential, non confidential, WGK (DE), TA-Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. GENERAL INFORMATION

1.0.1 Applicant and Company Information

Туре:	cooperating company	У
Name:	Bayer AG	
Town:	51368 Leverkusen 1	
Country:	Germany	

10-MAY-1994

1.0.2 Location of Production Site, Importer or Formulator

1.0.3 Identity of Recipients

1.0.4 Details on Category/Template

1.1.0 Substance Identification

1.1.1 General Substance Information

Substance type:	organic
Physical status:	liquid
Purity:	60 - 70 % w/w

Remark: technical mixture

1.1.2 Spectra

1.2 Synonyms and Tradenames

1,2-Bis-(2-aminoethylamino)-ethan

1,2-Di-(aminoethylamino)-ethan

1,4,7,10-Tetraazadecan

1,8-Diamino-3,6-diaza-octan

2,2'-(1.2-Ethylenbis-amino-)bis-ethanamin

3,6-Diazaoctan-1,8-diamin

N,N'-Bis-(2-aminoethyl)-1,2-ethanediamine

N,N'-Bis-(2-aminoethyl)-ethylendiamin

N,N'-Di-(2-aminoethyl)-1.2-ethandiamin

N,N'-Di-(2-aminoethyl)-1.2-ethylendiamin

TETA

Tetramin

Trien

Triethylentetramin

SUBSTANCE ID: 112-24-3

1. GENERAL INFORMATION

1.3 Impurities

EINECS-Name:	N,N¦-Bis-(2-aminoethyl)piperazin
Contents:	11 - 13 % w/w
EINECS-Name:	N-(Piperazin-1-ethyl)-ethan-1,2-diamin
Contents:	10 - 13 % w/w
EINECS-Name:	Tris-(2-aminoethyl)-amin
Contents:	4 - 6 % w/w
CAS-No:	111-40-0
EC-No:	203-865-4
EINECS-Name:	2,2'-iminodi(ethylamine)
Contents:	<= 3 - % w/w
EINECS-Name:	Water
Contents:	<= ,5 - % w/w

1.4 Additives

1.5 Total Quantity

Quantity:	1000 - 5000 tonnes produced	
Remark: 29-NOV-1994	in 1989-1991 (BRD)	(1)
Remark:	Netherland: ca. 6000 t/a USA: ca. 1100 t/a	
29-NOV-1994	Japan: ca. 1800 t/a	(1)

1.6.1 Labelling

Labelling: Symbols: R-Phrases:	(C) ((21) (34)	n Directive 67/548/EEC corrosive Harmful in contact with skin Causes burns May cause sensitization by skin contact
S-Phrases:		In case of contact with eyes, rinse immediately with plenty of water and seek medical advice Wear suitable protective clothing, gloves and eye/face protection
Country:		Germany

Country:

1.6.2 Classification

Classified: Class of danger:	as in Directive 67/548/EEC corrosive
R-Phrases:	(21) Harmful in contact with skin (34) Causes burns (43) May cause sensitization by skin contact
Country:	Germany

1.6.3 Packaging

1.7 Use Pattern

Туре:	industrial
Category:	Chemical industry: used in synthesis
Remark:	Intermediate for - hardeners for epoxy resins > 80 % - agents used in glues, paper industry and textile industry > 15 %
Туре:	use

Remark: TETA can also be used directly as hardener in epoxy resins (approx. 8 % of total production)

1.7.1 Detailed Use Pattern

1.7.2 Methods of Manufacture

1.8 Regulatory Measures

1.8.1 Occupational Exposure Limit Values

1.8.2 Acceptable Residues Levels

1.8.3 Water Pollution

Classified by: other: Bayer AG Labelled by: other: Bayer AG Class of danger: 2 (water polluting) Country: Germany

1.8.4 Major Accident Hazards

Substance listed: no

1.8.5 Air Pollution

Classified by: TA-Luft (DE) Labelled by: TA-Luft (DE) Number: 3.1.7 (organic substances) Class of danger: III

1.8.6 Listings e.g. Chemical Inventories

1.9.1 Degradation/Transformation Products

1.9.2 Components

1.10 Source of Exposure

Country: Germany

1. GENERAL INFORMATION

DATE: 24-JUL.-2002 SUBSTANCE ID: 112-24-3

Remark:	air: 6 kg/a at one processing site; no release into the atmosphere at all other production and processing sites
	water: 4,4 kg/a at all production and processing sites
	waste treatment:
	water: biological waste water treatment plant
	air: incineration
	There is no solid waste from production and processing.
	Possible emissionof very small amounts through migration out
	of epoxy resins (residual concentration of TETA in
	hardeners: at max. approx. 7.9 %)
29-NOV-1994	(1)

1.11 Additional Remarks

1.12 Last Literature Search

1.13 Reviews

2.PHYSICO-CHEMICAL DATA

2.1 Melting Point

Value:	= 12 degree C (2)
Remark: 26-APR-1994	Solidification point: approx35 degree C (technical product) (3)

2.2 Boiling Point

Value:	266 - 267 degree C	(4)
Value: Decomposition:	= 277,5 degree C yes	(1)
Remark:	93 - 96 % purity	(5)
Value:	= 277,9 degree C	(6)
Value: Decomposition:	= 278 degree C yes	
		(7)
Value:	ca. 280 degree C	
Remark: 26-APR-1994	technical product	(3)
2.3 Density		

Type: Value:	density = ,9739 g/cm³ at 20 degree C	(8)
Type: Value:	density ca. ,98 g/cm³ at 20 degree C	

Remark: technical product 26-APR-1994

Type:	density	
Value:	= ,9818 g/cm³ at 20 degree C	
		(5)
Type:	density	
Value:	= ,9839 g/cm³ at 20 degree C	

(6) **Type:** density **Value:** = ,977 g/cm³ at 25 degree C

2.3.1 Granulometry

(3)

(9)

2.PHYSICO-CHEMICAL DATA

DATE: 24-JUL2002
SUBSTANCE ID: 112-24-3

2.4 Vapour Pressure

Value:	= ,013 hPa at 20 degree C	
Value:	< ,1 hPa at 20 degree C	(6)
Remark: 26-APR-1994	technical product	(3)

2.5 Partition Coefficient

log Pow:	= -1,66	
Remark:	calculated (no further information)	
log Pow:	= -1,41 (10	')
Remark:	calculated (no further information)	\
log Pow:	= -1,4	.)
Method:	other (calculated): Leo, Hansch: A. Leo, CLOGP-3.63 (1991) Daylight, Chemical Information Systems, Inc. Irvine, CA, USA	
Remark:	undissociated form (12	2)

2.6.1 Solubility in different media

Remark:	completely miscible	
		(7)

2.6.2 Surface Tension

2.7 Flash Point

Value:	= 118 degree C	(13)
Value:	= 125 degree C	(6)
Value: Method: Remark: 26-APR-1994	ca. 129 degree C other: DIN 51758 technical product	(3)
Value:	= 135 degree C	(5)

2.8 Auto Flammability

2.PHYSICO-CHEMICAL DATA

2.9 Flammability

Remark:	LFL: 1.0 % v/v (180 deg. C)
	UFL: 3.6 % v/v (180 deg. C)
Source:	DOW Europe S.A., Switzerland
24-MAY-1994	

(14)

2.10 Explosive Properties

2.11 Oxidizing Properties

2.12 Dissociation Constant

2.13 Viscosity

2.14 Additional Remarks

Remark:	Henry-constant : 6.7x10E-11 Pa.m3/mol (at 25 degree C, calculated)	
29-NOV-1994		(1)
Remark: 26-APR-1994	Ignition-temperature : 335 Grad C (DIN 51794)	(3)
Remark:	Ignition-temperature : 338 Grad C	(5)
Remark:	UV-Spectrum in water : epsilon < 10 e/molxcm at lamda > 24	0 nm (15)

SUBSTANCE ID: 112-24-3

3. ENVIRONMENTAL FATE AND PATHWAYS

3.1.1 Photodegradation

<u>stiti instadgiu</u>	
Type: INDIRECT PHOTOLY: Sensitizer: Rate constant: Degradation:	OH ,00000000225 cm³/(molecule * sec)
Method:	other (calculated): according to Atkinson
29-NOV-1994	(16) (1)
3.1.2 Stability	in Water
Туре:	abiotic
Year: Test substance:	1985 other TS: technical grade (purity > 70 %)
Remark:	No hydrolysis in water during the experiment of 36 days. Tested concentrations: 1, 100 and 200 mg/l (17)
3.1.3 Stability	in Soil
3.2.1 Monitoring	Data (Environment)
3.2.2 Field Stud	lies
3.3.1 Transport	between Environmental Compartments
Remark:	Based on the physico-chemical properties transport from water to air is not to be expected (Henry-constant: H = 6.7 x 10E-11 Pa.m3/mol, 25 degree C, calculated)
29-NOV-1994	(1)
3.3.2 Distributio	
Remark:	Based on the physical-chemical data, the preferred environmental compartment of TETA is the hydrosphere
3.4 Mode of Degr	adation in Actual Use
3.5 Biodegradat:	on
Type: Inoculum: Concentration: Degradation: Result:	aerobic activated sludge, industrial 100 mg/l related to DOC (Dissolved Organic Carbon) 0 % after 28 day(s) under test conditions no biodegradation observed
Method:	OECD Guide-line 302 B "Inherent biodegradability: Modified Zahn-Wellens Test"

SUBSTANCE ID: 112-24-3

3. ENVIRONMENTAL FATE AND PATHWAYS

Year:	1989	
GLP:	no data	
Remark:	technical product	
		(18)
Type:	aerobic	
Inoculum:	predominantly domestic sewage, adapted	
Concentration:	related to Test substance	
Degradation:	0 % after 20 day(s)	
Result:	under test conditions no biodegradation observed	
Method:	other: in accordance with OECD Guide-line 301 D "Ready	
	Biodegradability: Closed Bottle Test"	
Year:	1977	
GLP:	no data	
Remark:	technical product;	
	Substance concentrations: 2.6, 8.5, 25.5, 85 mg/l	
		(18)
		/

3.6 BOD5, COD or BOD5/COD Ratio

3.7 Bioaccumulation

Remark: Bioaccumulation is not to be expected (logPow = -1,4; -1.66 calculated)

3.8 Additional Remarks

4. ECOTOXICITY

AQUATIC ORGANISMS

```
4.1 Acute/Prolonged Toxicity to Fish
```

Type: Species: Exposure period: Unit: LC0: LC50: LC100:	semistatic Poecilia reticulata (Fish, fresh water) 96 hour(s) mg/l Analytical monitoring: no 180 - 570 - 1800 -	
Method: Year: GLP: Test substance:	Directive 84/449/EEC, C.1 "Acute toxicity for fish" 1989 yes other TS: Triethylenetetramine, purity: 97.5%	
Remark: 10-MAY-1994	48h-LC50 = 1140 mg/l	(19)
Species: Exposure period: Unit: LC0:	Leuciscus idus (Fish, fresh water) 48 hour(s) mg/l Analytical monitoring: 200 -	
Method: GLP:	other: Bestimmung der akuten Wirkung von Stoffen auf Fisch Arbeitskreis "Fischtest" im Hauptausschuss "Detergentien" (15.10.73) no	le.
Remark:	open system; at 500 mg/l, all test organisms had died after 27 h; no further information on test conditions	(18)
Species: Exposure period: Unit: LC50:	Pimephales promelas (Fish, fresh water) 96 hour(s) mg/l Analytical monitoring: 495 -	
Remark: Source: 26-APR-1995	validation not possible DOW Europe S.A., Switzerland	(20)
4.2 Acute Toxicit	y to Aquatic Invertebrates	

Species: Exposure period:	Daphnia magna (Crustacea) 48 hour(s)
Unit:	mg/l Analytical monitoring: no
EC0:	18 -
EC50:	31,1 -
EC100:	56 -
Method:	Directive 84/449/EEC, C.2 "Acute toxicity for Daphnia"
Year:	1989
GLP:	yes
Test substance:	other TS: Triethylentetramine, purity: 97.5%

4. ECOTOXICITY

	DATE:	24-J U	L200	4
SUE	STANCE	ID: 1	12-24-	3

Remark:	static test
10-MAY-1994	24h-EC50: 75 mg/l (21)
Species: Exposure period: Unit: NOEC:	Daphnia magna (Crustacea) 21 day(s) mg/l Analytical monitoring: 1 -
Method:	OECD Guide-line 202
Remark:	EC50: > 3.2 - < 10 (Immobilization of parental organisms); a NOEC for the inhibition of the reproduction rate could not be determined
26-APR-1995	(18)
Species: Exposure period:	Daphnia magna (Crustacea) 24 hour(s)
Unit: EC0: EC50: EC100:	mg/l Analytical monitoring: no 22 - 92,4 - 354 - -
Method: Year: GLP:	other: Daphnien-Schwimmunfaehigkeits-Test, UBA-Verfahrensvorschlag Mai 1984, Bestimmung der Schwimmunfaehigkeit beim Wasserfloh Daphnia magna, ECO, EC5O, EC100 24h, statisches System 1989 yes
_	
Remark:	Distillate of technical product (18)
Species:	(18) Daphnia magna (Crustacea)
	(18)
Species: Exposure period: Unit:	(18) Daphnia magna (Crustacea) 48 hour(s) mg/l Analytical monitoring: no data
Species: Exposure period: Unit: EC50: Method: Year: GLP:	<pre>(18) Daphnia magna (Crustacea) 48 hour(s) mg/l Analytical monitoring: no data 33,9 - other: EEC, 1989, Methods for the determination of ecotoxicity. C.2 Acute toxicitty for Daphnia (Updated Version 11/89). EEC Directive 79(831, Annex V, Part C. Brussels, Belgium (static) 1994 no data</pre>
Species: Exposure period: Unit: EC50: Method: Year: GLP: Test substance:	<pre>(18) Daphnia magna (Crustacea) 48 hour(s) mg/l Analytical monitoring: no data 33,9 - other: EEC, 1989, Methods for the determination of ecotoxicity. C.2 Acute toxicitty for Daphnia (Updated Version 11/89). EEC Directive 79(831, Annex V, Part C. Brussels, Belgium (static) 1994 no data other TS: purity > 99 %</pre>
Species: Exposure period: Unit: EC50: Method: Year: GLP: Test substance: Remark:	<pre>(18) Daphnia magna (Crustacea) 48 hour(s) mg/l Analytical monitoring: no data 33,9 - other: EEC, 1989, Methods for the determination of ecotoxicity. C.2 Acute toxicitty for Daphnia (Updated Version 11/89). EEC Directive 79(831, Annex V, Part C. Brussels, Belgium (static) 1994 no data other TS: purity > 99 % Arithmetic mean of 3 test results (standard deviation was 5.3 mg/l).</pre>

4.3 Toxicity to Aquatic Plants e.g. Algae

Species: Endpoint: Exposure period: Unit: EC100 :	Chlorella pyrenoidosa (Algae) growth rate 5 day(s) mg/l Analytical monitoring: >= 146 -
Remark: Test condition:	Validity uncertain. Slow growth of the control culture. 25 degree C, pH 7
Species: Endpoint: Exposure period: Unit: EC10: EC50:	Scenedesmus subspicatus (Algae) biomass 72 hour(s) mg/l Analytical monitoring: no ,67 - 2,5 -
Method: Year: GLP:	other: Scenedesmus-Zellvermehrungs-Hemmtest, DIN 38412 Teil 9, Bestimmung der Hemmwirkung von Wasserinhaltsstoffen auf Gruenalgen 1989 yes
Test substance:	other TS: purity 98.04 %
Remark:	Due to the high growth rate, the pH rose to 10.2 - 10.3 after 72 hours in the control and for concentrations of TETA up to 1 mg/l (18)
Species:	Scenedesmus subspicatus (Algae)
Endpoint: Exposure period: Unit:	growth rate 72 hour(s)
EC10: EC50:	<pre>mg/l Analytical monitoring: no ,95 - >= 100 -</pre>
EC10:	,95 -
EC10: EC50: Method: Year:	,95 - >= 100 - other: Scenedesmus-Zellvermehrungs-Hemmtest, DIN 38412 Teil 9, Bestimmung der Hemmwirkung von Wasserinhaltsstoffen auf Gruenalgen 1989
EC10: EC50: Method:	,95 - >= 100 - other: Scenedesmus-Zellvermehrungs-Hemmtest, DIN 38412 Teil 9, Bestimmung der Hemmwirkung von Wasserinhaltsstoffen auf Gruenalgen
EC10: EC50: Method: Year: GLP:	<pre>,95 - >= 100 - other: Scenedesmus-Zellvermehrungs-Hemmtest, DIN 38412 Teil 9, Bestimmung der Hemmwirkung von Wasserinhaltsstoffen auf Gruenalgen 1989 yes other TS: purity 98.04 % Due to the high growth rate, the pH rose to 10.2 - 10.3 after 72 hours in the control and for concentrations of TETA up to 1 mg/l</pre>
EC10: EC50: Method: Year: GLP: Test substance:	<pre>,95 - >= 100 - other: Scenedesmus-Zellvermehrungs-Hemmtest, DIN 38412 Teil 9, Bestimmung der Hemmwirkung von Wasserinhaltsstoffen auf Gruenalgen 1989 yes other TS: purity 98.04 % Due to the high growth rate, the pH rose to 10.2 - 10.3 after 72 hours in the control and for concentrations of TETA</pre>
EC10: EC50: Method: Year: GLP: Test substance: Remark: Species:	<pre>,95 - >= 100 - other: Scenedesmus-Zellvermehrungs-Hemmtest, DIN 38412 Teil 9, Bestimmung der Hemmwirkung von Wasserinhaltsstoffen auf Gruenalgen 1989 yes other TS: purity 98.04 % Due to the high growth rate, the pH rose to 10.2 - 10.3 after 72 hours in the control and for concentrations of TETA up to 1 mg/l (18) Selenastrum capricornutum (Algae) biomass</pre>

OECD SIDS

4. ECOTOXICITY

SUBSTANCE ID: 112-24-3

other TS: Triethylenetetramine, purity 97.5%
For the enspoint growth rate , the same results were obtained
(24)
Selenastrum capricornutum (Algae) growth rate 96 hour(s) mg/l Analytical monitoring: no data 3,7 -
other: EEC, 1988, Methods for the determination of ecotoxicity. Algal inhibition test. Off J. Eur. Comm. L 133 1988-0530 1994 no data other TS: purity > 99 %
Arithmetic mean of 5 test results (standard deviation: 1.5 mg/l). The culture medium was modified by increasing the KH2PO4 conc. from 1.6 to 160 mg/l and the NaHCO3 conc. from 50 to 100 mg/l, to improve the growth of algae and the buffer capacity of the medium. (22)

4.4 Toxicity to Microorganisms e.g. Bacteria

Type: Species: Exposure period: Unit: EC0:	aquatic Pseudomonas fluorescens (Bacteria) 24 hour(s) mg/l Analytical monitoring: 500 -
Method:	other: Bestimmung der biologischen Schadwirkung toxischer Abwaesser gegen Bakterien. DEV, L 8 (1968) modifiziert
Remark:	technical product; no further information on test conditions

(18)

4.5 Chronic Toxicity to Aquatic Organisms

4.5.1 Chronic Toxicity to Fish

4.5.2 Chronic Toxicity to Aquatic Invertebrates

TERRESTRIAL ORGANISMS

4.6.1 Toxicity to Sediment Dwelling Organisms

4.6.2 Toxicity to Terrestrial Plants

Remark: no validated information

4.6.3 Toxicity to Soil Dwelling Organisms

4.6.4 Toxicity to other Non-Mamm. Terrestrial Species

Species:	other avian: Agelaius Phoenicus (redwinged blackbird)
Endpoint:	mortality
Unit:	mg/kg bw
LD50 :	> 101 -
Method:	other: no data
GLP:	no data
Test substance:	other TS: TETA (no information about purity)
Remark:	Estimated LD50 based on food consumption data over a 18 h period
29-NOV-1994	(25)

4.7 Biological Effects Monitoring

4.8 Biotransformation and Kinetics

4.9 Additional Remarks

Remark:	<pre>Sea-urchin: Inhibition of development Eggs of the species Paracentrotus lividus were incubated sea-water 30 min after impregnation (concentration TETA: - 7313 mg/l). No teratogenic effects observed. Depending on the developmental stage there was an effect</pre>	293
	larvae (293 mg/l), gastrula (731 mg/l), blastula (2925 mg/l), cleavage stage (7313 mg/l).	(26)
Remark:	Application of 1460 mg/l TETA (alcoholic solution) to 1-2 days old larval stages and 2 days old egg-stages of the species Dysdercus koenigii F. had no acute toxic effects no effects on the eggs as well as no sterilizing effects.	and

5. TOXICITY

5.0 Toxicokinetics, Metabolism and Distribution

5.1 Acute Toxicity

5.1.1 Acute Oral	Toxicity
Type: Species: Value:	LD50 rat = 2780 mg/kg bw
Method:	other: male rats, undiluted testsubstance (no further information) no data
GLP: Test substance:	no data
29-JUL-1996	(28)
Type: Species: Value:	LD50 rat ca. 3750 mg/kg bw
Method: GLP:	other: 3 animals per group; doses: 1000, 2500, 3750, 5000 mg/kg; test substance diluted in water no data
Test substance:	
17-OCT-1994	(29)
Type: Species: Value:	LD50 rat = 4340 mg/kg bw
Method: GLP: Test substance:	other: 5 animals per group, test substance diluted in water no data no data
Type: Species: Value:	(30) LD50 rat = 2500 mg/kg bw
GLP: Test substance:	no data no data
Remark:	method: no data (13)
Type: Species: Value:	LD50 rat = 4300 mg/kg bw
GLP: Test substance:	no data no data
Remark: 17-OCT-1994	method: no data (31)

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Type: Species: Value:	LD50 mouse = 1600 mg/kg bw	
GLP: Test substance:	no data no data	
Remark: 17-OCT-1994	method: no data	(
Type: Species: Value:	LD50 rabbit = 5500 mg/kg bw	
GLP: Test substance:	no data no data	
Remark: 17-OCT-1994	method: no data	(

Type: Species:	other: see method rat
Method: GLP:	other: saturated vapor at 21 degree C, 8 h exposure, 6 animals no data
Test substance:	no data
Remark: 17-OCT-1994	no symptoms (28)
Type: Species:	other: see method rat
Method: GLP: Test substance:	other: saturated vapor inhalation up to 8 h no data no data
Remark:	maximal time for no deaths 4 h (30)
Type:	other: see method
Species:	other: see method
Method:	other: 2 rats, 1 rabbit, 1 guinea pig, and 4 mice were exposed together to aerosol (10 ml of 40 (v/v) ethanol solution, 400 l chamber) for 1 h
GLP:	no data
Test substance:	no data
Remark:	effects: slight irratation of the mucous membranes and impeded respiration, effects reversible
17-OCT-1994	(29)

5.1.3 Acute Dermal Toxicity

TITRA	LD50	
Type:		
Species:	rabbit	
Value:	= 550 mg/kg bw	
Method:	other: 4 animals per dose, undiluted test substance	
GLP:	no data	
Test substance:	no data	
Remark:	no further information available	
17-OCT-1994		(20)
17-001-1994		(28)
Type:	LD50	
Species:	rabbit	
Value:	= 805 mg/kg bw	
Method:	other: occlusive application of undiluted test substance	
GLP:	no data	
Test substance:	no data	
Remark:	no further information available	
		(30)
5.1.4 Acute Toxi	city, other Routes	
Type:	LD50	
Species:	rat	
Route of admin.:		
Value:	= 200 mg/kg bw	
Method:	3-5 animals per group, test substance as aqueous solution	ı
GLP:	no data	
Test substance:	no data	
Remark:	impeded respiration	
17-OCT-1994		(29)
1, 001 1991		(2))
_		
Type:	LD50	
Species:	rat	
Route of admin.:	i.p.	
Value:	= 78,4 mg/kg bw	
Method:	no data	
GLP:	no data	
Test substance:	no data	
iebe bubbeunce.	no data	
D 1		
Remark:	symptoms like hyperemia, extravasations; regressive	
	changes in liver and kidneys; abstract	
		(32)
Type:	LD50	
Species:	mouse	
Route of admin.:		
Value:	= 604 mg/kg bw	
	ייע בא עביי בסט	
Nothod	tost substance noutralized with WGL 10 mins as	
Method:	test substance neutralized with HCl, 10 mice per group	
GLP:	no data	
Test substance:	no data	

Remark:

convulsions for max. 20 min, hyperemia of inner organs in the dead animals

(33)

5.2 Corrosiveness and Irritation

5.2.1 Skin Irritation Species: rabbit Method: other: non occlusive appl.; a) 0.01 ml undiluted b) 10% in water no data GLP: no data Test substance: effects: a) 2 out of 2 animals with necrosis Remark: b) no effects no further information available 17-OCT-1994 (28) Species: rabbit Method: other: 20 mg applied to skin no data GLP: no data Test substance: Remark: effects: necrotic foci and extravasations no further information available, abstract (32) Species: rabbit Method: other: undiluted drug applied to the skin of 5 animals; no further information available GLP: no data Test substance: no data effects: erythema, edema, necrosis Remark: (30) Species: guinea pig Method: other: intracutaneous injection of 0.1 ml 0.5-1% solution in water (non neutralized) or 2-3% solution in neutralized form GLP: no data Test substance: no data Remark: effects: slight necrosis no further information available (34) Species: rat Method: other: a) 1000 mg/kg undiluted; b) 50 mg/kg (25% in water); application on the shaved ventral skin; exposure time: 2 h GLP: no data Test substance: no data

Remark: 17-OCT-1994	effects: strong irritations in both cases	(29)
5.2.2 Eye Irrita	tion	
Species:	rabbit	
Method:	other: instillation of a) 0.005 ml undiluted or b) 0.5 ml 40% watery solution	of a
GLP:	no data	
Test substance:	no data	
Remark:	effects: a) severe damage of the cornea b) 15% of the cornea damaged	
17-OCT-1994		(28)
Species:	rabbit	
Method: GLP:	other: 20 mg applied to the conjunctival sac no data	
	no data	
Remark:	effects: inflammation and lymphatic exudation no further information available, abstract	

(32)

5.3 Sensitization

Type: Species: Result:	Guinea pig maximization test guinea pig sensitizing	
Method: GLP: Test substance:	other: 10 animals tested; induction concentration 0.5% intradermal and topical, challenge 2% no data other TS: purity 99.5 %	
Remark:	90% positive	(35)
Type: Species: Result:	Guinea pig maximization test guinea pig sensitizing	
Method: GLP: Test substance:	other: 15 animals tested; induction concentration 0.5% intradermal and topical, challenge 2% (in water) no data other TS: technical grade (no specification)	
Remark:	80% of guinea pigs with positive reaction	(36)
Type: Species: Result:	Mouse ear swelling test mouse sensitizing	
GLP: Test substance:	no data other TS: purity 99.5 %	

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Remark:	4/10 positive (significant), induction conc. 10%, chal- lenge 2.5%.
	(35)
Type: Species:	Open epicutaneous test human
Remark:	10 out of 22 workers exposed to araldite D and hardener TETA showed slight dermatosis, one worker serious allergic eczema. One of the 11 (the one with serious allergic eczema) showed allergic hypersensitivity in epicutaneous testing to TETA.
	(37)
Type: Species: Result:	Patch-Test guinea pig not sensitizing
Method:	other: no data
GLP:	no data
Test substance:	no data
Remark:	no further information available, abstract (32)
Type:	Patch-Test
Species:	human
Test substance:	no data
Remark:	4 out of 10 patients with dermatitis due to oil-based, amine containing drilling mud, showed allergic response to a 0.5% solution in the patch test.
	(38)
Type: Species:	Patch-Test human
species.	11010011
Remark:	In 23 out of 135 (18%) workers exposed to epoxy resins, a work-related dermatosis on the hands and/or forearms had been presented during the past 3 years. In all workers patch tests were performed and in 2 positive reactions to TETA were observed (2 out of 112 without dermatosis). (39)
Туре:	Patch-Test
Species:	human
Remark:	422 employees of 8 factories had contact to epoxy resins and hardener TETA. In the course of 7 years there were 126 cases of dermatitis, 99 of whom were patch tested. 55.1% were positive to 1% TETA in water. The mean period between starting work and occurrence of dermatitis was 18.5 months. (40)
Type: Species:	Patch-Test human
Remark:	1544 patients(dermatitis) without exposure to epoxy resin systems and 137 patients in occupational contact with epoxy resins were patch tested. 28 out of the 1544 patients were

5. TOXICITY	SUBSTANCE ID: 112-24-3
	positive to ethylenediamine; 12 of these were tested with TETA, 2 were positive. 400 out of the 1544 patients were also tested with TETA and re- sults were negative. Tests with 137 patients in occupational contact to resins resulted in coexistence of positive reactions to TETA and ethylenediamine and TETA and diethylenetriamine. (41)
Type: Species:	Patch-Test human
Remark:	A 58 years old woman with dermatitis due to exposure with epoxy resins showed positive reaction in the patch test to epoxy resin and TETA as well as to ethylenediamine. (42)
Type: Species:	Patch-Test human
Remark:	12 out of 32 ethylenediamine-sensitive patients showed cross-sensitivity reaction to TETA in the patch test. (43)
Type: Species:	Patch-Test human
Remark:	19 out of 71 patients with allergic epoxy resin dermatitis were also allergic to different hardeners. 3 of them showed positive reactions to TETA in epicutaneous testing. (44)
Type: Species:	Patch-Test human
Remark:	A shipwright''s yard worker complained a chronic dermatitis of the fingertips and palms. Beside other material he used epoxy resin SP 106. In the patch test a positive reaction to TETA was demonstrated after 48 and 96 h. (45)
Type: Species:	Patch-Test human
Test substance:	no data
Remark:	31 students and instructors at the same dental school were patch tested to contactants in dental components including TETA. None had any history of allergy. No positive allergic reactions were found. (46)
Type: Species:	Patch-Test human
Test substance:	no data
Remark:	2 out of 7 patients with airborn contact dermatitis of hands and face due to epoxy resins showed positive reactions in the patch test to TETA.

(47)

Type: Species:	Patch-Test human
Remark:	14 young female patients (12 of them were seborrhean) in occupational contact with araldite D and hardener 951 (mainly TETA) suffering from eczema were patch tested. 1 of the 14 women was positiv to 3% of the hardener in ethanol (48 h). (48)
Type: Species:	other human
Remark:	20 workers (6 without, 8 with slight and 6 with severe dermatosis) were patch tested with technical TETA (1% in water). 5 of the 6 workers with severe dermatosis showed a positive reaction. (34)
	(34)
Type: Species:	other: see remarks human
Remark:	<pre>164 out of 328 workers from 11 factories producing electrical equipment showed slight dermatosis (21%, erytamotous itching patches) or severe eczemas (22%) caused by direct contact to araldite resin D or hardener TETA. TETA concentration in air was below analytic limits of 0.00015 mg/l.</pre> (49) (50)
Type: Species:	other: see remarks human
Remark:	6 workers with diagnoses of occupational asthma were examined for sensitivity to epoxy resin systems and their components. In one worker asthma followed exposure to TETA fume in inhalation challenge testing. Skin sensitivity test was negative. (51)
Type:	other: see remarks
Species:	human
Remark:	447 patients suffering from eczema, occupationally exposed to epoxy resins, have been tested with Epidian 5 (resin) and five concentrations of the hardener TETA. In Poland these health damages were characterized by a considerable percentage of those sensitized to TETA. The calculation of eczema incubation period and testing the allergen by several allergen concentrations demonstrated that the sensitivity to TETA was sometimes very enhanced.

(52)

5.4 Repeated Dose Toxicity

rat. Species: Sex: male/female Strain: other: Harlan-Wistar Route of administration: oral feed Exposure period: 7 days Frequency of treatment: daily ad libitum Post exposure period: no data m: 0.5, 1.23, 2.98 g/kg b.w.; f: 0.47, 1.38, 2.63 g/kg Doses: b.w. Control Group: no data specified NOAEL: , 5 Method: other: 5 rats per dose and sex GLP: no data Test substance: no data Remark: LOEL: 1.23 (m) and 1.38 (f) mg/kg b.w./day remarks: no deaths occurred Result: highest dose: depression of body weight gain, decrease of relative and absolute liver weights, increase of relative kidney weights. medium dose: increase of relative kidney weights. 17-OCT-1994 (28) rat Species: Sex: male/female Strain: Fischer 344 Route of administration: drinking water Exposure period: 90 d Frequency of treatment: daily Post exposure period: no 0, 120, 600, 3000 ppm (see remarks) Doses: Control Group: other: concurrent no treatment (diet: cereal based NIH-31, purified AIN-76A, Cu-deficient AIN-76A) NOAEL: = 3000 ppm Method: other: 18 rats/sex and dose group, different diets: cereal based (NIH-31) or purified (AIN-76A) diet; hematology and plasma chemistry; necropsy and histopathology; statistical analyses Year: 1996 GLP: no data Test substance: other TS: trientine-2HCl: purity: > 99 % Remark: test substance consumption: NIH-31 diet: f:14, 70, 352 mg/kg bw; m:10, 55, 276 mg/kg bw AIN-76A diet: f:13, 60, 323 mg/kg bw; m:10, 53, 270 mg/kg bw Result: no death occurred; pobabely attributed to dosing with trien-2HCL: females: a significant trend toward an increased prevalence of uterine dilatation; no other findings 23-JUN-1997 (53) Species: Sex: female rat Strain: Wistar Route of administration: dermal 17 days Exposure period:

Frequency of trea Post exposure per Doses: Control Group:		once daily (3rd - 19th day of gestand no ca. 4 mg/rat and day yes	tion)
Method:		10 rats per group. One drop of the t bbed into the shaved skin	test substance
GLP: Test substance:	no data no data		
Remark: Result:	pregnam progress inflamm superfi sialic asperta serum; reduced leucylm decreass of serce activit alkalim	no data nt and nonpregnant rats: reduced weig ssive emaciation, apathy, lack of apy matory symptoms such as erythema, ed icial erosions. pregnant rats: increa acid; increased activity of lactate ate aminotransferase and acid phospha decreased plasma activity of alkalin d haptaglobin concentration; increas haphthylamidase in amniotic fluid. Increased sed total plasma protein and elevated bmucoid a. haptaglobin; in the serum ty of lactate dehydrogenase, leucylna ne phosphatase; inhibited activity o e aminotransferase.	petite, local ema and ase of plasma dehydrogenase, atase in the ne phosphatase; sed acti- vity of onpregnant rats: d concentrations increa- sed aphthylamidase and
Species: Strain: Route of administ Exposure period: Frequency of trea Post exposure per Doses: Control Group:	tment:	Wistar dermal 17 days	ex: female
Method: GLP: Test substance:		a	
Remark: Result:	pregnan weight subcuta pregnan liver i nonpreg gammag	no data nt and nonpregnant rats: loss, hyperemia of liver and kidney aneous tissue with inflammatory infi nt rats: aspartate aminotransferase inhibited. gnant rats: increased activity of lutamyltranspeptidase in the kidney a anine aminotransferases in the liver	ltrates. activity in the and aspartate
Species: Strain: Route of administ Exposure period: Frequency of trea Post exposure per	tment:	no data	ex: no data

Doses: Control Group:	a) 215 or 430 mg/kg b) 0.8 or 4 mg/kg no data specified
Method: GLP: Test substance:	other: no data no data no data
Remark:	LOEL: a) 215 mg/kg b.w. b) 0.8 mg/kg b.w./day, 10 months no dose effect relation; abstract, no further information available.
Result:	<pre>4 months both doses: Excitability of the central nervous system decreased. Plasma levels of hippuric acid, protein and hemaglobin were decreased. Inhibited activities of catalase and peroxidase. 10 months both doses: Increased excitability, stimulated tactile reflexes. Antitoxic, carbohydrate and protein function of the liver disturbed. Transient inhibition of nicotinamide coenzymes and stimulation of cytochrome oxidase.</pre>
17-OCT-1994	(31)
Species: Strain: Route of administ: Exposure period:	mouse Sex: male/female B6C3F1 ration: drinking water 90 d
Frequency of trea Post exposure per	-
Doses:	0, 120, 600, 3000 ppm (see remarks)
Control Group:	other: concurrent no treatment, (diet: cereal based NIH-31, purified AIN-76 A, Cu-deficient AIN-76A)
NOAEL:	= 600 ppm
Method:	other: 20 mice/sex and dose group; different diets: cereal based (NIH-31) or purified (AIN-76A); hematology and plasma chemistry; necropsy, histopathology, statistical analyses
Year:	1996
GLP: Test substance:	no data other TS: trientine-2HCl; purity: > 99 %
Remark:	test substance consumption: NIH-31 diet: f:22,107, 551 mg/kg bw; m:22,107, 487 mg/kg bw AIN-76A diet: f:19, 99, 483 mg/kg bw; m:17, 92, 443 mg/kg bw
Result:	diet AIN-76A, 3000 ppm: chronic interstititial inflammation and alveolar histocytic infiltration of the lung, spleen hemapoetic cell proliferation, liver periportal fatty change, kidney weight reduction, reduced renal cytoplasmatic vacuolization, body weight gain reduction
27-JAN-1998	(53)
Species: Strain: Route of administ: Exposure period: Frequency of trea Post exposure per Doses: Control Group:	55 days tment: once daily
	-

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Method: GLP: Test substance:	of ges	tation. One drop of the aved skin. a	pregnant guinea pigs on day 10 test substance was rubbed into
Remark:	LOEL:	no data	
	remark: expose	s: 6 out of 10 nonpregna	ant and 2 out of 9 pregnant re end of experiment. No ic effects available.
Result:	activi elevat nonpre signif	nt guinea pigs: ty of gammaglutamyltrans ed in kidney and blood. gnant guinea pigs: icantly increased activ: ransferase.	speptidase significantly ity of liver aspartate
			(56)
Species: Strain:		guinea pig no data	Sex: female
Route of administ Exposure period:	ration:		, then every second day for 45
Post exposure per Doses: Control Group:	iod:	no ca.4 mg/animal and day yes	
Method: GLP:		ion; one drop of the tes skin	sure started on day 10 of st substance was rubbed into the
Test substance:	no dat	a	
Remark: Result:	7 out w died w nonpre emacia by ery ainmal conges Pregna	ithin the first 10 days, gnant animals showed wei tion; skin revealed infl thema, edema and erosion s showed all fatty deger tion of the kidney and k	lammatory alterations indicated n. Surviving and nonsurviving heration of the liver, brain, and brain edema. tic changes in the placenta
Species		other: see remarks	Sex: no data
Species: Strain: Route of administ: Exposure period:	ration:	no data	
Post exposure per Doses: Control Group:	iod:	no data 0.4 ml in 5 ml ethanol no data specified	as aerosol in a 400 l chamber
Method:		ther: 1 guinea pig, 1 ra together in one chamber.	bbit, 2 rats, 4 mice were exposed.
GLP: Test substance:		no data no data	

Remark:	LOI	EL: no da	ata	
	no	further	information	available
Result:	no	effects		
17-OCT-1994				

(29)

5.5 Genetic Toxicity 'in Vitro'

Type: System of testing: Metabolic activation: Result:	Ames test Salmonella typhimurium, TA 100, TA 1535 with and without positive	
Method: GLP: Test substance:	other: no data no data no data	
Remark:	abstract, no further information available	(58)
Type: System of testing: Metabolic activation: Result:	Ames test Salmonella typhimurium, TA 100, no data positive	
Method: GLP: Test substance:	other: no data no data no data	
Remark:	0.07 revertants per nmole; abstract, no further information available (!	59)
Type: System of testing: Metabolic activation: Result:	Bacterial gene mutation assay Escherichia coli without positive	
Method: GLP: Test substance:	other: no data no data no data	(60)
Type: System of testing: Metabolic activation: Result:	Ames test Salmonella typhimurium, TA 92, 98, 100 without positive	
Method: GLP: Test substance:	other: no data no data no data	(60)
Type: System of testing: Metabolic activation: Result:	Ames test Salmonella typhimurium, TA 98, 100, 1535, 1537, 1538 with and without positive	
Method: GLP:	other: no data no data	

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other TS: purified TETA-2Hydrochloride Test substance: (61) Type: Ames test Salmonella typhimurium, TA 98, 100, 1535, 1537 System of testing: Metabolic activation: with and without Result: positive Method: other: preincubation assay GLP: no data Test substance: other TS: technical grade (68.1%) (62) Type: Ames test Salmonella typhimurium, TA 98, 100, 1535, 1537, 1538 System of testing: Metabolic activation: with and without Result: positive Method: other: no data GLP: ves other TS: techn. grade; 2 samples: 56.4 and 68.5% purity Test substance: (63) (64) Type: Mammalian cell gene mutation assay CHO cells System of testing: with and without Metabolic activation: Result: positive Method: other: no data GLP: no data Test substance: other TS: purity 79.15% Remark: no clear dose-response relationship (65) Type: Mammalian cell gene mutation assay System of testing: CHO cells with and without Metabolic activation: Result: negative Method: other: no data GLP: no data Test substance: other TS: purity 99.42% (66) Type: Sister chromatid exchange assay System of testing: CHO cells Metabolic activation: with and without positive Result: Method: other: no data GLP: no data Test substance: other TS: purity 99.42% (66) Unscheduled DNA synthesis Type: System of testing: rat hepatocytes Metabolic activation: without Result: positive

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Method: GLP:	other: no data no data	
Test substance:	other TS: purity 99.42%	
		(66)
Туре:	Sister chromatid exchange assay	
System of testing		
Metabolic activat		
Result:	positive	
Method:	other: no data	
GLP:	no data	
Test substance:	other TS: purity 79.15%	
		(65)
Type:	Unscheduled DNA synthesis	
System of testing	: rat hepatocytes	
Metabolic activat		
Result:	positive	
Method:	other: no data	
GLP:	no data	
Test substance:	other TS: purity 79.15%	
		(65)
Type:	Sister chromatid exchange assay	
System of testing	: CHO cells	
Metabolic activat	ion: with and without	
Result:	positive	
Method:	other: no data	
GLP:	no data	
	other TS: purity 56.4%, technical grade	
Remark:	with metab. activation only at the lowest concentration	
	(0.5 g/l) significant increase of SCEs/chromosome; no increase at 0.6 and 0.8 g/l.	
	no increase at o.o and o.o g/i.	(67)
5.6 Genetic Toxi	city 'in Vivo'	
Type:	Drosophila SLRL test	
Species:	Drosophila melanogaster Sex: no data	
Route of admin.:	-	
Exposure period: Doses:	no data no data	
20262.	no data	
Method:	other: no data	
GLP:	no data	
Test substance:	no data	
Result:	no effects	
		(68)
Туре:	Micronucleus assay	
Species:	mouse Sex: male/female	
Route of admin.:		
Exposure period:	single injection	
Doses:	185, 370, 600 mg/kg	

TRIETHYLENETETRAMINE DATE: 24-JUL.-2002

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Method: GLP: Test substance:	other: Bushy Run Research Center standard yes other TS: purity 68.5%, technical grade	d protocol
Result:	not clastogenic	(69)
Type: Species: Route of admin.: Exposure period: Doses:		o data
Method: GLP: Test substance:	other: according to Schmid, W., Mitt. II Mutagenitaetsfragen, 53 (1975) no data other TS: purified TETA-Dihydrochloride	I der Komm. fuer
Result:	not clastogenic	(61)
Type: Species: Route of admin.: Exposure period: Doses:	Micronucleus assay mouse Sex: no oral unspecified single application 1500, 3000, 6000 mg/kg	o data
Method: GLP: Test substance:	other: according to several published met no data other TS: purified TETA-2Hydrochloride	thods
Result:	not clastogenic	

(61)

5.7 Carcinogenicity

Species: Strain: Route of administration: Exposure period: Frequency of treatment: Post exposure period: Doses: Control Group:	life-time	Sex:	male
GLP: no dat	see remarks a IS: purity 79.15% (analytic)		
remark solutio skin i mortal:	: no further data available s: 50 animals per group; 0.025 ml on applied; dose highest one that rratation nor reduced weight gain. ity. Dosage very low compared to LI	resul No i D50.	ted in neither ncreased
	atment related skin tumors, no evic nce of any other tumor.	dence	of increased

(70)

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Species: Strain: Route of administration Exposure period: Frequency of treatment: Doses:	2 years	Sex: male
Result: No ef morta non-t	imals/group fects were observed on any paramete lity, body weights and incidence of umorous lesions. arope S.A., Switzerland	, 5
24-MAY-1994		(71)

5.8.1 Toxicity to Fertility

5.8.2 Developmental Toxicity/Teratogenicity

Species: Strain: Route of administ: Exposure period: Frequency of trea Doses: Control Group:		rat Sprague-Dawley gavage day 6-15 of gestation once daily 75, 325, 750 mg/kg yes	Sex:	female
Method: GLP: Test substance:	no data	est substance diluted in water : purity > 98%		
Remark: Result:	No subst	er information available ance related effects on dams or fetal body weight at 750 mg/kg ance).		
Species: Strain: Route of administ: Exposure period: Frequency of trea Doses: Control Group:		<pre>rat Sprague-Dawley oral feed day 0-21 of gestation daily ad libitum 0.17, 0.83, 1.66% in the diet b.w. and day) yes</pre>		female D, 1660 mg/kg
GLP: Test substance:	no data other TS	: purity > 99%, TETA-4Hydrochlo	ride	
Remark:	dose rel	ize unchanged, all described ef: ated. Authors comment: teratoge to induced Cu deficiency and Z	nicity of	f the drug in
Result:	0.17% dams(n=5 increase	<pre>(n=7): no resorbed or abnormal): no effects except reduced li d kidney zinc concentration. Fe whole fetus and liver Zn conc. duced.</pre>	ver copp tuses: 5	er and .8% resorbed

<pre>0.83% dams (n=9): reduced weight gai and plasma, Zn conc. increased Fetuses: 8.7% resorbed (7/93), like hemorrhage and edema, Cu liver and placenta, Zn concent and liver. 1.66% dams (n=5): reduced food consu highly signif. reduced weight in liver and plasma. Zn conc. nese conc. in muscle and iron Fetuses: 18.8% resorbed (9/48) like hemorrhages, edema, reduc vertebrae and phalanges; feta Trace elements same results as</pre>	in kidney and mus 25,6% abnormalit: decreased in who ration elevated is amption; gain and copper co in kidney and mus conc. in liver in ; 100% abnormalit: red ossification of weight and lengt in medium dose.	scle. ies (22/86) le body, n whole body oncentration cle, manga- creased. ies (39/39) f caudal
rat	Sex:	female

Species: Strain: Route of administ Exposure period: Frequency of trea Doses: Control Group:		rat Sprague-Dawley oral feed day 0-21 of gesta daily ad libitum 0, 0.83 or 1.67% Cu/kg diet yes			female 0.05 or 0.5 mg
Method:	other: 4	rats per group			
GLP:	no data				
Test substance:	other TS	: purity > 99%			
Remark:		ize not altered by	test substance o	r Cu	
	administ				
		comment: teratogen			
	-	to induced Cu def	1		
Result:	Maternal signific copper s in any g (69%, 27 with the high Cu edema, h skulls. correlat copper l Increase fetal zi	and to 830 or 1670 weight gain and f antly decreased at upplement. Frequen roup. Significant out of 39 fetuses low Cu concentration. Typ ydronephrotic kidn The lowered terato ed with an increas evels by Cu supple d maternal and nc levels due to t by Cu coadministra	etal weight and 1 1.67% without im cy of resorption of incidence of fetal due to 1.67% in on was lowered to es of abnormalitie eys, micrognathia ogenetic effect of e in maternal and ment. he test substance	engtl prove not c l abr com 6.5% es: f and 1.6° feta	h were ement by different normalities bination (3/46) by nemorrhage, domed 7% was al tissue e not
				(7)	7) (78) (79)
Species:		rabbit		Sex:	female

sex: female

rabbit
other: New Zealand
dermal
day 6-18 of gestation
6 h each day

OECD SIDS

5. TOXICITY

Doses: 5, 50, 125 mg/kg dissolved in 2 ml distilled water Control Group: yes NOAEL Teratogenicity: 125 mg/kg bw Method: other: 22 rabbits per group; application occlusive GLP: no data Test substance: other TS: purity 95% Result: No embryotoxic or teratogenic drug related effects at any dose. Maternal toxicity: 125 mg/kg induced delayed weight gain and death of 2 out of 22 rabbits. Strong local irritations of the skin at 50 and 125 mg/kg and slight reversible irratations at 5 mg/kg. No reduction of copper concentrations in urine and plasma. (80) Species: other: chicken Sex: no data Strain: other: White Leghorn Route of administration: other Exposure period: once in 3 days old embryos Doses: 0.051, 0.102, 0.204 or 0.408 mg per egg dissolved in 5 ul acetone other: solvent Control Group: Method: other: injection on the inner shell membrane GLP: no data Test substance: other TS: technical grade Result: deaths of embryos malformed survivors 0.051 mg 1 out of 30 2 out of 29 0.102 mg 3/30 3/27 0.204 mg 10/30 4/20 0.408 mg 20/20 _ _ _ _ acetone 1/100 0/100 Malformations occurred in the eyes, wings and abdominal wall. Oedema, enlarged lymph sacs and stunting and twisting of the backbone. ED50 for embryotoxicity: 0.155 mg per egg. (81)

5.8.3 Toxicity to Reproduction, Other Studies

5.9 Specific Investigations

5.10 Exposure Experience

Remark:	TETA-2Hydrochloride is used in the therapy of Wilson''s
	disease (inherited metabolic desease characterised by
	copper accumulation predominantly in liver, cornea, brain,
	and kidney) when the drug of choice (Penicillamine) is not
	tolerated. All authors reported no serious side effects.
	(82) (83) (84) (85) (86) (87) (88) (89) (90) (91)

Remark: In primary biliary cirrhosis treatment TETA is an unsuitable drug due to gastrointestinal side effects, skin rash and rhabdomyolysis (one out of 4 patients 48 h after 1. dose)

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	(92)	
Remark:	There was no evidence of teratogenicity in 4 patients who became pregnant while taking TETA-2Hydrochoride against Wilson''s disease (6 pregnancies).	
	(89)	
Remark:	6 out of 20 employees working with ethoxylin cast resin and the hardener TETA suffered from work related eczematous dematosis. 8/20 showed slight skin irratations like erythemaand itching. In epicutaneous skin test 5 out of 6 workers with strong dermatosis were sensitized to TETA (technical grade).	
	(93)	
Remark:	Serum monoamine oxidase activity in 15 workers handling with epoxy resin and hardener TETA was significantly elevated compared to a control group. Increased activity reflect possibly increased amine metabolism in the connective tissue.	
	(94)	
Remark:	12 workers exposed to araldite and hardener TETA were examined 2 to 4 times at intervals of 6 months. After 1 year there was a decrease in the relative percentage of lymphocytes and a corresponding increase in neutrophils. 5 workers reported subjective symptomes like drowsiness, headache, gastric pain, fatigue, weakness and decreased appetite. 7 showed dermatosis. (95)	
Remark:	No significant improvement occurred in hand eczema of 23 nickel-sensitive patients treated with 300 mg TETA/d in a double blind study.	
	(96)	
Remark:	Plasma levels were measured in 4 male and 4 female patients receiving treatment for excess copper. Maximal plasma levels of 0.3- 15 mg/l (male) and 1.0- 2.2 mg/l (female) were seen 3 h after oral administration of 8.3 mg/kg b.w	
	The free form of the drug was not detected, indicating chelation with metal ions (predominantly copper). test substance: TETA-2Hydrochloride	
	(97)	
Remark:	Using the oral copper loading test and the 24 h urine excretion test on patients with Wilson''s disease it could be	
	shown, that longterm therapy with 1.2 g/d TETA (more than 3 months) led to a decreased intestinal copper absorption and to an increased urine copper excretion. test substance: TETA-2Hydrochloride	
	- (98)	

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5.11 Additional Remarks

Type:	Biochemical or cellular interactions
Remark:	Female F-344 rats received i.m. 0.75 mmol/kg TETA prior to 0.068 or 0.10 mmol/kg nickeldichloride (i.p. or i.m.). In rats killed 6 h after injection of TETA and nickelchloride, Ni concentration in liver, kidney, spleen, lung and heart averaged 3.4, 0.72, 0.27, 0.22, and 0.12 times corresponding Ni concentrations in contol rats that received only nickelchlorid. Ni-induced hyperglycemia and hyperglucagonemia were not prevented. TETA markedly reduced plasma Ni conc. and increased urine Ni excretion during 6 h after injection. Test substance: purified TETA-4Hydrochloride
	(99)
Type:	Biochemical or cellular interactions
Remark:	Norwegian hooded rats received 100 mg TETA per rat with the diet for 3 days and the urine copper concentration was determined. The basal copper excretion of 65.1 nmol/24 h rose after drug application to 305.9 nmol/24 h. Test substance: TETA-2Hydrochloride (100)
Type:	Biochemical or cellular interactions
Remark:	Female mixed-breed dogs were administered 150 mg TETA orally in gelantine capsules twice daily for 23 days and serum and 24 h urine were analysed on day 0, 9, 15, and 23. Cu concentration in serum was unchanged but increased in urine from 0.119 to 0.663 mg/24 h. Zn and Fe concentration in plasma and urine were not changed. Predictive value reduced by low number of animals (n=3). Test substance: TETA-4Hydrochloride (101)
Туре:	Biochemical or cellular interactions
Remark:	Nickel-poisened rats survived at a nickel:TETA ratio of 1:1. Urinary and biliary excretion of nickel was significantly enhanced. (102)
Туре:	Biochemical or cellular interactions
Remark:	Sodium diethyldithiocarbamate and D- pencillamine are significantly more effective upon acute toxicity of nickel carbonyl in rats than TETA. (103)
Туре:	Biochemical or cellular interactions
Remark:	The distribution of radioactive nickel, iron, manganese, and tin in plasma was studied in rats which received i.p. injections of their salts with or without i.m. injection of TETA. TETA was most effective in reducing nickel, followed by iron, manganese and tin.

	test substance: no data
	(104)
Туре:	Biochemical or cellular interactions
Remark:	A single i.p. application of TETA decreased significantly the total body burden of zinc 24 h after i.v. injection of Zn chloride (0.14 mg/kg). Simultaneous peroral administration of TETA with Zn increased whole body burden of Zn, indicating possibly enhanced absorption of zinc. test substance: TETA-2Hydrochloride (105)
Туре:	Biochemical or cellular interactions
Remark:	In a comparative study on the effects of 7 chelating drugs on trace metal and biochem. alteration in the rat TETA is one of the drugs producing least effects on the levels of trace metals and biochem. parameters. test substance: no data
	(106)
Туре:	Biochemical or cellular interactions
Remark:	TETA is an effective antidote to acute nickel carbonyl poisoning (4.35 mg/l for 15 min) when it is administered 10 min after and not 10 min before exposure in rats. test substance: no data
	(107)
Туре:	Biochemical or cellular interactions
Remark:	In a comparative study with 16 chelating agents TETA has been shown to be one of the most effective drugs enhancing urinary excretion of copper in the rat. test substance: no data (108)
Туре:	Biochemical or cellular interactions
Remark:	6 daily i.p. injections of 146 mg/kg TETA enhanced significantly excretion of all essential trace metals in rats. In serum levels there were no significant changes indicating redistribution. test substance: no data (109)
Type:	Biochemical or cellular interactions
Remark:	In cadmium preexposed rats 500 mg/kg TETA reduced the hepatic Cd burden but did not elicit any influence on other tissues except pancreas. test substance: TETA-hydrochloride (110)
Туре:	Toxicokinetics
Remark:	The maximal plasma concentration 2 h after a single oral administration of 25 mg/kg was 8 microg/ml in fasted, 3 in nonfasted rats(max after 1h) and 24 microg/ml after

intraduodenal application. Bioavailability 4 h after administration was 6.6, 2.3, and 17.6%, respectively. Plasma levels after i.v. administration of 0.1 mg per rat were 0.0013 mg/ml 10 min. after injection and 0.00045 mg/ml after 4 h. The urinary excretion of unchanged TETA during 24 h was 3.1% of the oral dose and total urinary excretion including not identified metabolites amounted to 35.7% of the dose. Main absorption by permeation across the plasma membrane of intestinal epithelial cells. Binding to the brush border membran was totally inhibited by 0.05 mmol copper.

test substance: TETA-2Hydrochloride

(111)

- (1) BUA Report No. 89 "Triethylentetramin", VCH, Weinheim, (1992)
- (2) Beilsteins Handbuch der organischen Chemie, Band 4, Springer Ver- lag, Berlin, 255 (1922)
- (3) Bayer AG, DIN-Safety Data Sheet Triethylentetramin 18.06.93
- (4) Keith, L.H. & Walters, D.B., Compendium of safety data sheets for research and industrial chemicals, Part 3, Verlag Chemie, Wein- heim, 1670-1671 (1985)
- (5) Kuehn, R. & Birett, K., Merkblaetter gefaehrlicher Arbeitsstoffe, Band 6, Blatt Nr. T 43, Ecomed Verlag, Landsberg (1982)
- (6) Wilson, A.L., Ind. Eng. Chem. 27, 867-871 (1935)
- (7) Hommel, G., Handbuch der gefachrlichen Gueter, Springer Verlag, Berlin, Merkblatt 569 (1988)
- (8) Hann, R.W. & Jensen, P.A., Water quality characteristics of ha- zardous materials, Texas A & M University, 3 S. (1974)
- (9) Hart, A.W., In: Kirk-Othmer, Encycl. chem. technol., Bd. 7, New York, 22-39 (1965)
- (10) Leo, A. et al., Chem. Reviews 71, 525&575 (1971)
- (11) Fiedler, H., Chemical and physical properties worksheet (CAS Nr. 112-24-3), unveroeffentlicht (1989)
- (12) Calculation Bayer AG, WV-UWS/Produktsicherheit, 1992
- (13) Spitz, R.D., Diamines and higher amines, aliphatic, In: Kirk- Othmer, Encyclopedia of chemical technology, Vol.7, 580-602 (1979)
- (14) Safety Data Sheet Dow Europe S.A., August 1993
- (15) Baldwin, W.C.G, Pr. Roy. Soc. A, 167, 539-554 (1937)
- (16) Atkinson, R., Ed., AOP, Rate of hydroxyl radical and ozone reaction from chemical structure, Version SRC 1.31, Syracuse Res. Co. University of California, Riverside, USA (1990)
- (17) Hansen, E.B. et al., J. Anal. Toxicol. 9, 167-171 (1985)
- (18) Bayer AG data
- (19) unpublished report CRL 189039 from AKZO, 1989

- (20) DOW Chemical Company, unpublished data (1978)
- (21) unpublished report CRL 189044 from AKZO, 1989
- (23) Dombrowicz, E. et al., Acta Pol. Pharm. 42, 184-191 (1985)
- (24) unpublished report CRL F90088 from AKZO, 1990
- (25) Schafer, E.W. et al., Arch. Environm. Contam. Toxicol. 12, 355-382 (1983)
- (27) Chatterjee, M. et al., Int. J. Environm. Stud. 24, 87-95 (1985)
- (28) Chemical Hygiene Fellowship, Carnegie-Mellon University, Special Report 39-54 (1976); cited in BG Chemie, Toxikologische Bewertung Nr. 181, Triethylentetramin (1991)
- (29) Bayer AG, Untersuchung von E 570 und Haerter T, unveroeffentlichter Bericht (1957)
- (30) Smyth, H.F. et al., J. Ind. Hyg. Toxicol. 31, 60-62 (1949)
- (32) Kowalski, Z. et al., Chem. Abstracts 59, 6889 (1963)
- (33) Srivastava, A. et al., Chemosphere 17, 839-844 (1988)
- (34) Pletscher, A. et al., Z. Unfallmed. Berufskrankh. 47, 163-176 (1954)
- (35) Maisey, J. et al., Contact Dermatitis 18, 133-137 (1988)
- (36) Thorgeirsson, A., Acta Derm. Venereol. 58, 332-336 (1978)
- (37) Markicevic, A. et al., Chem. Abstracts 61, 16688 (1964)
- (38) Ormerod, A.D. et al., Contact Dermatitis 21, 326-329 (1989)
- (40) Krajewska, D. and Rudzki, E., Contact Dermatitis 2, 135-138 (1976)

(41) Rudzki, E. and Krajewska, D., Contact Dermatitis 2, 311-313 (1976)
(42) Rudzki, E., Contact Dermatitis 4, 53 (1978)
(43) Balato, N. et al., Contact Dermatitis 15, 263-265 (1986)
(44) Jolanki, R. et al., Acta Derm. Venereol., S 134, 90-94 (1987)
(45) Camarasa, J.G. and Serra-Baldrich, E., Contact Dermatitis 20, 382 (1989)
(46) Oshima, H. et al., Contact Dermatitis 24, 138-139 (1991)
(47) Tosti, A. et al., Contact Dermatitis 19, 220-222 (1988)
(48) Welcker, A., Dermatol. Wochenschr. 132, 871-876 (1955)
(49) Grandjean, E., Brit. J. Industr. Med. 14, 1-4 (1957)
(50) Grandjean. E., Z. Praeventivmed. 2, 77-98 (1957)
(51) Fawcett, I.W. et al., Clinic. Allergy 7, 1-14 (1977)
(52) Rudzki, E. et al., Med. Pr. 32, 59-62 (1981); article in polish, data from TOXLINE abstract
(53) Greenman D.L. et al., Fundam. Appl. Toxicol. 29, 185-193 (1996)
(54) Dobryszycka, W. et al., Arch. Toxicol. 33, 73-80 (1974)
(55) Woyton, J. et al., Toxicol. Appl. Pharmacol. 32, 5-10 (1975)
<pre>(56) Dobryszycka, W. et al., Arch. Immunol. Therap. Exp. 23, 867-870 (1975)</pre>
(57) Szacki, J. et al., Arch. Immunol. Therap. Exp. 22, 123-128 (1974)
(58) Hedenstedt, A., Mutat. Res. 53, 198-199 (1978)
(59) Hulla, J. E. et al., Environ. Mutagen. 3, 332-333 (1981)
(60) Warren, G. et al., Mutat. Res. 88, 165-173 (1981)
(61) Heinz, N. and Schroeder, H.F., Drug Res. 31, 950-953 (1981)
(62) Mortelmans, K. et al., Environ. Mutagen. 8, S7, 1-119 (1986)

- (63) Union Carbide Corporation, Bushy Run Res. Center, Report 50-43 (1987)
- (64) Union Carbide Corporation, Bushy Run Res. Center, Report 50-9 (1987); cited in BG Chemie, Toxikologische Bewertung Nr. 181, Triethylentetramin (1991)
- (65) Union Carbide Corporation, Bushy Run Res. Center, Report 43-127 (1981); cited in BG Chemie, Toxikologische Bewertung Nr. 181, Triethylentetramin (1991)
- (66) Union Carbide Corporation, Bushy Run Res. Center, Report 44-11 (1981); cited in BG Chemie, Toxikologische Bewertung Nr.181, Triethylentetramin (1991)
- (67) Union Carbide Corporation, Bushy Run Res. Center, Report 50-99 (1987); cited in BG Chemie, Toxikologische Bewertung Nr. 181, Triethylentetramin (1991)
- (68) National Toxicology Program, Annual Plan for Fiscal Year 1986, report NTP-86-086, 72 (1986)
- (69) Union Carbide Corporation, Bushy Run Res. Center, Report 50-122 (1987)
- (71) Young, J.T., Grandjean, M. and Swaim, L.D., unpublished report of the DOW Chemical Corporation (1986)
- (72) Ciba-Geigy Ltd., Report on TK 10458, report No. 830035 (1984), cited in BG Chemie, Toxikologische Bewertung Nr. 181, Triethyl- tetramin (1991)
- (73) Cohen, N.L. et al., Fed. Proc. Fed. Am. Soc. Exp. Biol. 41, 944 (1982)
- (74) Keen et al., Inflammatory Dis. Copper (Proc. Symp.) 109-121 (1982)
- (75) Keen, C.L. et al., Lancet I, 1127 (1982)

- (78) Hurley, L.S. et al., Teratology 25, 51A (1982)
- (79) Keen, C.L. et al., Teratology 25, 53A (1982)

TRITHYLENE TETRAMINE DATE: 24-JUL.-2002 SUBSTANCE ID: 112-24-3

(80)	Union Carbide Corporation, Bushy run research center, Project report 50-127 (1988); cited in BG Chemie, Toxikologische Bewer- tung Nr. 181, Triethylentetramin (1991)
(81)	Korhonen, A. et al., J. Appl. Toxicol. 3, 112-117 (1983)
(82)	Bachmann, H. et al., Eur. Neurol. 29, 301-305 (1989)
(83)	Brewer, G.J. et al., Semin. Neurol. 7, 209-219 (1987)
(84)	Dubois, R.S. et al., J. Pediat. Gastroent. Nutr. 10, 77-81 (1990)
(85)	Fromtling, R.A., Drugs of Today 23, 507-508 (1987)
(86)	Haslam, R.H.A. et al., Dev. Pharmacol. Ther. 1, 318-324 (1980)
(87)	Scheinberg, I.H. et al., New Engl. J. Med. 317, 209-213 (1987)
(88)	Walshe, J.M., Lancet II, 1401-1402 (1969)
(89)	Walshe, J.M., Lancet II, 643-647 (1982)
(90)	Walshe, J.M., Prog. Clin. Biol. Res. 34, 271-280 (1979)
(91)	Walshe, J.M., Quart. J. Med. 42, 441-452 (1973)
(92)	Epstein, O. and Sherlock, S. Gastroentrology 78, 1442-1445 (1980)
(93)	Pletscher, A. et al., Z. Unfallmed. Berufskrankh. 47, 163-176 (1954)
(94)	Yano, E., Toxicol. Letters 37, 27-32 (1987)
(95)	Zielhuis, R.L., J. Occup. Med. 3, 25-29 (1961)
(96)	Burrows, D. et al., Contact Dermatitis 15, 55-57 (1986)
(97)	Miyazaki, K. et al., Chem. Pharm. Bull. 38, 1035-1038 (1990)
(98)	Loessner, S.R. et al., Acta Neurol. Scand. 83, 364-366 (1991)
(99)	Sunderman, F.W. et al., Toxicol. Appl. Pharmacol. 38, 177-188 (1976)
(100)	Gibbs, K. and J.M. Walshe, Clinic. Sci. Mol. Med. 53, 317-320 (1977)
(101)	Allen, K.G.D. et al., Am. J. Vet. Res. 48, 28-30 (1987)
(102)	Athar, M. et al., Fundam. Appl. Toxicol. 9, 26-33 (1987)
(103)	Baselt, R.C., et al., Res. Communic. Chem. Pathol. Pharmac., 18, 677-689 (1977)

(104)	Dwivedi,R.S. et al., Chemosphere 11, 925-932 (1978)
(105)	Eybl, V. et al., Arch. Toxicol. Suppl. 13, 370-372 (1989)
(106)	Misra, M. et al., Bull. Environm. Contam. Toxicol. 41, 172-184 (1988)
(107)	Mitchell, J. et al., Clinical Toxicol. 12, 606-607 (1978)
(108)	Planas-Bohne, F., Toxicol. Appl. Pharmacol. 50, 337-345 (1979)
(109)	Tandon, S.K. et al., Environm. Res. 35, 237-245 (1984)
(110)	Tewari, P.C. et al., Clin. Exp. Pharmacol. Physiol. 15, 71-75 (1988)
(111)	Kobayashi, M. et al., Yakugaku Zasshi 110, 759-763 (1990)