FOREWORD

INTRODUCTION

2-DIETHYLAMINOETHANOL

CAS N[•]:100-37-8

SIDS Initial Assessment Report

For

SIAM 15

Boston, USA; 22-25 October 2002

1. Chemical Name: 2-Diethylaminoethanol 2. CAS Number: 100-37-8 **3.** Sponsor Country: Germany Contact Point: BMU (Bundesministerium für Umwelt, Naturschutz und Reaktorsicherheit) Prof. Dr. Ulrich Schlottmann Postfach 12 06 29 D- 53048 Bonn-Bad Godesberg BASF AG, Germany; Air Products and Chemicals Inc., USA; 4. Shared Partnership with: Atofina Chemicals Inc., USA; DOW Chemicals Company, USA; The Amines HPV Panel of the American Chemistry Council, USA 5. Roles/Responsibilities of the Partners: Name of industry sponsor BASF AG, Germany • /consortium Contact person: Dr. Hubert Lendle GUP/CL - Z570 D-67056 Ludwigshafen Process used see next page 6. Sponsorship History How was the chemical or by ICCA-Initiative category brought into the **HPV** Chemicals Programme? 7. Review Process Prior to last literature search (update): 16 May 2002 (Human Health): databases medline, toxline; the SIAM: search profile CAS-No. and special search terms 04 January 2002 (Ecotoxicology): databases CA, biosis; search profile CAS-No. and special search terms 8. Quality check process: As basis for the SIDS-Dossier the IUCLID was used. All data have been checked and validated by BUA 9. Date of Submission: 13 August 2002 **10.Comments:**

OECD/ICCA - The BUA* Peer Review Process

Qualified BUA personnel (toxicologists, ecotoxicologists) perform a quality control on the full SIDS dossier submitted by industry. This quality control process follows internal BUA guidelines/instructions for the OECD/ICCA peer review proc ess and includes:

- a full (or update) literature search to verify completeness of data provided by industry in the IUCLID/HEDSET
- Review of data and assessment of the quality of data
- Review of data evaluation
- Check of adequacy of selection process for key studies for OECD endpoints, and, where relevant, for non-OECD endpoints by checking original reports/publications
- Review of key study description according robust summaries requirements; completeness and correctness is checked against original reports/publications (if original reports are missing: reliability (4), i.e. reliability not assignable)
- Review of validity of structure-activity relationships
- Review of full SIDS dossier (including SIAR, SIAP and proposal for conclusion and recommendation for further work)
- In case of data gaps, review of testing plan or rationale for not testing

^{*} BUA (GDCh-Beratergremium für Altstoffe): Advisory Committee on Existing Chemicals of the Association of German Chemists (GDCh)

SIDS INITIAL ASSESSMENT PROFILE

CAS No.	100-37-8
Chemical Name	2-Diethylaminoethanol
Structural Formula	OH

SUMMARY CONCLUSIONS OF THE SIAR

Human Health

2-Diethylaminoethanol was rapidly absorbed via the oral route. It is presumably absorbed by dermal and inhalation routes of administration. In the rat it was widely distributed to many tissues. It was primarily excreted unchanged via the urine in rats. Excretion via the feces was also observed in rats, but to a lesser extent. Urinary excretion was also reported in humans. The major metabolites in rats were reported to be diethylaminoacetic acid and diethyl-(2-hydroxyethyl)-amino-oxide.

The LD50 for the rat after oral administration was 1320 mg/kg bw. The main clinical signs described were apathy and dyspnea. After inhalation of vapors of 2diethylaminoethanol an LC50 of ca. 4600 mg/m³/4 hour was estimated in rats using Haber's rule. Severe signs of irritation were observed, e.g. mucous membrane irritation and dyspnea. A dermal LD50 in guinea pigs was reported to be ca. 885 mg/kg bw.

2-Diethylaminoethanol was corrosive to the skin of rabbits; since the pH was measured to be 11.5 (100 g/l) at 20°C, the corrosive effects are not surprising. The potential for severe damage to the eyes can be expected based on the animal studies available and on the pH. 2-Diethylaminoethanol was not sensitizing to the skin in studies with guinea pigs.

Repeated exposure of rats to 2-diethylaminoethanol vapors (up to 365 mg/m³) for 14 weeks caused local toxicity (irritation) at the site of contact, namely, the upper respiratory tract and the eyes; however, systemic toxicity was not observed (NOAEC, systemic toxicity, 365 mg/m³ or 76 ppm). After inhalation exposure, the main symptom described was respiratory irritation which led to noises called rales and irritation of the eyes. The LOAEC for local toxicity (irritation) to the respiratory tract was 120 mg/m³ (25 ppm) and the NOAEC for local toxicity was 53 mg/m³ (10 ppm) based on histopathological effects in the nasal cavity. However, since an effect (rales) was seen at the lowest concentration a NOEC was not reached.

2-Diethylaminoethanol gave no evidence of *in vitro* mutagenic activity nor *in vivo* clastogenic potential.

Repeated exposure of rats to 2-diethylaminoethanol vapors (365 mg/m^3) for 14 weeks did not cause any adverse effects on the reproductive organs when administered by inhalation. In pregnant rats even the highest concentration tested of 486 mg/m³, which already produced maternally toxic effects, did not lead to adverse developmental effects.

In a limited study, 2-diethylaminoethanol was not carcinogenic to rats when given by feed (tested up to ca. 50-400 mg/kg/d).

An odor threshold of 0.011 ppm (approx. 0.053 mg/mg^3) has been reported. In a laboratory worker short-time exposure to approx. 100 ppm (480 mg/m³) 2-diethylaminoethanol caused nausea and vomiting. Subjects exposed to 2-diethylaminoethanol vapor by humidified air in office buildings complained about eye, nose and throat irritation, dizziness, nausea and vomiting. Also several cases of asthma were observed. However, these symptoms were more consistent with reactive airway dysfunction syndrome than with an allergic respiratory reaction. In one case

detectable amounts of 2-diethylaminoethanol were 0.05 and 0.04 mg/m^3 .

Environment

2-diethylaminoethanol is a colourless – light yellowish organic liquid. The hygroscopic substance is miscible with water in all proportions, has a vapor pressure of about 1.8 hPa at 20 °C. The density is 0.885 g/cm³. Melting point and boiling point are -68 °C and 162-163 °C (at 1013 hPa) respectively.

The distribution of the substance between the compartments of air, biota, sediment, soil and water was calculated according to Mackay Level I. The non-charged molecule distributes mainly to the water (99.1 %).

A soil adsorption coefficient (K_{OC}) of 5.98 was estimated for 2-diethylaminoethanol (DEAE). This Koc value suggests that this compound would be mobile in soil and adsorption to suspended solids would not be important. From the pKa-value of 9.87 it can be assumed that under environmental conditions the substance is available as a cation. Therefore, binding of the substance to the matrix of soils with high capacities for cation exchange (e.g. clay) cannot be excluded. However, no data was available for ionic-ionic interactions in soil. The calculated Henry's law constant ($3.16*10^4$ Pa m³ mol⁻¹ at 25 °C) and complete water solubility of 2-diethylaminoethanol suggest that volatilization from water would not be an important fate process. The substance has no considerable potential for bioaccumulation (logKow = 0.21, measured). The compound is readily biodegradable (OECD 301 A, 95% after 22 days 10d-window fulfilled). The EC20 (30 min) for activated sludge was determined to be >1000 mg/l. The photodegradation rate in the atmosphere is fast under environmental conditions (50% after 3.9 hours).

The following aquatic effect concentrations are available:

Leuciscus idus LC50 (96 h) = 147 mg/l (nominal concentration). The toxic effect may be (partly) due to the high pH of the non-neutralized test solutions, since the pH adjusted 1000 mg/l dose group tolerated the substance for 96 h without mortality.

Pimephales promelas LC50(96 h) = 1780 mg/l (measured concentration, adjustment of pH)

Daphnia magna: EC50 (48 h) = 83.6 mg/l (nominal concentration) (toxicity due to pH effects cannot be excluded)

Daphnia magna EC50 (48 h) = 165 mg/l (nominal concentration, adjustment of pH)

Scenedesmus subspicatus: EC50 = 44 mg/l, with a NOEC of 5 mg/l (corresponding values for biomass are 30 and 5 mg/l respectively; nominal concentration).

Using the aquatic toxic effect on the most sensitive species, *Scenedesmus subspicatus*, of 44 mg/l for the endpoint growth rate (30 mg/l endpoint biomass) a PNEC _{aqua} of 44 μ g/l is derived by applying an assessment factor of 1000. This factor is justified, because only short-term toxicity values were available.

The following terrestrial effect concentration was reported:

Chrysanthemum morifolium cultivar "Indianapolis white" EC50 (22 d) = 0.12 mg/l (in the nutrient solution; endpoint: chlorosis; nominal concentration). However, no PNECsoil can be derived from this result as no soil concentration is given.

Exposure

The production volume of this chemical at BASF, Germany, was more than 1000 tons in 2000. No information about the worldwide production volume is available.

The organic compound is used for the synthesis of pharmaceuticals and as a catalyst in the synthesis of polymers in the chemical industry. It is also used as a pH stabilizer. According to Swiss, Danish and Swedish Products Registers and the Hazardous Substances Data Bank, 2-diethylaminoethanol is contained in a large number of products. Some of them may be available to consumers.

Releases into the environment are likely to occur during the production and processing of 2-diethylaminoethanol as an intermediate, as well as from the use of the substance itself and use of products containing the substance.

Assuming worst case conditions, less than 9.5 kg of 2-diethylaminoethanol per day were released into the Rhine from an industrial site. During production and internal processing, less than 25 kg/a were emitted into the air from the same production site. From the reported use in consumer products, it can be concluded that most of the 2-diethylaminoethanol is released into wastewater, but part of it may also be released into the atmosphere.

RECOMMENDATION

The chemical is currently of low priority for further work.

RATIONALE FOR THE RECOMMENDATION AND NATURE OF FURTHER WORK RECOMMENDED

<u>Human Health</u> The chemical is currently of low priority for further work. Due to the corrosive potential, exposure to humans at the workplace and from consumer products has been regulated in the sponsor country. However, if this is not the case in other countries, further exposure assessment and, if necessary, risk assessment are recommended.

<u>Environment</u>: In addition to its use as chemical intermediate, European product registers indicate a wide dispersive use of 2-diethylaminoethanol. No information is available about the total production volume and about total environmental releases. However, the low aquatic toxicity, the low bioaccumulation potential and the ready biodegradability lead to the recommendation, that the chemical is currently of low priority for further work

SIDS Initial Assessment Report

1 IDENTITY

1.1 Identification of the Substance



Synonyms:	DEAE
	ethanol, 2-(diethylamino)- (8CI, 9CI)
	N,N -diethylethanolamine

1.2 Purity/Impurities/Additives

Substance type:	organic
Physical status:	liquid
Purity:	≥ 99.5 % (w/w)

1.3 Physico - Chemical properties

2-Diethylaminoethanol is a colorless – light yellowish organic flammable liquid with an amine-like odor (BASF AG, 2000). The hygroscopic substance is miscible with water in all proportions (BASF AG, 2000; Hazardous Substances Data Bank, 2001). The vapor pressure is about 1.8 hPa at 20 °C (BASF AG, 1984). A Henry's law constant of $3.16*10^4$ Pa*m³/mol at 25 °C was calculated via HENRYWIN v3.00 (BASF AG, 2001a). The measured partition coefficient (log Kow) was 0.21 (BASF AG, 1987). The density was determined to be 0.885 g/cm³ (BASF AG, 1985). Melting point and boiling point of the substance are – 68 °C (BASF AG, 1985) and 162-163 °C (at 1013 hPa; Beilstein, 2001), respectively.

2 **GENERAL INFORMATION ON EXPOSURE**

2-Diethylaminoethanol is produced by the thermal reaction of diethylamine with ethylenoxide.

The production volume of 2-diethylaminoethanol at BASF AG, Germany, exceeded 1000 tons in 2000. No information is available about the worldwide production volume; however, it is produced by at least two companies in the USA and at least two companies in Europe.

The substance was not imported into the European Union in 2000.

The compound is used for the synthesis of drugs in the pharmaceutical industry and as a catalyst for the synthesis of polymers in the chemical industry. It is also used as a pH stabilizer. 2-Diethylaminoethanol is listed in the provisional list of monomers and additives notified to the European Commission as substances which may be used in the manufacture of plastics intended to come into contact with foodstuffs (Ref.-No 48370; European Commission, Directorate D, D3; Introduction of "Synoptic Document"; 2002).

Additional applications are cited in the European Product registers.

According to the current Swedish Register, there were 119 products on the Swedish market containing 2-diethylaminoethanol in a total amount of 216 t/a. The main uses are rust preventives, various paints, pigments, polishing agents, paper manufacturing chemicals, anti-shell agents etc. (Swedish Products Register, 23 September 2002).

The Danish Product Register cited overall 392 products. These were estimated to account for about 15 tons/a. The types of product listed were process regulators, coloring agents, corrosion inhibitors, surface-active agents, cleaning/washing agents, cutting fluids, paint, laquers and varnishes, surface treatment (Danish Product Register, 26 Feb 2002).

According to the Swiss Product register, there were 375 registrations on the Swiss market of products containing 2-diethylaminoethanol, predominantly paints, varnishes and lacquers (approx. 50 % of the registrations), glues/adhesives, cement, filler or sealing compounds (approx. 10 %), car polishes (approx. 10 %), technical oils (approx. 10 %) and other uses like wood stains and inks. Most of these uses were for professional applications. Approx. 60 products, i.e. about 16 % of the total product spectrum, were available to consumers. Paints, lacquer, household cleaners and polishes (shoe, leather, car) accounted for more than 75 % of these products (Swiss Product Register, 2001).

In the Norwegian Product Register 243 products containing a total quantity of 27 t are registered. Use categories are paints and inhibitors.

The following uses are listed in the Hazardous Substances Data Bank: the manufacture of emulsifying agents and special soaps, chemical intermediates for petroleum and gas processing chemicals as well as for paints and lacquers, cosmetics, pharmaceuticals, crop protection agents, flocculants, surface coatings, textiles and fibers, procaine, chloroquinone, anti-rust additives, plastics, paper and leather chemicals (Hazardous Substances Data Bank, 2001).

Releases into the environment are likely to occur during the production and processing of 2diethylaminoethanol as an intermediate, as well as from the use of the substance itself and use of products containing the substance.

2-Diethylaminoethanol is measured in the influent and the effluent of the waste water treatment plant of BASF AG at regular intervals (24 h-mixing sample). Between the 1^{t} of January 2001 and the 30^{th} of June 2002 the concentration in the sewage as well as in the effluent was always found to be below the limit of quantitation (influent: 1 mg/l; effluent: 0.1 mg/l; BASF AG, 2002b). Additionally, the concentration in the effluent was always found to be below the limit of detection

(effluent: 0.025 mg/l) between the 1st and 30th of June in 2002. Based on the limit of detection and assuming worst case conditions, less than 9.5 kg of 2-diethylaminoethanol per day were released into the river Rhine in that period (BASF AG, 2002c).

In the 1970s 2-diethylaminoethanol was identified in an industrial effluent discharge in the USA (concentration: > 0.1 mg/L; Perry et al., 1978).

During production and internal processing at BASF AG, Ludwigshafen (Germany), less than 25 kg/a were emitted into the air in 2000 (BASF AG, 2002d).

Emission data from other production and processing sites was not available.

2.1 Environmental Exposure and Fate

For the uncharged molecule, modeling using Mackay, Level I indicates that water is the main target for environmental distribution (99.1 %; BASF AG, 2001b).

A soil adsorption coefficient (Koc) of 5.98 (log Koc = 0.78) was estimated for 2diethylaminoethanol via PCKOCWIN (BASF AG, 2001c). The Koc value suggests that adsorption to suspended solids is not to be expected. From the pKa-value of 9.87 it can be assumed that under environmental conditions the substance is available as cation. Therefore, binding of he substance to the matrix of soils with high capacities for cation exchange (e.g. clay) cannot be excluded. However, no data was available for ionic-ionic interactions in soil. The calculation of the Henry's law constant via HENRYWIN v3.00 yields a value of $3.16 \times 10^4 \text{ Pa} \times \text{m}^3/\text{mol}$ at 25 °C (BASF AG, 2001a). This low value and the complete water solubility of 2-diethylaminoethanol suggests that volatilization from water would not be an important factor in the environmental distribution process. In addition, due to the protonation of the substance under environmental conditions, the volatility is further reduced. In the air rapid degradation can be expected according to the half life calculated via AOP v1.87 (50 % after 3.9 hours; BASF AG, 2001d). Due to the measured and calculated partition coefficients (log Kow 0.21 and log Kow 0.047; BASF AG, 1987 and BASF AG, 2002a) an accumulation in organisms is not to be expected.

2-Diethylaminoethanol is readily biodegradable according to the results obtained in a test conducted according to OECD 301 A (95 % degradation after 22 days, 10d-window fulfilled; BASF AG, 2002f). Using the model Simpletreat, an elimination in sewage treatment plants of 87 % can be estimated (no volatilization, no adsorption, biodegradation rate constant $1 h^{-1}$). Based on the chemical structure of the substance hydrolysis is not likely to occur.

2.2 Human Exposure

Due to the information from European product registers, exposures to consumers and workers are likely. In the Swiss Product register, 375 products, among them 61 consumer products, containing 2-diethylaminoethanol are listed. The highest concentrations (up to 10%) are reported for cleaning agents, surface treatment, car and floor care products (Swiss Product Register, 2001). According to the current Swedish Product Register, there were 108 products, among them 6 consumer products, on the Swedish market containing 2-diethylaminoethanol (Swedish Product Register, 2002). Concentrations in the range between 0 and 2% may be found in paints, lacquers, varnishes and surface treatment products according to the Danish Product Register (Danish Product Register, 2002).

No data on human workplace exposure was available. 2-Diethylaminoethanol is produced by the thermal reaction of diethylamine with ethylenoxide in a closed system. There is no open handling of the product (BASF AG, 2002e). In the Safety Data Sheet of BASF AG, the workers should wear breathing protection if ventilation is inadequate and chemical resistant protective gloves and tightly

fitting safety goggles for personal protection. Body protection must be chosen depending on activity and possible exposure (BASF AG, 2000).

In Germany the current MAK value for 2-diethylaminoethanol is 5 ppm (24 mg/m^3) (DFG, 2002) and in the USA the current TLV value is 2 ppm (9,6 mg/m³) (ACGIH, 2002).

The following indoor air monitoring data was reported:

2-Diethylaminoethanol was used as a corrosion inhibitor in an open steam humidification system at the H.F. Johnson Museum of Cornell University (USA). Analyses of air samples from the museum environment were taken in January of 1983. The chemical was detected in only 2 air samples of the 14 collected. The measured DEAE concentrations were 0.04 and 0.05 mg/m³. In addition residues of condensed 2-diethylaminoethanol were identified on samples of plastic materials in the museum at concentrations of 30 mg/m². Condensation accumulation was found to be directly related to the time of deposition for airborne concentrations (NIOSH, Health Hazard Evaluation Report, 1983; Volent P. and Baer N.S., 1985).

During the winter season 2-diethylaminoethanol was measured in the indoor air in a study room at Battelle Columbus Division (USA). The average room concentration of 2-diethylaminoethanol at an average relative humidity of 42 % was about 0.6 ppb (approx. 2.9*10³ mg/m³) and at an average relative humidity of 61 % about 2.4 ppb (approx. 0.01 mg/m³). The primary fate of the amine that was introduced into the room air via steam humidification appeared to be condensation on surfaces (Edgerton et al., 1989).

This indoor concentration data is about 15 - 20 years old and may not be representative of, or comparable to, today's conditions. However, they indicate that inhalation is a possible route of exposure.

3 HUMAN HEALTH HAZARDS

3.1 Effects on Human Health

3.1.1 Toxicokinetics, Metabolism and Distribution

2-Diethylaminoethanol has been reported to be readily absorbed via the gastrointestinal tract in humans and rats (Rosenberg et al, 1949; Schulte et al., 1972). On the basis of the physico-chemical properties of a saturated aqueous 2-diethylaminoethanol solution, a skin penetration rate of 3.44 mg/cm² per hour was estimated for human skin, and therefore, the resulting body burden from exposure to 5 ml/m³ (the current MAK value) of 2-diethylaminoethalol by inhalation for 8 hours could potentially be increased by an additional three-fold factor via dermal absorption (Fiserova-Bergerova et al., 1990). However, this model was suspected to be too conservative and likely to overestimate percutaneous penetration (Guy and Russell, 1993). Absorption via inhalation has been mentioned (Toren, 1994), but the primary literature was not available for an assessment.

In a limited study with humans (Rosenberg et al, 1949), the plasma concentration peaked 3 hours after an oral administration of 5.6 g of 2-diethylaminoethanol-HCl, but was almost undetectable after 8 hours. About 25% of the 2-diethylaminoethanol was excreted unchanged in the urine within 48 hours. Similar excretion results were observed after intravenous administration. In the same publication it was reported that 2-diethylaminoethanol-HCl given to dogs by intravenous infusion (71 mg/kg bw), distributed rapidly. Three hours after infusion the level of 2-diethylaminoethanol was higher in the tissues examined (muscle, heart, brain, lung, liver and spleen) than in the plasma.

In a gavage study with rats (Schulte et al., 1972), ¹⁴C-labeled-2-diethylaminoethanol-HCl was reported to be rapidly absorbed into the blood stream (with a dose of 68 mg/kg the maximum concentration in the blood was reached in 30 minutes and with 679 mg/kg it was reached within 1 hour). Elimination occurred primarily via the kidney. Elimination via exhalation and the feces played only minor role. The kinetics of urinary elimination was affected by the dose. In this regard, by 6 hours after the application of a 679 mg/kg bw dose 40% was eliminated in the urine, and by 24 hours after application 58.5% was eliminated. When a 68 mg/kg dose was given, then after 6 and 24 hours 17.5% and 37.4% were excreted via the urine, respectively. In the experiment with 679 mg/kg 2-diethylaminoethanol, 90% of the test substance had been eliminated via the urine 10 days after treatment. Some radioactivity was still detectable in the urine 40 days after treatment. 2-Diethylaminoethanol was predominantly (> 60%) excreted unchanged over the first 96 hours. In the same period, the following metabolites were seen based on the recovery of radioactive compounds: 2-ethylaminoethanol (ca. 1%), phosphoric acid-mono-(2-diethylaminoethylester) (2-8%), diethylaminoacetic acid (ca. 10%) and the N-oxide of 2-diethylaminoethanol (ca. 15 - 19%). Incorporation into phospholipids was observed. In this study, autoradiography indicated that 2diethylaminoe thanol was widelv distributed throughout the body after gavaging. 2-Diethylaminoethanol was concentrated in the liver, reaching a maximum at 7 hours, but thereafter, it decreased. Initially, the central nervous system showed very low levels of activity, but by day 7 it had increased. For the oral dose of 679 mg/kg the biological half-life was 19 hours and for the 67.9 mg/kg dose it was 36 hours.

In a separate study ¹⁴C-labeled-2-diethylaminoethanol-HCl was given to rats by intravenous injection at doses of 2.9 μ mol/rat (ca. 1.94 mg/kg bw) (Michelot et al., 1981). Cumulative excretion of 19.9 and 42.2% of the radioactivity in the urine was observed after 24 and 48 hours, respectively. Additionally, 8.5 and 29.5% of the radioactivity was excreted via the feces during the same time interval. Excretion via the bile was only measured over the first 6 hours, and was reported to be 5%.

Since tertiary amines are poor substrates for monoamine oxidase, 2-diethylaminoethanol might presumably be metabolized by a P450 monoxygenase, or by a microsomal flavoprotein (reviewed in Cavender, 2001). Nitrosation would only be expected to occur very slowly in comparison to secondary amines (Mirvish, 1975).

Conclusion

2-Diethylaminoethanol was rapidly absorbed via the oral route. It is presumably absorbed by dermal and inhalation routes of administration. In the rat it was widely distributed to many tissues. It was primarily excreted unchanged via the urine in rats. Excretion via the feces was also observed in rats, but to a lesser extent. Urinary excretion was also reported in humans. The major metabolites in rats were reported to be diethylaminoacetic acid and diethyl-(2-hydroxyethyl)-amino-oxide.

3.1.2 Acute Toxicity

There were no studies available performed according to current guidelines, but studies were available which gave sufficient information to characterize the following endpoints.

Inhalation

 LC_{50} rat (inhalation): ca. 4600 mg/m³/4 hour; estimated by Haber's Rule from an Inhalation Hazard Test which used a highly enriched/saturated vapor exposure system at 20 °C, in which animals were exposed to 2-diethylaminoethanol vapor for 1, 3 and 8 hours (BASF AG, 1969). Clinical signs indicating severe irritation were noted, namely, attempts to escape, mucous membrane irritation, dyspnoea and gasping.

Dermal

In a non-guideline study with an exposure period of 4 days instead of 24 hours as prescribed by OECD Guideline 402, a dermal LD_{50} in guinea pigs was reported as ca. 885 mg/kg bw (Smyth and Carpenter, 1944). Normally, rats or rabbits are used for this endpoint, but the only available rabbit data was from secondary literature [LD_{50} rabbit (dermal): ca. 1100 mg/kg bw; from Smyth, 1964].

Oral

In a non-guideline study an LD_{50} in rats (oral) was determined to be ca. 1320 mg/kg bw (BASF AG, 1969). The clinical signs reported were described as apathy and dyspnea. This study was chosen as the key study since it was the one for which the most details are available. This LD50 is essentially the same as that reported by Smyth and Carpenter (1944). Higher values have been reported, but these were from tests conducted with the neutralized substance.

Conclusion

The LD₅₀ for the rat after oral administration was 1320 mg/kg bw. The main clinical signs described were apathy and dyspnea. After inhalation of vapors of 2-diethylaminoethanol an LC₅₀ of ca. 4600 mg/m³/4 hour was estimated in rats using Haber's rule. Severe signs of irritation were observed, e.g. mucous membrane irritation and dyspnea. A dermal LD₅₀ in guinea pigs was reported to be ca. 885 mg/kg bw.

3.1.3 Corrosiveness and Irritation

In guideline studies (OECD 404) 2-diethylaminoethanol was corrosive to the skin of rabbits after both occlusive and semi-occlusive 4 hour applications (BASF AG, 1982 and Potokar et al., 1985).

Several studies were available that examined irritation to the eye. They are difficult to assess since they did not follow guideline conditions, but they demonstrate that 2-diethylaminoethanol has the potential of being severely irritating to the eyes, or could cause serious damage to the eyes (e.g. in

one study, 50 μ l of the undiluted liquid was applied to the eye, and irreversible damage to corneal tissue was observed [staphyloma], as well as corrosion of the conjunctiva and eyelids; these finding were also not reversible after 8 days; BASF AG, 1969)

Conclusion

2-Diethylaminoethanol was corrosive to the skin of rabbits. Since its pH value was measured to be 11.5 (100 g/l) at 20 °C, these corrosive effects are not surprising. The potential for severe damage to occur to the eyes can be expected based on the animal studies available and on the pH.

3.1.4 Sensitisation

2-Diethylaminoethanol was tested for skin sensitization in guinea pigs using the method of Draize (TSCATS, 2/13/84) and the method of Magnusson and Kligman (Leung and Blaszcak, 1998; Nakamura et al., 1994) and was reported to be negative in all three studies. Taking all 3 studies into consideration, none of the 70 2-diethylaminoethanol-induced animals showed signs of sensitization when challenged with 2-diethylaminoethanol.

Conclusion

2-Diethylaminoethanol was not sensitizing to the skin of guinea pigs.

3.1.5 Repeated Dose Toxicity

A well documented 14 week inhalation study was available. In this study, 20 rats/dose/sex were exposed to 0, 11, 25 or 76 ppm (0, 0.053, 0.120 and 0.365 mg/l; or 0, 53, 120 and 365 mg/m3) of 2diethylaminoethanol for 6 h/day, 5 days/week for 14 weeks using a whole body exposure method (Hinz et al., 1992; Exxon, 1990) and are comparable to ca. 13, 29 and 88 mg/kg bw per day doses assuming 100% lung deposition and absorption. These concentrations were chosen based on a 2 week study. Half of the animals were terminated at the end of 14 week. Neurologic exams were preformed monthly using a modified Irwin Screen (Psychopharmacology, 13, 222-257, 1968) during the exposure period. The remaining animals were given a four week post-exposure recovery period prior to sacrifice. Full histological exams which included the gonads were conducted in the high dose group, but in the low and middle dose groups only the nasal cavity/turbinates (4 sections) were evaluated. No animals died as a result of exposure to 2-diethylaminoethanol. During exposure, dose-dependent transient signs of mild to moderate respiratory irritation (noises or rales) were noted. They usually cleared within one hour after exposure. In the high dose group, some animals continued to exhibit these signs overnight. Nasal discharge was observed at the beginning of the study, but this subsided as the study progressed. Corneal opacities were observed in control and 2diethylaminoethanol-treated animals. According to the authors aging F344 rats are genetically predisposed toward developing corneal lesions. Prolonged exposure to an alkaline compound such as 2-diethylaminoethanol could have accelerated an underlying predisposition toward corneal dystrophy. Since the opacity was thought to be due to a calcium precipitate, this problem may have also been exacerbated by the high vitamin D diet which would increase calcium absorption. In this regard, all rats in the high dose group had corneal opacities after 1 month of exposure, and most rats in the middle group had developed them by 2 months. By the end of the study all rats were confirmed to have corneal opacity. No outstanding effects on blood chemistry, urinalysis or neurobehavioural parameters were observed. Through the first 7 weeks of exposure the high dose group had a slight, but statistically significant decrease in body weight gain as compared to controls. Subsequently, the rate of growth paralleled the other groups, but the initial decrement was never regained. Mean body weights of the high concentration group never decreased more than about 7% from the controls. At week 14 there was a slight, but statistically significant increase in

the male liver and kidney weights in the high dose group (8.0 and 7.1% resp.), but histologic changes were not associated with these findings. Histomorphologic changes in nerve tissues were not observed. The low dose group was free of exposure-related histologic changes in the nasal cavities and turbinates, but changes were noted in the middle and high dose groups sacrificed in the 14th week of exposure [45% (50% M, 40% F) incidence in the middle and 95% (90% M, 100% F) in the high concentration group]. These changes consisted of an increased incidence and severity of focal hyperplasias alone or in association with squamous metaplasia of the respiratory epithelium, and multi-focal mixed infiltrations of inflammatory cells in the nasal mucosa. These changes were most evident in the anterior sections of the nasoturbinates and on the lateral wall of the nasal cavity. In the high dose groups after the 4 week recovery period were similar to those seen in the 14 week rats, however, the incidence of focal hyperplasia with squamous metaplasia was decreased. The incidence of focal hyperplasia alone, infiltrations of inflammatory cells and goblet cell were comparable to what was noted at 14 weeks.

<u>Summary</u>: This study indicated that 2-diethylaminoethanol lacked systemic toxic properties, and the point of contact was the site of action, namely, the upper respiratory tract and the eyes. Since no systemic toxicological effects were observed, the No Observed [Adverse] Effect Concentration (NO[A]EC) for systemic toxicity was 0.365 mg/l (365 mg/m³, 76 ppm), which was the highest dose tested. The NO[A]EC for local toxicity, based on the lack of an effect observed in the nasal cavity was 0.053 mg/l (53 mg/m³ or 11 ppm, rounded off to 10 ppm). The noises or rales at this concentration can be considered an adaptive effect, but not an adverse effect, since no histological changes were observed at this concentration. However, since an effect was seen at the lowest concentration, a NOEC was not reached. The Lowest Observed [Adverse] Effect Concentration (LO[A]EC) was the first dose where nasal lesions were observed, namely, 0.12 mg/l (120 mg/m³ or 25 ppm).

NO[A]EC, rat (inhalation) 14 weeks, systemic tox.: 0.365 mg/l (365 mg/m³ or 76 ppm)

NO[A]EC, rat (inhalation) 14 weeks, local toxicity: 0.053 mg/l (53 mg/m³ or 10 ppm)

LO[A]EC, rat (inhalation) 14 weeks, local toxicity: 0.12 mg/l (120 mg/m³ or 25 ppm)

Several repeated dose toxicity studies were performed in the 1960's, but their quality was limited in part due to a lack of detail or poor design. In one of these studies (TSCATS, 8/02/90), rats were fed 2-diethylaminoethanol-HCl at doses of 200, 500 and 1000 (10,000) ppm of the free base (i.e. ca. 11, 25, 50-400 mg/kg bw/day, respectively) for 2 years. The high dose group was progressively increased to 10,000 ppm (ca. 400 mg/kg/d). For 2-diethylaminoethanol treatment 35 rats/sex/group were used, and in the control there were 60 rats/sex/group. Males were reported to have low incidences of testicular atrophy, namely 0/34 (0%), 3/18 (17%), 2/17 (12%) and 4/15 (27%) in the control, 200 ppm, 500 ppm and 1000 (10,000) ppm dose groups, respectively. The atrophy sometimes occurred bilaterally. This finding was considered insignificant for the following reasons:

- 1) It was not dose-dependent;
- 2) Similar findings were not seen in 6 month old males (a more appropriate time point to assess the induction of gonadal lesions), and only one case was seen in 12 month old males (200 ppm dose group) from the same study;
- 3) It is not an unusual finding in aging rats (Glaister, 1986);
- 4) Anorexia was also reported in this study, but data on individual animal weight was not given except at the final sacrifice. Thus, anorexia could have been a / the causative factor for testicular atrophy. Furthermore, in the last 12 weeks of the study, the high dose male body

weight was on average 8.6% lower that the controls, and the terminal body weight of this group was significantly (11 %, p < 0.05) lower than controls;

5) And finally, the historical control data for testicular atrophy in Charles River albino rats was not available, but based on a comment by the sponsor, similar incidences were reported in control animals. No historical control data from the 1960s was available. But, in a study conducted nearly 35 years later, it was reported that the incidence of **e**sticular atrophy found in control 31 week old Charles River CrJ: (SD) rats was 20 % (Sugimoto et al., 2000). According to Charles River, the type of rat used in the 1960s was probably a Sprague-Dawley (personal communication, P.A. Mirley, 24 June 2002).

Thus, the testicular finding in this study with 2 year old Charles River albino rats is not surprising. The lack of a finding in the 34 control males in this could most likely be a result of the randomness of the group assignments.

Another study was conducted with dogs in the 1960s in which 2-diethylaminoethanol-HCl was given via feed at doses of ca. 20, 40, 200 (80) and 400 mg 2-diethylaminoethanol/kg/day, [i.e. 500, 1000, 5000 (2000) and 10000 ppm] (TSCATS, 8/02/90). According to the report, "weakness, tremors, convulsions and ataxia" were observed, however, this occurred at doses where animals were dying, namely at 5000(2000) and 10000 ppm. In a lower dose (1000 ppm based on the free base) "tremors and/or shaking of the head from side to side" were described. This data is difficult to assess due to the limited nature of the report. Furthermore, in the 2 year study in rats already mentioned, no such findings were seen in rats treated with similar doses. Cerebellar changes were observed in all males and one female of the 5000 (2000) ppm dose group.

In another study with rats, they were given 2-diethylaminoethanol neutralized with HCl via drinking water for up to six months at doses resulting in 2-diethylaminoethanol levels of ca. 150 and 300 mg/kg bw/day (Cornish, 1965). No significant adverse changes were observed, although a slight increase in kidney weight was described in both 2-diethylaminoethanol-treated groups, and decreased body weight was seen in the high dose group.

Conclusion

Repeated exposure of rats to 2-diethylaminoethanol vapors (up to 0.365 mg/l) for 14 weeks caused local toxicity (irritation) at the site of contact, namely, the upper respiratory tract and the eyes; however, systemic toxicity was not observed (NOAEC, systemic toxicity, 0.365 mg/l; i.e. 365 mg/m³ or 76 ppm). After inhalation exposure, the main finding described was respiratory irritation which led to noises called rales and irritation of the eyes. The LOAEC for local toxicity (irritation) to the respiratory tract was 0.120 mg/l (120 mg/m³ or 25 ppm). The NOAEC for local toxicity was 0.053 mg/l (53 mg/m³ or 10 ppm) based on a lack of histopathological effects in the nasal cavity at this dose. However, since an effect (rales) was seen at the lowest concentration a NOEC was not reached.

3.1.6 Mutagenicity

In vitro Studies

2-Diethylaminoethanol was tested at doses of up to 5000 µg/plate in *Salmonella typhimurium* strains TA98, TA100, TA1535, TA1537 and TA1538, both with and without metabolic activation using guideline and non-guideline conditions and consistently yielded negative results (BASF AG, 1989; Life Science Research, 1991a; Zeiger et al, 1987). In a guideline assay using the HPRT locus of V79 Chinese hamster cells, no evidence for mutagenic activity was seen when tested up to 3500 µg/ml (Life Science Research, 1991b). It was also negative in a DNA damage test using *E. coli* which used doses up to 3500 µg/ml (Life Science Research, 1991b).

In vivo Studies

In a detailed and well conducted study 2-diethylaminoethanol was tested for its ability to induce micronuclei in bone marrow erythrocytes in mice using doses up to 500 mg/kg bw under guideline conditions and was found to be negative (Life Science Research, 1991d). The report also indicates that the highest dose tested was adequate since animals showed a hunched posture, piloerection, rales, irregular respiration, a swollen abdomen and one animal was sacrificed *in extremis*. Data from the preliminary test indicated that the test substance can reach the bone marrow.

Conclusion

2-Diethylaminoethanol gave no evidence of *in vitro* mutagenic activity or *in vivo* clastogenic potential.

3.1.7 Carcinogenicity

In a study from the 1960's (TSCATS, 8/02/90) which does not meet the guidelines for carcinogenicity studies of today (see below), rats received 2-diethylaminoethanol-HCl at dietary concentrations of 200, 500 and 1000 (10,000) ppm of the free base (ca. 11, 25, 50-400 mg/kg bw/day, respectively) for two years. The high dose group was progressively increased to 10,000 ppm (ca. 400 mg/kg/d). For the 2-diethylaminoethanol treated groups 35 rats/sex/group were used, and the control group consisted of 60 rats/sex. Ten animals/sex from controls and the high dose group were examined histologically at the end of the 2 years. The following tumors were seen in the fourty animals which were examined completely: pituitary adenomas in 9 animals each in the control and high dose group group; mammary gland fibromas, adenomes or fibroadenomes in 8 control and 4 high dose group females; miscellaneous tumors which included one ganglioneuroma, one pheochromocytoma and two renal embryomas in the control group; adrenal cortical adenomas in one control and three high dose group females; and one pancreatic duct adenoma, one hepatoma and three granulosa cell tumors in the high dose group. Regarding the limitations of the study, in the 1960s international guidelines for cancer studies were not available. Even so, the following limitations should be noted: the number of animals was low, a maximum tolerated dose was not achieved and the rational for the doses chosen was not given.

Conclusion

2-Diethylaminoethanol was not carcinogenic to rats by feed in a limited 2-year study from the 1960s. While this data is limited, the lack of a carcinogenic effect of 2-diethylaminoethanol is supported by the negative data for genotoxicity.

3.1.8 Toxicity for Reproduction

Studies specifically designed to assess reproductive toxicity were not available for an assessment. However, an investigation of changes in the weight and/or morphology of the reproductive organs in a repeated dose toxicity studies can also be used to assess reproductive toxicity. Typically, a 28 day or 90 day study can be used for such an assessment (ECETOC, 2002). Several repeated dose studies were available for 2-diethylaminoethanol (see section 3.2.4); however, the best documented and most appropriate study was a 14 week repeated dose toxicity study which performed gonadal examinations.

In a range finding study for the 14 week inhalation study detailed in section 3.2.4, rats were exposed to 301 ppm $(1.44 \text{ mg/l or } 1440 \text{ mg/m}^3)$ of 2-diethylaminoethanol for 9 days, 6 h/day over a 2 week period (Hinz et al., 1992). At this concentration animals lost weight throughout the study

and a high rate of mortality occurred. In addition, undersized spleens, thymuses and gonads were reported in the surviving animals. This effect on the gonads should be considered as the result of secondary toxicity, given that mortality also occurred at this dose. In the main study (see section 3.2.4 for details), i.e. the 14 week inhalation study, the testes were weighed prior to fixation and epididymides, testes and ovaries were prepared and stained for histopathological assessment. No signs of gonadal toxicity were observed in either sex (Hinz et al., 1992). Since no effects on the gonads were noted at the highest concentration tested in the 14 week study (the two lower concentrations were not examined), the NO[A]EC for gonadal toxicity was the highest concentration tested, i.e. 0.365 mg/l (365 mg/m³, i.e. 76 ppm, or ca. 88 mg/kg/day assuming 100 % lung deposition and absorption).

In an older 2 year feed study in rats (see section 3.2.4 for details) a low incidence of gonadal degeneration was reported, but only (with one exception) in 2 year old males with an incidence of 0/34 (0%), 3/18 (17%), 2/17 (12%) and 4/15 (27%) in the 0, 200, 500 and 1000 (10,000) ppm dose groups (TSCATS, 8/02/90). Unilateral and bilateral cases were seen. Since testicular atrophy is not an unusual finding in aging rats (Glaister 1986), and since there was not a clear dose-dependency in their occurrence, these results were not considered to be substance-related. Furthermore, during the time frame where the model is suitable for detecting testicular toxicity, no changes were observed (see also discussion in section 3.2.4).

Conclusion

Based on the results of a well-documented 14 week animal study, the inhalation of 365 mg/m^3 (76 ppm) 2-diethylaminoethanol did not cause any adverse effects on the reproductive organs in rats. This data suggests that 2-diethylaminoethanol does not cause any adverse effects on the reproductive system under the conditions tested.

3.1.9 Developmental Toxicity/Teratogenicity

In a guideline-like study, pregnant rats were exposed to vapors of 2-diethylaminoethanol at concentrations of 0, 33, 66 and 100 ppm (i.e. 0.160, 0.320 and 0.486 mg/l; or 0, 160, 320 and 486 mg/m³) for 6 h/day on gestational days (GD) 6 - 15, and then the dams were sacrificed on GD 21 (Exxon, 1991; Leung and Murphy, 1998). These concentrations are comparable to ca. 38, 76 and 116 mg/kg bw per day doses, assuming 100 % lung deposition and absorption. The doses were chosen based on toxicity observed in a range finding study. No deaths were observed in the study. Maternal toxicity was seen in the high dose group and included reduced body weight (6%) on GD day 15 and reduced body weight gain (52%) during the entire exposure period (GD 6-15). Dry rales were observed in up to one third of the animals over GD 11 to 21 in the high dose group. Decreased body weight gain was also observed in the middle dose group during GD 12 to 15. Statistically significant (p < 0.01) decreases in maternal food consumption were observed during the exposure period in the middle and high dose group and during the post-exposure period in the high dose group. No treatment-related effects were observed in gestational parameters, including pre- and post-implantation loss or sex ratio. Mean fetal body weights in the treated groups were similar to control. There was no increase in the incidence of total malformations (external, visceral, or skeletal) or individually by category. The incidence of a single developmental variation (hypoplastic bones of the forepaw) was significantly decreased in the high dose group when compared to control. A statistically significant (p < 0.05) increasing dose-trend in the incidence of advanced ossification of the hindpaw was reported (9.7, 9.8, 12.6, 16.9 % control to high concentration group), but it was not significant when analyzed on a per fetus or per litter basis. Since this increased incidence of advanced hindpaw ossification was higher than expected in both treated and control groups compared to the laboratories' historical control range (0 - 2.3%), this finding was not considered treatment-related or biologically relevant.

The no-observed-adverse-effect concentration (NO[A]EC) for maternal toxicity was 33 ppm and for developmental toxicity the NO[A]EC was 100 ppm.

NO[A]EC, rats, maternal toxicity:	0.160 mg/l (160 mg/m ³ ; 33 ppm)
NO[A]EC, rats, developmental toxicity:	0.486 mg/l (486 mg/m ³ ; 100 ppm)

In an embryotoxicity screening study 2-diethylaminoethanol was administered by gavage to 5 pregnant female Crl:CD(SD)Br rats/dose at doses of 0, 10, 30, 100 and 250 mg/kg bw on days 0 to 11 of gestation (TSCATS, 11/12/97). The dams were sacrificed on day 12 of gestation. The only treatment-related clinical finding in the study was rales at the 250 mg/kg bw dose, indicating that a highly irritating dose was achieved. No internal findings were seen macroscopically. Body weights, body weight gain, food consumption and liver and kidney weights were not effected by treatment. No treatment-related microscopic findings were seen in the liver and kidneys of the 250 mg/kg dose group. In the 250 mg/kg bw dose group post implantation loss was increased by 16.6% (S.D. 20.90), and the number of viable embryos was decreased by 15 % (83.4 % in the 250 mg/kg group versus 98.6% in controls). The increase in postimplantation loss and the decrease in live litter size in this group was predominantly due to one female with nine early resorptions (52.9%). Intrauterine parameters were unaffected by treatment in the 10, 30 and 100 mg/kg dose groups. According to the report, the NOEL was 100 mg/kg bw for maternal toxicity and embryotoxicity. The basis for this was the observation of rales in 2/5 dams of the 250 mg/kg dose and the lack of this finding in the 100 mg/kg dose. Thus, the LOEL for the dams was 250 mg/kg bw. However, it is not clear if a true maternally toxic dose was achieved on the basis of rales. Histologic examinations were limited to the kidneys and liver. Due to the low number of animals in the study, the assignment of clear substance-related embryotoxic effect at 250 mg/kg bw is also difficult since the increase in postimplantation loss and the decrease in live litter size was due to one animal with 9 resorptions.

Conclusion:

In pregnant rats even the highest concentration tested of 0.486 mg/l (486 mg/m³ or 100 ppm), which already produced maternally toxic effects, did not lead to adverse developmental effects.

3.1.10 Experience with human exposure

An odor threshold of 0.011 ppm has been reported (Amoore and Hautala, 1983). A laboratory worker was briefly exposed (approx. 30 sec.) to approx. 100 ppm 2-diethylaminoethanol, which caused nausea and vomiting. No irritation of eyes or throat was noted. (Cornish, 1965). Subjects exposed to 2-diethylaminoethanol vapor by humidified air in office buildings complained about eye, nose and throat irritation, dizziness, nausea and vomiting. Also, several cases of asthma were observed (Fannick et al, 1983, Hills and Lushniak, 1989, Gadon et al., 1994); however, the symptoms were more consistent with a reactive airway dysfunction syndrome than with an allergic respiratory reaction. Detectable amounts of 2-diethylaminoethanol were 0.05 and 0.04 mg/m³. Since 2-diethylaminoethanol has a low vapor pressure and was detected on surfaces, skin contact with surfaces was a possible route of absorption (Fannick et al., 1983).

3.2 Initial Assessment for Human Health

2-Diethylaminoethanol was rapidly absorbed orally. It is presumably absorbed by dermal and inhalation routes of administration. In the rat it was widely distributed to many tissues. It was primarily excreted unchanged via the urine in rats. Excretion via the feces was also observed in rats, but to a lesser extent. Urinary excretion was also reported in humans. The major metabolites in rats were reported to be diethylaminoacetic acid and diethyl(2-hydroxyethyl)-amino-oxide.

2-Diethylaminoethanol had the following acute toxic effects in mammals:

LD50 rat (oral): 1320 mg/kg bw. The clinical signs reported were described as apathy and dyspnea.

LC50 rat (inhalation): ca. $4600 \text{ mg/m}^3/4$ hour (estimated by Haber's rule). Severe signs of irritation were observed, e.g. mucous membrane irritation and dyspnoea.

LD50 guinea pig (dermal): ca. 885 mg/kg bw [LD50 rabbit (dermal): ca. 1100 mg/kg bw, only available as a secondary citation]

2-Diethylaminoethanol was corrosive to the skin of rabbits. Since the pH was measured to be 11.5 (100 g/l) at 20°C, the corrosive effects are not surprising. The potential for severe damage to the eyes can be expected based on the animal studies available and on the pH.

2-Diethylaminoethanol was not sensitizing to the skin in studies with guinea pigs.

Repeated exposure to 2-diethylaminoethanol vapors up to 365 mg/m^3 for 14 weeks caused local toxicity (irritation) at the site of contact in rats, namely, in the upper respiratory tract and the eyes.

This study indicated that 2-diethylaminoethanol lacked systemic toxic properties up to 365 mg/m^3 . Since no systemic toxicological effects were observed, the NO[A]EC for systemic toxicity was the highest dose tested, i.e. 0.365 mg/l (76 ppm, or 365 mg/m^3). The NO[A]EC for local toxicity, based on the lack of observed effects in the nasal cavity/turbinates, was 0.053 mg/l (11 ppm, rounded off to 10 ppm, or 53 mg/m^3). A NOEC could not be derived since respiratory noises (rales) were observed in all concentration groups.

2-Diethylaminoethanol gave no evidence for *in vitro* mutagenic activity or *in vivo* clastogenic potential.

Based on the results of a well-documented animal study, 2-diethylaminoethanol did not cause any adverse effects on the reproductive organs of rats when administered by inhalation at concentrations of up to 365 mg/m^3 (0. 365 mg/l or 76 ppm) for 14 weeks.

In pregnant rats even the highest concentration tested of 0.486 mg/l (486 mg/m³ or 100 ppm), which already produced maternally toxic effects, did not lead to adverse developmental effects.

2-Diethylaminoethanol was not carcinogenic to rats when given by feed for 2 years in a limited 1960's study (tested up to ca. 50 - 400 mg/kg/d). While this data is limited, the lack of a carcinogenic effect of 2-diethylaminoethanol is supported by the negative data for genotoxicity.

An odor threshold of 0.011 ppm (approx. 0.053 mg/m^3) has been reported. In a laboratory worker short-time exposure to approx. 100 ppm (480 mg/m^3) 2-diethylaminoethanol caused nausea and vomiting. Subjects exposed to 2-diethylaminoethanol vapor by humidified air in office buildings complained about eye, nose and throat irritation, dizziness, nausea and vomiting. Also several cases of asthma were observed; however, the symptoms were more consistent with a reactive airway dysfunction syndrome than with an allergic respiratory reaction. In one case detectable amounts of 2-diethylaminoethanol were 0.05 and 0.04 mg/m³.

4 HAZARDS TO THE ENVIRONMENT

4.1 Aquatic Effects

The most sensitive studies available were considered to evaluate the aquatic toxicity of 2diethylaminoethanol.

Acute Toxicity Test Results

Fish

In a study with *Leuciscus idus*, following the German DIN 38 412, 4 concentrations, from 100-1000 mg/l (nominal) plus control and pH adjusted 1000 mg/l group, were tested. An LC50 (96 h) of 147 mg/l (nominal; geometric mean of LC₀ at 100 mg/l and LC100 at 215 mg/l) was calculated. The toxic effect may be, in part, due to the high pH of the non-neutralized test solutions, since the pH adjusted 1000 mg/l concentration group tolerated the substance for 96 h without mortality (BASF AG, 1987). This assumption is also confirmed by the result of other fish tests. Geiger et al. (1986) found a LC50 (96 h) of 1780 mg/l (measured) for *Pimephales promelas* exposed in a flow-through system. The pH of the test solution was adjusted to that of lake water with HCl.

Invertebrates

A test following Directive 79/831/EEC, C2 with Daphnia magna with 7 nominal concentrations ranging from 7.8 - 1000 mg/l, resulted in an EC50 (48 h, immobilisation) of 83.6 mg/l (BASF AG, 1988a). As the pH value at the concentration at which immobilization of the daphnids occurred was in the range of 8.6 to 10.7 it cannot be excluded that the toxicity was in part due to pH effects. This fact could be confirmed by a further study with *Daphnia magna* (Atofina, 1993) where an EC50 (48 h) of 165 mg/l was found using pH adjusted test solutions.

Algae

Acute Toxicity to Scenedesmus subspicatus was determined in a study following DIN 38 412 part 9, with 7 nominal concentrations ranging from 2-320 mg/l. The E_rC_{50} for growth rate (72 h) was 44 mg/l and the NOEC 5 mg/l; corresponding values for the endpoint biomass were 30 mg/l and 5 mg/l respectively (BASF AG, 1988b). As the reported pH of the test solutions was in the range of 7 to 8.5 it can be concluded that the observed effects are due to the inherent toxicity of the test substance itself and not due to pH effects.

Chronic Toxicity Test Results

No chronic aquatic toxicity data are available.

Toxicity to Microorganisms

The EC_{20} (30 min) for inhibition of the respiration of activated sludge (domestic) was >1000 mg/l (nominal concentration; BASF AG, 1994b).

4.2 Terrestrial Effects

In a study with the terrestrial plant *Chrysanthemum morifolium* cultivar "Indianapolis white" an EC_{50} (22 d, nominal concentration) of 0.12 mg/l in the nutrient solution for the endpoint chlorosis was obtained (Horst et al., 1983).

4.3 Initial Assessment for the Environment

Distribution modeling predicts water to be the main target compartment for 2-diethylaminoethanol. The substance tends not to accumulate in biota (log Kow = 0.21, measured). The calculated log Koc of 0.78 suggests that adsorption to suspended solids is not to be expected. From the pKa-value of 9.87 it can be assumed that under environmental conditions the substance is available as cation. Therefore, binding of the substance to the matrix of soils with high capacities for cation exchange (e.g. clay) cannot be excluded. However, no data was available for ionic-ionic interactions in soil. 2-Diethylaminoethanol was readily biodegradable in a test conducted according to OECD 301 A with domestic sludge (95% degradation after 22 days, 10d-window was fulfilled). Based on the chemical structure of the substance hydrolysis is not likely to occur. 2-Diethylaminoethanol entering the atmosphere is rapidly degraded by reaction with photochemically produced hydroxyl radicals ($t_{1/2} = 3.9$ h).

The following aquatic effect concentrations are available:

Leuciscus idus LC50 (96 h) = 147 mg/l (nominal concentration). The toxic effect may be (partly) due to the high pH of the non-neutralized test solutions, since the pH adjusted 1000 mg/l dose group tolerated the substance for 96 h without mortality.

Pimephales promelas LC50 (96 h) = 1780 mg/l (measured concentration, adjustment of pH).

Daphnia magna: EC50 (48 h) = 83.6 mg/l (nominal concentration) (toxicity due to pH effects cannot be excluded).

Daphnia magna: EC50 (48 h) = 165 mg/l (nominal concentration, adjustment of pH)

Scenedesmus subspicatus: ErC50 (72 h) = 44 mg/l, EbC50 (72 h) = 30 mg/l, NOEC = 5 mg/l for both endpoints.

Compared to daphnia and fish, aquatic plants are most sensitive to 2-ethylaminoethanol.

Using the aquatic toxic effect on the most sensitive species, *Scenedesmus subspicatus*, for the endpoint growth rate, a PNEC_{aqua} of 44 μ g/l is derived by applying an assessment factor of 1000.

No PNECsoil can be derived from the available terrestrial study with *Chrysanthemum* as no soil concentration is given (EC50 (22 d) = 0.12 mg/l; chlorosis).

5 RECOMMENDATIONS

Human Health:

The chemical is currently of low priority for further work. Due to the corrosive potential, exposure to humans at the workplace and from consumer products has been regulated in the sponsor country. However, if this is not the case in other countries, further exposure assessment and, if necessary, risk assessment are recommended.

Environment:

The chemical is currently of low priority for further work. In addition to its use as chemical intermediate European product registers indicate a wide dispersive use of 2-diethylaminoethanol. No information is available about the total production volume and about total environmental releases. However, the low aquatic toxicity, the low bioaccumulation potential and the ready biodegradability lead to the recommendation that the chemical is currently of low priority for further work.

6 **REFERENCES**

ACGIH (2002) Threshold Limit Values for Chemical Substances and Physical Agents, Biological Exposure Indices 2002. ACGIH worldwide, Cincinnati, USA

Amoore JE and Hautala E (1983), Odor as an aid to chemical safety: Odor thresholds compared with threshold limit values and volatilities for 214 industrial chemicals in air and water dilution. J Appl Toxicol 3, 272 - 290

Atofina (1993), Project No. 159142, Exxon Biomedical Sciences, Inc., 14 Dec 1993

BASF AG (1969), Department of Toxicology, unpublished studies (XVIII/320), 27 Jan 1969

BASF AG (1982), Department of Toxicology, unpublished studies (82/18), 26 Aug 1982

BASF AG (1984), ZET/FC, unpublished data, report 184.0448.1, 08 Nov 1984

BASF AG (1985), Physikalisch-chemische Konstanten, unpublished study, report BRU 85.219, 16 Dec 1985

BASF AG (1987), Analytisches Labor, unpublished report, report BRU 87.262, 18 Dec 1987

BASF AG (1987), Department of Toxicology, unpublished studies (87/267), 9 Nov 1987

BASF AG (1988a), Department of Ecology, unpublished data, 1/1130/2/87-1130/87, 15 Jan 1988

BASF AG (1988b), unpublished data, 3-BASF-ökolimna-12/88/-026, report of ÖKOLIMNA Gesellschaft für Ökologie und Gewasserkunde mbH

BASF AG (1989), Department of Toxicology, unpublished studies (88/956), 8 Mar 1989

BASF AG (1994a), Department of Ecology, unpublished data, report No. 94/0979/10/1, 1994

BASF AG (1994b), Department of Ecology, unpublished data, report No. 94/0979/08/1, 1994

BASF AG (2000), Safety Data Sheet N,N-Diethylethanolamine, 05 July 2000

BASF AG (2001a), Department of Product Safety, unpublished calculation (Henry's law constant; HENRYWIN v3.00), 10 July 2001

BASF AG (2001b), Department of Product Safety, unpublished calculation (Mackay Level I V 2.11 Model), 10 July 2001

BASF AG (2001c), Department of Product Safety, unpublished calculation (Koc; PCKOCWIN v1.63), 10 July 2001

BASF AG (2001d), Department of Product Safety, unpublished calculation (Photodegradation; AOP v1.87), 09 July 2001

BASF AG (2002a), Department of Product Safety, unpublished calculation (Log Kow; KOWWIN v1.66), 27 June 2002

BASF AG (2002b), GUU/WI, unpublished data, effluent concentration between 01 Jan 2001 – 30 June 2002, 04 July 2002

BASF AG (2002c), GUU/WU, unpublished data, effluent concentration between 01 June - 30 June 2002, 09 July 2002

BASF AG (2002d), GUU/LE, unpublished data (emission of 2-diethylaminoethanol in the German Emission Register 2000), 12 July 2002

BASF AG (2002e), GUP/PC, unpublished data (information about production procedure), 08 August 2002

BASF AG (2002f) Department of Product Safety, unpublished study, 02/0430/21/1, 01.10.2002

Beilstein (2001), Handbook of Organic Chemistry

Cavender FL (2001), Patty's Toxicology, 5th Edition, volume 4, Aliphatic and Alicyclic Amines (chapter 56), (eds. Bingham E, Cohrssen B, and Powell CH): 683 - 815, New York

Cornish HH (1965), Oral and Inhalation Toxicity of 2-Diethylaminoethanol, Am Ind Hyg Assoc J 26:479 - 484

Danish Product Register (2002), 26 Feb 2002

DFG (2002) List of MAK and BAT values 2002. Commission for the Investigation of Health Hazards of Chemical Compounds in the Work Area, No. 38. Wiley-VCH, Weinheim, Germany

Edgerton SA, Kenny DV, Joseph DW (1989), Determination of amines in indoor air from steam humidification, Environ Sci Technol 23 (4): 484 - 488

European Commission (2002), Directorate D, D3; Introduction of "Synoptic Document"; Provisional lists of monomers and additives notified to European Commission as substances which may be used in the manufacture of plastics intended to come into contact with foodstuffs, updated to 15 January 2002

Exxon Biomedical Sciences (1990), Project Number 208618, Sponsored research by Synthetic Organic Chemicals Manufacturing Association, 3 May 1990

Exxon Biomedical Sciences (1991), Project No. 108634, Sponsored research by Synthetic Organic Chemicals Manufacturing Association, 11 Sept 1991

Fannick N, Lipscomb J, McManus K (1983), Health Hazard Evaluation Report No. HETA-83-020-1351, NIOSH, Cincinnati

Fiserova-Bergerova V, Pierce JT and Droz PO (1990), Dermal Absorption Potential of Industrial Chemicals: Criteria for Skin Notation, Am J Ind Med 17: 617 - 635

Gadon ME, Melius JM, McDonald GJ, Orgel D (1994), New-onset asthma after exposure to the stream system additive 2-diethylaminoethanol. J Occup. Med. 36, 623 - 626

Geiger DL, Poirier SH, Brooke LT and Call DJ (1986), Acute Toxicities of Organic Chemicals to Fathead Minnows (Pimephales promelas), Vol. 3, Center for Lake Superior Environmental Studies, University of Wisconsin, Superior, WI: 149 - 150

Glaister JR (1986), Principles of Toxicological Pathology, Laboratory-animal pathology: 131 - 203, Taylor and Francis, London

Guy RH and Potts RO(1993), Penetration of Industrial Chemicals Across the Skin: A Predictive Model, Am J Ind Med 23: 711 - 719

Hills B, Lushniak B (1989), Health Hazard Evaluation Report No. HETA-89-057-2003, NIOSH, Cincinnati

Hinz JP, Thomas JA and Ben-Dyke R (1992), Evaluation of the Inhalation Toxicity of Diethylethanolamine (DEEA) in Rats, Fundam. Appl Toxicol 18: 418 - 424

Horst RK, Kawamoto SO, Schumann GL and Dietert MF (1983), Chlorosis in healthy and viroid-infected plants exposed to the steam additive diethanolamine; Scientia Horticulturae 19: 1 - 8

HSDB (2001), Hazardous Substances Data Bank, 2001

Leung HW and Blaszcak DL (1998), The Skin Sensitization Potential of Four Alkylalkanolamines, Vet Hum Toxicol 40: 65 - 67

Leung HW and Murphy SR (1998), Developmental Toxicity Study in Sprague-Dawley Rats by Whole-body Exposure to N,N -Diethylethanolamine Vapor, J Appl Toxicol 18: 191 - 196

Life Science Research (1991a), Report No. 91/SHG001/0251, Diethylaminoethanol: Assessment of mutagenic potential in histidine auxotrophs of Salmonella typhimurium (the Ames test), Sponsored research by Synthetic Organic Chemicals Manufacturing Association, 2 July 1991

Life Science Research (1991b), Report No. 91/SHG002/0302, Diethylaminoethanol: Investigation of mutagenic activity at the HPGRT locus in a Chinese hamster V79 cell mutation system, Sponsored research by Synthetic Organic Chemicals Manufacturing Association, 2 July 1991

Life Science Research (1991c), Report No. 91/SHG004/0263, Diethylaminoethanol: Assessment of its ability to cause lethal DNA damage to strains of Escherichia coli, Sponsored research by Synthetic Organic Chemicals Manufacturing Association, 2 July 1991

Life Science Research (1991d), Report No. 91/SHG003/0321, Diethylaminoethanol: Assessment of clastogenic action on bone marrow erythrocytes in the micronucleus test, Sponsored research by Synthetic Organic Chemicals Manufacturing Association, 2 July 1991

Michelot J, Madelmont JC, Jordan D, Mornex R and Meyniel G (1981), Metabolism of adiphenine. I. Absorption, distribution and excretion in rats and mice, Xenobiotica 11: 123 - 130

Mirvish SS (1975), Formation of N-Nitroso Compounds: Chemistry, Kinetics, and in Vivo Occurrence, Toxicol Appl Pharmacol 31: 325 - 351

Nakamura A, Momma J, Sekiguchi H, Noda T, Yamano T, Kaniwa MA, Kojima S, Tsuda M and Kurokawa Y (1994), A new protocol and criteria for quantitative determination of sensitization potencies of chemicals by guinea pig maximization test, Contact Dermatitis 31: 72 - 85

NIOSH (1983), Health Hazard Evaluation Report, HETA 83-020-1351, Aug 1983

Perry DL, Chuang, CC, Jungclaus, GA and Warner, JS (1978), Identification of Organic Compounds in Industrial Effluent Discharges, Battelle Columbus Labs, OH, prepared for EPA, Washington, DC, Office of Toxic Substances, PB-291 900, EPA-560-6/78-009; Nov 1978

Potokar M, Grundler OJ, Heusener A, Jung R, Mürmann P, Schöbel C, Suberg H and Zechel HJ (1985), Studies on the design of animal tests for the corrosiveness of industrial chemicals, Food Chem Toxico 23: 615 - 617

Rosenberg B, Kayden H, Lief PA, Mark LC, Steele JM and Brodie BB (1949), Studies on Diethylaminoethanol, J Pharm Exp Ther 95: 18 - 27

Schulte KE, Dreymann, H and Möllmann H (1972), Resorption, Verteilung in den Organen und Metabolisierung von Diäthylaminoäthanol nach oraler Applikation an Ratten, Arzneim-Forsch 22: 1381 - 1390

Smyth HF Jr (1964), Unpublished study, from personal communication with TLV committee member, as cited in ACGIH's Documentation of TLV, 5th ed., Cincinnati, Ohio, 1986, 2-Diethylaminoethanol, Nov. 1964: 198

Smyth HF Jr, Carpenter CP (1944), The Place of the Range Finding Test in the Industrial Toxicology Laboratory, J Ind Hyg Toxicol 26: 269 - 273

Sugimoto K Shibuya K, Ihara M, Saitoh T, Itabashi M and Nunoya T (2001), Background data on organ weights and histopathological lesions in the Crj:CD(SD)IGS rats for 4-, 13-, and 26 week repeated-dose toxicity studies, in Maeda, Y. and Inoue, H. (eds.) Biological Reference Data on CD(SD) IGS Study Group - 2000, Yokohama, 2001: 79 - 87

Swedish Products Register (2002), 23 September 2002

Swiss Product Register (2001)

Toren K (1994), NEG and NIOSH Basis for an Occupational Health Standard, Arbete och Haelsa, 25:1-18

TSCATS (1984), OTS 0001031, New Doc. I.D. FYI-OTS-0794-1031, Date produced 02/13/84, Penwalt Corp

TSCATS (1990), OTS 0530455, New Doc. I. D. 88-900000214, containing a report dated December 30, 1966 (1 year dog study) and July 31, 1967 (2 year rat study) for Pennsalt Chem. Corp., Date Produced 8/02/90, Atochem N. America, Inc.

TSCATS (1997), OTS0559210, New Doc. ID 89980000077, Date Produced 11/12/97, Elf Atochem North America

Volent P, Baer NS (1985), Volatile amines used as corrosion inhibitors in museum humidification systems, The International Journal of Museum Management and Curatorship 4: 359 - 364

Zeiger E, Anderson B, Haworth S, Lawlor T, Mortelmans K and Speck W (1987), Salmonella Mutagenicity Tests: III. Results From the Testing of 255 Chemicals, Environ Mutagen 9, Suppl 9: 1 - 110

ANNEX I

Details of the literature search used

The data bases searched are indicated below.

Toxicology: Date of last literature search: June 24, 2002

JETOC RTECS AGRICOLA CABA CANCERLIT TOXCENTER TOXLINE JICST-EPLUS LIFESCI TOXLIT EMBASE **ESBIOBASE EMBAL** HEALSAFE **CSNB** MEDLINE IRIS ATSDR TOX. PROFILES ATSDR TOX: FAQS CHEMFINDER CIVS **GESTIS** GINC NICNAS NTP

Ecology: Date of last literature search: 21June 2002

AQUASCI BIOSIS EMBASE **ESBIOBASE** LIFESCI **OCEAN** POLLUAB **SCISEARCH** TOXCENTER TOXLINE ULIDAT DATALOG CHEMFATE BIODEG AQUIRE **HSDB**

IUCLIDData Set

Existing Chemical CAS No. EINECS Name EC No. Index number Molecular Formula	ID: 100-37-8 100-37-8 2-diethylaminoethanol 202-845-2 603-048-00-6 C6H15NO
Producer Related Part Company: Creation date:	BASF AG 12-NOV-1992
Substance Related Part Company: Creation date:	BASF AG 12-NOV-1992
Memo:	master
Printing date: Revision date: Date of last Update:	07-MAR-2003 06-MAR-2003
Number of Pages:	116
Chapter (profile): Reliability (profile): Flags (profile):	Chapter: 1, 2, 3, 4, 5, 6, 7, 8, 10 Reliability: without reliability, 1, 2, 3, 4 Flags: without flag, SIDS

1. GENERAL INFORMATION

1.0.1 Applicant and Company Information

Type:	lead organisation
Contact Person:	Product Safety Date: c/o Dr. Hubert Lendle GUP/CL - Z570
Street:	Carl-Bosch-Str.
Town:	67056 Ludwigshafen
Country:	Germany
Phone:	+49 621 60 44712
Telefax:	+49 621 60 58043
Email:	hubert.lendle@basf-ag.de
Homepage:	www.basf.com
Flag: 24-JUN-2002	Critical study for SIDS endpoint
Type:	cooperating company
Name:	Air products and Chemicals Inc.
Country:	United States
Flag: 24-JUN-2002	Critical study for SIDS endpoint
Type:	cooperating company
Name:	Atofina Chemicals Inc.
Country:	United States
_	
Flag: 24-JUN-2002	Critical study for SIDS endpoint
Type:	cooperating company
Name:	DOW Chemical Company
Country:	United States
Flag: 24-JUN-2002	Critical study for SIDS endpoint
Type:	other: cooperating organisation
Name:	The Amines HPV Panel of the American Chemistry Council
Country:	United States
Flag:	Critical study for SIDS endpoint
24-JUN-2UU2	

1.0.2 Location of Production Site, Importer or Formulator

1.0.3 Identity of Recipients

<u>1.0.4 Details on Category/Template</u>

1. GENERAL INFORMATION

1.1.0 Substance Identification

Mol. Formula: Mol. Weight:	C6 H15 N O 117.19 g/mol					
Flag: 24-JUN-2002	non confidential,	Critical	study	for	SIDS	endpoint

1.1.1 General Substance Information

Substance type: Physical status: Purity: Colour: Odour:	organic liquid >= 99.5 - % w/w colourless - faint yellow amine - like	
Method: Flag: 24-JUN-2002	GC non confidential, Critical study for SIDS endpoint	(1)

1.1.2 Spectra

1.2 Synonyms and Tradenames

(2-Hydroxyethyl)diethylamine

Flag:	non	confidential,	Critical	study	for	SIDS	endpoint
02-DEC-1992							

(Diethylamino)ethanol

Flag:non confidential, Critical study for SIDS endpoint02-DEC-1992

beta.-(Diethylamino)ethanol

Flag: non confidential, Critical study for SIDS endpoint 02-DEC-1992

2-(Diethylamino)ethanol

Flag:non confidential, Critical study for SIDS endpoint02-DEC-1992

2-(Diethylamino)ethyl alcohol

Flag:non confidential, Critical study for SIDS endpoint02-DEC-1992

2-(N,N-Diethylamino)ethanol

Flag:non confidential, Critical study for SIDS endpoint02-DEC-1992

1. GENERAL INFORMATION

2-Hydroxytriethylamine non confidential, Critical study for SIDS endpoint Flag: 02-DEC-1992 DEAE Flag: non confidential, Critical study for SIDS endpoint 02-DEC-1992 Diethyl(2-hydroxyethyl)amine non confidential, Critical study for SIDS endpoint Flag: 02-DEC-1992 Diethylethanolamin Flag: non confidential, Critical study for SIDS endpoint 02-DEC-1992 Diethylethanolamine Flag: non confidential, Critical study for SIDS endpoint 02-DEC-1992 Diethylmonoethanolamine non confidential, Critical study for SIDS endpoint Flag: 02-DEC-1992 Ethanol, 2-(diethylamino)- (8CI, 9CI) Flag: non confidential, Critical study for SIDS endpoint 02-DEC-1992 MKS non confidential, Critical study for SIDS endpoint Flag: 02-DEC-1992 N,N-Diethyl-2-aminoethanol non confidential, Critical study for SIDS endpoint Flag: 02-DEC-1992 N,N-Diethyl-2-hydroxyethylamine non confidential, Critical study for SIDS endpoint Flag: 02-DEC-1992 N, N-Diethyl-N-(.beta.-hydroxyethyl)amine Flag: non confidential, Critical study for SIDS endpoint 02-DEC-1992 N,N-Diethylethanolamine Flag: non confidential, Critical study for SIDS endpoint 02-DEC-1992

1. GENERAL INFORMATION

N.N-Diethvlmonoethanol

Flag: 02-DEC-1992	non	confidential,	Critical	study	for	SIDS	endpoint
N-(2-Hydroxyethyl	diet	hylamine					
Flag: 02-DEC-1992	non	confidential,	Critical	study	for	SIDS	endpoint

Pennad 150

Flag: non confidential, Critical study for SIDS endpoint 02-DEC-1992

<u>1.3 Impurities</u>

CAS-No:	7732-18-5	
EC-No:	231-791-2	
EINECS -Name:	water	
Mol. Formula:	H2 O	
Contents:	<= .2 - % w/w	
Method:	DIN 51777	
Flag:	non confidential, Critical study for SIDS endpoint	
24-JUN-2002		(1)

1.4 Additives

<u>1.5 Total Quantity</u>

Remark:	Quantity produced
Flag:	> 1000 t/a in Germany 2000 Critical study for SIDS endpoint
13-JAN-2003	

1.6.1 Labelling

Labelling:	as in Directive 67/548/EEC	
Symbols:	(C) corrosive	
Specific limits:	no	
R-Phrases:	(10) Flammable	
	(34) Causes burns	
	(20/21/22) Harmful by inhalation, in contact with skin and if	
	swallowed	
S-Phrases:	(25) Avoid contact with eyes	
	(26) In case of contact with eyes, rinse immediately with	
	plenty of water and seek medical advice	
(36/37/39) Wear suitable protective clothing, gloves		
	eye/face protection	
	(45) In case of accident or if you feel unwell, seek medical	
	advice immediately (show the label where possible)	

1. GENERAL INFORMATION

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Flag: 23-OCT-2001	non confidential, Critical study for SIDS endpoint	(2)
1.6.2 Classification		
Classified: Class of danger: R-Phrases:	as in Directive 67/548/EEC corrosive (34) Causes burns	
Flag: 23-OCT-2001	non confidential, Critical study for SIDS endpoint	(2)
Classified: Class of danger: R-Phrases:	as in Directive 67/548/EEC flammable (10) Flammable	
Flag: 23-OCT-2001	non confidential, Critical study for SIDS endpoint	(2)
Classified: Class of danger: R-Phrases:	as in Directive 67/548/EEC harmful (20/21/22) Harmful by inhalation, in contact with skin and swallowed	l if
Flag: 23-OCT-2001	non confidential, Critical study for SIDS endpoint	(2)

1.6.3 Packaging

<u>1.7 Use Pattern</u>

Type:	industrial
Category:	Polymers industry
Remark: Flag: 05-OCT-2001	used as a catalyst in the synthesis of polymers non confidential, Critical study for SIDS endpoint
Type:	use
Category:	pH-regulating agents
Remark: Flag: 05-OCT-2001	pH stabilizer non confidential, Critical study for SIDS endpoint
Type:	use
Category:	Pharmaceuticals
Remark: Flag: 05-OCT-2001	used in the synthesis of drugs non confidential, Critical study for SIDS endpoint

1.7.1 Detailed Use Pattern

1. GENERAL INFORMATION

1.7.2 Methods of Manufacture

Orig. of Subst.: Type:	Synthesis Production
Remark:	2-diethylaminoethanol is produced by the thermal reaction of diethylamine with ethylenoxide.
Flag:	non confidential, Critical study for SIDS endpoint
27-FEB-2003	

1.8 Regulatory Measures

1.8.1 Occupational Exposure Limit Values

Type of limit: Limit value:	MAK (DE) 5 ml/m3	
Flag: 05-AUG-2002	non confidential, Critical study for SIDS endpoint	(3)
Type of limit: Limit value:	MAK (DE) 24 mg/m3	
Remark:	cutaneous resorption	
Flag: 05-AUG-2002	non confidential, Critical study for SIDS endpoint	(3)
Type of limit: Limit value:	TLV (US) 9.6 mg/m3	
Remark: Flag: 05-AUG-2002	equal to 2 ppm non confidential, Critical study for SIDS endpoint	(1)

1.8.2 Acceptable Residues Levels

1.8.3 Water Pollution

Classified by:	other: VwVwS (Germany) of 17.05.1999, Annex 2		
Labelled by:	other: VwVwS (Germany) of 17.05.1999, Annex 2		
Class of danger:	: 1 (weakly water polluting)		
Remark:	Identification number: 1288		
Flag:	non confidential, Critical study for SIDS endpoint		
03-JUL-2002		(4)	

1.8.4 Major Accident Hazards

<u>1.8.5 Air Pollution</u>

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

1. GENERAL INFORMATION

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003

SUBSTANCE ID: 100-37-8

Flag: 23-FEB-2001	non confidential, Critical study for SIDS endpoint (1)
<u>1.8.6 Listings e.g.</u>	Chemical Inventories
Type: Additional Info:	EINECS EINECS No. 202-845-2
Flag: 24-JUN-2002	non confidential, Critical study for SIDS endpoint (5)
Type: Additional Info:	ENCS ENCS No. 2-297X
Flag: 24-JUN-2002	non confidential, Critical study for SIDS endpoint (5)
Type: Additional Info:	ECL ECL Serial No. KE-20903
Flag: 24-JUN-2002	non confidential, Critical study for SIDS endpoint (5)
Type: Additional Info:	other: SWISS SWISS No. G-1873
Remark: Flag: 24-JUN-2002	SWISS classification: Giftliste 1 (List of Toxic Substances 1), 31 May 1999 Toxic Category 4: Acute oral lethal dose of 500 - 2000 mg/kg. non confidential, Critical study for SIDS endpoint (5)
Туре:	TSCA
Flag: 24-JUN-2002	non confidential, Critical study for SIDS endpoint (5)
Туре:	DSL
Flag: 24-JUN-2002	non confidential, Critical study for SIDS endpoint (5)
Туре:	AICS
Flag: 24-JUN-2002	non confidential, Critical study for SIDS endpoint (5)
Туре:	PICCS
Flag: 24-JUN-2002	non confidential, Critical study for SIDS endpoint (5)

1.9.1 Degradation/Transformation Products

1.9.2 Components

1.10 Source of Exposure

1. GENERAL INFORMATION

1.11 Additional Remarks

Memo:	Hazardous reactions: Exothermic reaction with acids	
Flag:	non confidential, Critical study for SIDS endpoint	
24-JUN-2002		(1)

1.12 Last Literature Search

Type of Search:	External
Remark:	<pre>date of last literature search: - toxicology: 24 June 2002 - ecology/environment: 21 June 2002 Critical study for SLDS endpoint</pre>
02-JUL-2002	clitical study for SIDS enupoint
Type of Search: Chapters covered: Date of Search:	Internal and External 3, 4, 5 13-NOV-2002
Remark: 13-JAN-2003	update 2003, no new data found
Type of Search: Chapters covered: Date of Search:	Internal and External 5.10 06-NOV-2002

06-FEB-2003

1.13 Reviews
2. PHYSICO-CHEMICAL DATA

2.1 Melting Point

Value:	< -70 degree C	
Reliability:	(4) not assignable	
09-MAY-2000	Manufacturer/producer data without proof	(1)
Value:	= -68 degree C	
Method:	other: measured (test procedure according to an internal standard, comparable to OECD 102)	l basf
Remark: Test substance: Reliability:	<pre>reason for flagging this data: experimental derived data 2-diethylethanolamine, purity > 99.5 % (GC) (2) valid with restrictions scientifically acceptable method</pre>	ł
Flag: 18-NOV-2002	Critical study for SIDS endpoint ((6) (7)

<u>2.2 Boiling Point</u>

Value:	= 161.5 - 163 degree C	
Reliability:	(4) not assignable Manufacturer/producer data without proof	
09-MAY-2000	nanaraocarer, producer auca wrenoue proor	(1)
Value:	= 162 - 163 degree C at 1013 hPa	
Remark:	reason for flagging this data: handbook (Beilstein) enjoys	a
Test substance: Reliability:	CAS 100-37-8 (2-diethylaminoethanol), purity not indicated (2) valid with restrictions	
Flag.	data from reliable handbook Critical study for SIDS endpoint	
02-JUL-2002	critical stady for SLDS chapoline	(8)
Value:	= 162.1 degree C at 1012.96 hPa	
Decomposition:	no	
Method: Year:	other: measured using a twin ebulliometric apparatus 2002	
GLP: Test substance:	no as prescribed by 1.1 - 1.4	
Remark:	Water was used in the reference ebulliometer. T is the condensation temperature of the sample. The press p was calculated considering the condensation temperature the reference substance.	ure of
Test substance:	N,N-Diethylethanolamine, purity >= 99.95 mol %	
Reliability:	(2) valid with restrictions	
14-NOV-2002	sciencifically acceptable method	(9)
Value:	= 163 degree C at 1010 hPa	

2. PHYSICO-CHEMICAL DATA

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Test substance: Reliability:	CAS 100-37-8 (2-diethylaminoethanol), purity not indicated (4) not assignable only secondary guotation
15-OCT-2001	(10)
2.3 Density	
Value:	= .8889 g/cm³ at 20 degree C
Reliability:	(4) not assignable Manufacturer/producer data without proof
09-MAY-2000	(1)
Type: Value:	density = .885 g/cm³ at 20 degree C
Method:	other: measured (test procedure according to an internal BASF standard)
Remark:	reason for flagging this data: reliable data available on this
Test substance: Reliability:	<pre>parameter 2-diethylethanolamine, purity > 99.5 % (GC) (2) valid with restrictions</pre>
Flag:	scientifically acceptable method Critical study for SIDS endpoint
18-NOV-2002	(7)
Type: Value:	density = .8809 g/cm³ at 22.9 degree C
Method:	other: measured using a high-pressure pycnometer
GLP:	no
Test substance:	other TS
Remark:	all measurements were done at atmospheric pressure in the present work
Test substance: Reliability:	N,N-Diethylethanolamine, purity 99 % (2) valid with restrictions
14-NOV-2002	(11)
Type: Value:	density = .8559 g/cm³ at 50 degree C
Method: Year:	other: comparable to OECD 109 2002
GLP: Test substance:	no as prescribed by 1.1 - 1.4
Method: Test substance:	vibrating tube densimeter N,N-Diethylethanolamine, purity >= 99.95 mol %
Reliability:	(2) valid with restrictions
18-NOV-2002	scientifically acceptable method (9)

2. PHYSICO-CHEMICAL DATA

2.3.1 Granulometry

2.4 Vapour Pressure

Value:	ca. 1.8 hPa at 20 de	gree C		
Method:	other (measured): dy according to an inte 104)	namic with nitrogen (test procedure rnal BASF standard, comparable to OECD		
Remark:	reason for flagging	chis data: experimentally derived data		
Result:	temperature (°C)	vapour pressure (hPa)		
	13.36	1.007		
	22.40	2.006		
	28.19	3.004		
	38.66	5.996		
	47.10	10.061		
	67.24	30.046		
	84.90	70.022		
	93.30	100.610		
	122.29	300.400		
	148.94	700.160		
	155.71	850.040		
	161.61	999.910		
	using the measured s	et of data and plotting the log of the		
	vapour pressure (in Pascals) against the reciprocal of the			
	temperature (in Kelv	ins) then drawing in the line of best fit,		
	to give the equation	y = -2505.4x + 10.801, were $y = vapour$		
	pressure (Pa) and x	= temperature (K) the vapour pressure at		
Test substance.	2-diethylethanolamin	and at 20°C it is ca. i.o nea p purity 99 68 area%		
Reliability.	(2) valid with rest	rictions		
ActiuDiticy.	ccientifically accept	table method		
Flag:	Critical study for S	IDS endpoint		
18-NOV-2002		(12)		
		(==)		
Value:	= 1.9 hPa at 20 degre	ee C		
Reliability:	(4) not assignable			
	Manufacturer/produce	r data without proof		
09-MAY-2000		(1)		
0, 1111 2000		(-)		
Value:	= 2 hPa at 22.4 degre	ee C		
Method:	other (measured): dy	namic with argon (test procedure according		
	to an internal BASF	standard, comparable to OECD 104)		
Remark:	reason for flagging	this data: experimentally derived data		
Result:	temperature (°C)	vapour pressure (hPa)		
	22.40	2.00		
	35.63	5.00		
	46.93	10.00		
	59.28	20.00		
	77.61	50.00		
	93.43	100.0		
	111.11	200.0		
	138.07	500.0		

OECD SIDS 2. PHYSICO-CHEMICAL DATA

SUBSTANCE ID: 100-37-8

Test substance: Reliability: Flag: 18-NOV-2002 Value: Decomposition:	<pre>162.30 2-diethyl (2) vali scientifi Critical = 20.022 no</pre>	1 ethanolamine, purit d with restrictions cally acceptable me study for SIDS endp hPa at 59.4 degree	011.5 y 99.7 % (GC) thod oint C	(13)
Method: Year:	other (me 2002	asured): using a tw	in ebulliometric apparatus	
GLP:	no			
Test substance:	as prescr	ibed by 1.1 - 1.4		
Result:	method d d d d d d d d d w w w w w w w w w w	temperature (T/K) 332.500 346.094 352.159 361.220 368.003 373.496 379.233 383.986 390.254 390.227 396.546 402.902 409.277 415.709 422.189 428.716 435.281	<pre>vapour pressure (p/kPa) 2.0022 4.0031 5.3331 8.0029 10.6642 13.325 16.669 19.937 25.037 25.011 31.178 38.576 47.366 57.801 70.099 84.517 101.296</pre>	
Test substance: Reliability:	Water (w) the refer T is the p was cal reference N,N-Dieth (2) vali scientifi	or n-decane (d) re rence ebulliometer. condensation temper culated from the co substance. ylethanolamine, pur d with restrictions cally acceptable me	fers to which material was use ature of the sample. The press ndensation temperature of the ity >= 99.95 mol % thod	d in ure
18-NOV-2002				(9)

2.5 Partition Coefficient

Partition Coeff.: log Pow:	octanol-water = .047
Method:	other (calculated): computer program KOWWIN v1.66
Remark:	reason for flagging this calculation: model accepted by the US-EPA
Reliability:	(2) valid with restrictions scientifically acceptable method
Flag:	Critical study for SIDS endpoint
23-JUL-2002	(14)
log Pow:	= .0551

Method:

Reliability:

28-FEB-2000

log Pow:

Method:

Year:

2. PHYSICO-CHEMICAL DATA

standard methods

1987

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

(15)

other (calculated): different methods (2) valid with restrictions Calculated value in accordance with generally accepted = .21 at 23 degree C other (measured): following OECD 107

GLP: no Method: TS concentration was measured titrimetrically in both phases reason for flagging this study: Inspite of using a Remark: non-buffered solution the result is comprehensible. Result: TS concentration (mol/L) Pow (1) log Pow (octanol) (water) (mean) 0.0266 0.21 0.0139 1.628 0.0309 0.0187 1.655 (1) = c(oct.)/c(H20)CAS Nr. 100-37-8 (2-diethylaminoethanol), purity 99.1 % Test substance: Reliability: (2) valid with restrictions following guideline with restrictions Flag: Critical study for SIDS endpoint 04-JUL-2002 (16) = .31 - .46 log Pow: Reliability: (2) valid with restrictions data from reliable handbook 03-JUL-2002 (17)= .333 log Pow: Method: other (calculated): incremental method of Rekker (computer program from CompuDrug Ltd.) Year: 1989 Reliability: (2) valid with restrictions

Scientifically acceptable method 09-OCT-2001 = .401 log Pow: Method: other (calculated): computer program CLOGP 3.3

Method: In the article calculated log Pow values were compared with experimentally derived values. Test substance: CAS 100-37-8 (2-diethylaminoethanol), purity not indicated Reliability: valid with restrictions (2) Scientifically acceptable method 23-JUL-2002 (19) (20)

(18)

2. PHYSICO-CHEMICAL DATA

(7)

2.6.1 Solubility in different media

Value: pH	value: Conc.:	at 20 degree C 11.5 100 g/l at 20 degree C
Descr.:		other: miscible in all proportions
Reliabil	ity:	(4) not assignable Manufacturer/producer data without proof.
02-JUL-2	002	(1)
рН	value: Conc.:	= 12 100 g/l at 20 degree C
Method:		other: DIN 19 267
Remark: Result:		reason for flagging this data: experimentially derived data pH values at different concentrations of DEAE:
		(wDEAE + (1-w) H20; w= mass proportion)
Reliabil	ity:	pH = 11.45 at 20 °C (w = 0.01) pH = 11.86 at 20 °C (w = 0.05) pH = 12.03 at 20 °C (w = 0.1) pH = 12.36 at 20 °C (w = 0.3) (4) not assignable
		Secondary quotation. Only test results are cited in the report BRU 85.219, BASF, Physikalisch-chemische Konstanten, 16.12.1985. Original reference (BASF, ZAM/C, JNo. 306025) is not available
Flag: 02-JUL-2	002	Critical study for SIDS endpoint (7)
Remark:		reason for flagging this data: important information on this parameter
Result: Test sub Reliabil	stance: ity:	soluble in all proportions CAS 100-37-8 (2-diethylaminoethanol), purity not indicated (4) not assignable
Flag: 15-OCT-2	001	only secondary quotation Critical study for SIDS endpoint (10)
<u>2.6.2 Sur</u>	<u>rface Tens</u>	<u>ion</u>
Test typ	e:	other: capillary method (test procedure according to an internal BASF standard)
Value:		= 27.9 mN/m at 20 degree C
Remark: Test sub Reliabil	stance: ity:	reason for flagging this data: experimentially derived data 2-diethylethanolamine, purity > 99.5 % (GC) (2) valid with restrictions
Flag:		Critical study for SIDS endpoint

2.7 Flash Point

2. PHYSICO-CHEMICAL DATA

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Value:	= 51.5 degree C	
Туре:	closed cup	
Method:	other: DIN 51 755	
Remark:	reason for flagging this data: experimentially derived da	ata
Reliability:	(1) valid without restriction	
	National standard specification	
Flag:	Critical study for SIDS endpoint	
09-OCT-2001		(21)

2.8 Auto Flammability

Value:	270 degree C	
Method:	other: DIN 51 794	
Remark:	Ignition temperature	
Reliability:	reason for flagging this data: experimentially derived dat (2) valid with restrictions National standard specification	a
Flag:	Critical study for SIDS endpoint	
24-JUN-2002		(21)

2.9 Flammability

Result:	flammable	
Remark:	reason for flagging this data: only statement available of this parameter	n
Reliability:	(4) not assignable	
	Manufacturer/producer data without proof	
Flag:	Critical study for SIDS endpoint	
09-OCT-2001		(1)

2.10 Explosive Properties

Result: not explosive

Remark:	because of chemical structure	
	reason for flagging this data: only statement available or	ı
	this parameter	
Reliability:	(2) valid with restrictions	
	Expert judgement	
Flag:	Critical study for SIDS endpoint	
09-OCT-2001		(22)

2.11 Oxidizing Properties

Result:	no oxidizing properties
Remark:	because of chemical structure reason for flagging this data: only statement available on this parameter

2. PHYSICO-CHEMICAL DATA

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Reliability:	(2) valid with restrictions Expert judgement	
Flag: 09-OCT-2001	Critical study for SIDS endpoint	(22)

2.12 Dissociation Constant

Acid-base Const.:	pKa = 9.87	
Method: GLP:	other no data	
Remark:	reason for flagging this data: only information available of this parameter	on
Test substance:	CAS 100-37-8 (2-diethylaminoethanol), purity not indicated	
Reliability:	(4) not assignable	
	only secondary quotion	
Flag:	Critical study for SIDS endpoint	
10-OCT-2002		(23)

2.13 Viscosity

Value:	= 4.6 mPa s (dynamic) at 25 degree C							
Reliability:	4) not assignable anufacturer/producer data without proof							
23-JUL-2002	(1)							
Test type: Value:	Capillary Method = 4.496 mPa s (dynamic) at 22.2 degree C							
Method: GLP:	other: similar OECD 114 no							
Test substance:	other TS							
Method:	Kinematic viscosities were measured using calibrated Cannon-Ubbelohde capillary viscometer. The measured kinematic viscosities were converted to absolute viscosity considering the measured densities							
Test substance: Reliability:	<pre>N,N-Diethylethanolamine, purity 99 % (2) valid with restrictions scientifically acceptable method</pre>							
14-NOV-2002	(11)							

2.14 Additional Remarks

Memo:	Hygroscopic liquid	
Remark:	reason for flagging this information: important information of this parameter	n
Reliability:	(4) not assignable only secondary quotion	
Flag:	Critical study for SIDS endpoint	
10-OCT-2002	(24)

2. PHYSICO-CHEMICAL DATA

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Remark:	reason for flagging this data: reliable data available on this parameter
Result:	Explosive limits in air: 0.7 vol.% (39 °C) - 10.1 vol.% (92.5 °C)
Reliability:	(2) valid with restrictions
Flag:	test procedure according to internal standard Critical study for SIDS endpoint
09-OCT-2001	(21)

3. ENVIRONMENTAL FATE AND PATHWAYS

3.1.1 Photodegradation

Type:	air	
INDIRECT PHOTOLYS	IS	
Sensitizer:	ОН	
Conc. of sens.:	500000 molecule/cm ³	
Rate constant:	= .00000000985519 cm ³ /(molecule * sec)	
Degradation:	= 50 % after 3.9 hour(s)	
Method:	other (calculated): AOP v1.87	
Remark:	reason for flagging this data: only information available	on
Reliability:	this parameter; model accepted by the US-EPA (2) valid with restrictions	
	scientifically acceptable method	
Flag:	Critical study for SIDS endpoint	
04-JUL-2002		(25)

3.1.2 Stability in Water

3.1.3 Stability in Soil

3.2.1 Monitoring Data (Environment)

Remark:	reason for flagging this information: important information
	about measured DEAE in industrial effluent discharges even
	data were evaluated nearly 25 years ago and may not be
	representative of, or comparable to today's situation.
	Samples of 63 effluent and 22 intake waters were collected
	from a wide range of chemical manufacturers in areas across
	the United States. The samples were analyzed for organic
	compounds (among them DEAE) in an effort to identify
	previously unknown and potentially hazardous organic
	pollutants.
	Each water sample was preconcentrated for analysis of organic
	compounds. All sample analyses involved a GC/MS/COMP system
	that used high-resolution glass capillary GC columns.
Result:	- Over 570 compounds were tentatively identified.
	- DEAE was analyzed in one sample (compound concentration:
	>100 µg/L)
Reliability:	(2) valid with restrictions
	acceptable publication which meets basic scientific principles
Flag:	Critical study for SIDS endpoint
16-JUL-2002	(26)
Type of measureme	ent: other
Remark:	In 1980, N,N-diethylethanolamine has been detected
	qualitatively by gas chromatography-mass spectroscopy in the
	effluent of the publicly-owned treatment works at Decatur
	(Illinois) in which industrial wastes of different origin were
	discharged.
	reason for non-flagging this information: the documentation is
	insufficient for assessment since the effluent concentration
	of DEAE is not reported

3. ENVIRONMENTAL FATE AND PATHWAYS

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Reliability:	(2) valid with restrictions			
08-JUL-2002	(27)			
Type of measuremen	ht: other			
Remark:	<pre>Emissions of volatile organic compounds (VOC) from different types of furniture coatings have been investigated by test chamber studies under dynamic conditions: - test chamber: 1 m³ glass chamber (Salthammer T. and Marutzky R., Kammerverfahren zur Bestimmung der Emissionen organischer Substanzen aus Materialien. In: Bagda E. (ed.). Emissionen aus Beschichtungsstoffen und deren Einfluß auf die Innenraumluft, Renningen, Expert-Verlag, 75-94, 1995) - temperature: 23 °C - relative humidity: 45 % - air exchange rate n: 1.0 h-1 (air velocities: 0.05 m/s - 0. - loading factor a (surface to volume ratio): 1.0 m²/m³ - samples: 44 furniture samples manufactured under industrial conditions. Samples were placed in the test chamber immediately after production (samples 1-17) and after preconditioning for 20 days at 23°C and 45 % relative humidity (samples 18-44), respectively. Reason for non flagging this study: the documentation is insufficient for assessment since the chamber concentration or</pre>			
Result:	emission rate of DEAE is not reported 2-diethylethanolamine was identified in 2 samples in the chamber air test, but no chamber concentrations c (μ g/m ³) or emission rate ER (μ g/(h*m ²)) are reported in the article			
25-JUL-2002	acceptable publication which meets basic scientific principles (2) (2) (2)			
Type of measuremen	nt: other			
Remark:	DEAE has been used as a corrosion inhibitor since 1979 in the open steam humidification system of the H. F. Johnson Museum at Cornell University. Analysis of air samples from the museum environment were undertaken in January of 1983. Reason for flagging this information: important information about measured indoor air concentration even data were evaluated nearly 20 years ago and may not be representative of, or comparable to today's situation.			
Result:	14 air samples were collected, 10 at a sampling rate of 0.2 l/min (detection limit: 0.4 mg/m ³) and 4 at 1.5 l/min (detection limit: 0.04 mg/m ³). DEAE was detected in only 2 air samples in concentrations of 0.05 and 0.04 mg/m ³ . Residues of condensed DEAE were identified on samples of plastic materials in the museum in concentrations of 30 mg/m ² . Condensation accumulation was found directly related to time of deposition for airborne concentrations of 0.05 mg/m ³ .			
Reliability:	(2) valid with restrictions acceptable publications, which meet basic scientific			
Flag: 25-JUL-2002	Critical study for SIDS endpoint (29) (30)			

Type of measurement: other

3. ENVIRONMENTAL FATE AND PATHWAYS

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003

SUBSTANCE ID: 100-37-8

Remark:	- During the winter season, buildings are often humidified
	with steam which may come from a boiler system that is being
	treated with volatile neutralizing amines to prevent
	corrosion. In this study, a room at Battelle Columbus
	Division, in Columbus, OH, was selected as a typical
	steam-humidified room.
	- The Battelle boiler system was treated with a mixture of
	cyclohexylamine (CAS-No. 108-91-8) and 2-diethylaminoethanol
	for corrosion control.
	- The concentrations of both chemicals were measured in indoor
	air in the study room.
	Reason for flagging this information: important information
	about measured indoor air concentration even data were
	evaluated nearly 15 years ago and may not be representative
	of, or comparable to today's situation.
Result:	- The concentration of DEAE (detection limit: 0.1 ppb)
	measured in indoor room air during normal operation of the
	boiler and humidification systems remained very low compared
	with any established health standards and did not present any
	hazard to health.
	- The average room concentration of DEAE at the average of 42
	<pre>% relative humidity was about 0.6 nph (approx 2.9*10-3)</pre>
	$m_{\rm cr}/m^3$)
	- At 61 % relative humidity the average DEAE-concentration
	was 2.4 ppb (approx 0.01 mg/m^3)
	- During the final hour of the monitoring period when the
	humidifier was shut off the concentration decayed to ~50 % of
	the steady-state value at 61 % relative humidity
	- When the humidifier was opened all the way at the end of the
	study the concentration of the DEAE increased up to 8.2 ppb
	$(approx 0.034 \text{ mg/m}^3)$ the highest value recorded
	- At 61 % relative humidity the removal rate was 14 ug/s for
	DEAE
	- The primary fate of the amine that was introduced into room
	air through steam humidification appears to be removal to
	surfaces.
Reliability:	(2) valid with restrictions
	acceptable, well documented study which meets basic scientific
	principles
Flag:	Critical study for SIDS endpoint
25-JUL-2002	(31)

3.2.2 Field Studies

3.3.1 Transport between Environmental Compartments

Type:	adsorption						
Media:	water - soil						
Method: other: calculated: PCKOWWIN v1.63							
Remark:	reason for flagging this data: model accepted by the US-EPA						
Result:	KOC = 5.979; Log KOC = 0.7767						
Reliability:	(2) valid with restrictions						
	scientifically acceptable method						
Flag:	Critical study for SIDS endpoint						
04-JUL-2002	(32)						

OECD SIDS 3. ENVIRONMENTAL FATE AND PATHWAYS

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Type: adsorption Media: water - soil other: model calculation Method: A soil sorption coefficient Koc = 30.99 (log Koc = 1.49) was Remark: estimated on the basis of the regression derived equation (log Koc = 0.544 log Pow + 1.377; log Pow = 0.21) Reliability: (2) valid with restrictions scientifically acceptable method 02-JUL-2002 (33) (34) Type: volatility water - air Media: Method: other: estimated value Remark: a Henry's law constant (H = 2.51E-4 (hPa*m3/mol)) was estimated on the basis of the equation H= p/S. S = water-solubility. The water-solubility of 2-diethylaminoethanol is unlimited (S = 1E6 ppm (= 1 m3/m3) respectively S = 7.560372E3 mol/m3. p = vapour pressure (p = 1.9 hPa)(2) valid with restrictions Reliability: scientifically acceptable method 09-OCT-2001 (35) Type: volatility Media: water - air Method: other: calculated: HENRYWIN v3.00 reason for flagging this data: calculation based on standard Remark: calculation program Result: Henry's law constant: H = 3.16E-4 Pa*m3/mol at 25 °C (bond method)Reliability: (2) valid with restrictions scientifically acceptable method Flag: Critical study for SIDS endpoint 02-JUL-2002 (32)

3.3.2 Distribution

Media:	air - biota - sediment(s) - soil - water							
Method:	Calculation according Mackay, Level I							
Method:	Level I, V 2.11							
Remark:	<pre>calculation is based on the following physical chemical properties of the substance: H = 0.0251 Pa*m3/mol (20 °C); molecular weight: 117 g/mol; vapour pressure: 190 Pa; melting point: -68 °C; water-solubility: 8.8E5 g/m3; log Kow: 0.21 reason for flagging this data: only information available on this orderaint</pre>							
Result:	The calculation is for the uncharged molecule:							
	air: 0.88 % water: 99.1 % soil: 0.01 % sediment: 0.01 %							

3. ENVIRONMENTAL FATE AND PATHWAYS

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Reliability:	(2)	valid	d with	n res	strict	cions	
	scien	tific	cally	acce	eptabl	Le me	thod
Flag:	Criti	cal s	study	for	SIDS	endp	oint
23-JUL-2002							

(32)

3.4 Mode of Degradation in Actual Use

3.5 Biodegradation

Type:	aerobic					
Inoculum:	activated sludge, domestic					
Concentration:	33 mg/l related to Test substance					
	20 mg/l related to DOC (Dissolved Organic Carbon)					
Contact time:	22 day(s)					
Degradation:	95 % after 22 day(s)					
Result:	readily biodegradable					
Kinetic:	7 day(s) 5 %					
	10 day(s) 82 %					
	22 dav(s) = 95%					
Control Subst.:	Aniline					
Kinetic:	3 day(s) 2 %					
	5 93 %					
	5 55 0					
Method:	OECD Guide-line 301 A (new version) "Ready Biodegradability:					
	DOC Die Away Test"					
Year:	1992					
GLP:	Ves					
Test substance:	other TS: N N-Diethylethanolamin purity: 99 5 % (area)					
1000 Dubbeance.	Scher is with siden field and a many party so is a carea,					
Kemark:	<pre>reason for flagging this study: only reliable test on ready biodegradability available test device: - 2 l Erlenmeyer flasks, liquid volume: 1000 ml incubation: - on a laboratory shaker, approx. 120 rpm - temperature: 22 +- 2 °C number of replicates: - test substance (TS): 2</pre>					
	- blank (BC): 2					
	- inhibition control (IH): 1					
	 assay to examine physico chemical (abiotic) elimination (PC): 1 adsorption control (AC): 1 					
	inoculum:					
	 source: activated sludge, domestic (sludge from laboratory wastewater treatment plants fed with municipal sewage) 					
	reference control:					
	- reference substance: aniline					
	- concentration: 20 mg/L related to DOC					
	- kinetic of reference substance: 3 dav(s) 2 % DOC-elimin					
	7 day(s) 89 % DOC-elimin.					

2-DIETHYLAMINOETHANOL

3. ENVIRONMENTAL FATE AND PATHWAYS

DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

```
14 day(s) 92 % DOC-elimin.
                                                    22 day(s) 96 % DOC-elimin.
                  inhibition control:
                   - substances: aniline + test substance
                  - concentration: aniline: 20 mg/L related to DOC
                                   test subst.: 20 mg/L related to DOC
                  - kinetic of inhibition control: 3 day(s)
                                                              2 % DOC-elimin.
                                                   7 day(s)
                                                             49 % DOC-elimin.
                                                   14 day(s) 96 % DOC-elimin.
                                                   22 day(s) 101 % DOC-elimin.
                  assay to examine physico-chemical (abiotic) elimination:
                  - concentration: 20 mg/L related to DOC
                  - w/o inoculum, w mercury chloride to avoid microbial
                  biodegradation
                  - kinetic of physico-chemical elimination:
                                                   1 day(s): -5 % DOC-elimin.
                                                              8 % DOC-elimin.
                                                   7 day(s):
                                                   14 day(s): 1 % DOC-elimin.
                                                   22 day(s): -4 % DOC-elimin.
                  adsorption control:
                  - concentration: 20 mg/L related to DOC
                  - w incoulum, w mercury chloride to avoid microbial
                  biodegradation
                  - kinetic of adsorption control:
                                                   1 day (s): -3 % DOC-elimin.
                                                   5 day (s): 0 % DOC-elimin.
Result:
                  - duration of the adaption phase (lag-phase): 7 days
                  - duration of the degradation phase (log-phase): 10 days
                  - physico-chemical (abiotic) elimination assay: <10 % (DOC) at
                  the end of the test
                  - adsorption control: < 10 % (DOC) after 5 days
                  - inhibition assay: 100 % (DOC) after 22 days
Reliability:
                  (1) valid without restriction
                  guideline study
                  Critical study for SIDS endpoint
Flag:
07-NOV-2002
                                                                             (36)
                  aerobic
Type:
Inoculum:
                  activated sludge, domestic
Concentration:
                  354 mg/l related to Test substance
                  200 mg/l related to DOC (Dissolved Organic Carbon)
Contact time:
                  14 \, day(s)
Degradation:
                  = 96 % after 14 day(s)
                  inherently biodegradable
Result:
Control Subst.:
                  Diethylene glycol
Method:
                  OECD Guide-line 302 B "Inherent biodegradability: Modified
                  Zahn-Wellens Test"
  Year:
                  1992
  GLP:
                  ves
                  other TS: 2-diethylaminoethanol, purity 99.9 %
Test substance:
Method:
                  Test mixture (total volume 1.5 L) containing test substance
                  (354 mg/L <> 200 mg/L DOC), mineral medium and activated
                  sludge from a laboratory wastewater treatment plant treating
                  municipal wastewater (1 g/L) was incubated for 14 days.
```

3. ENVIRONMENTAL FATE AND PATHWAYS

2-DIETHYLAMINOETHANOL

DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

The following controls were included: - Inoculum blanc: control without test substance but with					
	inoculum (1 flask)				
	- Positive	control: die 1 flack)	thylene glycol (200 m	ng/L DOC) with	
	- Abiotic (control: cont	rol with test substar	ice and mercury	
	chloride bu	ut without ind	oculum (1 flask)		
	Aliquots w 1, 2, 3, 6	ere removed f: , 7, 9, 10, 1	rom the flasks on day 3 and 14 and analyzed	y 0 (0 and 3 h), l for DOC	
Remark:	Control sul	bstance: diet	hylene glycol. DOC-el	imination after	
	reason for	flagging this	s data: test was perf	formed with non	
_ .	adapted in	oculum from m	unicipal wwtp		
Result:	time	elimina	ation (% DOC of day 0	values)	
	(day) TS	with inoculu	m Reference substand	ce abiotic control	
	0 (0 h)	0	0	U	
	0 (3 h)	-5	4	-6	
	1	-5	8	-4	
	2	-5	8	-4	
	3	-1	12	-3	
	6	-4	98	-6	
	7	8	100	6	
	9	38	99	0	
	10	97		3	
	13	98		-4	
	14	96		1	
Reliability:	- The sigmo indication - Test subs - Adaptatio - Biodegrao (1) valid	oidal shape or on biodegrada stance was no on phase: app dation phase: without restr	f the elimination cur ation t adsorbed by the slu rox. 8 days. approx. 2-3 days iction	rve is a strong udge	
_	GLP-study				
Flag:	Critical st	tudy for SIDS	endpoint		
23-JUL-2002				(37)	
Type:	aerobic				
Inoculum:	other: acclimated sewage sludge enrichment culture				
Remark:	Rothkopf an 2-diethyle: medium: 0.1 temperatury µl/ml. Cony yield/2-di	nd Bartha (19) thanolamin. T 05 % test subs e: 28 °C. Max version facto: ethylethanolas	84) observed biodegra hey measured cell gro stance as C- and N-so . protein yield: appr r of 35 % protein min. 70 µg/mL were me	adation of owth. Test ource. Incubation coximately 175 easured for the	
	test substa	ance in an ada	apted culture and aft	er an incubation	
	time of one	e week			
Reliability:	(2) valid	with restrict	tions		
	acceptable	publication v	which meets basic sci	entific principles	
02-JUL-2002		- 30110401011 V		(38)	
Thogulum	other hast	oria: inducto	ial gludge from write	of BACE	
Degradation:	= 100 % af	ter 11 day(s)	Tar Brudge IIOm wwtp	OT DADL	

3. ENVIRONMENTAL FATE AND PATHWAYS

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Method: GLP: Test substance:	other: Modified Zahn-Wellens Test no as prescribed by 1.1 - 1.4
Remark: Reliability:	easily eliminated, mainly by biodegradation (2) valid with restrictions comparable to quideline study with acceptable restrictions
09-OCT-2001	(39)

02-OCT-2002

3.6 BOD5, COD or BOD5/COD Ratio

Method: Year: Method:	
Pemark.	- inoqulum:
Kemark.	effluent from industrial wwtp (BASF AG, Ludwigshafen, Germany)
	- result: TOC = 571 mg/g COD = 760 mg/g BOD5 = 2 mg/g
	BOD5*100/COD: < 1%
	- method: BODx-determination, DEV H DIN 38409, part 51, Deutsche Einheitsverfahren zur Wasser-, Abwasser- und Schlammuntersuchung (determination of biochemical oxygen demand)
Reliability:	(2) valid with restrictions test procedure according to guideline with acceptable
24-JUN-2002 Method:	restrictions (39)
Year:	
Method:	
Remark:	- inoculum:
	acclimated microbial seed: a mixed microbial culture capable of using 2-diethylaminoethanol as sole carbon and energy sources was isolated by an enrichment culture technique. Microbial seed for the BOD test was prepared from the culture growth (1E5-1E6 cells/ml) in mineral salts medium containing 100 µl/l chemical substance. The culture was diluted (1:1) with physiological saline and incubated on a shaker for 24 h prior to its use.
Result:	<pre>- method: BOD technique according to American Public Health Association. Standard Methods for the Examination of Water and Wastewater. American Public Health Association, Washington, DC., 1980. incubation: 20 days at 21 °C +-3 °C; 300 ml BOD bottle. ThBOD: 9.25 mmol/mmol chemical BOD5 +- SD: 5.5+-0.24 mmol/mmol chemical</pre>
	BOD5*100/ThBOD = 59.5 %

3. ENVIRONMENTAL FATE AND PATHWAYS

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Reliability:	(2)	valid	with	restri	ctions				
	accer	ptable	publi	ication	which	meets	basic	scientific	principles
23-JUL-2002									(40)

3.7 Bioaccumulation

Method:	other: estimated value
Remark:	Based on a measured log Pow of 0.21, the bioconcentration factor (BCF) for 2-diethylaminoethanol can be estimated to be 0.85 from the recommended regression-derived equation log BCF = 0.76*log Pow - 0.23. reason for flagging this data: only data available on this
Reliability:	endpoint (2) valid with restrictions
	scientifically acceptable method
Flag:	Critical study for SIDS endpoint
27-JUN-2002	(33) (35)

3.8 Additional Remarks

AQUATIC ORGANISMS

4.1 Acute/Prolonged Toxicity to Fish

Type:	flow through					
Species:	Pimephales promelas (Fish, fresh water)					
Exposure period:	96 hour(s)					
Unit:	mg/l Analytical monitoring: yes					
LC50:	= 1780					
Method:	other					
GLP: Togt gubgtango.	no data					
lest substance.	<pre>> 00%</pre>					
Remark:	The TS was given in a solution in which the pH was adjusted					
	to that of lake water with HCl.					
Reliability:	(1) valid without restriction					
_	comparable to guideline study					
Flag:	Critical study for SIDS endpoint					
01-JUL-2002	(41)					
Type •	static					
Species.	Leuciscus idus (Fish fresh water)					
Exposure period:	96 hour(s)					
Unit:	mg/1 Analytical monitoring: no					
NOEC:	= 100					
LC50:	147					
Method:	other: German Industrial Standard DIN 38 412, Part 15					
Year:	1982					
GLP:	no					
Remark:	Closely followed the German Industrial Standard Guideline					
	Number DIN 38 412. Part 15 (June 1982) using a static					
	exposure procedure.					
	The Golden Orfe (L. idus), golden variety was used.					
	Aeration: slight, with oil free air					
	Duration of housing and adaptation: about 5 weeks					
	Duration of adaptation: 3 days					
	Withdrawal of food before exposure: 1 day					
	Body Length: 5.1 cm (4.8-5.7 range)					
	Body weight: 1.8 g (1.4-2.7 range)					
	Test design: 10 fish were used per concentration at					
	concentrations of 0, 100, 215, 464 and 1000 mg/l, and a pH					
	neutralized dose group of 1000 mg/l dose;					
	measured pH values:					
	concentration pH					
	(nominal, mg/L) beginning 24 h 48 h 72 h 96 h					
	464 10 5					
	1000 10.8					
	control 7.7 7.8 7.9 8.0 7.9					
	1000# 7.9 7.8 7.8 7.8 7.8					
	<pre># = test solution after pH-adjustment</pre>					

Result:	The D.O. concentration ranged between 8.1 and 8.4 mg/l at the beginning of the experiment. The doses used were chosen based on a range finding study. The product was added to the test water without any prior treatment. Subsequently, the fish were added to the water. Test vessel: All-glass aquarium non-sealed (30 x 22 x 24 cm) Dilution water chemistry: reconstituted freshwater was prepared from demineralized tap water that was resalted by the addition of 294.0 mg/l CaCl2.2H2O, 123.3 mg/l MgSO4.7H2O, 64.8 mg/l NaHCO3 and 5.8 mg/l KCl. Test water had a total hardness of 2.5 mmol/l, an acid capacity of 0.8 mmol/l and a pH of about 7.8. The water temperature was 20 degrees centigrade +/- 1 degree. The controls were the test water without the test substance and the pH adjusted control mentioned above. Reason for flagging this data: most sensitive study available on this endpoint Statistical evaluation: the number of doses tested did not allow for a Probit Analysis to be performed, since the data did not yield intermittent values between the LCO (~100 mg/L) and the LC100 (~215 mg/L). Therefore the geometric mean of the two concentrations was calculated: LC50 (96 h): 147 mg/L (nominal). The fish were found to respond to a positive control
	(chloroacetamide) with an LC50 of 32 mg/L after 48 hours. None of the fish in the test died in the 100 mg/L dose group. All fish in the 215, 464 and 1000 mg/L dose groups died. Thus, the LC50 was >100 and <220 mg/L (nominal), respectively 147 mg/L using the geometric mean of the two concentrations.
	Symptoms: apathy, tumbling.
Test substance: Reliability: Flag: 06-MAR-2003	The toxic effect may be due to the high pH of the test solutions: no adverse effects were observed, when testing a neutralized sample (1000 mg/L). CAS 100-37-8 (2-diethylaminoethanol), purity >99 % (1) valid without restriction guideline study Critical study for SIDS endpoint (42)
Species: Exposure period: Unit: LC50:	Pimephales promelas (Fish, fresh water) 96 hour(s) mg/l Analytical monitoring: no data = 1900
Method: GLP: Test substance:	other: not indicated no data other TS
Method:	The report consists of a QSAR study of the toxicity to the fathead minnow. In the report calculated results were compared with observed results

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Reliability:	(2) valid with restrictions			
	test method (observed 96 h LC50 to the fathead minnow			
	(Pimephales promelas)) not specified, but original reference			
	is well cited			
09-OCT-2001	(19) (20)			
Species: Exposure period:	Pimephales promelas (Fish, fresh water) 96 hour(s)			
Unit:	mmol/l Analytical monitoring:			
log LC50 :	1.18			
Polishility	(1) not assignable			
Reliability:	(4) Hot assignable			
01 TIT 2002	only secondary quotation (42)			
01-000-2002	(43)			
Species:	Pimephales promelas (Fish, fresh water)			
Result:	acute toxicity to the fathead minnow (Pimephales promelas):			
	-log (LC50; 96 h): 1.82			
Reliability:	(4) not assignable			
	only secondary quotation			
01-JUL-2002	(44)			

4.2 Acute Toxicity to Aquatic Invertebrates

Species: Exposure period: Unit: EC50:	Daphnia magna (Crusta 48 hour(s) mg/l = 165	cea) Analytical monitoring:	yes	
Method: Year: GLP: Test substance:	OECD Guide-line 202 1993 yes other TS: Diethylamino Code: 00129	ethanol, MRD-93-591, El	f Atochem Product	
Remark: Result:	reason for flagging this study: Although the complete test report is not available, the study can be regarded as valid, because the data for concentration-effect relationship were provided. The ECO is not clearly deducible, but the EC50 is comprehensible. The study was performed under pH-adjusted conditions at 1.6, 8, 40, 200, 600 1000 mg/l 2-diethylaminethanol.			
	<pre>pH measurements: da</pre>	y 0 (0h) day 7.3 7.3 7.3 7.3 7.3 7.3 7.3 7.3 7.3 7.3	<pre>2 (48h) 6.0 6.1 6.1 6.2 6.3 7.4 7.0</pre>	

The number of immobile daphnids were recorded at 24 h and 48 h: survival (n): 24 h 48 h control 20 19 1.6 mg/l 18 18 8 mg/l 19 19 40 mg/l 20 19 200 mg/l 14 7 600 mg/l 0 0 0 1000 mg/l Ω Test material was stable in test solutions with day 2 measurements being >/=83 % of day 0 measurements. All measured values were >/=88 % of nominal concentrations. Endpoint calculated using nominal concentrations. EC50 = 165 mg/L (95 % C.L. = 127-230 mg/L) Test substance: CAS 100-37-8 (2-diethylaminoethanol) Reliability: (2) valid with restrictions guideline study, but ECO not clearly deducible Critical study for SIDS endpoint Flag: 05-AUG-2002 (45) Species: Daphnia magna (Crustacea) **Exposure period:** 48 hour(s) Unit: mg/l Analytical monitoring: no EC0: = 62.5 EC50: = 83.6 EC100: = 250 Method: other: Directive 79/831/EEC, C2 Year: 1984 GLP: no Test substance: as prescribed by 1.1 - 1.4 reason for flagging this data: most sensitive study available Remark: on this endpoint Result: results after 24 h: EC0 = 62,5 mg/LEC50 = 92,4 mg/LEC100 = 250 mg/Lwater solubility: >500 mg/l at 293 K, 02-contents: > 2 mg/l, Test condition: illumination: day/night rhythm (16:8 h), light intensity: 5 uE at a wave of 400-700 nm, test volume: 10 ml, volume/animals: 2 ml, number of animals/vessel: 5, total number of animals/conc.: 20, age of animals: 2-24 h, check of the study: visually after 0, 3, 6, 24 and 48 h, concentration range: 7,81 - 500 mg/L Reliability: (1) valid without restriction Guideline study Flag: Critical study for SIDS endpoint

17-JUL-2002

(46)

<u>OECD SIDS</u>

4.3 Toxicity to Aquatic Plants e.g. Algae

Species: Endpoint: Exposure period:	Scenedesmus subspicatus (Algae) growth rate 72 hour(s)
Unit: NOEC:	mg/l Analytical monitoring: no = 5
LOEC:	= 10
EC10:	= 16
EC50:	= 44
Method:	other: Closely followed German Industrial Standard Guideline DIN 38 412, Part 9 using a static exposure
Year:	1988
GLP:	no
Remark:	reason for flagging this data: only study available on this endpoint
Result:	Endpoint: blomass
	EDCI0 $(72 \text{ II}) = 9.8 \text{ mg/L}$
	EbC90 (72 h) = 90.0 mg/L
	Growth factor control: 16
	Rises in pH of 1-2 units in the control treatments were probably associated with CO2 depletion from test media and do not invalidate the test, since in controls within 72 hours the minimum growth factor of 16 was achieved. In the highest test concentration a decrease in pH of 1.5 units was observed
Test condition:	A stem culture of Scenedsmus subspicatus was cultivated in nutrient medium of pH 8,2 according to DIN 38412, Part 9. The culture was reinoculated once a week. Three days before test initiation a preculture was inoculated at a cell density of 10000 cells/mL. Duration of the study test: 72 h; test temperature: 20 °C ± 1 °C; test flasks: test tubes containing 10 mL medium; inoculum density: 10000 exponential-growing cells/mL; test substance stock solution: 1 g/L sterile bidest. water; test concentrations: 5, 10, 20, 40, 80, 160 and 320 mg/L; replicates: per test concentration: 4, control: 8; illumination: permanent artificial light; light intensity: 10000 LUX white light or 120 μE*S-1m-2; test tubes were shaken twice a day to hold cells in suspension; measurements: fluorescence (interval: 24, 48 and 72 h; value of 0,12 ± 0,01 corresponds to 10000 cells/mL) and pH (interval: 0 and 72 h). pH-values: Control: 8.1 (0 h), 9.6 (72 h) 320 mg/L: 8.5 (0 h), 7.0 (72 h)
Test substance:	CAS 100-37-8 (2-diethylaminoethanol)
Reliability:	(1) valid without restriction
Flage	test procedure following national standards
19-JUL-2002	(47) (48)

4. ECOTOXICITY

4.4 Toxicity to Microorganisms e.g. Bacteria

Type: Species:	aquatic other bacteria: Acti plant treating munic	vated sludge fr ipal sewage.	com laboratory w	vaste water
Exposure period: Unit: EC20 :	30 minute(s) mg/l > 1000	Analytical r	nonitoring: no	
Method: Year: GLP:	Directive 87/302/EEC Activated sludge res 1994 yes	, part C, p. 11 piration inhib:	l8 "Biodegradat ition test"	ion:
Method:	Test mixture (total deg C for 30 min, af measured. Test mixtu sewage feed (as press from laboratory wast municipal wastewater The following controo - Inoculum blank: co inoculum (3 flasks) - Positive control: flasks, 1, 4, 10, 50	volume 250 mL) ter which the m re contained te cribed by OECD ewater treatmer ls were include ntrol without t 3,5-dichlorophe and 100 mg/L)	was incubated a respiration rate est substance (1 209) and activa it plants treati ed: test substance k enol with inocul	at 20 +/- 2 e was 2000 mg/L), ated sludge .ng put with Lum (5
Kemark:	No significant infib rate was observed ev mg/L). Disturbances in the sludge are not to be correctly introduced plants reason for flagging from municipal wwtp	biodegradation expected if th into adapted to this study: tes	process of actine test test concent test substance waste water treas st was performed	Avated tration (1000 vated tratment d with sludge
Result:	rat	e (mg O2/L.h)	Respiration	% inhibition
	Test substance Inoculum blank Reference subst. 1 mg/L	28 30 32	7 - -7 (stimulat	ion)
	4 mg/L 10 mg/L 50 mg/L 100 mg/L	28 22 8	7 27 73	
Test substance:	Limit test without r mg/L. At 1000 mg/L r to the control mean CAS 100-37-8 (2-diet	eplication (3 c espiration was value. hylaminoethanol	control replicat inhibited by 7 l), purity 99.9	tes) at 1000 % compared %
Flag: 24-JUN-2002	Guideline study Critical study for S	IDS endpoint		(49)
Species: Unit:	Pseudomonas putida mg/l	(Bacteria) Analytical r	monitoring: no	

LOEC :	= 375	
Method: GLP:	other: cell growth inhibition test no data	
Reliability:	(4) not assignable	
09-OCT-2001	Original reference not available (50)
Species: Exposure period: Unit: EC10:	<pre>other bacteria: industrial activated sludge from wwtp of BASF 30 minute(s) mg/l Analytical monitoring: > 1995</pre>	
Method:	other: Activated Sludge Respiration Inhibition Test	
Remark: Reliability:	no inhibition of the respiration was observed up to 1995 mg/L, but a stimulation (2) valid with restrictions comparable to guideline study with acceptable restrictions.	
09-OCT-2001	(39)

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

Species:	other terrestrial plant: Chrysanthemum morifolium CV "Indianapolis white"
Endpoint:	other: chlorosis
Expos. period:	22 day(s)
Unit:	mg/l
EC50:	= .12
Method:	other
GLP:	no data
Test substance:	other TS: 2-diethylaminoethanol (DEAE); purity: >99 %
Remark.	plants were grown individually in potting-soil consisting of 1
Result:	plants were grown individually in potting-soft consisting of 1 part peat moss : 2 parts vermiculite, amended with lime. A fertilizer was applied at each watering at the rate of 15:1 (water:fertilizer). All plants, with the exception of tomatoes, were grown in 725 ml volume plastic pots (tomatoes: 400 ml volume). Glasshouse experiments were performed at 28 °C +-2°C with a minimum light intensity (10.700 lux; incident light and cool white fluorescent) and 16-h photoperiod. DEAE was introduced into plants by three methods: (1) ten-fold aqueous dilutions of DEAE were prepared so that 100 ml of solution contained 1E0-1E-5 ml DEAE and each pot received a single application of 100 ml of solution (2) soil in which plants were grown was autoclaved with DEAE-containing steam (3) plants were grown in controlled-environment chambers in either non-sterile or DEAE-steam-sterilized soil and in which atmospheric humidity was enhanced by DEAE-containing steam. reason for flagging this publication: reliable data on this endpoint DEAE was found to cause chlorosis in young leaves of chrysanthemum, tomato, corn and bean. Sensitivity of corn and chrysanthemum to DEAE was cultivar-dependent: chrysanthemum cultivar "Indianapolis White" was sensitive to low concentrations of DEAE, while "Bonnie Jean", "Velvet Ridge" and "Mistletoe" were less so, and corn "3XD50" was sensitive to DEAE while "Ohio 28" was not. The presence of latent chrysanthemum chlorotic mottle viroid (ChCMV-L) increased the sensitivity of "Bonnie Jean" and "Velvet Ridge" to DEAE. Infection of tomatoes with mild or severe strains of potato spindle tuber viroid (PSTV) resulted in increased sensitivity to DEAE under conditions of environmental stress. Leaves of chrysanthemum exhibition DEAE-induced chlorosis contained higher, but non-toxic, levels of iron and other minerals than those from non DEAE-treated plants.
	Chrysanthemum cultivars varied in their sensitivity to DEAE. Symptoms were generally restricted to leaves which developed after initiation of DEAE exposure, and included chlorosis, epinasty, stunting and necrosis.

OECD SIDS2-DIETHYLAMINOETHANOL4. ECOTOXICITYDATE: 07-MARS-2003SUBSTANCE ID: 100-37-8

Reliability:	<pre>All "Indianapolis White" (4/4) exhibited chlorosis 7 days after exposure to 1E-2 ml DEAE added to non-sterile soil (725 ml pot). At 1E-4 ml/pot, only 50 % (2/4) of the plants were chlorotic after 22 days. (2) valid with restrictions acceptable, well-documented publication which meets basic scientific principles</pre>
Flag: 02-JUL-2002	Critical study for SIDS endpoint (51)
Species:	other terrestrial plant: Chrysanthemum morifolium
Method: GLP:	other no data
Result: Reliability:	Certain varieties of Chrysanthemum morifolium are particularly sensitive to DEAE; 1E-4 ml DEAE per 12.4 cm pot of soil caused chlorosis of the variety Indianapolis White in 22 days. Other plants found to be sensitive to DEAE are Licopersicum esculentum var. Rutgers, and Morus rubra. Ulmus americana, Zea mays, and Oryza sativa grown in controlled environment chambers displayed symptoms similar to those caused by DEAE in other plants. Symptoms usually restricted to new leaves produced after contact with DEAE, include interveinal chlorosis, mottling, complete chlorosis, albinism, deformation, and size reduction depending upon species, variety, dosage, and length of exposure. High concentrations of DEAE caused marginal necrosis of fully mature leaves and death of the growing point. Chrysanthemum plants infected with the symptomless strain of chrysanthemum chlorotic mottle were at least 10 times more sensitive to DEAE than are uninfected plants. (4) not assignable
01-JUL-2002	only abstract (52)
Species:	other terrestrial plant: Phaseolus vulgaris (pinto bean)
Remark:	during the winter of 1972, greenhouse and chamber grown pinto beans (Phaseolus vulgaris) began to display symptoms consisting of dark, bifacial necrotic spots, developing initially along the margins and later in the central portion of the leaves. Marginal spots often coalasced as severity increased. The uniform response suggested toxicant or nutrient involvement. Live steam used for humidity control in both greenhouse and growth chambers was suspect, and follow-up studies indicate that either of the two compounds (cyclohexylamine or 2 hydroxytriethylamine) used to control the pH of the steam
Reliability:	condensate were associated with the observed injury. (4) not assignable
08-JUL-2002	abstract only (53)

4.6.3 Toxicity to Soil Dwelling Organisms

4.6.4 Toxicity to other Non-Mamm. Terrestrial Species

4.7 Biological Effects Monitoring

4.8 Biotransformation and Kinetics

4.9 Additional Remarks

Memo:	toxicity on frog tadpoles (Rana bravipoda porosa)
Remark:	the toxicity of solvents used in prepn. of agrochem was tested by detn. of the LC50 (median lethal concn.) of tested solvents on frog (Rana bravipoda porosa) tadpoles in freshwater at 25 °C.
Result:	LC50 (3 h): 360 ppm LC50 (6 h): 230 ppm LC50 (12 h): 300 ppm LC50 (24 h): 230 ppm LC50 (48 h): 85 ppm
Test substance:	Diethylethanolamine
Reliability:	(4) not assignable only abstract in English available (test method and endpoint remains unclear)

12-JUL-2002

(54)

5.0 Toxicokinetics, Metabolism and Distribution

Type:

Toxicokinetics

Remark: See chapter 5.11 26-JUL-2002

5.1 Acute Toxicity

5.1.1 Acute Oral Toxicity

Type: Species: Sex: Vehicle: Doses: Value: Method: Year: GLP: Test substance:	LD50 rat male/female water 177, 708, 1106 and 1416 mg/kg ca. 1320 mg/kg bw other: BASF-Test 1969 no as prescribed by 1.1 - 1.4
Method:	The test article was administered by gavage. Ten male and 10 female rats were used per dose. The substance was dissolved in water and administered at concentrations of 2 to 20% test substance in water. The doses used were 177, 708, 1106 and 1416 mg/kg bw. Animals were observed over a period of 7 days after administration of the test article.
Remark:	Other studies were available reporting essentially similar values. Experiments conducted with the neutralized compound showed toxicity in the range of 5000 to 8000 mg/kg bw.
Result:	No animals died within the lowest dose group, but 1/20, 0/20 and 14/20 died in the 708, 1106 and 1416 mg/kg dose groups, respectively. No animals died in the first hour after administration of the test substance, but most deaths occurred within the first 24 hours. The clinical signs were described as apathy and dyspnea. No clinical signs were noted in the lowest dose group; after the first day the 708 mg/kg dose group was symptom free; after 3-4 days no symptoms were observed in the 1106 and 1416 mg/kg dose groups. Necropsy: Hemorrhaging of the stomach and intestines was observed in the animals that died. The animals that survived to the end had no unusual findings except for chronic bronchitis in 2 animals of the 708 mg/kg dose group. The original LD50 value was reported to be: LD50 ca. 1500 ul/kg.
Test substance:	2-diethylaminoethanol, purity: ca. 99%.
Reliability:	(2) valid with restrictions Chosen as key study for ICCA Robust Summaries since essential details were available.
riag: 10-JAN-2003	(55)

OECD SIDS 5. TOXICITY

T.D50 Type: Species: rat Sex: male Value: = 1300 mg/kg bw Method: other GLP: no Test substance: other TS Remark: the TS was not neutralized Test substance: 2-diethylaminoethanol, purity not indicated Reliability: (2) valid with restrictions essential details given 05-OCT-2001 (56) LD50 Type: Species: rat Value: ca. 2460 Method: other: no data GLP: no Test substance: other TS Method: test substance given as a 10% solution Test substance: 2-diethylaminoethanol, purity not indicated Reliability: (4) not assignable secondary literature, essential details lacking 05-OCT-2001 (57) LD50 Type: Species: rat Sex: no data Vehicle: water Value: ca. 1300 mg/kg bw Method: other: no data GLP: no Test substance: other TS Remark: Original value: LD50 was in the neighbourhood of 1.3 g/kg. Late deaths were observed. Deaths occurred between 24 h (for high doses) and 5 days (for lower doses) after dosing. Slight apathy was the only clinical sign. Considerable irritation of the intestinal tract, red bile, congested liver, and pale kidney with congested spleen were noted. The page of the TSCAT with this data was dated Feb. 11, 1942 and signed by Henry F. Smyth, Jr. Test substance: 2-diethylaminoethanol, purity not indicated Reliability: (4) not assignable Essential details not given, but the data fits with other existing data. 05-OCT-2001 (58) LD50 Type: Species: rat Sex: male Value: ca. 8000 mg/kg bw

OECD SIDS 5. TOXICITY

Method: GLP: Test substance:	other no other TS
Remark:	Observation period was 3 days. In one study 3/10 animals died, and in the other 6/10 died. Thus, 9/20 animals died in total.
Test substance:	2-diethylaminoethanol neutralized with HCl prior to administration.
Reliability:	<pre>(4) not assignable The study is limited since the observation time was only 3 days.</pre>
17-JUL-2002	(59)
Type: Species: Sex: Value:	LD50 rat male ca. 5600 mg/kg bw
Method: GLP: Test substance:	other no other TS
Method:	According to the report, the approx. LD50 was determined by "feeding" logarithmically graded doses to groups of 5 male rats. In the results section it clarifies that intubation was used. The observation period was 14 d. Calculations were made using the method of Weil (Biometrics, 8, 249-263, 1952).
Result: Test substance:	The 95% confidence interval was reported to be 3.5-9.1 g/kg. 2-diethylaminoethanol neutralized with HCl prior to
Reliability:	(2) valid with restrictions
02-OCT-2001	(60) (61)

5.1.2 Acute Inhalation Toxicity

Type: Species: Sex: Exposure time:	other: Inhalation Hazard Test rat male/female 8 hour(s)
Method:	other: BASF-Test
GLP:	no
Test substance:	as prescribed by 1.1 - 1.4
Method:	The animals were exposed to a 20°C highly saturated vapor-air-mixture for 1, 3 or 8 hours. A total of 6 animals per sex were used for the one hour experiment, and a total of 3 animals per sex were used for the 3 and 8 hour experiments.
Result:	No animals (0/12) died within a 7 day period following the 1 hour exposure. In the 3 hour test 1 animal died after 3 days, and 5/6 had died by 9 days after exposure. The last rat remained alive until the end of the observation period (14 days). In the 8 hour test, 2 animals died during exposure, and by day 7 only one animal remained alive. This animal was sacrificed on day 18.

5. TOXICITY	DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8
	Clinical signs: Severe signs of irritation were noted, namely, attempts to escape, mucous membrane irritation, dyspnoea, gasping.
	Necropsy: Corrosion of the snout, eyes, ears and front paws were noted in the animals that were exposed to the vapors for 3 hours.
	Since the vapor pressure of the substance at 20°C is 1.9 mbar, the saturated vapor concentration is about 9.2 mg/l. Worst case graphical extrapolation from a probability sheet using the lethality rates after 1 and 3 hours of exposure results in an estimated LT50 of about 2.4 hours. From this a 4 hour LC50 of ca. 4.6 mg/l (4600 mg/m3) can be calculated using Haber's rule.
Test substance: Reliability:	CAS Nr. 100-37-8 (2-diethylaminoethanol), purity: ca. 99% (2) valid with restrictions Essential details are given for an Inhalation Risk Test; chosen as the key study since a better study to determine the LC50 was not available. The only other acute inhalation toxicity available that reported an LC value in rats was an LCLo reported as 4.5 mg/l (4500 mg/m3) in 4 h and in mice an LC50 of 5 mg/l (5000 mg/m3) with no information on exposure time. Both values were reported in RTECS Update No. 9107 (through August 1991) and the original report (Gigiena Truda i Professional 'nye Zabolevaniya, 14 (11), E2. 70) was not available for an available
Flag: 10-JAN-2003	Critical study for SIDS endpoint (55)
Type: Species: Exposure time: Value:	LCLo rat 4 hour(s) 4.5 mg/l
Method: GLP: Test substance:	other: no data no other TS
Test substance: Reliability:	2-diethylaminoethanol, purity not indicated (4) not assignable
02-OCT-2001	secondary literature, essential details lacking (62)
Type: Species: Sex:	other: IHT rat male
Exposure time:	4 hour(s)
Method: GLP: Test substance:	other: no data no other TS
Method:	Groups of 6 albino rats were exposed to an atmosphere that had been saturated with the volatile component of the TS generated at room temperature. After exposure animals were held for 2 weeks.

2-DIETHYLAMINOETHANOL

No mortality occurred after a 4 hour exposure.

Result:

OECD SIDS

OECD SIDS 5. TOXICITY

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Test substance:	2-diethylaminoethanol, purity not indicated	
Reliability:	(2) valid with restrictions essential details are given for a so-called Inhalation Hazard Test	
27-JUN-2002	(5)	6)
Type: Species: Exposure time:	other: IHT rat 8 hour(s)	
Method: GLP:	other: no data no	
Test substance:	other TS	
Remark:	One fifth of the exposed rats (no data on number) died by 8 hours inhalation of an atmosphere that had been saturated at 25 degrees Centigrade with the volatile part of the compound No mortality occurred at 4 hours. Death was delayed, probably due to liver and kidney injury. Eye and nose irritation was seen, but no narcosis. The page of the TSCAT with this data was dated Feb. 11, 1942 and signed by Henry F Smyth Jr	•
Test substance: Reliability:	 2-diethylaminoethanol, purity not indicated (2) valid with restrictions essential details are given for a so-called Inhalation 	
05-OCT-2001	Hazard Test (5)	8)
Type: Species: Exposure time: Value:	LC50 mouse unspecified 5 mg/l	
Method: GLP: Test substance:	other: no data no other TS	
Test substance: Reliability: 05-OCT-2001	<pre>2-diethylaminoethanol, purity not indicated (4) not assignable secondary literature, essential details lacking (6)</pre>	2)
Type: Species: Exposure time: Value:	LC50 mouse 2 hour(s) ca. 5 mg/l	
Method: GLP: Test substance:	other: no data no data other TS	
Remark: Test substance: Reliability:	Original value: LC50 (2 h) = 5000 mg/m3 (1050 ppm). 2-diethylaminoethanol, purity not indicated (4) not assignable	
02-OCT-2001	secondary literature, essential details lacking (6	3)

5.1.3 Acute Dermal Toxicity

Type: Species:	LD50 guinea pig
Value:	ca. 885 mg/kg bw
Method: GLP:	other no
Test substance:	other TS
Method:	Penetration of Guinea Pig Skin Method: Doses differed by a multiple of 10. Six animals were used per dose. The sex was not specified. The sample was retained on the abdomen with absorbent cotton over the clipped area for 4 days. The animals were observed for a maximum of 14 days. The dose was retained by covering the cotton with a film of rubber, vinyl or "the like".
Remark:	Original value LD50 = 1.0 ml/kg.
Test substance:	2-diethylaminoethanol, purity not indicated
Reliability:	(2) valid with restrictionsEssential details given; not performed according to today's guidelines, but useful for acute dermal toxicity risk.Chosen as the key study since more details were available.
Flag:	Critical study for SIDS endpoint
13-JAN-2003	(56)
Type:	LD50
Species: Value:	ca. 1100 mg/kg bw
Method:	other
GLP: Test substance:	no other TS
reper pubbedneer	
Remark:	The TLV documentation reads, "Smyth comments that further study indicatesa rabbit skin penetration LD50 of 1.26 (0.85-1.87) ml/kg undiluted. This value is very close to the value reported in a test with guinea pigs performed by Smyth & Carpenter (J. Ind. Hyg. Toxicol., 26, 269-273, 1944), namely a LD50 = 1.0 ml/kg (ca. 885 mg/kg bw). This rabbit data is also cited in a TSCAT from Union Carbide (Date Produced: 3-02-84), which makes reference to Rpt.16-102, 1953.
Test substance: Reliability:	2-diethylaminoethanol, purity not indicated
	Secondary literature, essential details are missing. This value should be considered only in the light of the guinea pig data generated by Smyth and Carpenter (1944); this data is mentioned since the rabbit (or rat) is usually the species of choice.
Flag:	Critical study for SIDS endpoint
26-OCT-2001	(57) (58)
Type: Species: Sex: Vehicle:	LD50 guinea pig no data other: none

OECD SIDS 5. TOXICITY

Value:	ca. 885 mg/kg bw
Method:	other
GLP:	no
Test substance:	other TS
Remark:	Original value: LD50 = 1 g/kg. Application of the undiluted test substance for 4 days, local necrosis was observed. The page of the TSCAT with this data was dated Feb. 11, 1942 and signed by Henry F. Smyth, Jr.
Test substance:	2-diethylaminoethanol
Reliability:	(2) valid with restrictions
	Some essential details are missing, and this report would not be reliable standing alone. Necessary details are available when Smyth & Carpenter (J. Ind. Hyg. Toxicol. 26, 269-273, 1944) is considered.
05-OCT-2001	(58)

5.1.4 Acute Toxicity, other Routes

Туре:	LD50	
Species:	rat	
Route of admin.:	i.p.	
Value:	= 1220 mg/kg bw	
Method:	other: no data	
GLP:	no	
Test substance:	other TS	
Remark:	Application in neutralized form.	
Test substance: 27-JUN-2002	2-diethylaminoethanol, purity not indicated	(61)
Type:	LD50	
Species:	mouse	
Route of admin.:	i.p.	
Value:	ca. 160 mg/kg bw	
Method:	other: BASF-Test	
GLP:	no	
Test substance:	as prescribed by 1.1 - 1.4	
Remark:	The original data was given as: LD50 ca. 180 ul/kg	
Test substance:	CAS Nr. 100-37-8 (2-diethylaminoethanol), purity: ca. 99%	\
27 - JUN - 2002		(55)
Туре:	LD50	
Species:	mouse	
Route of admin.:	i.p.	
Value:	192 mg/kg bw	
Method:	other: no data	
GLP:	no	
Test substance:	other TS	
Test substance:	2-diethylaminoethanol, purity not indicated	
05-OCT-2001		(64)
Type:	LD50	

OECD SIDS 5. TOXICITY

Species: mouse Route of admin.: i.p. Value: 308 mg/kg bw other: no data Method: GLP: no data Test substance: other TS Test substance: 2-diethylaminoethanol, purity not indicated 05-OCT-2001 (65) LD50 Type: Species: mouse no data Sex: Vehicle: no data Route of admin.: s.c. Value: = 650 mg/kg bwother: no data Method: GLP: no Test substance: other TS Test substance: 2-diethylaminoethanol 16-AUG-2000 (66) LD50 Type: Species: mouse Sex: no data Vehicle: no data Route of admin.: s.c. Value: = 1561 mg/kg bw Method: other: no data GLP: no Test substance: other TS Test substance: 2-diethylaminoethanol 18-SEP-2000 (67) (65) Type: LD50 Species: mouse Route of admin.: i.m. Value: 416 mg/kg bw other: no data Method: GLP: no other TS Test substance: Test substance: 2-diethylaminoethanol, purity not indicated 05-OCT-2001 (68) Type: LD50 Species: mouse Route of admin.: i.v. Value: 188 mg/kg bw Method: other: no data GLP: no
Test substance: other TS

Test substance: 2-diethylaminoethanol, purity not indicated 05-OCT-2001

(68)

5.2 Corrosiveness and Irritation

5.2.1 Skin Irritation

Species: Exposure Time: Result:	rabbit 4 hour(s) corrosive
Method: Year: GLP:	OECD Guide-line 404 "Acute Dermal Irritation/Corrosion" 1981 no
Test substance:	as prescribed by 1.1 - 1.4
Method:	One day prior to the substance administration, the animals (New Zealand rabbits) were shaved to prepare sites for substance application. Dorsal and lateral sections of the trunk were used as the application sites (3x3 cm per application site). 0.5 ml/patch were used. Two males and four females were used for the 3 min application time. Three males and three females were used for 1hr and 4 hr application times. Each animal had 4 sites of application (except for the 3 min application: 2 application sites: occlusive and semi-occlusive), namely, a site for:
	1 hr application, occlusive 4 hr application, occlusive 1 hr application, semi-occlusive 4 hr application, semi-occlusive
	After treatment the patches were removed, and the treated area was rinsed with a polyethylene glycol 400 solution or a polyethylene glycol 400/water (1:1) solution and dried. Observations were made 1 hr after the removal of the patch, and at 24, 48 and 72 hr, as well as 7 days after start of application.
Result:	Note: In accordance to OECD Guideline 404 of 1981, "Acute Dermal Irritation/Corrosion", a 4 hour application time was used, but additional application times (3 min and 1 h) were also used. After a 3 min occlusive exposure, as well as semi-occlusive
	exposure only slight erythema was observed. It was reversible by the end of the experiment. Some scaling was observed on day 7.
	After 1 hr and 4 hr occlusive and semi-occlusive exposure severe erythema, eschar formation and necrosis were observed. Very slight to slight edema was also observed. By day 7 after application the findings were not reversed.

	The score for erythema after the 4 hr occlusive expose always a 4 at all times, with the exception of one re 3 at 1h. The scores for edema after the 4 hr occlusion exposure was 2 in all animals 1 hr after patch remove after the start scores of 2 were seen in 5 animals and of 3 was seen in 1 animal; 48 hr after the start score were seen in 2 animals, and scores of 2 were seen in animals; the same scores were seen at 72 hr after the and at 7 days 3 animals had a score of 1, and 3 had a 2.	eading of two al; 24 hr dd a score ores of 1 4 e start; a score of
	The scores for erythema after the 4 hr semi-occlusive were reported to be: 2-3 at 1 hr after patch removal; 3-4 at 24 hr after t and thereafter it was observed to have a score of 4. scores for edema after the 4 hr semi-occlusive exposu reported to be:	e exposure he start; The re were
	in 3 animals and 2 in 3 animals one in after patch i animals and 2 in 3 animals 24 hr after the start animals and 2 in 4 animals 48 hr after the start; 1 i animals and 2 in 2 animals 72 hr after the start; and after the start a score of 1 was reported in all 6 ar	imals.
	Necrosis was confirmed histologically. In this regar the semi-occlusive 1 hr application group full thickn necrosis was seen in 2/6 animals; in the 4 hr applica group full thickness necrosis was seen in 5/6 animals the occlusive 1 hr application group full thickness n was seen in 3/6 animals; in the 4 hr application grou thickness necrosis was seen in all animals.	rd, in less ation s. In lecrosis up full
Reliability:	(1) valid without restriction Well documented, guideline study, conducted under GLE conditions. Chosen as key study for ICCA Robust Summa since it was the best study available.	2-like aries
26-JUL-2002	clitical study for SIDS enupoint	(69) (70)
Species: Result:	rabbit corrosive	
Method:	Draize Test	
Test substance:	as prescribed by 1.1 - 1.4	
Reliability:	(2) valid with restrictions	
05-OCT-2001	Essential details given.	(71) (55)
Species:	rabbit	
Method: GLP: Test substance:	other no other TS	
Remark:	The TS was applied to shaved skin of the belly of an rabbit. Observations were made after 24 hours. Its was comparable to morpholine.	albino effect
Test substance:	2-diethylaminoethanol, purity not indicated	

Reliability:	(4) not assignable
05 075 0001	Essential details not given.
05-0CT-2001	(56)
Species.	rabbit
Exposure:	Occlusive
Exposure Time:	4 hour(s)
Result:	corrosive
Method:	other
GLP:	no data
Test substance:	other TS
Method:	Applied volume was 0.5 ml. Observations went out to 14
	days. Number of animals used:
	2 for the 50% solution
	2 for the 25% solution
	4 for the 10% solution
	6 for the 5% solution
Remark:	50% (w/w) and 25% (w/w): severe erythema and necrosis on 2
	of 2 from 0.5 ml; ulceration observed from the 50% dilution
	10% (w/w): minor to moderate erythema on 6 of 6 rabbits,
	minor edema on 5, ulceration and necrosis on one from 0.5
	ml; 5 rabbits normal at 3 days.
	5% (w/w): minor erythema on 4 of 6 rabbits, minor edema on 3
Test substance.	Irom U.5 ml; all rabbits normal at 2 days.
Peliability.	(2) valid with restrictions
Kerrubrite,.	Essential details given
05-OCT-2001	(72)
Species:	rabbit
Concentration:	undiluted
Exposure:	Occlusive
Result:	corrosive
Method.	other
GLP:	no data
Test substance:	other TS
Method:	Six animals (3 males and 3 females) were used for each
	duration. 0.5 ml was used per treatment. The sample was
	applied as received (i.e. undiluted).
Result:	One hour contact: Moderate to severe erythema and edema on 6
	of 6 rabbits, full thickness necrosis on 6, ulceration on 5,
	ecchymosis on 4 and scabs on 3 from 0.5 ml; animals
	sacrificed at 2 days for numane reasons.
	rabbita minor edema en 2 auperficial negregia en 2
	ulceration on 2 ecchymosis on 3 scabs on 3 desquamation
	on
	4 and alopecia on 3 from 0.5ml; irritation resisted on 4
	through 14 days.
Test substance:	2-diethylaminoethanol, purity not indicated
Reliability:	(2) valid with restrictions
	Essential details given.
05-OCT-2001	(73)
6	
species:	raddit

Concentration: undiluted Occlusive Exposure: Exposure Time: 4 hour(s) No. of Animals: 1 Result: corrosive Method: other: according to 29CFR191.11 GLP: no Test substance: other TS Remark: Five hundred milligrams of the test substance was applied onto the dry fur-clipped skin of one albino rabbit. After a 24-hour contact period, the application patch was removed and the skin site was washed. The treated site was examined for corrosive effects for at least three days. Necrosis was observed. Result presented only in tabular form. Date of original report: 6/28/72. Test substance: 2-diethylaminoethanol, purity not indicated Reliability: (4) not assignable Essential details of the results are not given. However, the data is plausible. 05-OCT-2001 (74)rabbit Species: Concentration: undiluted Result: irritating Method: other: no data GLP: no Test substance: other TS Remark: On the belly of the rabbit, the undiluted test substance produced erythema. No skin reaction was seen with a 10% solution. No further details were given. The page of the TSCAT with this data was dated Feb. 11, 1942 and signed by Henry F. Smyth, Jr. Test substance: 2-diethylaminoethanol, purity not indicated Reliability: (4) not assignable Essential details not given. 05-OCT-2001 (58) Species: rabbit Concentration: no data Exposure: no data 4 hour(s) Exposure Time: No. of Animals: 6 other: according to 49 CFR 173.240 Method: GLP: no Test substance: other TS Remark: The test substance was applied to six albino rabbits for a 4-hour contact period. Skin sites were observed for 48 hours. At the end of the 4-hour contact period, the treated skin sites were dark grey in color, with a "red surround" and with sloughed patches of surface epithelium. Subsequently, the sites became thick (hypertrophy) and dry, but remained supple. No signs of dermal destruction were observed.

SUBSTANCE ID: 100-37-8

Test substance:	According to the authors, the test substance was classified as noncorrosive (DOT). No further details given. The pages with the data for this test were dated 8/01/77. 2-diethylaminoethanol, purity not indicated
Reliability:	(4) not assignable
05-OCT-2001	(75)
Species: Concentration: Exposure: Exposure Time: No. of Animals: Result:	guinea pig 5 % Occlusive 24 hour(s) 10 not irritating
Method: GLP: Test substance:	other no data other TS
Method:	This investigation was a part of a guinea pig sensitization study (preliminary irritation test). Five male and 5 female Dunkin Hartley guinea pigs were treated for 24 hr with a patch containing the test substance at a concentration of 5%
Result: Test substance: Reliability:	<pre>Slight, well-defined erythema was seen in 1 animal, the remaining 9 animals exhibited no skin response. 2-diethylaminoethanol, purity not indicated (2) valid with restrictions Reliable for the purpose of the test; i.e. preliminary</pre>
05-OCT-2001	information for a sensitization test. (76)
Species: Result:	human irritating
Method: GLP: Test substance:	other: BASF-Test no as prescribed by 1.1 - 1.4
Reliability: 02-OCT-2001	(4) not assignable essential details are not given (77)
5.2.2 Eye Irritation	
Species: EC classificat.:	rabbit risk of serious damage to eyes
Method: GLP: Test substance:	other: BASF-Test no as prescribed by 1.1 - 1.4
Method:	50 µl of the undiluted liquid was applied to the eye. It was not washed out. Two animals were used. As a control, a

OECD SIDS	2-DIETHYLAMINOETHA	NOL
5. TOXICITY	DATE: 07-MARS-20	03
	SUBSTANCE ID: 100-3	7-8
Result:	Between one and 24 hours after application corrosion of th conjunctiva and eyelids was seen. It was not reversible after 8 days. Irreversible damage to corneal tissue was observed (staphyloma).	le
Test substance: Reliability:	CAS Nr. 100-37-8 (2-diethylaminoethanol), purity: ca. 99% (2) valid with restrictions Essential details are given. Chosen as key study for ICCA Robust Summaries since more details were available.	
10-JAN-2003	critical study for SIDS enaporne	(55)
Species: Result:	rabbit irritating	
Method:	other: no data	
Test substance:	other TS	
Remark: Test substance: Reliability:	5 mg; strong irritant 2-diethylaminoethanol, purity not indicated (4) not assignable Secondary literature; essential details not available.	
05-OCT-2001	-	(78)
Species:	rabbit	
Method: GLP: Test substance:	other no other TS	
Remark:	Applied to the cornea undiluted. The effect was compared	to
Test substance: Reliability:	ammonium hydroxide. 2-diethylaminoethanol, purity not indicated (4) not assignable Essential details of results not given.	
05-OCT-2001		(56)
Species:	rabbit	
Method: GLP:	other: no data no	
Test substance:	other TS	
Remark: Test substance:	The TLV documentation states, "he regards the major industrial hazard to be eye injury from the fluid (very severe in the rabbit from 0.005 ml undiluted, severe from 15% or more in glycol and not severe from 5% in glycol." 2-diethylaminoethanol, purity not indicated	
Reliability:	(4) not assignable Secondary literature, essential details lacking.	
05-OCT-2001		(57)
Species: EC classificat.:	rabbit risk of serious damage to eyes	
Method:	other	
GLP: Test substance:	no data other TS	
TODE DANBEATER.		

OECD SIDS	2-DIETHYLAMINOETHANOI
5. TOXICITY	DATE: 07-MARS-2003
	SUBSTANCE ID: 100-37-8
Result:	100% (undiluted), 50% (w/w) and 25% (w/w); 0.1ml: severe corneal injury (with vascularization and irregular shape), iritis and severe conjunctival irritation (with necrosis); injury persisted through 21 days.
	100% (undiluted), 50% (w/w) and 25% (w/w); 0.005ml: pinpoint pupils from undiluted sample; moderate to severe corneal injury (with instances of vascularization and irregular shape), iritis and severe conjunctival irritation (with necrosis); injury from undiluted sample persisted through 21 days; 9 of 10 eyes receiving 50% or 25% healed by 21 days.
	10% (w/w) and 5% (w/w); 0.005 ml: minor to moderate corneal injury, iritis and moderate to severe conjunctival irritation (with instances of necrosis from 10%); all eyes
Test substance: Reliability:	2-diethylaminoethanol, purity not indicated (2) valid with restrictions
25-JUL-2002	available and purity is not indicated. (72)
Species:	rabbit
Method.	other: no data
GLP:	no
Test substance:	other TS
Remark:	In the rabbit eye, 0.001 ml of the test substance caused necrosis. The date on the page where this data was given is Feb. 11, 1942, and it is signed by Henry F. Smyth, Jr.
Test substance: Reliability:	2-diethylaminoethanol, purity not indicated (4) not assignable
05-OCT-2001	Essential details not given. (58)
Species.	rabbit
Concentration:	undiluted
Dose:	.1 ml
NO: OI Animais.	
Method:	other: no data
Test substance:	other TS
Method:	One tenth ml of sample was instilled into the conjunctival sac of one eye of each of six albino rabbits. The treated eyes of three rabbits were washed at 20 to 30 seconds after
	instillation for ca. 1 minute; the treated eyes of the remaining three rabbits remained unwashed. Irritation reactions were scored for 7 days.
Remark:	Date of original report: 8/01/77, performed by Pharmacology Research, Inc.
Result:	In unwashed eyes the cornea opacified promptly and
	completely obscured the pupil and iris. The conjunctivae became completely necrotic (black) with minimal swelling. Patches of necrosis were seen on the eyelids. No signs of recovery were seen after 7 days. In washed eyes, corneal opacification developed more slowly. During the 1st 2 hours the pupil was constricted and the iris was severely

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003

SUBSTANCE ID: 100-37-8

	congested. At 3-4 hours the iris reacted slowly to light;
	subsequently, it failed to react at all. Initially, the
	conjunctivae were severely inflamed with minimal swelling and
	with patches of necrosis. They were completely necrotic
	after 24 hours. No signs of recovery were seen after 7 days.
	According to the authors, the TS was considered corrosive
	by both washed and unwashed applications.
Test substance:	2-diethylaminoethanol, purity not indicated
Reliability:	(2) valid with restrictions
	Some essential details given.
27-JUN-2002	(75)

5.3 Sensitization

Type: Species:	Guinea pig maximization test guinea pig		
Concentration 1st:	Induction 5 % intracutaneous		
2nd:	Induction 25 % occlusive epicutaneous		
3rd:	Challenge 5 % occlusive epicutaneous		
No. of Animals:	20		
Vehicle:	other: saline or ethanol		
Result:	not sensitizing		
Method:	other: according to Magnusson, B. and Kligman, A.M.: J. Invest. Derm. 52, 268		
Year:	1969		
GLP:	no data		
Test substance:	other TS		
Method:	The method met the main requirements of OECD Guideline 406. Ten male and 10 female Dunkin-Hartley guinea pigs were used. For induction, the animals were given intradermal injections into 2 sites each of clipped shoulder skin followed by a 48-hour application 7 days later. After removal of the 48-hour patches, the application sites were washed. Epicutaneous challenge was performed by a 24-hour patch at 14 days after epicutaneaous induction (i.e. 21 days after beginning of the study); the challenge patch was applied to a previously untreated site. Observation times after challenge: 24 and 48 h after removal of occlusive dressings. For the positive control, 10 guinea pigs were treated with 1-chloro-2,4-dinitro-benzene (DNCB) using a concentration of 0.1% for induction (intradermal and topical) and challenge (topical).		
Result:	None of the animals treated with the test substance (0/20) exhibited skin responses. All 10 positive control animals challenged with 0.1% DCNB showed a clear skin response. Except for one animal which had a score of 1 (meaning weak), all irritation control animals were free of skin responses.		
Test substance:	2-diethylaminoethanol, purity not indicated		
Reliability:	(2) valid with restrictions		
	Well documented study. Essential details were given, but purity is not indicated. This study was chosen as the key study for the ICCA Robust Summaries since it was the most recent. Other studies were also available which showed the substance to be non-sensitizing.		
Flag:	Critical study for SIDS endpoint		
ZO -u ull - ZUUZ	(76)		

Type:	Draize Test
Species:	guinea pig
Concentration 1st	: Induction .1 % intracutaneous
2nd	: Challenge .1 % intracutaneous
	10
No. of Animals:	10
Vehicle:	water
Result:	not sensitizing
Method:	other: according to Draize et al.: J. Pharmacol. Exp. Ther. 82, 377-390
Year:	1944
GLP:	no
Test substance:	other TS
Method:	Ten guinea pigs were used. Induction: intradermal injections of 0.1% aqueous solutions every other day for a total of 10 doses. Challenge: single intradermal injection of a 0.1% aqueous solution at 14 days after the last induction application. Evaluation: 24 hours after challenge injection.
Remark:	Original date of report: March 13, 1958 from Pharmacology
Result:	According to the authors, the test substance was not a sensitizer in this study. Zero out of 10 animals responded to DEAE
Test substance:	2-diethylaminoethanol, purity not indicated
Reliability.	(2) valid with restrictions
Kerrabilicy.	Not according to the guidelines of today, but guitable for
	its time
27 _ TIN _ 2002	(75)
27-00N-2002	(75)
Type:	Guinea pig maximization test
Species:	quinea pig
Result:	not sensitizing
Method:	other: Magnusson, B. and Kligman, A.M.: J. Invest. Derm. 52, 268
Year:	1969
GLP:	no data
Test substance:	other TS
Method:	Females of the Hartley strain were used. Induction was made via intradermal injection of 10,000 ppm in olive oil and topical treatment with a 50,000 ppm solution in olive oil. Twenty one days after the initial intradermal injection, 0.1 ml aliquots of various non-irritating concentrations of DEAE were applied in saline for the challenge (0, 1,250, 2,500, 5,000 and 10,000 ppm). The observation time after the challenge application was at 48 hours.
Result:	Sensitization was not observed in any of the DEAE-challenged animals (0/10 of the controls responded, 0/10 of the animals challenged with 1,250 ppm responded, 0/10 of the animals challenged with 2,500 ppm responded, 0/10 of the animals challenged with 5,000 ppm responded and 0/10 of the animals challenged with 10,000 ppm responded).
Test substance:	2-aletnylaminoethanol, purity not indicated
Reliability:	(2) valid with restrictions
01 7777 0000	Essential details given.
01 - 000 - 2002	(79)

5.4 Repeated Dose Toxicity

Species: Strain: Route of administr Exposure period: Frequency of treat Post exposure per: Doses: Control Group: NOAEL: LOAEL:	cation: cment: iod:	<pre>rat Fischer 344 inhalation 14 weeks 6 hr/day, 5 days/week 4 weeks 0.053, 0.120 and 0.365 mg/l (53, 5) original values: 11, 25 and 76 ppr yes, concurrent no treatment = .053 mg/l = .12 mg/l</pre>	Sex: male/female 120 and 365 mg/m3; n)
Method: GLP: Test substance:	other: guideli was not yes other 5	guideline-like; mentions conformin ines for particular parameters, but t specified TS	ng to U.S. EPA :, the exact guideline
Method:	Animals Twenty (approx days/we These of (Hinz of termina) observa water of examinal monthly and qua during assays hematol examinal prior to cavity/ (sterminal prior to cavity/ (sterminal in the dose gr The examinal was con Toxicol Nominal daily. were me Samples within	s were approx. 10 weeks old at the rats/dose/sex were exposed to 0, 1 k. 0, 53, 120 or 365 mg/m3) of DEA eek for 14 weeks using a whole body doses were chosen based on results et al., 1992; Exxon, 1990). Half ated at the end of week 14. In-cha ations were performed daily. Indivi- ations and body weights were detern consumption was determined monthly ations were performed prior to expo- y during the study. Urine for urin- antitative) was collected during the the last week in the recovery exp- , performed at both week 14 and af logy, serum chemistry, cholinester- ation, organ weight, and histopathe ogic exams were preformed monthly of Screen" during the exposure periods were given a four week post-expo- to sacrifice. Full histological es (turbinates, adrenal glands, bone at m), brain, epididymes, eyes, hear lung, cervical lymph node, gastro , parathyroid, pituitary, spinal co d, trachea, urinary bladder and ut high dose group and control, but at coups only the nasal cavity/turbina amination of the nasal cavity/turbina amination di the nasal cavity/turbina amination di turbina to the method of 1., 1, 309-312, 1981).	<pre>start of treatment. 11, 25 or 76 ppm E for 6h/day, 5 7 exposure method. from a 2 week study of the animals were amber group idual in-life nined weekly. Food and . Ophthalmological osure initiation and alysis (qualitative he 13th week and eriment. Terminal ter recovery included ase, gross necropsy ology evaluations. using a modified d. The remaining sure recovery period xams [nasal and bone marrow t, kidneys, larynx, cnemius muscle, ord, spleen, thymus, erus] were conducted in the low and middle ates were evaluated. inates (4 sections) Young (Fund. Appl.</pre>

	STATISTICAL METHODS: Bartlett's test, ANOVA, Dunnett's, Kruskall-Wallis test and several other tests if applicable.
Remark:	Temperature and relative humidity were monitored regularly and according to the report were within the ranges specified in U.S. EPA guidelines. The average analytical concentrations were 10.5, 25.5 and 76 ppm. These concentrations are comparable to ca. 13, 29 and 88 mg/kg bw per day doses assuming 100% lung deposition and absorption
Result:	No animals died as a result of exposure to DEAE. During exposure, dose-dependent transient signs of mild to moderate respiratory irritation (sneeze-like sounds or rales) were noted. They usually cleared within one hour after exposure. In the high dose group, some animals continued to exhibit these signs overnight. Nasal discharge was observed at the beginning of the study, but this subsided as the study progressed. Corneal opacities were observed in control and DEAE-treated animals. DEAE exposure appeared accelerate the appearance of this lesion in the middle and high concentration groups. According to the authors, aging F344 rats are genetically predisposed toward developing these corneal lesions. Prolonged exposure to an alkaline compound such as DEAE might have accelerated an underlying predisposition toward corneal dystrophy. Since the opacity was thought to be due to a calcium precipitate, this problem may have also been exacerbated by the high vitamin D diet which would increase calcium absorption.
	No outstanding effects on blood chemistry, urinalysis or neurobehavioural parameters were observed.
	Through the first 7 weeks of exposure the high dose group had a slight, but statistically significant decrease in body weight gain as compared to controls. Subsequently the rate of growth paralleled the other groups, but the initial decrement was never regained. Mean body weights of the high dose groups never decreased more than about 7% from the controls. At week 14 there was a slight, but statistically significant increase in the absolute male liver and kidney weights in the high dose group (8.0 and 7.1%, respectively), as well as in their relative weights (increased by 5.9 and 7.1%, respectively), but histologic changes were not associated with these findings. Histomorphologic changes in nervous tissues were not observed. In females, only the high dose relative kidney weight was slightly increased (7.9%, p < 0.05).
	The low dose group was free of exposure-related histologic changes in the nasal cavities and turbinates, but changes were noted in the middle and high dose groups sacrificed in the 14th week of exposure. These consisted of an increased incidence [45% (50% M, 40%F) at the middle concentration & 95% (90% M, 100% F) at the high concentration] and severity of focal hyperplasias alone or in association with squamous metaplasia of the respiratory epithelium, and multi-focal mixed infiltrations of inflammatory cells in the nasal mucosa. Changes were most evident in the anterior sections of the nasoturbinates and on the lateral wall of the nasal cavity. In the high dose group

	hypertrophic goblet cells were seen in the nasal septum, along with a low incidence of focal necrosis and exudate in the lumen of the nasal cavity. The findings in the middle and high dose groups after the 4 week recovery period were similar to those seen in the 14 week rats, however, the incidence of focal hyperplasia with squamous metaplasia was decreased. The incidence of focal hyperplasia alone, infiltrations of inflammatory cells and goblet cell hypertrophy were comparable to what was noted at 14 weeks. There were no exposure-related findings in the other areas of the respiratory system to indicate any irritating effect on the lower respiratory tract.
	This study indicated that DEAE lacked systemic toxic properties, and the point of contact (eyes and upper respiratory tract) was the site of action. Since no systemic toxicological effects were observed, the NO[A]EC for systemic toxicity was the highest dose tested, i.e. 0.365 mg/l (76 ppm, or 365 mg/m3). The NO[A]EC for local toxicity, based on the lack of observed effects in the nasal cavity/turbinates, was 0.053 mg/l (11 ppm, rounded off to 10 ppm, or 53 mg/m3). The noises or rales at this concentration were considered an adaptive effect, but not an adverse effect, since no histological changes were observed at this concentration. However, since an effect (rales) was seen at the lowest concentration, a NOEC was not reached.
	NO[A]EC, rat (inhalation) 14 weeks, systemic toxicity: 0.365 mg/l (76 ppm or 365 mg/m3)
	NO[A]EC, rat (inhalation) 14 weeks, local toxicity: 0.053 mg/l (10 ppm or 53 mg/m3)
Test substance: Reliability:	<pre>LO[A]EC, rat (inhalation) 14 weeks, local toxicity: 0.12 mg/l (25 ppm or 120 mg/m3) 2-diethylaminoethanol, purity: 99% pure, Pennwalt Corp. (1) valid without restriction Well documented, GLP certified, guideline-like study; mentions conforming to U.S. EPA guidelines for particular parameters, but, the exact guideline was not specified. Chosen as the key study for the ICCA Robust Summaries since more details were available.</pre>
Flag: 07-JAN-2003	Critical study for SIDS endpoint (80) (81) (82) (83)
Species: Strain: Route of administ Exposure period: Frequency of trea	<pre>rat Sex: male/female Fischer 344 ration: inhalation 2 weeks tment: 6 hr/day, 5 days of exposition followed by a 2 day</pre>
Post exposure per Doses:	<pre>pause, then 4 more days of exposure no 0.048, 0.272 and 1.463 mg/l (48, 272 and 1463 mg/m3, respectively; original data: 10, 56 and 301 ppm,</pre>
Control Group: NOAEL: LOAEL:	respectively) yes, concurrent no treatment = .048 mg/l = .269 mg/l

Method:	other					
GLP:	no data					
Test substance:	other TS					
Method:	Ten animals/dose/sex were used and were exposed a total of 9 times over a 2 week period. Histopathological examinations that included the gonads were performed in the control and 56 ppm group. In the 10 ppm group only the nasal turbinates were examined. Due to the high mortality, histological examinations were not performed in the 301 ppm group.					
Remark:	This experiment was conducted to determine the doses to be used for a subchronic study.					
Result:	Ten and 56 ppm exposures did not cause any major treatment-related changes. Generally, male rats in the 56 ppm group did not gain as much weight as the controls, however, this difference was not statistically significant. There was no mortality in either group. Urinalysis, hematology and serum chemistry failed to show any significant DEAE-induced differences. At necropsy, no treatment-related macroscopic changes were noted, and absolute and relative organ weight values were unremarkable. The principle histologic lesions found occurred only in the anterior sections of the nasal turbinate mucosa. Inflammation of the nasal turbinate and lateral wall mucosa were noted in half of the animals of the 56 ppm group. The epithelium in the infiltrated areas appeared to be flattened, with early squamous cell metaplasia evident in one male of the 56 ppm group. The histopathologic changes were dose-related and most evident in the 56 ppm group. The 10 ppm was basically free of abnormalities. There were no signs of systemic toxicity.					
	301 ppm exposure caused overt signs of ocular, nasal and respiratory distress during and immediately after exposure. Ocular discharge, opacities and ulcerations, minor skin sores, nasal discharge, rales, labored breathing and gasping, decreased activity and responsiveness, impaired coordination and reflexes, hypothermia, and increasing emaciation were noted as the study progressed. Group mean values for food and water consumption were significantly (p < 0.01) lower than controls. Animals lost weight throughout the study. Deaths occurred in 9/10 males and 5/10 females, with deaths occurring sooner in males than in females. These deaths made statistical analyses difficult. None-the-less, the urinalysis, hematology and serum chemistry of survivors was not comparable to controls (data not shown), and there were no consistent trends. Surviving animals exhibited a number of gross changes: under sized spleens, thymuses and gonads, nasal discharges, enlarged adrenals and intestinal gas. Although organ weights were lower than controls, due to the depressed body wt. their relative organ weights appeared to be comparable to controls. Autolytic changes precluded meaningful necropsy evaluations in animals that died as a result of exposure to DEAE.					

SUMMARY The NOAEC was the 10 ppm dose (48 mg/m3 or 0.048 mg/l) based on the lack of local toxicity to the upper respiratiory tract, and the LOAEC was the 56 ppm dose (269 mg/m3 or 0.269 mg/l) based on local toxicity. No systemic toxicity was observed at these two doses. Test substance: 2-diethylaminoethanol, purity: 99% pure, Pennwalt Corp. Reliability: (2) valid with restrictions Essential details given for a dose setting experiment. 03-JUL-2002 (81) Species: Sex: male rat Strain: Sprague-Dawley Route of administration: inhalation Exposure period: 1, 3 and 6 months Frequency of treatment: 6 hr/d, 5 d/wk Post exposure period: no data Doses: 0.97 mg/l (970 mg/m3, i.e. 200 ppm) Control Group: yes, concurrent no treatment Method: other: no data GLP: no Test substance: other TS Method: 50 males were exposed to DEAE vapors using a whole body method. 32 rats were exposed to the "air flow" in a similar chamber to serve as a control. Animals were exposed to the TS by metering DEAE into a vaporizing unit maintained slightly above room temperature. Vapors were picked up by the air stream passing through the vaporizer and carried into the top of the vapor chamber. Chamber concentrations of DEAE were measured daily by 30-minute sampling into glacial acetic acid and titration with perchloric acid. Animals were sacrificed and hemoglobin, hematocrit, RBCs, WBCs, differential cell counts, serum protein, serum glutamic oxaloacetic transaminase, ratios of liver and kidney wt to body wt and histopathologic observations were made. The sacrifice schedule was as follows: at 1 month 8 animals/group, at 3 months 12 animals/group, at 6 months 11 control animals and 23 DEAE-exposed animals. During the 1st month 7/50 animals lost weight (reduced by Result: 15% compared to controls) and died from bronchial pneumonia (based on histologic examination). Other parameters were not markedly different from controls. In the 2nd month one control rat died. By the end of 3 months the weight of both groups were nearly comparable. After 6 months there was no significant difference between exposed animals and the controls in regard to body weight and hematology, clinical chemistry and histopathology. Test substance: 2-diethylaminoethanol, purity not indicated Reliability: (4) not assignable Essential details not given (clinical symptoms, information on post exposure observation period, details of airway histological examinations, gonads were not examined).

02-JUL-2002

86

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Species: Strain: Route of administration: Exposure period: Frequency of treatment: Post exposure period: Doses: Control Group:		<pre>rat Sprague-Dawley inhalation 5 days 6 hr/day not data 0.97 mg/l (970 mg/m3 or 200 ppm) yes, concurrent no treatment</pre>	Sex: male/female
Method: GLP:	other: no	no data	
Test substance:	other 7	rs	
Method:	8 males vapors the TS slight: the ain into th of DEAD glacia Animal WBCs, o glutami kidney made. Mild i	s and 8 females (ca. 200-250 g) we using a whole body method. Anima by metering DEAE into a vaporizin by above room temperature. Vapors to stream passing through the vapor he top of the vapor chamber. Cham E were measured daily by 30-minute l acetic acid and titration with p s were sacrificed and hemoglobin, differential cell counts, serum pr c oxaloacetic transaminase, ratio wt to body wt and histopathologic	re exposed to DEAE ls were exposed to g unit maintained were picked up by izer and carried ber concentrations sampling into perchloric acid. hematocrit, RBCs, otein, serum s of liver and observations were
Result: Test substance: Reliability:	Mild i: day; si exposur develop the lux liver v 2-dietl (4) no Essent informa	Fritation to the eyes was observed light nasal irritation occurred at re. Both symptoms did not progres oment was similar to controls. The ng, brain, heart, spleen, adrenal were comparable to controls. hylaminoethanol, purity not indicate of assignable ial details not given (number of contion on post exposure observation	the end of 3rd the end of 3rd s. Body weight he histopathology of gland, kidney and ted control animals, h period, details of
02 - TITL - 2002	airway	histological examinations).	(60)
02 001 2002			(00)
Species: Strain: Route of administr Exposure period: Frequency of treat Post exposure per: Doses: Control Group:	ration: tment: iod:	<pre>rat Sprague-Dawley inhalation 5 days 6 hr/day yes, no details 2.43 mg/l (2430 mg/m3 or 500 ppm) yes, concurrent no treatment</pre>	Sex: male
Method: GLP: Test substance:	other: no	no data	
Method:	20 male whole 1 meterin above 1 stream top of were me	es (ca. 200 g) were exposed to DEA body method. Animals were exposed ng DEAE into a vaporizing unit mai coom temperature. Vapors were pick passing through the vaporizer and the vapor chamber. Chamber conce easured daily by 30-minute samplin	E vapors using a to the TS by ntained slightly ed up by the air carried into the entrations of DEAE of into glacial

OECD SIDS	2-DIETHYLAMINOETHANOL
5. TOXICITY	DATE: 07-MARS-2003
	SUBSTANCE ID: 100-37-8
Result:	acetic acid and titration with perchloric acid. Animals were sacrificed and hemoglobin, hematocrit, RBCs, WBCs, differential cell counts, serum protein, serum glutamic oxaloacetic transaminase, ratios of liver and kidney wt to body wt and histopathologic observations were made. During the 1st exposure day, irritation of the eyes and nose were seen; all animals exhibited tremors of the head and forlegs. These symptoms continued throughout the five-day exposure period. After the 3rd exposition corneal opacity was observed. Four animals died (one on exposure day 4 and 3 died 3 days after exposure had ceased). All animals had lost 40 to 80 g of body weight when compared to controls.
Test substance: Reliability:	<pre>Histopathological findings: acute purulent bronchiolitis and bronchopneumonia; other tissues were similar to controls. 2-diethylaminoethanol, purity not indicated (4) not assignable Essential details not given (number of control animals, length of post exposure observation period, details of airway histological examinations).</pre>
02-JUL-2002	(60)
Species: Strain: Route of administr Exposure period: Frequency of treat Post exposure per: Doses: Control Group:	rat Sex: no data no data cation: inhalation 5 months tment: 4 hr/day iod: no data 128 ppm (ca. 622 mg/m3 or 0.622 mg/l) no data specified
Method:	other: no data
GLP: Test substance:	no other TS
Result:	Treated rats exhibited symptoms of excitation of the CNS, clonicotonic convulsions, irritation of the respiratory tract, and depressed body weights.
Test substance:	2-diethylaminoethanol
Reliability:	(4) not assignable
02-JUL-2002	Secondary literature. Essential details are lacking. (84)
Species: Strain: Route of administr Exposure period: Frequency of treat Post exposure period Doses: Control Group:	<pre>rat Sex: male/female other: albino rats from Charles River Breeding Lab. ration: oral feed 6 months, 12 months and 2 years tment: continuously in the food iod: not indicated 200, 500 and 1000-10,000 ppm (the high dose group was gradually increased to 10,000 ppm); corresponding to ca. 11, 25, 50-400 mg/kg bw/day, respectively. These were the doses of the free base, but the TS was given as the HCl salt. yes, concurrent no treatment</pre>
Method:	other
GLP:	no data
Test substance:	other TS

OECD SIDS

5. TOXICITY

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Method: For the DEAE-treated groups, 35 rats/sex were used, and in the control there were 60 rats/sex. Observations were made 6 days/week for signs of toxicity. Body weights and food consumption were recorded weekly for the 1st 26 wks, and biweekly for the next 26 wks. In the 2nd year, body weights and food consumption were measured every 4 weeks. Hematocrits, hemoglobin determinations and total and differential leucocyte counts were made from 5 rats of each sex and group on day 30, 45, 90, 180, 360, 540 and 720. Complete blood counts were performed on 5 additional females from the control and high dose group on day 180, and total white cell counts were performed on 5 additional rats of each sex from all groups on day 360. Urine analyses (albumin, acetone, bilirubin, color, occult blood, sugar, pH, appearance and microscopic sediment examination) were made on day 30, 45, 60, 90, 180, 360, 540 and 720 from pooled sample groups of the same rats used for the blood tests. At 6 and 12 months, 5animals/sex/group were sacrificed and complete necropsies were performed. Representative tissues from each animal were fixed in formalin. These tissues were: 3 sections of the brain, two sections each of the stomach and small intestine, one section each of the pituitary, thyroid, heart, lung, liver, spleen, kidney, adrenal, pancreas, large intestine, urinary bladder, gonads, bone marrow and any unusual lesion. Animals which died or which were moribund were taken and when post-mortem autolysis was not advanced, target tissues as well as tumors were examined histologically. All of these tissues from the control group and the high dose group were examined microscopically from the 6 and 12 month sample groups. At the end of 2 years, all surviving animals were necropsied. Ten animals/sex from controls and the high dose group were examined histologically. The gonads of all groups of the 2 year sampling were eventually examined histologically. Organ weights and organ/terminal body weights were recorded from all scheduled sacrificed animals. The doses were based on the free base, but the Remark: substance was given as a HCl salt. The dietary levels of the free base were: 0, 200, 500 and 1000 ppm. After week 47 the 1000 ppm group was gradually raised to 10,000 ppm as follows: Week 48 - 56, 1500 ppm; Week 57 - 64, 2500 ppm; Week 65 - 72, 3500 ppm; Week 73 - 80, 5000 ppm; Week 81 - 84, 7500 ppm; Week 85 - 104, 10000 ppm. None of the treated animals displayed gross signs of Result: substance-induced toxicity. Adverse signs occurred mainly in the last 6 months in all groups and were associated with aging (this included general poorer health and an increase in mortality). According to the report, since the incidence of mortality of the DEAE-treated groups compared favorably with the controls, DEAE did not produce any earlier or greater numbers of deaths at any dose level. Instances of anorexia were also reported in this study, but

the data on individual animals was not given.

According to the report, the animals were said to have generally had a vigorous appetite and the the average weekly body weight and feed consumption values for males and females were generally comparable to controls at the corresponding interval. However, in the last 12 weeks of the study the high dose male body weight was on average 8.6% (maximum 11%) lower than the controls. No significant hematological changes were seen. In both sexes of the high dose group, the hematocrit and hemoglobin values were slightly decreased at the 720 day time point. At 6 months no testicular atrophy was observed. At 12 months one case of slight testicular atrophy was grossly observed in a 200 ppm dosed male. In the 24 month samples testicular atrophy was observed in: 0/34 control animals, 3/18 (17%) animals of the 200 ppm dose group, 2/17 (12%) animals of the 500 ppm dose group and 4/15 (27%) animals of the 1000-10000 ppm dose group. In the high dose group, 3 cases were grossly described as slight atrophy in one testis, and this was confirmed histologically; the fourth case was not grossly observed, but the histologic description indicated that 2/3 of the testis had a "moderately severe" generalized degree of atrophy present with only Sertoli cells which remained within most of the seminiferous tubules. According to the sponsor, the lack of testicular atrophy in the controls was surprising given the historical control range of this lesion (data not available). It should be noted that this is not an unusual lesion in ageing rats (Glaister, 1986). One high dose female had gonadal atrophy, but the report considered this fortuitous. It should be emphasized that in the six month samples, the more appropriate time point for examining reproductive toxicity, no testicular effects were observed. The atrophy often occurred bilaterally. Careful histologic examination of the cerebellums of the high dose animals revealed no anomalies. No other signs of toxicity were observed. Roundworms were found in the colon of 2 control and 3 high dose animals. Given the lack of a DEAE-related effect seen in the 6 month and 12 month samples, the NOAEL was the highest dose tested for these time points, i.e. 1000-1500 ppm/day, or ca. 50-70 mg/kg/day. The gonadal degeneration seen in 2 year old DEAE-treated animals was confounding since it was not dose-dependent, however, due to the presence of this lesion,

a NOAEL could not be assigned for the 2 year time point. See section 5.7 for a description of the results in regard to carcinogenicity.
 Test substance: According to the report, 1.000 ml contains 0.577-0.585 g of Diethylaminoethanol or 0.758-0.766 g Diethylaminoethanol hydrochloride; Pennsalt Chemicals Corp.
 Reliability: (2) valid with restrictions
 Some essential details are given. A limitation is the lack of a detail on the anorexia observed (i.e. was it associated with DEAE treatment?). A maximum tolerated dose was not

achieved, thus, there is no clear basis for the doses

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

chosen. For today's requirements, the number of animals used is a limitation (for the 2 year carcinogenicity endpoint); however, in the 1960s there were no international guidelines. No basis is given for the dose selection. 26-JUL-2002 (85) (86) (87) Species: rat Sex: male Strain: Spraque-Dawley Route of administration: drinking water Exposure period: 4 weeks Frequency of treatment: continuous Post exposure period: no data Doses: ca. 500 mg/kg/dControl Group: no data specified Method: other: no data GLP: no Test substance: other TS Remark: The dose level was described as 80-120 mg/rat/day, according to the report. Male rats were maintained on normal diets and received DEAE neutralized with HCl in the drinking water at a level of 4 mg/ml. The weight of the rats was given as 150-175 g at the start. Taking averages, the dose was estimated to be ca. 80 mg/0.162 kg/day, i.e. ca. 500 mg/kg. The only effect noted after 4 wk of DEAE treatment was a slight elevation of the kidney wt to body wt ratio (p < 0.05). Liver lipids, serum lipids, cholesterol levels and liver wt to body wt ratios were not effected by DEAE. No significant histological changes were noted in the lung, liver, kidney or spleen. No further information was available. Test substance: 2-diethylaminoethanol neutralized with HCl, purity not indicated Reliability: (4) not assignable Essential details not given. 19-OCT-2001 (59) Species: rat Sex: male Spraque-Dawley Strain: Route of administration: drinking water Exposure period: 1, 2 and 6 months Frequency of treatment: continuous Post exposure period: no data Doses: ca. 150 and 300 mg/kg bw/day (2000 and 4000 mg/l of neutralized DEAE) Control Group: yes, concurrent no treatment Method: other: no data GT.P . no Test substance: other TS Method: Weight at study initiation: 200-250 g Exposure period: up to 6 months Doses: 0, 50 and 100 mg/rat/day (the article states that the rats consumed 25-30 ml of a water solution containing 0, 200 $\,$ or 400 mg per 100 ml). That corresponds to approx. 0, 150 and 300 mg/kg bw/d. 15 males/dose were used. 5

OECD SIDS
5. TOXICITY

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

	animals/group were sacrificed after 1, 2 and 6 months.			
	Parameters investigated: Body weight, liver and kidney			
	weights, hemoglobin, clotting time, liver lipids, serum			
	total lipids cholesterol(esters) phospholipids and			
	bistonathology (liver kidney heart spleen) Statistical			
	methods were not indicated			
Domomire	According to the 1006 TIV Degumentation supplement rate			
Remark:	According to the 1996 TLV Documentation supplement, rats			
	were "fed" diethylaminoethanol at 50 and 100 mg/kg, however,			
	this is a misreading of the original report.			
Result:	Body weights of treated animals were comparable to control			
	values up to day 30. Thereafter, decreased body weights were			
	observed in treated animals. In this regard at 60 days, the			
	low dose group had lost about 30 g and the high dose had			
	lost about 50 g when compared to the control group (reduced			
	ca. 8 and 14% from the control value). After 6 months rats			
	receiving the low dose recovered from their decreased body			
	weight but high dose rats remained below control body			
	weight, but high dose lats lemained below control body			
	weights (by ca. 11%). No effects were seen in hematology			
	and clinical chemistry parameters at the 1, 2 or 6 month			
	time points. Nor was an effect on the liver to body weight			
	ratio seen. The only difference noted between DEAE-treated			
	animals and controls seen at autopsy was that the ratio of			
	kidney weight to body weight was slightly elevated in both			
	treated groups at the 1, 2 and 6 months time points. No			
	changes in the kidney were noted histologically in treated			
	animals at any of the scheduled sacrifices. Nor were			
	histologic changes noted in the other organs investigated			
	(liver, heart, spleen).			
Test substance:	2-diethylaminoethanol neutralized with HCl, purity not			
	indicated			
Reliability:	indicated (4) not assignable			
Reliability:	indicated (4) not assignable The study is limited by the low number of animals/group and			
Reliability:	indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically			
Reliability:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically. (60) (88)</pre>			
Reliability:	indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically. (60) (88)			
Reliability:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically. (60) (88) dog</pre>			
Reliability: 19-OCT-2001 Species:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of trea Post exposure per Doses:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of trea Post exposure per Doses:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method: GLP:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method: GLP: Test substance:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method: GLP: Test substance:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method: GLP: Test substance: Method:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method: GLP: Test substance: Method:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method: GLP: Test substance: Method:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method: GLP: Test substance: Method:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method: GLP: Test substance: Method:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method: GLP: Test substance: Method:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method: GLP: Test substance: Method:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method: GLP: Test substance: Method:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method: GLP: Test substance: Method: Result:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			
Reliability: 19-OCT-2001 Species: Strain: Route of administ Exposure period: Frequency of treat Post exposure per Doses: Control Group: Method: GLP: Test substance: Method: Result:	<pre>indicated (4) not assignable The study is limited by the low number of animals/group and only a few organs were examined histologically.</pre>			

occurred intermittently at first and eventually occurred continuously in a few animals. All dogs of the next two dose groups exhibited severe cases of weakness, tremors, convulsions and ataxia, with two animals in the 5000 ppm group dying (one on d 35 and one on d 41). All animals in the 10,000 ppm group died between days 18 and 39 of the study. After stopping treatment in the 5000 ppm group, the animals showed signs of improvement; however, the ataxia and tremors were still occurring when dosing (now with 2000 ppm) was resumed on day 134. This group displayed an increase in ill effects after dosing resumed, but some improvement occurred with time.

Body wt and food consumption appeared normal in the 2 lowest dose groups. No pronounced treatment-related findings were observed in the blood or urine in any group.

With the exception of the findings in moribund animals of the 2 highest dosed groups, clinical examinations showed no abnormalities with respect to pulse rate, reflexes or condition of the mucous membranes. ECG tracings showed no DEAE-related damage to the heart.

Gross necropsy of the animals that died in the 1st 180 days showed congestion and hemorrhages of the lungs, congestion of the kidneys, reddish mottled coloration of the spleen, hardness of the liver, and numerous enlarged and congested lymph nodes. Gross necropsy of the animals that survived to the termination revealed no gross pathological changes attributable to treatment. Terminal body and organ weights for control and DEAE treated animals showed no pronounced differences.

	Microscopic examination of the tissues of the 5000 (2000)
	ppm group showed atrophy of the thyroid gland (one male and
	s remarks) and gonads (three marks), which according to the
	report was interpreted as a non-specific secondary response to
	the metabolic or toxic insult of the TS. One female in this
	group had to a decrease in oogenesis. Cerebellar changes were
	also observed in this group and occurred in all males and one
	female. These changes consisted of irregular patchy
	degeneration and loss of small to moderate numbers of Purkinje
	cells, with occasional mild decreases in the cellularity of
	the granular layer. Foci of
	tissue calcification were present in the one female.
Test substance:	2-diethylaminoethanol neutralized with HCl; according to the
	report, 1.000 ml contains 0.577 g of Diethylaminoethanol or
	0.758 g Diethylaminoethanol hydrochloride; Pennsalt Chemicals
Reliability:	(4) not assignable
	The study was limited to summary information only Tables
	and individual animal data ways not evaluable for an
	and individual animal data were not available for an
0.000	assessment.
20-101-2002	(75) (87)

5.5 Genetic Toxicity 'in Vitro'

Type: System of testing: Concentration: Metabolic activation: Result:		Ames test Standard plate test and preincubation test with Salmonella typhimurium TA1535, TA100, TA1537, TA98 0, 20,100, 500, 2500 and 5000 µg/plate with and without negative			
Method: Year: GLP: Test substance:	OECD Gu 1983 no as pres	wuide-line 471 escribed by 1.1 - 1.4			
Method:	The S-9 Three p study w The fol with S- without 10 ug 4 9-amino A paral mix was spontan Evaluat followi - a dou - a dos	<pre>9 mix was prepared from Aroclor 1254 pretreated rats. plates per dose and per control were tested. The was performed under GLP-like conditions. ollowing positive controls were used (dissolved in DMSO): -9 mix, 10 ug 2-aminoanthracene, at S-9 mix, 5 ug MNNG for strains TA 100 and TA 1535, 4-nitro-o-phenylendiamine for TA 98, and 100 ug moacridine chloride monohydrate for TA 1537. cllel "negative" solvent control with and without S-9 as carried out for each tester strain to determine the neous mutation rate. tion Criteria-a positive result had to fulfill the ring: publing of the spontaneous mutation rate (control) ose-response relationship</pre>			
Remark: Result:	 N.B.: This study followed the OECD Guideline 471 of 1983 and not the current OECD guideline 471 of1997. That is why E. co and TA 102 were not included. Similar negative results were reported by Zeiger et al. (Environ. Mutagen. 9, Suppl.9, 1-110, 1987) and in another guideline-GLP study sponsored by the Synthetic Organic Chemicals Manufacturing Association (Life Science Research, Report No. 91/SHG001/0251, Diethylaminoethanol: Assessment of mutagenic potential in histidine auxotrophs of Salmonella typhimurium, 2 July 1991). No increase in the number of his+ revertants was seen in any 				
Test substance: Reliability:	of the test su range t observe with TA test) a expecte CAS No. (1) va Well con conditi Summari	substance was completely soluble in water in the dose substance was completely soluble in water in the dose s tested. A weak bacteriotoxic effect was occasionally rved using TA 1537 with S-9 mix (standard plate test) and TA 1535, TA 1537 and TA 98 without S-9 mix (preincubation) at 5000 ug/plate. All positive controls produced the cted effects. No. 100-37-8 (2-diethylaminoethanol), purity: > 99% valid without restriction conducted guideline study conducted under GLP-like itions. Chosen as the key study for the ICCA Robust aries.			

Critical study for SIDS endpoint Flag: 28-JUN-2002 (89) Type: Ames test System of testing: Standard plate test with Salmonella typhimurium TA1535, TA1537, TA1538, TA98 and TA100 Concentration: 50 to 5000 μ g/plate Metabolic activation: with and without Result: negative Method: OECD Guide-line 471 Year: 1983 GLP: ves Test substance: other TS Method: N.B.: This study followed the OECD Guideline 471 of 1983 and not the current OECD quideline 471 of1997. That is why E. coli and TA 102 were not included. Test substance: CAS No. 100-37-8 (2-diethylaminoethanol), purity: 99.8%, Atochem North America (1) valid without restriction Reliability: Well conducted guideline study conducted under GLP conditions. 28-JUN-2002 (90) Type: Ames test System of testing: Preincubation modification of the Salmonella/microsome test in the absence and presence of exogenous metabolic activation (S. typhimurium TA 98, TA 100, TA 1535, TA 1537) Concentration: 33, 100, 333, 1000, 2500 und 3333 µg/Platte Metabolic activation: with and without negative Result: Method: other: according to Haworth, S. et al.: Environ. Mutagen. 5, Suppl. 1, 3-142 1983 Year: GLP: no data other TS Test substance: Method: Preincubation modification of the Salmonella/microsome test in the absence of exogenous metabolic activation and in the presence of liver S-9 from Aroclor-induced male Sprague-Dawley rats and Syrian hamsters. S. typhimurium strains TA 98, TA 100, TA 1535 and TA 1537 were used. Test substance: CAS No. 100-37-8 (2-diethylaminoethanol), purity: "99+", Fluka (2) valid with restrictions Reliability: Essential details are given. 28-JUN-2002 (91) HGPRT assay Type: System of testing: Chinese hamster V79 cell mutation system Concentration: 1st experiment: 4.8 to 3000 $\mu\text{g/ml}\textsc{;}$ 2nd experiment: 5.6 to 3500 $\mu\text{g/ml}$ Metabolic activation: with and without negative Result:

Method:	OECD Guide-line 476
Year:	1984
GLP:	Ves
mant maket and a	
Test substance:	other is
Method:	The test conformed to the OECD Guideline 476 of 1984. The study was also carried out in accordance to EPA (TSCA) guidelines (1985, revised 1987).
	Cell type: V79 clone 6 Metabolic activation system: rat-liver derived S9-mix (Arochlor 1254-treated rats).
	Dosing: 1st expt.: 4.8, 24, 120, 600, 3000 ug/ml; 2nd expt.: 5.6, 28, 140, 700, 3500 ug/ml Number of replicates: 3 plates/dose group
	Negative control: distilled water (vehicle) Positive controls: -S9, ethylmethanesulphonate +S9, dimethylbenzanthracene
Result:	<pre>Selection agent: 6-thioguanine (6-TG) Treatment procedure: 7.5 x 10E5 cells/dose group were seeded in 25 cm2 flasks for 24 hours and then were treated for 3 hours. After treatment the cell sheet was washed and non-selective medium was added. A sample of cells was taken to measure survival after treatment. Cells were then passaged to maintain subconfluence for an expression time of 7 days. Then, cells were plated for the mutant selection (10E5 cells/plate/3-fold) and for the plating efficiency (200 cells/plate/3-fold); after 6 days the resultant colonies were scored. FOLLOW UP REPEAT STUDY: independent repeats were performed. CRITERIA FOR EVALUATING RESULTS: a dose dependent increase in the number of 6-TG resistant colonies compared to the solvent control. GENOTOXIC EFFECTS: With and without metabolic activation: negative Cultures exposed to diethyaminoethanol showed no increases in 6-TG resistant colony numbers and no significantly increased mutant frequencies compared to solvent controls. CYTOTOXIC CONCENTRATION: With and without metabolic activation: In both experiments in the highest dose level of 3000 - 3500 ug/ml, no effect on</pre>
Test substance: Reliability:	<pre>the surviving cells were observed in the plating efficiency. However, in both experiments at the end of the exposure time, changes in the cell morphology were observed. No precipitation was observed. DEAE caused a concentration-related increase in pH of the treatment medium. CAS No. 100-37-8 (2-diethylaminoethanol), purity: 99.8% (1) valid without restriction Well conducted guideline study conducted under GLP conditions. Chosen as the key study for the ICCA Robust Summaries</pre>
Flag: 26-JUL-2002	Critical study for SIDS endpoint (92)

Type: System of testing: Concentration: Metabolic activat: Result:	other: DNA Damage in E. Coli WP2, WP67 and CM871 35 to 3500 µg/ml on: with and without negative	
Method: Year: GLP: Test substance:	other: EPA OTS Sect. 798.5500 1985 yes other TS	
Test substance: Reliability:	CAS No. 100-37-8 (2-diethylaminoethanol), purity: 99.8%, Atochem North America (1) valid without restriction Well conducted guideline study conducted under GLP conditions.	
19-OCT-2001	(93)

5.6 Genetic Toxicity 'in Vivo'

Type: Species: Strain: Route of admin.: Doses: Result:	Micronucleus assay mouse ICR oral unspecified 20, 100 and 500 mg/kg bw negative	Sex: male/female
Method: Year: GLP: Test substance:	OECD Guide-line 474 "Genetic Toxi 1983 yes other TS	.cology: Micronucleus Test"
Method:	N.B.: The test conformed to the OE US EPA Guidelines of 1985.	CD Guideline 474 of 1983 and
	TEST ORGANISMS 4-5 week old mice, 18-28 g at stud 5/sex/dose/sampling time	ly initiation,
	ADMINISTRATION Vehicle: 0.9% saline Frequency of treatment: single ora Dosing volume: 10 ml/kg Sampling times: 24 hr (all groups) dose) and 72 hr (control + high do Controls: negative, 0.9% saline positive, 30 mg/kg chlorambucil N.B.: The doses were chosen based test using 2 animals/sex/group treated and 2500 mg/kg bw.	el dose (, 48 hr (control + high (on a preliminary toxicity (ated with 312.5, 625, 1250
	Three bone marrow smears per anima Clinical observations were made da measured on day 0 and at the termi	l were made. ily. Body weight was .nation.

	The frequency of micronucleated cells per at least 1000 polychromatic erythrocytes, as well as the frequency of micronucleated cells per at least 1000 mature erythrocytes was determined. The ratio of polychromatic to mature cells was also determined.
Result:	<pre>Statistical method: Mann-Whitney U-test. TOXIC RESPONSE/EFFECTS BY DOSE LEVEL Mortality and time to death: One male at 500 mg/kg was killed in extremis 42 hours after dosing. Clinical signs: no signs observed at 20 mg/kg; at 100 mg/kg hunched posture and piloerection (1 male); at 500 mg/kg 1 male was killed in extremis and showed rales, piloerection, hunched posture and a swollen abdomen, and 16 of the remaining animals showed rales, piloerection, hunched posture as well and/or irregular respiration. Body weight changes: body weight loss was noted in 12/30 mice at 500 mg/kg during the period before termination; other incidences of wt loss were noted throughout the study, but these were small and not dose-related.</pre>
	The PCE/NCE ratio in the preliminary study was 0.9, 0.8 and 0.5 in the 312.5, 625 and 1250 mg/kg dose groups, respectively. This indicated that the TS reached the bone marrow.
	EFFECT ON PCE/NCE RATIO PCE/NCE ratio (males and females combined): at 24 hours: 0.9 (all doses); 0.8 (saline control) at 48 hours: 0.9 (500 mg/kg); 0.9 (saline control) at 72 hours: 0.7 (500 mg/kg); 0.7 (saline control)
	<pre>GENOTOXIC EFFECTS Mean number of micronucleated PCE/1000 PCE (males and females combined) - 0.9% saline control: 1.2, 0.6 and 0.6 at 24, 48 and 72 hours, respectively; - 20 mg/kg: 0.8 at 24 hours; - 100 mg/kg: 0.7 at 24 hours; - 500 mg/kg: 0.6, 0.8 and 0.3 at 24, 48 and 72 hours, respectively.</pre>
	STATISTICAL RESULTS Frequencies of micronucleated polychromatic erythrocytes were not significantly different from controls at any dose or sampling time. Positive controls gave the expected response.
Test substance: Reliability:	CONCLUSIONS Under the conditions of this test, diethylaminoethanol did not exhibit a chromosome-damaging (clastogenic) effect, nor was there an indication of an impairment of mitotic chromosome distribution. CAS No. 100-37-8, diethylaminoethanol, purity: > 99% (1) valid without restriction Well conducted guideline study conducted under GLP conditions. Chosen as the key study for the ICCA Robust

Flag:	Critical	study	for	SIDS	endpoint
26-JUL-2002					

(94)

5.7 Carcinogenicity

Species:		rat Sex: male/female			
Strain:		other: albino rats from Charles River Breeding			
Pouto of administr	ation.	Laboratories			
Exposure period.	acion:	2 years			
Frequency of treat	ment:	continuously			
Post exposure per	Lod:	no			
Doses:		200, 500 and 1000 ppm (the high dose group was			
		gradually increased to 10,000 ppm); corresponding to			
		ca. 11, 25, 50-400 mg/kg bw/day, respectively			
Result:		negative			
Control Group:		yes			
Method:	other				
GLP:	no				
Test substance:	other 1	IS			
Method:	For the	e DEAE-treated groups, 35 rats/sex were used, and in			
	the co	ntrol there were 60 rats/sex. Observations were made			
	6 days,	/week for signs of toxicity. Body weights and food			
	consum	ption were recorded weekly for the 1st 26 wks, and			
	biweek!	ly for the next 26 wks. In the 2nd year, body weights			
	and for	od consumption were measured every 4 weeks.			
	Hematod	rits, hemoglobin determinations and total and			
	differe	ential leucocyte counts were made from 5 rats of each			
	Sex and	group on day 30, 45, 90, 180, 360, 540 and 720.			
	from th	te prood counts were performed on 5 additional remains			
	white of	cell counts were performed on 5 additional rats of each			
	sex fro	om all groups on day 360. Urine analyses (albumin,			
	acetone	e, bilirubin, color, occult blood, sugar, pH, appearance			
	and mid	croscopic sediment examination) were made on day 30, 45,			
	60, 90	, 180, 360, 540 and 720 from pooled sample groups of the			
	same ra	ats used for the blood tests. At 6 and 12 months, 5			
	animals	s/sex/group were sacrificed and complete necropsies were			
	in for	med. Representative tissues from each animal were fixed			
	section	main. These classes were, a sections of the blain, two			
	each of	f the pituitary, thyroid, heart, lung, liver, spleen,			
	kidney	, adrenal, pancreas, large intestine, urinary bladder,			
	gonads	, bone marrow and any unusual lesion. Animals which died			
	or which	ch were moribund were taken and when post-mortem			
	autoly	sis was not advanced, target tissues as well as tumors			
	were er	xamined histologically. All of these tissues from the			
	contro.	I group and the high dose group were examined			
	uncroso	Suprearry from the b and 12 month sample groups. At the			
	animal	2 years, arr surviving annuals were necropsied. Ten			
	examina	ed histologically. The gonads of all groups of the 2			
	year sa	ampling were eventually examined histologically. Organ			
	weight	s and organ/terminal body weights were recorded from all			
	schedu	led sacrificed animals.			

OECD SIDS

5. TOXICITY

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003

SUBSTANCE ID: 100-37-8

Remark: Result:	According to personal communication with Charles River (Patricia A. Mirley, 24 June 2002), the rats used were likely to be Sprague-Dawleys. The concentrations were based on the free base, but the substance was given as a HCl salt. The dietary levels of the free base were: 0, 200, 500 and 1000 ppm After week 47, the 1000 ppm group was periodically raised as to 10,000 ppm as follows: Week 48 - 56, 1500 ppm; Week 57 - 64, 2500 ppm; Week 65 - 72, 3500 ppm; Week 65 - 72, 3500 ppm; Week 81 - 84, 7500 ppm; Week 85 - 104, 10000 ppm. There was no increase in the number of tumors in the treated animals as compared to the controls. For information on the interim sacrifices, see section 5.4 (Repeated Dose Toxicity).
	CONTROLS AND HIGH DOSE GROUP LESIONS (10 ANIMALS/SEX/GROUP): Almost all rats of each sex of the control and high dose group which were examined completely showed a mild degree of chronic bronchitis and/or pneumonitis, occasionally of the aspiration variety, plus a mild chronic interstitial nephritis. Minimal squamous metaplasia of thyroid follicular epithelium was found in one male animal of the control and high dose group. Perivascular iron deposition in macrophages in the pancreas was found to a mild degree in four control males and one high dose group male. Roundworms were encountered within the colon of two control and high dose group rats. Patchy adrenal cortical degeneration was present in five control and eight high dose group animals. Cystitis was seen in one control and two high dose group males but was absent from the control group, and ovarian atrophy was similarly found in one experimental female only. Miscellaneous findings included one animal each with fatty metamorphosis of the liver, a chronic gastric ulcer, chronic sialadenitis, and spicarditis, all within the high dose group. Careful examination of the sections of the cerebellum in the high dose group animals showed no degenerative changes, cell loss, or other abnormality.
Test substance:	Tumor were numerous in the forty animals which were examined completely. These included pituitary ademonas in 9 animals each in the control and high dose group groups; mammary gland fibromas, adenomes or fibroadenomes in 8 control and 4 high dose group females; and miscellaneous tumors which included one ganglioneuroma, one pheochromocytoma and two renal embryomas in the control group; adrenal cortical adenomas in one control and three high dose group females; and one pancreatic duct adenoma, one hepatoma and three granulosa cell tumors in the high dose group. According to the report, 1.000 ml contains 0.577-0.585 g of Diethylaminoethanol or 0.758-0.766 g Diethylaminoethanol hydrochloride; Pennsalt Chemicals Corp.

OECD SIDS	2-DIETHYLAMINOETHANOL
5. TOXICITY	DATE: 07-MARS-2003
	SUBSTANCE ID: 100-37-8
Reliability:	(2) valid with restrictions
	Some essential details are given. A limitation is the lack of a detail on the anorexia observed (i.e. was it associated with DEAE treatment?). A maximum tolerated dose was not achieved, thus, there is no clear basis for the doses chosen. For today's requirements, the number of animals used is a limitation (for the 2 year carcinogenicity endpoint); however, in the 1960s there were no international guidelines. No basis is given for the dose selection.
03-JUL-2002	(95) (87)

5.8.1 Toxicity to Fertility

5.8.2 Developme ntal Toxicity/Teratogenicity

Species: Strain: Route of administr Exposure period: Frequency of treat Duration of test: Doses: Control Group: NOAEL Maternal Tox NOAEL Teratogenic:	ration: tment: tity:	<pre>rat Sprague-Dawley inhalation days 6 to 15 of gestation 6 h/d until day 21 of gestation 0.160, 0.320 and 0.486 mg/l (160, original values: 33, 66, 100 ppm) yes, concurrent no treatment = .16 mg/l = .486 mg/l</pre>	Sex: 320,	<pre>female and 486 mg/m3;</pre>
Method:	other			
GLP: Test substance:	yes other TS			
Method:	The concentrations used were selected based on results from range finding study using 8 pregnant females/group exposed 10, 50, 100, 150, 200 ppm of the TS on GD 6-15. In the main study, the developmental toxicity of the test substance was evaluated in groups of 25 pregnant rats. The rats were expose to vapours of the test substance at concentrations of 0 (control), 33, 66, and 100 ppm (ca. 0, 0.158, 0.316, and 0 mg/l, respectively). All dams were sacrificed on day 21 of gestation.		results from a oup exposed to In the main ubstance was is were exposed ons of 0 316, and 0.486 h day 21 of	
	Age at st Weight at Route of Air chang	art: 83-91 days day 0 of gestation: 290-297 g administration: inhalation (whole ges: 12/hour	body)	- tion was
	confirmed	by the presence of a vaginal plug	g or s	sperm.
	PARAMETER Mortality Body weig gestatic Examinati implantat resorptic	AS ASSESSED DURING STUDY: //clinical observations: daily ght/food consumption: day 0,6,9,12 on .on of uterine content: uterine we cion sites, number of corpora lutes ons	,15, 1 ight, a, ear	l8 and 21 of number of rly and late

Examination of fetuses: weight, number of live and dead fetuses, sex and eternal viscera (1/2 of fetuses) and skeletal abnormalities (1/2 of fetuses)

ORGANS EXAMINED AT NECROPSY: histopathology of abnormal tissues was performed.

Analytical Methods: analytical concentration were determined by an online GC. Nominal concentration were calculated daily from the amount of test substance used and the airflow through the exposure chamber. Analytical concentrations were measured hourly. Samples were drawn from 4 points around the horizontal center plane of each of the exposure chambers to determine homogeneity of the TS distribution.

STATISTICAL METHODS: Bartlett's test, Dunnnett's test, linear regression, Kruskal-Wallis test, Fisher's test, chi-square test, Armitage test

Remark: The average analytical concentrations were 33, 66, 100 ppm These concentrations are comparable to ca. 38, 76 and 116 mg/kg bw per day doses assuming 100% lung deposition and absorption.

Result: Actual dose levels: 0, 33, 66, 100 ppm (analytical); 0, 37, 72, 112 ppm (nominal). The TS remained at least 99.88% pure during the study.

No deaths occurred as a result of exposure to DEAE. Maternal toxicity was observed in the high dose group and included reduced body weights (up to 6% on GD 15), and reduced body weight gain (up to 52%) during the entire exposure period (GD 6-15). Dry rales were observed in up to one third of the animals at the high concentration over GD 11 to 21. Decreased body weight gain was also observed in the 66 ppm group during GD 12 to 15. Statistically significant decreases in mean maternal food consumption were observed during the exposure period in the mid and high dose group and during the post-exposure period in the high dose group.

No treatment-related effects were seen on gestational parameters, including pre- and post-implantation loss or sex ratio. Mean fetal body weights in the treated groups were similar to controls. A statistically significant decrease in mean resorptions was observed in the high dose group. There was no increase in the incidence of malformations (external, visceral, or skeletal) individually or by category. The incidence of a single developmental variation (hypoplastic bones of the forepaw) was significantly (statistically) decreased in the high dose group relative to controls. This was only significant when analyzed on a per fetus basis, and was not significant when analyzed on a per litter basis. [N.B.: A decreased incidence of a developmental variation was not considered an adverse effect.] A statistically significant (p < 0.05) increasing dose-trend in the incidence of advanced ossification of the hind paw was reported, but it was not significant when analyzed on a per fetus or per litter basis. In this regard, the number of fetuses effected were 18/185, 18/183, 23/183 & 33/195, and the litters effected were 9/24, 9/25, 14/24 & 13/25 in the 0, 33, 66 and 100 ppm concentration groups, respectively. This increased incidence of advanced

OECD SIDS			2-DIETHYLAMINOETHANOL		
5. TOXICITY			DATE: 07-MARS-2003		
			SUBSTANCE ID: 100-37-8		
	ossifica	tion was higher	than expected in all groups, including		
	the cont	rol, compared to	the historical control range (0 -		
	2.3%) fr	om this laborat	ory. Thus, this finding was not		
	consider	ed treatment rel	ated or biologically relevant.		
	The no o	bserved adverse	effect concentration (NO[A]EC) for		
	maternal the NO[A	. toxicity was 0]EC for develop	.160 mg/l (160 mg/m3, i.e. 33 ppm) and nental toxicity was the highest dose		
	tested,	i.e. 0.486 mg/l	(486 mg/mg3 or 100 ppm).		
Test substance:	2-diethy America	laminoethanol, p	ourity 99.88%, Elf Atochem North		
Reliability:	(1) val	id without restr	riction		
	Well doc	umented, guidel:	ne-like study (the major points of		
	OECD Gui	deline 414 are o	overed), conducted under GLP		
	conditio	ns. Chosen as t	he key study for the ICCA Robust		
	Summarie	s since it was t	he best study available.		
Flag:	Critical	study for SIDS	endpoint (OC) (OZ)		
10 - JAN - 2003			(96) (97)		
Species:		rat	Sex: female		
Strain:		Sprague-Dawley			
Route of administ	ration:	inhalation			
Exposure period:		days 6 to 15 of	gestation		
Frequency of trea	tment:	6 h/d			
Duration of test:		until day 21 of a_2 0 05 0 24	$0.48 0.73 0.97 mmmmmmmmmmmmmmmmmmmmml{mm}$		
Dobes.		240.480.730	and 970 mg/m3 respectively. or 10, 50,		
		100, 150, 200	ppm, respectively)		
Control Group:		yes, concurren	no treatment		
Method.	other: r	ange_finding stu	udar.		
GLP:	ves	ange-rinding set	lay		
Test substance:	other TS				
Result:	The aim	of this study wa	is to determine the dose-range for a		
	developm	ental toxicity :	study. Groups of 8 pregnant rats were		
	exposed to vapors of DEAE at concentrations of 0 (control),				
	10, 50, 100, 150, and 200 ppm on gestational days (GD) 6 to				
	Clinical	signs were lim:	ted to rales and nasal or ocular		
	discharg	e (150 and 200 p	opm). Dams exposed to 150 and 200 ppm		
	lost body weight. Body weight gains were markedly reduced at				
	100 ppm (GD 6-15) and slightly reduced at 50 ppm (GD 12-15).				
	Decrease	d food consumpt:	on was noted at 100 ppm and above.		
	Pregnanc	y rates, mean u	erine implantation data and retal body		
	post-imp	lantation loss s	aried slightly; according to the		
	authors,	this was not of	E biological significance. A relatively		
	high inc	idence of resor	ptions was seen in several groups,		
	includin	g the control.	According to the authors, this may have		
	been ind	icative of effe	ts due to general stress associated		
	with inh	alation exposure	e, including the absence of food and		
	any obse	rvable abnormal	ties upon external examination		
	External	fetal malformat	cions were limited to abnormal flexure		
	of the h	ind-limb in one	fetus each of the 50 and 100 ppm		
	groups.	Based on these i	cesults, concentrations of 0, 33, 66,		
	and 100	ppm were selecte	ed for the main study.		

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

Test substance: Reliability: 10-JAN-2003	CAS No. 100-37-8 (2-diethylaminoethanol), purity: 99.8%, Atochem North America (2) valid with restrictions Meets the necessary criteria of a range finding study. (97)
Species: Strain: Route of administr Exposure period: Frequency of treat Duration of test: Doses: Control Group:	rat Sex: female Sprague-Dawley ation: gavage days 0 to 11 of gestation ment: daily until day 12 of gestation 10, 30, 100 and 250 mg/kg bw yes
Method: GLP: Test substance:	other yes other TS
Method:	The TS was administered to 5 bred female Crl:CD(SD)Br rats/ group. To achieve the desired doses of 10, 30, 100 and 250 mg/kg bw, the dosage volume used was 1, 3, 3.3 and 8.3 ml/kg of 10, 10, 30 and 30 mg/ml solutions. The control group received 8.3 ml/kg of the vehicle. Clinical observations were made twice daily. Body weight and food consumption were measured on GD 0, 1, 3, 7, 9, 11 and 12. The doses were chosen based on a preliminary toxicity study (7 days of treatment in non-mated females). On day 12 the thoracic, abdominal and pelvic cavities were opened and examined. The aterus and ovaries were exposed and examined, and any abnormalities were recorded. The no. of corpora lutea in each ovary were recorded. The uterus was opened and the number of implantation sites were recorded. The liver and kidneys were weighed at necropsy from all groups, and in the control and high dose group these tissues were prepared for microscopic examination. Paired organs were weighed collectively. The liver, kidneys and gross lesions from all maternal animals were preserved in formalin. Uteri with no macroscopic evidence of nidation were opened and put in 10% ammonium sulfide for the detection of early implantations.
Remark:	According to the report, the NOEL was 100 mg/kg bw for maternal toxicity and embryotoxicity. The basis for this was the observation of rales in 2/5 dams of the 250 mg/kg dose and the lack of this finding in the 100 mg/kg dose. Thus, the LOEL for the dams was 250 mg/kg bw. Histologic examinations were limited to the kidneys and liver. Due to the low number of animals in the study, the assignment of clear substance-related embryotoxic effect at 250 mg/kg bw is also difficult since the increase in postimplantaion loss and the decrease in live litter size was due to one animal with 9
Result:	All animals survived to sacrifice. The only treatment-related clinical finding in was rales seen in 2/5 dams in the 250 mg/kg bw dose. No internal findings were seen macroscopically. Body weights, body weight gain, food consumption and organ weights of the dams were not effected by treatment. No treatment-related microscopic findings were seen in the liver and kidneys of the 250 mg/kg dosed

	dams. In the 250 mg/kg bw dose group post implantation loss was increased by 16.6% (S.D. 20.90), and the number of viable embryos was decreased by 15% (83.4% in the 250 mg/kg group versus 98.6%) in controls). The increase in postimplantation loss and the decrease in live litter size in this group was predominantly due to one female with nine early resorptions (52.9%). Intrauterine parameters were unaffected by treatment in the 10 mg/kg dose	e t
Test substance: Reliability:	<pre>groups. 2-diethylaminoethanol, "100% pure", Elf Atochem North America (2) valid with restrictions Performed under GLP conditions, the protocol was not widely used. Only early embryo-toxicity can be addressed due to the study design. The study is limited by the number of females used (n = 5).</pre>	a e
25-JUL-2002	98)	3)

5.8.3 Toxicity to Reproduction, Other Studies

5.9 Specific Investigations

5.10 Exposure Experience

Remark: Reliability:	An odor threshold of 0.011 ppm has been reported. (2) valid with restrictions 2.2; basic data given, restrictions Critical study for SIDS endpoint
12-AUG-2002	(99)
Remark: Reliability:	An attempt by a laboratory worker to remove animals from an inhalation chamber containing approx. 100 ppm 2-diethylaminoethanol resulted in nausea and vomiting within 5 minutes after a fleeting exposure; no irritation of the eyes or throat was noted during this brief exposure. Other persons in the same room also complained of a nauseating odor but showed no ill effects. (2) valid with restrictions 2.2; basic data given, restrictions
Flag:	Critical study for SIDS endpoint
12-AUG-2002	(60)
Remark:	Two boilers were prepared for operation by adding corrosion inhibiting chemicals, 2-diethylaminoethanol and cyclohexylamine. Steam produced by the boilers was used for humidity in the work area. Symptoms consistent with acute toxic effects of 2-diethylaminoethanol and cyclohexylamine were noted in 65 of the employees. These included nausea, dizziness, vomiting, and eye, nose, and throat irritation. Exposure concentration is not known.
Reliability:	(2) valid with restrictions2.2; basic data given, restrictions
Flag:	Critical study for SIDS endpoint
12-AUG-2002	(100)

OECD SIDS	2-DIETHYLAMINOETHANOL
5. TOXICITY	DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8
Remark: Reliability:	2-diethylaminoethanol was used to humidify the air in a building. Of 14 samples taken, only two had detectable amounts of 2-diethylamioethanol (0.05 and 0.04 mg/m3). Two bulk samples contained about 30 mg per square meter of the exposed area. A total of 16 of the 35 employees, who participated in medical interviews, complained of eye irritation, 13 of skin irritation, and 6 of headache, nose and throat irritation, or dizziness. Six females reported gynecological problems. Since 2-diethylaminoethanol has a low vapour pressure and was detected on surfaces, skin contact with surfaces was a possible route of absorption. (2) valid with restrictions 2.2; basic data given, restrictions
Flag: 12-AUG-2002	Critical study for SIDS endpoint (101)
Remark:	Through a leak in the steam heating system, the anticorrosive agent 2-diethylaminoethanol was released into the air of a large office building. Irritative symptoms of the respiratory tract, and ear, nose and throat were experienced by most of the 2500 employees, and 14 workers developed asthma within 3 months of exposure. Seven of the 14 cases were defined as "confirmed" and 7 of 14 as "suspect", using the NIOSH case definition of occupational asthma. Spirometry obtained in 12 cases was positive in 4 and peak flow testing in 10 of 11 tested persons. Three cases were diagnosed on the basis of work-related symptoms and physical examination alone.
Reliability:	(2) valid with restrictions
Flag: 12-AUG-2002	2.2; basic data given, restrictions Critical study for SIDS endpoint (102)
Remark:	Approximately 15 employees in the office support area of a production building had been experiencing rashes. A medical evaluation, consisting of interviews, skin examinations, and a review of medical records was conducted. 2-diethylaminoethanol was identified as the only volatile component. However, environmental samples did not reveal any 2-diethylamioethanol in air samples. Skin examinations revealed an irritant-type rash on the exposed areas of the faces, neck, and hands. The distribution of the rash was consistent with and suggestive of a phototoxic skin reaction. Both the environmental and medical evaluations indicated the source of the dermtitis to be the air-handling system. However, no specific etiologic agent has been identified.
26-JUN-2002	(103)
Remark:	Two subjects received 5.6 grams of diethylaminoethanol hydrochloride (CAS 14426-20-1) intravenously in 11.2 % solution. The same two subjects received 5.6 grams of diethylaminoethanol hydrochloride orally in aqueous solution. About 25 % of the drug was excreted in the urine; the remainder was metabolized by an unknown route. A single dose is almost completely metabolized or excreted in 8 hours. The effect of diethylaminoethanol hydrochloride was tested in 14 subjects with ventricular premature contractions in a dose ranging from 05. to 5 grams injected

<u>OECD SIDS</u>	2-DIETHYLAMINOETHANOL
5. TOXICITY	DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8
	at a rate of 1 gram per minute. The ectopic ventricular beats disappeared in 13 of 14 cases. In 10 cases it was transient, lasting from 3 to 20 minutes. In 3 cases the ectopic rhythm had not reappeared within a week. Shortly after the injection, most subjects noted a peculiar taste variously described as bitter, metallic, or peppermint-like, followed by a sensation of warmth, dizziness and fluttering in front of the eyes. Nauseas and vomiting were observed in about 10 to 15 min. in about 15 % of the cases but did not occur until the change in rhythm had been effected. These side effects were usually transient and disappeared 10 to 15 min. following injection. In some instances a transitory fall in both the diastolic and systolic blood pressure was
	noted which disappeared within 20 minutes. Diethylaminoethanol is a product of hydrolysis, in vivo, of procaine.
26-JUN-2002	(104) (105)
Remark:	Report of therapeutic i.v. administration (1 g in 5 ml, given once a day for up to 10 consecutive days) of diethyaminoethanol in small groups of patients with perioheral circulatory disturbances, hypertension, and bronchial asthma. Feeling of warmth, sweet taste, double vision, dizziness, and nausea were the reported side effects. No further information was given on the galenic of the drug (Debydasal)
26-JUN-2002	(106)
Remark:	Inhibition of lactation was observed in women receiving diethylamioethanol for eclampsia (1 g i.v. once a day for up to 11 days or given as suppository for up to 5 days, dosage of the suppository and the galenic of the drug `Dehydasal` not given). Feeling of warmth, dry mouth, flickering in front of the eyes, and dizziness were the reported side offects
26-JUN-2002	(107) (108)

5.11 Additional Remarks

Type: Biochemical or cellular interactions

Remark: To determine whether concentrations of DEAE and procaine below those that reduce the amplitude of action potentials might alter the excitability of brain cells, a single microelectrode intracellular recording technique was used to measure firing threshold and action potential amplitude of pyramidal cells in rat hippocampal slices. At low concentrations of both DEAE (less than or equal to 5 mM) and procaine (less than or equal to 0.5 mM), firing threshold was significantly increased (P < 0.01), whereas action potential spike amplitude was minimally altered. At higher concentrations, both drugs significantly decreased action potential spike amplitude (P < 0.025) as well as increased firing threshold (P < 0.001). DEAE tended to increase threshold relatively more than procaine, when drug concentrations that similarly reduced action potential amplitude were compared. All actions of DEAE and procaine

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

were reversible. Inhibition of action potentials by DEAE and procaine was clearly concentration-dependent (P <= 0.015). DEAE effects on threshold were marginally concentration-dependent (P = 0.08); procaine did not demonstrate clear concentration-dependent effects (P = 0.33) over the concentrations tested in this study. These similar actions of procaine and DEAE on brain cells suggest a mechanism by which intravenous local anesthetics may contribute to the general anesthetic state. Moreover, it appears possible that procaine metabolism and DEAE accumulation may underlie the prolonged effects sometimes seen after intravenous procaine administration. 2-diethylaminoethanol

29-APR-2002

Test substance:

(109)

Type: Biochemical or cellular interactions

Remark: To test whether the products of procaine hydrolysis have local anesthetic actions resembling those of procaine, the authors compared the ability of procaine and its metabolites DEAE and para-aminobenzoic acid (PABA) to block compound action potentials in excised, desheathed frog and rat sciatic nerves. Studies were performed in solutions of impermeant buffers at pH 7.4 (corresponding to mammalian physiologic pH) and at pH 9.2 (close to the pKa of procaine and DEAE) to test for extracellular pH-dependent increases in drug permeation and potency. Both procaine and DEAE inhibited compound action potentials at pH 7.4 and 9.2 in a reversible and dose-dependent manner, and both were approximately ten-fold more potent at pH 9.2 than at pH 7.4, procaine inhibiting the action potential height by 50% at 0.15 mM (pH 9.2) and 1.1 mM (pH 7.4), DEAE at 4 mM (pH 9.2) and 70 mM (pH 7.4). In contrast, PABA at concentrations up to 25 mM and at either pH failed to inhibit compound action potentials, and did not modify the effects of DEAE when both drugs were given together. Procaine produced greater use-dependent block at the higher pH and at higher stimulation rates (100 Hz vs. 40 Hz); DEAE produced almost no use-dependent block. These observations suggest: 1) that DEAE might account for some of the neuropharmacologic activity of procaine in techniques that favor the accumulation of metabolites (such as those requiring large doses or prolonged infusions); and 2) that alkalinization of procaine and DEAE solutions appears to increase their potency for both resting and use-dependent block of action potentials. Test substance: 2-diethylaminoethanol 01-OCT-2001 (110)Type: Biochemical or cellular interactions Remark: 3.1 mmol/l of DEAE inhibited in vitro cholinesterase

Test substance: 2-diethylaminoethanol solution neutralized to pH 6.8 to 7.2 prior to use

01-OCT-2001
OECD SIDS 5. TOXICITY

Type:	Biochemical or cellular interactions
Remark:	In male rats the daily oral application of DEAE at 100 mg/kg bw for 2 weeks led to a statistically significant increase in the liver protein synthesis rate (159% of the control value). Six animals were used per group.
Test substance:	2-diethylaminoethanol
01-OCT-2001	(111)
Туре:	Biochemical or cellular interactions
Remark: Test substance:	Rats given DEAE for 2 weeks at a level of 200 mg/kg of feed had a slight increase in the level of acetyl coenzyme A (increased from ca. 12 to ca 16 ug/g of tissue) and a decrease in the level of coenzyme A (reduced from ca. 38 ug/g to 30 ug/g tissue). According to the report, the changes were statistically significant. This effect was not seen in tissues from the cerebral cortex, heart, muscle or duodenum. 2-diethylaminoethanol
10-NOV-1993	(112)
Type:	Biochemical or cellular interactions
Demonito	
Test substance:	oxygen radicals (FOR) by polymorphonuclear (PMN) leukocytes obtained from rabbits and humans was studied. New Zealand rabbits were given daily i.m. injections of the test substance (15 mg/kg/d) for 35 days; another group of rabbits served as control. Further, human leukocytes (isolated from the peripheral blood of 10 healthy volunteers) were treated with the test substance. The FOR released by the PMN leukocytes were evaluated by in vitro chemoluminescence. Addition of the test substance had no effect on the FOR by PMN leukocytes of control rabbits. In rabbits treated with the test substance, the release of FOR by PMN leukocytes was much more reduced. In treated rabbits, the intensity of the emitted light was 3.46 mV (control: 6.74 mV). An inhibitory effect of the test substance on the FOR release was observed in the human PMN cells stimulated with opsonized zymosan (50 mV in treated cells vs. 67.7 mV in control cultures). According to the authors, the fact that the inflammation was associated with accumulation of free radicals suggested the opportunity to evaluate the test substance as an antioxidant agent. 2-diethylaminoethanol
01-OCT-2001	(113)
Туре:	Chemobiokinetics general studies
Remark:	In a gavage study with rats, 14C-labeled-2-diethylaminoethanol-HCl was reported to be rapidly absorbed into the blood stream. With a dose of 68 mg/kg the maximum concentration in the blood was reached in 30 minutes. With 679 mg/kg the maximum concentration in the blood was reached within 1 hour. Elimination occurred primarily via

OECD SIDS 5. TOXICITY

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

(115)

the kidney. Elimination via exhalation and the feces played only minor role. The kinetics of urinary elimination were affected by the dose. In this regard, by 6 hours after the application of a 679 mg/kg bw dose 40% was eliminated in the urine, and by 24 hours after application 58.5% was eliminated. When a 68 mg/kg dose was given, then after 6 and 24 hours 17.5% and 37.4% were excreted via the urine, respectively. Τn the experiment with 679 mg/kg 2-diethylaminoethanol, 90% of the test substance had been eliminated via the urine 10 days after treatment. Some radioactivity was still detectable in the urine 40 days after treatment. 2-Diethylaminoethanol was predominantly (> 60%) excreted unchanged over the first 96 hours. In the same period, the following metabolites were seen based on the recovery of radioactive compounds: 2-ethylaminoethanol (ca. 1%), phosphoric acid-mono-(2-diethylaminoethylester) (2-8%), diethylaminoacetic acid (ca. 10%) and the N-oxide of DEAE (ca. 15-19%). Incorporation into phospholipids was observed. In this study, autoradiography indicated that 2-diethylaminoethanol was widely distributed throughout the body after gavaging. 2-Diethylaminoethanol was concentrated in the liver, reaching a maximum at 7 hours, but thereafter, it decreased. Initially, the central nervous system showed very low levels of activity, but by day 7 it had increased. For the oral dose of 679 mg/kg the biological half-life was 19 hours and for the 67.9 mg/kg dose it was 36 hours. Test substance: 2-diethylaminoethanol Flag: Critical study for SIDS endpoint 10-JAN-2003 (114)Type: Chemobiokinetics general studies Remark: N.B.: The test substance was never clearly identified, but it was presumably 2-diethylaminoethanol. Experimental radiopharmacokinetic studies were carried out on female Wistar rats, 23 months and 5 months old, injected with 99mTc-DEAE 2.5 mg/kg b.w., via i.m. administration, and 7.5 mg/kg via p.o. The absorption and the biodistribution of the labeled product were followed in seven organs, including the brain (in thirteen areas of the brain), during the first 240 minutes after administration. The results allowed the conclusion that DEAE is rapidly absorbed and distributed in the different organs, irrespective of administration. The affinity of the different organs or brain areas to DEAE depends on age. Investigations regarding the fate of 99mTc-DEAE in the rat body point out that it undergoes a metabolic splitting leading to ethanolamine, glycine and urea. Elimination studies infer that DEAE is eliminated unchanged or as the mentioned metabolites, especially by the kidney, following first order elimination kinetics. The elimination time is about 72 hours and the half time about 28. Test substance: 99m-Tc-DEAE, radiochemical purity > 95%, and radionuclidic purity 99.9%; no further data.

07-AUG-2002

Type: Distribution

OECD SIDS	2-DIETHYLAMINOETHANOL
5. TOXICITY	DATE: 07-MARS-2003
	SUBSTANCE ID: 100-37-8
Remark:	In a study with 2 humans the plasma concentration peaked 3 hours after an oral administration of 5.6 g of 2-diethylaminoethanol-HCl, but was almost undetectable after 8 hours. About 25% of the 2-diethylaminoethanol was excreted unchanged in the urine within 48 hours. Similar excretion results were observed after intravenous administration.
Test substance:	In the same article it was reported that 2-diethylaminoethanol-HCl given to dogs by intravenous infusion (71 mg/kg bw), distributed rapidly. Three hours after infusion the level of 2-diethylaminoethanol was higher in the tissues examined (muscle, heart, brain, lung, liver and spleen) than in the plasma. 2-diethylaminoethanol-hydrochloride
26-JUL-2002	(116)
Type:	Excretion
Remark:	Male Sprague-Dawley rats received 15 mmol/kg via gavage. 24 hr later urine was collected, and measurements were made for several amines: approx. 1% of the dose applied was found in the form of trimethylaminoxide, 0.1 % as diethylamine and 0.05% monomethylamine.
Test substance:	2-diethylaminoethanol
29-APR-2002	(117)
Туре:	Excretion
Remark:	14C-Labeled-2-diethylaminoethanol-HCl was given to rats by intravenous injection at doses of 2.9 µmol/rat (ca. 1.94 mg/kg bw). Cumulative excretion of 19.9 and 42.2% of the radioactivity in the urine was observed after 24 and 48 hours, respectively. Additionally, 8.5 and 29.5% of the radioactivity was excreted via the feces during the same time interval. Excretion via the bile was only measured over the first 6 hours, and was reported to be 5%.
Test substance:	2-diethylaminoethanol
26-JUL-2002	(118)
Type:	Metabolism
Remark:	The article cites IARC (vol. 17, 1978) to say that on the basis of the chemical structure there is the potential for DEAE to be nitrosated to form N-nitrosodiethylamine and N-nitrosoethylethanolamine, both carcinogens in animals. It also mentions that nitrosation of tertiary amines is normally slower as compared to secondary amines, however, conditions exist where tertiary amines can be readily nitrosated. No data on the kinetics of such reactions was available.
Test substance:	2-diethylaminoethanol
02-OCT-2001	(119)
Type:	Metabolism

<u>OECD SIDS</u> 5. TOXICITY	<u>2-DIETHYLAMINOETHANOL</u> DATE: 07-MARS-2003	
	SUBSTANCE ID: 100-37-8	
Remark:	Abstract only: Unmetabolized diethylaminoethanol and diethylmethylethanolamine was found in the liver of rats fed the test substance.	
Test substance:	2-diethylaminoethanol, no further data	
27-JUN-2002	(120)	
Туре:	Metabolism	
Remark: Test substance:	The nitrosation potential of the test substance is discussed. Based on the chemical structure, the test substance is presumed to have the potential to be nitrosated to N-nitrosodiethylamine and N-nitrosoethylethanolamine. According to the author, both of these compounds are regarded as potent carcinogens in animals. Considering the nitrosation potential of the test substance, contamination with secondary amines, such as diethanolamine, ethylethanolamine, and diethylamine (which can be nitrosated to N-nitrosodiethanolamine, N-nitrosodiethylamine, and ethylhydroxy-ethylnitrosmine) should be taken into account. 2-diethylaminoethanol	
29-APR-2002	(121)	
Туре:	other: Absorption	
Remark: Test substance:	The dermal penetration rate (flux) of a saturated aqueous solution of the test substance was calculated to be 3.44 mg/cm3 per hour based on its physical properties. Based on these calculations, the test substance is expected to have a high dermal penetration rate and has a potential for dermal toxicity. According to the authors, this suggests that dermal absorption of the test substance would raise the biological levels above those occurring during inhalation of concentrations equal to the threshold limit value (0.05 mg/L). 2-diethylaminoethanol	
08-JAN-2003	(122)	
Туре:	other: Dermal toxicity	
Remark:	Based on modeling data, DEAE has been predicted to be readily absorbed by via the skin (Fiserova-Bergerova et al., 1990); however, this article reports that the model of Fiserova-Bergerova et al. was too conservative, and is likely	
Test substance:	to overestimate percutaneous penetration. 2-diethylaminoethanol	
29-APR-2002	(123)	
Туре:	other: MAK Value	
Remark:	The MAK value (1997) was given as 5 ml/m3 (ppm) which corresponds to 24 mg/m3.	
07-JAN-2003	(124)	
Туре:	other: NEG and NIOSH Basis for an Occupational Health Standard	

OECD SIDS 5. TOXICITY

Test substance:	2-diethylaminoethanol	
27-JUN-2002	(125)	(121)
Type:	other: OSHA PEL	
Remark: Test substance:	The current OSHA standard for DEAE is 10 ppm (50 mg/m3) averaged over an 8 hr work shift. This is also known as permisible exposure limit (PEL). 2-diethylaminoethanol	the
29-APR-2002		(126)
Туре:	other: TLV Citations	
Test substance:	2-diethylaminoethanol	
29-APR-2002	(127) (128) (129)	(130)
Туре:	other: Waltzing syndrome	
Remark:	The test substance did not produce the waltzing syndrome (hyperactivity and impaired coordination) in rats	
Test substance:	2-diethylaminoethanol	
29-APR-2002		(131)

6. ANALYT. METH.FOR DETECTION & IDENTIFICATION

DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

6.1 Analytical Methods

<u>6.2 Detection and Identification</u>

SUBSTANCE ID: 100-37-8

7. EFF. AGAINST TARGET ORG. AND INTENDED USES

7.1 Function

7.2 Effects on Organisms to be Controlled

7.3 Organisms to be Protected

<u>7.4 User</u>

7.5 Resistance

8.1 Methods Handling and Storing

- 8.2 Fire Guidance
- **8.3 Emergency Measures**
- 8.4 Possib. of Rendering Subst. Harmless
- **8.5 Waste Management**
- **<u>8.6 Side -effects Detection</u>**
- **8.7 Substance Registered as Dangerous for Ground Water**
- **8.8 Reactivity Towards Container Material**

- (1) BASF AG, Safety Data Sheet N,N-DIETHYLETHANOLAMINE, 05.07.2000
- (2) Commission Directive 2000/33/EC, 25 April 2000 (27th adaption to the technical progress of 67/548/EEC)
- (3) TRGS 900 (Technical guidance for hazardous substances -Technische Regeln für Gefahrstoffe) (Germany) of 09/2001
- (4) VwVwS (Administrative Regulation on the Classification of Substances Hazardous to Waters into Water Hazard Classes -Verwaltungsvorschrift wassergefährdende Stoffe - VwVwS) (Germany) of 17.05.1999
- (5) National Chemical Inventories, 2001 Issue 2
- (6) BASF AG, Analytik und Messtechnik, unpublished data, 126157, 05.11.1985
- (7) BASF AG, Physikalisch-chemische Konstanten, unpublished study, report BRU 85.219, 16.12.1985
- (8) Beilstein, 2001
- (9) Steele W.V., et al., Measurements of Vapor Pressure, Heat Capacity, and Density along the Saturation Line for Cyclopropane Carboxylic Acid, N,N-Diethylethanolamine, 2,3-Dihydrofuran, 5-Hexen-2-one, Perfluorobutanoic Acid, and 2-Phenylpropionaldehyde, J. Chem. Eng. Data 47, 715-724, 2002
- (10) Hazardous Substances Data Bank, 2001
- (11) DiGuilio R.M., et al., Densities and Viscosities of the Ethanolamines, J. Chem. Eng. Data 37, 239-242, 1992
- (13) BASF AG, Physikalische Chemie, unpublished report, report BRU 83.67, 14.04.1983
- (14) BASF AG, Department of Product Safety, unpublished calculation, 27.06.2002
- (15) Rekker R.F., The Hydrophobic Fragmental Constant, Elsevier Scientific Publishing Company, Amsterdam, 1977
- (16) BASF AG, Analytisches Labor; unpublished report, report BRU 87.262, 18.12.1987
- (17) Verschueren K., Handbook of Environmental Data on Organic Chemicals, 2nd ed., Van Nostrand Reinhold, New York, 1983
- (18) BASF AG, Labor fuer Umweltanalytik; unpublished study, 09.01.1989

- (19) Newsome L.D., A QSAR study of the toxicity of amines to the fathead minnow. The Science of the Total Environment, 109/110, 537-551, 1991
- (20) Newsome L.D., Validation and upgrade of a QSAR study of the toxicity of amines to freshwater fish. Env. Toxicol. and Risk Ass. ASTM 1179, 413-426, 1993 (230)
- (21) BASF AG, Sicherheitstechnische Kenndaten, unpublished report, SIK-No. 85/0729, 28.08.1985
- (22) BASF AG, Sicherheitstechnik, internal notice, 11.11.1999
- (23) Perrin D.D., Dissociation constants of organic bases in aqueous solution. IUPAC Chem Data Ser., Buttersworth, London, 1965 (as cited in Hazardous Substances Data Bank, 2001)
- (24) Sax N.I. and Lewis R.J. (eds.). Hawley's Condensed Chemical Dictionary. 11th ed. New York: Van Nostrand Reinhold Co., 1987. 387 (as cited in Hazardous Substances Data Bank, 2001)
- (25) BASF AG, Department of Product Safety, unpublished calculation, 09.07.2001
- (26) Perry D.L. et al., Identification of Organic Compounds in Industrial Effluent Discharges, Battelle Columbus Labs, OH, prepared for EPA, Washington, DC, Office of Toxic Substances, PB-291 900, EPA-560-6/78-009; Nov 1978
- (27) Ellis D.D.; Cyrenius M.J.; Larson R.A.; Schaeffer D.J.: Organic constituents of mutagenic secondary effluents from wastewater treatment plants. Arch. Environm. Contam. Toxicol. 11, 373-382, 1982
- (28) Salthammer T., Emission of volatile organic compounds from furniture coatings, Indoor Air, 7 (3), 189-197, 1997
- (29) NIOSH, Health Hazard Evaluation Report, HETA 83-020-1351, Aug 1983
- (30) Volent P., Baer N.S., Volatile amines used as corrosion inhibitors in museum humidification systems, The International Journal of Museum Management and Curatorship, 4, 359-364, 1985
- (31) Edgerton S.A., Kenny D.V., Joseph D.W., Determination of amines in indoor air from steam humidification, Environ. Sci. Technol., 23 (4), 484-488, 1989
- (32) BASF AG, Department of Product Safety, unpublished calculation, 10.07.2001
- (33) BASF AG, Department of Product Safety, unpublished calculation, 26.06.2002
- (34) Meylan W. et al., Environ. Sci. Technol., 26 (8), 1580-1587, 1992

9. REFERENCES

2-DIETHYLAMINOETHANOL DATE: 07-MARS-2003 SUBSTANCE ID: 100-37-8

- (35) Lyman W.J.; Reehl W.F.; Rosenblatt D.H.: Handbook of Chemical Property Estimation Methods. American Chemical Society, Washington, DC, 1990
- (36) BASF AG, Department of Product Safety, unpublished study, 02/0430/21/1, 01.10.2002
- (37) BASF AG, Department of Ecology, unpublished study, report No. 94/0979/10/1, 1994
- (38) Rothkopf G.S.; Bartha R.: Structure-biodegradability correlations among Xenobiotic industrial amines. J.Amer. Oil Chem. Soc. 61,977-80, 1984
- (39) BASF AG, Department of Ecology; unpublished study, 1981
- (40) Babeu L. and Vaishnav D.D., Prediction of biodegradability for selected organic chemicals. Journal Industrial Microbiology, 2, 107-115, 1987
- (41) Geiger, D.L. et al., Acute Toxicities of Organic Chemicals to Fathead Minnows (Pimephales promelas), Vol. 3, Center for Lake Superior Environmental Studies, University of Wisconsin, Superior, WI, 149-150, 1986
- (42) BASF AG, Department of Toxicology, unpublished study, 87/267, 09.11.1987
- (43) Cash G.G., Clements R.G., Comparison of structure-activity relationships derived from two methods for estimation octanol-water partition coefficients, SAR and QSAR in Environmental Research, 5, 113-124, 1996
- (44) Martin T.M., Young D.M., Prediction of the acute toxicity (96-h LC50) of organic compounds to the fathead minnow (Pimephales promelas) using a group contribution method; Chem. Res. Toxicol., 14, 1378-1385, 2001
- (45) Atofina: Project No. 159142, Exxon Biomedical Sciences, Inc., Dec 14 1993
- (47) BASF AG, Department of Ecology, unpublished study,
 3-BASF-ökolimna-12/88/-026, report of ÖKOLIMNA Gesellschaft für Ökologie und Gewässerkunde mbH, 1988
- (48) Dr. A. Weyers, Techniche Universität Dresden, Beratergremium für Altstoffe (BUA), unpublished statistical analysis of the algal growth inhibition data, 19.07.2002
- (49) BASF AG, Department of Ecology; unpublished study, report No. 94/979/8/1, 1994
- (50) BASF AG, Analytisches Labor; unpublished study, 307371/1988, 1988

OECD SIDS 9. REFERENCES

- (51) Horst R.K. et al., Chlorosis in healthy and viroid-infected plants exposed to the steam additive diethanolamine; Scientia Horticulturae, 19, 1-8, 1983
- (52) Kawamoto S.O., Horst R.K., Phytotoxicity caused by diethylaminoethanol, an anticorrosion compound used in steam lines, Proc. Am. Phytopathol. Soc., 4 190, 1977
- (53) Drummond D.B., Vasiloff G.N., Hill A.W., Live steam humidification as a source of phytotoxic contaminants in controlled environment growth chambers and greenhouse, Proc. Am. Phytopathol. Soc., 1, 135, 1974
- (54) Nishiuchi Yasuhiro, Toxicity of agrochemicals to freshwater organism. CIII. Solvents; Suisan Zoshoku, 32 (2), 115-119, 1984
- (55) BASF AG, Department of Toxicology, unpublished report (XVIII/320), 27 Jan 1969
- (56) Smyth, H.F., Jr. & Carpenter, C.P., J. Ind. Hyg. Toxicol. 26, 269-273, 1944
- (57) Smyth, H.F., Jr., unpublished data from personal communication given through a member of the TLV Committee (November 1964); cited in: Documentation of the Threshold Limit Values and Biological Exposure Indices, 2-diethylaminoethanol, pp. 140-141, 1980
- (58) TSCATS, OTS 0000996, Doc. ID. FYI-OTS-0794-0996, Mellon Institute of Industrial Research for Union Carbide Corp. 3-02-84
- (59) Cornish, H.H. and Adefuin, J., Fd. Cosmet. Toxicol. 5, 327-332, 1967
- (60) Cornish H.H, Am. Ind. Hyg. Assoc. J., 26, 479-484, (1965)
- (61) Hartung, R. and Cornish, H.H., Toxicol. Appl. Pharmacol., 12, 486-494, 1968
- (63) Izmerov, N.F. et al.: Toxicometric Parameters of Industrial Toxic Chemicals Under Single Exposure, United Nations Program (UNEP), Moscow (1982); cited in: TLV documentation "2-diethylaminoethanol",1996
- (64) RTECS, Update Code 9107 (August 1991): Journal of Pharmacology and Experimental Therapeutics 94, 249, 1948
- (65) Lewis, R.J. und Tatken, R.L. (eds.): Registry of Toxic Effects of Chemical Substances, Rockville, MD, U.S. Dept. of Health, Education, and Welfare, Public Health Service, Center for Disease Control, NIOSH, p. 763, 1982, cited in: Assessment of the Health Risks of Diethylaminoethanol and Morpholine, National Research Council, prepared for Army Medical Research and Development Command, PB85-122771, August 1983

- (67) Koch, R., Arzneimittel Forsch. 4, 649-654 (1954); cited in: TLV documentation "2-diethylaminoethanol", 1996
- (68) RTECS, Update Code 9107 (August 1991): Arzneimittel-Forschung 9, 31 (1959)
- (69) BASF AG, Department of Toxicology, unpublished report (82/18), 26.08.1982
- (70) Potokar, M. et al., Food Chem. Toxicol. 23, 615-617, 1985
- (71) BASF AG, Department of Toxicology, unpublished report (79/529), 5.5.1980
- (72) TSCATS, OTS0536389, Doc.I.D. 88-920002337, Bushy Run Res Ctr, for Union Carbide Corp., 8/26/86
- (73) TSCATS, OTS0537485, Doc.I.D. 88-920003529, Project Report 53-21; Bushy Run Res Ctr, for Union Carbide Corp, 4/06/90
- (74) TSCATS, OTS0558740, Doc. ID. 86960000539, Pharmacology Research Inc., for Elf Atochem North America Inc., 6-28-72
- (75) TSCATS, OTS 0001031, New Doc. ID. FYI-OTS-0794-1031, Penwalt Corp., 2/13/84
- (76) Leung, H.-W. and Blaszcak, D.L., Vet. Hum. Toxicol. 40 (2), 65-67,1998
- (77) BASF AG, Department of Toxicology, unpublished report (1628/U), 28.06.1937
- (78) RTECS, Update Code 9107 (August 1991): Union Carbide Data Sheet 6/11/63
- (79) Nakamura A. et al., Contact Dermatitis 31, 72-85, 1994
- (80) EXXON Biomedical Sciences, Inc.: Projekt No. 208618, Test Material : MRD-87-086, 03.05.1990
- (81) Hinz, J. P. et al., Fundam. Appl. Toxicol. 18, 418-424, 1992
- (82) TSCATS, OTS 0000699, Doc. I. D. FYI-OTS-0689-0699, Synthetic Organic Chemical Manufactures Assoc. (SOCMA), Date Produced: 6/29/89, (note: this TSCAT only has copies of a letter submitted to the EPA by SOCMA)
- (83) TSCATS, OTS 0000699-1, Doc. I. D. FYI-OTS-0790-0699, Synthetic Organic Chemical Manufactures Assoc. (SOCMA), Date Produced: 5/03/90

- (84) Lomonova, G.V.: Gig. Tr. Prof. Zabol. 14 (11), 52-53, 1970; cited in: Documentation of the Threshold Limit Values and Biological Exposure Indices, 2-diethylaminoethanol, Supplements to 6th Edition, 1996
- (85) Glaister, J.R., Principles of Toxicological Pathology, Laboratory-animal pathology, p. 131-203, Taylor and Francis, London, 1986
- (86) TSCATS, OTS0001257, Doc. I. D. FYI-OTS-0395-1257, Penwalt Corp. Date produced: 07/31/67, Date Received: 07/24/94
- (87) TSCATS, OTS0530455, Doc. I.D. 98-900000214, 8EHQ-0890-1043, Atochem N America Inc., 8/02/90
- (88) Cornish, H.H., Am. Ind. Hyg. Assoc. J. 26, 479-484, 1965, as cited in TLV documentation "2-diethylaminoethanol", Suppl. 1996
- (89) BASF AG, Department of Toxicology, unpublished report (88/956), 8 Mar 1989
- (90) Life Science Research, Report No. 91/SHG001/0251, Diethylaminoethanol: Assessment of mutagenic potential in histidine auxotrophs of Salmonella typhimurium (the Ames test), Sponsored research by Synthetic Organic Chemicals Manufacturing Association, 2 July 1991
- (92) Life Science Research, Report No. 91/SHG002/0302, Diethylaminoethanol: Investigation of mutagenic activity at the HPGRT locus in a Chinese hamster V79 cell mutation system, Sponsored research by Synthetic Organic Chemicals Manufacturing Association, 2 July 1991
- (93) Life Science Research, Report No. 91/SHG004/0263, Diethylaminoethanol: Assessment of its ability to cause lethal DNA damage to strains of Escherichia coli, Sponsored research by Synthetic Organic Chemicals Manufacturing Association, 2 July 1991
- (94) Life Science Research, Report No. 91/SHG003/0321, Diethylaminoethanol: Assessment of clastogenic action on bone marrow erythrocytes in the micronucleus test, Sponsored research by Synthetic Organic Chemicals Manufacturing Association, 2 July 1991
- (95) TSCATS, OTS0001257, Doc. I. D. FYI-OTS-0395-1257, Penwalt Corp. 07/31/67
- (96) Exxon Biomedical Sciences, Inc.: Project No. 108634, Test Material MR-87-086, Final Report, 11 Nov 1991
- (97) Leung, H.-W. and Murphy, S.R., J. Appl. Toxicol. 18, 191-196, 1998

OECD SIDS 9. REFERENCES

(98) TSCATS, OTS0559210, Doc. ID 89980000077, Date Produced 11/12/97, Wil Research for Elf Atochem North America Inc (99) Amoore, J., E., Hautala, E.; J. Appl. Toxicol. 3, 272-290, (1983) (100) Hills, B., Lushniak, B.; Health Hazard Evaluation Report No. HETA-89-057-2003, NIOSH, Cincinnati, (1989) (101) Fannick, N., Lipscomb, J., McManus, K.; Health Hazard Evaluation Report No. HETA-83-020-1351, NIOSH, Cincinnati, (1983) (102) Gadon, M., E., et al; J. Occup. Med. 36, 623-626, (1994) (103) McManus K., Baker D. B., Health Hazard Evaluation Report No. HETA-81-247-958, NIOSH, Cincinnati, (1981)(104) Papper E.M., et al, NY State J. Med., 48, 1711-1714, (1948) (105) Rosenberg B., et al, J. Pharmacol. Exp. Ter., 95, 18-27, (1949) (106) Grothe, I., Z. Aerztl. Fortb., 44, 1-8, (1950) (107) Rautenberg, O., Dt. Gesundheitsw., 10, 582-584, (1955) (108) Waitz, R., Dt. Gesundheitsw., 10, 573-574, (1955) (109) Butterworth, J.F. & Cole, L.R., Anesth. Analg. 71, 404-410 (1990)(110) Butterworth, J.F. IV et al., Anesthesiology, 68, 501-506, 1988 (111) Kaemmerer, K. & Kietzmann, M., Z. Alternsforsch. 44, 189-199, 1989 (112) Kietzmann, M. & Kaemmerer, K., Z. Alternsforsch. 44, 211-217, 1989 (113) Dolganiuc, A. et al.: Rom. Arch. Microbiol. Immunol. 57 (1), 23-32, 1998; cited in: DIMDI database, Doc. No. CA13008090213B (114) Schulte, K.E et al., Arzneimittel-Forsch., 22, 1381-1390 1972 (115) Dobre, V., et al., Rom. J. Gerontol. Geriatr., 13, 75-84, 1992 (116) Rosenberg B., et al, J. Pharmacol. Exp. Ter., 95, 18-27, (1949) (117) Zeisel, S.H. et al., Food Chem. Toxicol. 27, 31-34 (1989) (118) Michelot, J. et al., Xenobiotica 11, 123-130, 1981

OECD SIDS 9. REFERENCES

- (119) National Research Council, An Assessment of the Health Risks of Morpholine and Diethylaminoethanol, PB85-122661, Washington D.C., 1983
- (121) Toren, K., NEG and NIOSH Basis for an Occupational Health Standard, 2-Diethylaminoethanol, Arbete och Haelsa 25, 1-17, 1994
- (122) Fiserova-Bergerova, V. et al., Am. J. Ind. Med. 17, 617-635, 1990
- (123) Guy R.H. and Potts R.O.: Am. J.Ind. Med. 23: 711-719, 1993
- (124) Occupational Toxicants, Critical Data Evaluation for MAK Values and Classification of Carcinogens, Greim, H. (ed.), Deutsche Forschungsgemeinschaft, Wiley-VCH, Weinheim, vol.14, pp. 91-100, 2000
- (125) NEG and NIOSH Basis for an Occupational Health Standard, DHHS Publication No. 96-104, May 1996
- (126) OSHA, Occupational Health Guideline for Diethylaminoethanol, U.S. Department of Health and Human Services, U.S. Department of Labor, Occupational Safety and Health Administration (OSHA), 5 pages, September 1978
- (127) Documentation of the Threshold Limit Values and Biological Exposure Indices, 2-diethylaminoethanol, 5th edition, p. 198, 1986
- (128) Documentation of the Threshold Limit Values and Biological Exposure Indices, 2-diethylaminoethanol, 6th edition, pp. 462-463, 1991
- (129) Documentation of the Threshold Limit Values and Biological Exposure Indices, 2-diethylaminoethanol, pp. 140-141, 1980
- (130) Documentation of the Threshold Limit Values and Biological Exposure Indices, 2-diethylaminoethanol, Supplements to 6th Edition, 1996
- (131) Goldin, A. et al.: J. Pharmacol. Exp. Ther. 94, 249-261, 1948; cited in: Toren, K.: Arbete och Haelsa 25, 1-17, 1994

10. SUMMARY AND EVALUATION

<u>10.1 End Point Summary</u>

10.2 Hazard Summary

10.3 Risk Assessment