AR201-13451B

02 JAN - 3 AM 10: 54

IUCLID

Data Set

Existing Chemical CAS No. Generic name Synonym Product name	: ID: 1918-00-9 : 1918-00-9 : 2-methoxy-3,6-dichlorobenzoic acid : 3,6-dichloro-o-anisic acid : dicamba
Producer Related Part Company Creation date	: Toxicology and Regulatory Affairs : 25.12.2001
Substance Related Part Company Creation date	: Toxicology and Regulatory Affairs : 25.12.2001
Memo	:
Printing date Revision date Date of last Update	: 27.12.2001 : : 27.12.2001
Number of Pages	: 39
Chapter (profile) Reliability (profile) Flags (profile)	 Chapter: 1, 2, 3, 4, 5, 7 Reliability: without reliability, 1, 2, 3, 4 Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. General Information	1918-00-9 27.12.2001
1.0.1 OECD AND COMPANY INFORMATION	
1.0.2 LOCATION OF PRODUCTION SITE	
1.0.3 IDENTITY OF RECIPIENTS	
1.1 GENERAL SUBSTANCE INFORMATION	
1.1.0 DETAILS ON TEMPLATE	
1.1.1 SPECTRA	
1.2 SYNONYMS	
Source: Notox Hertogenbosch19.03.2001DicambaSource19.03.2001	
1.3 IMPURITIES	
1.4 ADDITIVES	
1.5 QUANTITY	
1.6.1 LABELLING	
1.6.2 CLASSIFICATION	
1.7 USE PATTERN	
1.7.1 TECHNOLOGY PRODUCTION/USE	
1.8 OCCUPATIONAL EXPOSURE LIMIT VALUES	
2 / 39	

1. General Information	ld 1918-00-9 Date 27.12.2001
1.9 SOURCE OF EXPOSURE	
1.10.1 RECOMMENDATIONS/PRECAUTIONARY MEASUR	ES
1.10.2 EMERGENCY MEASURES	
1.11 PACKAGING	
1.12 POSSIB. OF RENDERING SUBST. HARMLESS	
1.13 STATEMENTS CONCERNING WASTE	
1.14.1 WATER POLLUTION	
1.14.2 MAJOR ACCIDENT HAZARDS	
1.14.3 AIR POLLUTION	
1.15 ADDITIONAL REMARKS	
1.16 LAST LITERATURE SEARCH	
1.17 REVIEWS	
1.18 LISTINGS E.G. CHEMICAL INVENTORIES	

2.1 MELTING POINT

Value Sublimation Method Year GLP	:	87 - 108 ° C OECD Guide-line 102 "Melting Point/Melting Range" 1981 yes	
Test substance	:		
Method	:	Test was performed according to OECD 102, capillary method - metal block apparatus.	
		Two capillary tubes containing finely ground test substance were tested simultaneously (determination 1 and 2). Melting point of acetanilide was measured to determine the accuracy of the apparatus before the actual test.	
Result	:	determination 1 determination 2 beginning of 87 87 melting (deg C)	
		final stage of 108 108 melting	
Test substance	:	I, CAS 1918-00-9 (dicamba, technical), purity 85.9% (by HPLC)	
Conclusion Reliability	:	melting range is 87-108 deg C (1) valid without restriction No results for the reference substance are given. However, accuracy was estimated to be 0.5 deg C which is by far	
Flag 25.12.2001	:	exceeded by the length of the temperature range. Critical study for SIDS endpoint	(12)
2.2 BOILING POINT			
2.3 DENSITY			
2.3.1 GRANULOMETRY			
2.4 VAPOUR PRESSUR			
2.4 VAFOUR FRESSUR			
Value Decomposition Method	:	.0000167 hPa at 25° C ambiguous other (measured): US EPA Pesticide Assessment Guidelines (40 CFR 158), Subdivision D, No 63-9. Essentially OECD 104, gas saturation method.	
Year GLP Test substance Decomposition	:	yes other TS Ambiguous	

. Physico-Chem	Id 1918-00-9 Date 27.12.2001
Method	 VP was determined at 8 different temperatures between 95 and 111 deg C using a Dupont 916 Thermal Evolution Analyzer. Using this apparatus, test substance saturation in a carrier gas is achieved at a certain temperature. The gas chamber effluent is swept to an on-line coupled Flame lonization Detector, the response of which is proportional to the number of moles of TS reaching the detector per unit of time. TS (0.1061 g) was loaded on sea sand (0.9373 g). Nitrogen was used as carrier gas; VP was determined at 3 flow rates (0.680, 1.858 and 3.893 mL/min) for each temperature. Validity of the method was determined using dimethylphthalate as a reference substance. VP at 25 deg C was determined by extrapolation of a log VP vs. 1000/T line.
Remark	
Remark	 The vapor pressure is supported by the EPIWIN v3.05 calculated value of 0.0000075 hPa.
Result	: Temperature Average empirical VP (deg C) (mm Hg)
	95 0.1080
	97 0.1281
	99 0.1500
	100 0.1796 104 0.2558
	106 0.3209
	110 0.4512 111 0.5471
Test substance Conclusion Reliability Flag	 Log VP = -6145.6/T (K) + 15.7189 (mm Hg) with T(K) = t(deg C) + 273 (correlation coefficient = -0.9980) I, CAS 1918-00-9 (dicamba), purity 99.18% (HPLC) VP at 25 deg C = 1.25E-5 mm Hg (1.67E-5 hPa) (2) valid with restrictions Extrapolation from 95 deg C as lowest T to 25 deg C may cause a relative error since, at 95 deg C TS may be partially fluid, whereas at 25 deg C it is a solid. Extrapolation may therefore be problemetic. It is, however, the best possible option under these circumstances. Critical study for SIDS endpoint
25.12.2001	(12) (14
.5 PARTITION CO	EFFICIENT
Log pow	: = 2.21 at ° C
Test substance	: CAS 1918-00-9 (dicamba)
Reliability	 (2) valid with restrictions Score of 2 given to handbook or published values for physical constants. The measured value in the other listed study is for the partially ionized form of the TS.
Flag	: Critical study for SIDS endpoint
25.12.2001	(4
Log pow	: .545 at 25° C
Method	other (measured): EPA Pesticide Assessment Guidelines, Subdivision D, Product Chemistry, Section 63-11. Essentially OECD 107
Year	: 1982
GLP Test substance	: yes : other TS

phase, measu phases 0.497 r 1.28E6 dissolv 5 mL n were s centrifu were ta each p	Kow of non- red test subs and on pKa ng and 5.05 dpm/mg an ed in 5 mL b -octanol-pre haken in a w uged (2000 r sken from bo	tance dissociates in aqueous and octanol -dissociated TS was calculated on basis of stance concentrations and pH of the two a of the test substance (1.94). 4 mg test substance (specific activities and 1.26E5 dpm/mg, respectively) were each puffer-presaturated n-octanol after which saturated buffer was added. The mixtures vater bath at 25 deg C for 1 hour, rpm, 20 min) and duplicate 1.0 mL aliquots	
1.28E6 dissolv 5 mL n were sl centrifu were ta each p	dpm/mg an ed in 5 mL b -octanol-pre haken in a w uged (2000 r aken from bo	d 1.26E5 dpm/mg, respectively) were each ouffer-presaturated n-octanol after which saturated buffer was added. The mixtures vater bath at 25 deg C for 1 hour,	
each p were p The fra	H and each repared. Iction of und	oth phases and analyzed by LSC. The pH of easured. ons of pH 5.0, 7.0 and 9.0 were used. For TS concentration triplicate test mixtures issociated dicamba in each phase was	
	concentratio	on (mean of 3 replicates)	
5.0 7.0 9.0	4.58 4.58 4.58	6.86 +/- 0.60 0.54 +/- 0.01 8.95 +/- 0.06	
5.0 7.0 9.0	0.499 0.499 0.499	3.98 +/- 0.11 0.16 +/- 0.00 0.58 +/- 0.00	
Averag	e Kow: 3.51	+/- 3.73	
: I, CAS	1918-00-9 (dicamba), analytical reference standard	
: (2) vali 1. Mea results should at low p Howev low. Th methoo 2. Only	d with restrict surement was in deviations have been p bH. OECD 1 er, as pKa = lerefore, this d.	ctions as performed on ionized form of TS, which s from the partition law. Measurement performed on non-ionized TS and therefore 07 suggests pH at least one unit below pKa. : 1.94 pH should have been < 1 which is very s has to be considered best possible	
			(15
BILITY			
: soluble : at 25 ° : at a : other:e	(1000-1000 C and °C	00 mg/L)	
	 were p The fra calcula pH. Buffer p 5.0 7.0 9.0 5.0 7.0 9.0 5.0 7.0 9.0 Average Notox I, CAS Kow of substat Kow ra (2) valit 1. Mea results should at low p Howev low. Th method 2. Only and co 	 were prepared. The fraction of und calculated on basis pH. Buffer pH Initial TS concentration in n-octanol 5.0 4.58 7.0 4.58 9.0 4.58 5.0 0.499 7.0 0.499 9.0 0.499 Average Kow: 3.51 Notox Hertogenboo I, CAS 1918-00-9 (I, CAS 1918-00-9 (I	The fraction of undissociated dicamba in each phase was calculated on basis of measured ion concentration, pKa and pH. : Buffer pH Initial TS Kow concentration (mean of 3 replicates) in n-octanol (mM) 5.0 4.58 6.86 +/- 0.60 7.0 4.58 0.54 +/- 0.01 9.0 4.58 8.95 +/- 0.06 5.0 0.499 3.98 +/- 0.11 7.0 0.499 0.16 +/- 0.00 9.0 0.499 0.58 +/- 0.00 Average Kow: 3.51 +/- 3.73 : Notox Hertogenbosch : I, CAS 1918-00-9 (dicamba), analytical reference standard I, CAS 1918-00-9 (dicamba), radiochemical purity 98% : Kow of test substance strongly depends on pH and on test substance concentration. Kow ranged between 0.2 and 9.0. : (2) valid with restrictions 1. Measurement was performed on ionized form of TS, which results in deviations from the partition law. Measurement should have been performed on non-ionized TS and therefore at low pH. OECD 107 suggests pH at least one unit below pKa. However, as pKa = 1.94 pH should have been < 1 which is very low. Therefore, this has to be considered best possible method. 2. Only one n-octanol: water ratio was tested for each pH and concentration. SULITY : 8.24 g/l at 25 ° C : soluble (1000-10000 mg/L) : at 25° C : at and ° C : other: essentially OECD 105 (flask method)

2. Physico-Chem		ld 1918-00-9 te 27.12.2001
GLP Test substance	: yes : other TS	
Method	 25 mL water of Milli-Q reagent grade were added to 0 test substance. The mixture was shaken for about on and was then placed in a water bath (25 deg C) for at 48 hrs. With intervals of at least 24 h the mixture was centrifuged and returned to a waterbath (25 deg C) for temperature equilibration (at least 1 h). The test solut were analyzed in duplicate using HPLC against dicar calibration standards (dicamba in methanol, 1.028-10 mg/mL). Measurements were repeated until SD of the measurements was within the method reproducibility. 	e hour t least or ions nba 0.285 e two last
Remark	: This value is supported by a vlaue of 6500 mg/L at 25 C.D.S. (ed.). The Pesticide Manual - World Compend UK: The British Crop Protection Council, 1994. 298 (a Substance Data Base)	lium. 10th ed. Surrey,
Result Source Test substance Conclusion Reliability	 Solubility in water at 25 deg C: 0.824 g per 100 mL solution Notox Hertogenbosch I, CAS 1918-00-9 (dicamba, technical), purity 85.9% Solubility of test substance in water is 8.24 g/L. (2) valid with restrictions 	
Flag 25.12.2001	 Only the end result is reported, no individual result measurements are given. Results can therefore not b checked. Method is intended for essentially pure chemicals. Dicamba technical cannot be regarded as such. It should be noted that whereas technical dicamba tested, a reference standard of 99.18% purity was us calibration. Impurities have therefore been disregarded Critical study for SIDS endpoint 	was ed for
2.6.2 SURFACE TENS	ION	
2.7 FLASH POINT		
2.8 AUTO FLAMMA	BILITY	
2.9 FLAMMABILITY		
2.10 EXPLOSIVE PR	PERTIES	
2.11 OXIDIZING PRO	PERTIES	
2.12 ADDITIONAL RE	MARKS	

3.1.1 PHOTODEGRADATION

Type Light source Light spect. Rel. intensity Conc. of subst. Direct photolysis Halflife t1/2 Degradation Quantum yield Deg. Product Method Year GLP Test substance	 water Xenon lamp > 290 nm 1.32 based on Intensity of Sunlight 100.19 mg/l at 25 degree C 50.3 day 31.3 % after 30 day yes EPA Guide-line subdivision N 161-2 "Photodegradation studies in water" 1982 yes other TS
Method	 A 1000 mL test solution consisting of 100.19 mg dicamba with a specific activity of 412.2 dpm/ug (total 688 kBq) in aqueous buffer solution pH 7 containing 1% acetonitrile was prepared. The test solution was incubated at 25 +/- 1 deg C under contineous stirring for 30 days. Average incident radiation on the reactor surface was 7.704E2 W/m2 (measured before and after the study). The reaction solution was aerated and connected to a silica gel trap, an ethylene glycol trap (organic volatiles) and a 10% NaOH trap (supposed to collect CO2) in series. Before initiation of photolysis, a 50 mL sample was taken as dark control sample. 20 mL samples were taken before initiation of photolysis and on day 1, 3, 8, 15, 22 and 30. The samples were analyzed as follows: duplicate 1 mL samples were analyzed by LSC 15 mL was extracted twice at pH < 1 with ethyl acetate, both fractions were analyzed by LSC (duplicate 1 mL samples) ethyl acetate fraction was dried and concentrated, and analyzed by TLC using 4 solvent systems (cochromatographed with reference standards) extracted buffer solution of day 15, 22 and 30 were lyophilized followed by acetonitrile extraction; the extract was concentrated and analyzed by TLC using 4 solvent systems (cochromatographed with reference standards) duplicate 1 mL ethylene glycol and 10% NaOH trap samples were analyzed by LSC; residual radioactivity in the silica traps was determined by combustion identity of radioactivity supposed to be CO2 in 10% NaOH trap samples was confirmed for day 22 and 30 by precipitation as BaCO3 and subsequent evolution as CO2 after addition of HCI On day 30, the reactor was washed with methanol and with acetone. Volumes were measured and 1 mL duplicatealiquots were analyzed by LSC. Photodegradation was calculated using the SAS Regression Prog

Environmental	Fate and Pathways Id 1918-00- Date 27.12.200	
Result	: time point (days) 14C-dicamba (% of actually applied	
neoun	14C-dicamba (70 of doctary applied 14C-dicamba)*	
	0 100 (92.14% of applied 14C)	
	1 98.83	
	3 95.25	
	8 86.87	
	15 75.62	
	22 66.44	
	30 58.74 (degradation: 41.26%)	
	30 (dark control) 98.61	
	* calculated by reviewer from % of applied 14C Unchanged dicamba was confirmed by HPLC.	
	onchanged dicamba was confinitied by HFLC.	
	All other compounds in the different fractions, separated by	
	TLC, were <10% of applied 14C and did not match with	
	reference standards. CO2 in the 10% NaOH trap was 11.7% of	
	applied at day 22 and 16.6% of applied 14C at day 30. Radioactivity in the other traps was <10% of applied 14C at	
	all time points. Reactor wash yielded 0.3% of applied	
	activity. The mass balance was >99% and <103.5% at all time	
	points.	
	Under these conditions, the of disample was 20.1 down the	
	Under these conditions, t1/2 of dicamba was 38.1 days; the photolysis rate constant was 0.018 day-1. Based on the	
	spring sunlight intensity at 40 deg latitude at noon (5.83E2	
	W/m2) the corresponding photodegradation rate for natural	
	sunlight will be 0.0138 day-1; t1/2 will be 50.3 days.	
Test substance	: I, CAS 1918-00-9 (dicamba), purity 99.6% by IR	
Test substance	I, (14C-dicamba), radiochemical purity 100% by TLC	
Conclusion	: The photodegradation rate constant in spring sunlight at 40	
	deg latitude at noon is 0.0138 day-1; t1/2 is 50.3 days. The	
	major photodegradation product is CO2.	
Reliability	: (1) valid without restriction	
	1. In the calculation of $t1/2$, no correction for the	
	degradation in the dark control was made. However, this will	
	only slightly influence the results, as there was hardly any	
	degradation in the dark control.	
	Except for sterilization of the buffer solution, no	
	measures to guarantee sterility of the samples were	
	described. However, as there was hardly any degradation in	
	the dark control (which was a subsample of the sample to be	
	irradiated), it can be assumed biodegradation was negligible.	
Flag 25.12.2001	: Critical study for SIDS endpoint	(1
		()
Type	: air	
Light source	: Sun light	
Light spect. Rel. intensity	: nm : based on Intensity of Sunlight	
Indirect photolysis	. based on mensity of ournight	
Sensitizer	: ОН	
Conc. of sens.	: 1500000 molecule/cm3	
Rate constant	: = .0000000002985 cm3/(molecule*sec)	
Degradation	: = % after 43 hour(s)	
Deg. Product	:	
Method	: 2001	
Year	: 2001	
	9 / 39	

. Environmenta)18-00-9 7.12.2001
GLP Test substance Reliability Flag 25.12.2001	: no : : (2) valid with restrictions : Critical study for SIDS endpoint	
.1.2 STABILITY IN W	ATER	
Type t1/2 pH4 t1/2 pH7 t1/2 pH9 Degradation Deg. Product	 abiotic at degree C at degree C at degree C at degree C = 0 - 7.6 % after 30 day at pH and degree C 	
Method Year GLP	 other: essentially OECD 111 1981 no 	
Test substance Method	 Solutions of 10 ppm and 100 ppm dicamba (1.17% and 0. 14C-dicamba, respectively) in distilled water or aqueous buffer solutions of pH 5.0, 7.0 and 9.0 were incubated at 2 and 35 deg C for 30 days (volume 201 mL, in amber bottle shaking water baths). Acetone concentrations were 0.5%. After 1, 7, 14, 21 and 30 days, a duplicate 1-mL sample was taken for radioassay and a duplicate 15-mL sample was ta for extraction using diethyl ether (at pH < 1). Organic and aqueous layers were first radioassayed and then analyzed using TLC and radioautography detection, followed by quantification using LSC. Samples were cochromatograph with dicamba and three metabolite reference standards. 	5 s in as ken
Result	 There was no significant dicamba hydrolysis (i.e. equal to or less than 7.6%) at each pH value, both concentrations a both temperatures, except for 100 ppm, pH 7.0, 35 deg C t=14, 21 and 30 days in the 100 ppm, when degradation w to 18.5%. Total recovery was only 82.5-83.4% for these samples, whereas it was > 95 for all other samples. Radioactivity remaining in the aqueous phase after extraction was equal to or less than 1% of applied. Three unknown degradation products each constituted less than of applied. 	at as up
Source Test substance	 Notox Hertogenbosch I, CAS 1918-00-9 (14C-dicamba), purity not specified I, CAS 1918-00-9 (14C-dicamba), radiochemical purity gre than 98% 	ater
Conclusion	: Dicamba is stable with slight or no hydrolysis over 30 days under the conditions tested.	
Reliability	 (2) valid with restrictions 1. The fact that at 100 ppm, pH 7.0, 35 deg C up to 18.5% degradation occurred was disregarded because recoveries low. However, no explanation was given for the low recoveries. It cannot be excluded that loss of radioactivity is due to hydrolysis. 2. Section "Results and discussion" contained 2 values that were not in agreement with values in tables of results. 3. No measures to guarantee sterility of the samples or to exclude oxygen from the solutions were described. However 	t

8. Environmental	Fate and Pathways	ld 1918-00-9 Date 27.12.2001
	as measured degradation percentag 100 ppm, pH 7.0, 35 deg C), no sigr degradation or oxidation can have o 2. No duplicate samples at any pH. 3. pH 5.0 was tested, whereas OEC	nificant biotic ccurred.
25.12.2001	5. pri 5.0 was tested, whereas OLC	(19)
3.1.3 STABILITY IN SC	IL	
3.2 MONITORING DA	TA	
3.3.1 TRANSPORT BE	TWEEN ENVIRONMENTAL COMPARTME	INTS
Туре	: fugacity model level III	
Media	: other	
Air (level I)	:	
Water (level I) Soil (level I)		
Biota (level II / III)		
Soil (level II / III)	:	
Method Year	: 2001	
Remark	EPIWIN 3.05. Measured values we	urrent best estimate (from HSDB). Half POWIN program. Direct photolysis
Result	: Level III Fugacity Model (Full-Out	tput):
	Chem Name : Dicamba Molecular Wt: 221.04 Henry's LC : 2.18e-009 atm-m3/m Vapor Press : 1.26e-005 mm Hg Liquid VP : 6.95e-005 mm Hg Melting Pt : 100 deg C (user-er Log Kow : 2.21 (user-entere Soil Koc : 66.5 (calc by mod	(user-entered) (super-cooled) htered) ed)
	Concentration Half-Lif (percent) (hr) Air 0.0498 43 Water 29.9 500 Soil 70 500 Sediment 0.122 2e+06	(kg/hr) 1000 1000 1000
	Fugacity Reaction A (atm) (kg/hr) Air 9.61e-013 14.2 Water 2.6e-014 732 Soil 3.58e-013 1.72e+003 Sediment 2.06e-014 0.75	AdvectionReactionAdvection(kg/hr)(percent)(percent)8.80.4730.29352824.417.6057.200.04330.0250.00144
	Persistence Time: 590 hr Reaction Time: 718 hr Advection Time: 3.29e+003 hr Percent Reacted: 82.1 Percent Advected: 17.9	
	Half-Lives (hr), (based upon us Air: 43 Water: 500 Soil: 500	ser-entry):

3. Environmental	Fate and Pathways	ld 1918-00-9 Date 27.12.2001
	Sediment: 2000	
	Advection Times (hr): Air: 100 Water: 1000 Sediment: 5e+004	
Test substance Reliability Flag 25.12.2001	 CAS 1918-00-9 (dicamba) (2) valid with restrictions Critical study for SIDS endpoint 	(3
3.3.2 DISTRIBUTION		
3.4 MODE OF DEGR	ADATION IN ACTUAL USE	
3.4 MODE OF DEGR	ADATION IN ACTUAL USE	
3.5 BIODEGRADATI	ON	
Inoculum Remark Test substance	levels of 2,3-dihydroxy-3,6-dichlorosa under anaerobic conditions is similar except the rate of dicamba metabolis conditions. [Krueger JP et al; J Agric cited in HSDB update of 8-09-2001. AQUATIC FATE: Based on the result degradation appears to be the import natural water. Photolysis may contrib water(Scifres CJ et al; J Environ Qua update of 8-09-2001.	gricultural soil under aerobic nineralized to CO2 under aerobic icid as the only major metabolite. Low alicylic acid were detected. Metabolisn to that which occurred in aerobic soil is reduced under anaerobic Food Chem 39: 995-9 (1991)]. As ts of various studies, microbial tant dicamba removal process in bute to dicamba removal from
Test substance Conclusion Flag 25.12.2001	 CAS 1918-00-9 (dicamba) Dicamba biodegrades under both aer not know if it can be considered read criteria. Critical study for SIDS endpoint 	
20.12.2001		
3.6 BOD5, COD OR	BOD5/COD RATIO	
3.7 BIOACCUMULA	TION	
3.8 ADDITIONAL RE	MARKS	

4. Ecotoxicity

4.1 ACUTE/PROLONGED TOXICITY TO FISH

Type Species Exposure period Unit Analytical monitoring LC50 Method Year GLP Test substance Method		static Cyprinodon variegatus (Fish, estuary, marine) 96 hour(s) mg/l no > 180 other: EPA-660/3-75-00 1975 no other TS TEST ORGANISMS - Species: Cyprinodon variegatus - Supplier: commercial supplier in Florida - Size (mean)/weight (mean)/loading: 32 mm/480 mg/0.32 g/L - Feeding (pretreatment): disontinued 48 hours prior to test - Feeding (pretreatment): disontinued 48 hours prior to test - Feeding during test: none STOCK AND TEST SOLUTION AND THEIR PREPARATION - Vehicle, solvent: acetone - Concentration of vehicle/ solvent: 0.06-0.6 mL/L DILUTION WATER - Source: artificial seawater (origin well water) - Chemistry (Salinity;pH): 27 ppt; 8.18 TEST SYSTEM - Test type: static - Concentrations: 18, 32, 56, 100 and 180 mg/L, solvent treated and untreated controls - Exposure vessel type: 20 L glass vessel containing 15 L water - Number of fish: 10/treatment - Photoperiod: not indicated PHYSICAL MEASUREMENTS - Measuring times: 0, 48 (only O2), 96 h in controls, 18, 56 and 180 mg/L - Dis. oxygen: 101-104% (0 h), 74-83% (48 h), 51-78% (96 h) - pH: 7.5-8.2, for 180 mg/L 6.6-7.4 - Test temperature: 21 C DURATION OF THE TEST: 96 hours TEST PARAMETER: Mortality OBSERVATION TIMES: 24, 48 and 96 hours
Result Source Test substance Reliability	:	STATISTICAL METHOD: not applicable RESULTS: - Mortality: no mortality - Other effects: not reported Notox Hertogenbosch I, CAS 1918-00-9 (dicamba technical), purity 86.82% (2) valid with restrictions Since there is no specific guideline for saltwater fish, the test performance was checked with EPA OPPTS 850.1075 (1996): A) No analyses were performed to confirm the nominal test concentrations (EPA >80% of nominal) B) The dissolved oxygen concentration was lower than
		40.400

	ld 1918-00-9 Date 27.12.2001
	recommended in some test vessels at the end of the test only (51-78% at 96 hours, EPA >60%); the salinity was higher than recommended (27 ppt, EPA 20 +/- 5 ppt); vehicle concentration was higher than recommended in the highest tested concentration only (0.6 mL/L, EPA 0.5 mL/L); pH-values in the highest tested concentration only were lower than recommended (6.6-7.4, EPA 7.5-8.5), due to inherent properties of the test substance; the photoperiod
28.03.2001	was not indicated (EPA 12-16 h light). (20
2 ACUTE TOXICITY 1	TO AQUATIC INVERTEBRATES
Туре	: static
Species	: Daphnia magna (Crustacea)
Exposure period	: 48 hour(s)
Unit	: mg/l
Analytical monitoring EC50	: : m > 100
Method	: 11 > 100
Year	: 1980
GLP	: no data
Test substance	: The study of stated is the UODD as and for discussion of fully one
Method	: The study was reported in the HSDB record for dicamba as follows:
Test substance Reliability Flag 27.12.2001	 EC50 Daphnia magna greater than 100 mg/l/48 hr @ 21 deg c, first instar /technical material, 88%/. effect: immobilization. static bioassay without aeration, ph 7.2-7.5, water hardness 40-50 mg/l as calcium carbonate and alkalinity of 30-35 mg/l. CAS 1918-00-9 (dicamba, technical), purity 88% (2) valid with restrictions Critical study for SIDS endpoint (17)
27.12.2001	
3 TOXICITY TO AQU	ATIC PLANTS E.G. ALGAE
3 TOXICITY TO AQU Species Endpoint	 ATIC PLANTS E.G. ALGAE Selenastrum capricornutum (Algae) other: biomass/growth rate
Species Endpoint Exposure period	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s)
Species Endpoint Exposure period Unit	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l
Species Endpoint Exposure period Unit Analytical monitoring	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l yes
Species Endpoint Exposure period Unit Analytical monitoring NOEC	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l yes 3.7
Species Endpoint Exposure period Unit Analytical monitoring	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l yes
Species Endpoint Exposure period Unit Analytical monitoring NOEC EC0 EC10 EC50	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l yes 3.7 3.7 > 3.7 > 3.7 > 3.7 > 3.7
Species Endpoint Exposure period Unit Analytical monitoring NOEC EC0 EC10 EC50 Method	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l yes 3.7 3.7 > 3.7 > 3.7 > 3.7 > 3.7 > 1.7 > 1.7 > 2.7 > 3.7 > 3.7 > 3.7 > 3.7 > 3.7
Species Endpoint Exposure period Unit Analytical monitoring NOEC EC0 EC10 EC50 Method Year	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l yes 3.7 3.7 > 3.7 > 3.7 > 3.7 > 120 hour(s)
Species Endpoint Exposure period Unit Analytical monitoring NOEC EC0 EC10 EC50 Method	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l yes 3.7 3.7 > 3.7 > 3.7 > 3.7 > 3.7 > 1.2 > 3.7 > 3.7 > 3.7
Species Endpoint Exposure period Unit Analytical monitoring NOEC EC0 EC10 EC50 Method Year GLP	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l yes 3.7 3.7 > 3.7 > 3.7 > other: EPA 122-2, 123-2 1982 yes other TS TEST ORGANISMS
Species Endpoint Exposure period Unit Analytical monitoring NOEC EC0 EC10 EC50 Method Year GLP Test substance	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l yes 3.7 3.7 > 3.7 > 3.7 > other: EPA 122-2, 123-2 1982 yes other TS TEST ORGANISMS Species: Selenastrum capricornutum, strain 1648, family
Species Endpoint Exposure period Unit Analytical monitoring NOEC EC0 EC10 EC50 Method Year GLP Test substance	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l yes 3.7 3.7 > 3.7 > 3.7 other: EPA 122-2, 123-2 1982 yes other TS TEST ORGANISMS Species: Selenastrum capricornutum, strain 1648, family Chlorophyceae
Species Endpoint Exposure period Unit Analytical monitoring NOEC EC0 EC10 EC50 Method Year GLP Test substance	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l yes 3.7 3.7 > 3.7 > 3.7 other: EPA 122-2, 123-2 1982 yes other TS TEST ORGANISMS Species: Selenastrum capricornutum, strain 1648, family Chlorophyceae Source/supplier: Carolina Biological Supply Company,
Species Endpoint Exposure period Unit Analytical monitoring NOEC EC0 EC10 EC50 Method Year GLP Test substance	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l yes 3.7 3.7 > 3.7 > 3.7 > other: EPA 122-2, 123-2 1982 yes other TS TEST ORGANISMS Species: Selenastrum capricornutum, strain 1648, family Chlorophyceae
Species Endpoint Exposure period Unit Analytical monitoring NOEC EC0 EC10 EC50 Method Year GLP Test substance	 Selenastrum capricornutum (Algae) other: biomass/growth rate 120 hour(s) mg/l yes 3.7 3.7 > 3.7 > 3.7 other: EPA 122-2, 123-2 1982 yes other TS TEST ORGANISMS Species: Selenastrum capricornutum, strain 1648, family Chlorophyceae Source/supplier: Carolina Biological Supply Company, Burlington, North Carolina

4. Ecotoxicity	Id 1918-00-9
_	Date 27.12.2001
	fresh medium ~twice weekly. - Pretreatment: at least 2 days prior to test initiation algae were maintained under test conditions (culture medium, 100 rpm, 25 C, continuous illumination (3200-4300 lux) - Initial cell concentration: 0.3 E4 cells/mL
	STOCK AND TEST SOLUTION AND THEIR PREPARATION - Vehicle, solvent: none
	GROWTH/TEST MEDIUM CHEMISTRY - Chemistry (Hardness (Mg+Ca) 0.4 mmol/L;TOC 2.1 mg/L;P 1.6 mg/L;N 14 mg/L;EDTA 12E-2 mmol/L) - pH: 7.5 (after adjustment)
	TEST SYSTEM - Test type: static - Concentrations: 4 mg a.i./L and controls - Exposure vessel: 125 mL erlenmeyer flasks containing 50 mL of test medium (shaken at 100 rpm) - Number of replicates: 3 - Photoperiod (intensity of irradiation): continuous (3200-4800 lux) PHYSICAL MEASUREMENTS - Measuring times: 0 and 120 h - Test temperature: 25 C - pH: 7.3-7.5 (0 h); 10.4 (120 h)
	DURATION OF TEST: 120 hours
	TEST PARAMETER: algal growth (cell counts), measured by a haemacytometer OBSERVATION TIMES: 0, 24, 48, 72, 96, 120 h ANALYSES:
	- Method: direct HPLC-UV - Sampling times: 0 and 120 h
Result	 STATISTICAL METHOD: t-test RESULTS: Nominal concentrations (mg a.i./L): 0, 4 Measured concentrations (mg a.i./L): <loq, (="93%" 3.7="" li="" nominal)<="" of=""> Cell density data after 0, 24, 48, 72, 96 and 120 h (x E4 cells/mL) : 0: 0.3, 3, 18, 39, 54, 258 4: 0.3, 3, 17, 44, 51, 260 Growth rate/ biomass(AUC) (% of control): 100/99 </loq,>
	GROWTH FACTOR CONTROL: 130 after 72 hours
	ANALYTICAL RESULTS: validated at 0.025-2.5 mg/L (recovery 101+/-2%, LOQ 14 ug/L. QCs fortified at 4 mg/L showed a recovery of 83-119%.
Source Test substance Reliability	 STATISTICAL RESULTS: no significant differences between control and treatments Notox Hertogenbosch I, CAS 1918-00-9 (Dicamba technical), purity 89.5% (1) valid without restriction Minor remark. The test medium was not in accordance with OECD 201. The pH-increase observed during the test was probably associated with the strong cell growth (factor 130 15 / 39

4. Ecc	otoxicity	1918-00-9 27.12.2001	
28.03	after 72 hours). 3.2001		(9)
4.4	TOXICITY TO MICROORGANISMS E.G. BACTERIA		
4.5.1	CHRONIC TOXICITY TO FISH		
4.5.2	CHRONIC TOXICITY TO AQUATIC INVERTEBRATES		
4.6.1	TOXICITY TO SOIL DWELLING ORGANISMS		
4.6.2	TOXICITY TO TERRESTRIAL PLANTS		
4.6.3	TOXICITY TO OTHER NON-MAMM. TERRESTRIAL SPECIES		
4.7	BIOLOGICAL EFFECTS MONITORING		
4.8	BIOTRANSFORMATION AND KINETICS		
4.9	ADDITIONAL REMARKS		

5. Toxicity

5.1.1 ACUTE ORAL TOXICITY

Type Species Strain Sex Number of animals Vehicle Value Method Year GLP Test substance Method	 other: corn oil = 1465 mg/kg bw other: not specified no other TS TEST ORGANISMS: Source: not specified Age: not specified Number: 5/sex/dose Weight at study initiation: 200-248 g Controls: no ADMINISTRATION:
	 Doses: 500, 794, 1250, 1984, 3150 and 5000 mg/kg bw Doses per time period: single Volume administered: 10 ml/kg bw for all dosage levels except for the 5000 mg/kg level where 20 ml/kg bw was administered. Post dose observation period: 14 days food was withheld overnight EXAMINATIONS: for mortality (at least daily).
Result	 BODY WEIGHT: at dosing and at 14 days. STATISTICAL METHOD: Thompson (1947) MORTALITY: Number of deaths at each dose: 500, 794, 1250, 1984, 3150, 5000 mg/kg bw 0/10, 1/10, 4/10, 4/10, 10/10, 10/10 Time of death: within 48 hours after dosing
	CLINICAL SIGNS: no data on decendents BODY WEIGHT: all surviving rats exhibited normal body weight gains during the observation period NECROPSY FINDINGS: no data POTENTIAL TARGET ORGANS: no data
Source Test substance Conclusion Reliability	 SEX-SPECIFIC DIFFERENCES: LD50 males= 1879 mg/kg bw LD50 females= 1581 mg/kg bw Notox Hertogenbosch I, CAS 1918-00-9 (Dicamba 85.8%), purity 85.8% LD50 1707 mg/kg bw = 1465 mg a.i./kg bw (2) valid with restrictions The information was essentially confined to what is included in the current summary. no data were presented for effects other than mortality.

Toxicity	ld 1918-00-9 Date 27.12.2001	
	3. The dose volume used at the 5000 mg/kg bw was higher than recommended (20 ml/kg, OECD 401 =< 10 ml/kg). Since at 3150 mg/kg all rats died already, the reliability is not lowered because of this.	
04.04.2001		(18
1.2 ACUTE INHALATIO	ON TOXICITY	
Туре	: LC50	
Species Strain	: rat . other: Spartan	
Sex	: other: Spartan : male/female	
Number of animals	: 10	
Vehicle	: other: no vehicle	
Exposure time	: 4 hour(s)	
Value	: > 8.2 mg/l	
Method Year	other: not specified	
GLP	: no	
Test substance	: other TS	
Method	: TEST ORGANISMS:	
	- Source: not specified	
	- Age: not specified	
	- Weight at study initiation: 206-245 g - Number of animals: 5/sex/dose	
	- Number of animals. 5/sex/dose - Controls: no	
	ADMINISTRATION:	
	- Type of exposure: whole body exposure to dust of test	
	material	
	- Exposure duration: 4 hours	
	- Concentrations(nominal/measured): approx. nominal conc. of	
	9.6 mg/l or 8.2 mg a.i./l - Particle size: not specified	
	- Type or preparation of particles: control by Wright Dust	
	Feeder	
	- Air changes: no data	
	EXAMINATIONS: during exposure: changes in behavior and	
	appearance, after exposure: pharmacodynamic and/or toxic	
	signs; 14 days observation period	
	BODY WEIGHTS: not specified	
	ANALYSES:	
	- Method: no data	
	- Sampling times: no data	
	STATISTICAL METHOD: no data	
Result	: MORTALITY:	
	- Number of deaths at each dose: no deaths	
	CLINICAL SIGNS: during exposure: increased, then decreased	
	motor activity, and nasal porphyrin discharge. 14 day	
	observation period decreased motor activity (1/10), corneal	
	opacity (few rats).	
	BODY WEIGHTS: gains were normal during the study.	
	NECROPSY FINDINGS: no data	
	18 / 39	

Toxicity	ld 1918-00-9 Date 27.12.2001
	POTENTIAL TARGET ORGANS: no data
Source Test condition Conclusion Reliability	 SEX-SPECIFIC DIFFERENCES: no data Notox Hertogenbosch I, CAS 1918-00-9 (Dicamba 85.8%), purity 85.8% LC50 > 9.6 mg/l = > 8.2 mg a.i./l (2) valid with restrictions The information was essentially confined to what is included in the current summary As this is a limit test, the LC50 value was derived by the reviewer.
04.04.2001	3. no individual data were present. (18
1.3 ACUTE DERMAL	ΤΟΧΙΟΙΤΥ
Type Species Strain Sex Number of animals Vehicle Value Method Year GLP Test substance Method	 LD50 rabbit New Zealand white male/female 4 other: not specified > 1716 mg/kg bw other: not specified no other TS TEST ORGANISMS: Source: not specified Age: not specified Age: not specified Weight at study initiation: 2324-2454 g Controls: no ADMINISTRATION: Area covered: not specified Occlusion: yes Vehicle: not specified Concentration in vehicle: not specified Total volume applied: not specified Doses: 2000 mg/kg bw Removal of test substance: washed with tepid tap water after 24 hours
Result	 EXAMINATIONS: observed for mortality over 14 days. BODY WEIGHT: pre-dosing and at day 14 STATISTICAL METHOD: not specified MORTALITY: Number of deaths at each dose: no deaths CLINICAL SIGNS: not specified BODY WEIGHTS: normal gains during study period NECROPSY FINDINGS: no data POTENTIAL TARGET ORGANS: no data

5. Toxicity	ld 1918-00-9 Date 27.12.2001	
Source Test substance Conclusion Reliability	 SEX-SPECIFIC DIFFERENCES: no data Notox Hertogenbosch I, CAS 1918-00-9 (Dicamba 85.8%), purity 85.8% LD50 > 2000 mg/kg bw = > 1716 mg a.i./kg bw (4) not assignable The information was essentially confined to what is included in the current summary. As this is a limit test, the LD50 value was derived by the reviewer. Only 4 animals were used (OECD 402 5) of which 2 had an abraded skin, which could alter the permeability of the test substance. no individual data were present. 	
04.04.2001		(18)
5.1.4 ACUTE TOXICIT	Y, OTHER ROUTES	
5.2.2 EYE IRRITATION		
5.3 SENSITIZATION		
Species Sex Strain Route of admin. Exposure period Frequency of treatment Post obs. period Doses Control group NOAEL Method Year GLP Test substance Method	 rat male/female other: CD oral feed 21 weeks none 1000, 5000 and 10000 ppm yes = 342 mg/kg bw EPA OPP 82-1 1978 yes other TS TEST ORGANISMS: Species: Charles River CD rat Source: Charles River CD rat Source: Charles River Laboratories, Portage, Michigan Age: exact age was not mentioned 	
	 Weight at study initiation: male (122-164 g) female (111-145 g) Number of animals: 20/sex/dose group ADMINISTRATION / EXPOSURE Exposure period: 21 days Route of administration: diet Post exposure period: none Doses: 1000, 5000 and 10000ppm, resulting in 69.4, 342 and 682 mg/kg bw/day for males and 79.5, 392 and 751 mg/kg 	

ld 1918-00-9

Date 27.12.2001

5. Toxicity

bw/day for females

CLINICAL OBSERVATIONS AND FREQUENCY: - Mortality/clinical signs: twice daily, detailed observations weekly - Body weight: weekly Individueal food consumption: weekly CLINICAL LABORATORY TESTS In 10 rats/sex/dose group at baseline and in week 6 and 13. - Haematology: hemoglobin, hematocrit, erythrocyte count, yotal and differential leukocyte counts, platelet count, mean corpuscular volume (MCV), mean corpuscular hemoglobin (MCH), mean corpuscular hemoglobin concentrations (MCHC), and reticulocyte count. - Biochemistry: sodium, potassium, chloride, alkaline phosphatase, blood urea nitrogen (BUN), serum glutamic pyruvate transaminase (SGPT), serum glutamic oxaloacetate transaminase (SGOT), calcium, creatinine, phosphorous, lactic dehydrogenase (LDH), glucose, total bilirubin total cholesterol, albumin, globulin, total protein. - Urinalysis: specific gravity, volume, color and appearance, occult blood, protein, pH, bilirubin, urobilinogen, ketones, glucose, microscopic examination sediment, nitrites, urobilinogen, ketones. ORGANS EXAMINED AT NECROPSY (MACROSCOPIC AND MICROSCOPIC): - Organ weights: brain, heart, kidnevs, liver, gonads, - Microscopic (control animals and 10000 ppm, heart, liver, kidneys and gross lesions in all groups): all gross lesions, adrenals, eye, trachea, esophagus, stomach, duodenum, jejenum, ileum, caecum, colon, liver (2 sections), spleen, urinary bladder, testes/ ovaries, pancreas, brain (3levelsforebrain, midbrain, hindbrain), heart, lungs+mainstem bronchi, pituitary, thyroid and parathyroid, thymus, lymph node (mesenteric), sternum (bone marrow), spinal cord), salivary gland, (submaxillary), skeletal muscle (thigh), kidneys, prostate/ corpus and cervix uteri, peripheral nerve (sciatic). ANALYSES: - homogeneity of diet before study inititation - stability of test article at weeks 1,3,4,8 and 13 by GC/ECD STATISTICAL METHODS: - analyses of variance, Bartlett and t-test as described by Steel and Torrie Result CLINICAL SIGNS/MORTALITY - Mortality (dweek): 1 female control (6), 1 female 5000 ppm (2), 1 female 10000 ppm (13); Three female rats died during the course of the study. - Clinical signs: No changes were seen in general behavior and appearance; incidental findings in treated rats: rales, yellow material on the anogenital region, mouth ulcer, pale exposed skin areas, black material on or around the eye, nose, mouth or anogenital region, corneal opacity, dilated pupil, eye enlarged and protruded, increased distance between pupil and cornea, nose malaligned, swollen foot, portion of the ear

21/39

5. Toxicity	ld 1918-00-9	
	Date 27.12.2001	
	missing, and portion of the tail black or missing. These	
	signs were noted randomly among the treated rats.	
	One mid-dose male rat had a subcutaneous mass in the	
	anogenital region. Incidental findings in both treated and control rats:	
	malaligned upper incisors, red areas around the eyes,	
	scabbing, excessive lacrimation and hair loss.	
	- Body weight gain: slightly decreased at 10000 ppm in both sexes, significantly in week 13.	
	 Food consumption: at 10000 ppm decreased consumption in both sexes 	
	CLINICAL CHEMISTRY	
	- hematology: no abnormalities; one female at 10000 ppm had	
	elevated leucocyte, reticulocyte and platelet counts and slightly decreased hemoglobin, hematocrit and erythrocyte count	
	- Biochemistry: slightly elevated ALP activity at 10000 ppm	
	(weeks 6 and 13) significance at group means level; at week	
	13 (2 males at 5000 and 2 females and 1 male at 10000 ppm) decreased glucose in both sexes at 5000 and 10000 ppm (but	
	within biological range) significance at group means level	
	- Urinalysis: no abnormalities	
	MACRO- AND MICROSCOPIC FINDINGS:	
	No gross leasion were seen.	
	- Organ weights: no treatment related variations	
	 Histopathology: absence or reduction in cytoplasmic vacuolation in hepatocytes at all dose levels (and so a 	
	reduction of liver glycogen)	
	ANALYSES:	
	 stability of test substance: after 7 day storage values 	
	ranged from 79-87% of target concentration, samples taken in	
	week 1-4, 8 and 13 had mean concentrations of 84, 96 and 83%	
	of target concentration for 1000, 5000 and 10000 ppm respectively.	
Source	: Notox Hertogenbosch	
Test substance	: CAS 1819-00-9 (2-methoxy-3,6-dichlorobenzoic acid), purity 86.8%	
Conclusion	: NOAEL 342 mg/kg bw based on effects on body weight, food	
	consumption and elavated ALP	
Reliability 21.05.2001	: (1) valid without restriction	(6)
		(0)
Species Sex	: rabbit : male/female	
Strain	: New Zealand white	
Route of admin.	: dermal	
Exposure period	: 3 weeks	
Frequency of	: 5 days a week	
treatment Bost obs. poriod	- nono	
Post obs. period Doses	: none : 100, 500, 2500	
Control group	: yes	
Method	:	
Year	:	
GLP	: yes	
* = ·		
Test substance Method	: other TS : TEST ORGANISMS:	

5. Toxicity	ld 1918-00-9 Date 27.12.2001
	Date 21.12.2001
	- Species: New Zealand white rabbits - Age: no data - Weight at study initiation: males: 1.9 - 2.6 kg, females:
	- Weight at study initiation. males: 1.9 - 2.0 kg, lemales. 2.1-2.7 kg - Number of animals: 4/sex/dose group
	ADMINISTRATION / EXPOSURE
	- Doses: 100, 500 and 2500 mg/kg/day - Exposure period: 21 days
	- Duration of exposure: 6 hours - Route of administration: dermal - Post exposure period: none
	- Vehicle: 0.9% saline - Total volume applied: no details given. Maximum vehicle
	amount used was 5ml. - Area exposed: 10% of body surface
	 Occlusion: not specified Removal of test substance: by wiping
	CLINICAL OBSERVATIONS AND FREQUENCY:
	 pre- and post-test determination of hematological and biochemical blood parameters (total and differential loukooste counts, on throaste count, hematogrit, hemaglobin
	leukocyte counts, erythrocyte count, hematocrit, hemoglobin, alkaline phosphatase, blood urea nitrogen, glutamic pyruvate transaminase, glutamic oxaloacetate transaminase, calcium,
	inorganic phosphorus, fasting blood glucose, albumin, total protein)
	- pre- and post-test urinalysis (volume, specific gravity, color and appearance, pH, albumin, glucose, occult blood and
	billirubin) - Clinical signs and mortality: daily observations, scoring
	of dermal irritation - Body weight: weekly
	ORGANS EXAMINED AT NECROPSY (MACROSCOPIC AND MICROSCOPIC):
	 Organ weights: The spleen, liver, adrenals, ovaries/ testes, thyroid (parathyroid), brain and kidneys were
	weighed fresh. - Microscopic: skin (treated and untreated), gallbladder,
	lung, trachea, liver, kidneys, large intestine, small intestine, stomach, pancreas, urinary bladder, spleen,
	heart, regional lymph node, mesenteric lymph node, prostate/uterus, testes/ovaries, pituitary, thymus, thursid/page_adreade_thursid_ave_page, muscle_heare
	thyroid/pars, adrenals, thyroid, eye, nerve, muscle, bone marrow, spinal cord, brain, any unusual lesions
	STATISTICAL METHODS: analysis of variance (one-way classification), Bartlett's
Result	test, Dunnett's multiple comparison tables : TOXIC RESPONSE/EFFECTS BY DOSE LEVEL:
	- Mortality and time of death: males: 1(9) control, 1(17) 100 mg/kg; females 1(18) 100 mg/kg, 2(6&10) 500 mg/kg, 2(6&7) 2500 mg/kg
	- Clinical signs: Animals that died: diarrhea, hypoactivity, distended
	abdomen, anorexia and slight cyanosis. Surviving animals diarrhea and soft stools, erythema, desquamation, atonia, coriaceousness, fissuring
	- Body weight gain: no abnormalities
	23 / 39

5. Toxicity	Id 1918-00-9	
_	Date 27.12.2001	
	- Clinical chemistry: blood glucose in females at 2500 mg/kg significantly higher than controls, but within biological range	
	- Haematology: no abnormalities	
	 Urinalysis: Significant difference in pH for males at 2500 and females at 100 mg/kg compared to controls, but values were within biological range 	
	NECROPSY FINDINGS - Organ weights: increased adrenal weight (not toxicologically significant)	
	- Gross pathology: skin thickening and erythema of the application site in 2 rabbits at 2500 mg/kg/day	
Source	 Histopathology: at application site: acanthotic epidermal thickening and hyperkeratosis, slight parakeratosis. No dose response Notox Hertogenbosch 	
Test substance	 CAS 1918-00-9, (2-methoxy-3,6,-dichlorobenzoic acid), purity 86.8% 	
Reliability	 (3) invalid Too many animals died. From 8 control and 24 dosed rabbits one control and 6 exposed rabbits died during the study. Five of the six animals that died were female rabbits. Therefore 43% of the dosed female rats did not survive the study. This was not considered in the discussion of the data. The purity, stability and composition of the compound 	
21.05.2001	were not determined. 4. The food consumption was not monitored.	(7)
5.5 GENETIC TOXICIT	'Y 'IN VITRO'	
Type System of testing Concentration Cycotoxic conc. Metabolic activation Result	 Ames test TA98, TA100, TA1535, TA1537 and TA102 8-5000 ug/plate 1500 ug/plate with and without negative 	
Method Year	 OECD Guide-line 471 "Genetic Toxicology: Salmonella thyphimurium Reverse Mutation Assay" 1983 	
GLP	: yes	
Test substance Method	 other TS SYSTEM OF TESTING: Species/cell type: Salmonella typhimurium TA98, TA100, TA1535, TA1537 and TA102. Deficiences/Proficiences: histidine-requiring strains Metabolic activation system: rat S-9 mix, Arochlor 1254 induced 	
	ADMINISTRATION: - Dosing:	

Toxicity	ld 1918-00-9	
-	Date 27.12.2001	
	Mutation experiment 1 (without preincubation): 8, 40, 200,	
	1000, 5000µg/plate;	
	Mutation experiment 2: TA98, TA100, TA1535, and TA1537:	
	187.5, 375, 750, 1500 and 3000 ug/plate. TA102: 46.875, 93.75, 187.5, 375 and 750µg/plate.	
	- Number of replicates: 3	
	- Application: solution in DMSO	
	 Positive and negative control groups and treatment: 	
	Positive controls: -S9: 2-nitrofluorene (TA98), sodium azide	
	(TA100, TA1535), 9-aminoacridine (TA1537), gluturaldehyde (TA102).	
	+S9: 2-aminoanthracene (at least one strain).	
	Negative controls: DMSO (vehicle)	
	- Pre-incubation time: Mutation experiment 2; 1h incubation	
	at 37°C of S9 with the test compound prior to addition to	
	the tester strain.	
	CRITERIA FOR EVALUATING RESULTS:	
	- Statistical method: Dunnett's test	
.	- Method of calculation: linear regression analysis	
Result	: GENOTOXIC EFFECTS:	
	 With metabolic activation: none Without metabolic activation: none 	
	PRECIPITATION CONCENTRATION: no precipitation was observed	
	CYTOTOXIC CONCENTRATION: 1500 ug/plate with and without	
-	metabolic activation	
Source	: Notox Hertogenbosch	
Test substance	: CAS 1918-00-9 (3,6-dichloro-2-methoxybenzoic acid), purity 88.5%	
Reliability	: (1) valid without restriction	
16.05.2001		(2)
Туре	: Chromosomal aberration test	
System of testing	: CHO cells	
Concentration	: 300-2330 ug/ml	
Cycotoxic conc. Metabolic activation	: : with and without	
Result	: negative	
Method	:	
Year	:	
GLP Toot outpoteneo	: yes	
Test substance Method	 other TS - Species/cell type: Chinese hamster ovary (CHO-K1) cells 	
Method	- Metabolic activation system: rat S9 mix (Aroclor 1254	
	induced)	
	- No. of metaphases analyzed: 100	
	ADMINISTRATION:	
	- Dosing: 2330, 1170, 590 and 300 µg/ml.	
	- Number of replicates: 2	
	- Application: solution in DMSO	
	- Exposure time: 8 hours (-S9) or 2 hours (+S9)	
	 Positive and negative control groups and treatment: Positive controls: with S-9: triethylene melamine; without 	
	S-9: cyclophophamide	
	Negative controls: DMSO	
	CRITERIA FOR EVALUATING RESULTS:	
	- Statistical method: Student's t test	
	- method of calculation: linear regression analysis	
Result	: GENOTOXIC EFFECTS:	
	- With metabolic activation: none	
	25 / 39	

5. Toxicity	ld 1918-00-9 Date 27.12.2001
	- Without metabolic activation: none
	PRECIPITATION CONCENTRATION: No precipitation was observed
	CYTOTOXIC CONCENTRATION: No cytotoxicity was observed
	STATISTICAL RESULTS: no significant increase in number of aberrations in test group compared to control group.
	Positive control triethylene melamine gave 0.45 structural aberrations per cell, positive control Cyclophosphamide induced 0.69 aberrations per cell. This was in both cases a significant increase above the untreated control
Source Test substance	 Notox Hertogenbosch CAS 1918-00-9, (3,6-dichloro-2-methoxybenzoic acid), purity
	88.5%
Reliability	 (2) valid with restrictions 1. Only 100 metafases are scored (OECD 473: at least 200)
21.05.2001	(8)
5.6 GENETIC TOXICI	TY 'IN VITRO'
Туре	: Micronucleus assay
Species Sex	: mouse
Strain	: ICR
Route of admin.	: i.p.
Exposure period	: single dose
Doses Result	: 450, 900 and 1800 mg/kg bw : negative
Method	
Year	:
GLP	: yes
Test substance Method	: other TS : TEST ORGANISMS:
Method	- Species: ICR mice
	- Source: Harlan Sprague Dawley Inc., Frederick, MD.
	- Age: 6 to 8 weeks
	- Weight at study initiation: males (29.5 - 36.6g), females (25.5 - 32.0g)
	- No. of animals per dose: 15/sex/dose
	ADMINISTRATION:
	- Vehicle: deionized distilled water
	- Doses: 0, 450, 900, 1800 mg/kg bw.
	- Doses: 0, 450, 900, 1800 mg/kg bw. - Duration of test: Five animals of each dose group were
	 Doses: 0, 450, 900, 1800 mg/kg bw. Duration of test: Five animals of each dose group were killed after 24, 48, and 72 hr dosing.
	- Doses: 0, 450, 900, 1800 mg/kg bw. - Duration of test: Five animals of each dose group were
	 Doses: 0, 450, 900, 1800 mg/kg bw. Duration of test: Five animals of each dose group were killed after 24, 48, and 72 hr dosing. Frequency of treatment: single dose by i.p. injection Sampling times and number of samples: 24, 48 and 72 hours; 2-4 slides per animal
	 Doses: 0, 450, 900, 1800 mg/kg bw. Duration of test: Five animals of each dose group were killed after 24, 48, and 72 hr dosing. Frequency of treatment: single dose by i.p. injection Sampling times and number of samples: 24, 48 and 72 hours; 2-4 slides per animal Control groups and treatment:
	 Doses: 0, 450, 900, 1800 mg/kg bw. Duration of test: Five animals of each dose group were killed after 24, 48, and 72 hr dosing. Frequency of treatment: single dose by i.p. injection Sampling times and number of samples: 24, 48 and 72 hours; 2-4 slides per animal
	 Doses: 0, 450, 900, 1800 mg/kg bw. Duration of test: Five animals of each dose group were killed after 24, 48, and 72 hr dosing. Frequency of treatment: single dose by i.p. injection Sampling times and number of samples: 24, 48 and 72 hours; 2-4 slides per animal Control groups and treatment: Negative control group: vehicle 15 animals per sex. Positive control: cyclophosphamide, 5 animals per sex.
	 Doses: 0, 450, 900, 1800 mg/kg bw. Duration of test: Five animals of each dose group were killed after 24, 48, and 72 hr dosing. Frequency of treatment: single dose by i.p. injection Sampling times and number of samples: 24, 48 and 72 hours; 2-4 slides per animal Control groups and treatment: Negative control group: vehicle 15 animals per sex. Positive control: cyclophosphamide, 5 animals per sex.
	 Doses: 0, 450, 900, 1800 mg/kg bw. Duration of test: Five animals of each dose group were killed after 24, 48, and 72 hr dosing. Frequency of treatment: single dose by i.p. injection Sampling times and number of samples: 24, 48 and 72 hours; 2-4 slides per animal Control groups and treatment: Negative control group: vehicle 15 animals per sex. Positive control: cyclophosphamide, 5 animals per sex. EXAMINATIONS: mortality and clinical signs number of micronucleated Polychromatic erythrocytes
	 Doses: 0, 450, 900, 1800 mg/kg bw. Duration of test: Five animals of each dose group were killed after 24, 48, and 72 hr dosing. Frequency of treatment: single dose by i.p. injection Sampling times and number of samples: 24, 48 and 72 hours; 2-4 slides per animal Control groups and treatment: Negative control group: vehicle 15 animals per sex. Positive control: cyclophosphamide, 5 animals per sex. EXAMINATIONS: mortality and clinical signs number of micronucleated Polychromatic erythrocytes (PCE)/1000 PCE
	 Doses: 0, 450, 900, 1800 mg/kg bw. Duration of test: Five animals of each dose group were killed after 24, 48, and 72 hr dosing. Frequency of treatment: single dose by i.p. injection Sampling times and number of samples: 24, 48 and 72 hours; 2-4 slides per animal Control groups and treatment: Negative control group: vehicle 15 animals per sex. Positive control: cyclophosphamide, 5 animals per sex. EXAMINATIONS: mortality and clinical signs number of micronucleated Polychromatic erythrocytes

5. Toxicity	ld 1918-00-9
	Date 27.12.2001
Remark Result	 Evaluation of Test Results: statistical: Kastenbaum-Bowman The DMA salt of dicamba is the test substance. Mortality: males 4/20 and 1/15, females 3/20 and 0/15 at 1800 and 900 mg/kg resp.
	Clinical signs: lethargy at all dose levels
	 EFFECT ON PCE/NCE RATIO: number of micronucleated PCE per 1000 PCE: 450 mg/kg bw: 0.8, 0.3 and 0.2 at 24, 48 and 72 hours resp. 900 mg/kg bw: 0.9, 0.1 and 0.2 at 24, 48 and 72 hours resp. 1800 mg/kg bw: 1.4, 0.6 and 0.3 at 24, 48 and 72 hours resp. PCE/total erythrocytes 450 mg/kg bw: 0.65, 0.60 and 0.56 at 24, 48 and 72 hours resp. 900 mg/kg bw: 0.60, 0.58 and 0.56 at 24, 48 and 72 hours resp. 1800 mg/kg bw: 0.59, 0.52 and 0.62 at 24, 48 and 72 hours resp.
Source Test substance Reliability 21.05.2001	 Statistical results: micronucleated PCE/1000 PCE was not significantly increased at any dose level at any collection time in either males or females. The positive control induced a significant increase in micronucleated PCE/1000 PCE Notox Hertogenbosch Dicamba DMA salt, purity 40.3% (3) invalid Purity of the test substance is unknown. It is not mentioned what DMA (DMA salt of dicamba) stands for. Only 1000 erythrocytes are scored for incidence of micronucleated PCE (OECD 474, 1997: at least 2000) Sampling at 72 hours is too late. However 2 sampling times remain (24 and 48 hours), which is sufficient according to OECD 474, 1997.
5.7 CARCINOGENITY	1
5.8 TOXICITY TO REI	
Type Species Sex Strain Route of admin. Exposure period Frequency of treatment	 Two generation study rat male/female other: CrI:CD-(SD) BR VAF/Plus oral feed Parent-generation (males/females): 10 weeks prior to mating until weaning of the litters (day 21 post-partum); F1-generation 12 weeks prior to mating until weaning of the litters (day 21 post-partum) continuous
Premating exposure period Male Female Duration of test	 10 weeks (parental generation) or 12 weeks (F1-generation) 10 weeks (parental generation) or 12 weeks (F1-generation) 50 weeks

Toxicity	ld 1918-00-9 Date 27.12.2001
ooses Control group IOAEL Parental IOAEL F1 Offspr. IOAEL F2 Offspr. Iethod	 = 500 ppm OECD Guide-line 416 "Two-generation Reproduction Toxicity Study" 1983
GLP Test substance Method	 yes other TS TEST ORGANISMS (PARENTAL GENERATION): Age: males/females 6 weeks at start treatment Weight at study initiation: At start treatment males
	180-271g and females 137-190g - Source: Charles River UK Ltd - Number of animals: 32/sex/treatment (parental), 28/sex/treatment (F1)
	ADMINISTRATION / EXPOSURE - Test duration: maximum 50 weeks - Exposure period: males and females 10 weeks (parent generation) or 12 weeks (F1-generation) prior to mating and until weaning of the F1 or F2 generation, respectively - Route of administration: oral via the diet - Doses: 0, 500, 1500 and 5000 ppm in the diet
	MATING PROCEDURES (PARENTAL AND F1-GENERATION): - Mating: 1 female / 1 male (or occasionally 2 females / 1 male) during 20 days - Day 0 of gestation: presence of vaginal plugs and/or spermatozoa in the vaginal smear of females
	PARAMETERS ASSESSED DURING STUDY (PARENTAL AND F1-GENERATION):
	 Mortality/clinical observations: regularly Body weight gain: weekly (males/females) or daily for females during mating and until parturition Food consumption: weekly during the premating treatment
	phases - Water consumption: daily during initial and final two weeks of the premating treatment periods - Female oestrous cycle: vaginal cytology examination 7 days
	prior to mating (parental generation) and the first mate of the F1-generation and during the 20-day mating period - Male sperm analysis: at necropsy samples from both vas deferens were analysed for total count, motility and
	morphology (1 every 4 male rat/cage). Left testis examined for spermatid counts - Mating and fertility data (males/females): number and days of successful matings, time between pairing and mating (with
	 1rst or 2nd male, F1-generation) Maternal delivery data: duration of gestation, number pregnant, litter size (live pups) and number of implant sites
	- Pup viability: number of live pups at birth and post-partum days 4, 8, 12, 16, 21 (culling on day 4 post-partum to 8 pups/litter)
	 Pup observations: clinical signs, sex and external examinations; body weights on days 1 (birth), 4, 8, 12, 16 and 21 post-partum; sexual maturation of female pups by the unset of vaginal opening (as of day 28 post-partum) and of

5. Toxicity	ld 1918-00-9 Date 27.12.2001
	balanopreputial skinfold (as of day 35 post-partum)
	 ORGANS EXAMINED AT NECROPSY (PARENTAL AND F1-GENERATIONS): Macroscopy: all males and females (parental generation), those selected for pairing (F1-generation) and one male and one female pup from each litter (day 21 post-partum) were necropsied and gross findings recorded. The following organs were weighed; adrenals. brain, heart, kidneys, liver, lungs, pituitary prostate (with seminal vesicles and coagulating gland) tests with epididymides and thymus. Additionally, a full range of tissues (see microscopy) was preserved for histopathology. Remaining pups were examined externally and internally and the sex was confirmed by gonadal inspection. Gross findings were preserved (when considered usefull) for possible histopathology Microscopy: histopathology examinations were preformed on the adrenals, aorta, bone and joint, bone marrow, brain, cranial vault, caecum, colon, duodenum, eyes, heart, ileum, jejunum, kidneys, liver, lungs, lymph nodes, mammary gland, oesophagus, ovaries, pancreas, pituitary, prostate (for F1 weanlings with seminal vesicles and coagulating gland), rectum, salivary gland, seminal vesicles (with coagulating gland) sciatic nerve, skeletal muscle, skin, spinal column, spleen, stomach, testes, epididymides, thymus, thyroids (with parathyroids), tongue, trachea (with larynx and pharynx), urinary bladderuterus (with cervix) vagina and vas deference
	ANALYSES: - Method: High Performance Liquid Chromatography (HPLC) with UV detection - Sampling time: prior to start of the first premating treatment (500 ppm and 12000 ppm dietary inclusion levels) for analysis of stability and homogeneity. Samples for accuracy of exposure concentrations for each generation were taken at start of the premating treatment and at start of the mating and end of gestation/start lactation
Result	 STATISTICAL METHODS: analysis of variance, Williams' test, Kruskal-Wallis test, Analysis of covariance, Shirley's test, Fisher's exact test ANALYSES: Actual dose level: the accuracy of all test diets was acceptable (94-112% of nominal) Stability: stable for at least 18 days (within 91-93%) Homogeneity: homogeneous (all samples 91-99% of nominal) Actual intake during week 1-10 at 500, 1500 and 5000 ppm: F0: males 35, 105 and 347 mg/kg bw resp., females 41, 125 and 390 mg/kg bw resp. F1: males 40, 121 and 432 mg/kg bw resp., females 44, 35 and 458 mg/kg bw resp.
	TOXIC EFFECTS BY DOSE LEVEL
	PARENTAL GENERATION:
	 Mortality: at 500 and 5000 ppm one female Body weight gain: at 5000 ppm decreased in females during pregnancy and the first week of lactation Food consumption/water consumption: no treatment-related
	29 / 39

ld 1918-00-9

Date 27.12.2001

5. Toxicity

findings

- Clinical signs: incidental hairless and scabbing, but no treatment-related findings

- Mating and fertility data (males/females): no differences between the dose groups (sperm motility, morphology and number normal); pregnant females at 500, 1500 and 5000 ppm 27, 28, 29 and 27 resp.

- Maternal delivery data: at 5000 ppm slight shift of the duration of pregnancy from 22/23 to 21 days and decreased litter and pup weights

 Macroscopic examinations: pale subpleural foci on the lungs of males at 5000 ppm (parent); increased incidence of pelvic dilations in pups (without relationship to dose)
 Organ weights:

parents: at 5000 ppm increased rel. liver weights in females, decreased epididymides, prostate and rel. kidney weight in males; at all treatments decreased pituitary weight (rel.)

pups: at 1500 ppm increased liver and decreased lung weights (both relative); at 5000 ppm decreased absolute brain weight and relative heart and lung and increased relative liver weight

Microscopic examinations: no treatment-related findings
 Pup viability/observations: at 5000 ppm decreased pup weights and delayed sexual maturation of the males, no effects on sex ratio.

F1 GENERATION:

- Mortality: at 0, 500, 1500 and 5000 ppm, 2 males/1 female, 1 male/1 female, 1 male and 1 male, respectively

- Body weight: decreased in males at 5000 ppm and females at 5000 ppm during the first weeks after weaning

- Food consumption/water consumption: at 5000 ppm in males and females decreased (food weeks 5-8/water weeks 5-6 of premating treatment)

- Clinical signs: at 5000 ppm increased incidence of tense/stiff body tone and slow righting reflex at the latter part of lactation

- Mating and fertility data (males/females): first mate gave pregnancy rate of 56-75%; second mate 56-68%; sperm motility, morphology and number normal

- Maternal delivery data: at 5000 ppm decreased pregnancy rate (first mate), decreased litter weights; slightly higher pup loss (second mate) resulting in slightly lower litter sizes at 1500 and 5000 ppm

- Macroscopic examinations: dose related increase of the number of pale foci on the lungs in parents

- Organ weights:

parents: at 5000 ppm increased liver weights (absolute females, relative males); at all treatments kidney weight decreased relative to body weight

pups: at 5000 ppm increased relative liver weight, decreased rel. kidney and heart weight

Microscopic examinations: no treatment-related findings
 Pup viability/observations: at 5000 ppm decreased pup

weights and associated delayed male and female sexual maturation

F2 GENERATION:

- Clinical signs: no treatment-related findings

30 / 39

. Toxicity	ld 1918-00-9 Date 27.12.2001
Source	 Pup viability/observations: at 1500 slightly decreased pup weights and at 5000 ppm decreased pup weights and increased liver weights Notox Hertogenbosch
Test substance	: I, CAS 1918-00-9 (dicamba technical, 3,6-dichloro-o-anisic
Conclusion	 acid), purity 86.9% NO(A)EL (parents): 1500 ppm, based on decreased female body weight gain during pregnancy and increased liver weights in both sexes in the 5000 ppm group. NO(A)EL (F1-generation): 1500 ppm, based on a marked impairment of growth of the F1-offspring and associated reduced food and water consumption, slightly delayed sexual maturation of males and increased liver weights. Additionally F1-females showed slightly lower body weight gain during pregnancy and signs of increased bodytone and
Reliability	 slow righting reflex during late lactation NO(A)EL (F2 generation): 500 ppm, based on reduced body weight gain of F1-females during pregnancy and slghtly reduced growth of F2-pups (1) valid without restriction
21.05.2001	. (1) valid without restriction (5
Sex Strain Route of admin. Exposure period Frequency of treatment Duration of test Doses Control group NOAEL Maternalt. NOAEL Teratogen NOAEL Teratogen NOAEL Fetotoxicity Method Year GLP Test substance Method	 Crj: CD(SD) gavage gestation days 6-19 Once daily Caesarean sections on gestation day 20 64, 160 and 400 mg/kg/day yes, concurrent vehicle <= 160 mg/kg bw <= 400 mg/kg bw <= 400 mg/kg bw other: US 43 FR 37336, Part 163.83-3 1981 yes other TS TEST ORGANISMS Age: females not indicated (sexually mature) Weight at study initiation: 196-251g (gestation day 0) Number of animals: 25 (treatment/control groups) Source: Stone Ridge, N.Y. facilities of Charles River, Breeding Laboratories, Inc. USA
	ADMINISTRATION / EXPOSURE

5. Toxicity	ld 1918-00-9
-	Date 27.12.2001
	 PARAMETERS ASSESSED DURING STUDY: Mortality: twice daily Clinical observations: twice daily (early morning, late afternoon) Body weight gain: gestation days 0, 6 and 20 Food consumption: daily (gestation days 0-19) Examination of uterine content: number and distribution of implantations, early and late resorptions and live and dead foetuses Examination of fetuses: sex; weight; external, visceral (1/3) and skeletal (2/3 foetuses) findings ORGANS EXAMINED AT NECROPSY (MACROSCOPIC AND MICROSCOPIC): Macroscopy: not indicated
	- Microscopy: no tissues retained
	OTHER EXAMINATIONS:
	No
	ANALYSES: - Method: Liquid Chromatograph (HPLC) - Sampling time: samples taken from all preparations (1 interval subjected to analysis)
Result	 STATISTICAL METHODS: Scheffe's or Turkey's ANALYSES: Actual dose level: dose preparations were confirmed to be accurate Stability: Stable during at least 1 week
	MATERNAL TOXIC EFFECTS BY DOSE LEVEL:
	 Mortality and day of death: at 400 mg/kg 3 females died on gestation days 7 or 8 Body weight: at 400 mg/kg decreased on gestation day 20 Food consumption: at 400 mg/kg decreased during exposure (gestation days 6-19) Clinical signs: at 400 mg/kg females showed increased incidence of crusty nose/muzzle, wheezing, ataxia, stiffening of the body when held, urine soaked fur, salivation and decreased motor activity Number pregnant per dose level: at 0, 64, 160 and 400 mg/kg, 23, 24, 23 and 17, respectively Number of resorptions (early/late): at 0, 64, 160 and 400 mg/kg, 6.4%, 3.0%, 5.3% and 8.7%, respectively (percent of implantation sites) Number of implantations: at 0, 64, 160 and 400 mg/kg, 14.2, 12.3, 14.3 and 13.1, respectively Post implantation loss: idem number of resorptions Number of corpora lutea: not recorded Duration of Pregnancy: scheduled sacrifice on gestation day 20 Gross pathology incidence and severity: no findings
	FETAL DATA:
	There were no gross external, soft tissue or skeletal alterations that were considered effects of the test
1	32 / 39

. Toxicity	ld 1918-00-9	
	Date 27.12.2001	
	substance. Foetal body weight and sex were comparable between all groups	
	- Litter weights (gravid uterus): at 0, 64, 160 and 400 mg/kg, 73g, 66g, 75g and 62g, respectively - Number viable: at 0, 64, 160 and 400 mg/kg, 13.3, 11.9,	
	- Number viable, at 0, 64, 160 and 400 mg/kg, 13.3, 11.9, 13.6 and 11.8, respectively - Sex ratio (percentage of males): at 0, 64, 160 and 400	
	mg/kg, 49.2%, 49.0%, 49.5% and 52.0%, respectively - Body weight: at 0, 64, 160 and 400 mg/kg, for males 3.5g,	
	3.5g, 3.4g and 3.3g, respectively and for females 3.3g, 3.3g, 3.2g and 3.1g, respectively. - Grossly visible abnormalities: at 160 mg/kg one foetus	
	showed a shortened body and anurous - Visceral abnormalities: at 400 mg/kg increased incidence renal pelvic cavitation (one litter)	
	 Skeletal abnormalities: at 400 mg/kg percentage incomplete frontal(s) and/or parietal(s) ossification 	
Source Test substance	 Notox Hertogenbosch I, CAS 1918-00-9 (dicamba technical, 3,6-dichloro-o-anisic acid), purity 86.9% 	
Conclusion	I, CAS 1918-00-9 (technical Dicamba), purity: technical grade	
Conclusion	 NOAEL (maternal): 160 mg/kg based on decreased body weights and food consumption and clinical symptoms such as ataxia stiffening of the body when held and decreased motor 	
	activity at 400 mg/kg NOAEL (teratogenicity): 400 mg/kg based on the absence of any significantly increased malformation or variation	
Reliability	 NOAEL (foetotoxicity): 400 mg/kg based on the absence of any effects on foetal growth or deaths (1) valid without restriction 	
-	No corpora lutea recorded Post implantation loss not calculated	
15.05.2001		(16)
Species Sex	: rabbit : female	
Strain	: New Zealand white	
Route of admin.	: other: oral via capsules	
Exposure period Frequency of	: gestation days 6-18 : Once daily	
treatment Duration of test	: Caesarean sections on gestation day 29	
Doses Control group	: 30, 50 and 300 mg/kg : yes, concurrent vehicle	
NOAEL Maternalt.	= 30 mg/kg bw	
NOAEL Teratogen Method	: <= 300 mg/kg bw :	
Year	: 1984	
GLP	: yes	
Test substance Method	: other TS : TEST ORGANISMS	
Method	- Age: females (at insemination) 26 weeks	
	- Weight at study initiation: 3.05-4.14 kg - Number of animals: 20 (treatment groups), 19 (control	
	group	
	- Source: Hazelton Research Products, Inc., Denver Pennsylvania, USA	
	ADMINISTRATION / EXPOSURE - Test duration: 29 days	
	33 / 39	

5. Toxicity	ld 1918-00-9 Date 27.12.2001
	 Exposure period: gestation days 6-18 Route of administration: oral (via capsules) Doses: 0, 30, 150 and 300 mg/kg Vehicle: opaque white gelatin capsules
	MATING PROCEDURES: - Artificial insemination: Semen collected from 4 proven donor bucks of the same strain and source as the females. 3 hours before insemination females were intravenously injected with 20 USP units of Human Chorionic Gonadotropin. Insemination of 0.25 mL of diluted (with saline) semen sample (6.0 million spermatozoa/0.25 mL) - Day 0 of gestation: day of insemination
	PARAMETERS ASSESSED DURING STUDY: - Mortality: twice daily - Clinical observations: once daily or on gestation days 6-19 immediately before dosage and within 60 minutes after dosage - Body weight gain: once weekly before insemination and on gestation days 0 and 6-29 - Food consumption: daily - Examination of uterine content: number of corpora lutea;
	number and distribution of implantations, early and late resorptions and live and dead foetuses - Examination of fetuses: sex; weight; external, visceral (all foetuses) and skeletal (all foetuses) findings; brains free-hand cross-sectioned and examined for hydrocephaly
	ORGANS EXAMINED AT NECROPSY (MACROSCOPIC AND MICROSCOPIC): - Macroscopy: findings all dams recorded, all gross lesions (except commonly found parovarian cysts) were fixed for possible histopathology - Microscopy: not performed
	OTHER EXAMINATIONS: - Uterus staining: uteri from non-pregnant rabbits were stained with 10% ammonium sulfide to comfirm absence of implantation sites
	ANALYSES: - Method: Not indicated (samples not analysed) - Sampling time: Bulk test substance sampled on day 2 and the end of the dosing period for possible analysis
Result	 STATISTICAL METHODS: Bartlett's Test, Dunnett's Test, Kruskal-Wallis Test, Dunn's Test and Fisher's Exact Test ANALYSES: No analyses performed. Test substance dosed via capsules. Data on the identity, composition, strength, purity and stability of the test substance are kept on file with the sponsor
	MATERNAL TOXIC EFFECTS BY DOSE LEVEL:
	There were no differences noted among the dose groups in the number of corpora lutea, implantations, litter sizes, early and late resorptions, foetal sex ratio, foetal body weights, percent resorbed conceptuses and number of does with any resorptions
	34 / 39

5. Toxicity	ld 1918-00-9
	Date 27.12.2001
	 Mortality and day of death: One female dosed at 300 mg/kg died due to an intubation error on gestation day 12. Abortion and subsequent sacrifice occurred in the 150 mg/kg dose group for 1 female on gestation day 22 and in the 300 mg/kg dose group for four females on gestation days 19 (one female), 21 (one female) and 24 (two females) Body weight: at 300 mg/kg body weight loss on gestation days 19 (one female), 20 (one
	days 6-7, 6-9, 9-12, 12-15, 15-19 and overall loss during gestation days 6-19. Decreased overall body weight gain during gestation days 6-19 (loss), 6-29 and 0-29 - Food consumption: at 300 mg/kg often during the dosing period resulting in a reduced overall food consumption during gestation days 6-10, 6-20 and 0-20
	during gestation days 6-19, 6-29 and 0-29 - Clinical signs: at 150 and 300 mg/kg females showed ataxia (and decreased motor activity). In addition, females receiving 300 mg/kg incidentally showed rales, laboured breathing, perinasal substance (red or yellow), dried
	faeces, impaired righting reflex, no faeces and a red substance in the cage pan - Number pregnant per dose level: 16 (80% of number inseminated) in the 30 mg/kg group and 18 in all other groups (90-94.7% of number inseminated) - Number aborting: at 150 mg/kg 1 and at 300 mg/kg 4
	 Number abouting. at 150 mg/kg 1 and at 500 mg/kg 4 Number of resorptions (early/late): at 0, 30, 150 and 300 mg/kg, 0.5, 0.5, 1.0 and 0.5, respectively Number of implantations: at 0, 30, 150 and 300 mg/kg, 6.8, 5.9, 6.4 and 6.3, respectively Post implantation loss: at 0, 30, 150 and 300 mg/kg, 6.4%,
	 4.8%, 10.1% and 7.6%, respectively Number of corpora lutea: at 0, 30, 150 and 300 mg/kg, 9.6, 8.4, 8.9 and 9.2, respectively Duration of Pregnancy: scheduled sacrifice on gestation day 29
	- Gross pathology incidence and severity: no findings other then those related to intubation error (thick, hard and gray oesophagus and trachea containing white mucoid substance) or commonly found parovarian cysts
	FETAL DATA:
	There were no gross external, soft tissue or skeletal alterations that were considered effects of the test substance
	- Litter size and weights: at 0, 30, 150 and 300 mg/kg, 6.3, 5.4, 5.4 and 5.8, respectively - Number viable: at 0, 30, 150 and 300 mg/kg, 6.3, 5.4, 5.4 and 5.8, respectively
	 Sex ratio (percentage of males): at 0, 30, 150 or 300 mg/kg, 49.4%, 64.4%, 54.7% and 54.6%, respectively Body weight: at 0, 30, 150 and 300 mg/kg, 44.55g, 47.11g, 44.20g and 42.47g, respectively Grossly visible abnormalities: incidentally observed
	findings consisted of umbilical hernia, menigocele, medially rotated hindlimbs, flexed hindpaws and shortened tail - Visceral abnormalities: incidental findings comprised protrusion of the liver through the abdominal wall, agenesis of the intermediate lobe of the lungs, agenesis of the gall bladder and caudally displaced right kidney.
	 Skeletal abnormalities: incidentally observed finding consisted of vertabral malformations (irregular shaped left

. Toxicity	ld 1918-00-9 Date 27.12.2001	
Source Test substance Conclusion	 arch of the 3rd lumbar vertebra and fosion of the left arches of the 3rd and 4th lumbar vertebrae), tail malformation (14 vertebrae present) and variations in skull and sternal ossification (displaced nasal suture, internasal ossification site and fused 3rd and 4th sternebrae) Notox Hertogenbosch I, 1918-00-9 (Technical dicamba), purity (not reported) NOAEL (maternal): 30 mg/kg based on the abortions, clinical signs (viz. decreased motor activity, ataxia, rales, laboured breathing, perinasal substance red/yellow, dried faeces, impaired righting reflex, no faeces, red substance in the cage pan), reduced body weight gains and reduced feed consumption NOAEL (teratogenicity): 300 mg/kg based on the absence of any significantly increased malformation or variation NOAEL (foetotoxicity): 300 mg/kg based on the absence of any effects on foetal growth or deaths 	
Reliability	: (1) valid without restriction	
19.04.2001		(1

5.10 OTHER RELEVANT INFORMATION

5.11 EXPERIENCE WITH HUMAN EXPOSURE

6. Refer	ences Id 1918-00-9 Date 27.12.2001	
(1)	ARGUS RESEARCH LABORATORIES INC, DEVELOPMENTAL TOXICITY (EMBRYO-FETAL TOXICITY AND TERATOGENIC POTENTIAL) STUDY OF TECHNICAL DICAMBA ADMINISTERED ORALLY VIA CAPSULE TO NEW ZEALAND WHITE RABBITS, 1992 (103)	
(2)	Ballantyne, M., Dicamba Technical: Reverse mutation in five histidine-requiring strains of Salmonella typhimurium	
(3)	EPIWIN v3.05, Syracuse Research Corporation, Syracuse, NY (July 12, 2000)	
(4)	Hansch, C., Leo, A., D. Hoekman. Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. Washington, DC: American Chemical Society., 1995. 37 (as cited in Hazardous Substance Data Base)	
(5)	Huntingdon Research Centre Ltd., Huntingdon, England, Technical dicamba A study on the reproductive function of two generations in the rat, 1993	
(6)	International Research and Development Corporation, 13-week dietary toxicity study in rats with Dicamba, 1980	
(7)	International Research and Development Corporation, 3-week dermal toxicity study in rabbits, 1979	
(8)	Microbiological Associates Inc., Chromosome aberrations in Chinese hamster ovary cells, 1986	
(9)	Sandoz Agro Inc, Dicamba technical - toxicity to the freshwater green alga, Selenastrum capricornutum (BASF 93/5221), 1993 (98)	
(10)	Sandoz Agro, Dicamba: Photodegradation Study in pH 7 Aqueous Solution (1993) (95) unpublished study	
(11)	Sandoz Agro, Inc., Micronucleus cytogenetic assay in mice, 1994	
(12)	Sandoz Agro, Melting Point of Dicamba, Technical (1993) (89)	
(13)	Sandoz Agro, Solubility of Technical Dicamba in Solvents, unpublished report (1993) (91)	
(14)	Sandoz Agro, Vapor Pressure of Dicamba Using the Thermal Evolution Analyzer, unpublished report (1994) (92)	
(15)	Sandoz Crop Protection Corporation, Determination of the n-octanol/water partition coefficient for dicamba, 1987	
(16)	ToxiGenics, Inc., Decatur, USA. Teratology study in albino rats with Technical Dicamba. 1981 (102)	
(17)	U.S. Department of Interior, Fish and Wildlife Service. Handbook of Acute Toxicity of Chemicals to Fish and Aquatic Invertebrates. Resource Publication No. 137. Washington, DC: U.S. Government PrintingOffice, 1980. 27, as cited in HSDB record for dicamba.	
(18)	Velsicol Chemical Corporation, Acute Toxicity Studies in rats and rabbits, 1974 (99)	
(19)	Velsicol Chemical Corporation, Hydrolysis of 14C-dicamba, 1981	

6. Refere	ences	ld 1918-00-9 Date 27.12.2001	
(20)	Velsicol Chemical Corporation, The acute toxicity of banvel technical to the sheepshead minnow Cyprinodon variegatus (BASF 77/5078), 1977 (97)		

7. R	isk Assessment	ld	1918-00-9
		Date	27.12.2001
7.1	END POINT SUMMARY		
7.2	HAZARD SUMMARY		
7.3	RISK ASSESSMENT		

IUCLID

Data Set

Existing Chemical CAS No. Generic name Tag name	 ID: 1982-69-0 1982-69-0 3,6-dichloro-2-methoxybenzoic acid, sodium salt dicamba, sodium
Producer Related Part Company Creation date	Toxicology and Regulatory Affairs25.12.2001
Substance Related Part Company Creation date	: Toxicology and Regulatory Affairs : 25.12.2001
Memo	:
Printing date Revision date Date of last Update	: 26.12.2001 : : 26.12.2001
Number of Pages	: 17
Chapter (profile) Reliability (profile) Flags (profile)	 Chapter: 1, 2, 3, 4, 5, 7 Reliability: without reliability, 1, 2, 3, 4 Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. 0	General Information	1982-69-0 26.12.2001
1.0.1	OECD AND COMPANY INFORMATION	
1.0.2	LOCATION OF PRODUCTION SITE	
1.0.3	IDENTITY OF RECIPIENTS	
1.1	GENERAL SUBSTANCE INFORMATION	
1.1.0	DETAILS ON TEMPLATE	
1.1.1	SPECTRA	
1.2	SYNONYMS	
1.3	IMPURITIES	
1.4	ADDITIVES	
1.5	QUANTITY	
1.6.1	LABELLING	
1.6.2	CLASSIFICATION	
1.7	USE PATTERN	
1.7.1	TECHNOLOGY PRODUCTION/USE	
1.8	OCCUPATIONAL EXPOSURE LIMIT VALUES	
1.9	SOURCE OF EXPOSURE	

1. General Information	1982-69-0 26.12.2001
1.10.1 RECOMMENDATIONS/PRECAUTIONARY MEASURES	
1.10.2 EMERGENCY MEASURES	
1.11 PACKAGING	
1.12 POSSIB. OF RENDERING SUBST. HARMLESS	
1.13 STATEMENTS CONCERNING WASTE	
1.14.1 WATER POLLUTION	
1.14.2 MAJOR ACCIDENT HAZARDS	
1.14.3 AIR POLLUTION	
1.15 ADDITIONAL REMARKS	
1.16 LAST LITERATURE SEARCH	
1.17 REVIEWS	
1.18 LISTINGS E.G. CHEMICAL INVENTORIES	

2. Physico-Chemical Data

2.1 MELTING POINT

Value Sublimation Method Year GLP Test substance Method Remark Result	 ca. 225 ° C other: Estimation 2001 no Estimation using MPBPWIN v1.40 in EPIWIN 3.05 As a salt of a substance melting about 100 C, this material will have a higher MP and be solid at temperaturec below 100 C. SUMMARY MPBPWIN v1.40 	
Test substance Reliability Flag 25.12.2001	 Boiling Point: 525.94 deg C (Adapted Stein and Brown Method) Melting Point: 349.84 deg C (Adapted Joback Method) Melting Point: 193.43 deg C (Gold and Ogle Method) Mean Melt Pt : 271.64 deg C (Joback; Gold,Ogle Methods) Selected MP: 224.71 deg C (Weighted Value) CAS 1982-69-0 Sodium salt of dicamba (2) valid with restrictions Critical study for SIDS endpoint 	1)

2.2 BOILING POINT

2.3 DENSITY

2.3.1 GRANULOMETRY

2.4 VAPOUR PRESSURE

Value Decomposition Method Year GLP Test substance Remark Result		<.00001 hPa at ° C other (calculated) 2001 no Estimation using MPBPWIN v1.40 in EPIWIN 3.05 Vapor Pressure Estimations (25 deg C): (Using BP: 525.94 deg C (estimated)) (Using MP: 224.71 deg C (estimated)) (Using MP: 224.71 deg C (estimated)) VP: 2.44E-013 mm Hg (Antoine Method) VP: 4.36E-011 mm Hg (Modified Grain Method) VP: 1.36E-010 mm Hg (Mackay Method) Selected VP: 4.36E-011 mm Hg (Modified Grain Method)
Source Test substance	:	Toxicology and Regulatory Affairs, Freeburg IL CAS 1982-69-0 Sodium salt of dicamba

	2001
: (2) valid with restrictions: Critical study for SIDS endpoint	(1)
FFICIENT	
 =9 at ° C other (calculated) 2001 Log Kow(version 1.66 estimate): -0.90 SMILES : c1(CL)ccc(CL)c(OC)c1C(=O)O[Na] CHEM : Dicamba, Sodium salt MOL FOR: C8 H5 CL2 O3 Na1 MOL WT : 243.02 	
TYPE NUM LOGKOW FRAGMENT COEFF VALUE + Frag 1 -CH3 0.5473 0.5473 Frag 6 Aromatic Carbon 0.2940 1.7640 Frag 2 -CL 0.6445 1.2890 Frag 1 -O- -0.4664 -0.4664 Frag 1 -C(=O)O -0.7121 -0.7121 Factor 1 C(=O)-O-{Na -3.5500 -3.5500 Const Equation Constant 0.2290 + Log Kow = -0.8992 * Toxicology and Regulatory Affairs, Freeburg IL * CAS 1982-69-0 Sodium salt of dicamba * (2) valid with restrictions * Critical study for SIDS endpoint	(1)
	(')
 ca. 150 g/l at 25 ° C at 25 ° C at and ° C other: calculated 2001 Estimation using WSKOW v1.40 in EPIWIN 3.05 WSKOW v1.40 Results Log Kow (estimated) : -0.90 Log Kow (estimated) : -0.90 Log Kow (experimental): not available from database Log Kow used by Water solubility estimates: -0.90 Equation Used to Make Water Sol estimate: Log S (mol/L) = 0.796 - 0.854 log Kow - 0.00728 MW + Correction 	on
	<pre>: Critical study for SIDS endpoint EFFICIENT : =9 at ° C other (calculated) : 2001 : : : Log Kow(version 1.66 estimate): -0.90 SMILES : c1(CL)ccc(CL)c(OC)c1C(=O)O[Na] CHEM : Dicamba, Sodium salt MOL FOR: C8 H5 CL2 O3 Na1 MOL WT : 243.02</pre>

2. Physico-Chem	ical Data		1982-69-0 26.12.2001	
	Correction(s): Value			
	No Applicable Correction Factors			
Source Test substance Reliability Flag	Log Water Solubility (in moles/L): -0.20 Water Solubility at 25 deg C (mg/L): 1.51 Toxicology and Regulatory Affairs, Freebur CAS 1982-69-0 Sodium salt of dicamba (2) valid with restrictions Critical study for SIDS endpoint	5e+005		
25.12.2001				(1)
2.6.2 SURFACE TENSIO)N			
2.7 FLASH POINT				
2.8 AUTO FLAMMABI	LITY			
2.9 FLAMMABILITY				
2.10 EXPLOSIVE PROF	PERTIES			
2.11 OXIDIZING PROPE	ERTIES			
	IARKS			

3.1.1 PHOTODEGRADATION

Type Light source Light spect. Rel. intensity Conc. of subst. Direct photolysis Halflife t1/2	: 100.19 mg/l at 25 degree C : 50.3 day
Degradation Quantum yield Method	 31.3 % after 30 day A 1000 mL test solution consisting of 100.19 mg dicamba with a specific activity of 412.2 dpm/ug (total 688 kBq) in aqueous buffer solution pH 7 containing 1% acetonitrile was prepared. The test solution was incubated at 25 +/- 1 deg C under contineous stirring for 30 days. Average incident radiation on the reactor surface was 7.704E2 W/m2 (measured before and after the study). The reaction solution was aerated and connected to a silica gel trap, an ethylene glycol trap (organic volatiles) and a 10% NaOH trap (supposed to collect CO2) in series. Before initiation of photolysis, a 50 mL sample was taken as dark control sample. 20 mL samples were taken before initiation
	of photolysis and on day 1, 3, 8, 15, 22 and 30. The samples were analyzed as follows: - duplicate 1 mL samples were analyzed by LSC - 15 mL was extracted twice at pH < 1 with ethyl acetate, both fractions were analyzed by LSC (duplicate 1 mL samples) - ethyl acetate fraction was dried and concentrated, and analyzed by TLC using 4 solvent systems (cochromatographed with reference standards) - extracted buffer solution of day 15, 22 and 30 were lyophilized followed by acetonitrile extraction; the extract was concentrated and analyzed by TLC using 4 solvent systems (cochromatographed with reference standards) - duplicate 1 mL ethylene glycol and 10% NaOH trap samples were analyzed by LSC - silica gel traps were extracted with with methanol, which was then analyzed by LSC; residual radioactivity in the silica traps was determined by combustion - identity of radioactivity supposed to be CO2 in 10% NaOH trap samples was confirmed for day 22 and 30 by precipitation as BaCO3 and subsequent evolution as CO2 after addition of HCI
	On day 30, the reactor was washed with methanol and with acetone. Volumes were measured and 1 mL duplicatealiquots were analyzed by LSC.
Remark	 Photodegradation was calculated using the SAS Regression Program.A 1000 mL test solution consisting of 100.19 mg dicamba with The test substance for this study was dicamba (acid form) rather than the salt. In solution, at pH 7 it does not matter if the salt or acid form is used to prepare the solution.

3. Environmenta	al Fate and Pathways	ld 1982-69-0 Date 26.12.2001
Result	: time point (days) 14C-dicamba (% of ac 14C-dicamba)*	ctually applied
	0 100 (92.14% of applied 140 1 98.83 3 95.25 8 86.87 15 75.62 22 66.44 30 58.74 (degradation: 41.26% 30 (dark control) 98.61 * calculated by reviewer from % of applie	%)
	Unchanged dicamba was confirmed by F All other compounds in the different fract TLC, were <10% of applied 14C and did reference standards. CO2 in the 10% Na applied at day 22 and 16.6% of applied 1 Radioactivity in the other traps was <10% all time points. Reactor wash yielded 0.3 activity. The mass balance was >99% ar	ions, separated by not match with IOH trap was 11.7% of 4C at day 30. 6 of applied 14C at % of applied
Test substance Conclusion	 points. Under these conditions, t1/2 of dicamba photolysis rate constant was 0.018 day-1 spring sunlight intensity at 40 deg latitude W/m2) the corresponding photodegradat sunlight will be 0.0138 day-1; t1/2 will be CAS 1918-00-9 (dicamba), purity 99.6% The photodegradation rate constant in spring sunlight will be 0.0138 day-1; t1/2 will be 	. Based on the e at noon (5.83E2 ion rate for natural 50.3 days. by IR
Reliability	 deg latitude at noon is 0.0138 day-1; t1/2 major photodegradation product is CO2. (2) valid with restrictions In the calculation of t1/2, no correction degradation in the dark control was made only slightly influence the results, as ther degradation in the dark control. Except for sterilization of the buffer so measures to guarantee sterility of the sai described. However, as there was hardly the dark control (which was a subsample irradiated), it can be assumed biodegrad 	for the e. However, this will e was hardly any lution, no mples were y any degradation in e of the sample to be
Flag	negligible. Critical study for SIDS endpoint	

3.1.2 STABILITY IN WATER

abiotic
at degree C
at degree C
at degree C
= 0 - 7.6 % after 30 day at pH and degree C
other: essentially OECD 111
1981

ld 1982-69-0 Date 26.12.2001

GLP	:
Test substance	:
Method	: Solutions of 10 ppm and 100 ppm dicamba (1.17% and 0.12%
	14C-dicamba, respectively) in distilled water or aqueous
	buffer solutions of pH 5.0, 7.0 and 9.0 were incubated at 25
	and 35 deg C for 30 days (volume 201 mL, in amber bottles in
	shaking water baths). Acetone concentrations were 0.5%.
	After 1, 7, 14, 21 and 30 days, a duplicate 1-mL sample was
	taken for radioassay and a duplicate 15-mL sample was taken
	for extraction using diethyl ether (at $pH < 1$). Organic and
	aqueous layers were first radioassayed and then analyzed
	using TLC and radioautography detection, followed by
	quantification using LSC. Samples were cochromatographed
	with dicamba and three metabolite reference standards.
Remark	: The test substance for this study was dicamba (acid form) rather than t
	salt. In solution, at specific pH levels it does not matter if the salt or aci
- <i>v</i>	form is used to prepare the solution.
Result	: There was no significant dicamba hydrolysis (i.e. equal to
	or less than 7.6%) at each pH value, both concentrations and
	both temperatures, except for 100 ppm, pH 7.0, 35 deg C at
	t=14, 21 and 30 days in the 100 ppm, when degradation was up
	to 18.5%. Total recovery was only 82.5-83.4% for these
	samples, whereas it was > 95 for all other samples.
	Radioactivity remaining in the aqueous phase after
	extraction was equal to or less than 1% of applied. Three
	unknown degradation products each constituted less than 4%
Testevbeteves	of applied.
Test substance	: CAS 1918-00-9 (14C-dicamba), purity not specified
Conclusion	: Dicamba is stable with slight or no hydrolysis over 30 days
Poliability	under the conditions tested.
Reliability	: (2) valid with restrictions
	1. The fact that at 100 ppm, pH 7.0, 35 deg C up to 18.5% degradation occurred was disregarded because recoveries were
	low. However, no explanation was given for the low
	recoveries. It cannot be excluded that loss of radioactivity
	is due to hydrolysis.
	2. Section "Results and discussion" contained 2 values that
	were not in agreement with values in tables of results.
	3. No measures to guarantee sterility of the samples or to
	exclude oxygen from the solutions were described. However,
	as measured degradation percentages were very low (except at
	100 ppm, pH 7.0, 35 deg C), no significant biotic
	degradation or oxidation can have occurred.
	2. No duplicate samples at any pH.
Flag	3. pH 5.0 was tested, whereas OECD 111 prescribes pH 4.
Flag 25.12.2001	

3.1.3 STABILITY IN SOIL

3.2 MONITORING DATA

ld 1982-69-0 Date 26.12.2001

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

Type Media Air (level I) Water (level I) Soil (level I) Biota (level II / III) Soil (level II / III) Method	fugacity model level III
Year Remark	 2001 The Fugacity was determined using the EQC Level III model as found in EPIWIN 3.05. Estimated values were used for physical constants. Biodegradation was based on the current best estimate for dicamba (from HSDB). Half life in air was determined from the APOWIN program for dicamba (acid) as this would be the likely volatile species. Direct photolysis was not considered in this model. Emissions were restricted to water and soil as it is not volatile. Other parameters used the default values found in EPIWIN.
Result	: Level III Fugacity Model (Full-Output):
	Fugacity (atm) Reaction (kg/hr) Advection (kg/hr) Reaction (percent) Advection (percent) Air 6.47e-014 0.945 0.586 0.0472 0.0293 Water 2.79e-013 638 460 31.9 23 Soil 2.64e-012 900 0 45 0 Sediment 2.23e-013 0.601 0.0347 0.0301 0.00174 Persistence Time: 556 hr r r Reaction Time: 722 hr Advection Time: 7.22 hr Advection Time: 2.41e+003 hr Percent Reacted: 77 Percent Advected: 23 Half-Lives (hr), (based upon user-entry): Air: 43 Water: 500 Sediment: 2000 Advection Times (hr): Air: Air: 1000 Sediment: 5e+004 Sediment: 5e+004
Test substance Reliability Flag	 CAS 1982-69-0 Sodium salt of dicamba (2) valid with restrictions Critical study for SIDS endpoint

ld 1982-69-0 Date 26.12.2001

26.12.2001

3.3.2 DISTRIBUTION

3.4 MODE OF DEGRADATION IN ACTUAL USE

3.5 **BIODEGRADATION**

Type Inoculum Remark	::	aerobic Dicamba has a half life of 31 days with a first-order rate constant of 0.0224/day in a typical midwestern agricultural soil under aerobic conditions. Dicamba is completely mineralized to CO2 under aerobic conditions with 3,6-dichlorosalicylic acid as the only major metabolite. L levels of 2,3-dihydroxy-3,6-dichlorosalicylic acid were detected. Metabounder under anaerobic conditions is similar to that which occurred in aerobic sexcept the rate of dicamba metabolism is reduced under anaerobic conditions. [Krueger JP et al; J Agric Food Chem 39: 995-9 (1991)]. As cited in HSDB update of 8-09-2001.	olism soil
Test substance Conclusion Reliability	:::::::::::::::::::::::::::::::::::::::	AQUATIC FATE: Based on the results of various studies, microbial degradation appears to be the important dicamba removal process in natural water. Photolysis may contribute to dicamba removal from water(Scifres CJ et al; J Environ Qual 2: 306 (1973) As cited in HSDB update of 8-09-2001. CAS 1982-69-0 Sodium salt of dicamba Dicamba (and its soluble salts) biodegrades under both aerobic and anaerobic conditions, it is not know if it can be considered readily biodegradable by the OECD criteria. (2) valid with restrictions	
Flag 26.12.2001	:	Critical study for SIDS endpoint	(2)

3.6 BOD5, COD OR BOD5/COD RATIO

3.7 BIOACCUMULATION

3.8 ADDITIONAL REMARKS

(1)

4. E	Ecotoxicity	1982-69-0 26.12.2001
4.1	ACUTE/PROLONGED TOXICITY TO FISH	
4.2	ACUTE TOXICITY TO AQUATIC INVERTEBRATES	
4.3	TOXICITY TO AQUATIC PLANTS E.G. ALGAE	
4.4	TOXICITY TO MICROORGANISMS E.G. BACTERIA	
4.5.1	CHRONIC TOXICITY TO FISH	
4.5.2	CHRONIC TOXICITY TO AQUATIC INVERTEBRATES	
4.6.1	TOXICITY TO SOIL DWELLING ORGANISMS	
4.6.2	TOXICITY TO TERRESTRIAL PLANTS	
4.6.3	TOXICITY TO OTHER NON-MAMM. TERRESTRIAL SPECIES	
4.7	BIOLOGICAL EFFECTS MONITORING	
4.8	BIOTRANSFORMATION AND KINETICS	
4.9	ADDITIONAL REMARKS	

5. Toxicity

5.1.1 ACUTE ORAL TOXICITY

Type Species Strain Sex Number of animals Vehicle Value Method Year GLP Test substance Method	LD50 rat Sprague-Dawley male/female 10 water > 1000 mg/kg bw other: not specified no other TS TEST ORGANISMS: - Source: Charles River Breeding Laboraties, Kingst York - Age: young adult - Number: 5/sex/dose - Weight at study initiation: 188-269 g - Controls: no ADMINISTRATION: - Doses per time period: single - Volume administered or concentration: 50% (w/v d water); dose volume 10 ml/kg - Post dose observation period: 14 days - food withheld 24 hour pre-dosing till 1 hour after de	listilled
	EXAMINATIONS: gross signs of systemic toxicity an (at least twice daily for 14 days). Gross necropsy on visceral and thoracic cavities. BODY WEIGHT: pre-dosing, days 0, 7 and 13	
Result	STATISTICAL METHOD: Litchfield and Wilcoxon MORTALITY: - Number of deaths at each dose: no deaths CLINICAL SIGNS: on the day of dosing: lethargy, at inactivity, salivation, limbs extended and bodies bed rigid at touch or sound stimulus and slowed respirat loose faeces and urine stains. On day 2 after dosing animals appeared normal.	ame ion,
	NECROPSY FINDINGS: no significant gross pathol SEX-SPECIFIC DIFFERENCES: on day 1, all males mildly lethargic, ataxic and inactive while females or appeared slightly affected.	s appeared
Source Test substance	Notox Hertogenbosch I, 1982-69-0 (sodium salt of Dicamba), puity 20%, ir not indicated	npurities
Conclusion Reliability	LD50 > 5000 mg/kg bw (= > 1000 mg a.i./kg bw) (1) valid without restriction	
	13 / 17	

5. Toxicity	ld 1982-69-0 Date 26.12.2001
09.04.2001	1. The study was conducted in compliance with GLP. However, no compliance statement was present. (5)
5.1.2 ACUTE INHALATI	ΟΝ ΤΟΧΙΟΙΤΥ
5.1.3 ACUTE DERMAL	ΤΟΧΙΟΙΤΥ
Type Species Strain Sex Number of animals Vehicle Value Method Year GLP Test substance Method	 LD50 rabbit New Zealand white male/female 10 physiol. saline > 400 mg/kg bw other: not specified no other TS TEST ORGANISMS: Source: Kings Wheel Rabbitry, Mt. Vernon, Ohio Age: young adult Number: 5/sex/dose Weight at study initiation: 1.65-3.05 kg Controls: no ADMINISTRATION: Area covered: 10% of body surface area Occlusion: yes Vehicle: slightly moistened with physiological saline Doses: 2000 mg/kg bw
Result	 Removal of test substance: wiped with physiological saline EXAMINATIONS: signs of systemic toxicity and mortality (at least twice daily for 14 days). Gross necropsy on visceral and thoracic cavities. BODY WEIGHT: pre-dosing, days 0, 6 and 13 STATISTICAL METHOD: Litchfield and Wilcoxon MORTALITY: Number of deaths at each dose: no deaths CLINICAL SIGNS: Moderate to slight erythema and edema (10/10), a brown cast (10/10), slight scaling (10/10), and slight atonia (1/10). BODY WEIGHTS: changes appeared normal. NECROPSY FINDINGS: no significant findings
Source Test substance	 SEX-SPECIFIC DIFFERENCES: no data Notox Hertogenbosch I, CAS 1982-69-0 (sodium salt of Dicamba), pellets, purity
	14 / 17

5. T	oxicity	ld 1982-69-0 Date 26.12.2001
Rel	iability : (2) valid with rest 1. The skin was a permeability of th 2. The study was no compliance st	/kg bw (= > 400 mg a.i./kg bw)
09.0	04.2001	(4)
5.1.4	ACUTE TOXICITY, OTHER ROUTES	
5.2.1	SKIN IRRITATION	
5.2.2	EYE IRRITATION	
5.3	SENSITIZATION	
5.4	REPEATED DOSE TOXICITY	
5.5	GENETIC TOXICITY 'IN VITRO'	
5.6	GENETIC TOXICITY 'IN VITRO'	
5.7	CARCINOGENITY	
5.8	TOXICITY TO REPRODUCTION	
5.9	DEVELOPMENTAL TOXICITY/TERATOG	
0.0		
5.10	OTHER RELEVANT INFORMATION	
5.11	EXPERIENCE WITH HUMAN EXPOSURE	Ε

6. Refe	Id 1982-69-0 Date 26.12.2001
(1)	EPIWIN v3.05, Syracuse Research Corporation, Syracuse, NY (July 12, 2000)
(2)	Krueger JP et al; J Agric Food Chem 39: 995-9 (1991)]. As cited in HSDB update of 8-09-2001.
(3)	Sandoz Agro, Dicamba: Photodegradation Study in pH 7 Aqueous Solution (1993) (95) unpublished study
(4)	Velsicol Chemical Corporation, Acute Dermal Toxicity Study in Albino Rabbits with 20% sodium salt of Dicamba, 1982 (58)
(5)	Velsicol Chemical Corporation, Acute Oral Toxicity Study in Albino Rats with 20% sodium salt of Dicamba, 1982 (57)
(6)	Velsicol Chemical Corporation, Hydrolysis of 14C-dicamba, 1981

7.	Risk Assessment	ld Date	1982-69-0 26.12.2001
7.1	END POINT SUMMARY		
7.2	HAZARD SUMMARY		
7.3	RISK ASSESSMENT		

IUCLID

Data Set

Existing Chemical CAS No. Generic name	 ID: 68938-79-4 68938-79-4 3,6-Dichloro-2-hydroxybenzoic acid, sodium potassium salt
Producer Related Part Company Creation date	Toxicology and Regulatory Affairs26.12.2001
Substance Related Part Company Creation date	Toxicology and Regulatory Affairs26.12.2001
Memo	:
Printing date Revision date Date of last Update	: 27.12.2001 : : 26.12.2001
Number of Pages	: 14
Chapter (profile) Reliability (profile) Flags (profile)	 Chapter: 1, 2, 3, 4, 5, 7 Reliability: without reliability, 1, 2, 3, 4 Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. General Information	ld 68938-79-4 Date 27.12.2001
1.0.1 OECD AND COMPANY INFORMATION	
1.0.2 LOCATION OF PRODUCTION SITE	
1.0.3 IDENTITY OF RECIPIENTS	
1.1 GENERAL SUBSTANCE INFORMATION	
1.1.0 DETAILS ON TEMPLATE	
1.1.1 SPECTRA	
1.2 SYNONYMS	
1.3 IMPURITIES	
1.4 ADDITIVES	
1.5 QUANTITY	
1.6.1 LABELLING	
1.6.2 CLASSIFICATION	
1.7 USE PATTERN	
1.7.1 TECHNOLOGY PRODUCTION/USE	
1.8 OCCUPATIONAL EXPOSURE LIMIT VALUES	
1.9 SOURCE OF EXPOSURE	
1.10.1 RECOMMENDATIONS/PRECAUTIONARY MEASUR	ES

1. General Information	68938-79-4 27.12.2001
1.10.2 EMERGENCY MEASURES	
1.11 PACKAGING	
1.12 POSSIB. OF RENDERING SUBST. HARMLESS	
1.13 STATEMENTS CONCERNING WASTE	
1.14.1 WATER POLLUTION	
1.14.2 MAJOR ACCIDENT HAZARDS	
1.14.3 AIR POLLUTION	
1.15 ADDITIONAL REMARKS	
1.16 LAST LITERATURE SEARCH	
1.17 REVIEWS	

1.18 LISTINGS E.G. CHEMICAL INVENTORIES

2.1 MELTING POINT

Value	:	ca. 220 ° C	
Sublimation	:		
Method	:	other: calculated	
Year GLP	÷	2001 no	
Test substance	-	10	
Method	:	Estimation using MPBPWIN v1.40 in EPIWIN 3.05	
Result	:	MPBPWIN (v1.40) Program Results:	
		SMILES : c1(CL)ccc(CL)c(OK)c1C(=O)O[Na] CHEM : 3,6-Dichloro-2-hydroxybenzoic acid, sodium, potassium salt MOL FOR: C7 H2 CL2 O3 Na1 K1 MOL WT : 267.09	
		SUMMARY MPBPWIN v1.40	
		Boiling Point: 515.41 deg C (Adapted Stein and Brown Method)	
		Melting Point: 349.84 deg C (Adapted Joback Method) Melting Point: 187.28 deg C (Gold and Ogle Method) Mean Melt Pt : 268.56 deg C (Joback; Gold,Ogle Methods) Selected MP: 219.80 deg C (Weighted Value)	
Test substance	:	3,6-Dichloro-2-hydroxybenzoic acid, sodium, potassium salt CAS 68938- 79-4	
Reliability	:	(2) valid with restrictions	
Flag	:	Critical study for SIDS endpoint	
26.12.2001		(*	1)
2.2 BOILING POINT			
2.3 DENSITY			
2.3.1 GRANULOMETRY			
2.4 VAPOUR PRESSUR	RE		
Value	:	< .000001 at 25° C	
Decomposition Method	:	other (coloulated)	
Method Year		other (calculated) 2001	
GLP	÷	no	
Test substance	:		
Method	:	Estimation using MPBPWIN v1.40 in EPIWIN 3.05	
Result	:	MPBPWIN (v1.40) Program Results:	
		SMILES : c1(CL)ccc(CL)c(OK)c1C(=O)O[Na] CHEM : 3,6-Dichloro-2-hydroxybenzoic acid, sodium, potassium salt	

. Physico-Chen	ical Data Id 689 Date 27.1	
	MOL FOR: C7 H2 CL2 O3 Na1 K1 MOL WT : 267.09	
	Vapor Pressure Estimations (25 deg C): (Using BP: 515.41 deg C (estimated)) (Using MP: 219.80 deg C (estimated)) VP: 7.85E-013 mm Hg (Antoine Method) VP: 9.27E-011 mm Hg (Modified Grain Method) VP: 2.81E-010 mm Hg (Mackay Method) Selected VP: 9.27E-011 mm Hg (Modified Grain Method)	
Test substance	: 3,6-Dichloro-2-hydroxybenzoic acid, sodium, potassium salt 79-4	CAS 68938-
Reliability Flag	(2) valid with restrictionsCritical study for SIDS endpoint	
26.12.2001		(1
.5 PARTITION CO	FFICIENT	
Log pow Method Year GLP Test substance Method Test substance Reliability Flag 26.12.2001	 ca4.15 at 25° C other (calculated) 2001 no Estimation using KOWWIN v1.66 in EPIWIN 3.05 3,6-Dichloro-2-hydroxybenzoic acid, sodium, potassium salt 79-4 (2) valid with restrictions Critical study for SIDS endpoint 	CAS 68938- (1
.6.1 WATER SOLUB	LITY	
Value Qualitative Pka PH Method Year GLP Test substance Method	 ca. 1000 g/l at 25 ° C at 25 ° C at and ° C other: calculated from Ko/w estimate 2001 no Estimation using WSKOW v1.40 in EPIWIN 3.05 	
Result	 SMILES : c1(CL)ccc(CL)c(OK)c1C(=O)O[Na] CHEM : 3,6-Dichloro-2-hydroxybenzoic acid, sodium, potas MOL FOR: C7 H2 CL2 O3 Na1 K1 MOL WT : 267.09 WSKOW v1.40 Results Log Kow (estimated) : -4.15 Log Kow (estimated) : -4.15 Log Kow (experimental): not available from database Log Kow used by Water solubility estimates: -4.15 Equation Used to Make Water Sol estimate: Log S (mol/L) = 0.796 - 0.854 log Kow - 0.00728 MW + Co 	
	(used when Melting Point NOT available) Correction(s): Value	

2. Physico-	-Chemical Data	ld 68938-79-4 Date 27.12.2001
	No Applicable Correction Facto	rs
	Log Water Solubility (in moles Log Water Solubility (in moles	/L): 2.393 /L): 0.573 (Applied Upper Limit)
	Water Solubility at 25 deg C (m	ng/L): 1e+006
Test substan	nce : 3,6-Dichloro-2-hydroxybenzoic a 79-4	cid, sodium, potassium salt CAS 68938-
Reliability Flag	(2) valid with restrictionsCritical study for SIDS endpoint	
26.12.2001		(1)
2.7 FLASH F	POINT	
	POINT	
	LAMMABILITY	
2.8 AUTO FI 2.9 FLAMMA	LAMMABILITY	
2.8 AUTO FI 2.9 FLAMMA 2.10 EXPLOS	LAMMABILITY ABILITY	

3.1.1 PHOTODEGRADATION

Type Light source Light spect. Rel. intensity Indirect photolysis Sensitizer Conc. of sens. Rate constant Degradation Method	 air nm based on Intensity of Sunlight OH 1500000 cm3/(molecule*sec) % after Estimation using APOWIN v1.90 in EPIWIN 3.05
Result	: AOP Program (v1.90) Results: ====================================
Test substance Reliability Flag 26.12.2001	 3,6-Dichloro-2-hydroxybenzoic acid CAS 3401-80-7. This is the form that is expected to be present in air as a vapor. (2) valid with restrictions Critical study for SIDS endpoint (1)
3.1.2 STABILITY IN WAT	ER
Type t1/2 pH4 t1/2 pH7 t1/2 pH9 Deg. Product Method Year GLP Test substance Method Result	 abiotic > 1 year at 25 degree C > 1 year at 25 degree C > 1 year at 25 degree C other: estimated 2001 Estimated on chemical principles based on absence of groups susceptible to hydrolysis. This material has no groups that are susceptible to hydrolysis in the pH 4 to 9 range; therefore, it is considered stable to hydrolysis in surface and groundwater. It is estimated to have a hydrolysis half-life of greater than one year between pH 4 and pH 9.
	The estimation program in EPIWIN has no capability to estimate hydrolysis

8. Environmental	Fate and PathwaysId68938-7Date27.12.20	
	rates for this compound.	
Test substance	: 3,6-Dichloro-2-hydroxybenzoic acid, sodium, potassium salt CAS 79-4	68938-
Reliability	: (2) valid with restrictions	
Flag 26.12.2001	: Critical study for SIDS endpoint	(3
3.1.3 STABILITY IN SO	IL	
3.2 MONITORING DA	ТА	
3.3.1 TRANSPORT BE	TWEEN ENVIRONMENTAL COMPARTMENTS	
Туре	: fugacity model level III	
Media	:	
Air (level I) Water (level I)		
Soil (level I)	:	
Biota (level II / III) Soil (level II / III)		
Method Year	: : : 2001	
Method	The Fugacity was determined using the EQC Level III model as f EPIWIN 3.05. Estimated values were used for physical constant Biodegradation was based on the current best estimate for dican HSDB). Half life in air was determined from the APOWIN progra unionized species as this would be the likely volatile species. Di photolysis was not considered in this model. Emissions were res water and soil as it is not volatile. Other parameters used the def found in EPIWIN.	s. nba (from m for the rect tricted to
Result	: Level III Fugacity Model (Full-Output):	
	Chem Name : 3,6-Dichloro-2-hydroxybenzoic acid, sodium, salt Molecular Wt: 267.09 Henry's LC : 3.26e-017 atm-m3/mole (calc VP/Wsol) Vapor Press : 33.6 mm Hg (Mpbpwin program) Liquid VP : 2.84e+003 mm Hg (super-cooled) Melting Pt : 220 deg C (Mpbpwin program) Log Kow : -4.15 (Kowwin program) Soil Koc : 2.9e-005 (calc by model)	potassium
	Concentration Half-Life Emissions (percent) (hr) (kg/hr) Air 6.52e-020 40 0 Water 56.1 500 1000 Soil 43.8 500 1000 Sediment 0.0978 2e+003 0	
	(atm) (kg/hr) (kg/hr) (percent) (pe Air 6.13e-031 1.16e-017 6.7e-018 5.81e-019 3 Water 3.51e-022 799 576 39.9 2 Soil 1.02e-020 625 0 31.2 0	ection rcent) .35e-019 8.8 .00101
	Persistence Time: 514 hr Reaction Time: 722 hr Advection Time: 1.78e+003 hr Percent Reacted: 71.2 Percent Advected: 28.8	

3. Environmental Fa	ate and Pathways Id 68938-79-4 Date 27.12.2001
	Half-Lives (hr), (based upon user-entry): Air: 40 Water: 500 Soil: 500
	Sediment: 2000 Advection Times (hr): Air: 100 Water: 1000 Sediment: 5e+004
Test substance	: 3,6-Dichloro-2-hydroxybenzoic acid, sodium, potassium salt CAS 68938- 79-4
Reliability Flag 26.12.2001	 (2) valid with restrictions Critical study for SIDS endpoint
3.3.2 DISTRIBUTION	
3.5 BIODEGRADATION	: aerobic
Type Inoculum Test substance	 aerobic 3,6-Dichloro-2-hydroxybenzoic acid, sodium, potassium salt CAS 68938-
Conclusion	 79-4 Dicamba (and its soluble salts) biodegrades under both aerobic and anaerobic conditions. 3,6-Dichloro-2-hydroxybenzoic acid has been identified as an intermediate degradation product; therefore, its soluble salts will also biodegrade. It is not known if it can be considered readily
Reliability	biodegradable by the OECD criteria.(2) valid with restrictions
Flag 26.12.2001	: Critical study for SIDS endpoint (
3.6 BOD5, COD OR BO	D5/COD RATIO
3.7 BIOACCUMULATIO	Ν
3.7 BIOACCUMULATIO 3.8 ADDITIONAL REMA	

4. Ec	cotoxicity	68938-79-4 27.12.2001	
4.1	ACUTE/PROLONGED TOXICITY TO FISH		
4.2	ACUTE TOXICITY TO AQUATIC INVERTEBRATES		
4.3	TOXICITY TO AQUATIC PLANTS E.G. ALGAE		
4.4	TOXICITY TO MICROORGANISMS E.G. BACTERIA		
4.5.1	CHRONIC TOXICITY TO FISH		
4.5.2	CHRONIC TOXICITY TO AQUATIC INVERTEBRATES		
4.6.1	TOXICITY TO SOIL DWELLING ORGANISMS		
4.6.2	TOXICITY TO TERRESTRIAL PLANTS		
4.6.3	TOXICITY TO OTHER NON-MAMM. TERRESTRIAL SPECIES		
4.7	BIOLOGICAL EFFECTS MONITORING		
4.8	BIOTRANSFORMATION AND KINETICS		
4.9	ADDITIONAL REMARKS		

5. Toxicity

5.1.1 ACUTE ORAL TOXICITY

Туре	: LD50
Species	: rat
Strain	:
Sex	:
Number of animals	:
Vehicle	:
Value	: ca. 1562 mg/kg bw
Method	
Year	: 1981
GLP	: no data
Test substance	:
Remark	 This value comes from the literature for 2-hydroxy-3,6-dichlorobenzoic acid which is expected to have similar acute toxicity as its soluble salts.
Test substance	: 3,6-Dichloro-2-hydroxybenzoic acid CAS 3401-80-7.
Reliability	: (2) valid with restrictions
Flag	: Critical study for SIDS endpoint
26.12.2001	(4
	,
1.2 ACUTE INHALAT	ION TOXICITY

5.1.3 ACUTE DERMAL TOXICITY

5.1.4 ACUTE TOXICITY, OTHER ROUTES

5.2.1 SKIN IRRITATION

5.2.2 EYE IRRITATION

5.3 SENSITIZATION

5.4 REPEATED DOSE TOXICITY

5.5 GENETIC TOXICITY 'IN VITRO'

5.6 GENETIC TOXICITY 'IN VITRO'

5.7 CARCINOGENITY

5.8 TOXICITY TO REPRODUCTION

5. Toxicity

ld 68938-79-4 Date 27.12.2001

5.9 DEVELOPMENTAL TOXICITY/TERATOGENICITY

5.10 OTHER RELEVANT INFORMATION

5.11 EXPERIENCE WITH HUMAN EXPOSURE

6. Refer	Id 68938-79-4 Date 27.12.2001
(1)	EPIWIN v3.05, Syracuse Research Corporation, Syracuse, NY (July 12, 2000)
(2)	Krueger JP et al; J Agric Food Chem 39: 995-9 (1991)]. As cited in HSDB update of 8-09- 2001.
(3)	Lyman, W. J. et al. (1990). Handbook of Chemical PropertyEstimation Methods, pp. 7-4, Amer. Chem. Society,Washington, DC
(4)	Pis'ko, GT, Tolstopjatova, GV, and AI Tovstenko AI Comparative study of the toxicity of 2- hydroxy-3,6-dichlorobenzoic acid by various routes of administration Gigiena truda i professional'nye zabolevanija Sep. 1981, No.9, p.55-56.

7. Risk Assessment

ld 68938-79-4 Date 27.12.2001

7.1 END POINT SUMMARY

- 7.2 HAZARD SUMMARY
- 7.3 RISK ASSESSMENT

IUCLID

Data Set

Existing Chemical CAS No. Generic name	 ID: 68938-80-7 68938-80-7 3,6-dichloro-2-hydroxybenzoic acid, dipotassium salt
Producer Related Part Company Creation date	Toxicology and Regulatory Affairs25.12.2001
Substance Related Part Company Creation date	Toxicology and Regulatory Affairs25.12.2001
Memo	:
Printing date Revision date Date of last Update	: 26.12.2001 : : 26.12.2001
Number of Pages	: 15
Chapter (profile) Reliability (profile) Flags (profile)	 Chapter: 1, 2, 3, 4, 5, 7 Reliability: without reliability, 1, 2, 3, 4 Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. 0	eneral Information	68938-80-7 26.12.2001
1.0.1	OECD AND COMPANY INFORMATION	
1.0.2	LOCATION OF PRODUCTION SITE	
1.0.3	IDENTITY OF RECIPIENTS	
1.1	GENERAL SUBSTANCE INFORMATION	
1.1.0	DETAILS ON TEMPLATE	
1.1.1	SPECTRA	
1.2	SYNONYMS	
1.3	IMPURITIES	
1.4	ADDITIVES	
1.5	QUANTITY	
1.6.1	LABELLING	
1.6.2	CLASSIFICATION	
1.7	USE PATTERN	
1.7.1	TECHNOLOGY PRODUCTION/USE	
1.8	OCCUPATIONAL EXPOSURE LIMIT VALUES	
1.9	SOURCE OF EXPOSURE	

1. General Information	68938-80-7 26.12.2001
1.10.1 RECOMMENDATIONS/PRECAUTIONARY MEASURES	
1.10.2 EMERGENCY MEASURES	
1.11 PACKAGING	
1.12 POSSIB. OF RENDERING SUBST. HARMLESS	
1.13 STATEMENTS CONCERNING WASTE	
1.14.1 WATER POLLUTION	
1.14.2 MAJOR ACCIDENT HAZARDS	
1.14.3 AIR POLLUTION	
1.15 ADDITIONAL REMARKS	
1.16 LAST LITERATURE SEARCH	
1.17 REVIEWS	
1.18 LISTINGS E.G. CHEMICAL INVENTORIES	

2. Physico-Chemical Data

2.1 MELTING POINT

Met Yea GLF	limation hod r s t substance hod		ca. 220 ° C other: estimated 2001 no Estimation using MPBPWIN v1.40 in EPIWIN 3.05 MPBPWIN (v1.40) Program Results: ====================================	
			Mean Melt Pt : 268.56 deg C (Joback; Gold,Ogle Methods) Selected MP: 219.80 deg C (Weighted Value)	
Reli Flag	t substance ability J 2.2001		3,6-Dichloro-2-hydroxybenzoic acid, dipotassium salt CAS 68938-80-7 (2) valid with restrictions Critical study for SIDS endpoint	(1)
2.2	BOILING POINT			
2.3	DENSITY			
2.3.1	GRANULOMETRY			
2.4	VAPOUR PRESSUR	E		
Met Yea GLF	omposition hod r b t substance hod		< .0001 hPa at ° C other (calculated) 2001 no Estimation using MPBPWIN v1.40 in EPIWIN 3.05 MPBPWIN (v1.40) Program Results:	

2. Physico-Chemical	Id 68938-80-7 Date 26.12.2001	
	SMILES : c1(CL)ccc(CL)c(OK)c1C(=O)OK CHEM : 3,6-Dichloro-2-hydroxybenzoic acid, dipotassium salt MOL FOR: C7 H2 CL2 O3 K2 MOL WT : 283.19 SUMMARY MPBPWIN v1.40	
	Boiling Point: 515.41 deg C (Adapted Stein and Brown Method) Melting Point: 349.84 deg C (Adapted Joback Method) Melting Point: 187.28 deg C (Gold and Ogle Method) Mean Melt Pt : 268.56 deg C (Joback; Gold,Ogle Methods) Selected MP: 219.80 deg C (Weighted Value)	
	Vapor Pressure Estimations (25 deg C): (Using BP: 515.41 deg C (estimated)) (Using MP: 219.80 deg C (estimated)) VP: 7.85E-013 mm Hg (Antoine Method) VP: 9.27E-011 mm Hg (Modified Grain Method) VP: 2.81E-010 mm Hg (Mackay Method) Selected VP: 9.27E-011 mm Hg (Modified Grain Method)	
Reliability :	3,6-Dichloro-2-hydroxybenzoic acid, dipotassium salt CAS 68938-80-7 (2) valid with restrictions Critical study for SIDS endpoint	(1)
2.5 PARTITION COEFFICIE	INT	

Log pow Method	: ca4.15 at 25° C other (calculated)
Year	: 2001
GLP	: no
Test substance	:
Method	: Estimation using KOWWIN v1.66 in EPIWIN 3.05
Test substance	: 3,6-Dichloro-2-hydroxybenzoic acid, dipotassium salt CAS 68938-80-7
Reliability	: (2) valid with restrictions
Flag	: Critical study for SIDS endpoint
26.12.2001	(1)

2.6.1 WATER SOLUBILITY

Qualitativo	ca. 1000 at 25 ° C
Pka	: at 25 ° C
PH	at and °C
Method	other: estimated
Year	: 2001
GLP	no
Method	Estimation using WSKOW v1.40 in EPIWIN 3.05
	Water Sol from Kow (WSKOW v1.40) Results:

2. Physico-Chemi	cal Data Id 68938-80-7 Date 26.12.2001	
	Water Sol: 1e+006 mg/L	
	SMILES : c1(CL)ccc(CL)c(OK)c1C(=O)OK CHEM : 3,6-Dichloro-2-hydroxybenzoic acid, dipotassium salt MOL FOR: C7 H2 CL2 O3 K2 MOL WT : 283.19 WSKOW v1.40 Results	
	Log Kow (estimated) : -4.15 Log Kow (experimental): not available from database Log Kow used by Water solubility estimates: -4.15	
	Equation Used to Make Water Sol estimate: Log S (mol/L) = 0.796 - 0.854 log Kow - 0.00728 MW + Correction (used when Melting Point NOT available)	
	Correction(s): Value	
	No Applicable Correction Factors	
	Log Water Solubility (in moles/L) : 2.275 Log Water Solubility (in moles/L) : 0.548 (Applied Upper Limit) Water Solubility at 25 deg C (mg/L): 1e+006	
Test substance Reliability Flag	 3,6-Dichloro-2-hydroxybenzoic acid, dipotassium salt CAS 68938-80-7 (2) valid with restrictions Critical study for SIDS endpoint 	7
25.12.2001		(1)
2.6.2 SURFACE TENSION	Ν	
2.7 FLASH POINT		
2.8 AUTO FLAMMABIL	ΙΤΥ	
2.9 FLAMMABILITY		
2.10 EXPLOSIVE PROPE		
2.10 EXPLOSIVE PROPE		
2.11 OXIDIZING PROPE	RTIES	
2.12 ADDITIONAL REMA	ARKS	

3. Environmental Fate and Pathways

3.1.1 PHOTODEGRADATION

Type Light source Light spect. Rel. intensity Indirect photolysis Sensitizer Conc. of sens. Rate constant Degradation Method Result	air nm based on Intensity of Sunlight OH 1500000 molecule/cm3 cm3/(molecule*sec) % after Estimation using APOWIN v1.90 in EPIWIN 3.05 AOP Program (v1.90) Results:	
	SMILES : c1(CL)ccc(CL)c(O)c1C(=O)O CHEM : 3,6-Dichloro-2-hydroxybenzoic acid MOL FOR: C7 H4 CL2 O3 MOL WT : 207.01 	ec :
Test substance Reliability	3,6-Dichloro-2-hydroxybenzoic acid. CAS 3401-80-7 This is the form of test material that would be present in air as (2) valid with restrictions	a vapor.
Flag 25.12.2001	Critical study for SIDS endpoint	(1)

3.1.2 STABILITY IN WATER

Туре	: abiotic
t1/2 pH4	: > 1 year at 25 degree C
t1/2 pH7	: > 1 year at 25 degree C
t1/2 pH9	: > 1 year at 25 degree C
Deg. Product	
Method	: other: estimated
Year	: 2001
GLP	: no
Test substance	:
Method	 Estimated on chemical principles based on absence of groups susceptible to hydrolysis
Result	:
	This material has no groups that are susceptible to hydrolysis in the pH 4 to

9 range; therefore, it is considered stable to hydrolysis in surface and groundwater. It is estimated to have a hydrolysis half-life of greater than one year between pH 4 and pH 9. The estimation program in EPIWIN has no capability to estimate hydrolysis rates for this compound. Test substance : 3,6-Dichloro-2-hydroxybenzoic acid, dipotassium salt CAS 68938-80-7 Reliability : (2) valid with restrictions Flag : Critical study for SIDS endpoint 3.1.3 STABILITY IN SOIL 3.2 MONITORING DATA	3. Ei	nvironmenta	Fate and Pathways	ld 68938-80-7 Date 26.12.2001
	Relia Flag	ability	 groundwater. It is estimated to have a hydrolys one year between pH 4 and pH 9. The estimation program in EPIWIN has no cap rates for this compound. 3,6-Dichloro-2-hydroxybenzoic acid, dipotassis (2) valid with restrictions 	sis half-life of greater than bability to estimate hydrolysis um salt CAS 68938-80-7
3.2 MONITORING DATA	3.1.3	STABILITY IN SO	L	
3.2 WONTORING DATA	o 0			
	J.Z	WONTOKING DA		
	3.3.1	TRANSPORT BET	WEEN ENVIRONMENTAL COMPARTMENTS	

Type	fugacity model level III
Media	2001
Air (level I)	The Fugacity was determined using the EQC Level III model as found in
Water (level I)	EPIWIN 3.05. Estimated values were used for physical constants.
Soil (level I)	Biodegradation was based on the current best estimate for dicamba (from
Biota (level II / III)	HSDB). Half life in air was determined from the APOWIN program for
Soil (level II / III)	dicamba (acid) as this would be the likely volatile species. Direct
Method	photolysis was not considered in this model. Emissions were restricted to
Year	water and soil as it is not volatile. Other parameters used the default values
Method	found in EPIWIN.
Result	: Level III Fugacity Model (Full-Output): The second sec

3. Environment	3. Environmental Fate and Pathways			ld 68938-80-7 Date 26.12.2001		
	Air 6.5e-031 1.41e-0 Water 3.51e-022 799 Soil 1.02e-020 625 Sediment 3.07e-022 0.348	15 8.74e-016 576 0 0.0201	7.04e-017 39.9 31.2 0.0174	4.37e-017 28.8 0 0.00101		
	Persistence Time: 514 hr Reaction Time: 722 hr Advection Time: 1.78e+003 Percent Reacted: 71.2 Percent Advected: 28.8	hr				
	Half-Lives (hr), (based upo Air: 43 Water: 500 Soil: 500 Sediment: 2000	ı user-entry):				
	Advection Times (hr): Air: 100 Water: 1000 Sediment: 5e+004					
Test substance Reliability Flag 26.12.2001	 3,6-Dichloro-2-hydroxybenzoic a (2) valid with restrictions Critical study for SIDS endpoint 	cid, dipotassiun	n salt CAS 68	3938-80-7 (1)		
3.3.2 DISTRIBUTION						
3.4 MODE OF DEG	RADATION IN ACTUAL USE					
3.5 BIODEGRADAT	ION					
Type Inoculum Tost substanco	: aerobic : : 3.6 Dichloro 3 hydroxybonzoic c	oid dipotoooius		2039 90 7		
Test substance Conclusion	 3,6-Dichloro-2-hydroxybenzoic a Dicamba (and its soluble salts) b 	·				

anaerobic conditions. 3,6-Dichloro-2-hydroxybenzoic acid has been identified as an intermediate degradation product; therefore, its soluble salts will also biodegrade. It is not known if it can be considered readily biodegradable by the OECD criteria.

Reliability: (2) valid with restrictionsFlag: Critical study for SIDS endpoint26.12.2001

(2)

3.6 BOD5, COD OR BOD5/COD RATIO

3.7 BIOACCUMULATION

3. Environmental Fate and Pathways

ld 68938-80-7 Date 26.12.2001

3.8 ADDITIONAL REMARKS

	cotoxicity	68938-80-7 26.12.2001
4.1	ACUTE/PROLONGED TOXICITY TO FISH	
4.2	ACUTE TOXICITY TO AQUATIC INVERTEBRATES	
4.3	TOXICITY TO AQUATIC PLANTS E.G. ALGAE	
4.4	TOXICITY TO MICROORGANISMS E.G. BACTERIA	
4.5.1	CHRONIC TOXICITY TO FISH	
4.5.2	CHRONIC TOXICITY TO AQUATIC INVERTEBRATES	
4.6.1	TOXICITY TO SOIL DWELLING ORGANISMS	
4.6.2	TOXICITY TO TERRESTRIAL PLANTS	
4.6.3	TOXICITY TO OTHER NON-MAMM. TERRESTRIAL SPECIES	
4.7	BIOLOGICAL EFFECTS MONITORING	
4.8	BIOTRANSFORMATION AND KINETICS	
	ADDITIONAL REMARKS	

5. Toxicity

5.1.1 ACUTE ORAL TOXICITY

Type Species Strain Sex Number of animals Vehicle Value Method Year GLP Test substance Remark Test substance Reliability Flag	 LD50 rat ca. 1562 ml/kg bw 1981 no data This value comes from the literature for 2-hydroxy-3,6-dichlorobenzoic ac which is expected to have similar acute toxicity as its soluble salts. 3,6-Dichloro-2-hydroxybenzoic acid. CAS 3401-80-7 (2) valid with restrictions Critical study for SIDS endpoint 	id
26.12.2001		(4)
5.1.2 ACUTE INHALATIO	Ν ΤΟΧΙΟΙΤΥ	
5.1.3 ACUTE DERMAL TO	OXICITY	
5.1.4 ACUTE TOXICITY, C	OTHER ROUTES	

5.2.1 SKIN IRRITATION

5.2.2 EYE IRRITATION

- 5.3 SENSITIZATION
- 5.4 REPEATED DOSE TOXICITY

5.5 GENETIC TOXICITY 'IN VITRO'

5.6 GENETIC TOXICITY 'IN VITRO'

5. 1	Foxicity ^{Id}	68938-80-7
		26.12.2001
5.7	CARCINOGENITY	
5.8	TOXICITY TO REPRODUCTION	
5.9	DEVELOPMENTAL TOXICITY/TERATOGENICITY	
5.10	OTHER RELEVANT INFORMATION	

5.11 EXPERIENCE WITH HUMAN EXPOSURE

6. References Id 68938-80-7 Date 26.12.2001 (1) EPIWIN v3.05, Syracuse Research Corporation, Syracuse, NY (July 12, 2000) (2) Krueger JP et al; J Agric Food Chem 39: 995-9 (1991)]. As cited in HSDB update of 8-09-2001. (3) Lyman, W. J. et al. (1990). Handbook of Chemical PropertyEstimation Methods, pp. 7-4, Amer. Chem. Society,Washington, DC (4) Pis'ko, GT, Tolstopjatova, GV, and Al Tovstenko Al Comparative study of the toxicity of 2-hydroxy-3,6-dichlorobenzoic acid by various routes of administration Gigiena truda i professional'nye zabolevanija Sep. 1981, No.9, p.55-56.

7. Risk Assessment		68938-80-7 26.12.2001
7.1	END POINT SUMMARY	
7.2	HAZARD SUMMARY	
7.3	RISK ASSESSMENT	

IUCLID

Data Set

Existing Chemical CAS No. Molecular Formula Generic name	: 583-78-8
Producer Related Part Company Creation date	Toxicology and Regulatory Affairs26.12.2001
Substance Related Part Company Creation date	: Toxicology and Regulatory Affairs : 26.12.2001
Memo	:
Printing date Revision date Date of last Update	: 26.12.2001 : : 26.12.2001
Number of Pages	: 26
Chapter (profile) Reliability (profile) Flags (profile)	 Chapter: 1, 2, 3, 4, 5, 7 Reliability: without reliability, 1, 2, 3, 4 Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. G	eneral Information	583-78-8 26.12.2001
1.0.1	OECD AND COMPANY INFORMATION	
1.0.2	LOCATION OF PRODUCTION SITE	
1.0.3	IDENTITY OF RECIPIENTS	
1.1	GENERAL SUBSTANCE INFORMATION	
1.1.0	DETAILS ON TEMPLATE	
1.1.1	SPECTRA	
1.2	SYNONYMS	
1.3	IMPURITIES	
1.4	ADDITIVES	
1.5	QUANTITY	
1.6.1	LABELLING	
1.6.2	CLASSIFICATION	
1.7	USE PATTERN	
1.7.1	TECHNOLOGY PRODUCTION/USE	
1.8	OCCUPATIONAL EXPOSURE LIMIT VALUES	
1.9	SOURCE OF EXPOSURE	

1. 0	General Information	583-78-8 26.12.2001
1.10.1	RECOMMENDATIONS/PRECAUTIONARY MEASURES	
1.10.2		
1.11	PACKAGING	
1.12	POSSIB. OF RENDERING SUBST. HARMLESS	
1.13	STATEMENTS CONCERNING WASTE	
1.14.1	WATER POLLUTION	
1.14.2	MAJOR ACCIDENT HAZARDS	
1.14.3		
1.15	ADDITIONAL REMARKS	
1.16	LAST LITERATURE SEARCH	
1.17	REVIEWS	
1.18	LISTINGS E.G. CHEMICAL INVENTORIES	

2. Physico-Chemical Data

2.1 MELTING POINT

Value	: 59 °C	
Sublimation	:	
Method	: other: no data	
Year	:	
GLP	: no data	
Test substance	:	
Test substance	: CAS 583-78-8 (2,5-dichlorophenol), purity not specified	
Reliability	: (2) valid with restrictions Handbook data	
Flag	: Critical study for SIDS endpoint	
26.12.2001		(13)

2.2 BOILING POINT

Value	: 211 °C at	
Decomposition	:	
Method	: other: no data	
Year	:	
GLP	: no data	
Test substance	:	
Test substance	: CAS 583-78-8 (2,5-dichlorophenol), purity not specified	
Reliability	: (2) valid with restrictions	
	Handbook data	
Flag	: Critical study for SIDS endpoint	
26.12.2001		(13)

2.3 DENSITY

2.3.1 GRANULOMETRY

2.4 VAPOUR PRESSURE

Value	: = .08 hPa at 25° C	
Decomposition	:	
Method		
Year	:	
GLP	: no data	
Test substance	:	
Remark	: Supported by EPIWIN calculated value value of 0.06 hPa	
Reliability	: (2) valid with restrictions	
-	Literature value	
Flag	: Critical study for SIDS endpoint	
26.12.2001	· ·	(4)

2. Physico-Chemic	al Data		33-78-8 3.12.2001
2.5 PARTITION COEFFIC	CIENT		
Remark	 = 3.06 at 25° C Supported by EPIWIN calculated value value of 2.8 2,5-dichlorophenol, CAS 583-78-8 (2) valid with restrictions Literature value Critical study for SIDS endpoint 	30	(6)
Value Qualitative Pka PH Method Year GLP Test substance Remark	 = 2000 mg/l at 25 ° C other: slightly soluble at 25 ° C at and ° C other: no data no data Remarks: Secondary literature. No source or method of determination is given. There is an experimental database match given in EPIWIN 3.05 Experimental Water Solubility Database Match: Name : 2,5-DICHLOROPHENOL CAS Num : 000583-78-8 Exp WSol : 2000 mg/L (25 deg C) Exp Ref : CHEM INSPECT TEST INST (1992) 	WSKOW	v1.40 in
Test substance Reliability Flag 26.12.2001	 CAS 583-78-8 (2,5-dichlorophenol), purity not specified (4) not assignable secondary literature (remark 1) Critical study for SIDS endpoint 	cified	(3) (5)
2.6.2 SURFACE TENSION			
2.7 FLASH POINT			
2.8 AUTO FLAMMABILIT	Y		
2.9 FLAMMABILITY			

2. Physico-Chemical Data Id 583-78-8 Date 26.12.2001	
2.10 EXPLOSIVE PROPERTIES	
2.11 OXIDIZING PROPERTIES	
2.12 ADDITIONAL REMARKS	

3. Environmental Fate and Pathways

3.1.1 PHOTODEGRADATION

Type Light source Light spect. Rel. intensity Indirect photolysis Sensitizer Conc. of sens. Rate constant Degradation Deg. Product Method Year GLP Test substance Method Result	air nm based on Intensity of Sunlight OH 1500000 molecule/cm3 ca00000000007 cm3/(molecule*sec) = 50 % after 18 hour(s) other (calculated) 2001 no Estimation using APOWIN v1.90 in EPIWIN 3.05 AOP Program (v1.90) Results: ====================================	
Test substance Reliability Flag 26.12.2001	2,5-dichlorophenol, CAS 583-78-8(2) valid with restrictionsCritical study for SIDS endpoint	(5)

3.1.2 STABILITY IN WATER

Туре	:	abiotic
t1/2 pH4	:	> 1 year at 25 degree C
t1/2 pH7	:	> 1 year at 25 degree C
t1/2 pH9	:	> 1 year at 25 degree C
Deg. Product	:	
Method	:	
Year	:	2001
GLP	:	
Test substance	:	

3. Environment	al Fate and Pathways	ld 583-78-8 Date 26.12.2001
Method	: Estimated on chemical principles to hydrolysis	based on absence of groups susceptible
Remark		N has no capability to estimate hydrolysis
Result	9 range; therefore, it is considered	re susceptible to hydrolysis in the pH 4 to I stable to hydrolysis in surface and ve a hydrolysis half-life of greater than
Test substance	: 2,5-dichlorophenol, CAS 583-78-8	3
Reliability	: (2) valid with restrictions	
Flag	: Critical study for SIDS endpoint	
26.12.2001		(14)

3.1.3 STABILITY IN SOIL

3.2 MONITORING DATA

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

Type Media Air (level I) Water (level I) Soil (level I) Biota (level II / III) Soil (level II / III) Method Year Method	 fugacity model level III 2001 The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined using the EQC Level III model as found in The Fugacity was determined u	
Result	EPIWIN 3.05. Measured values were used for physical constants. Biodegradation was based on the current best estimate (from HS DB). Hall life in air was determined from the APOWIN program. Direct photolysis was not considered in this model. Other parameters used the default values found in EPIWIN : Level III Fugacity Model (Full-Output): Chem Name : 2, 5-Di chl orophenol Mol ecul ar Wt: 163	f
	Molecular w.103Henry's LC : 4.77e-007 atm-m3/mole (Henrywin program)Vapor Press : 0.06 mm Hg (user-entered)Liquid VP : 0.13 mm Hg (super-cooled)Melting Pt : 59 deg C (user-entered)Log Kow : 3.06 (user-entered)Soil Koc : 471 (calc by model)Concentration Half-Life Emissions (percent) (hr) (kg/hr)Air 4.4724Air 4.471000Water 31.5125Soil 63.9200Soil 63.9200Sediment 0.136400	
	FugacityReactionAdvectionReactionAdvection(atm)(kg/hr)(kg/hr)(percent)(percent)Air3.34e-01164422321.57.43	
	0 / 00	

	al Fate and Pathways	6			83-78-8 6.12.2001
	Water 2. 3e-012 Soil 4. 47e-012 Sediment 4. 03e-013	1. 1e+003 0	57)). 0136	29 36. 8 0. 0392	5. 23 0 0. 000452
		190 hr 1.31e+003 hr 87.3			
	Half-Lives (hr), (Air: 24 Water: 125 Soil: 200 Sediment: 400	based upon user	r-entry):		
	Advection Times (h Air: 100 Water: 1000 Sediment: 5e+00				
Test substance Reliability Flag 26.12.2001	 2,5-dichlorophenol, CA (2) valid with restriction Critical study for SIDS 	าร			(5)
3.3.2 DISTRIBUTION					
	RADATION IN ACTUAL USE				
3.4 MODE OF DEGI 3.5 BIODEGRADAT Type Inoculum Contact time Degradation		oted			
3.4 MODE OF DEGI 3.5 BIODEGRADAT Type Inoculum Contact time	FION : aerobic : activated sludge, adap : 4 day	oted			
3.4 MODE OF DEGI 3.5 BIODEGRADAT Type Inoculum Contact time Degradation Result Deg. Product Method Year GLP	FION : aerobic : activated sludge, adap : 4 day : = 52 % after 4 day : : : : 1966	ed to undergo 54 annot be seterm	ined if this	test substa	

[USEPA; Ambient Water Quality Criteria Doc: Chlorinated Phenols p.C-29

3. Environme	ntal Fate and Pathways	ld 583-78-8 Date 26.12.2001
Reliability Flag 26.12.2001	 (1980) EPA 440/5-80-032]**PEER RE of 8-09-2001 (2) valid with restrictions Critical study for SIDS endpoint 	VIEWED** As cited in HSDB update (8)
3.6 BOD5, COD	OR BOD5/COD RATIO	
3.7 BIOACCUMU	ILATION	
3.8 ADDITIONAL	REMARKS	

4. L	cotoxicity	583-78-8 26.12.2001
4.1	ACUTE/PROLONGED TOXICITY TO FISH	
4.2	ACUTE TOXICITY TO AQUATIC INVERTEBRATES	
4.3	TOXICITY TO AQUATIC PLANTS E.G. ALGAE	
4.4	TOXICITY TO MICROORGANISMS E.G. BACTERIA	
4.5.1	CHRONIC TOXICITY TO FISH	
4.5.2	CHRONIC TOXICITY TO AQUATIC INVERTEBRATES	
4.6.1	TOXICITY TO SOIL DWELLING ORGANISMS	
4.6.2	TOXICITY TO TERRESTRIAL PLANTS	
4.6.3	TOXICITY TO OTHER NON-MAMM. TERRESTRIAL SPECIES	
4.7	BIOLOGICAL EFFECTS MONITORING	
4.8	BIOTRANSFORMATION AND KINETICS	
4.9	ADDITIONAL REMARKS	

5. Toxicity

5.1.1 ACUTE ORAL TOXICITY

Type Species Strain Sex Number of animals Vehicle Value Method Year GLP Test substance Method	 LD50 rat Wistar female 10 other: sesame oil = 2475 mg/kg bw other: not specified no other TS TEST ORGANISMS: Source:no data Age: no data Number:10/dose Weight at study initiation: 80-97 g Controls: no ADMINISTRATION: Doses: 1600, 2500, 4000 mg/kg bw Doses per time period: single (gavage) Volume administered not indicated Post dose observation period: 14 days food withheld 16 hr before to 2 hr after dosing
	EXAMINATIONS: Necropsy of all animals with macroscopic examination. Body weight (pre-dosing, days 7 and 14)
Result	 STATISTICAL METHOD: probit (Linder and Weber) MORTALITY: Number of deaths at each dose: 1600, 2500 and 4000 mg/kg bw 1/10, 4/10 and 10/10 Time of death: deaths within 24 hours after dosing
	CLINICAL SIGNS: in dying animals: excessive breathing, equilibrium disturbance and tremor, moreover tonic clonic spasms in the ventral region. In the highest dose, these signs occurred immediately after dosing.
	NECROPSY FINDINGS: No abnormal findings were noted in surviving animals. In decendents: clear dilated bloodvessels on the intestines
	BODY WEIGHT: normal body weight gain in surviving animals No data on decendents
Source Test substance	 POTENTIAL TARGET ORGANS: intestines Notox Hertogenbosch II, CAS 583-78-8 (2,5-Dichlorphenol), purity not indicated, cristalline form
Conclusion Reliability	 LD50 2475 mg/kg bw (95% CI 2101-2916 mg/kg bw) (2) valid with restrictions 12 / 26

5. Toxicity	ld 583-78-8	
	Date 26.12.2001	
	 The information was essentially confined to what is included in the current summary only females were tested no individual data were present 	
02.04.2001		(7)
Type Species Strain Sex Number of animals Vehicle Value Method Year GLP Test substance Method	 LD50 mouse other: CD-1 ICR male/female 100 other: corn oil 946 - 1600 ml/kg bw other: not indicated ino data other TS TEST ORGANISMS: Age: adult Number: 10 males, 10 females per dosage level Weight at study initiation: Controls: no data 	
	ADMINISTRATION: - by gavage - Doses: 5 levels, levels not indicated - Volume administered or concentration: 10 mL/kg body weight - food withheld for 2 h after dosing - Post dose observation period: 14 days EXAMINATIONS: behavior and visible health, time of death, necropsy of animals that died during the test	
Remark	 STATISTICAL METHOD: Log probit analysisof Finney; Litchfield, Wilcoxon. Remarks: Remarks: Remarks: The article contains a summary rather than a full report. Information is essentially confined to what is mentioned in this summary. Especially no detailed results are given. 	
Result	 LD50 male: 1600 mg/kg bw (confidence limits: 1233-2075 mg/kg bw); LD50 female: 946 mg/kg bw (confidence limits: 623-1438 mg/kg bw) 	
Source Test substance Reliability	 Notox Hertogenbosch II, CAS 583-78-8 (2,5-dichlorophenol), purity 98% (4) not assignable secondary literature (remark 1) 	
Flag 15.03.2001	: Critical study for SIDS endpoint	(2)

5.1.2 ACUTE INHALATION TOXICITY

Туре	:	LC50
Species	:	rat
Strain	:	other: Spartan
Sex	:	male/female

5. Toxicity	ld 583-7 Date 26.12	
Number of animals	: 10	
Vehicle	:	
Exposure time	: 4 hour(s)	
Value Method	: > 185000 mg/m ³	
Year		
GLP	: no	
Test substance	: other TS	
Method	: TEST ORGANISMS:	
	- Source: no data	
	- Age: no data	
	- Weight at study initiation: 216-243 g	
	- Number of animals: 10 (5 male, 5 female)	
	ADMINISTRATION:	
	- Type of exposure: inhalation (whole body)	
	- Exposure duration: 4 hours	
	- Concentrations: 50000 mg/m3; 185000 mg/m3	
	- Particle size: no data	
	- Type or preparation of particles: no data	
	- Air changes: no data	
	EXAMINATIONS: clinical signs during and immediately	
	following exposure; macroscopy	
Result	: MORTALITY:	
	- Number of deaths at each dose:50000 mg/m3: none; 185000	
	mg/m3: 2 (females)	
	- Time of death: during exposure (both)	
	CLINICAL SIGNS: 50000 mg/m3, (all rats): increased/decreased	l
	motor activity, eye squint, erythema, lacrimation,	
	salivation, clear nasal discharge, ocular and nasal	
	porphyrin discharge, slight dispnoea. The symptoms	
	disappeared in all rats 24 hours after exposure	
	185000 mg/m3, (all rats): The same symptoms as at 50000	
	mg/m3, with addition of marked dispnoea, corneal opacity, ataxia, sedation and body jerking. The symptoms disappeared	
	72 hours after exposure (one rat exhibiting nasal porphyrin	
	discharge at day 10)	
	NECROPSY FINDINGS: congested lungs and liver, slight corne	al
_	opacity (in the animals that died)	
Source	: Notox Hertogenbosch	
Test condition	: II, CAS 583-78-8 (2,5-dichlorophenol), purity not specified	
Reliability	(2) valid with restrictions1. The information included in the report was confined to	
	what is included in the current summary	
	2. No information on body weight was presented	
09.04.2001		(*

5.1.3 ACUTE DERMAL TOXICITY

Type Species Strain Sex	: LD50 : rabbit : New Zealand white
Strain	: New Zealand white
Sex	: male/female

5. Toxicity		583-78-8 26.12.2001
Number of animals	: 4	
Vehicle		
Value	: > 8000 mg/kg bw	
Method	:	
Year	:	
GLP	: no	
Test substance	: other TS	
Method	: TEST ORGANISMS:	
	- Source: no data	
	- Age: no data	
	- Weight at study initiation: 2387-2970 g	
	- Controls: no data	
	ADMINISTRATION:	
	- Area covered: no data	
	- Occlusion: yes	
	- Vehicle: not applicable (applied as powder)	
	- Doses: 1000, 2000, 4000 and 8000 mg/kg bw	
	- Removal of test substance: washed with tepid tap water	
	······································	
	EXAMINATIONS: observations for mortality during 14 day	s;
	body weight at start and day 14	
	STATISTICAL METHOD: Thompson, W.R., Bact. Rev.: 11	5-145
	1947	0 1 10,
Result	: MORTALITY:	
	- Number of deaths at each dose: none	
	CLINICAL SIGNS: no data	
	BODY WEIGHT: decreased body weight in both females a	at 2000
	mg/kg bw , in one male and one female at 4000 mg/kg bw	
	in males at 8000 mg/kg	
Source	: Notox Hertogenbosch	
Test substance	: II, CAS 583-78-8 (2,5-dichlorophenol), purity not specified	
Reliability	: (2) valid with restrictions	
	1. The information included in the report was confined to	
	what is included in the current summary	
	2. Only 4 animals per group (animals not of one sex only),	
	of which one underwent skin abrasion (OECD 402: at leas	t
	five animals per dosage group, no abrading of the skin)	
09.04.2001	3. The size of the application area was not indicated	(1
03.04.2001		(1

5.2.1 SKIN IRRITATION

5.2.2 EYE IRRITATION

5. Toxicity

ld 583-78-8 Date 26.12.2001

5.3 SENSITIZATION

5.4 REPEATED DOSE TOXICITY

Species Sex Strain Route of admin. Exposure period Frequency of treatment Post obs. period Doses Control group LOAEL Method Year GLP Test substance Method	 rat male/female Sprague-Dawley inhalation 4 weeks 5 days/week, 6 hours/day 1 0.1, 0.3 and 1.0 mg/L yes, concurrent no treatment = .1 mg/l other: not indicated no other TS TEST ORGANISMS Age: 8 weeks Weight at study initiation: males 206-230 g,females 192-224 g Number of animals: 10/sex/treatment
	 Exposure period: 4 weeks, 6 hours/day, 5 days/week Route of administration: inhalation (whole body) Doses: 0.1, 0.3 and 1.0 mg/L Particle size: not applicable (vapour) Air changes: 2-16/hour
	CLINICAL OBSERVATIONS AND FREQUENCY: - Mortality/clinical signs: twice daily - Body weight: pre-treatment and weekly thereafter - Haematology: after 4 weeks: haematocrit, Hb, erythrocyte count, (differential) leucocyte count, MCV, MCH(C). - Biochemistry: after 4 weeks: glucose, BUN, ALP, ALAT, ASAT - Urinalysis: after 4 weeks accoridng to OECD 407
	ORGANS EXAMINED AT NECROPSY (MACROSCOPIC AND MICROSCOPIC): - Organ weights: liver, spleen, kidneys, heart, lungs, brain, adrenals, thyroid, pituitary - Macroscopic: all tissues (see microscopy) from all animals - Microscopic: from controls and high dose group: nasal turbinates, trachea, lung, spleen, pancreas, stomach, duodenum, uterus, prostate, kidneys, urinary bladder, ovaries, testes, bone marrow, heart, mediastinal and mesenteric lymphnodes, colon, liver, adrenals, olfactory bulb, thyroid, parathyroid, brain , eye, pituitary, gross lesions from other dose groups: nasal turbinates, trachea, lung,
	liver

5. Toxicity	ld 583-78-8
	Date 26.12.2001
	ANALYSES: - Method: nominal concentrations by weighing of the vaporisation flask before and after exposure
Result	 STATISTICAL METHODS: ANOVA, Bartlett's test, Dunnett's test ANALYSES: Nominal concentration: at 0.1, 0.3 and 1.0 mg/L 0.07-0.28, 0.07-1.09 and 0.45-1.36 mg/L respectively.
	TOXIC RESPONSE/EFFECTS BY DOSE LEVEL: - Mortality: none - Clinical signs: Nasal irritation with or without discharge in all treatment groups and controls Ocular irritation and discharge in all treatment groups
	Salivation in 8 males and 4 females at 0.3 mg/L and in 7 males and 7 females at 1.0 mg/L Dyspnoea in one male and 7 females at 0.3 mg/L Incidental findings respiratory distress, skin irritation, cloudy spots on eyes, decreased activity and soaked abdomen - Body weight gain: decreased at 0.3 mg/L during week 2-4 and at 1.0 during week 1-4.
	 Haematology: Hb increased at the high dose group, No. of leucocytes increased in females at 0.3 and 1.0 mg/L Clinical chemistry: ASAT increased in high dose males and females Urinalysis: no treatment related effects
	- Organ weights: Decreased absolute liver and brain weight in males at 0.3 and 1.0 mg/L Increased relative lung weight in females at 1.0 mg/L Decreased absolute heart weight in males at 0.3 mg/L
	Increased relative kidney weight in all treated males - Gross pathology: Brown cyanotic/discolored areas, foci and atelectasis in the lungs were seen in 1-2 animals/sex/treatment and in
	controls. At 1.0 mg/L the incidence was slightly increased in females. Other incidental effects included haemorrhagic/hyperemic lymphnodes, effects on stomach mucosa, pale/discolored liver areas/foci and haemorrhagic foci and discoloration of the kidneys. - Histopathology:
	Inflammatory cell and lymphocyte infiltrate, macrophage aggregation and septal fibrosis in the lungs of all treated animals Inflammation of the nasal cavity (mucosa) in animals at 1.0 mg/L
	Lymphocytic infiltrate, inflammation, foci and necrosis of the liver in treated and control animals. The incidence in control animals was slightly lower (9/20) compared to treatd animals (14-16/20).
	STATISTICAL RESULTS: The effects on body weight, organ weight and bloodparameters were statistically significant. None of the effects showed a clear concentration-response
	17 / 26

5. Toxicity	ld 583-78-8
-	Date 26.12.2001
	relationship.
Source	: Notox Hertogenbosch
Test substance	: II, CAS 583-78-8 (2,5-dichlorophenol), purity not specified
Conclusion	: LOAEL 0.1 mg/L based on liver effects.
	Other effects seen were related to a weight decrease (organ
	weights) or could be attributed to irritant properties of
	the test substance (effects in the respiratory tract).
Reliability	: (2) valid with restrictions
	1 No analyses for actual concentration, homogeneity and
	stability were performed.
	2 The effects on organ weights are expected to be related to the decreased body weight.
	3 No blood clotting parameters were determined
09.04.2001	(12)
Species	: rabbit
Sex	: male/female
Strain	: New Zealand white
Route of admin.	: dermal
Exposure period	: 21 days
Frequency of treatment	: 5 days/week, 6 hours/day
Post obs. period	
Doses	1.0, 10 and 100 mg/kg bw
Control group	: other: distilled water
Method	: other: not indicated
Year	:
GLP	: no
Test substance	: other TS
Method	: TEST ORGANISMS
	- Weight at study initiation: 2171-2921 g (males), 2028-3146
	g (females)
	- Number of animals: 4/sex/treatment
	- Source: HARE Rabbits Research, Hewitt, NJ
	ADMINISTRATION / EXPOSURE
	- Exposure period: 21 days, 5 days/week, 6 hours/day
	- Route of administration: dermal
	- Doses: 1.0, 10.0 and 100 mg/kg bw; water control
	- Vehicle: not applicable (substance was melted at 60 C
	before application)
	- Total volume applied: =<0.1 mL/kg
	- Area treated: 10% of body surface (at 1.0 and 10 mg/kg bw
	every day another area was treated)
	 Occlusion: no (a collar was applied to prevent oral ingestion of the test substance)
	- Removal of test substance: washed with tepid water after 6
	hours
	CLINICAL OBSERVATIONS AND FREQUENCY:
	- Mortality/clinical signs: daily
	- Dermal effects: before and after exposure
	- Body weight: weekly
	- Haematology/biochemistry: pre-test and after 21 days:
	haematocrit, Hb, erythrocyte count, (differential) leucocyte
	count, MCV, MCH(C) glucose, BUN, ALP, ALAT, ASAT

5. Toxicity	ld 583-78-8 Date 26.12.2001
	- Urinalysis: pre-test and after 21 days according to OECD 410
	ORGANS EXAMINED AT NECROPSY (MACROSCOPIC AND MICROSCOPIC):
	 Organ weights: liver, spleen, kidneys, brain, adrenals, thyroid, testes, ovaries
	 Macroscopic: all tissues (see microscopy) from all animals Microscopic: from all animals: skin, brain, lung, spleen,
	pancreas, stomach, small and large intestines, kidneys,
	urinary bladder, gallbladder, ovaries, testes, bone marrow,
	heart, prefemorral and mesenteric lymphnodes, liver, adrenals, thyroid, parathyroid, eye, pituitary, sciatic
	nerve, spinal cord, thymus, skeletal muscle, gross lesions
	STATISTICAL METHODS: ANOVA, Bartlett's test, t-test (Steel), Dunnett's test
Result	: TOXIC RESPONSE/EFFECTS BY DOSE LEVEL:
	- Mortality and time to death: one male at 10 mg/kg bw on
	day 20 and 3 females at 100 mg/ kg bw during week 3
	 Clinical signs: In males at 100 mg/kg bw red swollen eye, ocular and/or nasal discharge were seen.
	In animals that died diarrhoea was apparent on the day
	before death
	- Dermal effects:
	Skin effects were seen at all dose groups with increasing incidence and severity. At 1.0 mg/kg bw effects were
	restricted to erythema and oedema in all animals. At 10
	mg/kg bw atonia and corisceousness were seen next to
	erythema and oedema. At 100 mg/kg bw fissuring of the skin
	and desquamation was seen in addition to erythema, oedema, atonia and corisceousness
	- Body weight gain: no treatment related effects
	- Haematology:
	At 10 and 100 mg/kg bw the number of erythrocytes was
	increased in males. At 100 mg/kg bw an increased haemoglobin level was reported in males. Leucocyte counts were increased
	in males and females at 10 mg/kg bw and in males at 100
	mg/kg bw
	- Clinical chemistry:
	BUN and ALAT were decreased in the surviving female at 100 mg/kg bw
	- Urinalysis:
	A decreased volume was reported in males at 1.0 and 100
	mg/kg bw; specific gravity was increased at 1.0 mg/kg bw
	- Organ weights: Liver weight was decreased in females at 1.0 and 10 mg/ kg
	bw (both absolute and relative)
	Relative spleen weight was decreased in mid and high dosed
	females
	Absolute kidney weight and absolute and relative adrenal weight were decreased in females at 10 mg/kg bw
	- Gross pathology:
	Skin lesionss at the application site consisting of
	thickening, encrustation, sloughing, necrosis, leatherness,
	foci in the dermis and epidermis were reported in all

5. Toxicity	ld 583-78-8	
or reality	Date 26.12.2001	
	treated animals	
	 Histopathology: Skin effects (application site) included inflammatory cell 	
	infiltrate, acanthosis, hyperkeratosis and necrotic exudate	
	on the epidermal surface at 1.0 mg/kg bw. At 10 and/or 100	
	mg/kg bw additionally dermal fibroplasia and ulceration was	
	reported.	
	At 100 mg/kg hyperplasia of the lymphnodes was seen. Other incidental findings included areas of asperm and	
	ectatic tubuli in the testes, lung congestion, lymphoid	
	infiltrate in the liver, meningitis, nodules in the brain,	
	cysts in the thyroid.	
	Several animals showed an infection of coccidia in their small intestine	
	Sman mestine	
	STATISTICAL RESULTS: Effects on RBC and HB and liver weight	
0	reached a level of statistical significance	
Source Test substance	 Notox Hertogenbosch II, CAS 583-78-8 (2,5-dichlorophenol), purity not specified 	
Conclusion	: Based on local effects the LOAEL is 1.0 mg/kg bw.	
	For systemic effects a NOAEL of 100 mg/kg bw can be derived.	
	The lymphnode hyperplasia was considered secondary to skin	
B. P. I. W.	effects.	
Reliability	 (2) valid with restrictions 1 No analyses were performed to check the actual amount of 	
	test substance applied.	
	2 The number of animals/treatment was too small. Abrasion of	
	the skin of half of the animals did not seem to influence	
	the results, but is not requested by the OECD guideline 3 Effects on blood parameters remained within historical	
	values.	
	4 The liver effects were only seen in females and showed no	
	relationship with dose or microscopic changes. Therefore	
09.04.2001	they were considerd to be not related to treatment.	(9)
		(-)
5.5 GENETIC TOXICIT	TY 'IN VITRO'	
Туре	: HGPRT assay	
System of testing	: CHO-cells (K1-BH4)	
Concentration	: 62.5-250 ug/mL	
Cycotoxic conc. Metabolic activation	: 200 ug/mL : with and without	
Result	: negative	
Method	: other: not indicated	
Year	:	
GLP	: no data	
Test substance	: other TS	

- : no data : other TS
- : SYSTEM OF TESTING

Method

- Species/cell type: CHO-K1-BH4Proficiences: HGPRT
- Metabolic activation system: Arochlor-1254-induced male rat liver homogenate

ADMINISTRATION:

- Dosing: with and without S9 100, 125, 150, 200 and 250 $\,$

5. Toxicity	ld 583-78-8 Date 26.12.2001
Result	ug/mL; additionally with S9 62.5 and 75 ug/mL - Number of replicates: one - Positive and negative control: 5-Bromo 2'deoxyuridine (-S9), 3-methylcholanthrene (+S9) and DMSO (vehicle) Exposure time: 1.5E06 cells were exposed for 4 h followed by 6-7 day expression time CRITERIA FOR EVALUATING RESULTS: - Statistical method: Kastenbaum and Baumann : GENOTOXIC EFFECTS: - With metabolic activation: negative - Without metabolic activation: negative
	FREQUENCY OF EFFECTS: number of mutants remained within (negative) control ranges with the exception of the number of mutants in the lowest dose tested with S9-mix. Positive controls gave the expected results PRECIPITATION CONCENTRATION: not observed
	CYTOTOXICITY (% of control survival) at the highest tested concentration: - With metabolic activation: 0.4% at 250 ug/mL - Without metabolic activation: 20% at 250 ug/mL
Source Test substance Reliability	 STATISTICAL RESULTS: The increase of the number of mutants at 62.5 ug/mL (+S9) was statistically significant Notox Hertogenbosch II, CAS 583-78-8 (2,5-dichlorophenol), purity >98% (2) valid with restrictions The report is limited to the above mentioned. The increased number of mutants seen at 62.5 ug/mL in the assay with metabolic activation is considered to be not relevant, since no concentration effect relationship was observed.
06.04.2001	(1) (15)

Type Species Sex Strain Route of admin. Exposure period Doses Result Method Year GLP Test substance Method	 Micronucleus assay mouse male/female NMRI gavage single dose 1500 mg/kg bw negative other: not indicated no data other TS TEST ORGANISMS:
	 Age: 8-12 weeks Weight at study initiation: not indicated No. of animals: 10/treatment

5. Toxicity	ld 583-78-8 Date 26.12.2001
Result	ADMINISTRATION: • Vehicle: corn oil • Frequency of treatment: single dose by oral gavage (volume 5 ml/kg) • Sampling times: 24, 48 and 72 hours after treatment (samples from 10 animals each time, number of bone marrow smears not indicated) • Control groups and treatment: negative: corn oil (5 ml/kg) positive: cyclophosphamide (20 mg/kg bw in deionised water) EXAMINATIONS: • % of polychromatic erythrocytes (PCE) in 1000 erythrocytes • Number of micronucleated PCE/1000 PCE CRITERIA FOR EVALUATING RESULTS: • Statistical method: Wilcoxon's non-parametric rank sum test : TOXIC RESPONSE/EFFECTS BY DOSE LEVEL: Not reported EFFECT ON PCE/NCE RATIO: % PCE 44.6, 32.0 and 27.6 at 24, 48 and 72 hours, resp.
Source Test substance Conclusion Reliability	 GENOTOXIC EFFECTS: Mean number of micronucleated PCE: 0.6, 1.4 and 0.9 at 24, 48 and 72 hours sampling time, resp. STATISTICAL RESULTS: % PCE significantly decreased at the 72-hours sampling time Notox Hertogenbosch II, CAS 583-78-8 (2,5-dichlorophenol), purity >98% not clastogenic (2) valid with restrictions 1. The report was limited to the above mentioned. 2. The proportion of micronucleated PCE was determined for 1000 PCE. This is in agreement with OECD 474 (1983); OECD 474 (1997) requires evaluation of 2000 PCE.
06.04.2001	(1) (15)
5.7 CARCINOGENITY	
5.8 TOXICITY TO REP	RODUCTION
5.9 DEVELOPMENTA	L TOXICITY/TERATOGENICITY
5.10 OTHER RELEVAN	TINFORMATION
	22 / 26

5. Toxicity	ld	583-78-8
	Date	26.12.2001
5.11 EXPERIENCE WITH HUMAN EXPOSURE		

6. Refe	Id 583-78-8 Date 26.12.2001
(1)	Bayer, Investigations on the mutagenicity of 1,4-dichlorobenzene and its main metabolite 2,5-dichlorophenol in vivo and in vitro, 2000
(2)	Borzelleca J.F., Condie L.W. & Hayes J.R. Toxicological evaluation of selected chlorinated phanols Water chlorination: Chem. Envirn. Impact Health eff. Proc. Conf. 5K (1985) (1)
(3)	Borzelleca J.F., Condie L.W. & Hayes J.R. Toxicological evaluation of selected chlorinated phanols Water chlorination: Chem. Envirn. Impact Health eff. Proc. Conf. 5K (1985) (25)
(4)	Dolfing J, Harrison BK; Environ Sci Technol 26: 2213-93 (1991), As cited in HSDB update 8-09-2001
(5)	EPIWIN v3.05, Syracuse Research Corporation, Syracuse, NY (July 12, 2000)
(6)	Hansch, C., Leo, A., D. Hoekman. Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. Washington, DC: American Chemical Society., 1995. 15, As cited in HSDB update of 8-09-2001
(7)	Hoechst Aktiengesellschaft, Akute orale Toxizitaet von 2,5-Dichlorphenol an weiblichen SPF-Wistar-Ratten, 1976 (3)
(8)	Ingols RS et al; J Water Pollut Control Fed 38: 629-35 (1966) As cited in HSDB update of 8 09-2001
(9)	International Research and Development Corporation, 2,5-dichlorophenol: 3-week dermal toxicity study in rabbits, 1980 (1)
(10)	International Research and Development Corporation, 2,5-dichlorophenol: acute toxicity studies in rats and rabbits, 1974
(11)	International Research and Development Corporation, 2,5-dichlorophenol: acute toxicity studies in rats and rabbits, 1974 (108)
(12)	International Research and Development Corporation, 2,5- Dichlorophenol Four-week inhalation study in rats, 1980 (2)
(13)	Lide, D.R. (ed.). CRC Handbook of Chemistry and Physics. 76th ed. Boca Raton, FL: CRC Press Inc., 1995-1996.,p. 3-254
(14)	Lyman, W. J. et al. (1990). Handbook of Chemical PropertyEstimation Methods, pp. 7-4, Amer. Chem. Society,Washington, DC
(15)	Tegethoff K., Investigations on the mutagenicity of 1,4-dichlorobenzene and its main metabolite

6. References

ld 583-78-8 Date 26.12.2001

7.	Risk Assessment	ld	583-78-8
		Date	26.12.2001
7.1	END POINT SUMMARY		
7.2	HAZARD SUMMARY		
7.3	RISK ASSESSMENT		

IUCLID

Data Set

Existing Chemical CAS No. Generic name	: ID: 52166-72-0 : 52166-72-0 : 2,5-dichlorophenol, sodium salt
Producer Related Part Company Creation date	: Toxicology and Regulatory Affairs : 26.12.2001
Substance Related Part Company Creation date	Toxicology and Regulatory Affairs26.12.2001
Memo	:
Printing date Revision date Date of last Update	: 26.12.2001 : : 26.12.2001
Number of Pages	: 14
Chapter (profile) Reliability (profile) Flags (profile)	 Chapter: 1, 2, 3, 4, 5, 7 Reliability: without reliability, 1, 2, 3, 4 Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. 0	eneral Information	52166-72-0 26.12.2001
1.0.1	OECD AND COMPANY INFORMATION	
1.0.2	LOCATION OF PRODUCTION SITE	
1.0.3	IDENTITY OF RECIPIENTS	
1.1	GENERAL SUBSTANCE INFORMATION	
1.1.0	DETAILS ON TEMPLATE	
1.1.1	SPECTRA	
1.2	SYNONYMS	
1.3	IMPURITIES	
1.4	ADDITIVES	
1.5	QUANTITY	
1.6.1	LABELLING	
1.6.2	CLASSIFICATION	
1.7	USE PATTERN	
1.7.1	TECHNOLOGY PRODUCTION/USE	
1.8	OCCUPATIONAL EXPOSURE LIMIT VALUES	
1.9	SOURCE OF EXPOSURE	

1. General Information	bl	52166-72-0
1. General mormation		26.12.2001
1.10.1 RECOMMENDATIONS/PRECAUTIONARY MEASURES		
1.10.2 EMERGENCY MEASURES		
1.11 PACKAGING		
1.12 POSSIB. OF RENDERING SUBST. HARMLESS		
1.13 STATEMENTS CONCERNING WASTE		
1.14.1 WATER POLLUTION		
1.14.2 MAJOR ACCIDENT HAZARDS		
1.14.3 AIR POLLUTION		
1.15 ADDITIONAL REMARKS		
1.16 LAST LITERATURE SEARCH		
1.17 REVIEWS		
1.18 LISTINGS E.G. CHEMICAL INVENTORIES		

2. Physico-Chemical Data

(1)

2.1 MELTING POINT

Value Sublimation Method Year GLP Test substance Method Result		ca. 202 ° C 2001 no Estimation using MPBPWIN v1.40 in EPIWIN 3.05 SUMMARY MPBPWIN v1.40
		Boiling Point: 476.56 deg C (Adapted Stein and Brown Method) Melting Point: 349.84 deg C (Adapted Joback Method) Melting Point: 164.60 deg C (Gold and Ogle Method) Mean Melt Pt : 257.22 deg C (Joback; Gold,Ogle Methods) Selected MP: 201.65 deg C (Weighted Value)
Test substance Reliability Flag 26.12.2001	::	Sodium 2,5-dichlorophenol CAS 52166-72-0 (2) valid with restrictions Critical study for SIDS endpoint

2.2 BOILING POINT

2.3 DENSITY

2.3.1 GRANULOMETRY

2.4 VAPOUR PRESSURE

Decomposition:Methodother (calculated)Year: 2001GLP: noTest substance:Method: Estimation using MPBPWIN v1.40 in EPIWINResult:Vapor Pressure Estimations (25 deg C): (Using BP: 476.56 deg C (estimated)) (Using MP: 201.65 deg C (estimated)) VP: 1.46E-009 mm Hg (Modified Grain Me

VP: 4.04E-009 mm Hg (Mackay Method) Selected VP: 1.46E-009 mm Hg (Modified Grain Method) Test substance :: Reliability :: Flag :: Critical study for SIDS endpoint 26.12.2001 2.5 PARTITION COEFFICIENT Log pow :: ca12 at 25° C Method other (calculated) Year : 2001 GLP :: no Test substance :: Sodium 2,5-dichlorophenol CAS 52166-72-0 Reliability :: 2001 GLP :: no Test substance :: Sodium 2,5-dichlorophenol CAS 52166-72-0 Reliability :: (2) valid with restrictions Flag :: Critical study for SIDS endpoint 26.12.2001 : Critical study for SIDS endpoint 26.12.2001 : : Value :: ca. 40000 mg/l at 25 ° C Qualitative : : PH :: at and ° C Method : : Test substance : : <th></th>	
Reliability : (2) valid with restrictions Flag : Critical study for SIDS endpoint 26.12.2001 : 2.5 PARTITION COEFFICIENT Log pow : ca12 at 25° C Method other (calculated) Year : 2001 GLP : no Test substance : Sodium 2,5-dichlorophenol CAS 52166-72-0 Reliability : (2) valid with restrictions Flag : Critical study for SIDS endpoint 26.12.2001 : Ca. 40000 mg/l at 25 ° C Value : ca. 40000 mg/l at 25 ° C Qualitative : Pka : at 25 ° C PH : at	
Log pow : ca12 at 25° C Method other (calculated) Year : 2001 GLP : no Test substance : Method Method : Estimation using KOWWIN v1.66 in EPIWIN 3.05 Test substance : Sodium 2,5-dichlorophenol CAS 52166-72-0 Reliability : (2) valid with restrictions Flag : Critical study for SIDS endpoint 26.1 WATER SOLUBILITY Value : ca. 40000 mg/l at 25 ° C Qualitative : Pka : at 25 ° C PH : at and ° C Method : other: calculated Year : 2001 GLP : no Test substance : Method : Estimation using WSKOW v1.40 in EPIWIN 3.05 Result : :	(1)
Methodother (calculated)Year: 2001GLP: noTest substance:Method: Estimation using KOWWIN v1.66 in EPIWIN 3.05Test substance: Sodium 2,5-dichlorophenol CAS 52166-72-0Reliability: (2) valid with restrictionsFlag: Critical study for SIDS endpoint26.12.2001: Critical study for SIDS endpoint26.1WATER SOLUBILITYValue: ca. 40000 mg/l at 25 ° CQualitative:Pka: at 25 ° CPH: at and ° CMethod: Other: calculatedYear: 2001GLP: noTest substance:Method: Estimation using WSKOW v1.40 in EPIWIN 3.05Result:WSKOW v1.40 Results Log Kow (estimated) : 0.12Log Kow (estimated) : 0.12Log Kow (estimated) : 0.12Log Kow used by Water solubility estimates: 0.12Equation Used to Make Water Sol estimate: Log S (mol/L) = 0.796 - 0.854 log Kow - 0.00728 MW + Correction (used when Melting Point NOT available)	
Value : ca. 40000 mg/l at 25 ° C Qualitative : Pka : at 25 ° C PH : at and ° C Method : other: calculated Year : 2001 GLP : no Test substance : Method : Estimation using WSKOW v1.40 in EPIWIN 3.05 Result : WSKOW v1.40 Results Log Kow (estimated) : 0.12 Log Kow (estimated) : 0.12 Log Kow (experimental): not available from database Log Kow used by Water solubility estimates: 0.12 Equation Used to Make Water Sol estimate: Log S (mol/L) = 0.796 - 0.854 log Kow - 0.00728 MW + Correction (used when Melting Point NOT available)	(1)
Qualitative:Pka:at 25 ° CPH:at and ° CMethod:other: calculatedYear:2001GLP:noTest substance:Method:Estimation using WSKOW v1.40 in EPIWIN 3.05Result: WSKOW v1.40 ResultsLog Kow (estimated) : 0.12Log Kow (estimated) : 0.12Log Kow used by Water solubility estimates: 0.12Equation Used to Make Water Sol estimate:Log S (mol/L) = 0.796 - 0.854 log Kow - 0.00728 MW + Correction(used when Melting Point NOT available)	
Correction(s): Value No Applicable Correction Factors Log Water Solubility (in moles/L) : -0.649 Water Solubility at 25 deg C (mg/L): 4.147e+004	
Test substance:Sodium 2,5-dichlorophenol CAS 52166-72-0Reliability:(2) valid with restrictionsFlag:Critical study for SIDS endpoint26.12.2001::	(1)

2. F	Physico-Chemical Data	52166-72-0 26.12.2001
2.6.2	SURFACE TENSION	
2.7	FLASH POINT	
2.8	AUTO FLAMMABILITY	
2.9	FLAMMABILITY	
2.10	EXPLOSIVE PROPERTIES	
2.11	OXIDIZING PROPERTIES	
2.12	ADDITIONAL REMARKS	

3.1.1 PHOTODEGRADATION

Type Light source Light spect. Rel. intensity Indirect photolysis Sensitizer Conc. of sens. Rate constant Degradation Deg. Product Method Year GLP Test substance Method Remark Result	 air nm based on Intensity of Sunlight OH 1500000 molecule/cm3 cm3/(molecule*sec) % after 2001 no Estimation using APOWIN v1.90 in EPIWIN 3.05 The indirect photolysis rate was estimated using 2,5-dichlorophenol as t is the species most likely to exist in the vapor state. AOP Program (v1.90) Results: ====================================	
Test substance Reliability Flag 26.12.2001	 2,5-Dichlorophenol CAS 583-79-8 (2) valid with restrictions Critical study for SIDS endpoint 	(1)
3.1.2 STABILITY IN WAT	R	
Type t1/2 pH4 t1/2 pH7	: > 1 year at 25 degree C > 1 year at 25 degree C	

τι/z pπ4		> 1	year at 25 degree C
t1/2 pH7	:	> 1	year at 25 degree C
t1/2 pH9	:	> 1	year at 25 degree C
Deg. Product	:		
Method	:	othe	r (calculated)
0	-	othe	r (calculated)

Id 52166-72-0 Date 26.12.2001

Year GLP Test substance Method Remark	 2001 no Estimated on chemical principles based on absence of groups susceptible to hydrolysis The estimation program in EPIWIN has no capability to estimate hydrolysis
Result	rates for this compound This material has no groups that are susceptible to hydrolysis in the pH 4 to 9 range; therefore, it is considered stable to hydrolysis in surface and groundwater. It is estimated to have a hydrolysis half-life of greater than one year between pH 4 and pH 9.
Test substance Reliability Flag 26.12.2001	 Sodium 2,5-dichlorophenol CAS 52166-72-0 (2) valid with restrictions Critical study for SIDS endpoint (3)

3.1.3 STABILITY IN SOIL

3.2 MONITORING DATA

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

Type Media Air (level I) Water (level I) Soil (level I) Biota (level II / III) Soil (level II / III) Method Year Method	fugacity model level III
Result	The Fugacity was determined using the EQC Level III model as found in EPIWIN 3.05. Estimated values were used for physical constants. Biodegradation was based on the current best estimate for 2,5-dichlorophenol (from HSDB). Half life in air was determined from the APOWIN program for 2,5-dichlorophenol as this would be the likely volatile species. Direct photolysis was not considered in this model. Emissions were restricted to water and soil as this test substance it is not volatile. Other parameters used the default values found in EPIWIN.
Kesuit	: Level III Fugacity Model (Full-Output):

Id 52166-72-0

Date 26.12.2001

	Air Water Soil Sedime	Concentration (percent) 0.131 44 55.8 ent 0.0522	Hal f - Li (hr) 24 125 200 400	fe Emiss (kg/ 0 10 10 0	hr) 00		
	Air Water Soil	(atm) (k 5.92e-014 2.68e-012 1.21e-010	action g/hr) 15.6 1e+003 795 0.371	Advection (kg/hr) 5.4 181 0 0.00429	Reacti on (percent) 0. 779 50. 1 39. 8 0. 0186	Advecti on (percent) 0. 27 9. 04 0 0. 000214	
	Reacti Advect Percer		7 hr 21e+003 hr . 7				
	Air Wat Soi	ter: 125	sed upon u	ser-entry):			
	Air Wat	tion Times (hr) r: 100 ter: 1000 diment: 5e+004	:				
Test substance Reliability Flag 26.12.2001	: (2) valid	2,5-dichloropher with restrictions tudy for SIDS er		166-72-0		(1)

3.3.2 DISTRIBUTION

3.4 MODE OF DEGRADATION IN ACTUAL USE

3.5 **BIODEGRADATION**

Type Inoculum Contact time Degradation Result Remark	: aerobic : : 4 day : = 54 % after 4 day
	The free phenol form of this material is reported to undergo 54% ring degradation in 4 days with acclimated sludge, it cannot be determined if this test substance is considered readily biodegradable by OECD criteria
Result	The biological degradation of chlorophenols in activated sludge was studied. 2,5-Dichlorophenol was more resistent to degradation than 2,4- dichlorophenol. While 2,4-dichlorophenol was 100% degraded, including ring degradation, in five days, 2,5-dichlorophenol was only 52% ring- degraded in four days. [USEPA; Ambient Water Quality Criteria Doc: Chlorinated Phenols p.C-29 (1980) EPA 440/5-80-032]**PEER REVIEWED** As cited in HSDB record for 2,5-dichlorophenol, update of 8-

3. Environmental Fate and Pathways		52166-72-0 26.12.2001	
	09-2001	 	
Test substance Reliability	 2,5-Dichlorophenol CAS 583-79-8 (2) valid with restrictions 		
Flag 26.12.2001	: Critical study for SIDS endpoint		(2)
.6 BOD5, COD OR	BOD5/COD RATIO		
.7 BIOACCUMULA	TION		
.8 ADDITIONAL RI	EMARKS		

4. C	cotoxicity	52166-72-0 26.12.2001
4.1	ACUTE/PROLONGED TOXICITY TO FISH	
4.2	ACUTE TOXICITY TO AQUATIC INVERTEBRATES	
4.3	TOXICITY TO AQUATIC PLANTS E.G. ALGAE	
4.4	TOXICITY TO MICROORGANISMS E.G. BACTERIA	
4.5.1	CHRONIC TOXICITY TO FISH	
4.5.2	CHRONIC TOXICITY TO AQUATIC INVERTEBRATES	
4.6.1	TOXICITY TO SOIL DWELLING ORGANISMS	
4.6.2	TOXICITY TO TERRESTRIAL PLANTS	
4.6.3	TOXICITY TO OTHER NON-MAMM. TERRESTRIAL SPECIES	
4.7	BIOLOGICAL EFFECTS MONITORING	
4.8	BIOTRANSFORMATION AND KINETICS	
	ADDITIONAL REMARKS	

5. T	oxicity	52166-72-0 26.12.2001
544		
5.1.1	ACUTE ORAL TOXICITY	
5.1.2	ACUTE INHALATION TOXICITY	
5.1.3	ACUTE DERMAL TOXICITY	
5.1.4	ACUTE TOXICITY, OTHER ROUTES	
5.2.1	SKIN IRRITATION	
5.2.2	EYE IRRITATION	
5.3	SENSITIZATION	
5.4	REPEATED DOSE TOXICITY	
5.5	GENETIC TOXICITY 'IN VITRO'	
5.6	GENETIC TOXICITY 'IN VITRO'	
5.7	CARCINOGENITY	
5.8	TOXICITY TO REPRODUCTION	
5.9	DEVELOPMENTAL TOXICITY/TERATOGENICITY	
5.10	OTHER RELEVANT INFORMATION	
5.11	EXPERIENCE WITH HUMAN EXPOSURE	

6. Ref	erences Id 52166-72-0 Date 26.12.2001
(1)	EPIWIN v3.05, Syracuse Research Corporation, Syracuse, NY (July 12, 2000)
(2)	Ingols RS et al; J Water Pollut Control Fed 38: 629-35 (1966) As cited in HSDB update of 09-2001
(3)	Lyman, W. J. et al. (1990). Handbook of Chemical PropertyEstimation Methods, pp. 7-4, Amer. Chem. Society,Washington, DC

7.1	Risk Assessment	ld	52166-72-0	
		Date	26.12.2001	
7.1	END POINT SUMMARY			
7.2	HAZARD SUMMARY			
7.3	RISK ASSESSMENT			

IUCLID

Data Set

Existing Chemical CAS No. Generic name	 ID: 68938-81-8 68938-81-8 2,5-dichlorophenol, potassium salt
Producer Related Part Company Creation date	Toxicology and Regulatory Affairs26.12.2001
Substance Related Part Company Creation date	: Toxicology and Regulatory Affairs : 26.12.2001
Memo	:
Printing date Revision date Date of last Update	: 26.12.2001 : : 26.12.2001
Number of Pages	: 14
Chapter (profile) Reliability (profile) Flags (profile)	 Chapter: 1, 2, 3, 4, 5, 7 Reliability: without reliability, 1, 2, 3, 4 Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. General Information		d 68938-81-8 e 26.12.2001	
1.0.1	OECD AND COMPANY INFORMATION		
1.0.2	LOCATION OF PRODUCTION SITE		
1.0.3	IDENTITY OF RECIPIENTS		
1.1	GENERAL SUBSTANCE INFORMATION		
1.1.0	DETAILS ON TEMPLATE		
1.1.1	SPECTRA		
1.2	SYNONYMS		
1.3	IMPURITIES		
1.4	ADDITIVES		
1.5	QUANTITY		
1.6.1	LABELLING		
1.6.2	CLASSIFICATION		
1.7	USE PATTERN		
1.7.1	TECHNOLOGY PRODUCTION/USE		
1.8	OCCUPATIONAL EXPOSURE LIMIT VALUES		
1.9	SOURCE OF EXPOSURE		

1. 0	eneral Information	68938-81-8 26.12.2001
1.10.1	RECOMMENDATIONS/PRECAUTIONARY MEASURES	
1.10.2	EMERGENCY MEASURES	
1.11	PACKAGING	
1.12	POSSIB. OF RENDERING SUBST. HARMLESS	
1.13	STATEMENTS CONCERNING WASTE	
1.14.1	WATER POLLUTION	
1.14.2	MAJOR ACCIDENT HAZARDS	
1.14.3	AIR POLLUTION	
1.15	ADDITIONAL REMARKS	
1.16	LAST LITERATURE SEARCH	
1.17	REVIEWS	
1.18	LISTINGS E.G. CHEMICAL INVENTORIES	

2. Physico-Chemical Data

2.1 MELTING POINT

Value Sublimation Method Year GLP Test substance Method Result	ca. 201 ° C other: Calculated 2001 no Estimation using MPBPWIN v1.40 in EPIWIN 3.05 MPBPWIN (v1.40) Program Results: ====================================	
Test substance Reliability Flag 26.12.2001	Potassium 2,5-dichlorophenol CAS 68938-81-8 (2) valid with restrictions Critical study for SIDS endpoint	(1)
2.2 BOILING POINT		
2.3 DENSITY		
2.3.1 GRANULOMETRY		
2.4 VAPOUR PRESSUR		
2.4 VAFOUR FRESSUR		
Value Decomposition Method Year GLP Test substance	 < .00001 hPa at ° C other (calculated) 2001 no	
	4 / 14	

		58938-81-8 26.12.2001
Method Result	: Estimation using MPBPWIN v1.40 in EPIWIN 3.05	
Result	MPBPWIN (v1.40) Program Results:	
	Experimental Database Structure Match: no data	
	SMILES : c1(CL)ccc(CL)c(OK)c1 CHEM : Potassium 2,5-Dichlorophenol MOL FOR: C6 H3 CL2 O1 K1 MOL WT : 201.09	
	SUMMARY MPBPWIN v1.40	
	Vapor Pressure Estimations (25 deg C): (Using BP: 476.56 deg C (estimated)) (Using MP: 201.65 deg C (estimated)) VP: 4.71E-011 mm Hg (Antoine Method) VP: 1.46E-009 mm Hg (Modified Grain Method) VP: 4.04E-009 mm Hg (Mackay Method) Selected VP: 1.46E-009 mm Hg (Modified Grain Method))
Test substance	: Potassium 2,5-dichlorophenol CAS 68938-81-8	
Reliability Flag	(2) valid with restrictionsCritical study for SIDS endpoint	
26.12.2001 2.5 PARTITION COE	FFICIENT	(1)
Log pow	: ca12 at ° C	
Method	other (calculated)	
Method Year GLP		
Method Year GLP Test substance	other (calculated) 2001 no :	
Method Year GLP Test substance Method	other (calculated) 2001 no Estimation using KOWWIN v1.66 in EPIWIN 3.05	
Method Year GLP Test substance	other (calculated) 2001 no :	
Method Year GLP Test substance Method Test substance Reliability Flag	other (calculated) 2001 no Estimation using KOWWIN v1.66 in EPIWIN 3.05 Potassium 2,5-dichlorophenol CAS 68938-81-8	
Method Year GLP Test substance Method Test substance Reliability	other (calculated) 2001 no Estimation using KOWWIN v1.66 in EPIWIN 3.05 Potassium 2,5-dichlorophenol CAS 68938-81-8 (2) valid with restrictions	(1)
Method Year GLP Test substance Method Test substance Reliability Flag 26.12.2001	other (calculated) 2001 no Estimation using KOWWIN v1.66 in EPIWIN 3.05 Potassium 2,5-dichlorophenol CAS 68938-81-8 (2) valid with restrictions Critical study for SIDS endpoint	(1)
Method Year GLP Test substance Method Test substance Reliability Flag 26.12.2001 2.6.1 WATER SOLUBI	other (calculated) 2001 no Estimation using KOWWIN v1.66 in EPIWIN 3.05 Potassium 2,5-dichlorophenol CAS 68938-81-8 (2) valid with restrictions Critical study for SIDS endpoint	(1)
Method Year GLP Test substance Method Test substance Reliability Flag 26.12.2001 2.6.1 WATER SOLUBI Value Qualitative	 other (calculated) 2001 no Estimation using KOWWIN v1.66 in EPIWIN 3.05 Potassium 2,5-dichlorophenol CAS 68938-81-8 (2) valid with restrictions Critical study for SIDS endpoint LITY ca. 34 g/l at 25 ° C : 	(1)
Method Year GLP Test substance Method Test substance Reliability Flag 26.12.2001 2.6.1 WATER SOLUBI Value Qualitative Pka	 other (calculated) 2001 no Estimation using KOWWIN v1.66 in EPIWIN 3.05 Potassium 2,5-dichlorophenol CAS 68938-81-8 (2) valid with restrictions Critical study for SIDS endpoint LITY ca. 34 g/l at 25 ° C at 25 ° C 	(1)
Method Year GLP Test substance Method Test substance Reliability Flag 26.12.2001 2.6.1 WATER SOLUBI Value Qualitative	<pre>other (calculated) 2001 no Estimation using KOWWIN v1.66 in EPIWIN 3.05 Potassium 2,5-dichlorophenol CAS 68938-81-8 (2) valid with restrictions Critical study for SIDS endpoint LITY ca. 34 g/l at 25 ° C at 25 ° C at and ° C</pre>	(1)
Method Year GLP Test substance Method Test substance Reliability Flag 26.12.2001 2.6.1 WATER SOLUBI Value Qualitative Pka PH	 other (calculated) 2001 no Estimation using KOWWIN v1.66 in EPIWIN 3.05 Potassium 2,5-dichlorophenol CAS 68938-81-8 (2) valid with restrictions Critical study for SIDS endpoint LITY ca. 34 g/l at 25 ° C at 25 ° C at and ° C Estimation using WSKOW v1.40 in EPIWIN 3.05 	(1)
Method Year GLP Test substance Method Test substance Reliability Flag 26.12.2001 2.6.1 WATER SOLUBI Value Qualitative Pka PH Method	<pre>other (calculated) 2001 no Estimation using KOWWIN v1.66 in EPIWIN 3.05 Potassium 2,5-dichlorophenol CAS 68938-81-8 (2) valid with restrictions Critical study for SIDS endpoint LITY ca. 34 g/l at 25 ° C at 25 ° C at and ° C</pre>	(1)
Method Year GLP Test substance Method Test substance Reliability Flag 26.12.2001 2.6.1 WATER SOLUBI Value Qualitative Pka PH Method	 other (calculated) 2001 no Estimation using KOWWIN v1.66 in EPIWIN 3.05 Potassium 2,5-dichlorophenol CAS 68938-81-8 (2) valid with restrictions Critical study for SIDS endpoint LITY ca. 34 g/l at 25 ° C at 25 ° C at and ° C Estimation using WSKOW v1.40 in EPIWIN 3.05 Water Sol from Kow (WSKOW v1.40) Results: 	(1)

2. P	hysico-Chemic	al Data	ld 68938-81-8 Date 26.12.2001
		CHEM : Potassium 2,5-Dichloroph MOL FOR: C6 H3 CL2 O1 K1 MOL WT : 201.09	enol
		WSKOW v1.40 Results Log Kow (estimated) : 0.12 Log Kow (experimental): not availa Log Kow used by Water solubility es	ble from database
		Equation Used to Make Water Sol e Log S (mol/L) = 0.796 - 0.854 log (used when Melting Point NOT	Kow - 0.00728 MW + Correction
		Correction(s): Value	
		No Applicable Correction Factor	rs
		Log Water Solubility (in moles/L) Water Solubility at 25 deg C (mg/l	
	substance	: Potassium 2,5-dichlorophenol CAS	68938-81-8
Flag	ability I	(2) valid with restrictionsCritical study for SIDS endpoint	
26.1	2.2001		(1)
2.6.2	SURFACE TENSION		
2.7	FLASH POINT		
2.8	AUTO FLAMMABILI	тү	
2.9	FLAMMABILITY		
2.9			
2.10	EXPLOSIVE PROPE	RTIES	
2.11	OXIDIZING PROPER	TIES	
0.40	ADDITIONAL REMA	RKS	
2.12			
2.12			

3.1.1 PHOTODEGRADATION

Type Light source Light spect. Rel. intensity Indirect photolysis Sensitizer		air nm based on Intensity of Sunlight OH	
Conc. of sens. Rate constant Degradation Method	:	1500000 molecule/cm3 cm3/(molecule*sec) % after Estimation using APOWIN v1.90 in EPIWIN 3.05	
Remark Result	:	The indirect photolysis rate was estimated using 2,5-dichlorophenol as is the species most likely to exist in the vapor state. AOP Program (v1.90) Results:	that
		SMILES : c1(CL)ccc(CL)c(O)c1 CHEM : 2,5-Dichlorophenol MOL FOR: C6 H4 CL2 O1 MOL WT : 163.00 SUMMARY (AOP v1.90): HYDROXYL RADICALS Hydrogen Abstraction = 0.0000 E-12 cm3/molecule-sec Reaction with N, S and -OH = 0.1400 E-12 cm3/molecule-sec Addition to Triple Bonds = 0.0000 E-12 cm3/molecule-sec Addition to Olefinic Bonds = 0.0000 E-12 cm3/molecule-sec Addition to Olefinic Bonds = 0.0000 E-12 cm3/molecule-sec Addition to Fused Rings = 6.8451 E-12 cm3/molecule-sec Addition to Fused Rings = 0.0000 E-12 cm3/molecule-sec Addition to Fused Rings = 0.0000 E-12 cm3/molecule-sec Addition to Fused Rings = 0.0000 E-12 cm3/molecule-sec HALF-LIFE = 1.531 Days (12-hr day; 1.5E6 OH/cm3) HALF-LIFE = 18.375 Hrs	
Test substance Reliability Flag 26.12.2001	:	2,5-Dichlorophenol CAS 583-79-8 (2) valid with restrictions Critical study for SIDS endpoint	(1)

3.1.2 STABILITY IN WATER

Туре	: abiotic
t1/2 pH4	: > 1 year at 25 degree C
t1/2 pH7	: >1 year at 25 degree C
t1/2 pH9	: >1 year at 25 degree C
Deg. Product	:
Method	: other (calculated)
Year	: 2001
GLP	: no
Test substance	:
Method	: Estimated on chemical principles based on absence of groups susceptible

3. Environmental	Fate and Pathways	ld 68938-81-8	
		Date 26.12.2001	
	to hydrolysis		
Remark	: The estimation program in EPIWIN has rates for this compound.	no capability to estimate hydrolysis	
Result	: This material has no groups that are sus 9 range; therefore, it is considered stable groundwater. It is estimated to have a hy one year between pH 4 and pH 9.	e to hydrolysis in surface and	
Test substance Reliability	 Potassium 2,5-dichlorophenol CAS 6893 (2) valid with restrictions 	38-81-8	
Flag	: Critical study for SIDS endpoint		
26.12.2001		(3)	
.1.3 STABILITY IN SOII			

3.2 MONITORING DATA

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

Type Media Air (level I) Water (level I) Soil (level I) Biota (level II / III) Soil (level II / III) Method Year	fugacity model level III
Method Result	 The Fugacity was determined using the EQC Level III model as found in EPIWIN 3.05. Estimated values were used for physical constants. Biodegradation was based on the current best estimate for 2,5-dichlorophenol (from HSDB). Half life in air was determined from the APOWIN program for 2,5-dichlorophenol as this would be the likely volatile species. Direct photolysis was not considered in this model. Emissions were restricted to water and soil as this test substance it is not volatile. Other parameters used the default values found in EPIWIN. Level III Fugacity Model (Full-Output):
	$\begin{array}{rcl} \hline & & \\ \hline \hline & & \\ \hline & & \\ \hline \hline \\ \hline & & \\ \hline \hline & & \\ \hline \hline \\ \hline \hline \hline \hline \\ \hline \hline \hline \\$
	8 / 14

Id 68938-81-8

Date 26.12.2001

		ent 0.0517	400	0		
	Air Water Soil Sediment	Fugacity (atm) 4.82e-026 5.06e-020 2.32e-018 2.96e-020	Reaction (kg/hr) 1.38e-011 1.01e+003 813 0.373	Advection (kg/hr) 4.77e-012 181 0 0.0043	Reacti on (percent) 6. 89e-013 50. 3 40. 6 0. 0186	Advecti on (percent) 2. 39e-013 9. 07 0 0. 000215
	Reacti Advect Percer	stence Time: on Time: tion Time: nt Reacted: nt Advected:	229 hr 2.29e+003 hr 90.9	r		
	Ain Wat Soi	r: 24 ter: 125	(based upon t	user-entry):		
	Ai ı Wat	tion Times (r: 100 ter: 1000 diment: 5e+0)			
Test substance Reliability Flag 26.12.2001	: (2) valid	m 2,5-dichlo with restrictio tudy for SID		68938-81-8		(1
						(1
3.3.2 DISTRIBUTION						(1
3.3.2 DISTRIBUTION 3.4 MODE OF DEGR	ADATION IN AC	CTUAL USE				(1
		CTUAL USE				(1
3.4 MODE OF DEGR	ON : aerobic : activated : 4 day	CTUAL USE d sludge, ada after 4 day	apted			
 3.4 MODE OF DEGR 3.5 BIODEGRADATI Type Inoculum Contact time Degradation 	ON : aerobic : activated : 4 day : = 54 % : : The free degradat	l sludge, ada after 4 day phenol form tion in 4 days	apted of this materia s with acclimation	ted sludge, it	cannot be de	4% ring etermined if

3. Environmental Fate and Pathways			ld 68938-81-8 Date 26.12.2001		
Relia Flag	t substance ability 2.2001	:	2,5-Dichlorophenol CAS 583-79-8 (2) valid with restrictions Critical study for SIDS endpoint		(2)
3.6	BOD5, COD OR	BOD5/	COD RATIO		
3.7	BIOACCUMULA	TION			
3.8	ADDITIONAL RI	EMARK	S		

4. E	cotoxicity	68938-81-8 26.12.2001
4.1	ACUTE/PROLONGED TOXICITY TO FISH	
4.2	ACUTE TOXICITY TO AQUATIC INVERTEBRATES	
4.3	TOXICITY TO AQUATIC PLANTS E.G. ALGAE	
4.4	TOXICITY TO MICROORGANISMS E.G. BACTERIA	
4.5.1	CHRONIC TOXICITY TO FISH	
4.5.2	CHRONIC TOXICITY TO AQUATIC INVERTEBRATES	
4.6.1	TOXICITY TO SOIL DWELLING ORGANISMS	
4.6.2	TOXICITY TO TERRESTRIAL PLANTS	
4.6.3	TOXICITY TO OTHER NON-MAMM. TERRESTRIAL SPECIES	
4.7	BIOLOGICAL EFFECTS MONITORING	
4.8	BIOTRANSFORMATION AND KINETICS	
4.9	ADDITIONAL REMARKS	

5. Toxic	ity	68938-81-8 26.12.2001
5.1.1 ACU	TE ORAL TOXICITY	
5.1.2 ACU		
5.1.3 ACU	TE DERMAL TOXICITY	
5.1.4 ACU	TE TOXICITY, OTHER ROUTES	
5.2.1 SKIN	IRRITATION	
5.2.2 EYE	RRITATION	
5.3 SENS	SITIZATION	
5.4 REPE	EATED DOSE TOXICITY	
5.5 GENI	ETIC TOXICITY 'IN VITRO'	
5.6 GEN	ETIC TOXICITY 'IN VITRO'	
5.7 CAR	CINOGENITY	
5.8 TOXI	CITY TO REPRODUCTION	
5.9 DEVE	LOPMENTAL TOXICITY/TERATOGENICITY	
5.10 OTH	R RELEVANT INFORMATION	
5.11 EXPE	RIENCE WITH HUMAN EXPOSURE	

6. Ref	erences	ld Date	68938-81-8 26.12.2001
(1)	EPIWIN v3.05, Syracuse Research Corporation, Syracuse, N	IY (July 12, 2	2000)
(2)	Ingols RS et al; J Water Pollut Control Fed 38: 629-35 (1966) 09-2001) As cited in I	HSDB update of 8-
(3)	Lyman, W. J. et al. (1990). Handbook of Chemical PropertyE Amer. Chem. Society,Washington, DC	stimation Me	thods, pp. 7-4,

7. Risk Assessment		Id 68938-81-8	
		Date	26.12.2001
7.1	END POINT SUMMARY		
7.2	HAZARD SUMMARY		
7.3	RISK ASSESSMENT		

IUCLID

Data Set

Existing Chemical CAS No. Generic name	: ID: 1984-58-3 : 1984-58-3 : 2,5-dichloroanisole
Producer Related Part Company Creation date	: Toxicology and Regulatory Affairs : 26.12.2001
Substance Related Part Company Creation date	Toxicology and Regulatory Affairs26.12.2001
Memo	:
Printing date Revision date Date of last Update	: 26.12.2001 : : 26.12.2001
Number of Pages	: 14
Chapter (profile) Reliability (profile) Flags (profile)	 Chapter: 1, 2, 3, 4, 5, 7 Reliability: without reliability, 1, 2, 3, 4 Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. 0	eneral Information	1984-58-3 26.12.2001
1.0.1	OECD AND COMPANY INFORMATION	
1.0.2	LOCATION OF PRODUCTION SITE	
1.0.3	IDENTITY OF RECIPIENTS	
1.1	GENERAL SUBSTANCE INFORMATION	
1.1.0	DETAILS ON TEMPLATE	
1.1.1	SPECTRA	
1.2	SYNONYMS	
1.3	IMPURITIES	
1.4	ADDITIVES	
1.5	QUANTITY	
1.6.1	LABELLING	
1.6.2	CLASSIFICATION	
1.7	USE PATTERN	
1.7.1	TECHNOLOGY PRODUCTION/USE	
1.8	OCCUPATIONAL EXPOSURE LIMIT VALUES	
1.9	SOURCE OF EXPOSURE	

1. G	eneral Information	1984-58-3 26.12.2001
1.10.1	RECOMMENDATIONS/PRECAUTIONARY MEASURES	
1.10.2	EMERGENCY MEASURES	
1.11	PACKAGING	
1.12	POSSIB. OF RENDERING SUBST. HARMLESS	
1.13	STATEMENTS CONCERNING WASTE	
1.14.1	WATER POLLUTION	
1.14.2	MAJOR ACCIDENT HAZARDS	
1.14.3	AIR POLLUTION	
1.15	ADDITIONAL REMARKS	
1.16	LAST LITERATURE SEARCH	
1.17	REVIEWS	
1.18	LISTINGS E.G. CHEMICAL INVENTORIES	

2. Physico-Chemical Data

2.1 MELTING POINT

Value Sublimation Method Year GLP Test substance Method		ca. 21 ° C 2001 no Estimation using MPBPWIN v1.40 in EPIWIN 3.05	
Result	:	MPBPWIN (v1.40) Program Results:	
		Experimental Database Structure Match: no data	
		SMILES : c1(CL)ccc(CL)c(OC)c1 CHEM : 2,5-Dichloroanisole MOL FOR: C7 H6 CL2 O1 MOL WT : 177.03	
		SUMMARY MPBPWIN v1.40	
		Boiling Point: 215.67 deg C (Adapted Stein and Brown Method)	
		Melting Point:29.02 deg C (Adapted Joback Method)Melting Point:12.27 deg C (Gold and Ogle Method)Mean Melt Pt :20.65 deg C (Joback; Gold,Ogle Methods)Selected MP:20.65 deg C (Mean Value)	
Test substance Reliability Flag 26.12.2001	:	2,5-Dichloroanisole CAS 1984-58-3(2) valid with restrictionsCritical study for SIDS endpoint	(1)
2.2 BOILING POINT			
Value Method	:	ca. 216 °C at 1013 hPa Estimation using MPBPWIN v1.40 in EPIWIN 3.05	
Result	:	MPBPWIN (v1.40) Program Results:	
		Experimental Database Structure Match: no data	
		SMILES : c1(CL)ccc(CL)c(OC)c1 CHEM : 2,5-Dichloroanisole MOL FOR: C7 H6 CL2 O1 MOL WT : 177.03	
		SUMMARY MPBPWIN v1.40	
		Boiling Point: 215.67 deg C (Adapted Stein and Brown Method)	
		4 / 14	

2. Physico-Chem	ical Data		1984-58-3 26.12.2001
Test substance Reliability Flag 26.12.2001	 2,5-Dichloroanisole CAS 1984-58-3 (2) valid with restrictions Critical study for SIDS endpoint 		(1)
2.3 DENSITY			
2.3.1 GRANULOMETRY			
2.4 VAPOUR PRESSU	RE		
Value Decomposition Method Year GLP Test substance Method Result	 ca22 hPa at 25° C other (calculated) 2001 no Estimation using MPBPWIN v1.40 in E MPBPWIN (v1.40) Program Results: Experimental Database Structure Mate SMILES : c1(CL)ccc(CL)c(OC)c1 CHEM : 2,5-Dichloroanisole MOL FOR: C7 H6 CL2 O1 MOL WT : 177.03 SUMMARY MPBPWIN v1.40 Vapor Pressure Estimations (25 deg C (Using BP: 215.67 deg C (estimated)) (MP not used for liquids) VP: 0.176 mm Hg (Antoine Method) VP: 0.152 mm Hg (Modified Grain II) VP: 0.253 mm Hg (Mackay Method) Selected VP: 0.164 mm Hg (Mean or 100) 	:== ch: no data c):)) l) Method) l)	nethods)
Test substance Reliability Flag 26.12.2001	 2,5-Dichloroanisole CAS 1984-58-3 (2) valid with restrictions Critical study for SIDS endpoint 		(1)
2.5 PARTITION COEF	FICIENT		
Log pow Method	: ca. 3.36 at 25° C		
Year GLP	: 2001 : no		

Test substance

Method

	nical Data Id 1984-58- Date 26.12.20	
Test substance Reliability Flag 26.12.2001	 2,5-Dichloroanisole CAS 1984-58-3 (2) valid with restrictions Critical study for SIDS endpoint 	('
2.6.1 WATER SOLUB	LITY	
Value Qualitative Pka PH Method Year GLP Test substance Method Result	 ca. 75 mg/l at 25 ° C at 25 ° C at and ° C 2001 no Estimation using WSKOW v1.40 in EPIWIN 3.05 Water Sol from Kow (WSKOW v1.40) Results: ====================================	ı
	Correction(s): Value No Applicable Correction Factors Log Water Solubility (in moles/L) : -3.365	
Test substance Reliability Flag	 Water Solubility at 25 deg C (mg/L): 76.44 2,5-Dichloroanisole CAS 1984-58-3 (2) valid with restrictions Critical study for SIDS endpoint 	

2.7 FLASH POINT

2. P	Physico-Chemical Data	ld Date	1984-58-3 26.12.2001	
2.8	AUTO FLAMMABILITY			
2.9	FLAMMABILITY			
2.10	EXPLOSIVE PROPERTIES			
2.11	OXIDIZING PROPERTIES			
2.12	ADDITIONAL REMARKS			

(1)

3.1.1 PHOTODEGRADATION

Type Light source Light spect. Rel. intensity Indirect photolysis Sensitizer Conc. of sens. Rate constant Degradation Deg. Product Method Year GLP Test substance Method	air nm based on Intensity of Sunlight OH 1500000 cm3/(molecule*sec) % after 2001 Estimation using APOWIN v1.90 in EPIWIN 3.05	
Result	AOP Program (v1.90) Results: ====================================	
Test substance Reliability Flag 26.12.2001	2,5-Dichloroanisole CAS 1984-58-3 (2) valid with restrictions Critical study for SIDS endpoint	

3.1.2 STABILITY IN WATER

Туре	:	abiotic
t1/2 pH4	:	> 1 year at 25 degree C
t1/2 pH7		> 1 year at 25 degree C
t1/2 pH9	:	> 1 year at 25 degree C
Deg. Product	:	
Method	:	
Year	:	2001
GLP	:	no
Test substance	:	

3. Environment	al Fate and Pathways	ld 1984-58-3 Date 26.12.2001
Method	: Estimated on chemical principles to hydrolysis	based on absence of groups susceptible
Remark	: The estimation program in EPIWII rates for this compound	N has no capability to estimate hydrolysis
Result	9 range; therefore, it is considered	are susceptible to hydrolysis in the pH 4 to d stable to hydrolysis in surface and ve a hydrolysis half-life of greater than
Test substance Reliability Flag 26.12.2001	 2,5-Dichloroanisole CAS 1984-58 (2) valid with restrictions Critical study for SIDS endpoint 	8-3 (2)
3.1.3 STABILITY IN S	OIL	

3.2 MONITORING DATA

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

Type Media Air (level I) Water (level I) Soil (level I) Biota (level II / III) Soil (level II / III) Method Year	fugacity model level III fugacity fugacity model level III fugacity fugacity fugacit
Method	: The Fugacity was determined using the EQC Level III model as found in EPIWIN 3.05. Estimated values were used for physical constants. Biodegradation was based on the EPIWIN derived estimates that were assessed for reasonableness compared with similar compounds. Half life in air was determined from the APOWIN program for 2,5-dichlorophenol as this would be the likely volatile species. Direct photolysis was not considered in this model. Emissions were calculated from air water and soil as this test substance it is volatile. Other parameters used the default values found in EPIWIN
Result	: Level III Fugacity Model (Full-Output): Chem Name : 2, 5-Dichloroanisole Molecular Wt: 177.03 Henry's LC : 0.00315 atm-m3/mole (Henrywin program) Vapor Press : 0.164 mm Hg (Mpbpwin program) Log Kow : 3.36 (Kowwin program) Soil Koc : 939 (calc by model) Concentration Half-Life Emissions (percent) (hr) (kg/hr) Air 7.6 48.9 1000 9/14

Id 1984-58-3

Date 26.12.2001

		Water Soil Sedime	22. 8 68. 8 nt 0. 812	900 900 3. 6e		000 000		
		Ai r Water Soi l	Fugaci ty (atm) 1. 14e- 010 2. 19e- 008 3. 22e- 008 1. 66e- 008	Reacti on (kg/hr) 1. 16e+003 190 573 1. 69	Advection (kg/hr) 823 247 0 0.176	Reacti on (percent) 38. 8 6. 34 19. 1 0. 0564	Advecti on (percent) 27.4 8.23 0 0.00586	
		Reacti Advect Percen	tence Time: on Time: ion Time: t Reacted: t Advected:	561 hr 1. 01e+003 hr 64. 3				
		Air Wat Soi Sed	: 48.9 er: 900 1: 900 iment: 3600				opwin) :	
		Ai r Wat						
Rel Fla	iability : g : 12.2001	(2) valid v	oroanisole (with restrictio udy for SIDS		3			(1)
3.3.2	DISTRIBUTION							
3.4	MODE OF DEGRADA		TIIAI IISE					
5.7								
3.5	BIODEGRADATION							
3.6	BOD5, COD OR BOD5	COD RATI	0					

3.7 BIOACCUMULATION

3.8 ADDITIONAL REMARKS

	cotoxicity	1984-58-3 26.12.2001
4.1	ACUTE/PROLONGED TOXICITY TO FISH	
4.2	ACUTE TOXICITY TO AQUATIC INVERTEBRATES	
4.3	TOXICITY TO AQUATIC PLANTS E.G. ALGAE	
4.4	TOXICITY TO MICROORGANISMS E.G. BACTERIA	
4.5.1	CHRONIC TOXICITY TO FISH	
4.5.2	CHRONIC TOXICITY TO AQUATIC INVERTEBRATES	
4.6.1	TOXICITY TO SOIL DWELLING ORGANISMS	
4.6.2	TOXICITY TO TERRESTRIAL PLANTS	
4.6.3	TOXICITY TO OTHER NON-MAMM. TERRESTRIAL SPECIES	
4.7	BIOLOGICAL EFFECTS MONITORING	
4.8	BIOTRANSFORMATION AND KINETICS	
4.9	ADDITIONAL REMARKS	

5.1	oxicity	1984-58-3 26.12.2001
5.1.1	ACUTE ORAL TOXICITY	
5.1.2	ACUTE INHALATION TOXICITY	
5.1.3	ACUTE DERMAL TOXICITY	
5.1.4	ACUTE TOXICITY, OTHER ROUTES	
5.2.1	SKIN IRRITATION	
5.2.2	EYE IRRITATION	
5.3	SENSITIZATION	
5.4	REPEATED DOSE TOXICITY	
5.5	GENETIC TOXICITY 'IN VITRO'	
5.6	GENETIC TOXICITY 'IN VITRO'	
5.7	CARCINOGENITY	
5.8	TOXICITY TO REPRODUCTION	
5.9	DEVELOPMENTAL TOXICITY/TERATOGENICITY	
5.10	OTHER RELEVANT INFORMATION	
5.11	EXPERIENCE WITH HUMAN EXPOSURE	

 EPIWIN v3.05, Syracuse Research Corporation, Syracuse, Lyman, W. J. et al. (1990). Handbook of Chemical Property Amer. Chem. Society, Washington, DC 	NY (July 12, 2000)
	Estimation Methods, pp. 7-4

7. F	Risk Assessment	ld	1984-58-3
		Date	26.12.2001
7.1	END POINT SUMMARY		
7.2	HAZARD SUMMARY		
1.2			
7.3	RISK ASSESSMENT		

IUCLID

Data Set

Existing Chemical CAS No. Generic name	: ID: 50594-66-6 : 50594-66-6 : Acifluorfen
Producer Related Part Company Creation date	Toxicology and Regulatory Affairs26.12.2001
Substance Related Part Company Creation date	Toxicology and Regulatory Affairs26.12.2001
Memo	:
Printing date Revision date Date of last Update	: 27.12.2001 : : 27.12.2001
Number of Pages	: 23
Chapter (profile) Reliability (profile) Flags (profile)	 Chapter: 1, 2, 3, 4, 5, 7 Reliability: without reliability, 1, 2, 3, 4 Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. 0	eneral Information	50594-66-6 27.12.2001
1.0.1	OECD AND COMPANY INFORMATION	
1.0.2	LOCATION OF PRODUCTION SITE	
1.0.3	IDENTITY OF RECIPIENTS	
1.1	GENERAL SUBSTANCE INFORMATION	
1.1.0	DETAILS ON TEMPLATE	
1.1.1	SPECTRA	
1.2	SYNONYMS	
1.3	IMPURITIES	
1.4	ADDITIVES	
1.5	QUANTITY	
1.6.1	LABELLING	
1.6.2	CLASSIFICATION	
1.7	USE PATTERN	
1.7.1	TECHNOLOGY PRODUCTION/USE	
1.8	OCCUPATIONAL EXPOSURE LIMIT VALUES	
1.9	SOURCE OF EXPOSURE	

1. General Information	50594-66-6 27.12.2001
1.10.1 RECOMMENDATIONS/PRECAUTIONARY MEASURES	
1.10.2 EMERGENCY MEASURES	
1.11 PACKAGING	
1.12 POSSIB. OF RENDERING SUBST. HARMLESS	
1.13 STATEMENTS CONCERNING WASTE	
1.14.1 WATER POLLUTION	
1.14.2 MAJOR ACCIDENT HAZARDS	
1.14.3 AIR POLLUTION	
1.15 ADDITIONAL REMARKS	
1.16 LAST LITERATURE SEARCH	
1.17 REVIEWS	
1.18 LISTINGS E.G. CHEMICAL INVENTORIES	

2. Physico-Chemical Data

2.1 MELTING POINT

	blimation thod ar	: : : : : : : : : : : : : : : : : : : :	= 150 ° C no data	
• = -	t substance	:		
Rer	nark	:	Published data found in EPIWIN. SRC data base Supported by Estimation using MPBPWIN v1.40 in EPIWIN 3.05 - SUMMARY MPBPWIN v1.40	
			Boiling Point: 442.92 deg C (Adapted Stein and Brown Method)	
			Melting Point: 349.84 deg C (Adapted Joback Method) Melting Point: 144.96 deg C (Gold and Ogle Method) Mean Melt Pt : 247.40 deg C (Joback; Gold,Ogle Methods) Selected MP: 185.94 deg C (Weighted Value)	
Res	sult	:	CAS Number : 050594-66-6 Chem Name : ACIFLUORFEN Mol Formula: C14H7CIF3NO5 Mol Weight : 361.66 Melting Pt : 150 deg C	
	t substance iability		Acifluorfen CAS 50594-66-6 (2) valid with restrictions Data from handbooks and standard reference sources assigned a 2	
Fla 26.4	g 12.2001	:	Critical study for SIDS endpoint	(7)
2.2	BOILING POINT			
2.3	DENSITY			
2.3.1	GRANULOMETRY			
2.4	VAPOUR PRESSUR	RE		
Met Yea GLI	composition thod ar P	::	= .00000002 hPa at 25° C other (calculated) 1985 no data	
Ies	st substance	:		

2. Physico-Che	nical Data		50594-66-6 27.12.2001	
Remark	: Published data found in EPIWIN. SRC data base	e		
Result	: Vapor Pressure: Value : 1.53E-008 mm Hg Temp : 25 deg C Type : EST Ref : NEELY,WB & BLAU,GE (1985)			
Reliability Flag	(2) valid with restrictionsCritical study for SIDS endpoint			
26.12.2001			(7)
Value	: ca000000052 hPa at 25° C			
Decomposition Method	: other (calculated)			
Year	: 2001			
GLP Test substance	: no :			
Method	Estimation using MPBPWIN v1.40 in EPIWIN 3.	05		
Result	: SUMMARY MPBPWIN v1.40			
Test substance Reliability Flag	 (Using BP: 442.92 deg C (estimated)) (Using MP: 150.00 deg C (exp database)) VP: 3.26E-009 mm Hg (Antoine Method) VP: 3.94E-008 mm Hg (Modified Grain Method) VP: 8.94E-008 mm Hg (Mackay Method) Selected VP: 3.94E-008 mm Hg (Modified Grain Acifluorfen CAS 50594-66-6 (2) valid with restrictions Critical study for SIDS endpoint 		od)	
26.12.2001			(5	5)
2.5 PARTITION CO	FFICIENT			
Log pow Method	: = 3.7 at ° C			
Year	: 1992			
GLP Test substance	: no data :			
Result	Log P (octanol-water): Value : 3.70 Type : EXP Ref : NANDIHALLI UB ET AL. (1992)			
Test substance	: Acifluorfen CAS 50594-66-6			
Reliability Flag	(2) valid with restrictionsCritical study for SIDS endpoint			

2. Physico-Chemical Data

ld 50594-66-6 Date 27.12.2001

2.6.1 WATER SOLUBILITY

Pka PH Meth Year GLP	itative	= 120 mg/l at 25 ° at 25 ° C at and ° C 1994 no data	C	
Resi	ılt	Water Solubility: Value : 120 mg/L Temp : 25 deg C Type : EXP Ref : TOMLIN,C	;	
	substance ability	Acifluorfen CAS 50 (2) valid with restric		
Flag	-	Published value Critical study for SI		
	2.2001	- · · · · · · , · · ·		(7)
2.6.2	SURFACE TENSION			
2.7	FLASH POINT			
2.8	AUTO FLAMMABILI			
2.9	FLAMMABILITY			
2.10	EXPLOSIVE PROPE	IES		
2.11		ES		
2.12	ADDITIONAL REMA			

3.1.1 PHOTODEGRADATION

Type Light source Light spect. Rel. intensity Conc. of subst. Deg. Product Method Year GLP Test substance	 water Xenon lamp > 290 nm based on Intensity of Sunlight at 25 degree C EPA Guide-line subdivision N 161-2 "Photodegradation studies in water" yes other TS
Method	 Photolysis of acifluorfen 14C-labelled in the nitrobenzoate moiety {5-[2-chloro-4-(trifluoro-methyl)-phenoxy]-2-nitro benzoic acid-UL-14C (N-label)) and in the phenoxy trifluoromethyl moiety {5-[2-chloro-4-(trifluoro-methyl)-phenoxy-UL-14C]-2-nitro benzoic acid (F-label)) was studied at 25 deg C. Hereto, TS (N- or F-label) was dissolved in sterile 0.025M phosphate buffer (1% acetonitrile) at concentrations in the range 4 - 5 ppm. Volatiles were trapped in ethylene glycol (1 trap), 0.1N sulfuric (1 trap) acid and 1N NaOH (2 traps). Light source was a xenon lamp of intensity 1900 uE.m-2.s-1 (equivalent to summer noon time sun). Radiation < 290 nm was filtered out. Quantitation and identification/characterization was performed using LSC, TLC (two solvent systems), UV-vis spectroscopy and HPLC with 14C-detection (quantitation by scintillation of the column effluent). Intermediates and reference substances were derivatized by methylation using diazomethane and compared by 2D-HPLC. The following reference substances were available: Acifluorfen Amine Desnitro acifluorfen Acifluorfen Methyl Ester Descarboxy Acifluorfen Acifluorfen Acid Amine Acifluorfen Acetamide Amino Acifluorfen ME Acifluorfen Acetamide Antino Acifluorfen ME Acifluorfen Acid Antiranilic Acid Acifluorfen Dark controls and adsorption controls were included. Samples were taken in N-label test mixture at 0, 0.94, 1.8, 3.8, 18.0, 22.4, 30.2, 41.7, 64.3, 70.0, 87.1, 92.7, 110.7, 111.8, 116.1, 134.4, 134.5, 140.3, 157.8, 158.0, 162.8, 182.0 and 204.5 hrs. Samples in F-label test mixture were taken at 0, 64.3, 87.1, 110.7, 134.5 and 158 hrs; dark

3. Environmenta	al Fate and Pathways	ld 50594-66-6 Date 27.12.2001
	controls at 0, 64.3 and 110.7 hrs.	
Result	 Degradation could be described by 1st lives measured for N-label TS were in th half-life measured for F-label TS was 95% degradation N-label TS at 205 hrs: 87% degradation F-label TS at 158 hrs: 70% 	ne range 78-100 hrs, 5 (conc. 4-5 ppm). 1.4%
	Maximal concentration of metabolites (% radioactivity) measured during irradiatio	
	Meta- N-label test mixture F-label te bolite*	st mixture
	Max. % of Max. % of applied	
	P1 35.4 24.0 P2 5.1 7.4 P3 7.8 5.3 P4 6.8 5.6 P11 1.6 1.9 P12 1.8 1.6 Volatiles 0.2 3.3 0.0 0.0 (Sulf. acid) 9.4 5.1 (NaOH)	
	Remarks: Concentration range (N-label): 4.42-4.80 Concentration (F-label): 3.98 ppm Irradiation period: 205 hrs (N-label); 158 Mass balance: 85.6-101.6%.	
	 Hydrolysis of volatile recovered in ethy yielded one major intermediate and one HPLC retention time identical to that of trapped in NaOH. This suggests that the trap is the hydrolysis product of the vola trapped in ethylene glycol. Metabolites could not be identified. Ba isotope dilution experiments formation of acid and anthranilic acid could be excluinot yield distinct reaction products. Major metabolite (P1) appears to actual complex mixture of compounds (TLC arrest No adsorption or degradation in dark conserved. 	final moiety with an the compound e volatile in the NaOH atile incompletely sed on reverse of 2-nitrobenzoic ded. Methylation did ally consist of a nd derivatization).
Test substance	 III, CAS 50594-66-6 (acifluorfen), actual 5-[2-chloro-4-(trifluoro-methyl)-phenoxy benzoic acid, radiopurity 95.27% (HPLC III, CAS 50594-66-6 (acifluorfen), radio- 5-[2-chloro-4-(trifluoro-methyl)-phenoxy acid-UL-14C, radiochemical purity 99.64 5-[2-chloro-4-(trifluoro-methyl)-phenoxy benzoic acid, radiochemical purity 95.27 	-UL-14C]-2-nitro ;) labelled:]-2-nitro benzoic % (HPLC) and -UL-14C]-2-nitro
Conclusion	: t1/2 = 78-100 hrs	

3. Environment	al Fate and Pathways		50594-66-6 27.12.2001
Reliability Flag 26.12.2001	: (1) valid without restriction: Critical study for SIDS endpoint		(
3.1.2 STABILITY IN V	ATER		
Type t1/2 pH4 t1/2 pH7 t1/2 pH9 Deg. Product Method Year GLP Test substance	 abiotic > 1 year at 25 degree C 2001 no 		
Remark	 Estimated on chemical principles based on absence to hydrolysis The estimation program in EPIWIN has no capability rates for this compound. 	-	
Result	9 range; therefore, it is considered stable to hydrolys	9 range; therefore, it is considered stable to hydrolysis in surface and groundwater. It is estimated to have a hydrolysis half-life of greater than	
Test substance Reliability Flag 26.12.2001	 Acifluorfen CAS 50594-66-6 (2) valid with restrictions Critical study for SIDS endpoint 		(

3.2 MONITORING DATA

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

Type Media Air (level I) Water (level I) Soil (level I) Biota (level II / III) Soil (level II / III) Method Year	fugacity model level III
Method	: The Fugacity was determined using the EQC Level III model as found in EPIWIN 3.05. Measured and estimated values were used for physical constants. Biodegradation was based on information in the EPA Reregistration Documentation and data in HSDB. The aquatic soil and
	0.4.00

3. Environmenta	I Fate and Pathways	ld 50594-66-6 Date 27.12.2001
Result	sediment estimates are estimates of an a biodegradation and photolysis. As sedim life estimate for water was used in the m default rapid loss since this material is n calculated from using only water and soi volatile. Other parameters used the defa	ent distribution was low the half odel. Half life in air was set at a ot volatile. Emissions were I as this test substance it is not
	Level III Fugacity Model (Full-Output)	
	Chem Name : Acifluorfen Molecular Wt: 361.66 Henry's LC : 6.03e-011 atm-m3/mole Vapor Press : 3.94e-008 mm Hg (Mpbp Liquid VP : 1.54e-006 mm Hg (supe Melting Pt : 186 deg C (Mpbpwin pro Log Kow : 3.7 (Kowwin program) Soil Koc : 2.05e+003 (calc by mo	win program) er-cooled) gram)
	Concentration (percent) Half-Life (hr) Air 4.41e-009 296 Water 14.1 3.6e+003 Soil 83.8 3.6e+003 Sediment 2.09 1.44e+004	Emi ssi ons (kg/hr) 0 1000 1000 0
	Fugacity Reaction Advection (atm) (kg/hr) (kg/ Air 1.13e-019 6.22e-007 2.6 Water 7.09e-016 164 853 Soil 9.45e-016 974 0 Sediment 1.05e-015 6.08 2.5	(hr) (percent) (percent) 66-006 3. 11e-008 1. 33e-007 8 21 42. 7 48. 7 0
	Persistence Time:3.02e+003 hrReaction Time:5.28e+003 hrAdvection Time:7.06e+003 hrPercent Reacted:57.2Percent Advected:42.8	
	Half-Lives (hr), (based upon Biowir Air: 296.4 Water: 3600 Soil: 3600 Sediment: 1.44e+004 Biowin estimate: 1.541 (recal	· · · ·
	Advection Times (hr): Air: 100 Water: 1000 Sediment: 5e+004	
Test substance Reliability Flag	g : Acifluorfen CAS 50594-66-6 : (2) valid with restrictions : Critical study for SIDS endpoint	
27.12.2001		(

3.3.2 DISTRIBUTION

3.4 MODE OF DEGRADATION IN ACTUAL USE

3.5 **BIODEGRADATION**

ld 50594-66-6

Date 27.12.2001

Type Inoculum	:	aerobic	
Remark	:	Studies are reported in the EPA RED documentation. This material undergoes aquatic biodegradation with and estimated (EPA) half-life of 1 days.	17
Test substar	ice :	CAS 62476-59-9 (acifluorfen sodium)	
Reliability Flag	:	Expected to biodegrade at essentially the same rate in the environment. (2) valid with restrictions Critical study for SIDS endpoint	
27.12.2001			(4)
3.6 BOD5, C	OD OR BOD5/	COD RATIO	

3.7 BIOACCUMULATION

3.8 ADDITIONAL REMARKS

4. Ecotoxicity

4.1 ACUTE/PROLONGED TOXICITY TO FISH

4.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES

4.3 TOXICITY TO AQUATIC PLANTS E.G. ALGAE

 Anabaena flos-aquae (Algae) other: biomass/growth rate 120 hour(s) µg/l yes 355 other: EPA FIFRA 123-2 1982 yes other TS TEST ORGANISMS Species: Anabaena flos-aquae Source/supplier: Carolina Biological Supply Company, Burlington, North Carolina Method of cultivation: stock cultures were maintained under test conditions and transferred to fresh medium once or twice a week. The inoculum used in the tests was extracted from a 5 day old stock culture. Initial cell concentration: 0.3E4 cells/mL
STOCK AND TEST SOLUTION AND THEIR PREPARATION - Vehicle, solvent: none DILUTION WATER
 Source: MBL medium GROWTH/TEST MEDIUM CHEMISTRY Chemistry (P = 1.55 mg/L, N = 14 mg/L, Ca+Mg = 0.40 mmol/L, no EDTA) pH 7.5
 TEST SYSTEM Test type: static Concentrations: 370 ug a.i./L, control Exposure vessel type: 125 mL flask containing 50 mL test solution (covered; shaken at 100 rpm) Number of replicates: 3 Photoperiod: continuous illuminated at 1700-2000 lux PHYSICAL MEASUREMENTS Measuring times: 0 and 120 h Test temperature: 25-26 C pH: 7.4 at 0 hours, 9.2-9.4 at 120 hours

4. Ecotoxicity	ld 50594-66- Date 27.12.200 ⁻	
		1
	TEST PARAMETER: cell counts by a haematocytometer OBSERVATION TIMES: 24, 48, 72, 96 and 120 hours	
	ANALYSES - Method: direct HPLC	
	- Sampling times: 0 and 120 hours	
	STATISTICAL METHOD: t-test, one-way analysis of variance, Dunnett's test, Chi-Square test, Hartley's test, Kruskal-Wallis test	
Result	: RESULTS:	
	- Nominal concentrations (ug a.i./L): 0, 370 - Meas. concentrations (ug a.i./L): 0, 355	
	- Cell density data: see attached document	
	- Inhibition-growth rate: 0, -11%	
	- Inhibition-biomass(AUC): 0, -3%	
	GROWTH FACTOR CONTROL: 100 after 72 hours	
	STATISTICAL RESULTS: no statistical differences in cell densities.	
	ANALYTICAL METHOD:	
	The analytical method was validated by fortifying water samples with 0.025, 0.25 and 3.0 mg/L. The recoveries of	
	this samples (3x3) were 81-103%.	
	QCs (filtered (n=2) and unfiltered (n=2)) fortified at 25,	
	101, 202 ug a.i./L showed recoveries of respectively <loq-159%, 25="" 92-119%.="" 96-106%,="" a.i.="" for="" l="" td="" the="" the<="" ug=""><td></td></loq-159%,>	
	unfiltered samples showed recoveries of 159% (0 h) and 105%	
	(120 h), the filtered samples showed recoveries of 67% (0 h)	
Source	and <loq (120="" h).<br="">: Notox Hertogenbosch</loq>	
Test substance	: III, CAS 50594-66-6 (acifluorfen), purity 43,9%, impurities	
	not specified	
Attached doc. Conclusion	: BASF ref 80A.xls : 120 h EC50 >370 mg a.i./L (nominal)	
	120 h EC50 >355 mg a.i./L (measured)	
Reliability	: (1) valid without restriction	
	 Anabaena is not one of the recommended test species of OECD 203, it is a recommendedn test species of the EPA. 	
	Light intensity was not in accordance with the guidelines	
	(1700-2000 lux, OECD 201 8000 lux, EPA 2200 lux). 2. The medium used was not in accordance with OECD 201 (P:	
	1.55 mg/L, OECD 201 <=0.7 mg/L, N: 14 mg/L, OECD 201 <=10	
	mg/L). Higher P and N values may lead to stronger cell	
	growth during the test. 2. Rises in pH of 2 units were probably associated with	
	strong cell growth due to CO2 depletion from test media and	
	do not invalidate the test, since in controls within 72 hours an adequate growth factor of 60 was determined.	
09.05.2001	nours an adequate growth factor of oo was determined.	(2
		(2)
09.05.2001 Species Endpoint Exposure period	 Navicula pelliculosa (Algae) other: biomass/growth rate 120 hour(s) 	(2)

4. Ecotoxicity			50594-66-6 27.12.2001
Unit Analytical monitoring NOEC EC50 Method Year GLP Test substance Method	 Burlington, North Carolina Method of cultivation: sto under test conditions and to or twice a week. The inocule extracted from a 8 day old Initial cell concentration: STOCK AND TEST SOLU Vehicle, solvent: none DILUTION WATER Source: MBL medium GROWTH/TEST MEDIUM Chemistry (P = 1.55 mg/l mmol/L, no EDTA) 	a Biological Supply Company, ck cultures were maintained transferred to fresh medium on ilum used in the tests was stock culture. 0.3E4 cells/mL TION AND THEIR PREPARAT	
	solution (covered; shaken - Number of replicates: 3	25 mL flask containing 50 mL te at 100 rpm) illuminated at 4000-5000 lux NTS I 20 h C	est
	TEST PARAMETER: cell of	counts by a haematocytometer 4, 48, 72, 96 and 120 hours	
Result	 STATISTICAL METHOD: 1 Dunnett's test, Chi-Square Kruskal-Wallis test RESULTS: Nominal concentrations (Meas. concentrations (ug) Cell density data: see att 	ug a.i./L): 0, 370 j a.i./L): 0, 345	ance,

4. Ecotoxicity	ld 50594-66-6 Date 27.12.2001	
	- Inhibition-growth rate: 0, -3% - Inhibition-biomass(AUC): 0, -7%	
	GROWTH FACTOR CONTROL: 87 after 72 hours	
	STATISTICAL RESULTS: no statistical differences in cell densities.	
	ANALYTICAL METHOD: The analytical method was validated by fortifying water samples with 0.025, 0.25 and 3.0 mg/L. The recoveries of this samples (3x3) were 81-103%.	
Source Test substance	 QCs (filtered (n=2) and unfiltered (n=2)) fortified at 25, 101, 202 ug a.i./L showed recoveries of respectively <loq-159%, (0="" (120="" 105%="" 159%="" 25="" 67%="" 92-119%.="" 96-106%,="" <loq="" a.i.="" and="" filtered="" for="" h)="" h),="" h).<="" l="" li="" of="" recoveries="" samples="" showed="" the="" ug="" unfiltered=""> Notox Hertogenbosch III, CAS 50594-66-6 (acifluorfen), purity 43,9%, impurities </loq-159%,>	
Attached doc.	not specified BASF ref 80B.xls	
Conclusion	: 120 h EC50 370 ug/L (nominal) 120 h EC50 345 ug/L (measured)	
Reliability	 (1) valid without restriction 1. Navicula pelliculosa is not one of the recommended test species of OECD 203, it is a recommended test species of the EPA. Light intensity was not in accordance with the OECD guideline (4000-5000 lux, OECD 201 8000 lux, EPA 4300 lux). 2. The medium used was not in accordance with OECD 201 (P: 1.55 mg/L, OECD 201 <=0.7 mg/L, N: 14 mg/L, OECD 201 <=10 mg/L). Higher P and N values may lead to stronger cell growth during the test. 	
09.05.2001	giowin during the test.	(2
Species Endpoint Exposure period Unit Analytical monitoring NOEC EC50 Method Year GLP Test substance Method	 Selenastrum capricornutum (Algae) other: growth rate, biomass 120 hour(s) µg/l yes 260 > 260 other: EPA FIFRA 123-2 1982 yes other TS TEST ORGANISMS Species: Selenastrum capricornutum Source/supplier: Carolina Biological Supply Company, Burlington, North Carolina Method of cultivation: stock cultures were maintained under test consitions and transferred to fresh medium once or twice a week. The inoculum used in the tests was extracted from a 7 day old stock culture. 	

4. Ecotoxicity	Id 50594-66-6
	Date 27.12.2001
	STOCK AND TEST SOLUTION AND THEIR PREPARATION - Vehicle, solvent: none
	DILUTION WATER - Source: MBL medium
	GROWTH/TEST MEDIUM CHEMISTRY - Chemistry (P = 1.55 mg/L, N = 14 mg/L, Ca+Mg = 0.40 mmol/L, no EDTA) - pH 7.5 TEST SYSTEM - Test type: static - Concentrations: 24, 47, 93, 185, 370 ug a.i./L, control - Exposure vessel type: 125 mL flask containing 50 mL test solution (covered; shaken at 100 rpm) - Number of replicates: 3 - Photoperiod: continuous illuminated at 4000-5000 lux PHYSICAL MEASUREMENTS - Measuring times: 0 and 120 h
	- Test temperature: 25-26 C - pH: 7.4 at 0 hours, 9.7-10.4 at 120 hours
	DURATION OF TEST: 120 hours
	TEST PARAMETER: cell counts by a haematocytometer OBSERVATION TIMES: 24, 48, 72, 96 and 120 hours
	ANALYSES - Method: direct HPLC - Sampling times: 0 and 120 hours
Result	 STATISTICAL METHOD: t-test, one-way analysis of variance, Dunnett's test, Chi-Square test, Hartley's test, Kruskal-Wallis test RESULTS: Nominal concentrations (ug a.i./L): 0, 24, 47, 93, 185,
	370 - Meas. concentrations (ug a.i./L): 0, 19, 38, 88, 160, 260 - Cell density data: see attached document - Inhibition-growth rate [%]: 0, -2, 0, 0, 0, 0 - Inhibition-biomass(AUC) [%]: 0, -12, -3, -3, -1, 0
	GROWTH FACTOR CONTROL: 144 after 72 hours
	STATISTICAL RESULTS: no statistical differences in cell densities.
	ANALYTICAL METHOD: The analytical method was validated by fortifying water samples with 0.025, 0.25 and 3.0 mg/L. The recoveries of this samples (3x3) were 81-103%.
	QCs (filtered (n=2) and unfiltered (n=2)) fortified at 25, 101, 202 ug a.i./L showed recoveries of respectively <loq-159%, 25="" 92-119%.="" 96-106%,="" a.i.="" for="" l="" the="" the<br="" ug="">unfiltered samples showed recoveries of 159% (0 h) and 105% (120 h), the filtered samples showed recoveries of 67% (0 h)</loq-159%,>
	16 / 23

4. Ecotoxicity	ld 50594-66-6 Date 27.12.2001
	and $d OO (120 h)$
Source	and <loq (120="" h).<br="">: Notox Hertogenbosch</loq>
Test substance	: III, CAS 50594-66-6 (acifluorfen), purity 43,9%, impurities
	not specified
Attached doc.	: BASF ref 80.xls
Conclusion	: 120 h EC50 >370 mg a.i./L (nominal) 120 h EC50 >260 mg a.i./L (measured)
Reliability	: (1) valid without restriction
Ronability	1. The medium used was not in accordance with OECD 201 (P:
	1.55 mg/L, OECD 201 <=0.7 mg/L, N: 14 mg/L, OECD 201 <=10
	mg/L). Higher P and N values may lead to stronger cell
	growth during the test. Light intensity was lower than
	recommended (4000-5000 lux, OECD 201 8000 lux), which could decrease the cell growth.
	2. Rises in pH of 2-3 units were probably associated with
	strong cell growth due to CO2 depletion from test media and
	do not invalidate the test, since in controls within 72
	hours an adequate growth factor of 144 was determined.
09.05.2001	(2)
Species	: Skeletonema costatum (Algae)
Endpoint	: other: biomass/growth rate
Exposure period	: 120 hour(s)
Unit	: µg/l
Analytical monitoring	: yes
NOEC	: 300
EC50 Method	: > 300 : other: EPA FIFRA 123-2
Year	: 1982
GLP	: Yes
Test substance	: other TS
Method	: TEST ORGANISMS
	- Species: Skeletonema costatum
	- Source/supplier: Bigelow marine Laboratory, West Boothbay,
	Maine - Method of cultivation: stock cultures were maintained
	under test conditions and transferred to fresh medium once
	or twice a week. The inoculum used in the tests was
	extracted from a 9 day old stock culture.
	- Initial cell concentration: 1.0E4 cells/mL
	STOCK AND TEST SOLUTION AND THEIR PREPARATION - Vehicle, solvent: none
	DILUTION WATER
	- Source: Artificially Enriched Seawater prepared with
	filtered natural seawater
	GROWTH/TEST MEDIUM CHEMISTRY - Chemistry (P = 0.44 mg/L, N = 8.2 mg/L, no EDTA, salinity
	not indicated)
	- pH 8.0
	TEST SYSTEM
	- Test type: static
	 Concentrations: 370 ug a.i./L, control Exposure vessel type: 125 mL flask containing 50 mL test
	Exposure vesser type. The mask containing of the test

4. Ecotoxicity	ld 50594-66-6 Date 27.12.2001	
	solution (covered; shaken at 60 rpm) - Number of replicates: 3 - Photoperiod: 16 hours light (4000-5000 lux) PHYSICAL MEASUREMENTS - Measuring times: 0 and 120 h - Test temperature: 20-23 C - pH: 8.2-8.9	
	DURATION OF TEST: 120 hours	
	TEST PARAMETER: cell counts by a haematocytometer OBSERVATION TIMES: 24, 48, 72, 96 and 120 hours	
	ANALYSES - Method: direct HPLC - Sampling times: 0 and 120 hours	
Result	 STATISTICAL METHOD: t-test, one-way analysis of variance, Dunnett's test, Chi-Square test, Hartley's test, Kruskal-Wallis test RESULTS: Nominal concentrations (ug a.i./L): 0, 370 Meas. concentrations (ug a.i./L): 0, 300 Cell density data: see attached document Inhibition-growth rate: 0, 0% Inhibition-biomass(AUC): 0, 1% 	
	GROWTH FACTOR CONTROL: 59 after 72 hours	
	STATISTICAL RESULTS: no statistical differences in cell densities.	
	ANALYTICAL METHOD: The analytical method was validated by fortifying water samples with 0 and 379 ug/L. The recoveries of this samples (n=3) were 100-101%.	
Source Test substance	 QCs (n=2x2) fortified at 101, 202 and 303 mg a.i./L showed recoveries of 96-107% (filtered) and 69-84% (unfiltered). Notox Hertogenbosch III, CAS 50594-66-6 (acifluorfen), purity 43,9%, impurities not specified 	
Attached doc. Conclusion	: BASF ref 80C.xls : 120 h EC50 370 ug/L (nominal)	
Reliability	 120 h EC50 300 ug/L (measured) (1) valid without restriction 1. Skeletonema costatum is not one of the recommended test species of OECD 203, but a marine diatom recommended by the EPA. Light intensity was not in accordance with the OECD guideline (4000-5000 lux, OECD 201 8000 lux, EPA 4300 lux). 2. Salinity was not indicated, but since natural seawater was used for the preparation of the test medium, the reliability was not lowered. 	
	3. The QCs were reported to be fortified at 101-303 mg a.i./L. Probably this is a reporting error and the actual	
09.05.2001	fortification was 101-303 ug a.i./L.	(2)

		اما	50504 66 6
4. E	cotoxicity		50594-66-6
		Date	27.12.2001
4.4	TOXICITY TO MICROORGANISMS E.G. BACTERIA		
4.5.1	CHRONIC TOXICITY TO FISH		
4.5.2	CHRONIC TOXICITY TO AQUATIC INVERTEBRATES		
4.6.1	TOXICITY TO SOIL DWELLING ORGANISMS		
4.6.2	TOXICITY TO TERRESTRIAL PLANTS		
4.6.3	TOXICITY TO OTHER NON-MAMM. TERRESTRIAL SPECIES		
. –			
4.7	BIOLOGICAL EFFECTS MONITORING		
4.8	BIOTRANSFORMATION AND KINETICS		
4.0			
4.9	ADDITIONAL REMARKS		

5. Toxicity	ld 50594-66-6 Date 27.12.2001
5.1.1 ACUTE ORAL TOX	ICITY
5.1.2 ACUTE INHALATIO	
5.1.3 ACUTE DERMAL 1	ΟΧΙCΙΤΥ
5.1.4 ACUTE TOXICITY,	OTHER ROUTES
5.2.1 SKIN IRRITATION	
5.2.2 EYE IRRITATION	
5.3 SENSITIZATION	
5.4 REPEATED DOSE	ΤΟΧΙCΙΤΥ
5.5 GENETIC TOXICIT	Y 'IN VITRO'
Type System of testing Concentration Cycotoxic conc. Metabolic activation Result Method Year GLP Test substance Method	 Ames test TA98, TA100, TA1535 and TA1537 20-5000 ug/plate 5000 ug/plate with and without negative OECD Guide-line 471 "Genetic Toxicology: Salmonella thyphimurium Reverse Mutation Assay" other TS SYSTEM OF TESTING:
Wethod	 SYSTEM OF TESTING: Species/cell type: Salmonella typhimurium TA98, TA100, TA1535, TA1537. Deficiences/Proficiences: histidine Metabolic activation system: rat S9 mix (Arochlor 1254 induced) ADMINISTRATION: Dosing: 0, 20, 100, 500, 2500 and 5000 µg/plate: Number of replicates: 3 Application: DMSO Positive and negative control groups and treatment:
	i ostavo ana nogativo ostitioi groupo ana trodunonti.

5. Toxicity	ld 50594-66-6
or reality	Date 27.12.2001
	Without S-9: 2-N-methyl-N'-nitroso-guanidine (MNNG) (TA100 and TA1535); 4-nitro-o-phenylenediamine (TA98); 9-aminoacridine chloride monohydrate (TA1537) With S-9: 2-aminoantharacene Negative controls: DMSO - type of test: direct plate assay
Result	CRITERIA FOR EVALUATING RESULTS: number of revertant coloniesNo precipitation was observed.
Source Test substance Reliability 16.05.2001	 Slight toxicity to strains TA1535 and TA100 at 5000 ug/plate. Notox Hertogenbosch CAS 50594-66-6, (5-(2-chloro-4-trifluoromethylphenoxy) -2-nitrobenzoic acid), purity 99.5% (2) valid with restrictions Test results for the purity and stability of the compound are not included in the report. Only 4 strains of bacteria are used (OECD 471: at least 5 strains) 2-aminoanthracene alone as positive control is not sufficient according to OECD guideline 471. However, as the positive control induced a sufficient number of revertant colonies, reliability is not lowered.
5.6 GENETIC TOXIC	ICITY 'IN VITRO'
5.7 CARCINOGENI	ITY REP RODUCTION
5.8 TOXICITY TO RE	
	TAL TOXICITY/TERATOGENICITY
	TAL TOXICITY/TERATOGENICITY : Notox Hertogenbosch
 5.9 DEVELOPMENT Source 02.04.2001 	
 5.9 DEVELOPMENT Source 02.04.2001 5.10 OTHER RELEVA 	: Notox Hertogenbosch

6. Refere	Id 50594-66-6 Date 27.12.2001
(1)	BASF Aktiengesellschaft, Report on the study of Acifluoren-Reinwirkstoff in the Ames Test, 1990
(2)	BASF, Acifluorfen (BAS 9048 H): toxicity to the growth and reproduction of aquatic plants, 1990 (80)
(3)	BASF, Artificial Sunlight Photolysis of Acifluorfen in Aqueous Media at pH 7.0 (1993) (87).
(4)	EFED Ecological Risk Assessment for sodium acifluorfen. US EPA, Registration Process Documents, June 2000. http://www.epa.gov/pesticides/reregistration/acifluorfen/efedchapter.pdf
(5)	EPIWIN v3.05, Syracuse Research Corporation, Syracuse, NY (July 12, 2000)
(6)	Lyman, W. J. et al. (1990). Handbook of Chemical PropertyEstimation Methods, pp. 7-4, Amer. Chem. Society,Washington, DC
(7)	SRC PHYSPROP Database. http://esc.syrres.com/interkow/physdemo.htm

7.	Risk Assessment	ld	50594-66-6
		Date	27.12.2001
7.1	END POINT SUMMARY		
7.2	HAZARD SUMMARY		
7.3	RISK ASSESSMENT		

IUCLID

Data Set

Existing Chemical CAS No. Generic name	 ID: 63734-62-3 63734-62-3 benzoic acid, 3-[2-chloro-4-(trifluoromethyl)phenoxy]
Producer Related Part Company Creation date	: Toxicology and Regulatory Affairs : 27.12.2001
Substance Related Part Company Creation date	Toxicology and Regulatory Affairs27.12.2001
Memo	:
Printing date Revision date Date of last Update	: 27.12.2001 : : 27.12.2001
Number of Pages	: 24
Chapter (profile) Reliability (profile) Flags (profile)	 Chapter: 1, 2, 3, 4, 5, 7 Reliability: without reliability, 1, 2, 3, 4 Flags: without flag, confidential, non confidential, WGK (DE), TA -Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. Ge	eneral Information	63734-62-3 27.12.2001
1.0.1	OECD AND COMPANY INFORMATION	
1.0.2	LOCATION OF PRODUCTION SITE	
1.0.3	IDENTITY OF RECIPIENTS	
1.1	GENERAL SUBSTANCE INFORMATION	
1.1.0	DETAILS ON TEMPLATE	
1.1.1	SPECTRA	
1.2	SYNONYMS	
1.3	IMPURITIES	
1.4	ADDITIVES	
1.5	QUANTITY	
1.6.1	LABELLING	
1.6.2	CLASSIFICATION	
1.7	USE PATTERN	
1.7.1	TECHNOLOGY PRODUCTION/USE	
1.8	OCCUPATIONAL EXPOSURE LIMIT VALUES	
1.9	SOURCE OF EXPOSURE	
	RECOMMENDATIONS/PRECAUTIONARY MEASURES	

1. General Information

1.10.2 EMERGENCY MEASURES

- 1.11 PACKAGING
- 1.12 POSSIB. OF RENDERING SUBST. HARMLESS
- 1.13 STATEMENTS CONCERNING WASTE
- 1.14.1 WATER POLLUTION
- 1.14.2 MAJOR ACCIDENT HAZARDS
- 1.14.3 AIR POLLUTION
- 1.15 ADDITIONAL REMARKS
- 1.16 LAST LITERATURE SEARCH

1.17 REVIEWS

1.18 LISTINGS E.G. CHEMICAL INVENTORIES

2.1 MELTING POINT

Value Sublimation Method Year GLP Test substance Method Result		ca. 146 °C 2001 no Estimation using MPBPWIN v1.40 in EPIWIN 3.05 MPBPWIN (v1.40) Program Results: 	(1)
2.2 BOILING POINT			
2.3 DENSITY			
2.3.1 GRANULOMETRY			
2.4 VAPOUR PRESSUR	Е		
Value Decomposition Method Year GLP Test substance Method Result		 = .0000029 hPa at ° C other (calculated) 2001 no Estimation using MPBPWIN v1.40 in EPIWIN 3.05 MPBPWIN (v1.40) Program Results: 	
		Experimental Database Structure Match: no data SMILES : c1(CL)cc(C(F)(F)(F))ccc1Oc2cccc(C(=O)O)c2 4 / 24	

	CHEM : Trifluorobenzoic acid CAS 63734-62-3 MOL FOR: C14 H8 CL1 F3 O3 MOL WT : 316.67	
	SUMMARY MPBPWIN v1.40	
	Vapor Pressure Estimations (25 deg C): (Using BP: 387.24 deg C (estimated)) (Using MP: 146.30 deg C (estimated)) VP: 2.66E-007 mm Hg (Antoine Method) VP: 9.96E-007 mm Hg (Modified Grain Method) VP: 2.18E-006 mm Hg (Mackay Method) Selected VP: 9.96E-007 mm Hg (Modified Grain Method)	
Test substance Reliability Flag 27.12.2001	 3-[2-chloro-4-(trifluoromethyl)phenoxy]benzoic acid CAS 63734-62-3 (2) valid with restrictions Critical study for SIDS endpoint 	(1)
2.5 PARTITION COEFF	ICIENT	
Log pow Method Year GLP Test substance Method Test substance Reliability Flag 27.12.2001	 ca. 4.7 at 25° C 2001 no Estimation using KOWWIN v1.66 in EPIWIN 3.05 3-[2-chloro-4-(trifluoromethyl)phenoxy]benzoic acid CAS 63734-62-3 (2) valid with restrictions Critical study for SIDS endpoint 	(1)
2.6.1 WATER SOLUBILIT	~	
2.0.1 WATER SOLUBILIT		
Value Qualitative Pka PH Method Year GLP Test substance Method	ca. 1 mg/l at 25 ° C at 25 ° C at and ° C 2001 no Estimation using WSKOW v1.40 in EPIWIN 3.05	
Result	: Water Sol from Kow (WSKOW v1.40) Results:	
	Water Sol: 0.9521 mg/L SMILES : c1(CL)cc(C(F)(F)(F))ccc1Oc2cccc(C(=O)O)c2 CHEM : Trifluorobenzoic acid CAS 63734-62-3 MOL FOR: C14 H8 CL1 F3 O3 MOL WT : 316.67 - WSKOW v1.40 Results Log Kow (estimated) : 4.70	

ld 63734-62-3 Date 27.12.2001

Log Kow (experimental): not available from database Log Kow used by Water solubility estimates: 4.70 Equation Used to Make Water Sol estimate: Log S (mol/L) = 0.796 - 0.854 log Kow - 0.00728 MW + Correction (used when Melting Point NOT available) Correction(s): Value 0.000 Acid, aromatic Log Water Solubility (in moles/L) : -5.522 Water Solubility at 25 deg C (mg/L): 0.9521 Test substance : 3-[2-chloro-4-(trifluoromethyl)phenoxy]benzoic acid CAS 63734-62-3 Reliability (2) valid with restrictions : : Critical study for SIDS endpoint 27.12.2001 (1) 2.6.2 SURFACE TENSION **FLASH POINT**

2.8 AUTO FLAMMABILITY

2.9 FLAMMABILITY

Flag

2.7

2.10 EXPLOSIVE PROPERTIES

2.11 **OXIDIZING PROPERTIES**

2.12 ADDITIONAL REMARKS

3.1.1 PHOTODEGRADATION

Type Light source Light spect. Rel. intensity Indirect photolysis Sensitizer Conc. of sens. Rate constant Degradation Method	 air nm based on Intensity of Sunlight OH 1500000 cm3/(molecule*sec) % after Estimation using APOWIN v1.90 in EPIWIN 3.05 	
Result	: AOP Program (v1.90) Results:	
	======================================	
	- SUMMARY (AOP v1.90): HYDROXYL RADICALS	
	Hydrogen Abstraction = 0.0000 E-12 cm3/molecule-sec Reaction with N, S and -OH = 0.5200 E-12 cm3/molecule-sec Addition to Triple Bonds = 0.0000 E-12 cm3/molecule-sec Addition to Olefinic Bonds = 0.0000 E-12 cm3/molecule-sec **Addition to Aromatic Rings = 1.3056 E-12 cm3/molecule-sec Addition to Fused Rings = 0.0000 E-12 cm3/molecule-sec	
	OVERALL OH Rate Constant = 1.8256 E-12 cm3/molecule-sec HALF-LIFE = 5.859 Days (12-hr day; 1.5E6 OH/cm3) HALF-LIFE = 70.306 Hrs	
	** Designates Estimation(s) Using ASSUMED Value(s)	
Test substance Reliability Flag 27.12.2001	 3-[2-chloro-4-(trifluoromethyl)phenoxy]benzoic acid CAS 63734-62-3 (2) valid with restrictions Critical study for SIDS endpoint 	(1)
3.1.2 STABILITY IN WA	TER	
Type t1/2 pH4 t1/2 pH7 t1/2 pH9 Deg. Product Method	 > 1 year at 25 degree C 	
Year	- 2001	

3.

туре	
t1/2 pH4	: > 1 year at 25 degree C
t1/2 pH7	: > 1 year at 25 degree C
t1/2 pH9	: > 1 year at 25 degree C
Deg. Product	:
Method	:
Year	: 2001
GLP	: no
Test substance	:
Method	: Estimated on chemical principles based on absence of groups susceptible to hydrolysis
Remark	: The estimation program in EPIWIN has no capability to estimate hydrolysis rates for this compound.

Result	: This material has no groups that are susceptible to hydrolysis in the pH 4 to 9 range; therefore, it is considered stable to hydrolysis in surface and groundwater. It is estimated to have a hydrolysis half-life of greater than one year between pH 4 and pH 9.
Test substance Reliability Flag 27.12.2001	 3-[2-chloro-4-(trifluoromethyl)phenoxy]benzoic acid CAS 63734-62-3 (2) valid with restrictions Critical study for SIDS endpoint

3.1.3 STABILITY IN SOIL

3.2 MONITORING DATA

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

Method : The Fugacity was determined using the EQC Level III model as found in EPIWIN 3.05. Estimated values were used for physical constants. Biodegradation was based on the EPIWIN derived estimates (Biowin, Ultimate) that were assessed for reasonableness compared with similar compounds. Half life in air was determined from the APOWIN program. Direct photolysis was not considered in this model. Emissions were calculated from only water and soil as this test substance it is almost non volatile. Other parameters used the default values found in EPIWIN. Result : Level III Fugacity Model (Full-Output): Chem Name : Trifluorobenzoic acid CAS 63734-62-3 Molecular Wt: 316.67 Henry's LC : 1.53e-008 atm-m3/mole (Henrywin program) Vapor Press 9.96e-007 mm Hg (Mpbpwin program) Liquid VP : 1.58e-005 mm Hg (super-cooled) Melting Pt : 146 deg C (Mpbpwin program) Log Kow : 4.7 (Kowwin program) Soil Koc : 2.05e+004 (cal c by model)
Chem Name : Trifluorobenzoic acid CAS 63734-62-3 Molecular Wt: 316.67 Henry's LC : 1.53e-008 atm-m3/mole (Henrywin program) Vapor Press : 9.96e-007 mm Hg (Mpbpwin program) Liquid VP : 1.58e-005 mm Hg (super-cooled) Melting Pt : 146 deg C (Mpbpwin program) Log Kow : 4.7 (Kowwin program) Soil Koc : 2.05e+004 (calc by model) Concentration Half-Life Emissions
Air2.57e-0051410Water191.44e+0031000Soil63.41.44e+0031000Sediment17.75.76e+0030
Fugacity (atm)Reaction (kg/hr)Advection (kg/hr)Reaction (percent)Advection (percent)Air6.15e-0160.004150.008420.0002070.000421Water1.45e-01329962114.931.1Soil1.13e-014999049.90Sediment1.41e-01369.611.63.480.578
Persistence Time: 1.64e+003 hr Reaction Time: 2.4e+003 hr

ld 63734-62-3 Date 27.12.2001

Advection Time: 5.18e+003 hr Percent Reacted: 68.4 Percent Advected: 31.6 Half-Lives (hr), (based upon Biowin (Ultimate) and Aopwin): Ai r: 140.6 Water: 1440 1440 Soil: Sediment: 5760 Biowin estimate: 1.810 (months) Advection Times (hr): 100 Ai r: 1000 Water: Sediment: 5e+004 Test substance : 3-[2-chloro-4-(trifluoromethyl)phenoxy]benzoic acid CAS 63734-62-3 Reliability : (2) valid with restrictions : Critical study for SIDS endpoint Flag 27.12.2001 (1) 3.3.2 DISTRIBUTION 3.4 MODE OF DEGRADATION IN ACTUAL USE 3.5 BIODEGRADATION BOD5, COD OR BOD5/COD RATIO 3.6 3.7 BIOACCUMULATION

3.8 ADDITIONAL REMARKS

4.1 ACUTE/PROLONGED TOXICITY TO FISH

Type Species Exposure period Unit Analytical monitoring NOEC LC50 Method Year GLP Test substance Method		static Lepomis macrochirus (Fish, fresh water) 96 hour(s) mg/l 180 > 1000 other: EPA 1975 no other TS TEST ORGANISMS - Species: Lepomis macrochirus Rafinesque - Supplier: commercial hatchery in Nebraska - Age;size;weight;loading: ~4 months; 28-44 mm; 0.20-1.10 g; 0.3-0.4 g/L - Feeding during test: none, feeding was discontinued 48 hours prior to test initiation STOCK AND TEST SOLUTION AND THEIR PREPARATION - Vehicle, solvent: none - Other procedures: direct addition of the test substance to the test vessels DILUTION WATER - Source: Well water (Tarrytown site) - Chemistry (Alkalinity 32 mg CaCO3/L;Hardness 46 mg CaCO3/L/pH 7.70/Conductance 150 umhos/cm) TEST SYSTEM - Test type: static - Concentrations: 0, 100, 180, 320, 560 and 1000 mg/L - Exposure vessel type: 20 L glass vessels containing 15 L of water - Number of fish: 10 per treatment - Photoperiod: not indicated PHYSICAL MEASUREMENTS - Measuring times: 0, 48, 96 hours - Test temperature: 22-23 C - Dissolved oxygen: 61-101% - pH: 7.3-7.7 DURATION OF THE TEST: 96 hours TEST PARAMETER: mortality/symptoms
Result	:	OBSERVATION TIMES: daily STATISTICAL METHOD: not indicated RESULTS: - Mortality: no mortality
Source Test substance	:	 Other effects: irritated, exhibited abnormal sounding behaviour and/or dark discolouration at 320-1000 mg/L. REFERENCE SUBSTANCE: 96 h LC50 4.03 ug/L (3.59-4.52 ug/L) Notox Hertogenbosch III, CAS 63734-62-3: TD 77-373 (RH-41,833 W. Liq. (2.6 eq.)), purity not indicated

Reliability	:	 (2) valid with restrictions 1. No analyses were performed to confirm the nominal test concentrations. The study reliability was lowered because of this. 2. Fish were fasted longer than recommended (48 h, OECD 203 24 h). This may have increased the susceptibility of the fish. 3. The used fish were larger than recommended by the guideline of the OECD, but acceptable according to the EG-guideline (28-44 mm, OECD 20+/-10 mm, EG 50+/-20 mm). 4. The test substance was specified as TD 77 -373 (RH-41,833 W. Liq. (2.6 eq.)). No information was available on the composition of this compound. 	
09.05.2001			(9)
Type Species Exposure period Unit Analytical monitoring NOEC LC50 Method Year GLP Test substance Method		static Lepomis macrochirus (Fish, fresh water) 96 hour(s) mg/l no data 180 > 1000 other: EPA 660/375-009 1975 no other TS TEST ORGANISMS - Species: Lepomis macrochirus Rafinesque - Supplier: commercial hatchery in Nebraska - Age;size;weight;loading: -4 months; ? mm; ~0.68-0.78 g; 0.5 g/L - Feeding during test: none, feeding was discontinued 48 hours prior to test initiation STOCK AND TEST SOLUTION AND THEIR PREPARATION - Vehicle, solvent: none - Other procedures: direct addition of the test substance to the test vessels DILUTION WATER - Source: Well water (Tarrytown site) - Other procedures: direct addition of the test substance to the test vessels DILUTION WATER - Source: Well water (Tarrytown site) - Otherprocedures: 0, 100, 180, 320, 560 and 1000 mg/L - Exposure vessel type: 20 L glass vessels containing 15 L of water - Number of fish: 10 per treatment - Photoperiod: not indicated PHYSICAL MEASUREMENTS - Measuring times: 0, 48, 96 hours at control, low, middle and high dose - Test temperature: 22-23 C - Dissolved oxygen: Control: 101/61/56 at respectively 0/24/48 h 300 & 1000 mg/L: 100/20/16-18 at respectively 0/24/48 h 320 & 1000 mg/L: 100/20/16-18 at respectively 0/24/48 h	
 		11/24	

	TEST PARAMETER: mortal OBSERVATION TIMES: dail		
	REFERENCE SUBSTANCE	: p,p'-DDT	
Result	STATISTICAL METHOD: no RESULTS: - Mortality: no mortality - Other effects: quiescence, a swimming and/or gulping of a	abnormal surfacing, erratic	
Source Test substance	Notox Hertogenbosch	: 96 h LC50 4.03 ug/L (3.59-4.52 ug/L) 370 (RH-41,833 HOAc ppt (2.6	
	eq.)), purity not indicated		
Reliability	concentrations. The study re this.	ned to confirm the nominal test liability was lowered because of s dropped to minimal 16% at the	
	 end of the test (OECD 203 > fasted longer than recommendations may have increased 3. There was no information organisms, since table 3 of the information) was missing. 4. The test substance was specified to the set of the se	60%). Further the fish were nded (48 h, OECD 203 24 h). Both the susceptibility of the fish. on the length of the test he report (containing this pecified as TD 77-370 (RH-41,833 rmation was available on the	
09.05.2001	composition of this compoun	d.	(8)
Turne	atatia		
Type Species Exposure period	static Pimephales promelas (Fish, 96 hour(s)	fresh water)	
Species	Pimephales promelas (Fish,	fresh water)	
Species Exposure period Unit Analytical monitoring NOEC	Pimephales promelas (Fish, 96 hour(s) mg/l no data 1.4	fresh water)	
Species Exposure period Unit Analytical monitoring	Pimephales promelas (Fish, 96 hour(s) mg/l no data	fresh water)	
Species Exposure period Unit Analytical monitoring NOEC LC50 Method Year	Pimephales promelas (Fish, 96 hour(s) mg/l no data 1.4 2.6 other: EPA 660/3-75-009 1975	fresh water)	
Species Exposure period Unit Analytical monitoring NOEC LC50 Method	Pimephales promelas (Fish, 96 hour(s) mg/l no data 1.4 2.6 other: EPA 660/3-75-009	fresh water)	
Species Exposure period Unit Analytical monitoring NOEC LC50 Method Year GLP	Pimephales promelas (Fish, 96 hour(s) mg/l no data 1.4 2.6 other: EPA 660/3-75-009 1975 no other TS TEST ORGANISMS - Species: Pimephales prom - Supplier: commercial fish fa - Size;weight;loading: 44+/-3	elas armer in Arkansas	
Species Exposure period Unit Analytical monitoring NOEC LC50 Method Year GLP Test substance	Pimephales promelas (Fish, 96 hour(s) mg/l no data 1.4 2.6 other: EPA 660/3-75-009 1975 no other TS TEST ORGANISMS - Species: Pimephales prom - Supplier: commercial fish fa - Size;weight;loading: 44+/-3 - Feeding during test: not fee hours prior to the test	elas armer in Arkansas 3.9 mm;0.75+/-0.30 g;0.5 g/L	
Species Exposure period Unit Analytical monitoring NOEC LC50 Method Year GLP Test substance	Pimephales promelas (Fish, 96 hour(s) mg/l no data 1.4 2.6 other: EPA 660/3-75-009 1975 no other TS TEST ORGANISMS - Species: Pimephales prom - Supplier: commercial fish fa - Size;weight;loading: 44+/-3 - Feeding during test: not fee hours prior to the test STOCK AND TEST SOLUTI	elas armer in Arkansas 8.9 mm;0.75+/-0.30 g;0.5 g/L d (feeding was discontinued 48 ON AND THEIR PREPARATION	
Species Exposure period Unit Analytical monitoring NOEC LC50 Method Year GLP Test substance	Pimephales promelas (Fish, 96 hour(s) mg/l no data 1.4 2.6 other: EPA 660/3-75-009 1975 no other TS TEST ORGANISMS - Species: Pimephales prom - Supplier: commercial fish fa - Size;weight;loading: 44+/-3 - Feeding during test: not fee hours prior to the test STOCK AND TEST SOLUTI - Vehicle, solvent: acetone DILUTION WATER - Source: Well water - Chemistry (Alkalinity/Hardn TEST SYSTEM - Test type: static - Concentrations: 0 (untr), 0 4.2, 6.5, 10, 18 mg/L	elas armer in Arkansas 3.9 mm;0.75+/-0.30 g;0.5 g/L d (feeding was discontinued 48 ON AND THEIR PREPARATION ess 35 mg CaCO3/L/pH 7.1)	

4.3 TOXICITY TO AQUATIC PLANTS E.G. ALGAE

4.4 TOXICITY TO MIC ROORGANISMS E.G. BACTERIA

4.5.1 CHRONIC TOXICITY TO FISH

4. Ecotoxicity

4.5.2 CHRONIC TOXICITY TO AQUATIC INVERTEBRATES

- 4.6.1 TOXICITY TO SOIL DWELLING ORGANISMS
- 4.6.2 TOXICITY TO TERRESTRIAL PLANTS
- 4.6.3 TOXICITY TO OTHER NON-MAMM. TERRES TRIAL SPECIES
- 4.7 BIOLOGICAL EFFECTS MONITORING
- 4.8 BIOTRANSFORMATION AND KINETICS
- 4.9 ADDITIONAL REMARKS

5.1.1 ACUTE ORAL TOXICITY

Type Species Strain Sex Number of animals Vehicle Value Method Year GLP Test substance Method	 LD50 rat other: CF Nelson male 5 other: oil = 1170 mg/kg bw other: not indicated no other TS TEST ORGANISMS: Source: not indicated Age: not indicated Number: 5/dose Weight at study initiation: 189-199 g (mean) Controls: no 	
	 Doses: 625, 1250 and 2500 mg/kg bw Doses per time period: single concentration: 20% w/v Post dose observation period: 14 days food withheld for 24 hours pre-dosing EXAMINATIONS: signs for toxicity and gross necropsy 	
	BODY WEIGHT: pre-dosing and at termination of study	
Result	 STATISTICAL METHOD: not indicated MORTALITY: Number of deaths at each dose: 625, 1250 and 2500 mg/kg bw: 0/5, 3/5 and 5/5, respectively Time of death: for the highest dose: within 24 hours; for 1250 mg/kg bw: within 4 days 	
	CLINICAL SIGNS: lethargy, prostration at 2500 mg/kg bw (0-6 hours)	
	BODY WEIGHT: survivors increased bw	
Source Test substance	 NECROPSY FINDINGS: survivors normal, at 2500 mg/kg decendents were normal, at 1250 mg/kg one decendent had blood in small intestines. Notox Hertogenbosch III, 63734-62-3 (3-[2-chloro-4-(trifluoromethyl)phenoxy]benzoic acid), purity 86.5%, used as 20% dispersion with oil 	
Conclusion Reliability	 LD50 1170 mg/kg bw (3) invalid The information available in the report on the study findings is essentially confined to what is included in the above summary. There is no information on the individual toxicity data. The study is not reliable because the LD50 cannot be back-calculated to the amount of a.i./kg body weight (dosing 	
	45 / 04	

5. Toxicity	ld 63734-62-3 Date 27.12.2001	-
10.04.2001	was done with a 20% weight/volume oil dispersion and no data are available on the density of the oil).	(3)
Type Species Strain Sex Number of animals Vehicle Value Method Year GLP Test substance Method	 LD50 rat other: Charles River CD male 6 other: 0.5% methylcellulose in water solution > 50 mg/kg bw other: not specified no other TS TEST ORGANISMS: Source: not indicated Age: not indicated Number: 6/dose Weight at study initiation: 227-230 g Controls: no 	
	ADMINISTRATION: - Doses: 50 and 500 mg/kg bw - Doses per time period: single - concentration: 10% w/v - Post dose observation period: 14 days - food withheld for 24 hours pre-dosing EXAMINATIONS: signs for toxicity and gross necropsy	
	BODY WEIGHT: pre-dosing and at termination of study	
Result	STATISTICAL METHOD: not indicated MORTALITY: - Number of deaths at each dose: no deaths	
	CLINICAL SIGNS: lethargy, ataxia at both doses	
	BODY WEIGHT: no effects	
Source Test substance	 NECROPSY FINDINGS: no visible lesions Notox Hertogenbosch III, 63734-62-3 (3-[2-chloro-4-(trifluoromethyl)phenoxy]benzoic acid), purity 97%, used as 10% (w/v) dispersion 	
Conclusion Reliability	 LD50 > 500 mg/kg bw (> 50 mg a.i./kg bw) (2) valid with restrictions The information available in the report on the study findings is essentially confined to what is included in the above summary. There is no information on the individual toxicity data. The LD50 is back-calculated to the amount of a.i./kg body weight (dosing was done with a 10% weight/volume dispersion of 0.5% methylcellulose in water) using a density of about 1 g/ml. 	
10.04.2001		(4)

5. Toxicity

ld 63734-62-3 Date 27.12.2001

Type Species Strain Sex Number of animals Vehicle Exposure time Value Method Year GLP Test substance Method	 LC50 rat other: CrI:CD(SD)BR male/female 24 other: none 4 hour(s) > 3.4 mg/l other: not specified yes other TS TEST ORGANISMS: Source: Charles River Breeding Laboraties (Portage, MI) Age: not specified Weight at study initiation: not included in the report Number of animals: 12/sex/dose Controls: yes (12/sex) ADMINISTRATION: Type of exposure: whole body exposure to test substance dust Exposure duration: 4 hours Half of the rats (6m/6f) were killed immediately after exposure, the other half on day 14 post-exposure Type or preparation of particles: with dust generator Air changes: 15/hour EXAMINATIONS: for toxic signs once every hour during exposure, and twice daily during the post-exposure period. Haematology: hemoglobin, hematocrit, red cell count, white cell count, clot time, platelet count, prothrobin time, partial tromboplastin time. Necropsy for macroscopic abnormalities of organs (cervical lymph nodes, salivary glands, throids, trachea, lungs, heart and aorta, thymus, liver, stomach, nasal turbinates, pancreas, spleen, intestines, kidneys, adrenals, bladder, testes/ovaries, uterus and eyes). Those organs which showed abnormalities were examined histopathologically (trachea, lungs and nasal turbinates, pancreas, spleen, intestines, kidneys, adrenals, bladder, testes/ovaries, uterus and eyes). BODY WEIGHT: on days 0 (pre-dosing), 1, 3, 5, 7, 11, and 14 ANALYSES: chamber analytical concentration and particle size distribution Method: gravimetry Sampling times: analytical concentration: no data, PSD: twice (110 and 197 min) Concentrations(nominal/measured): 102.46 mg/l / 3.39 +/-0.56 mg/l (n=13)
Result	 8.5 (+/- 1.8) microns at 110 and 197 minutes into the exposure, resp. STATISTICAL METHOD: PSD by log-probit regression analysis (Hagan, 1980) MORTALITY: Number of deaths at each dose: no deaths in the control 2 deaths in the dose group Time of death: 2 days post-exposure

			CLINICAL SIGNS: during exposure of treated animals: dyspnea, gasping, eye squint, lacrimation, salivation, red exudate around the eyes. post-exposure of treated animals: thriftless appearance, red exudates around the eyes and muzzle, yellow-stained anal-genital area, alopecia around the eyes and muzzle, ptosis, exophthalmus, corneal opacities, lacrimation, nasal discharge, dyspnea, rales, ataxia, decreased motor activity, and prostration.	
			BODY WEIGHT: control animals no body weight losses treated animals: body weight losses on day 1 and 2, followed by body weight gains on day 7 to 11.	
			HEMATOLOGY: reduced white blood cell counts and increased platelet counts.	
Source	ndition	:	NECROPSY FINDINGS: control group: no gross lesions (8M,9F), hardened and/or enlarged salivary glands (4M,1F), hardened and/or enlarged cervical lymph nodes (2M,1F), diffuse brown areas on the lung (1M,1F), and dilated kidney medulla (1M). treated group: decendents: redness of lungs (2F), yellow-stained anal-genital area (2F), and red-stained muzzle (2F); surviving animals (0 and 14 days): no gross lesions (4M,5F), corneal opacities (6M,2F), red-spotted cervical lymph nodes (1F), hardened salivary glands (1F), dilated kidney medulla (1M) and alopecia around the eyes (1F). Histopathology reveals degeneration of the respiratory and olfactory epithelium and congestion of the mucosa of the nasal cavity. Notox Hertogenbosch	
Test co		:	III, 63734-62-3 (3-[2-chloro-4-(trifluoromethyl)phenoxy]benzoic acid), purity 100%	
Conclus Reliabil			LC50 > 3.4 mg/l (3) invalid 1. This report did not contain tables, nor figures. So, no individual data were present. 2. There is a great difference in nominal versus measured concentration of the test substance dust. 3. The study is not reliable because all animals showed a viral infection "Sialodacryoadenitis (SDA)" during the test. The interpretation of in-life observations is complicated by this fact and especially the hematology is obscured. 4. Due to the use of an out-of-date lot of Vacutainer tubes, the determination of the coagulation parameters was prevented	
10.04.20	001		prevented.	(

5.1.3 ACUTE DERMAL TOXICITY

Туре	: LD50
Species	: rabbit
Strain	: other: Albino
Sex	: male
Number of animals	: 5
Vehicle	: water
Value	: > 5000 mg/kg bw
Method	: other: not specified

(10)

Year GLP Test substance Method	: no other TS TEST ORGANISMS: - Source: not indicated - Age: not indicated - Weight at study initiation: 2.23-2.32 kg (mean) - Controls: no	
	ADMINISTRATION: - Area covered: not specified - Occlusion: yes - Vehicle: aqueous paste - Concentration in vehicle: not specified - Doses: 2500 and 5000 mg/kg bw - Removal of test substance: no data - contact time: 24 hours	
	EXAMINATIONS: signs of intoxication, skin irritation and gross autopsy	
	BODY WEIGHT: pre-dosing and at end of the test	
Result	 STATISTICAL METHOD: no data MORTALITY: Number of deaths at each dose: 2500 and 5000 mg/kg bw: 0/5 and 1/5, respectively Time of death: between days 8 and 14 	
	CLINICAL SIGNS: no signs of intoxication, very slight erythema, no edema observed	
	BODY WEIGHT: normal	
Source Test substance	 NECROPSY FINDINGS: normal in both decendents and survivors Notox Hertogenbosch III, 63734-62-3 (3-[2-chloro-4-(trifluoromethyl)phenoxy]benzoic acid), purity 86.5%, aqueous paste 	
Conclusion	: LD50 > 5000 mg/kg bw	
Reliability	 : (4) not assignable 1. The information was essentially confined to what is included in the current summary. No data were present on body area covered, concentration a.i. in the paste. This lowers the reliability of the study. 2. only males are included 	
10.04.2001		(3)
Type Species Strain Sex Number of animals Vehicle Value Method Year	 LD50 rabbit New Zealand white male 6 physiol. saline > 200 mg/kg bw other: not specified 	
GLP Test substance Method	: no : other TS : TEST ORGANISMS: - Source: not indicated Age: not indicated	
	- Age: not indicated 19 / 24	

	00101020
Date	27.12.2001

		- Weight at study initiation: 2.76 kg (mean) - Controls: no	
		ADMINISTRATION: - Area covered: not specified - Occlusion: yes - Vehicle: paste with saline - Concentration in vehicle: not specified - Doses: 200 mg/kg bw - Removal of test substance: no data - contact time: 24 hours	
		EXAMINATIONS: signs of intoxication, skin irritation and gross autopsy	
		BODY WEIGHT: pre-dosing and at end of the test	
Result	:	STATISTICAL METHOD: no data MORTALITY: - Number of deaths at each dose: no deaths	
		CLINICAL SIGNS: no signs of intoxication; no skin irritation observed on the intact skin; well defined erythema and slight edema observed on abraded skin.	
		BODY WEIGHT: normal	
		NECROPSY FINDINGS: no visible lesions; 1 rabbit indentation in surface of kidneys	
Source Test substance		Notox Hertogenbosch III, 63734-62-3	
		(3-[2-chloro-4-(trifluoromethyl)phenoxy]benzoic acid), purity 97%, used as saline paste	
Conclusion	:	LD50 > 200 mg/kg bw	
Reliability	:	 (4) not assignable 1. The information was essentially confined to what is included in the current summary. No data were present on body area covered, concentration a.i. in the paste. This lowers the reliability of the study. 2. Abrasion of the skin can influence the permeability of the test substance. 	
10.04.2001			(4)
5.1.4 ACUTE TOXICITY	ΟΤΗ	ER ROUTES	
	,		
5.2.1 SKIN IRRITATION	ł		

5.2.2 EYE IRRITATION

5.3 SENSITIZATION

5.4 REPEATED DOSE TOXICITY

5.5 GENETIC TOXICITY 'IN VITRO'

 TA1535, TA1537, TA98 and TA100 75-7500 ug/plate with and without negative no data other TS SYSTEM OF TESTING: Species/cell type: Salmonella typhimurium TA98, TA100, TA1535, TA1537. Deficiences/Proficiences: histidine Metabolic activation system: rat S9 mix (Arochlor 1254 induced) 	
ADMINISTRATION: - Dosing: 0, 75, 250, 750, 2500, 7500 μ g/plate - Number of replicates: unknown - Application: DMSO or saline buffer - Positive and negative control groups and treatment: Positive controls: \pm S -9: 2-anthramine for TA1535, TA1537 and TA100, \pm S -9 2-Acetaminofluorene for TA98. Negative controls: DMSO - type of test: no data	
 Notox Hertogenbosch CAS 63734-62-3, (3-(2-chloro-4-trifluoromethylphenoxy)benzoic acid), purity 88.5% 	
 : (4) not assignable 1. The information given in the report was essentially confined to what is included in the current summary. 2. No strain with an AT basepair at the primary reversion site is tested. 	
	(7)
 Ames test TA1535, TA1537, TA98 and TA100 75-7500 ug/plate with and without negative no data other TS SYSTEM OF TESTING: Species/cell type: Salmonella typhimurium TA98, TA100, TA1535, TA1537. Deficiences/Proficiences: histidine Metabolic activation system: rat S9 mix (Arochlor 1254 induced) ADMINISTRATION: Dosing: 0, 75, 250, 750, 2500, 7500µg/plate Number of replicates: unknown Application: DMSO or saline buffer Positive and negative control groups and treatment: 	
	 75-7500 ug/plate with and without negative no data other TS SYSTEM OF TESTING: Species/cell type: Salmonella typhimurium TA98, TA100, TA1535, TA1537. Deficiences/Proficiences: histidine Metabolic activation system: rat S9 mix (Arochlor 1254 induced) ADMINISTRATION: Dosing: 0, 75, 250, 750, 2500, 7500µg/plate Number of replicates: unknown Application: DMSO or saline buffer Positive controls: ±S -9: 2-anthramine for TA1535, TA1537 and TA100, ±S -9: 2-Actaminofluorene for TA98. Negative controls: DMSO type of test: no data Notox Hertogenbosch CAS 63734-62-3, (3-(2-chloro-4-trifluoromethylphenoxy)benzoic acid), purity 88.5% (4) no tassignable The information given in the report was essentially confined to what is included in the current summary. No strain with an AT basepair at the primary reversion site is tested. Ames test TA1535, TA1537, TA98 and TA100 75-7500 ug/plate with and without negative no data other TS SYSTEM OF TESTING: SySTEM OF TESTING: SySTEM OF TESTING: Species/cell type: Salmonella typhimurium TA98, TA100, TA1535, TA1537. Deficiences/Proficiences: histidine Metabolic activation system: rat S9 mix (Arochlor 1254 induced) ADMINISTRATION: Dosing: 0, 75, 250, 750, 2500, 7500µg/plate Number of replicates: unknown Application: DMSO or saline buffer

21 / 24

5. To	oxicity		3734-62-3 7.12.2001
		Positive controls: \pm S-9: 2-anthramine for TA1535, TA1537 TA100, \pm S-9 2-Acetaminofluorene for TA98. Negative controls: DMSO - type of test: no data	′ and
Sou		Notox Hertogenbosch	
Tes	t substance :	CAS 63734-62-3, (3-(2-chloro-4-trifluoromethylphenoxy)benzoic acid), purity 88.5%	,
Rel	iability :	(4) not assignable1. The information given in the report was essentially confined to what is included in the current summary.2. No strain with an AT basepair at the primary reversion site is tested.	
17.(05.2001	3. The report is incomplete.	(6)
5.6	GENETIC TOXICITY 'IN	N VITRO'	
5.7	CARCINOGENITY		
5.8	TOXICITY TO REP ROD	DUCTION	
5.9	DEVELOPMENTAL TO	XICITY/TERATOGENICITY	
		FORMATION	

5.11 EXPERIENCE WITH HUMAN EXPOSURE

- (1) EPIWIN v3.05, Syracuse Research Corporation, Syracuse, NY (July 12, 2000)
- (2) Lyman, W. J. et al. (1990). Handbook of Chemical PropertyEstimation Methods, pp. 7-4, Amer. Chem. Society,Washington, DC
- Rohm & Haas Co, Acute toxicity studies with 3-(2-chloro-4-(trifluoromethyl)phenoxy)benzoic acid in rats and rabbits, 1976 (48)
- Rohm & Haas Co, Acute toxicity studies with 3-(2-chloro-4-(trifluoromethyl)phenoxy)benzoic acid in rats and rabbits, 1978 (49)
- (5) Rohm and Haas Company, Acute toxicity of RH-41,833 to fathead minnow (Pimephales promelas), 1976 (47)
- (6) Rohm and Haas Company, RH-41, 833 microbial mutagen test (final report) with cover letter dated 06.09.93
- (7) Rohm and Haas Company, RH-41, 833 microbial mutagen test (final report) with cover letter dated 07.17.84
- (8) Rohm and Haas Company, The acute toxicity of TD-77-370 to Bluegill sunfish, 1978 (52)
- Rohm and Haas Company, The acute toxicity of TD-77-373 to the Bluegill sunfish Lepomis macrochirus Rafinesque, 1978 (50)
- (10) Rohm and Haas Company, Toxicology Department, Acute Inhalation Toxicity Study in Rats, 1985 (46)

7.1 END POINT SUMMARY

- 7.2 HAZARD SUMMARY
- 7.3 RISK ASSESSMENT

IUCLID

Data Set

Existing Chemical CAS No. Generic name	 ID: 72252-48-3 72252-48-3 Benzoic acid, 3-[2-chloro-4-(trifluoromethyl)phenoxy], potassium salt
Producer Related Part Company Creation date	Toxicology and Regulatory Affairs27.12.2001
Substance Related Part Company Creation date	: Toxicology and Regulatory Affairs : 27.12.2001
Memo	:
Printing date Revision date Date of last Update	: 27.12.2001 : : 27.12.2001
Number of Pages	: 13
Chapter (profile) Reliability (profile) Flags (profile)	 Chapter: 1, 2, 3, 4, 5, 7 Reliability: without reliability, 1, 2, 3, 4 Flags: without flag, confidential, non confidential, WGK (DE), TA -Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. Ge	eneral Information	72252-48-3 27.12.2001
1.0.1	OECD AND COMPANY INFORMATION	
1.0.2	LOCATION OF PRODUCTION SITE	
1.0.3	IDENTITY OF RECIPIENTS	
1.1	GENERAL SUBSTANCE INFORMATION	
1.1.0	DETAILS ON TEMPLATE	
1.1.1	SPECTRA	
1.2	SYNONYMS	
1.3	IMPURITIES	
1.4	ADDITIVES	
1.5	QUANTITY	
1.6.1	LABELLING	
1.6.2	CLASSIFICATION	
1.7	USE PATTERN	
1.7.1	TECHNOLOGY PRODUCTION/USE	
1.8	OCCUPATIONAL EXPOSURE LIMIT VALUES	
1.9	SOURCE OF EXPOSURE	
1.10.1	RECOMMENDATIONS/PRECAUTIONARY MEASURES	

1. General Information

ld 72252-48-3 Date 27.12.2001

1.10.2 EMERGENCY MEASURES

- 1.11 PACKAGING
- 1.12 POSSIB. OF RENDERING SUBST. HARMLESS
- 1.13 STATEMENTS CONCERNING WASTE
- 1.14.1 WATER POLLUTION
- 1.14.2 MAJOR ACCIDENT HAZARDS
- 1.14.3 AIR POLLUTION
- 1.15 ADDITIONAL REMARKS
- 1.16 LAST LITERATURE SEARCH

1.17 REVIEWS

1.18 LISTINGS E.G. CHEMICAL INVENTORIES

2.1 MELTING POINT

Value Sublimation Method Year GLP Test substance Method Result	 ca. 251 ° C 2001 no Estimation using MPBPWIN v1.40 in EPIWIN 3.05 MPBPWIN (v1.40) Program Results:
	 Experimental Database Structure Match: no data SMILES : c1(CL)cc(C(F)(F)(F))ccc1Oc2cccc(C(=O)OK)c2 CHEM : Potassium Trifluorobenzoic acid CAS 72252-48-3 MOL FOR: C14 H7 CL1 F3 O3 K1 MOL WT : 354.76 SUMMARY MPBPWIN v1.40 Boiling Point: 583.20 deg C (Adapted Stein and Brown Method) Melting Point: 349.84 deg C (Adapted Jobac k Method) Melting Point: 226.87 deg C (Gold and Ogle Method) Mean Melt Pt : 288.36 deg C (Joback; Gold,Ogle Methods) Selected MP: 251.47 deg C (Weighted Value)
Test substance Reliability Flag 27.12.2001	 Potassium salt of benzoic acid, 3-[2-chloro-4-(trifluoromethyl)phenoxy CAS 72252-48-3 (2) valid with restrictions Critical study for SIDS endpoint (1)

- 2.2 BOILING POINT
- 2.3 DENSITY

2.3.1 GRANULOMETRY

2.4 VAPOUR PRESSURE

Value Decomposition	< .0000000	0001 hPa at ° C
Method		
Year	2001	
GLP	no	
Test substance		
Method	Estimation	n using MPBPWIN v1.40 in EPIWIN 3.05
Result		-
	MPBPWIN	N (v1.40) Program Results:

	Experimental Database Structure Match: no data
	SMILES : c1(CL)cc(C(F)(F)(F))ccc1Oc2cccc(C(=O)OK)c2 CHEM : Potassium Trifluorobenzoic acid CAS 72252-48-3 MOL FOR: C14 H7 CL1 F3 O3 K1 MOL WT : 354.76
	- SUMMARY MPBPWIN v1.40
	Vapor Pressure Estimations (25 deg C): (Using BP: 583.20 deg C (estimated)) (Using MP: 251.47 deg C (estimated)) VP: 2.57E-016 mm Hg (Antoine Method) VP: 6.93E-013 mm Hg (Modified Grain Method) VP: 2.46E-012 mm Hg (Mackay Method)
	Selected VP: 6.93E-013 mm Hg (Modified Grain Method)
Test substance	: Potassium salt of benzoic acid, 3-[2-chloro-4-(trifluoromethyl)phenoxy CAS 72252-48-3
Reliability	: (2) valid with restrictions
Flag 27.12.2001	: Critical study for SIDS endpoint (1)

2.5 PARTITION COEFFICIENT

Log pow	: ca56 at ° C
Method	
Year	: 2001
GLP	: no
Test substance	:
Method	: Estimation using KOWWIN v1.66 in EPIWIN 3.05
Test substance	: Potassium salt of benzoic acid, 3-[2-chloro-4-(trifluoromethyl)phenoxy CAS 72252-48-3
Reliability	: (2) valid with restrictions
Flag	: Critical study for SIDS endpoint
27.12.2001	(1)

2.6.1 WATER SOLUBILITY

Value Qualitative Pka PH Method Year GLP Test substance Method	: ca. 1900 mg/l at 25 ° C : at 25 ° C : at and ° C : 2001 : no : Estimation using WSKOW(v1.40 in EBIW(N 2.05
Result	 Estimation using WSKOW v1.40 in EPIWIN 3.05 Water Sol from Kow (WSKOW v1.40) Results: ====================================

SMILES : c1(CL)cc(C(F)(F)(F))ccc1Oc2cccc(C(=O)OK)c2 CHEM : Potassium Trifluorobenzoic acid CAS 72252-48-3

		MOL FOR: C14 H7 CL1 F3 O3 K1 MOL WT : 354.76	
		- WSKOW v1.40 Results	
		Log Kow (estimated) : 0.56 Log Kow (experimental): not available from database Log Kow used by Water solubility estimates: 0.56	
		Equation Used to Make Water Sol estimate: Log S (mol/L) = 0.796 - 0.854 log Kow - 0.00728 MW + Correction (used when Melting Point NOT available)	
		Correction(s): Value	
		No Applicable Correction Factors	
		Log Water Solubility (in moles/L) : -2.261 Water Solubility at 25 deg C (mg/L): 1946	
Tes	t substance :	Potassium salt of benzoic acid, 3-[2-chloro-4-(trifluoromethyl)phenoxy C. 72252-48-3	AS
Reli Flag	-	(2) valid with restrictions Critical study for SIDS endpoint	
	2.2001		(1)
			(1)
27.1	2.2001		(1)
27.1 2.6.2	SURFACE TENSION		(1)
27.1 2.6.2	SURFACE TENSION		(1)
27.4 2.6.2 2.7 2.8	2.2001 SURFACE TENSION FLASH POINT AUTO FLAMMABILIT		(1)
27.1 2.6.2 2.7	SURFACE TENSION		(1)
27.4 2.6.2 2.7 2.8	2.2001 SURFACE TENSION FLASH POINT AUTO FLAMMABILITY FLAMMABILITY	· · · · · · · · · · · · · · · · · · ·	(1)
27.4 2.6.2 2.7 2.8	2.2001 SURFACE TENSION FLASH POINT AUTO FLAMMABILIT	· · · · · · · · · · · · · · · · · · ·	(1)
27.4 2.6.2 2.7 2.8 2.9	2.2001 SURFACE TENSION FLASH POINT AUTO FLAMMABILITY FLAMMABILITY	Υ	(1)

2.12 ADDITIONAL REMARKS

3.1.1 PHOTODEGRADATION

Type Light source Light spect. Rel. intensity Indirect photolysis Sensitizer Conc. of sens. Rate constant Degradation Deg. Product Method Year GLP Test substance Method Remark	 air nm based on Intensity of Sunlight OH 1500000 cm3/(molecule*sec) % after 2001 Estimation using APOWIN v1.90 in EPIWIN 3.05 Due to the low volatility, this reaction unlikely in practice.
Result	 AOP Program (v1.90) Results: SMILES : c1(CL)cc(C(F)(F)(F))ccc1Oc2cccc(C(=O)OK)c2 CHEM : Potassium Trifluorobenzoic acid CAS 72252-48-3 MOL FOR: C14 H7 CL1 F3 O3 K1 MOL WT : 354.76 SUMMARY (AOP v1.90): HYDROXYL RADICALS
Test substance Reliability	 Potassium salt of benzoic acid, 3-[2-chloro-4-(trifluoromethyl)phenoxy CAS 72252-48-3 (2) valid with restrictions Critical study for SIDS and point
Flag 27.12.2001	: Critical study for SIDS endpoint (1)

3.1.2 STABILITY IN WATER

: : : : : : : : : : : : : : : : : : : :	abiotic > 1 year at 25 degree C > 1 year at 25 degree C > 1 year at 25 degree C 2001
:	no
	: : : : :

ld 72252-48-3 Date 27.12.2001

Test substance Method	: Estimated on chemical principles based on absence of groups susceptible to hydrolysis
Remark	: The estimation program in EPIWIN has no capability to estimate hydrolysis rates for this compound.
Result	: This material has no groups that are susceptible to hydrolysis in the pH 4 to 9 range; therefore, it is considered stable to hydrolysis in surface and groundwater. It is estimated to have a hydrolysis half-life of greater than one year between pH 4 and pH 9.

Test substance	:	Potassium salt of benzoic acid, 3-[2-chloro-4-(trifluoromethyl)phenoxy 72252-48-3	CAS
Reliability Flag 27.12.2001		(2) valid with restrictions Critical study for SIDS endpoint	(2)

3.1.3 STABILITY IN SOIL

3.2 MONITORING DATA

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

Type Media Air (level I) Water (level I) Soil (level I) Biota (level II / III) Soil (level II / III) Method Year	fugacity model level III 2 2 2 2001
Method	: The Fugacity was determined using the EQC Level III model as found in EPIWIN 3.05. Estimated values were used for physical constants. Biodegradation was based on the EPIWIN derived estimates (Biowin, Ultimate) that were assessed for reasonableness compared with similar compounds. Half life in air was determined from the APOWIN program. Direct photolysis was not considered in this model. Emissions were calculated from only water and soil as this test substance it is non-volatile. Other parameters used the default values found in EPIWIN.
Result	<pre>: r Level III Fugacity Model (Full-Output): </pre>

ld 72252-48-3

Date 27.12.2001

		Air Water Soil Sediment	Fugaci ty (atm) 2.55e-029 3.53e-021 8.27e-020 3.44e-021	Reaction (kg/hr) 1.39e-012 290 205 0.146	Advecti on (kg/hr) 2.76e-012 1.5e+003 0 0.0607	Reaction (percent) 6.94e-014 14.5 10.3 0.00731	Advection (percent) 1.38e-013 75.2 0 0.00304
		Reacti Advect Percer	stence Time: ion Time: tion Time: nt Reacted: nt Advected:				
		Ain Wat Soi See	r: 138 ter: 3600 il: 3600 diment: 1.44)			owi n) :
		Ai Wat	tion Times (r: 100 ter: 100(diment: 5e+()			
Tes	t substance :	Potassiu		nzoic acid, 3-[2	2-chloro-4-(tr	fluoromethyl)phenoxy CAS
Rel	iability :	. ,	with restriction				
Fla g 27.2	g : 12.2001	Critical s	tudy for SID	S endpoint			(1)
3.3.2	DISTRIBUTION						
3.4	MODE OF DEGRADA		TUAL USE				
3.5	BIODEGRADATION						
3.6	BOD5, COD OR BOD5	/COD RAT	10				
3.7	BIOACCUMULATION						
3.8	ADDITIONAL REMARI	KS					

- 4.1 ACUTE/PROLONGED TOXICITY TO FISH
- 4.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES
- 4.3 TOXICITY TO AQUATIC PLANTS E.G. ALGAE
- 4.4 TOXICITY TO MIC ROORGANISMS E.G. BACTERIA
- 4.5.1 CHRONIC TOXICITY TO FISH
- 4.5.2 CHRONIC TOXICITY TO AQUATIC INVERTEBRATES
- 4.6.1 TOXICITY TO SOIL DWELLING ORGANISMS
- 4.6.2 TOXICITY TO TERRESTRIAL PLANTS
- 4.6.3 TOXICITY TO OTHER NON-MAMM. TERRES TRIAL SPECIES
- 4.7 BIOLOGICAL EFFECTS MONITORING
- 4.8 BIOTRANSFORMATION AND KINETICS
- 4.9 ADDITIONAL REMARKS

 5.1.1 ACUTE ORAL TOXICITY 5.1.2 ACUTE INHALATION TOXICITY 5.1.3 ACUTE DERMAL TOXICITY 5.1.4 ACUTE TOXICITY, OTHER ROUTES 5.2.1 SKIN IRRITATION 5.2.2 EYE IRRITATION 5.3 SENSITIZATION 5.4 REPEATED DOSE TOXICITY 5.5 GENETIC TOXICITY 'IN VITRO' 5.6 GENETIC TOXICITY 'IN VITRO' 5.7 CARCINOGENITY 5.8 TOXICITY TO REP RODUCTION 5.9 DEVELOPMENTAL TOXICITY/TERATOGENICITY 5.10 OTHER RELEVANT INFORMATION 5.11 EXPERIENCE WITH HUMAN EXPOSURE 		
 5.1.3 ACUTE DERMAL TOXICITY 5.1.4 ACUTE TOXICITY, OTHER ROUTES 5.2.1 SKIN IRRITATION 5.2.2 EYE IRRITATION 5.3 SENSITIZATION 5.4 REPEATED DOSE TOXICITY 5.5 GENETIC TOXICITY 'IN VITRO' 5.6 GENETIC TOXICITY 'IN VITRO' 5.7 CARCINOGENITY 5.8 TOXICITY TO REP RODUCTION 5.9 DEVELOPMENTAL TOXICITY/TERATOGENICITY 5.10 OTHER RELEVANT INFORMATION 	5.1.1	ACUTE ORAL TOXICITY
 5.1.4 ACUTE TOXICITY, OTHER ROUTES 5.2.1 SKIN IRRITATION 5.2.2 EYE IRRITATION 5.3 SENSITIZATION 5.4 REPEATED DOSE TOXICITY 5.5 GENETIC TOXICITY 'IN VITRO' 5.6 GENETIC TOXICITY 'IN VITRO' 5.7 CARCINOGENITY 5.8 TOXICITY TO REP RODUCTION 5.9 DEVELOPMENTAL TOXICITY/TERATOGENICITY 5.10 OTHER RELEVANT INFORMATION 	5.1.2	ACUTE INHALATION TOXICITY
 5.2.1 SKIN IRRITATION 5.2.2 EYE IRRITATION 5.3 SENSITIZATION 5.4 REPEATED DOSE TOXICITY 5.5 GENETIC TOXICITY 'IN VITRO' 5.6 GENETIC TOXICITY 'IN VITRO' 5.7 CARCINOGENITY 5.8 TOXICITY TO REP RODUCTION 5.9 DEVELOPMENTAL TOXICITY/TERATOGENICITY 5.10 OTHER RELEVANT INFORMATION 	5.1.3	ACUTE DERMAL TOXICITY
 5.2.2 EYE IRRITATION 5.3 SENSITIZATION 5.4 REPEATED DOSE TOXICITY 5.5 GENETIC TOXICITY 'IN VITRO' 5.6 GENETIC TOXICITY 'IN VITRO' 5.7 CARCINOGENITY 5.8 TOXICITY TO REP RODUCTION 5.9 DEVELOPMENTAL TOXICITY/TERATOGENICITY 5.10 OTHER RELEVANT INFORMATION 	5.1.4	ACUTE TOXICITY, OTHER ROUTES
 5.3 SENSITIZATION 5.4 REPEATED DOSE TOXICITY 5.5 GENETIC TOXICITY 'IN VITRO' 5.6 GENETIC TOXICITY 'IN VITRO' 5.7 CARCINOGENITY 5.8 TOXICITY TO REP RODUCTION 5.9 DEVELOPMENTAL TOXICITY/TERATOGENICITY 5.10 OTHER RELEVANT INFORMATION 	5.2.1	SKIN IRRITATION
 5.4 REPEATED DOSE TOXICITY 5.5 GENETIC TOXICITY 'IN VITRO' 5.6 GENETIC TOXICITY 'IN VITRO' 5.7 CARCINOGENITY 5.8 TOXICITY TO REP RODUCTION 5.9 DEVELOPMENTAL TOXICITY/TERATOGENICITY 5.10 OTHER RELEVANT INFORMATION 	5.2.2	EYE IRRITATION
 5.5 GENETIC TOXICITY 'IN VITRO' 5.6 GENETIC TOXICITY 'IN VITRO' 5.7 CARCINOGENITY 5.8 TOXICITY TO REP RODUCTION 5.9 DEVELOPMENTAL TOXICITY/TERATOGENICITY 5.10 OTHER RELEVANT INFORMATION 	5.3	SENSITIZATION
 5.6 GENETIC TOXICITY 'IN VITRO' 5.7 CARCINOGENITY 5.8 TOXICITY TO REP RODUCTION 5.9 DEVELOPMENTAL TOXICITY/TERATOGENICITY 5.10 OTHER RELEVANT INFORMATION 	5.4	REPEATED DOSE TOXICITY
 5.7 CARCINOGENITY 5.8 TOXICITY TO REP RODUCTION 5.9 DEVELOPMENTAL TOXICITY/TERATOGENICITY 5.10 OTHER RELEVANT INFORMATION 	5.5	GENETIC TOXICITY 'IN VITRO'
 5.8 TOXICITY TO REP RODUCTION 5.9 DEVELOPMENTAL TOXICITY/TERATOGENICITY 5.10 OTHER RELEVANT INFORMATION 	5.6	GENETIC TOXICITY 'IN VITRO'
5.9 DEVELOPMENTAL TOXICITY/TERATOGENICITY 5.10 OTHER RELEVANT INFORMATION	5.7	CARCINOGENITY
5.10 OTHER RELEVANT INFORMATION	5.8	TOXICITY TO REP RODUCTION
	5.9	DEVELOPMENTAL TOXICITY/TERATOGENICITY
5.11 EXPERIENCE WITH HUMAN EXPOSURE	5.10	OTHER RELEVANT INFORMATION
	5.11	EXPERIENCE WITH HUMAN EXPOSURE

- (1) EPIWIN v3.05, Syracuse Research Corporation, Syracuse, NY (July 12, 2000)
- (2) Lyman, W. J. et al. (1990). Handbook of Chemical PropertyEstimation Methods, pp. 7-4, Amer. Chem. Society,Washington, DC

7.1 END POINT SUMMARY

- 7.2 HAZARD SUMMARY
- 7.3 RISK ASSESSMENT

IUCLID

Data Set

Existing Chemical CAS No. Generic name	:	ID: 62476-59-9 62476-59-9 Sodium 5-(2-chloro-4-trifluoro-methylphenoxy) 2-nitrobenzoate
Producer Related Part Company Creation date		Toxicology and Regulatory Affairs 26.12.2001
Substance Related Part Company Creation date		Toxicology and Regulatory Affairs 26.12.2001
Memo	:	
Printing date Revision date Date of last Update	:	26.12.2001 26.12.2001
Number of Pages	:	44
Chapter (profile) Reliability (profile) Flags (profile)	:	Chapter: 1, 2, 3, 4, 5, 7 Reliability: without reliability, 1, 2, 3, 4 Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. 0	General Information	62476-59-9 26.12.2001
1.0.1	OECD AND COMPANY INFORMATION	
1.0.2	LOCATION OF PRODUCTION SITE	
1.0.3	IDENTITY OF RECIPIENTS	
1.1	GENERAL SUBSTANCE INFORMATION	
1.1.0	DETAILS ON TEMPLATE	
1.1.1	SPECTRA	
1.2	SYNONYMS	
1.3	IMPURITIES	
1.4	ADDITIVES	
1.5	QUANTITY	
1.6.1	LABELLING	
1.6.2	CLASSIFICATION	
1.7	USE PATTERN	
1.7.1	TECHNOLOGY PRODUCTION/USE	
1.8	OCCUPATIONAL EXPOSURE LIMIT VALUES	
1.9	SOURCE OF EXPOSURE	

1. General Information	62476-59-9 26.12.2001
1.10.1 RECOMMENDATIONS/PRECAUTIONARY MEASURES	
1.10.2 EMERGENCY MEASURES	
1.11 PACKAGING	
1.12 POSSIB. OF RENDERING SUBST. HARMLESS	
1.13 STATEMENTS CONCERNING WASTE	
1.14.1 WATER POLLUTION	
1.14.2 MAJOR ACCIDENT HAZARDS	
1.14.3 AIR POLLUTION	
1.15 ADDITIONAL REMARKS	
1.16 LAST LITERATURE SEARCH	
1.17 REVIEWS	
1.18 LISTINGS E.G. CHEMICAL INVENTORIES	

2.1 MELTING POINT

Value Decomposition Sublimation Method Year GLP Test substance Method		172 °C yes at ca. 240 °C OECD Guide-line 102 "Melting Point/Melting Range" 1981 no other TS capillary method/metal block apparatus	
Result	:	determination 1 beginning of melting (shrink point) (deg C)determination 2 172collapse point (deg C)178collapse point (deg C)178No other melt transitions were noted. Samples were heated to 240 deg C when sample degradation was noted by discoloration and offgassing	
Source	:	Notox Hertogenbosch	
Test condition	:	Duplicate dried powder samples were charged into a capillary column (resulting height about 2 mm). Samples were initially heated in the melting point apparatus at about 5 deg C/min, and at about 1 deg C/min within 10 deg C of the transition. Method was validated using a reference substance of known melting point (sulfanilamide).	
Test substance	:	III, CAS 62476-59-9 (acifluorfen-sodium, purified technical), purity 89.3%	
Conclusion Reliability Flag 26.12.2001	:	Melting starts at 172 deg C. Melting is not complete; test substance decomposes at about 240 deg C. (1) valid without restriction Critical study for SIDS endpoint	(18)
Value Decomposition Sublimation Method Year GLP Test substance Method		176 ° C yes at ca. 240 ° C OECD Guide-line 102 "Melting Point/Melting Range" 1981 no other TS capillary method/metal block apparatus	
Result	:	determination 1 determination 2	
		beginning of melting 176 176 (shrink point) (deg C)	
		No other melt transitions were noted. Samples were heated to 240 deg C when sample degradation was noted by discoloration and	
		4 / 44	

2. Physico-Che	mical Data	ld 62476-59-9 Date 26.12.2001	
	offgassing		
Source Test condition	column (resulting height abo heated in the melting point a and at about 1 deg C/min w	ples were charged into a capillary out 2 mm). Samples were initially apparatus at about 5 deg C/min, ithin 10 deg C of the transition. a reference substance of known	
Test substance	: III, CAS 62476-59-9 (acifluo 74.4%	rfen-sodium, technical), purity	
Conclusion	substance decomposes at a	Melting is not complete; test bout 240 deg C.	
Reliability	(1) valid without restrictionCritical study for SIDS endp	oint	
Flag 26.12.2001	. Chica study for SIDS enup		18)
		·	
2.2 BOILING POINT			
2.3 DENSITY			
2.3.1 GRANULOMETI	२४		
2.4 VAPOUR PRES	SUDE		
2.4 VAFOUR FRES	JURE		
Value	: < .000000133 hPa at 25° (2	
Decomposition	: no	,	
Method		y OECD 104 (gas saturation method)	
Year	: 1981		
GLP			
-	: yes		
Test substance	: other TS		
Decomposition	: no		
Result	: In all cases, acifluorfen sodi		
		re was < 1.33E-5 Pa, which is	
0	the lower limit of detection.		
Source	Material Instance I I		
	: Notox Hertogenbosch		
Test condition	: Vapor pressure was measu using 8 or 9 flow rates in the	red at 25, 35 and 45 +/- 0.5 deg C e range 7-140 cc/min. At 25 and	
	 Vapor pressure was measu using 8 or 9 flow rates in the 45 deg C two experiments v sodium was packed into 5 n wool plugs (sample length 6 	e range 7-140 cc/min. At 25 and vere performed. Hereto, acifluorfen nm glass tubing between 2 glass 0 mm) and connected to 2 XAD-2	
	 Vapor pressure was measu using 8 or 9 flow rates in the 45 deg C two experiments v sodium was packed into 5 n wool plugs (sample length 6 sorbent sections separated mm). The system was place 	e range 7-140 cc/min. At 25 and vere performed. Hereto, acifluorfen nm glass tubing between 2 glass 0 mm) and connected to 2 XAD-2 by glass wool (about 15 and 10 ed in a constant temperature box and	
	 Vapor pressure was measu using 8 or 9 flow rates in the 45 deg C two experiments v sodium was packed into 5 n wool plugs (sample length 6 sorbent sections separated mm). The system was place nitrogen gas was passed th the sorbent traps were extra water (shaking for 2 hrs). Th 	e range 7-140 cc/min. At 25 and vere performed. Hereto, acifluorfen nm glass tubing between 2 glass 0 mm) and connected to 2 XAD-2 by glass wool (about 15 and 10 ed in a constant temperature box and rough it. After at least 473 hrs, incted with 2 mL methanol and 1 mL ne extracts were analyzed by	
	 Vapor pressure was measu using 8 or 9 flow rates in the 45 deg C two experiments v sodium was packed into 5 n wool plugs (sample length 6 sorbent sections separated mm). The system was place nitrogen gas was passed th the sorbent traps were extra water (shaking for 2 hrs). Th HPLC; quantitation was per 	e range 7-140 cc/min. At 25 and vere performed. Hereto, acifluorfen m glass tubing between 2 glass 0 mm) and connected to 2 XAD-2 by glass wool (about 15 and 10 ed in a constant temperature box and rough it. After at least 473 hrs, locted with 2 mL methanol and 1 mL	
	: Vapor pressure was measu using 8 or 9 flow rates in the 45 deg C two experiments v sodium was packed into 5 n wool plugs (sample length 6 sorbent sections separated mm). The system was place nitrogen gas was passed th the sorbent traps were extra water (shaking for 2 hrs). Th HPLC; quantitation was per acifluorfen sodium (prepare	e range 7-140 cc/min. At 25 and vere performed. Hereto, acifluorfen nm glass tubing between 2 glass 0 mm) and connected to 2 XAD-2 by glass wool (about 15 and 10 ed in a constant temperature box and rough it. After at least 473 hrs, ucted with 2 mL methanol and 1 mL ne extracts were analyzed by formed using standard solutions of	

2. Physico-Che	mical Data	ld 62476 Date 26.12.	
	Blank sample t	ubes were included for each temperature.	
Test substance Conclusion Reliability	VP < 1.33E-5 F (2) valid with re 1. For all blank (or a contamina Therefore, the with 5 blanks (3 tubes), but blar only one of the exceed the app	estrictions sample tubes TS appeared to be recovered ant with an identical retention time). experiment was repeated at 25 and 45 deg C 3 tubes containing glass wool, 2 empty glass iks contained TS again (or contaminant). In 39 sample tubes did the compound detected parent concentrations found in the blanks.	
Flag 26.12.2001	: Critical study for	or SIDS endpoint	(4)
5 PARTITION CO	EFFICIENI		
Log pow Method Year GLP Test substance	: at 25° C other (measure : 1995 : yes : other TS	ed): essentially OECD 107	
Method		of acifluorfen sodium in octanol/aqueous	
	buffer at a ratio were prepared. (CAS 50594-66 octanol) and so octanol-saturat addition of octa levels (appr. 8 were prepared for pH 7, high o The samples w centrifugated, a with mobile pha	 of approximately 1:1 (v/v) (pH 5, 7 and 9) Hereto, equimolar amounts of acifluorfen acid 6-6, purity 99.4%, dissolved in buffer-saturated bodium hydroxide (dissolved in ed buffer) were mixed, followed by the nol. Triplicate samples of two concentration mM and 0.8 mM in the original octanol phase) for each pH. Total volume was 0.02 L, except concentration level (total volume 0.05 L). rere shaken at 25 +/- 1 deg C for 16 hours, and each octanol and water phase was diluted ase and analyzed by liquid chromatography n acid (purity 99.5%) as a reference 	
Result	: Buffer pH Initia concenti in n-octa		
	5 8	15.6 +/- 0.17	
	7 8 9 8	1.88 +/- 0.04 1.46 +/- 0.05	
	5 0.8	15.6 +/- 0.81	
	7 0.8 9 0.8	1.21 +/- 0.06 1.12 +/- 0.03	
		s no concentration dependence of Kow.	
Source	: Notox Hertoge	·	
Test substance	: III, CAS 62476-	-59-9 (acifluorfen sodium), purity 99.4% as nversion to sodium salt.	

2. Physico-Che	nical Data	ld 62476-59-9 Date 26.12.2001
Conclusion	: Kow* log Kow* pH 5 15.6 1.19 pH 7 < 2 < 0.3 pH 9 < 1.5 < 0.2 *(mean of two concentrations	s)
Reliability	 (2) valid with restrictions Remarks: TS is in the ionized form, which may caus from the partition law. Method is not suitable substances. OECD 107 advises adjustment below or above the pK, but in this case this i applicable as TS is a salt and should therefor protonated. 	e for ionized of pH to 1 unit is not
	Test was performed at only one water:oct each pH and TS concentration.	anol ratio for
Flag 26.12.2001	: Critical study for SIDS endpoint	(6)
.6.1 WATER SOLUB	ITY	
Value Qualitative Pka PH Method Year GLP Test substance	 405 other: mg/g at 25 ° C moderately soluble (100-1000 mg/L) at 25 ° C at and ° C other: essentially OECD 105 1981 yes other TS 	
Method	 Six centrifuge tubes with test mixture (approx TS/10 mL in HPLC grade water) and two bla interference in the analysis) were shaken in 35 +/- 1 deg C for about 4 hrs, followed by tr +/- 1 deg C water bath (continueous shaking 7 days aliquots were removed after centrifug 31,300 x G or 41,300 x G (3 replicates and 1 30 min. at 25 +/- 1 deg C. About 0.5 mL was by a factor 1000 and analyzed by LC (duplic Standard solutions in the range 0.370-0.685 included for quantification, as well as a refer acifluorfen acid control solution to check received 	anks (to check for a water bath of ransfer to a 25 g). After 3, 6 and gation at appr. 1 blank each) for s weighed, diluted cate injection). 5 mg/mL were rence
Result	: Day Acifluorfen sodium (mg/g) at centrifuge 31,300xG* 41,300xG* Mean	speed:
	3 411.6 405.7 409 +/- 6 6 404.3 407.4 406 +/- 5 7 396.0 407.2 402 +/- 8	
	* mean of three replicates, calculated by rev summarize data	iewer to

	o-Chemical Data	ld 62476-59-9 Date 26.12.2001
	Overall mean: 405 +/- 6.3 mg/ Statistical analysis indicated n difference between days 3, 6 a been established.	o statistically significant
Source Test substand Conclusion Reliability	 Notox Hertogenbosch III, CAS 62476-59-9 (acifluorfe Water solubility of acifluorfen s (1) valid without restriction minor remark: Purity of the test substance may influence the solubility of information on the identity of the 	sodium = 405 +/- 6.3 mg/g. was only 78.2%. Impurities acifluorfen sodium. No
Flag 14.05.2001	substance was given. Critical study for SIDS endpoir	
2.6.2 SURFAC	ETENSION	
2.7 FLASH P	OINT	
	AMMABILITY	
2.8 AUTO FL		
2.8 AUTO FL 2.9 FLAMMA	\BILITY	
2.9 FLAMMA	ABILITY IVE PROPERTIES	
2.9 FLAMMA 2.10 EXPLOSI	IVE PROPERTIES	
2.9 FLAMMA 2.10 EXPLOSI		
2.9 FLAMMA 2.10 EXPLOSI 2.11 OXIDIZIN	IVE PROPERTIES	

3. Environmental Fate and Pathways

3.1.1 PHOTODEGRADATION

Туре	: water
Light source	: Sun light
Light spect.	: nm
Rel. intensity	: based on Intensity of Sunlight
Remark	: Indirect photolysis is not considered as this material is not volatile. Several studies are reported in the EPA RED documentation. It is apparent that this material undergoes primary photodegradation; however, the exact rate and spectrum of degradation products is not fully understood.
Result	: Half life values ranged from 21 hours to 352 hours depending on concentrations and conditions. Near neutrality a mid estimate is 90 hours.
Reliability	: (2) valid with restrictions
Flag	: Critical study for SIDS endpoint
26.12.2001	(9)

3.1.2 STABILITY IN WATER

Type t1/2 pH4 t1/2 pH7 t1/2 pH9 Degradation Deg. Product Method Year GLP Test substance	 abiotic at degree C at degree C at degree C 0 % after 28 day at pH and degree C other: essentially OECD 111 1981 no other TS 	
Method	 Test solutions (1.0 ppm and 50.0 ppm TS; buffered to pH 4.5, 7.2 and 9.7) were incubated at 25 deg C in complete darkness for 28 (1.0 ppm samples) and 56 days (50.0 ppm samples). No cosolvent was used. Samples were taken on day 0,1,3,7,14 ar 28 (1.0 ppm samples) and on day 0,1,3,7,14,28 and 56 (50 pp samples). 0.1 N H3PO4 was added to samples (conversion of sodium acifluorfen to free acid) followed by extraction with benzene. Both aqeous and benzene fractions were analyzed b LSC, benzene fractions were also subjected to TLC. 	o nd m
Result	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	

3. Environment	al Fate and Pathways	ld 62476-59-9 Date 26.12.2001
	28 1 1.09 1.12 1.12	
Source Test substance Conclusion Reliability	 Mass balances were in the range 84.5-98 points, except at for samples and time points for these, mass balances were < 17%, will low extraction efficiencies. Extraction efficiencies. Extraction efficiencies by addition of 1 mL 0.1 N H3PO with benzene from day 7 onwards. Notox Hertogenbosch III, CAS 62476-59-9 (sodium acifluorfen), purity 99%, specific activity 4706 dpm/ug Test substance is stable in water. (2) valid with restrictions 	ints marked with *. hich is explained by ciency was 04 before extraction
Flag	 Volatiles were not measured (no traps), be of no concern because of high mass be an increase with time of TS concentration which is explained by evaporation of solve are only quantified for day 7 (no reference exact mass balance can therefore not be 2. The report consisted of a summary rath report. In this summary, only testing at 25 described, whereas results for 2 other tem 48 deg C) are also given. Results for the of temperatures support the conclusion of th 3. Sterility was not measured, nor was the buffers included in the study. However, as degradation was observed, biotic degrada excluded. Critical study for SIDS endpoint 	alance. In addition, was observed, ent. TLC results e standard). An calculated. her than a full deg C is hperatures (36 and other 2 e test at 25 deg C. e sterility of the s hardly any
10.05.2001		

3.1.3 STABILITY IN SOIL

3.2 MONITORING DATA

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

Type Media Air (level I) Water (level I) Soil (level I) Biota (level II / III) Soil (level II / III) Method Year		fugacity model level III 2001
Method	:	The Fugacity was determined using the EQC Level III model as found in EPIWIN 3.05. Measured values were used for most physical constants. Biodegradation was based on information in the EPA Reregistration Documentation and data in HSDB. The aquatic soil and sediment estimates are estimates of an average half life from biodegradation and
		10 / 44

photolysis. As sediment distribution was low the half life estimate for wat was used in the model. Half life in air was set at a default rapid loss sincuts is not volatile. Emissions were calculated from using only water and soil as this test substance it is not volatile. Chief and the default values found in EPIWIN. Result : Level III Fugacity Model (Full-Output): Chem Name : Sodium Acifluorfen Note in the intervent of the intervent		al Fate and Pathways Id 62476-59-9 Date 26.12.2001
Level III Fugacity Model (Full-Output): Chem Name : Sodium Acifluorfen Molecular W: 383.65 Henry's LC : 1.25e-012 atm-m3/mole (calc VP/Wsol) Vapor Press : 1e-007 mm Hg (user-entered) Liquid VP : 2.84e-006 mm Hg (user-cooled) Melting Pt : 172 deg C (user-entered) Log Kow : 0.37 (Kowwin program) Soil Koc : 0.951 (calc by model) Concentration Half-Life Emissions (percent) (hr) (kg/hr) Air 1.05e-010 24 0 Water 60.4 1.44e+003 00 Fugacity Reaction Advection Reaction Advection (atm) (kg/hr) (kg/hr) (percent) (percent) Air 8.62e-022 5.14e-008 1.78e-008 2.57e-009 8.91e-01 Water 1.66e-017 493 1.02e+003 24.5 5.12 Soil 3.74e-015 493 0.02e+003 24.5 5.12 Soil 3.74e-016 493 0.024+0 0.00174 Persistence Time: 847 hr Reaction Time: 1.74e+003 hr Advection Time: 1.74e+003 hr Advection Time: 1.74e+003 hr Advection Time: 1.65e+003 hr Percent Reacte: 48.8 Percent Advected: 51.2 Half-Lives (hr), (based upon user-entry): Air: 140 Soil: 960 Sediment: 1440 Advection Times (hr): Air: 100 Water: 1000 Sediment: 2e+004 Zeiment: 2e+004 Xeter: 1000 Sediment: 2e+004 Xeter: 2e+004 Xeter: 1000 Sediment: 2e+004 Xeter: 1000 Sediment: 2e+004 Xeter: 2e+004		was used in the model. Half life in air was set at a default rapid loss sinc this material is not volatile. Emissions were calculated from using only water and soil as this test substance it is not volatile. Other parameters
Chem Name : Sodium Acifluorfen Molecular Wt: 383.65 Henry's LC : 1.25e-012 atm-m3/mole (calc VP/Wsol) Vapor Press : 1e-007 mm Hg (user-entered) Liquid VP : 2.84e-006 mm Hg (user-entered) Log Kow : 0.37 (Kowwin program) Soil Koc : 0.961 (calc by model) Concentration Half-Life Emissions (percent) (hr) (Kg/hr) Air 1.05e-012 24 0 Water 60.4 1.44e+003 1000 Soil 39.5 960 1000 Sediment 0.103 1.44e+003 0 Fugacity Reaction Advection Reaction Advection (atm) (Kg/hr) (Kg/hr) (percent) (percent) Water 1.66e-017 493 1.42e+008 2.57e-009 8.91e-01 Water 1.66e-017 493 0.0348 0.0419 0.00174 Persistence Time: 8.47 hr Reaction Time: 1.74e+003 hr Advection Time: 1.74e+003 hr Advection Time: 1.74e+003 hr Advection Time: 1.65e+03 hr Advection Time: 1.62e+03 hr	Result	:
Chem Name : Sodium Acifluorfen Molecular Wt: 383.65 Henry's LC : 1.256-012 atm-m3/mole (calc VP/Wsol) Vapor Press : 1e-007 mm Hg (user-entered) Liquid VP : 2.84e-006 mm Hg (user-entered) Log Kow : 0.37 (Kowwin program) Soil Koc : 0.961 (calc by model)Concentration Half-Life Emissions (percent) (hr) (kg/hr) Air 1.05e-010 24 0 Water 60.4 1.44e+003 1000 Soil 39.5 960 1000 Sediment 0.103 1.44e+003 0Fuggity Reaction Advection Reaction Advection (atm) (kg/hr) (kg/hr) (percent) (percent) Air 8.62e-022 5.14e-008 1.78e-008 2.57e-008 8.91e-01 Water 1.66e-017 433 0 2.25e-008 8.91e-01 Water 1.66e-017 433 0 2.24.6 51.2 Soil 3.74e-016 443 0 24.1 0 Sediment 1.38e-017 0.838 0.0348 0.0419 0.00174Persistence Time: 847 hr Reaction Time: 1.73e+003 hr Advection Time: 1.65e+081 hr Percent Advected: 51.2Half-Lives (hr), (based upon user-entry): Air: 24 Water: 1440 Soil: 960 Sediment 1.440Advection Times (hr): Air: 1000 Water: 1000 Sediment : 5e+004Test substance Flag E : CAS 62476-59-9 (acifluorfen sodium) Flag E : Critical study for SIDS endpoint 26.12.2001		
(percent) (hr) (kg/hr) Air 1.05e-010 24 0 Water 60.4 1.44e+003 1000 Soil 39.5 960 1000 Sediment 0.03 1.44e+003 0 Fugacity Reaction Advection Reaction Advection (atm) (kg/hr) (kg/hr) (percent) (percent) Air 8.62e-022 5.14e-008 1.78e-003 2.57e-009 8.91e-01 Water 1.66e-017 493 1.02e+003 24.1 0 5 Soil 3.74e-016 483 0 24.1 0 0.00174 Persistence Time: 847 hr Reaction Time: 1.74e+003 hr Advection Time: 1.74e+003 hr Advection Time: 1.55e+003 hr Advection Time: 1.64e+03 hr Advection Time: 1.64e+003 hr Advection S 1.72e Half-Lives (hr) (based upon user-entry): Air: 24 Air: 24 40 30 30 30 30 30 30		Chem Name : Sodium Acifluorfen Molecular Wt: 383.65 Henry's LC : 1.25e-012 atm-m3/mole (calc VP/Wsol) Vapor Press : 1e-007 mm Hg (user-entered) Liquid VP : 2.84e-006 mm Hg (super-cooled) Melting Pt : 172 deg C (user-entered) Log Kow : 0.37 (Kowwin program)
(atm) (kg/hr) (kg/hr) (kg/hr) (percent) (percent) Air 8.62e-022 5.14e-008 1.78e-008 2.57e-009 8.91e-01 Water 1.66e-017 493 1.02e+003 24.6 51.2 Soil 3.74e-016 483 0 24.1 0 Sediment 1.38e-017 0.838 0.0348 0.0419 0.00174 Persistence Time: 847 hr Reaction Time: 1.74e+003 hr Advection Time: 1.65e+003 hr Advection Time: 1.65e+003 hr Percent Reacted: 48.8 Percent Advected: 51.2 Half-Lives (hr), (based upon user-entry): Air: 24 Air: 24 Water: 1440 Soil: 960 Sediment: 1440 Advection Times (hr): Air: 100 Water: 1000 Sediment: 5e+004 Test substance : CAS 62476-59-9 (acifluorfen sodium) (2) valid with restrictions Flag : Critical study for SIDS endpoint (4)		(percent)(hr)(kg/hr)Air1.05e-010240Water60.41.44e+0031000Soil39.59601000
Reaction Time: 1.74e+003 hr Advection Time: 1.65e+003 hr Percent Reacted: 48.8 Percent Advected: 51.2 Half-Lives (hr), (based upon user-entry): Air: Air: 24 Water: 1440 Soil: 960 Sediment: 1440 Advection Times (hr): Air: Air: 100 Water: 1000 Sediment: 5e+004 Test substance : CAS 62476-59-9 (acifluorfen sodium) Reliability : Y : Pag : Critical study for SIDS endpoint 26.12.2001 (1)		(atm) (kg/hr) (kg/hr) (percent) (percent) Air 8.62e-022 5.14e-008 1.78e-008 2.57e-009 8.91e-01 Water 1.66e-017 493 1.02e+003 24.6 51.2 Soil 3.74e-016 483 0 24.1 0
Air: 24 Water: 1440 Soil: 960 Sediment: 1440 Advection Times (hr): Air: Air: 100 Water: 1000 Sediment: 5e+004 Test substance : CAS 62476-59-9 (acifluorfen sodium) Reliability : Flag : Critical study for SIDS endpoint 26.12.2001 (1)		Reaction Time: 1.74e+003 hr Advection Time: 1.65e+003 hr Percent Reacted: 48.8
Air:100 Water:Water:1000 Sediment:Sediment:5e+004Test substance:CAS 62476-59-9 (acifluorfen sodium)Reliability:(2) valid with restrictionsFlag:Critical study for SIDS endpoint26.12.2001		Air: 24 Water: 1440 Soil: 960
Reliability: (2) valid with restrictionsFlag: Critical study for SIDS endpoint26.12.2001(Air: 100 Water: 1000
.3.2 DISTRIBUTION	Reliability Flag	(2) valid with restrictionsCritical study for SIDS endpoint
.3.2 DISTRIBUTION		
	3.3.2 DISTRIBUTION	

3.5 **BIODEGRADATION**

3. Environmental Fate and Pathways

ld 62476-59-9 Date 26.12.2001

Type	 aerobic Studies are reported in the EPA RED documentation. This material
Inoculum	undergoes aquatic biodegradation with and estimated (EPA) half-life of 117
Remark	days.
Reliability	: (2) valid with restrictions
Flag	: Critical study for SIDS endpoint
26.12.2001	(8)

3.6 BOD5, COD OR BOD5/COD RATIO

3.7 BIOACCUMULATION

3.8 ADDITIONAL REMARKS

4. Ecotoxicity

4.1 ACUTE/PROLONGED TOXICITY TO FISH

Type Species Exposure period Unit Analytical monitoring LC50 Method Year GLP Test substance Method		static Lepomis macrochirus (Fish, fresh water) 96 hour(s) mg/l yes 62 other: EPA 660/3-75-009 1975 no other TS TEST ORGANISMS - Species: Lepomis macrochirus - Supplier: Commercial fish supplier in Missouri - Size;weight;loading: 30-38 mm; 0.31-0.73 g; <0.5 g/L - Feeding (pretreatment): dry pelleted food daily, ad libitum; discontinued 48 hours prior to test initiation - Feeding during test: none STOCK AND TEST SOLUTION AND THEIR PREPARATION - Vehicle, solvent: none DILUTION WATER - Source: deionized, reconstituted water - Chemistry (Alkalinity 32-34 mg/L;Hardness 42 mg CaCO3/L;pH 7.4;Conductance 130-160 umhos/cm) TEST SYSTEM - Test type: static - Concentrations: 0, 22, 36, 60, 100 and 170 mg a.i/L - Exposure vessel type: 20 L glass jars containing 15 L of test water - Number of fish: 10 per treatment - Photoperiod: 16 hours PHYSICAL MEASUREMENTS - Measuring times: 0, 24, 48, 72, 96 hours - Test temperature: 22-23 C - Dissolved oxygen: 73-100% (0-24 h), 52-68% (48 h), 45-73% (72 h), 40-77% (96 h) - pH: 6.6-7.3 DURATION OF THE TEST: 96 hours TEST SADMITED: madefilt/accenters
		TEST PARAMETER: mortality/symptoms OBSERVATION TIMES: 24, 48, 72, 96 hours
		ANALYSES - Method: not specified - Sampling times: 0, 96 hours
Result	:	STATISTICAL METHOD: moving average angle analysis RESULTS: - Nominal concentrations (mg a.i./L): 0, 22, 36, 60, 100, 170
		13 / 44

4. Ecotoxicity	ld	62476-59-9
4. Lootoxicity	Date	26.12.2001
Source : Test substance : Conclusion : Reliability :	 Mortality [%]: 0, 0, 10, 20, 100, 100 Other effects: fish at surface, dark discoloured, respiring rapidly and /or swimming erratically at 60 and 10 mg a.i./L Effect concentration vs. test substance solubility: At 100 and 170 mg a.i./L the test solution had a cloudy appearant which could indicate undissolved substance Notox Hertogenbosch III, CAS 62476-59-9 (Sodium acifluorfen), purity 25% (impurities not specified) 96 h LC50 62 mg a.i./L (95% CI 49-80 mg a.i./L) (2) valid with restrictions 1. No analytical results were presented in this report. It cannot be excluded that the actual concentration differed from the nominal, at least at the highest test concentrations (cloudy appearance indicating undissolved substance). The study reliability is lowered because of this. 2. Fish may have been more sensitive due to the low oxys concentration during the test (40-100%, OECD 203 >60% the long fasting (48 hours, OECD 203 24 hours). 3. The used fish were larger than recommended by OECI but acceptable according to the EG-guideline (30-38 mm, 	d gen 5) and D 203,
09.05.2001	202 20+/-10 mm, EG 50+/-20 mm).	(15)
Type : Species : Exposure period : Unit : Analytical monitoring : LC50 : Method : Year : GLP : Test substance : Method :	static Salmo gairdneri (Fish, estuary, fresh water) 96 hour(s) mg/l yes 17 other: EPA 660/3-75-009 1975 no other TS TEST ORGANISMS - Species: Salmo gairdneri - Supplier: Commercial fish supplier in Nebraska - Size;weight;loading: 30-45 mm; 0.18-0.67 g; 0.3 g/L - Feeding (pretreatment): dry pelleted food daily, ad libitum; discontinued 48 hours prior to test initiation - Feeding during test: none STOCK AND TEST SOLUTION AND THEIR PREPARAT	ΊΟN
	 - Vehicle, solvent: none DILUTION WATER - Source: deionized, reconstituted well water - Chemistry (Alkalinity 32 mg/L;Hardness 40 mg CaCO3/I 7.2;Conductance 110 umhos/cm) TEST SYSTEM - Test type: static - Concentrations: 0, 4.6, 7.8, 13, 22 and 36 mg a.i./L - Exposure vessel type: 20 L glass jars containing 15 L of test water 	_;рН
	14 / 44	

4. Ecotoxicity		62476-59-9 26.12.2001
	Date	20.12.2001
	 Number of fish: 10 per treatment Photoperiod: 16 hours PHYSICAL MEASUREMENTS Measuring times: 0, 24, 48, 72, 96 hours Test temperature: 12 C Dissolved oxygen: 69-99% (0-72 h), 50-64% (96 h) pH: 6.8-7.2 	
	DURATION OF THE TEST: 96 hours	
	TEST PARAMETER: mortality/symptoms OBSERVATION TIMES: 24, 48, 72, 96 hours	
	ANALYSES - Method: not specified - Sampling times: 0, 96 hours	
Result	 STATISTICAL METHOD: binomial probability RESULTS: Nominal concentrations (mg a.i./L): 0, 4.6, 7.8, 13, 22, 36 Measured concentrations (mg/L): not reported Mortality [%]: 0, 0, 0, 0, 90, 100 Other officiate: autimating arrationally, dark coloured 	
	- Other effects: swimming erratically, dark coloured, staying at the surface and/or lethargic at 13-36 mg a.i./L	
Source Test substance	 Notox Hertogenbosch III, CAS 62476-59-9 (Sodium acifluorfen), purity 25% 	
Conclusion	 (impurities not specified) 96-h LC50 17 mg a.i./L (95% CI 13-22 mg a.i./L) 96-h NOEC 7.8 mg a.i./L 	
Reliability	 (2) valid with restrictions 1. No analytical results were presented in this report, so it cannot be excluded that the actual concentration differe from the nominal. The study reliability is lowered because of this. 2. Fish may have been more sensitive due to the long fas (48 hours, OECD 203 24 hours) and due to their small siz (30-45 mm, OECD 203 50+/-10 mm). 	ting
09.05.2001	· · · /	(14)

4.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES

Туре	: static	
Species	: Daphni	a magna (Crustacea)
Exposure period	. 48 hour	(S)
Unit	: mg/l	
Analytical monitoring	: yes	
EC50	: 77	
Method	:	
Year	:	
GLP	: no	
Test substance	: other T	S
Method	: TEST C	DRGANISMS
	-	es: Daphnia magna
	- Sourc	e/supplier: Bionomics culture facility
		ing method: Culture of Daphnia in water with hardness

4. Ecotoxicity	ld 62476-59-9
	Date 26.12.2001
	of 165 mg CaCO3/L, pH 7.9-8.3, temperature 22+/-1 C, Oxygen >60% (same as test water) - Age: <= 20 hours - Feeding before and during test: not specified
	STOCK AND TEST SOLUTION AND THEIR PREPARATION - Vehicle, solvent: none
	DILUTION WATER - Source: Deionized, reconstituted well water - Chemistry (Alkalinity 120 mg/L;Hardness 160-170 mg/L/pH 8.0-8.2/Conductance 440-450 umhos/cm)
	TEST SYSTEM - Test type: static - Concentrations: 0, 13, 22, 36, 60, 100 mg a.i./L - Exposure vessel type: 250 mL beakers containing 200 mL test solution - Number of individuals: 5 per replicate, 4 replicates/treatment - Photoperiod (intensity of irradiation): illuminated at 538-753 lux PHYSICAL MEASUREMENTS - Measuring times: 0, 24 (only temperature) and 48 hours - Test temperature: 21 C - Dissolved oxygen: 94-100% - pH: 8.0-8.2 DURATION OF THE TEST: 48 hours
	TEST PARAMETER: mortality/symptoms OBSERVATION TIMES: 0, 24, 48 hours ANALYSES
	- Method: not specified - Sampling times: 0 and 48 hours
Result	 STATISTICAL METHOD: moving average angle method RESULTS: Nominal concentrations (mg a.i./L): 0, 13, 22, 36, 60, 100 Measured concentrations (mg/L): not reported Immobility [%]: 0, 0, 0, 13, 13, 90 Other effects: lethargic at 36-100 mg/L
Source Test substance	 Notox Hertogenbosch III, CAS 62476-59-9 (Sodium acifluorfen), purity 25% (impurities not specified)
Conclusion Reliability	 48 h EC50: 77 mg a.i./L (95% CI 66-94 mg a.i./L) (1) valid without restriction 1. Analyses were performed, but the results were not included in the report. Since analyses were not recommended by OECD 202, the study reliability was not lowered. 2. There was no information on the feeding of the Daphnia.
	 Analyses were performed, but the results were not included in the report. Since analyses were not recommended by OECD 202, the study reliability was not lowered. There was no information on the feeding of the Daphnia.

Id 62476-59-9 4. Ecotoxicity Date 26.12.2001 4.3 TOXICITY TO AQUATIC PLANTS E.G. ALGAE 4.4 **TOXICITY TO MICROORGANISMS E.G. BACTERIA** 4.5.1 CHRONIC TOXICITY TO FISH 4.5.2 CHRONIC TOXICITY TO AQUATIC INVERTEBRATES 4.6.1 TOXICITY TO SOIL DWELLING ORGANISMS Type : artificial soil **Species** Eisenia fetida (Worm (Annelida), soil dwelling) : Endpoint : mortality Exposure period : 14 day mg/kg soil dw Unit : LC50 > 1800 : Method : OECD Guide-line 207 "Earthworm, Acute Toxcity Test" Year : 1984 GLP : no Test substance : other TS Method TEST ORGANISMS : - Species: Eisenia foetida - Age/weight: 2-5 months/450-530 mg (mean) - Keeping/breeding conditions: cultures of worms were maintained in jars with horse manure/sphagnum peat (2:1) at 20+/-2 C under continuous illumination. Test animals were overnight conditioned to artificial soil medium. TEST SOLUTION AND THEIR PREPARATION - Vehicle, solvent: distilled deionised water - Application procedures: test substance in water was added to partly moistured soil, mixed carefully and subsequently the moisture level was adjusted to 35% of dry weight with water ARTIFICIAL SOIL

- Kaolin clay: 20% - Fine sand: 70% - Calcium carbonate: 0.25-1% - pH: 6+/-0.5
TEST SYSTEM - Test type: artificial soil test - Concentrations: 0, 180, 320, 560, 1000, 1800 mg/kg dw - Exposure vessel type: 1 L covered glass beaker containing 750 g soil (wet weight) - Number of worms: 10 per replicate, 4 replicates/treatment

- Spahagnum peat: 10%

4. Ecotoxicity	ld 62476-59 Date 26.12.200	-
Result	 Photoperiod (light intensity): not indicated, but it was reported that all worms stayed below the soil surface during the test PHYSICAL MEASUREMENTS Measuring times: start and end Moisture level (% of dw): 35% pH: 5.0-5.6 temperature: 17-24 C DURATION OF THE TEST: 14 days TEST PARAMETER: mortality/symptoms OBSERVATION TIMES: 1, 3, 7 and 14 days REFERENCE SUBSTANCE: 2-chloroacetamide STATISTICAL METHOD: Litchfield and Wilcoxon, Probit analysis, Thompson's moving average procedure RESULTS: Nominal concentrations (mg a.i./L): 0, 180, 320, 560, 1000, 1800 Mortality (%): 0, 0, 0, 2.5, 2.5, 30 Body weight: no dosis related effects 	
	 Body weight: no dosis related effects Other effects: at 560, 1000 and 1800 mg a.i./kg the worms were found clustered together near the surface Dose related effects: yes 	
Source Test substance Reliability	 RESULTS: TEST WITH REFERENCE SUBSTANCE Concentrations: 0, 17.8, 26.7, 40, 60, 90 mg/kg Results: 14-d LC50 25-31 mg/kg Notox Hertogenbosch III, CAS 62476-59-9 (TACKLE 2AS formulation), purity 21.6% (2) valid with restrictions There is a discrepancy in the report concerning the mortality of the worms. At 560 mg a.i./L it is not clear whether none of the worms died, or 1 worm died. Therefore in this summary it is assumed that 1 worm died in this dose level. 	
10.05.2001	2. non-GLP study	(17)
	RRESTRIAL PLANTS HER NON-MAMM. TERRESTRIAL SPECIES	
4.7 BIOLOGICAL EF	FECTS MONITORING	
4.8 BIOTRANSFORM	IATION AND KINETICS	
	18 / 44	

4. Ecotoxicity	Ecotoxicity Id 62476-59-9 Date 26.12.2001	
4.9 ADDITIONAL REMARKS		

5. Toxicity

5.1.1 ACUTE ORAL TOXICITY

Type Species Strain Sex Number of animals Vehicle Value Method Year GLP Test substance Method		LD50 rat other: CF Nelson male 10 water = 122 mg/kg bw other: not indicated no other TS TEST ORGANISMS: - Source: not indicated - Age: not indicated - Age: not indicated - Age: not indicated - Number: 10/dose - Weight at study initiation: 196-201 g (mean) - Controls: no ADMINISTRATION: - Doses: 625, 1250, 2500 and 5000 mg/kg - Doses per time period: single - Volume administered or concentration: 20% (w/v) - Post dose observation period: 14 days - food withheld for 24 hours pre-dosing EXAMINATIONS: signs of intoxication and gross necropsy BODY WEIGHT: pre-dosing and at the end of the test
Result	:	STATISTICAL METHOD: not indicated MORTALITY: - Number of deaths at each dose: 625, 1250, 2500 and 5000 mg/kg bw: 0/10, 3/10, 9/10 and 10/10, resp. - Time of death: for the highest dose: within 6 hours, for the other doses: within two days. CLINICAL SIGNS: lethargy, prostration and ataxia at 2500 and 5000 mg/kg bw
		BODY WEIGHT: no effects
Source Test substance Conclusion Reliability	:	NECROPSY FINDINGS: no visible lesions in the survivors Notox Hertogenbosch III, CAS 62476-59-9 (sodium 5-(2-chloro-4-trifluoro-methylphenoxy)2-nitrobenzoate, purity 39.6%, used as 20% (w/v) aqueous dispersion LD50 1540 mg/kg bw (2) valid with restrictions 1. The information was essentially confined to what is included in the current summary. No individual data were present.
18.04.2001		

5. Toxicity	ld 62476-59-9 Date 26.12.2001
5.1.2 ACUTE INHALATION T	OXICITY
Type:Species:Strain:Sex:Number of animals:Vehicle:Exposure time:Value:Method:Year:GLP:Test substance:Method:	LC50 rat other: albino King (Kng:(SD)BR) male/female 10 other: no vehicle 4 hour(s) > 1.38 mg/l other: not indicated no other TS TEST ORGANISMS: - Source: King Animal Laboratories, Inc., Oregon, WI - Age: not specified - Weight at study initiation: males (246-291 g) and females (217-248 g) - Number of animals: 5/sex/dose - Controls: yes ADMINISTRATION: - Type of exposure: whole body exposure to aerosol - Exposure duration: 4 hours - Concentrations(nominal/measured): 17.9 / 6.91 mg/l (analytical conc.) or 2.6 mg/l (gravimetric conc.) - Particle size: mass median diameter: 2.11 micrometer with standard deviation 2.59 micrometer (first sample) and 3.65 micrometer with standard deviation of 2.20 micrometer (second sample).
	 Type or preparation of particles: air atomizing nozzle assembly Air changes: >= 15/hr EXAMINATIONS: for pharmacotoxic signs (during exposure and twice daily during 14 days post-exposure time); gross necropsy
	BODY WEIGHT: pre-exposure and at days 7 and 14
	ANALYSES: - Method: gravimetry and analytical concentration by extraction/spectrophotometry - Sampling times: 4 times/4 hours - Particle size determination at 1 and 3 hours
Result :	STATISTICAL METHOD: not specified MORTALITY: - Number of deaths at each dose: no deaths
	CLINICAL SIGNS: during exposure: squinting, nasal discharge, dyspnea and lacrimation; shortly after exposure: nasal discharge, dyspnea, crusty nose and yellow/brown stained fur; during the 14-day observation period: nasal discharge, crusty nose, yellow/brown stained fur, crusty mouth and poor

5. Toxicity	ld 62476-59-9
	Date 26.12.2001
	coat quality. The control group did not show any clinical signs
	BODY WEIGHT: no treatment-related effects
	NECROPSY FINDINGS: one treated rat with focal depressions of the lung; for the control animals: 2 rats with lung lesions and 1 rat with diaphramatic hernia of the liver.
Source Test condition	 SEX-SPECIFIC DIFFERENCES: no data Notox Hertogenbosch III, CAS 62476-59-9 (TACKLE 2AS formulation), 20% w/w aqueous solution
Conclusion Reliability	 LC50 > 6910 mg/m³ (2) valid with restrictions The obtainment of the results for the exposure chamber (nominal concentration, airchanges/hr) are unclear. The gravimetric measured concentration of 2.6 mg/l is less reliable than the analytical measured concentration. Only a QA statement was included, but no GLP statement
23.04.2001	signed by the study director. (25)
5.1.3 ACUTE DERMAL	TOXICITY
Type Species Strain Sex Number of animals Vehicle Value Method Year	: LD50 : rabbit : other: Albino : male : 5 : other: no vehicle : = 1457 mg/kg bw : other: not specified
fear GLP Test substance Method	: no conter TS TEST ORGANISMS: - Source: not indicated - Age: not indicated

ADMINISTRATION: - Area covered: not specified

Result

- Occlusion: yes
 Vehicle: no vehicle, test substance is an aqueous solution
 Doses: 2500, 3540 and 5000 mg/kg bw
- Removal of test substance: not indicated

EXAMINATIONS: signs of intoxication, skin irritation and gross necropsy

BODY WEIGHT: pre-dosing and at end of the test

STATISTICAL METHOD: not indicated : MORTALITY:

5. Toxicity	ld 62476-59-9 Date 26.12.2001
	 Number of deaths at each dose: 2500, 3540 and 5000 mg/kg bw: 1/5, 2/5 and 4/5, resp. Time of death: at 2500 and 3540 mg/kg bw, within 4 days; at 2500 mg/kg bw, between days 8 and 14.
	CLINICAL SIGNS: lethargy, ataxia, shallow respiration and prostration; well defined to moderate erythema, slight edema, followed by desiccation and flaking of skin at 3540 and 5000 mg/kg bw.
	BODY WEIGHT: increased bw for the lowest dose survivors; decreased bw for the two highest doses survivors.
	NECROPSY FINDINGS: no visible lesions for the decendents at 3540 and 2500 mg/kg bw; no visible lesions for the survivors.
Source Test substance	 Notox Hertogenbosch III, CAS 62476-59-9 (sodium 5-(2-chloro-4-trifluoro-methylphenoxy)2-nitrobenzoate, purity 39.6% aqueous technical
Conclusion Reliability	 LD50 3680 mg/kg bw (2) valid with restrictions The information was essentially confined to what is included in the current summary. No individual data were present. Protocols were attached to the document, but they were not related to this test.
18.04.2001	3. Only males were tested. (20)

5.1.4 ACUTE TOXICITY, OTHER ROUTES

5.2.1 SKIN IRRITATION

5.2.2 EYE IRRITATION

5.3 SENSITIZATION

5.4 REPEATED DOSE TOXICITY

Species	: rat
Sex	: male/female
Strain	: Fischer 344
Route of admin.	: oral feed
Exposure period	: 90 days
Frequency of	: daily
treatment	
Post obs. period	: None
Doses	: 1.7-422 mg/kg bw/day

5. Toxicity	ld 62476-59-9 Date 26.12.2001
Control group	: yes : = 23.7 mg/kg bw
Method	: other: FIFRA 83-2
Year	: 1978
GLP	: yes
Test substance	: other TS
Method	: TEST ORGANISMS:
	- Species/strain: Fischer 344 rats
	- Source: Charles River Breeding Laboratories Inc.
	- Age: six weeks - Weight at study initiation: male (130g), female (100g)
	- Number of animals: 30/sex/dose group
	ADMINISTRATION / EXPOSURE
	- Exposure period: 90 days
	 Route of administration: diet Post exposure period: none
	- Post exposure period: none - Doses: 0, 20, 80, 320, 1250, 2500, and 5000 ppm. which
	resulted in actual intakes of 1.5, 6.1, 23.7, 92.5, 191.8
	and 401.7 mg/kg bw/day in males and 1.8, 7.4, 29.7, 116.0,
	237.1 and 441.8 mg/kg bw/day in females
	CLINICAL OBSERVATIONS AND FREQUENCY:
	- Clinical observation and mortality: Twice daily, detailed
	examination weekly - Body weight: at baseline and weekly therafter
	- Food consumption: weekly
	CLINICAL CHEMISTRY:
	In 10 animals/sex/dose group, at day 30 and at study
	termination;
	 Hematology, hematocrit, hemoglobin, erythrocyte, count, mean corpuscular volume, total and differential leukocyte
	counts, platelet count, reticulocyte count.
	- Biochemistry (in 10 animals/sex/dose group): at day 30 and
	at study termination; Serum lactate dehydrogenase (LDH),
	serum glutamic oxaloacetic transaminase (SGOT), serum
	glutamic pyruyic transaminase (SGPT), serum alkaline phosphatase, albumin, creatinine phosphokinase (CPK),
	glucose, blood urea nitrogen (BUN), direct bilirubin, total
	bilirubin, total cholesterol, globulin, indirect bilirubin,
	triglyceride, total protein, creatinine, calcium, uric acid,
	sodium, inorganic phosphorous, chloride, potassium.
	- Urinalysis: specific gravity, pH, protein, glucose,
	ketones, bilirubin, urobilinogen, nitrite, hemoglobin and microscopic examination for cells or formed elements.
	ORGANS EXAMINED AT NECROPSY (MACROSCOPIC AND MICROSCOPIC):
	- Organ weights (at day 30 (10 animals/sex/dose) and at
	termination): liver, kidneys, heart, testes, and brain,
	including entire brain system.
	- Macroscopic and microscopic (control and high dose group):
	eyes and the contiguous Harderian glands; heart; thyroid
	(with parathyroid); trachea; esophagus; stomach; adrenal
	24 / 44

5. Toxicity	ld 62476-59-9 Date 26.12.2001
S. IOXICITY	
	at 30 days; elevated albumin/globulin ratio at three highest doses (males) and highest dose (females); depressed serum calcium levels at 5000 ppm and increased phosphorus in males, in females to a lesser extent; elevated alkaline phosphatase and serum G/P transaminase at 5000 ppm in both sexes indications of reduced renal function: significant increase in blood urea nitrogen in both sexes at 30 days for males at
	2500 and 5000 persistent at 90 days; increased BUN/creatinine ration in males at 30 days but not at 90

5. Toxicity	ld 62476-59-9 Date 26.12.2001
	days; significantly different values of uric acid for both sexes (without consistent trend)
	- Urinalysis: at 30 days: increased urobilinogen in males at 5000 ppm (other measures of bilirubin showed little deviation); slightly diminished protein excretion in both sexes at 5000 ppm; increased frequency of trace amounts of nitrite in males above 320 ppm at 90 days: increased urobilinogen in both sexes at 2500 and 5000 ppm; decreased protein excretion with increasing dose in females for males only at 5000 ppm; increased frequency of trace amounts of nitrite in females at 2500 and 5000 ppm
	MACRO- AND MICRSCOPIC FINDINGS - Organ weights: significantly increased liver and kidney weight, both absolute and relative, in males above 320 ppm at 30 and 90 days (except at day 30 for 2500 ppm), females to a lesser extent at 2500 and 5000 ppm on day 30 and at 5000 ppm on day 90); sporadic deviation in heart and brain weight (no toxicological pattern); increased relative testis weight (not considered significant) were a function of reduced overall body weight and are not considered significant. - Macroscopy: Interim kill - 30 Days: control animals: diffuse brown discoloration of the kidney (1 male); enlargement of left mandibular lymph node (1 male); 5000 ppm: liver (diffuse dark staining) and kidney (cortex darkening or diffuse discoloration) discoloration in both males and females 90 days: no abnormalities in controls, at 5000 ppm dark brown discoloration of the liver and kidney (dark brown
	 cortexes) in both males and females (females less affected) - Histopathology: Interim Kill - Day 30: Presence of mononuclear cells in the lungs in both control and treatment group (not test substane related) 5000 ppm: increased liver cell hypertrophy in both sexes; increased mitotic figures in males and females (but to a lesser extent); liver tissue damage in both sexes Terminal Kill - Day 90: Both control and treatment group showed presence of mononuclear cells and vascular mineralization in the lung and cysts in various organs (all considered not treatment related); Controls: cell death in liver in part of the males 5000 ppm: cell death and hypertrophy in liver cells of all males, in females only hypertrophy in part of the animals and no cell death; increased proliferation of oval cells and bile duct in majority of males; yellow pigmentation of Kupfer cells in all treated males ANALYSES: In all cases diet formulation concentrations and test

5. Toxicity	ld 62476-59-9	
-	Date 26.12.2001	
	substance concentrations were within 10% tolerance limits	
Source	: Notox Hertogenbosch	
Test substance	: III, CAS 62476-59-9 (TACKLE 2AS formulation), purity	
	20-21.6%	
Conclusion	: NOAEL 320 ppm (23.7 mg/kg bw) based on the presence of liver	
Deliebility	damage with concomittant changes in blood chemistry	
Reliability 21.05.2001	: (1) valid without restriction	(12)
21.03.2001		(12)
Species	: rabbit	
Sex	: male/female	
Strain	: New Zealand white	
Route of admin.	: dermal	
Exposure period	: 21 days	
Frequency of	: 5 days/week	
treatment		
Post obs. period	: None	
Doses	: 92, 277 and 923 mg/kg bw	
Control group	: yes, concurrent vehicle	
NOAEL	: = 277 mg/kg bw	
LOAEL	= 92 mg/kg bw	
Method	: EPA OPP 82-2	
Year GLP		
Test substance	: yes : other TS	
Method	: TEST ORGANISMS:	
motriou	- Species/strain: New Zealand white rabbits	
	- Source: H.A.R.E., Hewitt, NJ.	
	- Age: no data	
	- Mean Weight at study initiation: 2.59-2.64 (females),	
	2.65-2.68 (males)	
	- Number of animals: 10/sex/dose group	
	ADMINISTRATION / EXPOSURE	
	- Exposure period: 21 days	
	- Route of administration: dermal	
	- Post exposure period: none	
	- Doses: 92, 277 and 923 mg/kg bw, at day 4 highest dose was	
	reduced to 4.62 mg/kg bw - Vehicle: A NaOH solution (not specified) pH 7.5-7.6	
	- Total volume applied: 1ml, 3ml, 10ml (5ml after day 4)	
	- Area covered: 130cm2	
	- Occlusion: two layers of clean gauze plus occlusive	
	binders for six hours	
	- Removal of test substance: after 6 hours	
	CLINICAL OBSERVATIONS AND FREQUENCY:	
	- Clinical signs: daily observation for external signs of	
	toxicity. Dermal irritation readings according the method of	
	Draize (1965) daily prior to application.	
	- Mortality: twice daily	
	- Body weight: day -1 thereafter the 4th and 7th day of the	
	week, at sacrifice	
	- Food consumption: on day 1, 4, 7, 11, 14 and 21	
	CLINICAL CHEMISTRY	
	- Haematology: Total and differential leukocyte counts,	
	- Haematology. Total and unreferitial leukocyte counts,	

5. Toxicity	Id 62476-59-9
-	Date 26.12.2001
	erythrocyte count, hematocrit, hemoglobin, platelet count - Biochemistry: alkaline phosphatase, urea nitrogen, glutamic pyruvate transaminase, glutamic oxaloacetate transaminase, calcium, potassium, lactic dehydrogenase, glucose, bilirubin (total and direct), total cholesterol, albumin, globulin, total protein - Urinalysis: appearance, specific gravity, occult blood, protein, pH, bilirubin, urobilinogen, ketones, glucose, microscopic examination of formed elements
	ORGANS EXAMINED AT NECROPSY (MACROSCOPIC AND MICROSCOPIC): - Organ weights: adrenal glands, brain, heart, kidneys, liver, gonads, pituitary gland, thyroid and parathyroid. - Macroscopic: abdominal cavity, abdominal wall, adipose tissue, adrenals, bladder, diaphragm, epidydimes, gallblader, heart, large and small intestine, kidneys, liver, lungs, lymph nodes, mouth, nose, ovaries, pancreas, pituitary, salivary glands, sciatic nerve, skeletal muscle, treated and untreated skin, spleen, stomach, testes, thoracic cavity, thymus, thyroid, ureters, uterus and vagina - Microscopic: treated and untreated skin, liver, kidneys and grossly abnormal tissue
	ANALYSES: -Method: HPLC analysis of test compound: isocratic 65% methanol/35% water 2ml/min on a waters radial compression system radial-pak A C18, detector 280nm.
	 Sampling times: at study initiation and during week 1 and 2
Remark	 STATISTICAL METHODS: one-way analyses of variance (continuous data), Least Significant Difference (differences among groupes), Mantel-Haenszel chi-square test (score data), chi-square with Yates correction (pathology data) Tables with individual histopathological data are partly
Result	 missing. CLINICAL OBSERVATIONS AND MORTALITY Mortality and time to death (day): at 923 mg/kg bw 19/20 died ore were sacrificed before day 8, one male survived
	until sacrifice; at 92 mg/kg bw 1 male (8); at 277 mg/kg bw 1 male (13); controls one male and one female (21) - Clinical signs: at highest dose ataxia, decreased activity, nasal discharge, respiratory distress and salivation was seen in both sexes, males showed incidently diarrhoea and tremors; at 277 mg/kg bw incidental nasal discharge, hair loss, soft stool, tremors, diarrhoea and bloating was seen; at the lowest dose incidental signs were confined to diarrhoea and bloating; in all dose groups a white chrystaline substance at the application site was observed.
	Severe dermal irritation with eschar formation was seen in males and females from day 2-3 to day 21 of exposure. A relationship with amount of applied material was evident. - Body weight gain: decreased body weight in highest dose

5. Toxicity	ld 62476-59-9	
	Date 26.12.2001	
	group (significant in females) - Food/water consumption: individual low daily food consumption in high dose animals, significantly decreased on days 1-4 CLINICAL CHEMISTRY No treatment related effects MACRO- AND MICROSCOPIC FINDINGS - Organ weights: at 277 mg/kg bw significant increase in mean relative adrenal weight in females (toxicological significance questionable) - Macroscopy: marked dermatitis with epithelial necrosis and eschar formation at the exposure site for all exposure levels. - Histopathology: microscopic changes indicative of macroscopic findings, all other findings were incidental and	
	not related with treatment. Effects on intestinal epithelium were attributed to coccidal infections ANALYSES: - Actual dose was 87-106% of nominal value - Stability: ok - Homogeneity: ok	
Source Test substance	 Notox Hertogenbosch CAS 62476-59-9 (Acifluorfen, sodium salt), purity: technical acifluorfen was dissolved in 0.82 M NaOH yielding a preparation of 240 mg/ml liquid 	
Conclusion	 Tackle 2S was acutely toxic when administered at the high dose. Body weight gain and food consumption were decreased in high dose animals. Nineteen of 20 animals receiving the high dose did not survive past day eight of the study. In addition Tackle 2S was a severe cumulative dermal irritant at all dose levels. No toxicologically significant changes in body weight, food consumption, hematological and clinical chemistry parameters, or urinalysis data were observed among control, low dose, and mid dose groups. NOAEL systemic 277 mg/kg based on survival and body weight LOAEL local effects 92 mg/kg 	
Reliability	 (2) valid with restrictions 1. limited histopathology 2. effect on adrenal weight is questionable 	
21.05.2001	(11)	

5.5 GENETIC TOXICITY 'IN VITRO'

Туре	: Cytogenetic assay
System of testing	: CHO cells
Concentration	: 0.5-5.0 ul/ml
Cycotoxic conc.	:
Metabolic activation	: without
Result	: negative
Method	: other
Year	:
GLP	: no data
Test substance	: other TS

5. Toxicity	ld 62476-5 Date 26.12.2	
Method	 Species/cell type: Chinese hamster ovary cells Metabolic activation system: none No. of anaphases analyzed: 300 	
	ADMINISTRATION: - Doses: 0.5, 1.0 and 5.0 ul/ml - Exposure period: 3 hours - Positive and negative control groups and treatment: Positive control was Ethylmethanesulfonate (EMS) and was added at 0.5 μ1/ml; spontaneous controls were also maintained.	
Result	 CRITERIA FOR EVALUATING RESULTS: - assesment of mitotic spindle damage by screening cells microscopically for multinuclei or anaphase bridges - Statistical method: Chi square analysis : GENOTOXIC EFFECTS: - Without metabolic activation: none 	
	PRECIPITATION CONCENTRATION: no details given. CYTOTOXIC CONCENTRATION: no information available	
	STATISTICAL RESULTS: There was no significant difference between controls and test samples regarding mititic spindle damage.	
Source Test substance	 Notox Hertogenbosch CAS 62476-59-9 (sodium 5-(2-chloro-4-trifluoro-methylphenoxy) 2-nitrobenzoate), purity not indicated 	
Reliability	: (3) invalid 1. No standard study type; pilot study	
21.05.2001	The standard study type, pilot study	(13)

5.6 GENETIC TOXICITY 'IN VITRO'

Cytogenetic assay
mouse
male/female
CD-1
gavage
single dose
0, 100, 500, 1000 mg/kg.
negative
OECD Guide-line 475 "Genetic Toxicology: In vivo Mammalian Bone
Marrow Cytogenetic Test - Chromosomal Analysis"
1986
yes
other TS
TEST ORGANISMS:
- Strain: Crl:CD-1(ICR)BR mice
- Source: Charles River Kingston Breeding Laboratories
(Stoneridge, New York)
- Age: no data
- Weight at study initiation: 18.5 - 28.5 g
- No. of animals per dose: 15/sex/dosage

ld 62476-59-9 Date 26.12.2001
ADMINISTRATION: - Vehicle: distilled water - Doses: Test compound: 0, 100, 500, 1000 mg/kg by gavage. The corresponding dose levels based on active ingredient are 0, 42.8, 214, 428 mg/kg, respectively. - Duration of test: The in-life portion of the study was 3 days. Ten animals of each dose group were killed 6, 27, and 51 hr after dosing. - Frequency of treatment: single dose by oral gavage - volume 10 ml/kg. - Control groups and treatment: Negative control: vehicle 15 animals per sex. Positive control: Triethylmelamine, ip 0.3 mg/kg (5 animals per sex). - number of metaphases scored: 50/animal
EXAMINATIONS: - Clinical signs and mortality: daily. - Body weight: daily for 4 days (separate group of 8 animals)
CRITERIA FOR EVALUATING RESULTS: - no. of cells with aberrations per 5 animals
STATISTICAL ANALYSIS: The Beta-binomial model (Stiratelli et al., 1985)MORTALITY: none
CLINICAL SIGNS: Yellow stained anogenital area, passiveness, ruffled fur, and abdominal breathing were observed after treatment with 428 mg/kg test material and at a lower incidence at 214 mg/kg test material. Recovery was observed. Abnormal toxic signs were not observed in the animal positive control, distilled water control groups or test material 42.8 mg/kg treatment group prior to sacrifice.
BODY WEIGHT CHANGES: no effect
GENOTOXIC EFFECTS: No. of cells with aberrations at 6, 27 and 51 hours 11, 11 and 12 respectively (12, 11 and 5 in vehicle controls)
 POSITIVE CONTROL: A significant increase in the frequency of bone marrow chromosomal aberrations and an increase in translocations and rearrangements Notox Hertogenbosch III, CAS 62476-59-9 (Acifluorfen, sodium salt), purity 42.8% Negative, solvent and positive controls were within the expected ranges. (2) valid with restrictions Only slides of 1000 mg/kg were scored for genotoxic effects. Slides of the lower dose groups were not examined because an effect did not occur at the highest dose group. Only 50 metaphases per animal were scored (100 according

5. Toxicity

ld 62476-59-9 Date 26.12.2001

21.05.2001

(19)

5.7 CARCINOGENITY

5.8 TOXICITY TO REPRODUCTION

Туре	:	Two generation study
Species		rat
Sex	:	male/female
	:	
Strain		other: Crl:COBS-CD-(SD)BR
Route of admin.	:	oral feed
Exposure period	:	Parent/F1-generation (males/females): 12 weeks before cohabitation for mating until completion of a 3-week cohabitation period for males or until day 25 of presumed pregnancy (non-pregnant females) or day 21 of lactation (pregnant females)
Frequency of	:	continuous
treatment		
Premating exposure		
• •		
period		
Male		12 weeks
Female		12 weeks
Duration of test	:	42 weeks (maximum): Parent/F1-generation; 12 weeks
		premating/treatment, 3 weeks cohabitation, 3 weeks pregnancy, 3 weeks
		lactation
Doses	:	25, 500 and 2500 ppm in the diet
Control group		other: diet without the test substance
NOAEL Parental	:	
NOAEL F1 Offspr.	:	= 500 ppm
NOAEL F2 Offspr.	:	= 500 ppm
Method	:	other: US EPA, Pesticide Assessment Guideluines, Subdivision F, Hazard
		Evaluation: Human and Domestic Animals.
Year	:	1982
GLP	:	yes
Test substance	:	other TS
Method		
Method	•	TEST ORGANISMS
		- Age: males/females (parental generation) 7 weeks at start
		treatment
		- Source: Charles River Breeding Laboratories Inc.,
		Kingston, NY
		 Weight at study initiation: At start treatment males
		177-238g and females 123-169g
		 Number of animals: 35/sex/treatment (parent),
		40/sex/treatment (F1)
		ADMINISTRATION / EXPOSURE
		- Test duration: maximum 39 weeks
		 Exposure period: males (parent/F1 generation) 12 weeks
		prior to mating and maximal 3 weeks cohabitation;
		Females (parent generation) 12 weeks prior to mating,
		maximal 3 weeks cohabitation, 3 weeks pregnancy and 3 weeks
		lactation Females (F1-generation) after weaning 12 weeks
		prior to mating, maximal 3 weeks cohabitation, 3 weeks
		pregnancy and 3 weeks lactation
		22/11
		377777

5. Toxicity	ld 62476-59-9 Date 26.12.2001
	 Route of administration: oral via the diet Doses: 0, 25, 500 and 2500 ppm in the diet (actual exposure in terms of the average mg/kg/day dosage was calculated to be higher in females than in males for each generation and within each sex the second generation received higher mg/kg/day dosages than the first generation)
	MATING PROCEDURES: - Mating: 1 female / 1 male - Day 0 of gestation: presence of copulation plug and/or spermatozoa in the vaginal smear of females
	 PARAMETERS ASSESSED DURING STUDY: Mortality: mimimum of twice each day Clinical observations: daily during exposure Body weight gain: at least once weekly during exposure, during gestation on day 0, 6, 10, 15, 20 and 25, during lactation on day 1, 4, 7, 11, 14, 16, 18 and 21 Food consumption: at least once weekly during exposure, during gestation on day 0, 6, 10, 15, 20 and 25, during lactation on day 1, 4, 7, 11, 14, 16, 18 and 21 Female oestrous cycle: vaginal cytology examination during cohabitation and until confirmation of pregnancy (maximum 3 weeks) Mating and fertility data (males/females): days in cohabitation, number of males/females): days in cohabitation period when examining the pups Maternal behaviour (dams which delivered): during the 3-week lactation period when examining the pups Maternal delivery data: duration of gestation, number pregnant and surviving delivery, number surviving with still borns, litter size (live and dead pups), number and placements of implants at sacrifice (day 21 of lactation) Pup viability: vital status at birth (live or stillborn) and at least twice daily viability until culling (day 4 post-partum for the parent generation, maximum 8 pups/litter) or weaning (day 21 post-partum for the parent generation, maximum 8 pups/litter) or weaning (bays 1 (birth), 4, 7, 14 and 21 of lactation; body weights on days 1 (birth), 4, 7, 14 and 21 of lactation;
	ORGANS EXAMINED AT NECROPSY (MACROSCOPIC AND MICROSCOPIC): - Macroscopy: all males and females (parental generation) and those selected for pairing (F1-generation) were
	necropsied and gross findings recorded and all gross lesions, target organs (liver, kidney and stomach), pituitary gland and reproductive organs (males: testes, epididymides, seminal vesicles, prostate and coagulation gland and females: vagina, uterus, cervix, ovaries and mammary gland) were removed and preserved in fixative. All pups, except those precluded by autolysis or cannibalism, were necropsied and examined for gross lesions. Additionally, at weaning the heads of pups (except those
	22 / 44

5. Toxicity	ld 62476-59-9
	Date 26.12.2001
	selected for pairing) were cross-sectioned for examination of hydrocephaly - Microscopy: histopathology examinations were preformed on the kidney, stomach and gross lesions of rats of the parental generation and on gross lesions of pups of the F1 and F2 generations). The reproductive organs, liver and pituitary gland were examined from 20 selected males and females of the control and high dosage groups of the parental and F1 generations
	ANALYSES: - Method: HPLC/UV - Sampling time: weekly (accuracy of preparation) and on days 0, 1, 4, 7, 10, 14 and 21 (stability and homogeneity)
Result	 STATISTICAL METHODS: Bartlett's test, Analysis of variance, Dunnett's test, Kruskal-Wallis test, Dunn's test, Analysis of covariance, Covariance Analysis T-test, Variance test for the Homogeneity of the Binomial Distributio ANALYSES: Actual dose level: the accuracy of all test diets was acceptable (within 15% of nominal concentrations); in control diet from week 26 onwards significant amounts of test substance (compared to the low dose level) were found Stability: stable for at least 21 days (mean recovery 82-91%) Homogeneity: homogeneous (first batch recovery 72-120%, two samples at 250 ppm (mid) 131 and 184%; second batch
	84-113%) ACTUAL INTAKE (mg/kg bw): Males premating (P/F1): at 50, 500 and 2500 ppm, 1.5-1.7, 29-33 and 147-169 mg/kg bw resp Females premating (P/F1): at 50, 500 and 2500 ppm, 1.8-1.9, 29-38, 153-199 mg/kg bw resp Females pregnancy (P/F1) at 50, 500 and 2500 ppm, 1.5-1.6, 29-30, 153-157 mg/kg bw resp Females Lactation (P/F1): at 50, 500 and 2500 ppm, 2.9-3.2, 57-61, 252-287 mg/kg bw resp
	TOXIC EFFECTS BY DOSE LEVEL PARENTAL GENERATION:
	 Mortality: at 25 ppm one female and at 2500 ppm one male Body weight: at 2500 ppm decreased in males and females and at 500 ppm increased in females during lactation only Food consumption: at 500 ppm decreased in females (day 6-15 of gestation) and at 2500 ppm decreased in males and in females during lactation Clinical signs: at 2500 ppm increased chromodacryorrhoea and urine stained abdominal fur in males and emaciation in
	34 / 44

females	
---------	--

- Mating and fertility data (males/females): no differences between the dose groups; at 0, 50, 500 and 2500 ppm 30, 29, 31 and 32 females pregnant

- Maternal delivery data: no treatment related effects on duration of gestation, surviving dams/pups; at 2500 ppm decreased number of implantations sites

- Pup data: no differences between the dose groups considering viability and sex ratio: at 2500 ppm decreased pup weights between birth and day 21 post-partum

 Macroscopic examinations: very low incidences of mottled appearance of the renal pelvis in males at 500 and 2500 ppm; stomach with dark red to black areas in females at 2500 ppm
 Microscopic examinations: at 500 and 2500 ppm kidney lesions characterised by dilation of tubules in the outer medulla of females

F1 GENERATION:

- Mortality: at 0 ppm one female, 25 ppm one male and 2500 ppm one male and 5 females

- Body weight: at 2500 ppm decreased in males and females - Food consumption: at 2500 ppm increased in males and

females

- Clinical signs: at 2500 ppm thin or emaciated and/or weak appearance, chromorrhinorrhoea and urine stained fur among males and thin appearance among females

- Mating and fertility data (males/females): no differences between the dose groups; no of mated/pregnant femles 35/28, 36/29, 37/27 and 39/35 at 0, 50, 500 and 2500 ppm resp.

- Maternal delivery data: at 2500 ppm decreased duration of gestation; no effects on implantation sites and number of surviving dams/pups

- Pup data: no differences between the dose groups considering sex ratio; at 500 and 2500 decreased viability on days 1 and 4 post-partum

- Macroscopic examinations: at 2500 ppm kidney lesions consisting of dilated renal pelvis in males and white/brown raised areas in females and gastric lesions (black areas) in females

- Microscopic examinations: at 500 and 2500 ppm kidney lesions characterised by dilation of tubules in the outer medulla in females and an increased incidence of pelvic dilatation in males

F2 GENERATION:

- Clinical signs: at 2500 ppm thin and weak appearance and cannibalism of ears (partially) and tail tip

- Pup effects: at 500 and 2500 ppm one litter died after day 2 or day 5 post-partum, respectively; at 2500 ppm body weight was decreased

- Macroscopic examinations: at 2500 ppm gross kidney lesions consisting of slight/moderate dilation of the kidney pelvis Notox Hertogenbosch

Source Test substance

III, CAS 62476-59-9 (Acifluorfen sodium salt, technical grade), purity not reported

5. Toxicity	ld 62476-59-9 Date 26.12.2001
Conclusion	 NO(A)EL (parental): 25 ppm, based on an increased incidence of kidney lesions (dilated tubules in the outer medulla) in the 500 and 2500 ppm group. Additional findings in the 2500 ppm group consisted of decreased body weight NO(A)EL (developmental): 500 ppm, based on reduced pup body weights and an increased incidence of kidney pelvic dilatation
Reliability 21.05.2001	: (1) valid without restriction (1)
.9 DEVELOPMENTA	L TOXICITY/TERATOGENICITY
Species Sex Strain Route of admin. Exposure period Frequency of treatment Duration of test Doses Control group NOAEL Maternalt. NOAEL Teratogen NOAEL Fetotoxicity Method Year GLP Test substance Method	 rat female other: CrI:COBS-CD-(SD)BR gavage gestation days 6-19 daily Caesarean sections on gestation day 20 20, 90 and 180 mg/kg yes, concurrent vehicle = 20 mg/kg bw = 180 mg/kg bw = 180 mg/kg bw other: EPA; Hazard Evaluation: Humans and Domestic Animals, Federal Register. Part II, Vol. 43, no. 163.83-3 1978 yes other: TS TEST ORGANISMS Age: females 12 weeks (at start mating procedures) Weight at study initiation: 211-255g (gestation day 0) Number of animals: 25 (treatment/control groups) Source: Charles River, Breeding Laboratories, Inc. ADMINISTRATION / EXPOSURE Test duration: 20 days Exposure period: gestation days 6-19 Route of administration: roal gavage Doses: 0, 20, 90 and 180 mg/kg Total volume applied: 10 ml/kg Vehicle: water (reverse osmosis) MATING PROCEDURES: Mating: 1 female / 1 male Day 0 of gestation: gestation days 0 and 20 and several times per day on gestation days 6-19 Body weight gai: gestation: gestation days 0 and 20 and several times per day on gestation days 6-19 Body weight gai: gestation color plug PARAMETERS ASSESSED DURING STUDY: Mortality/clinical observations: gestation days 0 and 20 and several times per day on gestation days 6-19 Body weight gai: gestation days 6-19 Body weight gai: gestation days 6-19 Pod consumption: not measured Maternal reproduction parameters (general): Number of pregnancies and corpora lutea

5. Toxicity	ld 62476-59-9 Date 26.12.2001
	 Examination of uterine content: number and distribution of implantations, early and late resorptions and live and dead foetuses Examination of fetuses: sex; weight; external, visceral (1/3) and skeletal (2/3 foetuses) findings
	ORGANS EXAMINED AT NECROPSY (MACROSCOPIC AND MICROSCOPIC): - Macroscopy: all females - Microscopy: gross lesions (preliminary deaths) preserved for possible histopathology
	ANALYSES: - Method: HPLC - Sampling time: weekly samples taken for possible analysis
Result	 STATISTICAL METHODS: Bartlett's test, Analysis of Variance, Analysis of Covariance, aproximate test of equality of means, Dunnett's test, Kruskal-Wallis, Dunn's method of multiple comparisons ANALYSES: Actual dose level: not reported Stability: Not reported
	MATERNAL TOXIC EFFECTS BY DOSE LEVEL:
	 Mortality and day of death: at 180 mg/kg 3 females died on gestation days 10 or 17 Body weight: at 180 mg/kg decreased during treatment (gestation days 9-19) and overall gestation days 6-19 and 0-20 Clinical signs: females showed at 90 mg/kg excessive salivation and at 180 mg/kg excessive salivation, vocalization, hyperactivity, impaired/lost righting reflex, decreased motor activity, chromodacryorrhoea, rales, urine stained abdominal fur and chromorrhinorrhoea Number pregnant per dose level: at 0, 20, 90 and 180 mg/kg, 22, 21, 19, 24, respectively Number of resorptions (early/late): at 0, 20, 90 and 180 mg/kg, 0.95 (7.3%), 0.90 (6.6%), 1.42 (10.4%) and 2.20 (16.2%), respectively (percent of implantation sites) Number of implantations: at 0, 20, 90 and 180 mg/kg, 13.1, 13.6, 13.7 and 13.6, respectively Number of corpora lutea: at 0, 20, 90 and 180 mg/kg, 14.7, 14.7, 15.4 and 14.6, respectively Duration of Pregnancy: scheduled sacrifice on gestation day 20 Gross pathology incidence and severity: no findings in surviving females. In 2 out of 3 females found dead (180 mg/kg) erosions in the mucosa of the stomach or haemorrhagic lungs were noted
	FETAL DATA:
	There were no gross external, soft tissue or skeletal alterations that were considered effects of the test

5. Toxicity	ld 62476-59-9	
	Date 26.12.2001	
substance. Variations noted in soft tissue examinations and in skeletal ossification were correlated with lower foetal body weights		
	- Litter weights (gravid uterus): not recorded	
	- Number viable: at 0, 20, 90 and 180 mg/kg, 12.2, 12.7,	
	12.3 and 11.4, respectively - Sex ratio (percentage of males): at 0, 20, 90 and 180	
	mg/kg, 51.1%, 54.3%, 48.1% and 46.9%, respectively	
	- Body weight (gain): at 0, 20, 90 and 180 mg/kg, for males	
	3.8g, 3.87g, 3.5g and 3.09g, respectively and for females	
	3.62g, 3.64g, 3.30g and 2.97g, respectively Grossly visible abnormalities: no findings associated with	
	treatment	
	- Visceral abnormalities: at 90 and 180 mg/kg increased	
	incidence of slight dilation of the lateral ventricles of the brain	
	- Skeletal abnormalities: at 90 and 180 mg/kg delayed	
	ossification of metacarpals, forepaw phalanges and hindpaw	
	phalanges and additionally in 180 mg/kg group litters	
	delayed ossification of the caudal vertebrae, sternebrae and metatarsals	
Source	: Notox Hertogenbosch	
Test substance	: III, CAS 62476-59-9, purity 91.2%	
Conclusion	 NOAEL (maternal): 20 mg/kg, based on decreased body weights, clinical signs such as excessive salivation in the 90 and 	
	180 mg/kg groups and mortality and clinical signs including	
	vocalization, huperactivity, impaired righting reflezx,	
	decreased motor activity, chromodacryorrhoea, rales, urine	
	stained abdominal fur, chromorrhinorrhoea in the 180 mg/kg group	
	NOAEL (teratogenicity): 180 mg/kg	
Poliobility	NOAEL (foetotoxicity): 180 mg/kg	
Reliability 21.05.2001	: (1) valid without restriction	(
21.00.2001		``
Species	: rabbit	
Sex Strain	: female : New Zealand white	
Route of admin.	: gavage	
Exposure period	: gestation days 6-29	
Frequency of treatment	: Once daily	
Duration of test	: Caesarean sections on gestation day 30	
Doses	: 3, 12 and 36 mg/kg	
Control group	: yes, concurrent vehicle	
NOAEL Maternalt. NOAEL Teratogen	: = 12 mg/kg bw : = 36 mg/kg bw	
NOAEL Fetotoxicity	= 30 mg/kg bw = 12 mg/kg bw	
Method	: other: EPA, federal register, 1978, Part II, Vol. 43, No. 163, 163.83-3	
Year GLP	: 1978 . ves	
Test substance	: yes : other TS	
Method	: TEST ORGANISMS	
	- Age: females (at insemination) 26 weeks - Weight at study initiation: 3.06-5.13 kg	

5. Toxicity		62476-59-9 26.12.2001
	- Source: Dutchland Laboratories Inc., Denver Pennsylva USA	nia,
	ADMINISTRATION / EXPOSURE - Test duration: 309 days - Exposure period: gestation days 6-29 - Route of administration: oral gavage - Doses: 0, 3, 12 and 36 mg/kg/day - Vehicle: water (revers osmosis) - Dose volume: 10 mg/kg/day	
	MATING PROCEDURES: - Artificial insemination: Semen collected from 4 proven donor bucks of the same strain and source as the female hours before insemination females were intravenously injected with 20 USP units/kg of Human Chorionic Gonadotropin. Insemination of 0.25 mL of diluted (with saline) semen sample (6.0 million spermatozoa/0.25 mL - Day 0 of gestation: day of insemination	
	 PARAMETERS ASSESSED DURING STUDY: Mortality: several times/day during treatment (gestation days 6-29) and on gestation day 30 Clinical observations: On gestation day 0 and several times/day during treatment (gestation days 6-29) and on gestation day 30 Body weight gain: once daily on gestation days 0 and 6- Food consumption: once daily on gestation days 0 and 6- Examination of uterine content: number of corpora lutea number and distribution of implantations, early and late resorptions and live and dead foetuses Examination of fetuses: sex; weight; external, visceral (all foetuses) and skeletal (all foetuses) findings; brains being subjected to a variation of Staple's technique 	6-30
	ORGANS EXAMINED AT NECROPSY (MACROSCOPIC MICROSCOPIC): - Macroscopy: findings all dams recorded, all gross lesion (except commonly found parovarian cysts) were fixed for possible histopathology - Microscopy: not performed	IS
	ANALYSES: - Method: not indicated (analysis separately by the spons - Sampling time: weekly samples taken	or)
Result :	STATISTICAL METHODS: Bartlett's Test, Kruskal-Wallis and Fisher's Exact Test ANALYSES:	Test
	Data on the accuracy and stability of preparations were k on file with the sponsor	ept
	 Actual dose levels: reported as being correct Stability: no results presented Homogeneity: not determined (solutions) 	

5. Toxicity		62476-59-9 26.12.2001
	MATERNAL TOXIC EFFECTS BY DOSE LEVEL:	
	 Mortality and day of death: at 0 mg/kg three females die or were sacrificed (2 following an intubation error, 1 following abortion), at 3 mg/kg one female was sacrificed because of a back injury and at 12 mg/kg one female was sacrificed following abortion Body weight: at 36 mg/kg slightly inhibited body weight gain on gestation days 6-18 and overall inhibition of body weight gain during gestation days 6-30 Food consumption: at 36 mg/kg marked inhibition of foo consumption during gestation days 23-24. Recovery of foc consumption during gestation days 23-24. Recovery of foc consumption during gestation days 29-30 Clinical signs: no treatment-related signs Number pregnant per dose level: 13 (81.2% of number inseminated), 13 (81.2%), 12 (75.0%) and 11 (68.8%) in 10, 3, 12 and 36 mg/kg group, respectively Number aborting: at 0 mg/kg one female and at 12 mg/k female Natural deliveries: at 0, 3, 12 and 36 mg/kg, 1, 2, 2 and 2, respectively Number of resorptions (early/late): at 0, 3, 12 and 36 mg/kg, 0.6, 0.4, 0.7 and 0.7, respectively Number of implantations: at 0, 3, 12 and 36 mg/kg, 6.8, 7.2, 7.3 and 9.0, respectively Post implantation loss: not calculated Number of corpora lutea: at 0, 3, 12 and 36 mg/kg, 9.3, 9.7, 10.7 and 11.1, respectively Duration of Pregnancy: scheduled sacrifice on gestation day 30 Gross pathology incidence and severity: at 36 mg/kg, increased incidence of involuted ovaries combined with congested uterus in 4 females 	s d ood the (g one
	FETAL DATA:	
	There were no gross external, soft tissue or skeletal alterations that were considered effects of the test substance.	
	 Litter size: 0, 3, 12 and 36 mg/kg, 6.2, 6.8, 6.7 and 8.3, respectively Number viable: at 0, 3, 12 and 36 mg/kg, 6.2, 6.8, 6.7 and 8.3, respectively Sex ratio (percentage of males): at 0, 3, 12 or 36 mg/kg 50.0%, 51.5%, 55.9% and 48.0%, respectively Body weight: at 0, 3, 12 and 36 mg/kg, 51.3g, 47.4g, 53 and 43.1g, respectively Grossly visible abnormalities: no treatment related findings Visceral abnormalities: incidental findings comprised accessory spleen, agenesis of the gall bladder and malformation of the diaphragm with atelectasis Skeletal abnormalities: incidentally observed findings consisted of rudimentary rib (between R5-6), fused rib (L6-7), 1 or more fused sternebrae, 1-4 asymmetric sternebrae, stubbed tail and split xiphoid vertebral 	

5. Toxicity	Id 62476-59-9)
-	Date 26.12.2001	
Source	: Notox Hertogenbosch	
Test substance	: III, CAS 62476-59-9, Concentration 240 mg/ml in water (activity 22.4%), purity 81.2%	
Conclusion	 NOAEL (maternal): 12 mg/kg, based on slight inhibition of body weight gain and marked inhibition of food consumption NOAEL (teratogenicity): 36 mg/kg NOAEL (foetotoxicity): 12 mg/kg, based on possible interference with implantations and slight decrease of foetal body weights 	
	There were no differences noted among the dose groups in the number of corpora lutea, implantations, litter sizes, early and late resorptions, foetal sex ratio, number of resorbed conceptuses and number of does with any resorptions. The increased number of involuted corpora lutea and congested mucosa in the uteri may be attributed to interference of the test substance with implantation after fertilization (nidation of fertilized eggs in rabbits approximately gestation day 8)	
Reliability	: (2) valid with restrictions Only 9-10 litters per dose group evaluated	
21.05.2001		(2)

5.10 OTHER RELEVANT INFORMATION

5.11 EXPERIENCE WITH HUMAN EXPOSURE

6. Ref	erences Id 62476-59-9 Date 26.12.2001
(1)	Argus Research laboraties, Inc., Reproductive Effects of Tackle Administered orally in Feed to Crl:COBS-CD-(SD)BR Rats for Two Generations, 1986
(2)	Argus Research Laboratories, Inc, Teratogenic potential of TACU 06238001 in New Zealand White Rabbits (segment II Evaluation), 1980 (76)
(3)	Argus Research Laboratories, Inc., Teratogenicity Study of TACU 06238001 in Pregnant Rats, 1981
(4)	BASF, Acifluorfen-sodium - determination of vapor pressure (1990) (84)
(5)	BASF, Determination of acifluorfen sodium solubility in water and organic solvents (1991) (83)
(6)	BASF, Determination of aciflurofen sodium octanol/water partition coefficient (1991) (82)
(7)	BASF, Phase 3 Summary of Accession #095735 A Hydrolysis Study with 14C-RH-6201: Technical Report #3423-75-66 (1990) (86)
(8)	EFED Ecological Risk Assessment for sodium acifluorfen. US EPA, Registration Process Documents, June 2000. http://www.epa.gov/pesticides/reregistration/acifluorfen/efedchapter.pdf
(9)	EFED Ecological Risk Assessment for sodium acifluorfen. US EPA, Registration Process Documents, June 2000. http://www.epa.gov/pesticides/reregistration/acifluorfen/efedchapter.pdf p 71
(10)	EPIWIN v3.05, Syracuse Research Corporation, Syracuse, NY (July 12, 2000)
(11)	Food and Drug Research Laboratories, Subchronic 21-day dermal toxicity study in rabbits, 1981
(12)	Gulf South Research Institute, Evaluation of ninety day subchronic toxicity to 'Tackle' in Fischer 344 rats, 1981
(13)	Mobil Environmental and Health Science Department, Anaphase analysis of CHO cells treated in vitro with Tackle 2S, 1981.
(14)	Mobil Oil Corporation, Acute toxicity of 10318001 to rainbow trout (Salmo gairdneri), 1981 (79)
(15)	Mobil Oil Corporation, Acute toxicity of 10318001 to the bluegill (Lepomis macrochirus), 1981 (78)
(16)	Mobil Oil Corporation, Acute toxicity of 10318001 to the water flea (Daphnia magna), 1981 (77)
(17)	Rhone Poulenc, Acute toxicity of Tackle 2AS formulation to the earthworm Eisenia fetida, 1990 (88)

6. Refere	nces	ld	62476-59-9
		Date	26.12.2001
(18)	Rhone-Poulenc Ag Company, Acifluorfen-sodium - determination of melting point (1990) (81)		
(19)	Rohm and Haas Company, BLAZER herbicide in vivo cytogenetic study in mice, 1987 (69)		
(20)	Rohm and Haas, Research Division, Single dermal dose with (experimental) Herbicide RH 6201, Aqueous technical, 39.6% a.i., 1976 (67)		
(21)	Rohm and Haas, Research Division, Single oral dose (range-finding dogs) with (experimental) Herbicide RH 6201, Aqueous technical, 39.6% a.i., 1976 (67)		
(22)	Rohm and Haas, Research Division, Single oral dose (Beagle dogs) with (experimental) Herbicide RH 6201, Aqueous technical, 39.6% a.i., 1976 (67)		
(23)	Rohm and Haas, Research Division, Single oral dose (rabbits) with (experimental) Herbicide RH 6201, Aqueous technical, 39.6% a.i., 1976 (67)		
(24)	Rohm and Haas, Research Division, Single oral dose with (experimental) Herbicide RH 6201, Aqueous technical, 39.6% a.i., 1976 (67)		
(25)	Toxigenics, Inc., Four-hour acute aerosol inhalation toxicity study in rats of Tackle 2AS Herbicide, 1980 (68)		

7. Risk Assessment		ld	62476-59-9
		Date	26.12.2001
7.1	END POINT SUMMARY		
7.2	HAZARD SUMMARY		
7.3	RISK ASSESSMENT		