

201-15133B

Robust Summaries for

C.I. Acid Yellow 23

CAS No. 1934-21-0

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Consortium Registration Number

**Submitted to the EPA under the HPV Challenge Program by:
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Table of Contents

1	CHEMICAL AND PHYSICAL PROPERTIES	1
1.1	MELTING POINT	1
1.2	BOILING POINT	2
1.3	VAPOR PRESSURE	3
1.4	N-OCTANOL/WATER PARTITION COEFFICIENTS.....	3
1.5	WATER SOLUBILITY.....	4
2	ENVIRONMENTAL FATE AND PATHWAYS.....	5
2.1	PHOTODEGRADATION.....	5
2.2	BIODEGRADATION.....	7
2.3	FUGACITY	9
3	ECOTOXICITY.....	13
3.1	ACUTE TOXICITY TO FISH	13
3.2	ACUTE TOXICITY TO AQUATIC INVERTEBRATES.....	17
3.3	ACUTE TOXICITY TO AQUATIC PLANTS	19
4	HUMAN HEALTH TOXICITY.....	22
4.1	ACUTE TOXICITY	22
4.2	GENETIC TOXICITY	24
4.2.1	<i>In vitro Genotoxicity</i>	24
4.2.2	<i>In vivo Genotoxicity</i>	30
4.3	REPEATED DOSE TOXICITY	33
4.4	DEVELOPMENTAL TOXICITY	37
4.5	REPRODUCTIVE TOXICITY.....	39

Robust Summaries

for C.I. Acid Yellow 23

The evaluation of the quality of the following data uses a systematic approach described by Klimisch [Klimisch *et al.*, 1996]. Based on criteria relating to international testing standards for categorizing data reliability, four reliability categories have been established. The following categories are:

- Reliability code 1. Reliable without restrictions
- Reliability code 2. Reliable with restrictions
- Reliability code 3. Not reliable
- Reliability code 4. Not assignable

1 CHEMICAL AND PHYSICAL PROPERTIES

1.1 MELTING POINT

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
-----------------------	---------------------

Remarks for substance FD&C Yellow 5

Method/guideline Calculated

GLP

Year

Remarks for Test Conditions

Melting Point 350 °C

Decomposition

Sublimation

Remarks for Results

Conclusion Remarks

Remarks for General Remarks

Data Qualities Reliabilities	Reliability code 4. Not assignable.
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Remarks for Data Reliability	Code 4. Calculated.
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References	MPBPVPWIN EPI Suite (2000) US Environmental Protection Agency.
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1.2 BOILING POINT

CAS Numerical	1934-21-0
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Substance Name	C.I. Acid Yellow 23
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Remarks for Substance	FD&C Yellow 5
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Method/guideline	Calculated
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GLP

Year

Remarks for Test Conditions

Boiling Point	870 °C
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Pressure

Pressure Unit

Decomposition

Remarks for Results

Conclusion Remarks

Remarks for General Remarks

Data Qualities Reliabilities	Reliability code 4. Not assignable.
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Remarks for Data Reliability	Code 4. Calculated.
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References	MPBPVPWIN EPI Suite (2000) US Environmental Protection Agency.
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1.3 VAPOR PRESSURE

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for substance	FD&C Yellow 5
Method/guideline	Calculated/Mean of Antoine & Grain
GLP	No
Year	
Remarks for Test Conditions	
Vapor Pressure	7.43 X 10 ⁻²² mm Hg
Temperature	25 °C
Decomposition	
Remarks for Results	
Conclusion Remarks	
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	MPBPVPWIN EPI Suite (2000) US Environmental Protection Agency.

1.4 N-OCTANOL/WATER PARTITION COEFFICIENTS

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for substance	FD&C Yellow No. 5
Method/guideline	Calculated
GLP	
Year	
Remarks for Test Conditions	
Log Pow	-10.17

Temperature**Remarks for Results****Conclusion Remarks****Data Qualities Reliabilities** Reliability code 4. Not assignable.**Remarks for Data Reliability** Code 4. Calculated.**References** KOWWIN EPI Suite (2000) US Environmental Protection Agency.

1.5 WATER SOLUBILITY

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
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Remarks for Substance Purity not given**Method/guideline** Not given**GLP** Ambiguous**Year** 1991**Remarks for Test Conditions** Not given**Value (mg/L) at temperature** 38,000 mg/ml at 2 °C; 200,000 mg/ml at 25 °C; 200,000 mg/ml at 60 °C**Description of Solubility** Not given**pH value and concentration at temp****pKa value at 25 Celsius****Remarks for Results****Conclusion Remarks****Data Qualities Reliabilities** Reliability code 4. Not assignable.**Remarks for Data Reliability** Code 4. Only secondary literature (review, tables, books, etc.).**References** Marmion D.M. (1991) Handbook of U.S. Colorants: Foods, Drugs, and Cosmetics and Medical Devices. 3rd Ed. New York, John Wiley & Sons, Inc.

2 ENVIRONMENTAL FATE AND PATHWAYS

2.1 PHOTODEGRADATION

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
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Remarks for Substance Data are for structurally related substance 2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulphophenyl)azo]-, disodium salt (FD&C Red 40)

Method/guideline Not given

Test Type Experimental

GLP Ambiguous

Year 1991

Light Source 15-watt General Electric germicidal lamps

Light Spectrum (nm) Ultraviolet

Relative Intensity

Spectrum of Substance

Remarks for Test Conditions The concentration of the dye solution was measured before and after the photolysis using the Hewlett-Packard 8452A diode-array UV/Visible Spectrophotometer. Red 40 was prepared in an initial concentration of 5 mg/l. In the first part of the study, photolysis experiments were conducted using two 15-W (30 Watts total) General Electric germicidal lamps as the ultraviolet light source. The distance between the light source and the reaction vessels was approximately 2.5 cm. Both direct photolysis and indirect photolysis experiments were conducted. The indirect photolysis experiment used acetone as the sensitizer for indirect photodegradation.

Concentration of Substance 5 mg/L

Temperature

Direct photolysis 7% degradation after 50 minutes

Half-life $t_{1/2}$

Degradation % after

Quantum yield

Indirect photolysis 99% degradation after 20 minutes

Sensitizer acetone

Concentration of sensitizer 5 mg/L

Rate constant

Degradation %after

Breakdown products

Remarks field for results

Conclusion remarks

Data Qualities Reliabilities Reliability code 2. Reliable with restriction.

Remarks for Data Reliability Code 2. Basic data given: comparable to guidelines/standards.

References Pasin B. and Rickabaugh J. (1991) Destruction of Azo Dyes by Sensitized Photolysis. Hazard. Ind. Wastes, 359-367.

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
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Remarks for Substance FD&C Yellow 5

Method/guideline Calculation

Test Type AOPWIN

GLP

Year

Light Source

Light Spectrum (nm)

Relative Intensity

Spectrum of Substance

Remarks for Test Conditions

Concentration of Substance

Temperature

Direct photolysis

Half-life t_{1/2} 3.5 hours

Degradation % after

Quantum yield

Indirect photolysis

Sensitizer**Concentration of sensitizer****Rate constant****Degradation %after****Breakdown products****Remarks field for results****Conclusion remarks****Data Qualities Reliabilities** Reliability code 4. Not assignable.**Remarks for Data Reliability** Code 4. Calculated.**References** AOPWIN EPI Suite (2000) US Environmental Protection Agency.**2.2 BIODEGRADATION****CAS Numerical** 1934-21-0

Substance Name	C.I. Acid Yellow 23
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Remarks for Substance Data are for structurally related sulfonic acid C.I. Acid Red No. 14.**Method** Not given**Test Type****GLP** Ambiguous**Year** 1993**Contact time (units)** 24 hour**Innoculum** Activated sludge**Remarks for Test Conditions** Screened raw wastewater was used as the influent in three pilot scale activated sludge biological treatment systems. Each water soluble dye was tested at doses of 1 mg/L for low spike systems and 5 mg/L for high spike systems of influent flow. Before the data collection, dye analytical recovery studies were conducted by dosing the purified dye compound into organic free water, influent wastewater, and mixed liquor. These studies were run in duplicate and each recovery study was

repeated at least once to ensure that the dye compound could be extracted. Purified dye standards were analytically prepared from the commercial dye product by repeated recrystallization.

The INF, primary effluent (PE), and ASE were filtered and the filtrate was passed through a column packed with resin. The filter paper and resin were soaked in an ammonia acetonitrile solution and then Soxhlet extracted with ammonia-acetonitrile. The extract was concentrated and brought up to 50 mL volume with a methanol/dimethylformamide solution. The mixed liquor samples were separated into two components, the filtrate or soluble fraction (SOL) and the residue (RES) fraction. The SOL fraction was processed similar to these samples but the resin adsorption step was omitted. All extracted samples were analyzed by HPLC with an ultraviolet-visible detector. Total suspended solids analyses were also performed on the INF, PE, ML, and ASE samples.

All systems were operated for at least three times the solids retention time to ensure acclimation prior to initiation of data collection. All samples were 24 hour composites made up of 6 grab samples collected every 4 hours and stored at 4 °C.

Results

Percent recovery as measured: Organic Free Water: 101% at 1 mg/L and 90% at 5 mg/L; Wastewater: 98% at 1mg/L and 97% at 5 mg/L; Mixed Liquor: 88% at 1mg/L and 92% at 5 mg/L
Mass Balance Data Summary: Low spike: 116% recovered, 1% adsorbed; High spike: 148% recovered, less than 1% adsorbed.

Classification

Remarks fields for results

Since the majority of the test substance was recovered, the authors assumed that this compound was not biodegraded. The authors based this assumption on preliminary data indicating little or no problems in recovering the compounds from the sample matrix. Additionally the results also indicate that the material was not adsorbed. The authors attributed the high sulfonic acid substitution on the test substance as the reason why the material was not removed by the microbial cells or cell byproducts and subject to aerobic biodegradation.

Conclusion remarks

Data Qualities Reliabilities

Reliability code 1. Reliable without restriction.

Remarks for Data Reliability

Code 1. Comparable to guideline study.

References

Shaul G.M., Holdsworth T.J., Dempsey C.R., and Dostal K.A. (1990) Fate of water soluble azo dyes in the activated sludge process. *Chemosphere* 22, p107-119.

CAS Numerical

1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	FD&C Yellow 5

Method	
Test Type	Calculated
GLP	
Year	
Contact time (units)	
Innoculum	
Remarks for Test Conditions	
Results	
Classification	Not readily biodegradable
Remarks fields for results	
Conclusion remarks	
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	BIOWIN EPI Suite (2000) US Environmental Protection Agency.

2.3 FUGACITY

CAS Numerical	1934-21-0
Substance Name	C.I. Acid Yellow 23
Remarks for Substance	FD&C Yellow No. 5
Model Conditions	25 C, 100,000 lbs.
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used (title, version, date)	EQC V 2.70 Level III
Input parameters	MW, log Kow, water solubility, MP & VP

Year

Remarks for Test Conditions

Media Air

absorption coefficient

Desorption

Volatility

Model data and results

Estimated Distribution and Media Concentration 3.05E-13%
Remarks

Conclusion remarks

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References ECOSAR EPI Suite (2000) US Environmental Protection Agency. Level III. Fugacity.

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
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Remarks for Substance	FD&C Yellow No. 5
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Model Conditions 25 C, 100,000 lbs.

Test Type Environmental Equilibrium Partitioning Model

Method Mackay

Model Used (title, version, date) EQC V 2.70 Level III

Input parameters MW, log Kow, water solubility, MP & VP

Year

Remarks for Test Conditions

Media Water

absorption coefficient

Desorption

Volatility

Model data and results

Estimated Distribution and Media Concentration Remarks	51.8%
Conclusion remarks	
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	ECOSAR EPI Suite (2000) US Environmental Protection Agency. Level III. Fugacity.

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
-----------------------	---------------------

Remarks for Substance	FD&C Yellow No. 5
Model Conditions	25 C, 100,000 lbs.
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used (title, version, date)	EQC V 2.70 Level III
Input parameters	MW, log Kow, water solubility, MP & VP
Year	

Remarks for Test Conditions

Media Soil

absorption coefficient

Desorption

Volatility

Model data and results

Estimated Distribution and Media Concentration Remarks	48.1%
Conclusion remarks	
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	ECOSAR EPI Suite (2000) US Environmental Protection Agency. Level III. Fugacity.

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
-----------------------	---------------------

Remarks for Substance FD&C Yellow No. 5

Model Conditions 25 C, 100,000 lbs.

Test Type Environmental Equilibrium Partitioning Model

Method Mackay

Model Used (title, version, date) EQC V 2.70 Level III

Input parameters MW, log Kow, water solubility, MP & VP

Year

Remarks for Test Conditions

Media Sediment

absorption coefficient

Desorption

Volatility

Model data and results

Estimated Distribution and Media Concentration 0.0981%
Remarks

Conclusion remarks

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References ECOSAR EPI Suite (2000) US Environmental Protection Agency. Level III. Fugacity.

3 ECOTOXICITY

3.1 ACUTE TOXICITY TO FISH

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	Data are for sulfonic acid derivative 2,2'-(1,2-ethene-diyl)bis(5-amino)-benzenesulfonic acid
Method/guideline	
Test Type	Experimental
GLP	Ambiguous
Year	Not given
Species/Strain/Supplier	Fish
Analytical monitoring	
Exposure period (unit)	48 hour
Remarks for Test Conditions	
Observations on precipitation	
Nominal concentrations as mg/L	
Measured concentrations as mg/L	
Unit	
Endpoint value	LC50 = 200 mg/L
Reference substances (if used)	
Remarks fields for results	
Conclusion remarks	
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Only secondary literature (review, tables, books, etc.).
References	Alstoffe, (1992) Daten zur Beurteilung der Wirkung auf Mensch and Umwelt-Satensätze, Verband der Chemischen Industrie, Frankfurt 1992. Schön N. (1991) Altsoff-Grunddatensätze-Liste der bisher publizierten Grunddatensätze UWSF-Z. Umwelchem. Ökotox, 3(3), 183-185.

Schön N. (1992) Altsoff-Grunddatensätze-Liste der bisher publizierten Grunddatensätze UWSF-Z. Umwelchem. Ökotox, 4(6), 343-345.

CAS Numerical	1934-21-0
Substance Name	C.I. Acid Yellow 23
Remarks for Substance	Data are for sulfonic acid derivative 2,2'-(1,2-ethene-diyl)bis(5-amino)-benzenesulfonic acid, disodium salt
Method/guideline	
Test Type	Experimental
GLP	Ambiguous
Year	Not given
Species/Strain/Supplier	Fish
Analytical monitoring	
Exposure period (unit)	72 hour
Remarks for Test Conditions	
Observations on precipitation	
Nominal concentrations as mg/L	
Measured concentrations as mg/L	
Unit	
Endpoint value	LC50 greater than 1000 mg/L
Reference substances (if used)	
Remarks fields for results	
Conclusion remarks	
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4.Only secondary literature (review, tables, books, etc.).
References	<p>Alstoffe, (1992) Daten zur Beurteilung der Wirkung auf Mensch and Umwelt-Satensätze, Verband der Chemischen Industrie, Frankfurt 1992.</p> <p>Schön N. (1991) Altsoff-Grunddatensätze-Liste der bisher publizierten Grunddatensätze UWSF-Z. Umwelchem. Ökotox, 3(3), 183-185.</p> <p>Schön N. (1992) Altsoff-Grunddatensätze-Liste der bisher publizierten Grunddatensätze UWSF-Z. Umwelchem. Ökotox, 4(6), 343-345.</p>

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	Data are for sulfonic acid derivative 2,2'-(1,2-ethene-diyl)bis(5-amino)-benzenesulfonic acid, dipotassium salt
Method/guideline	
Test Type	Experimental
GLP	Ambiguous
Year	Not given
Species/Strain/Supplier	Fish
Analytical monitoring	
Exposure period (unit)	96 hour
Remarks for Test Conditions	
Observations on precipitation	
Nominal concentrations as mg/L	
Measured concentrations as mg/L	
Unit	
Endpoint value	LC50 greater than 10,000 mg/L
Reference substances (if used)	
Remarks fields for results	
Conclusion remarks	
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4.Only secondary literature (review, tables, books, etc.).
References	Alstoffe, (1992) Daten zur Beurteilung der Wirkung auf Mensch and Umwelt-Satensätze, Verband der Chemischen Industrie, Frankfurt 1992. Schön N. (1991) Altsoff-Grunddatensätze-Liste der bisher publizierten Grunddatensätze UWSF-Z. Umwelchem. Ökotox, 3(3), 183-185. Schön N. (1992) Altsoff-Grunddatensätze-Liste der bisher publizierten Grunddatensätze UWSF-Z. Umwelchem. Ökotox, 4(6), 343-345.

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
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Remarks for Substance FD&C Yellow 5

Method/guideline ECOSAR

Test Type Calculated

GLP

Year

Species/Strain/Supplier

Analytical monitoring

Exposure period (unit) 96 hour

Remarks for Test Conditions Input parameters: Molecular weight, Water solubility, 200,000 mg/L at 25 °C

Observations on precipitation

Nominal concentrations as mg/L

Measured concentrations as mg/L

Unit

Endpoint value LC50 = 1.14 E+14 mg/L

Reference substances (if used)

Remarks fields for results

Conclusion remarks

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

3.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	Data are for sulfonic acid derivative 2,2'-(1,2-ethene-diyl)bis(5-amino)-benzenesulfonic acid, disodium salt
Method/guideline	
Test Type	Experimental
GLP	
Year	
Analytical procedures	
Species/Strain	<i>Daphnia magna</i>
Test details	24 hour
Remarks for Test Conditions	
Nominal concentrations as mg/L	
Measured concentrations as mg/L	
Unit	
EC50, EL50, LC0, at 24,48 hours	EC50 = 100 mg/L
Biological observations	
Control response satisfactory?	
Appropriate statistical evaluations?	
Remarks fields for results	
Conclusion remarks	
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Only secondary literature (review, tables, books, etc.).
References	<p>Alstoffe, (1992) Daten zur Beurteilung der Wirkung auf Mensch and Umwelt-Satensätze, Verband der Chemischen Industrie, Frankfurt 1992.</p> <p>Schön N. (1991) Altsoff-Grunddatensätze-Liste der bisher publizierten Grunddatensätze UWSF-Z. Umwelchem. Ökotox, 3(3), 183-185.</p> <p>Schön N. (1992) Altsoff-Grunddatensätze-Liste der bisher publizierten Grunddatensätze UWSF-Z. Umwelchem. Ökotox, 4(6), 343-345.</p>

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
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Remarks for Substance FD&C Yellow 5

Method/guideline ECOSAR

Test Type Calculated

GLP

Year

Analytical procedures

Species/Strain *Daphnia magna*

Test details 48 hours

Remarks for Test Conditions Input parameters: Water solubility, 200,000 mg/L at 25 °C;
Molecular weight 556.34

Nominal concentrations as

mg/L

Measured concentrations as

mg/L

Unit

EC50, EL50, LC0, at 24,48 hours EC50 = 5.25 E+13 mg/L

Biological observations

Control response satisfactory?

Appropriate statistical evaluations?

Remarks fields for results

Conclusion remarks

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

3.3 ACUTE TOXICITY TO AQUATIC PLANTS

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	The test substance was an unidentified sulfonic acid substituted azo dye.
Method/guideline	
Test Type	Experimental
GLP	Ambiguous
Year	1996
Species/Strain/Supplier	Green algae, <i>Selenastrum capricornutum</i>
Endpoint basis	
Exposure period (duration)	96 hour
Analytical monitoring	
Remarks for Test Conditions	Algal chronic toxicity test were performed according the method of EPA, 1988. Three replicates were performed for each dye at a nominal concentration of 1 mg/l for the active colorant. One ml of dye stock solution was added to 50 mg/l of algal assay medium in 125 ml Erlenmeyer flasks. <i>S. capricornutum</i> in continuous culture provided the initial inoculum (10,000 algal cells/ml). The cells were incubated in the solution for 96 hours. The diluent and negative control were algal assay medium. AAM was prepared by adding 1 ml from each of five stock solutions to 900 ml of deionized water. After spiking, the total volume was brought to 1 liter with deionized water. Population growth was used to establish potential toxicity. If the dye inhibited algal growth by more than 50% of that of the negative controls, a definitive test using several dilutions of the dye was performed to allow for determination of an EC50 concentration.
Nominal concentrations as mg/L	
Measured concentrations as mg/L	
Unit	
Endpoint value	Average yield: 36.6% with 95% C.I. (34.9-38.4).
NOEC, LOEC or NOEL, LOEL	
Biological observations	26.4% stimulation of population growth compared to control.
Control response satisfactory?	Yes
Appropriate statistical evaluations?	Yes, Dunnett's test
Remarks fields for results	Not statistically significant.

Conclusion remarks

Data Qualities Reliabilities Reliability code 2. Reliable with restriction.

Remarks for Data Reliability Code 1. Comparable to guideline study.

References Greene J. C. and Baughman G.L. (1996) Effects of 46 dyes on population-growth of fresh-water green-alga *selenastrum-capricornutum*. Textile Chemist And Colorist, 28, 23-30.

Green J.D. et al. (1988) Protocols for short term toxicity screening of hazardous waste sites. Report to EPA 600/3-88-029. U.S. Environmental Protection Agency. Corvallis, Oregon.

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
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Remarks for Substance FD&C Yellow 5

Method/guideline ECOSAR

Test Type Calculated

GLP

Year

Species/Strain/Supplier Green algae

Endpoint basis

Exposure period (duration) 96 hour

Analytical monitoring

Remarks for Test Conditions Input parameters: Water solubility - 200,000 mg/L at 25 °C; Molecular weight 556.34

Nominal concentrations as

mg/L

Measured concentrations as

mg/L

Unit

Endpoint value EC50 = 1.63 E+13 mg/L

NOEC, LOEC or NOEL, LOEL

Biological observations

Control response

satisfactory?

Appropriate statistical

evaluations?

Remarks fields for results

Conclusion remarks

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References ECOSAR EPI Suite (2000) US Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

4 HUMAN HEALTH TOXICITY

4.1 ACUTE TOXICITY

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	Not given
Method/guideline	Not given
Test Type	Acute Toxicity LD50
GLP	No
Year	1957
Species/Strain	Rat
Sex	Not reported
# of animals per sex per dose	Not given
Vehicle	Not given
Route of administration	Intraperitoneal
Remarks for test conditions	
Value LD50 or LC50 with confidence limits	2,000 mg/kg bw
Number of deaths at each dose level	
Remarks for results	
Conclusion remarks	
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Only secondary literature (review, tables, books, etc.).
References	Deutsche Forschungsgemeinschaft, Bad Godesberg, Federal Republic of Germany, Farbstoff Kommission (1957) Mitteilung 6.
CAS Numerical	1934-21-0
Substance Name	C.I. Acid Yellow 23
Remarks for Substance	Not given
Method/guideline	Not given

Test Type	Acute Toxicity LD50
GLP	No
Year	1957
Species/Strain	Rat
Sex	Not reported
# of animals per sex per dose	Not given
Vehicle	Not given
Route of administration	Intravenous
Remarks for test conditions	
Value LD50 or LC50 with confidence limits	1,000 mg/kg bw
Number of deaths at each dose level	
Remarks for results	
Conclusion remarks	
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4.Only secondary literature (review, tables, books, etc.).
References	Deutsche Forschungsgemeinschaft, Bad Godesberg, Federal Republic of Germany, Farbstoff Kommission (1957) Mitteilung 6.
CAS Numerical	1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	Not given
Method/guideline	Not given
Test Type	Acute Toxicity LD50
GLP	No
Year	1964
Species/Strain	Mice
Sex	Not reported
# of animals per sex per dose	Not given
Vehicle	1% gum arabic

Route of administration	Oral
Remarks for test conditions	
Value LD50 or LC50 with confidence limits	12,750 mg/kg bw
Number of deaths at each dose level	
Remarks for results	
Conclusion remarks	
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Only secondary literature (review, tables, books, etc.).
References	National Institute of Hygienic Sciences of Japan. Unpublished data submitted to WHO, 1964 cited in ILSI report on FD&C Yellow 5 6/2/83.

4.2 GENETIC TOXICITY

4.2.1 *In vitro* Genotoxicity

CAS Numerical	1934-21-0
Substance Name	C.I. Acid Yellow 23
Remarks for Substance	FD&C Yellow No. 5; Purity not given
Method/guideline	Ames plate incorporation and liquid pre-incubation
Test Type	Reverse mutation
System of Testing	Bacterial
GLP	Ambiguous
Year	1981
Species/Strain	<i>Salmonella typhimurium</i> TA1535, TA 1537, TA1538, TA98, TA100
Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced rats
Doses/concentration levels	0.005- 5.0 mg/plate

Statistical Methods	Not given
Remarks for test conditions	Reverse mutation tests were carried out using <i>S. typhimurium</i> strains TA1535, TA 1537, TA1538, TA98, TA100. Plate incorporation tests were conducted according to Ames <i>et al.</i> , with the Andrews et al. modifications. Duplicates were performed at each of the six concentrations of the test substance. Mutagenic compounds were assayed using duplicate plates. A substance was considered positive when the number of revertants above background was at least twice the value of the historical control mean or twice the value of the current control mean, whichever was greater and a dose response curve could be generated.
Result	<p>Positive controls without metabolic activation were sodium azide (TA1535 and TA100), 9-aminoacridine (TA97 and TA1535), and 4-nitro-o-phenylenediamine (TA98). The positive controls were sodium azide, 9-aminoacridine, 2-nitrofluorene, and 2-aminoanthracene.</p> <p>Negative</p>
Cytotoxic concentration	5.0 mg/plate for plate-incorporation, and .5 mg/ml for pre-incubation test.
Genotoxic effects	Negative
Appropriate statistical evaluations?	None given
Remarks for results	Negative
Conclusion remarks	The test substance was negative in the AMES assay for reverse mutation using <i>Salmonella typhimurium</i> TA1535, TA 1537, TA1538, TA98, TA100.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.
References	Chung K.T., Fulk G.E., & Andrews A.W. (1981) Mutagenicity testing of some commonly used dyes. Applied and Environmental Microbiology 42, 641-648.

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	FD&C Yellow No. 5; Purity not given
Method/guideline	Ames
Test Type	Reverse mutation
System of Testing	Bacterial
GLP	No
Year	1979

Species/Strain	<i>Salmonella typhimurium</i> TA1535, TA 1537, TA98, TA100
Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced rats
Doses/concentration levels	10-250 mg/plate
Statistical Methods	Not given
Remarks for test conditions	The test substance was dissolved in DMSO. The test was considered positive if 2 fold increase in revertants was observed. Positive controls included 9-aminoacridine; 2-aminoflourine; N-methyl-N-nitrosoguanidine.
Result	Negative
Cytotoxic concentration	Not given
Genotoxic effects	Negative
Appropriate statistical evaluations?	None given
Remarks for results	Negative
Conclusion remarks	No evidence of genotoxicity was reported.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Muzzall J.M. and Cook W.I. (1979) Mutagenicity test of dyes used in cosmetics with the Salmonella/mammalian microsome test. Mutations Research 67, 1-8.a.
CAS Numerical	1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	FD&C Yellow No. 5; Purity not given
Method/guideline	Ames
Test Type	Reverse mutation
System of Testing	Bacterial
GLP	Ambiguous
Year	1984
Species/Strain	<i>Salmonella typhimurium</i> TA1535, TA 1537, TA98, TA100, TA92, TA94
Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced rats
Doses/concentration levels	Up to 5.0 mg/ml
Statistical Methods	Not given
Remarks for test conditions	Reverse mutation tests were carried out using <i>S. typhimurium</i> strains TA92, TA1535, TA100, TA1537, TA94 and TA98. Cells

cultured overnight were pre-incubated with the test substance and the S-9 mix for twenty minutes at 37 degrees Celsius prior to plating. Duplicates were performed at each of the six concentrations of the test substance. The number of revertant colonies were counted following incubation for two days. Negative controls were either untreated plates or solvent. Positive results were determined if the number of colonies found was twice the number in the control. If the test was positive and a dose response relationship was not detected, additional experiments at different doses or induced mutation frequency assays were performed.

Result	Negative
Cytotoxic concentration	5.0 mg/ml was the highest non-cytotoxic dose used in the experiment.
Genotoxic effects	Negative
Appropriate statistical evaluations?	None given
Remarks for results	Negative
Conclusion remarks	The test substance was negative in the AMES assay for reverse mutation using <i>Salmonella typhimurium</i> TA1535, TA 1537, TA98, TA100, TA92, TA94.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Ishidate, M., Sofuni, T., Yoshikawa, K., Hauashi, M., Nohmi, T., Sawada, M. and Matsuoka. (1984). Primary Mutagenicity Screening of Food Additives Currently Used in Japan. <i>Fd. Chem. Toxic.</i> 22(8) 623-636.

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	FD&C Yellow No. 5; Purity not given
Method/guideline	Chromosomal aberration test was carried out using a Chinese hamster fibroblast cell line, CHL. The cells were exposed to 3 different doses for 24 and 48 hours. No metabolic activation system was applied.
Test Type	Chromosomal aberration test
System of Testing	Chinese hamster fibroblast cell line CHL.
GLP	Ambiguous
Year	1984
Species/Strain	Chinese hamster fibroblast cell line CHL.
Metabolic Activation	None
Doses/concentration levels	up to 2.5 mg/ml

Statistical Methods	Not available
Remarks for test conditions	<p>Chromosomal aberration tests were carried out using the Chinese hamster fibroblast line. Cells were exposed to the test substance at three different doses for 24 and 48 hour. No metabolic activation was employed. The maximum dose used for each test substance was found in a preliminary test to determine the dose required for 50% cell-growth inhibition. Colcemid at a final concentration of 0.2 ug/ml was added to the culture two hours prior to cell harvesting. The cells were prepared for viewing on slides. One hundred visible metaphases were observed under the microscope and the incidence of polyploid cells and structural chromosomal aberrations (including chromosome and chromatid gaps, breaks, exchanges, ring formations, fragmentations and others) were recorded. Negative controls included untreated cells and solvent treated cells. The incidence of aberrations in the negative controls was generally less than 3.0%. The results were considered negative if less than 4.9%, equivocal if between 5.0-9.9%, and positive if more than 10%. If dose response relationships were not observed, additional experiments were carried out at similar dose levels.</p> <p>The maximum dose for positive results represents the dose at which the maximum effect was obtained.</p> <p>For quantitative evaluation of the clastogenic potential, the D20 was calculated, which is the dose (mg/ml) at which structural aberrations (including gaps) were detected in 20% of the metaphases observed. In addition, the TR value was calculated, which indicates the frequency of cells with exchange-type aberrations per unit dose (mg/ml). These values are relatively high for chemicals that show carcinogenic potential in animals.</p>
Result	<p>The test substance was shown to be positive (23% total incidence of cells with aberrations) in chromosomal aberration test at 48 hours. TR value was 3.5 and D20 = 1.8. Weakly positive at 24 hour (11.0%, total incidence of cells with aberrations) The results were considered positive if the total incidence of cells with aberrations (including gaps) was 10.0% or more. Two percent (2%) reported as polyploid.</p>
Cytotoxic concentration	Not given
Genotoxic effects	Positive
Appropriate statistical evaluations?	None given
Remarks for results	Positive
Conclusion remarks	C.I. Acid Yellow 23 tested positive in the chromosomal aberration test using Chinese hamster fibroblasts.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Acceptable, well-documented publication/study report which meets basic scientific principles.

References Ishidate, M., Sofuni, T., Yoshikawa, K., Hauashi, M., Nohmi, T., Sawada, M. and Matsuoka. (1984) Primary Mutagenicity Screening of Food Additives Currently Used in Japan. *Fd. Chem. Toxic.* 22(8) 623-636.

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
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Remarks for Substance FD&C Yellow No. 5; 94% purity

Method/guideline Williams, 1977

Test Type Unscheduled DNA Synthesis

System of Testing Rat hepatocytes

GLP Ambiguous

Year 1985

Species/Strain Rat/Sprague-Dawley

Metabolic Activation None

Doses/concentration levels 2 X 10⁻³
2 X 10⁻⁴
2 X 10⁻⁵
2 X 10⁻⁶

Statistical Methods None given

Remarks for test conditions Hepatocytes from rats were isolated and cultured according to the two step in situ liver perfusion model (Malansky and Williams, 1982). Viable hepatocytes (2 X 10⁺⁵) were seeded in wells and incubated for 4 hours with [H³]-thymidine (10 uCi/ml) and the test substance (prepared in either DMSO or water) according to a procedure similar to Williams, 1977. Control incubations were conducted with and without DMSO. The authors state that DMSO had no effect on DNA repair. Two experiments were conducted.

DNA repair was quantified by the autoradiographic determination of incorporated [H³]-thymidine. Net nuclear grains (NNG) were determined by counting the number of grains in each nuclei and subtracting the average number of grains present in the three equal size adjacent cytoplasmic areas. Average NNG counts of 5 or more were assumed to constitute a positive response, because these differed from the control response by greater than 2 standard deviations. In the negative controls, NNG counts ranged from -0.6- to -2.8 and from -0.9 to -2.1 for no solvent and 1% DMSO incubations, respectively. The proportion of cells with greater than or equal to 5 NNG was less than or equal to 8.1% for all control incubations. Therefore NNG below zero were considered negative responses. Concentrations of dyes producing 90% or greater detachment of the hepatocytes from the coverslips

were assumed to be toxic and not counted.

Result	The positive control was Solvent Yellow 3 (o-aminoazotoluene). Experiment 1 Conc Avg NNG % >5NNG 2 X 10 ⁻³ -1.7 (+/-2.6) 5 2 X 10 ⁻⁴ -2.4 (+/-3.3) 5 2 X 10 ⁻⁵ -2.4 (+/-3.2) 2 2 X 10 ⁻⁶ -2.0 (+/-2.8) 3 Experiment 2 Conc Avg NNG % greater than 5NNG 2 X 10 ⁻³ -2.2 (+/- Greater than 2 X 10 ⁻³
Cytotoxic concentration	
Genotoxic effects	Negative
Appropriate statistical evaluations?	None given
Remarks for results	Negative
Conclusion remarks	C.I. Acid Yellow 23 did not induce unscheduled DNA synthesis in an <i>in vitro</i> assay using rat hepatocytes isolated from the livers of Sprague-Dawley rats.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Kornbrust D. and Barfknecht T. (1985) Testing Dyes in HPC/DR systems. Environmental Mutagenesis 7, 101-120.

4.2.2 *In vivo* Genotoxicity

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	Data are for structurally related substance FD&C Yellow No. 6
Method/guideline	Rodent Micronucleus Test
Test Type	Rodent Micronucleus
GLP	Ambiguous

Year	1991
Species/Strain	Rat/PVG
Sex	Male
Route of administration	Oral-Gavage
Doses/concentration levels	10 ml/kg bw
Exposure period	Single dose
Remarks for test conditions	Male PVG rats received a single oral dose of 500, or 1000 mg/kg of the test substance. Bone marrow samples were taken at 24 and 48 hours later.
Effect on mitotic index or PCE/NCE ratio by dose level and sex	
Genotoxic effects	No significant increase in the frequency of micronucleated polychromatic erythrocytes at either time point and in either species was reported. Additionally, there was reported increase in the % PE (polychromatic erythrocytes).
NOEL (C)/ LOEL (C)	
Appropriate statistical evaluations?	Yes.
Remarks for results	No effects.
Conclusion remarks	
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Acceptable, well-documented publication/study report which meets basic scientific principles.
References	Westmoreland C. and Gatehouse D.G. (1991) The differential clastogenicity of Solvent Yellow 14 and FD & C Yellow No. 6 in vivo in the rodent micronucleus test (observations on species and tissue specificity). Carcinogenesis 12 (8), 1403-8.
CAS Numerical	1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	94% purity
Method/guideline	Mirsalis and Butterworth, 1980
Test Type	Unscheduled DNA Synthesis
GLP	Ambiguous
Year	1985
Species/Strain	Rat/Sprague Dawley
Sex	Male

Route of administration Oral-Gavage

Doses/concentration levels 500 mg/kg bw

Exposure period 2 hr; 15 hr

Remarks for test conditions Six to eight male Sprague-Dawley rats weighing 200-300 g were administered 500 mg acid yellow 23/kg bw *via* gavage. The control animal was administered corn oil only. Animals were killed at two time points, 2 hours and 15 hours. If negative results were obtained at time point 1 and time point 2, the *in vivo* testing was terminated and considered to be negative. If the initial test at time point 1 yielded a positive response, the test substance was retested at that time point. If another positive response was observed, the test was considered positive. Time points are the time the test substance was administered prior to the start of liver perfusion and isolation of hepatocytes.

Hepatocytes from rats were isolated and cultured according to the two step *in situ* liver perfusion model (Malansky and Williams, 1982). Viable hepatocytes (2×10^5) were seeded in wells and incubated for 4 hours with [H3]-thymidine (10 uCi/ml) and the test substance (prepared in either DMSO or water) according to a procedure similar to Williams, 1977. Control incubations were conducted with and without DMSO. The authors state that DMSO had no effect on DNA repair.

DNA repair was quantified by the autoradiographic determination of incorporated [3H]-thymidine. Net nuclear grains (NNG) were determined by counting the number of grains in each nuclei and subtracting the average number of grains present in the three equal size adjacent cytoplasmic areas. Average NNG counts of 5 or more were assumed to constitute a positive response, because these differed from the control response by greater than 2 standard deviations. In the negative controls, NNG counts ranged from -0.6- to -2.8 and from -0.9 to -2.1 for no solvent and 1% DMSO incubations, respectively. The proportion of cells with greater than or equal to 5 NNG was less than or equal to 8.1% for all control incubations. Therefore NNG below zero were considered negative responses. Concentrations of dyes producing 90% or greater detachment of the hepatocytes from the coverslips were assumed to be toxic and not counted.

Effect on mitotic index or PCE/NCE ratio by dose level and sex The positive control was Solvent Yellow 3 (o-aminoazotoluene). Experiment 1

Dose (mg/kg bw)	Time	Avg NNG	% >5NNG
500	2 hr	-2.6 (+/-3.7)	2
	15 hr	-1.3 (+/-2.6)	2

Genotoxic effects Negative

NOEL (C)/ LOEL (C)	Greater than 500 mg/kg bw
Appropriate statistical evaluations?	None given
Remarks for results	Negative
Conclusion remarks	C.I. Acid Yellow 23 did not induce unscheduled DNA synthesis in an invivo assay using rat hepatocytes isolated from the livers of Sprague Dawley rats.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Kornbrust D. and Barfknecht T. (1985) Testing Dyes in HPC/DR systems. Enviromental Mutagenesis 7, 101-120.

4.3 REPEATED DOSE TOXICITY

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	FD&C Yellow 5; 90% purity; 10% intermediates or volatile matter
Method/guideline	Chronic Toxicity/Carcinogenicity Study
GLP	Yes
Year	1988
Species/Strain	Rat/Charles River CD
Sex	Male and Female
Route of administration	Oral-Diet
Doses/concentration levels	0, 0.1, 1.0, or 2.0% (original study) 0, 5.0% (high dose study)
Exposure period	113 weeks (males) or 114 weeks (females) (original study); 122 weeks (males) or 125 weeks (females) high-dose study
Frequency of treatment	Daily
Control Group	Yes, 2 concurrent controls (original study); 1 concurrent control (high-dose study)
Post exposure observation period	
Remarks for test conditions	In the <i>in utero</i> phase, groups of rats (60/sex/group) were administered 0, 0, 0.1, 1.0 or 2.0% FD & C Yellow No. 5 in the

diet daily for approximately 2 months prior to mating. In the high-dose study, 60/sex/group received 0 or 5% FD&C Yellow 5 for approximately 2 months prior to mating. The 3 control groups received the basal diet only. A maximum of 2 rats/sex/litter were randomly selected for the chronic phase of the study. There were 70/sex/group at the initiation of the chronic phase and these offspring were exposed to the same dietary levels as their parents.

Animals were housed individually and fed the test diet *ad libitum*. Clinical observations were recorded twice daily with at least 5 hours between observations. Detailed physical examinations and palpation for masses were performed weekly. Body weights and food consumption were determined weekly for the first fourteen weeks, bi-weekly for the next 12 weeks and every 4 weeks thereafter until the end of the study. The intake of the test substance was determined from body weight, food consumption and dietary concentration. Hematology tests, including hemoglobin, hematocrit, erythrocyte and total and differential leukocyte counts, and erythrocyte morphology, were conducted on ten randomly selected animals at months 3, 6, 12, 18 and 24 of the study. Necropsies were conducted on all animals dying prior to study termination, killed in a moribund condition or killed on schedule. Histological examinations were conducted on all animals from both control groups, the highest dose group (2.0 or 5.0%) from each study and also on 10 rats randomly selected from each group for an interim sacrifice at 12 months. Histology was also performed on any animal with gross lesions or masses.

Tissues examined included adrenal glands, aorta, blood smear, brain, cecum, colon, duodenum, epididymus or uterus, esophagus, eyes, femur including marrow, tissue masses, gallbladder, heart, ileum, jejunum, duodenum, kidneys, liver, lungs and bronchi, mammary gland, nerves (sciatic), ovaries, lymph nodes, pancreas, parathyroids, pituitary gland, prostate, rectum, skin, spleen, seminal vesicles, skeletal muscle, testes with epididymides, stomach, thymus, thyroid gland including parathyroid, trachea, urinary bladder, uterus.

NOAEL(NOEL)

5.0 % (Males: 2641 mg/kg/d and Females: 3348 mg/kg/day)

LOAEL(LOEL)

Not determined

**Actual dose received by dose level and sex
Toxic response/effects by dose level**

Males: 48, 491, 984 or 2641 mg/kg/day; Females: 58, 589, 1225 or 3348 mg/kg/day
In utero

There were no compound-related effects on fertility, gestation, parturition, lactation, pup survival through weaning or number of live and still-born pups. Slight decreases in body weight (4-5%) and slight increases in food consumption were noted in the F0 rats treated at dietary level of 5.0%. Two F0 female control rats died during the *in utero* phase of the original study and one male and one female from the control and 5.0% group, respectively, died during the *in utero* phases of the high-dose study. There were no compound-related effects on pup

survival.

In the F1 generation, a yellow tint was reported at all intake levels above 0.1%. At the 1.0% dietary level, group mean body weights at termination for both sexes were lower than the control animals, but the difference was only statistically significant for the females. In the high dose study (5.0% dietary level), group mean body weights were significantly lower in both sexes at termination. Food consumption was similar for control and treated animals at the 0.01, 1 or 2% dietary levels, but was slightly higher at the 5% level in the high-dose study, although not statistically significant. Hematological, clinical chemistry and urinalysis parameters did not differ significantly from the controls. Necropsies at one year did not reveal any treatment-related gross or microscopic changes.

At study termination, no treatment-related effects were reported on survival. No treatment-related changes were reported at gross necropsy. Histological evaluation revealed a variety of lesions, including neoplasms, present at similar incidences in control and treated animals. The authors considered the lesions to be spontaneous and not related to administration of the test material.

Yes, F-test, Anova

Appropriate statistical evaluations?

Remarks for results

The decrease in mean body weight at the 5.0% treatment level was not considered toxicologically significant give the non-nutritive character of FD & C Yellow No. 5.

Conclusion remarks

The NOAEL of 5.0% providing an average daily intake of 2641 mg/kg/d and 3348 mg/kg/d for male and female rats, respectively, under the conditions of this study.

Data Qualities Reliabilities

Reliability code 1. Reliable without restriction.

Remarks for Data Reliability

Code 1. Comparable to guideline study.

References

Borzelleca J. and Hallagan J. (1988a) A chronic toxicity/carcinogenicity study of FD & C Yellow No. 5 (Tartazine) in rats. *Fd Chem Toxic* 26, 179-187.

CAS Numerical

1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	FD&C Yellow 5; 90% purity; 10% intermediates or volatile matter
Method/guideline	Chronic Toxicity/Carcinogenicity Study
GLP	Yes
Year	1988
Species/Strain	Mice/Charles River CD-1
Sex	Male and Female

Route of administration	Oral-Diet
Doses/concentration levels	0, 0.5, 1.5, or 5.0%
Exposure period	104 weeks
Frequency of treatment	Daily
Control Group	Yes
Post exposure observation period	
Remarks for test conditions	<p>Groups of sixty male and sixty female mice each were administered 0, 0, 0.5, 1.5 or 5.0% FD & C Yellow No. 5 in the diet daily for 104 weeks. Animals were housed individually and fed the test diet <i>ad libitum</i>. Clinical observations were recorded twice daily, detailed physical examinations and palpations for masses were performed weekly. Body weights and food consumption were determined weekly for the first fourteen weeks, bi-weekly for weeks 16-26 and monthly from week 26 until the end of the study. The intake of the test substance was determined from body weight, food consumption and dietary concentration. Hematology tests, including hemoglobin, hematocrit, erythrocyte and total and differential leukocyte counts, and erythrocyte morphology, were conducted on randomly selected animals at months 3, 6, 12, 18 and 24 of the study. Necropsies were conducted on all animals dying prior to study termination, killed in a moribund condition or killed on schedule. Histological examinations were conducted on all animals from both control groups, the highest dose group (5.0%) and any animals with gross lesions or masses.</p> <p>Tissues examined included adrenal glands, brain, cecum, colon, duodenum, epididymus or uterus, esophagus, eyes, femur including marrow, tissue masses, gallbladder, heart, ileum, jejunum, kidneys, liver, lungs and bronchi, mammary gland, nerves (sciatic), ovaries, lymph nodes, pancreas, parathyroids, pituitary gland, prostate, rectum, skin, spleen, seminal vesicles, skeletal muscle, testes, stomach, thymus, thyroid gland including parathyroid, trachea, and urinary bladder.</p>
NOAEL(NOEL)	5.0 % (8103 mg/kg/day)
LOAEL(LOEL)	Not determined
Actual dose received by dose level and sex	M: 714, 2173 or 8103; F: 870, 2662 or 9735 mg/kg/day
Toxic response/effects by dose level	Physical observations included hair loss, lacrimation, nasal discharge, staining of hair in the anogenital region and soft stools. None of these observations was attributed to administration of the test substance. Discolored urine and feces was reported at all treatment levels within one week of the study initiation. Mean body weights of both sexes were slightly lower than controls at the 5.0% treatment group for a number of sampling intervals, and male mice at the 1.5% treatment group were lower than controls for a number of sampling intervals. These differences were significantly lower in

	some intervals. Mean food consumption was significantly increased in male mice at the 5.0% treatment level. No statistically significant differences were reported for any of the hematological parameters. Common neoplastic, inflammatory, and degenerative lesions were reported amongst treated and control animals but were not considered to be treatment related.
Appropriate statistical evaluations?	Yes, F-test, Anova
Remarks for results	The decrease in mean body weight at the 5.0% treatment level was not considered toxicologically significant give the non-nutritive character of FD & C Yellow No. 5.
Conclusion remarks	The NOAEL of 5.0% providing an average daily intake of 8103 or 9753 mg/kg/d was established for male and female mice under the conditions of this study.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Comparable to guideline study.
References	Borzelleca J. and Hallagan J. (1988b) A chronic toxicity/carcinogenicity study of FD & C Yellow No. 5 (Tartazine) in mice. Fd Chem Toxic 26, 189-194.

4.4 DEVELOPMENTAL TOXICITY

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
Remarks for Substance	FD&C Yellow No. 5; 92.7% purity
Method/guideline	FDA Teratology Study
Test Type	
GLP	Yes
Year	1990
Species/Strain	Rat/Osborne-Mendel (FDA strain)
Sex	Female
Route of administration	Oral-Gavage

Duration of test	19 days
Doses/concentration levels	0, 60, 100, 200, 400, 600 or 1000 mg/kg bw/day
Exposure period	19 days
Frequency of treatment	Daily
Control Group and treatment	Yes
Remarks for test conditions	Female Osborne-Mendel (FDA strain) rats (40-41 per group) were administered FD & C Yellow No. 5 via gavage at dose levels of 0, 60, 100, 200, 400, 600 or 1000 mg/kg bw/day for the first 19 days of gestation. On day 19, the animals were examined for gross abnormalities followed by euthanization. Caesarean sections were performed. The uterus was examined for presence and position of resorption sites and fetuses, number of corpora lutea and implantation sites. All live fetuses were promptly weighed, sexed, and examined. Crown-rump lengths were measured. Fetuses were divided and assigned to skeletal or soft tissue examination.
NOAEL(NOEL) maternal toxicity	Greater than 1000 mg/kg bw/day
LOAEL(LOEL) maternal toxicity	Not determined
NOAEL (NOEL) developmental toxicity	Greater than 1000 mg/kg bw/day
LOAEL (LOEL) developmental toxicity	Not determined
Actual dose received by dose level and sex	0, 60, 100, 200, 400, 600 or 1000 mg/kg bw/day
Maternal data with dose level	No unusual behavior or external findings were reported. One female at the 60 mg/kg bw/day dose level died on gestation day 13 due to gavage difficulties. The mean daily food consumption of rats administered the 1000 mg/kg bw/day dose level was significantly greater than the controls. Initial body weight and maternal weight gain during gestation did not significantly differ between treated animals and controls. Pregnancy rate was similar among all groups.
Fetal data with dose level	No dose related findings were reported on fetal viability or fetal development. The incidence of sternebral variations was similar for all groups.
Appropriate statistical evaluations?	Yes, ANOVA, Fisher's Exact Test, t-test.
Remarks for results	The authors commented that the significant increase in food consumption observed in the highest dose group without a corresponding effect on body weight indicated an effect on food utilization.
Conclusion remarks	The authors concluded that FD&C Yellow No. 5 was not developmentally toxic or teratogenic under the conditions of the study. The NOAEL's for maternal and fetal toxicity were greater than 1000 mg/kg bw/day.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.

References

Collins T., Black T.N., Brown L.H., and Bulhack P. (1990) Study of the teratogenic potential of FD & C Yellow No. 5 when given by gavage to rats. *Fd. Chem. Toxic.* Vol 28, pp 821-827.

4.5 REPRODUCTIVE TOXICITY

CAS Numerical 1934-21-0

Substance Name	C.I. Acid Yellow 23
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Remarks for Substance FD&C Yellow 5; 90% purity; 10% intermediates or volatile matter

Method/guideline Lifetime Toxicity/Carcinogenicity study

Test Type

GLP Ambiguous

Year 1988

Species/Strain Rats/Charles River CD

Sex Male and Female

Route of administration Oral-Diet

Duration of test 114 weeks

Doses/concentration levels 0, 0.1, 1.0, or 2.0% (original study) 0, 5.0% (high dose study)

Premating Exposure period for males 2 months

Premating Exposure period for females 2 months

Frequency of treatment Daily

Control Group and treatment Yes.

Remarks for test conditions In the *in utero* phase, groups of rats (60/sex/group) were administered 0, 0, 0.1, 1.0 or 2.0% FD & C Yellow No. 5 in the diet daily for approximately 2 months prior to mating. In the high-dose study, 60/sex/group received 0 or 5% FD&C Yellow 5 for approximately 2 months prior to mating. The 3 controls groups received the basal diet only. A maximum of 2 rats/sex/litter were randomly selected for the chronic phase of the study. There were 70/sex/group at the initiation of the chronic phase and these offspring were exposed to the same dietary levels as their parents.

Animals were housed individually and fed the test diet ad libitum. Clinical observations were recorded twice daily with at least 5 hours between observations. Detailed physical examinations and palpation for masses were performed weekly. Body weights and food consumption were determined weekly for the first fourteen weeks, bi-weekly for the next 12 weeks and every 4 weeks thereafter until the end of the study. The intake of the test substance was determined from body weight, food consumption and dietary concentration. Hematology tests, including hemoglobin, hematocrit, erythrocyte and total and differential leukocyte counts, and erythrocyte morphology, were conducted on ten randomly selected animals at months 3, 6, 12, 18 and 24 of the study. Necropsies were conducted on all animals dying prior to study termination, killed in a moribund condition or killed on schedule. Histological examinations were conducted on all animals from both control groups, the highest dose group (2.0 or 5.0%) from each study and also on 10 rats randomly selected from each group for an interim sacrifice at 12 months. Histology was also performed on any animal with gross lesions or masses.

Tissues examined included adrenal glands, aorta, blood smear, brain, cecum, colon, duodenum, epididymus or uterus, esophagus, eyes, femur including marrow, tissue masses, gallbladder, heart, ileum, jejunum, duodenum, kidneys, liver, lungs and bronchi, mammary gland, nerves (sciatic), ovaries, lymph nodes, pancreas, parathyroids, pituitary gland, prostate, rectum, skin, spleen, seminal vesicles, skeletal muscle, testes with epididymides, stomach, thymus, thyroid gland including parathyroid, trachea, urinary bladder, uterus.

NOAEL(NOEL)

5.0 % (Males: 2641 mg/kg/d and Females: 3348 mg/kg/day)

LOAEL(LOEL)

Not determined

Actual dose received by dose level and sex Parental data and F1 as appropriate

Males: 48, 491, 984 or 2641 mg/kg/day Females: 58, 589, 1225 or 3348 mg/kg/d

In utero

There were no compound-related effects on fertility, gestation, parturition, lactation, pup survival through weaning or number of live and still-born pups. Slight decreases in body weight (4-5%) and slight increases in food consumption were noted in the F0 rats treated at dietary level of 5.0%. Two F0 female controls rats died during the in utero phase of the original study and one male and one female from the control and 5.0% group, respectively, died during the in utero phases of the high-dose study. There were no compound-related effects on pup survival.

Offspring toxicity F1 and F2

In the F1 generation, a yellow tint was reported at all intake levels above 0.1%. At the 1.0% dietary level, group mean body weights at termination for both sexes were lower than the control animals, but the difference was only statistically significant for the females. In the high dose study (5.0% dietary level), group mean body weights were significantly lower in both sexes at termination. Food consumption was similar for control and treated animals at the 0.01, 1 or 2% dietary levels,

but was slightly higher at the 5% level in the high-dose study, although not statistically significant. Hematological, clinical chemistry and urinalysis parameters did not differ significantly from the controls. Necropsies at one year did not reveal any treatment-related gross or microscopic changes.

At study termination, no treatment-related effects were reported on survival. No treatment-related changes were reported at gross necropsy. Histological evaluation revealed a variety of lesions, including neoplasms, present at similar incidences in control and treated animals. The authors considered the lesions to be spontaneous and not related to administration of the test material.

Yes, F-test, Anova

Appropriate statistical evaluations?

Remarks for results

The decrease in mean body weight at the 5.0% treatment level was not considered toxicologically significant given the non-nutritive character of FD & C Yellow No. 5.

Conclusion remarks

Data Qualities Reliabilities

Reliability code 1. Reliable without restriction.

Remarks for Data Reliability

Code 1. Comparable to guideline study.

References

Borzelleca J. and Hallagan J. (1988a) A chronic toxicity/carcinogenicity study of FD & C Yellow No. 5 (Tartazine) in rats. *Fd Chem Toxic* 26, 179-187.