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**NATIONAL INDUSTRIAL CHEMICALS NOTIFICATION AND ASSESSMENT SCHEME
(NICNAS)**

FULL PUBLIC REPORT

**Rape Oil, Polymer with Tung Oil (INCI name: Brassica Campestris/Aleurites Fordi
Oil Copolymer)**

This Assessment has been compiled in accordance with the provisions of the *Industrial Chemicals (Notification and Assessment) Act 1989* (Cwlth) (the Act) and Regulations. This legislation is an Act of the Commonwealth of Australia. The National Industrial Chemicals Notification and Assessment Scheme (NICNAS) is administered by the Department of Health and Ageing, and conducts the risk assessment for public health and occupational health and safety. The assessment of environmental risk is conducted by the Department of the Environment, Water, Heritage and the Arts.

For the purposes of subsection 78(1) of the Act, this Full Public Report may be inspected at our NICNAS office by appointment only at 334-336 Illawarra Road, Marrickville NSW 2204.

This Full Public Report is also available for viewing and downloading from the NICNAS website or available on request, free of charge, by contacting NICNAS. For requests and enquiries please contact the NICNAS Administration Coordinator at:

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**Director
NICNAS**

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FULL PUBLIC REPORT**Rape Oil, Polymer with Tung Oil (INCI name: Brassica Campestris/Aleurites Fordi Oil Copolymer)****1. APPLICANT AND NOTIFICATION DETAILS**

APPLICANT(S)

Ingredients Plus Pty Ltd (ABN: 25 112 469 619)

Unit 8, 9-11 South Street

Rydalmere, NSW, 2116

NOTIFICATION CATEGORY

Limited-small volume: Chemical other than polymer (1 tonne or less per year).

EXEMPT INFORMATION (SECTION 75 OF THE ACT)

No details are claimed exempt from publication.

VARIATION OF DATA REQUIREMENTS (SECTION 24 OF THE ACT)

Variation to the schedule of data requirements is claimed as follows: Hydrolysis as a Function of pH, Partition Coefficient, Absorption/Desorption, Dissociation Constant, Flammability Limits and Explosive Properties.

PREVIOUS NOTIFICATION IN AUSTRALIA BY APPLICANT(S)

None.

NOTIFICATION IN OTHER COUNTRIES

EU, USA and Japan prior to 2002

2. IDENTITY OF CHEMICAL

MARKETING NAME(S)

Glossamer L6600

CAS NUMBER

185323-46-0

CHEMICAL NAME

Rape oil, polymer with tung oil

OTHER NAME(S)

Brassica Campestris/Aleurites Fordi Oil Copolymer (INCI name)

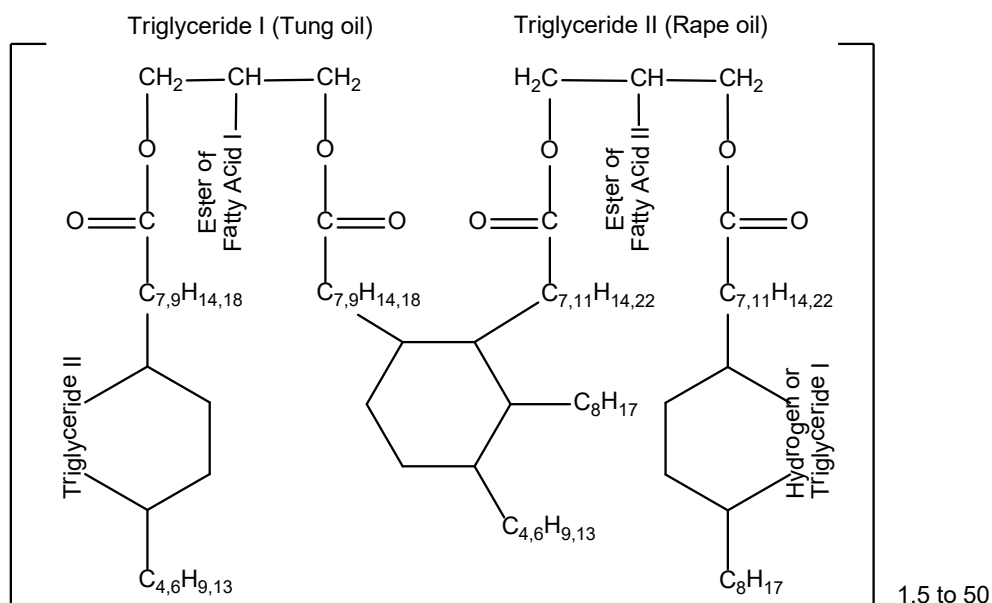
MOLECULAR FORMULA

unspecified (UVCB substance)

STRUCTURAL FORMULA

Note: UVCB Representative Structure.

- Rape oil consists primarily of the glycerides of the fatty acids erucic (18:1), linoleic (18:2) and oleic (18:1).
- Tung oil consists primarily of the glycerides of the fatty acid eleostearic (18:3)
- The notified chemical has a reported iodine value (indicating the degree of unsaturation) of 103.5.



MOLECULAR WEIGHT

Mn 2,200 [Note: The notified chemical consists of *ca.* 40 wt% of high molecular weight polymerised product (and low molecular weight oligomers) and *ca.* 60 wt% of a mixture of tung oil (Mn: 1,230) and rapeseed oil (Mn: 1,430)]. The percentages of low molecular weight species are 9.8% <1000 and 1.2% <500.

ANALYTICAL DATA

Reference IR and GPC spectra were provided.

3. COMPOSITION

DEGREE OF PURITY UVCB Substance. The notified chemical consists of *ca.* 40 wt% of high molecular weight polymerised product (and low molecular weight oligomers) and *ca.* 60 wt% of a mixture of tung oil (Mn: 1,230) and rapeseed oil (Mn: 1,430).

HAZARDOUS IMPURITIES/RESIDUAL MONOMERS None identified

NON HAZARDOUS IMPURITIES/RESIDUAL MONOMERS (>1% by weight) None identified

ADDITIVES/ADJUVANTS Tocopherol (natural): 0.1 wt%

4. PHYSICAL AND CHEMICAL PROPERTIES

APPEARANCE AT 20°C AND 101.3 kPa: yellow liquid

| Property | Value | Data Source/Justification |
|---|--------------------|--|
| Pour point | -21 °C | Measured |
| Boiling Point | 291.7 °C | Measured |
| Relative Density | 0.9393 at 60/60 °F | Measured |
| Vapour Pressure | 0.013 kPa at 25°C | Measured |
| Water Solubility | ~0.04 g/L | Measured |
| Hydrolysis as a Function of pH | Not determined | The notified chemical contains hydrolysable functionality. However, hydrolysis is expected to be slow in the environmental pH range (4-9). |
| Partition Coefficient (n-octanol/water) | Not determined | The notified chemical is expected to partition from water to oil based on its hydrophobic structure. |
| Adsorption/Desorption | Not determined | The notified chemical is expected to partition to soil, sediment and sludge based on its hydrophobic structure. |
| Dissociation Constant | Not determined | The notified chemical does not contain dissociable functionality. |
| Flash Point | 268 °C | Measured |
| Autoignition Temperature | 424 °C | Measured |
| Explosive Properties | Not determined | Expected to be stable under normal conditions of use. The notified chemical contains no functional groups that would imply explosive properties. |

DISCUSSION OF PROPERTIES

For full details of tests on physical and chemical properties, refer to Appendix A.

Reactivity

The reactivity of the notified chemical is predicted to be low and there are no known hazardous decomposition products. In addition, the high auto-ignition temperature indicates that it is not liable to spontaneous combustion. The notified chemical is incompatible with strong oxidising agents and will burn if involved in a fire, evolving noxious fumes. Therefore, it should be kept away from oxidising agents, excessive heat and ignition sources.

Dangerous Goods classification

Based on the submitted physical-chemical data in the above table the notified chemical is not classified according to the Australian Dangerous Goods Code (NTC, 2007). However the data above does not address all Dangerous Goods endpoints. Therefore consideration of all endpoints should be undertaken before a final decision on the Dangerous Goods classification is made by the introducer of the chemical.

5. INTRODUCTION AND USE INFORMATION

MODE OF INTRODUCTION OF NOTIFIED CHEMICAL (100%) OVER NEXT 5 YEARS

The notified chemical will be introduced into Australia (at 100% concentration) in HDPE buckets (Nett weight 18.14 kg).

MAXIMUM INTRODUCTION VOLUME OF NOTIFIED CHEMICAL (100%) OVER NEXT 5 YEARS

| Year | 1 | 2 | 3 | 4 | 5 |
|--------|----|----|----|----|----|
| Tonnes | ≤1 | ≤1 | ≤1 | ≤1 | ≤1 |

PORT OF ENTRY

The notified chemical will be imported into Sydney, NSW.

IDENTITY OF MANUFACTURER/RECIPIENTS

Following its introduction into Australia, the notified chemical will initially be delivered to Ingredients Plus Pty Ltd, Rydalmere, NSW, and then delivered to reformulation sites within Australia.

TRANSPORTATION AND PACKAGING

The notified chemical will be contained in HDPE buckets (Nett weight 18.14 kg). Shrink-wrapped pallets of the buckets will be transported by road to reformulation sites. Following reformulation, end-use products containing the notified chemical will be distributed within Australia *via* road or freight-train. End-use products may also be transported overseas *via* sea-freight or air-freight.

USE

The notified chemical is proposed to be used as a waterproofing and/or conditioning agent in cosmetic and personal care products at concentrations $\leq 10\%$. The notified chemical may be used in rinse-off and leave-on products, including secondary sunscreens, lip products and hair-care products (for example, in hair conditioner at 0.3%).

OPERATION DESCRIPTION

The operation description details will likely vary depending on the nature of the cosmetic and personal care products formulated, and may involve both automated and manual transfer steps. In general, it is expected that following distribution to reformulation sites and quality-assurance analysis, the notified chemical will be added to mixing vessels at the specified concentration and then transferred to appropriate containers for distribution to retail stores and sale to the general public. The following, more-detailed operation description refers to the formulation of a hair conditioner product containing the notified chemical at 0.3%.

Following quality-assurance analysis, the notified chemical will be manually added to mixing vessels (final concentration 0.3%). The product containing the notified chemical will be transferred *via* automated processes to palletcon boxes, which will in-turn be transferred to product filling lines (*via* forklift). Automated processes will then be used to transfer the product to 750 mL HDPE bottles for end-use (open system). During this process, the end-use product will be exposed to the atmosphere until the containers are sealed for distribution to retail stores.

6. HUMAN HEALTH IMPLICATIONS

6.1 Exposure assessment

6.1.1 Occupational exposure

NUMBER AND CATEGORY OF WORKERS

| <i>Category of Worker</i> | <i>Number</i> | <i>Exposure Duration (hours/day)</i> | <i>Exposure Frequency (days/year)</i> |
|---------------------------|---------------|--|---|
| Transport and storage | 10-20 | 1-2 | 50 |
| Mixing/weighing | 10-20 | ≤ 8 | 240 |
| Quality control samplers | 1-2 | 0.5 | 240 |
| Cleaners/maintenance | 5-10 | ≤ 8 | 240 |

EXPOSURE DETAILS

Transport and storage workers may come into contact with the notified chemical (at 100% or as a component of end-use products) only in the event of accidental rupture of containers.

During formulation, exposure to the notified chemical (100% and/or as a component of end-use products) may occur during weighing and transfer stages, quality control analysis and cleaning and maintenance of equipment. Exposure is expected to be minimised through the use of mechanical ventilation and personal protective equipment (MSDS of the notified chemical recommends the use of eye/face protection and chemical resistant gloves).

Exposure to the notified chemical in end-use products may occur in professions where the services provided involve the application of personal care products to clients (e.g. hair dressers, workers in beauty salons). Such professionals may use some personal protective equipment to minimise repeated exposure, and good hygiene practices are expected to be in place. Exposure of such workers is expected to be of either a similar or higher level than that experienced by consumers using products containing the notified chemical.

6.1.2. Public exposure

There will be widespread and repeated dermal exposure of the public to the notified chemical through the use of the rinse-off and leave-on cosmetic and personal care products.

6.2. Human health effects assessment

The results from toxicological investigations conducted on the notified chemical are summarised in the table below. Details of these studies can be found in Appendix B.

| <i>Endpoint (concentration of notified chemical tested)</i> | <i>Result and Assessment Conclusion</i> |
|---|---|
| Rat, acute oral toxicity (50%) | LD50 >5,000 mg/kg bw; low toxicity |
| Rabbit, skin irritation (≤20%) | slightly irritating |
| Rabbit, eye irritation (10%) | minimally irritating |
| Guinea pig, skin sensitisation – non-adjuvant test (10%) | no evidence of sensitisation |
| Mutagenicity – bacterial reverse mutation (10%) | non mutagenic |
| Comedogenicity (10%) | Non-comedogenic. |

Toxicokinetics, metabolism and distribution.

Based on the high molecular weight (>1000 Da) and low water solubility (~0.04 g/L) of the notified chemical, the potential to cross the gastrointestinal (GI) tract by passive diffusion or to be dermally absorbed after exposure is limited. However, the chemical contains a significant proportion of low molecular weight species (*ca.* 10% <1000 Da) that may be absorbed.

Acute toxicity.

The notified chemical (tested at 50% concentration) was found to be of low acute oral toxicity (LD50 >5,000 mg/kg bw)

Irritation and Sensitisation.

The notified chemical (tested at 3 concentrations: 5%, 10%, 20% notified chemical) was a slight skin irritant in rabbits. However, only 5% concentration showed slight dermal irritation whereas 10% and 20% concentrations showed no irritation effects.

The notified chemical (tested at 10% concentration) was found to be minimally irritating to the eyes of rabbits. These results suggest that the notified chemical (100%) may be a slight eye irritant.

The notified chemical (tested at 10% concentration) was not a skin sensitiser in guinea pigs (Buehler method).

The notified chemical (tested at 10% concentration) was found to be non-comedogenic in a rabbit comedogenicity assay.

Mutagenicity.

The notified chemical (tested at 10% concentration) was not mutagenic in a bacterial reverse mutation study.

Additional information

No repeat dose toxicity studies were conducted on the notified chemical.

Information regarding the potential toxicity of tung oil is, in general, sparse. The available information on this chemical is also conflicting in terms of the health effects of the chemical, and in terms of the component of the tung nut that is responsible for any observed toxicity (*i.e.* it may be components of the nut and the meal and/or the oil).

Health hazard classification

Based on the data provided the notified chemical is not classified as hazardous according to the *Approved Criteria for Classifying Hazardous Substances* (NOHSC, 2004).

6.3. Human health risk characterisation

6.3.1. Occupational health and safety

Exposure of workers to the notified chemical at up to 100% may occur during formulation of cosmetics. At such concentrations there is a possibility of slight skin and/or eye irritancy effects. However, given the measures in place to lower exposure (use of PPE), the risk of irritancy is not expected.

The risk for beauty care professionals who regularly use products containing the notified chemical is expected to be of a similar or perhaps higher level than that experienced by members of the public who use such products on a regular basis. This is because the duration of exposure will be longer for workers applying products in many clients. Considering the concentrations of notified chemical in end-use products (up to 10%), acute toxicity effects, such as skin and eye irritations, are not expected. As no repeat dose toxicity data were provided, no repeat dose risk assessment was conducted.

6.3.2. Public health

At the proposed use concentration of up to 10% notified chemical in rinse-off and leave-on cosmetic products, skin or eye irritation is not expected.

While the notified chemical was found to be of low acute oral toxicity, information is not available on the effects of long-term repeated exposure to the notified chemical.

Therefore, based on the available data, when used in the proposed manner, the risk to the public associated with the use of the notified chemical at up to 10% concentration in rinse-off and leave-on cosmetic products is not considered to be unacceptable.

7. ENVIRONMENTAL IMPLICATIONS

7.1 Environmental Exposure & Fate Assessment

7.1.1 Environmental Exposure

RELEASE OF CHEMICAL AT SITE

The notified chemical will be imported as a raw material for blending. It is expected to be released to landfill as residue in import containers (estimated to be <1% of the annual import volume). Blending will be executed in closed automated systems and notified chemical residue remaining in blending equipment is estimated to be <1%. Washings from blending equipment are anticipated to be included in the next formulation batch or treated by means of on-site waste treatment plants.

Accidental spills during transport or reformulation are expected to be collected with inert material and disposed of to landfill.

RELEASE OF CHEMICAL FROM USE

The notified chemical is a component in cosmetic products. Therefore, it is expected that the majority of the imported quantity will be released to the sewer.

RELEASE OF CHEMICAL FROM DISPOSAL

Residue of the notified chemical in the empty containers (<1%) is likely either to share the fate of the container and be disposed of to landfill, or to be washed to the sewer when containers are rinsed before recycling.

7.1.2 Environmental fate

No environmental fate data were submitted. The majority of the notified chemical will be disposed of to the sewer and, as it is a high molecular weight non-ionic chemical, it is estimated to be removed by up to 90% in sewage treatment plant from adsorption to sediment and sludge (Boethling & Nabholz, 1996). Notified chemical in the aquatic compartment is unlikely to bioaccumulate based on its high molecular weight. In landfill, the notified chemical is likely to adsorb to soil and be immobile. It is expected to degrade biotically and abiotically to form water and oxides of carbon.

7.1.3 Predicted Environmental Concentration (PEC)

Assuming that most of the notified chemical will be washed to the sewer, the following Predicted Environmental Concentration (PEC) in sewage effluent on a nationwide basis was calculated.

| <i>Predicted Environmental Concentration (PEC) for the Aquatic Compartment</i> | | |
|--|--------|--------------|
| Total Annual Import/Manufactured Volume | 1,000 | kg/year |
| Proportion expected to be released to sewer | 100% | |
| Annual quantity of chemical released to sewer | 1,000 | kg/year |
| Days per year where release occurs | 365 | days/year |
| Daily chemical release: | 2.74 | kg/day |
| Water use | 200.0 | L/person/day |
| Population of Australia (Millions) | 21.161 | million |
| Removal within STP | 0% | |
| Daily effluent production: | 4,232 | ML |
| Dilution Factor - River | 1.0 | |
| Dilution Factor - Ocean | 10.0 | |
| PEC - River: | 0.65 | µg/L |
| PEC - Ocean: | 0.06 | µg/L |

The notified chemical is expected to adsorb to sludge and sediment, hence the removal of the notified chemical from influent by sewage treatment plant (STP) processes is expected. However, in this worst case model, the majority of the notified chemical is assumed to be released in effluent. STP effluent re-use for irrigation occurs throughout Australia. The agricultural irrigation application rate is assumed to be 1000 L/m²/year (10 ML/ha/year). The notified chemical in this volume is assumed to infiltrate and accumulate in the top 10 cm of soil (density 1500 kg/m³). Using these assumptions, irrigation with a concentration of 0.647 µg/L may potentially result in a soil concentration of approximately 4.316 µg/kg. Assuming accumulation of the notified chemical in soil for 5 and 10 years under repeated irrigation, the concentration of notified chemical in the applied soil in 5 and 10 years may be approximately 21.58 µg/kg and 43.16 µg/kg, respectively. However, given the adsorptive nature of the notified chemical, these values should be considered as theoretical maximum concentrations only.

7.2. Environmental effects assessment

No ecotoxicity data were submitted. Nonionic chemicals of high molecular weight and limited water solubility are generally of low concern for the environment (Boethling & Nabholz, 1996).

7.2.1 Predicted No-Effect Concentration

The predicted no-effect concentration (PNEC) was not calculated as no ecotoxicity data were submitted and nonionic chemicals of high molecular weight and limited water solubility are generally of low concern for the environment.

7.3. Environmental risk assessment

The Risk Quotient (PEC/PNEC) was not calculated as the PNEC was not determined. However, the majority of notified chemical disposed of to the sewer is expected to be removed by adsorption to sludge and sediment in sewage treatment plant processes, and is unlikely to bioaccumulate based on its high molecular weight. Therefore, the notified chemical is not expected to pose a risk to the environment on the basis of the reported use pattern.

8. CONCLUSIONS AND REGULATORY OBLIGATIONS

Hazard classification

Based on the data provided, the notified chemical is not classified as hazardous according to the *Approved Criteria for Classifying Hazardous Substances* [NOHSC:1008(2004)].

Human health risk assessment

Under the conditions of the occupational settings described, the notified chemical is not considered to pose an unacceptable risk to the health of workers.

When used in the proposed manner, the notified chemical is not considered to pose an unacceptable risk to public health.

Environmental risk assessment

On the basis of the reported use pattern, the notified chemical is not expected to pose a risk to the environment.

Recommendations

REGULATORY CONTROLS

CONTROL MEASURES

Occupational Health and Safety

- Employers should implement the following safe work practices to minimise occupational exposure during handling of the notified chemical (100%) during formulation of products:
 - Avoid contact with skin and eyes.
- Employers should ensure that the following personal protective equipment is used by workers to minimise occupational exposure to the notified chemical (100%) during formulation of products:
 - Gloves, overalls and goggles.

Guidance in selection of personal protective equipment can be obtained from Australian, Australian/New Zealand or other approved standards.

- A copy of the MSDS should be easily accessible to employees.
- If products and mixtures containing the notified chemical are classified as hazardous to health in accordance with the *Approved Criteria for Classifying Hazardous Substances* [NOHSC:1008(2004)] workplace practices and control procedures consistent with provisions of State and Territory hazardous substances legislation must be in operation.

Disposal

- The notified chemical should be disposed of to landfill.

Emergency procedures

- Spills or accidental release of the notified chemical should be handled by physical containment, collection and subsequent safe removal.

Regulatory Obligations

Secondary Notification

This risk assessment is based on the information available at the time of notification. The Director may call for the reassessment of the chemical under secondary notification provisions based on changes in certain circumstances. Under Section 64 of the *Industrial Chemicals (Notification and Assessment) Act (1989)* the notifier, as well as any other importer or manufacturer of the notified chemical, have post-assessment regulatory obligations to notify NICNAS when any of these circumstances change. These obligations apply even when the notified chemical is listed on the Australian Inventory of Chemical Substances (AICS).

Therefore, the Director of NICNAS must be notified in writing within 28 days by the notifier, other importer or manufacturer:

- (1) Under Section 64(1) of the Act; if
 - the importation volume exceeds one tonne per annum notified chemical;
 - the notified chemical is to be used in rinse-off or leave-on cosmetic products at concentrations >10%;

or

- (2) Under Section 64(2) of the Act; if
- the function or use of the chemical has changed from a component in rinse-off and leave-on cosmetic products, or is likely to change significantly;
 - the amount of chemical being introduced has increased from 1 tonne per annum, or is likely to increase, significantly;
 - the chemical has begun to be manufactured in Australia;
 - additional information has become available to the person as to an adverse effect of the chemical on occupational health and safety, public health, or the environment.

The Director will then decide whether a reassessment (i.e. a secondary notification and assessment) is required.

Material Safety Data Sheet

The MSDS of the notified chemical provided by the notifier was reviewed by NICNAS. The accuracy of the information on the MSDS remains the responsibility of the applicant.

APPENDIX A: PHYSICAL AND CHEMICAL PROPERTIES**Pour point** -21 °C

Method ASTM D 97a
Remarks Full report not provided
Test Facility SGS (2004)

Boiling Point 291.7 °C

Method ASTM D 1120
Remarks Full report not provided
Test Facility SGS (2004)

Relative Density 0.9393 @ 60/60 °F

Method ASTM D 1298
Remarks Full report not provided
Test Facility PLTL (2004)

Vapour Pressure 0.013 kPa at 25°C

Method ASTM D 2879
Remarks Full report not provided
Test Facility PLTL (2004)

Water Solubility ~0.04 g/L

Method In house method.
Remarks The notified chemical (1.0 g) was added to distilled water (50 mL), in triplicate. The sample preparations were agitated vigorously, and excess test substance was visually observed on the surface of the water. A blank was run in parallel. All samples were centrifuged to separate the oily phase to the surface. The water phases were filtered through Whatman filter paper (#541, 150 mm diameter), and 10 mL aliquots were added to tared aluminium weighing dishes. The samples were evaporated to dryness at 70 °C, cooled in a dessicator and reweighed. Pure test substance was dried concurrently, to confirm that it was not volatile at 70 °C.
Test Facility IAL (2004)

Flash Point 268 °C

Method ASTM D 93
Remarks Full report not provided
Test Facility PLTL (2004)

Autoignition Temperature 424 °C

Method ASTM E-659
Remarks Full report not provided
Test Facility PLTL (2004)

APPENDIX B: TOXICOLOGICAL INVESTIGATIONS

B.1. Acute toxicity – oral

| | |
|-------------------|---|
| TEST SUBSTANCE | Notified chemical (50% in corn oil) |
| METHOD | Similar to OECD TG 401 Acute Oral Toxicity – Limit Test. |
| Species/Strain | Rat/Sprague-Dawley derived, albino |
| Vehicle | None |
| Remarks - Method | No significant protocol deviations |
| RESULTS | |
| Remarks - Results | There were no mortalities observed. |
| LD50 | >5,000 mg/kg bw |
| Signs of Toxicity | None |
| Effects in Organs | None |
| CONCLUSION | The test substance is of low toxicity via the oral route. |
| TEST FACILITY | PSL (1996a) |

B.2. Irritation – skin

| | |
|--------------------|---|
| TEST SUBSTANCE | Notified chemical (5%, 10% and 20% in corn oil) |
| METHOD | Similar to OECD TG 404 Acute Dermal Irritation/Corrosion. |
| Species/Strain | Rabbit/New Zealand albino |
| Number of Animals | 3 male, 3 female |
| Vehicle | None. |
| Observation Period | ca. 1, 24, 48 and 72 hours after patch removal |
| Type of Dressing | Semi-occlusive. |
| Remarks - Method | 0.5 mL of the test substances (and 100% corn oil control) were applied to 1 of 4 sites on each animal (3 male, 3 female). After 4 hours of exposure, the pads were removed and the sites wiped to remove residual test article. Individual dose sites were scored according to the Draize scoring system at ca. 1, 24, 48 and 72 hours after patch removal. |
| RESULTS | |
| Remarks - Results | For the 10% and 20% test substances, no dermal irritation was observed. For the 5% test substance, very slight edema and/or erythema was recorded at 3/6 sites. The irritation cleared from the affected sites within 24 hours. |
| CONCLUSION | The test substances are slightly irritating to the skin. |
| TEST FACILITY | PSL (1996b) |

B.3. Irritation – eye

| | |
|--------------------|---|
| TEST SUBSTANCE | Notified chemical (10% in corn oil) |
| METHOD | Similar to OECD TG 405 Acute Eye Irritation/Corrosion. |
| Species/Strain | Rabbit/New Zealand White |
| Number of Animals | 4 male, 5 female |
| Observation Period | ca. 1, 24, 48 and 72 hours post-instillation |
| Remarks - Method | 0.1 mL of the test substance was instilled into the right eye of each animal. The treated eyes of three rabbits were rinsed with saline, the eyes of the remaining six were not rinsed. The left eye served as a control. Ocular irritation was evaluated based on scores determined at ca. 1, 24, 48 and 72 hours post-instillation. |

RESULTS

Remarks - Results

No corneal opacity or iritis was noted during the study. At 1 and 24 hours post-instillation, all treated eyes exhibited conjunctivitis. All rabbits were free of ocular irritation by 72 hours.

CONCLUSION

The test substance is minimally irritating to the eye.

TEST FACILITY

PSL (1996c)

B.4. Skin sensitisation

TEST SUBSTANCE

Notified chemical (10% in corn oil)

METHOD

Similar to OECD TG 406 Skin Sensitisation – Buehler test method.

Species/Strain

Guinea pig/Hartley albino

PRELIMINARY STUDY

Maximum Non-irritating Concentration: 100%

MAIN STUDY

Number of Animals

Test Group: 10

Positive control Group: 10

INDUCTION PHASE

Induction Concentration: 100% (topical)

Signs of Irritation

Very faint erythema was observed in two treated animals following the first and second inductions.

CHALLENGE PHASE

1st challenge

topical: 100%

Remarks - Method

- 1-Chloro-2,4-dinitrobenzene [DNCB, 0.08% in 80% aqueous ethanol (induction phase) and 0.04% w/w in acetone (challenge phase)] was used as the positive control.

- In addition to the 10 animals each in the Test and Positive Control Groups, there were 5 animals each in the Test Naïve and Positive Naïve Control Groups (treated at challenge only).

RESULTS

| <i>Animal</i> | <i>Challenge Concentration</i> | <i>Number of Animals Showing Skin Reactions after:</i> | |
|-------------------------------|--------------------------------|--|-------------|
| | | <i>1st challenge</i> | |
| | | <i>24 h</i> | <i>48 h</i> |
| <i>Test Group</i> | 100% | 2/10 | 0/10 |
| <i>Positive Control Group</i> | 0.04% | 7/10 | 5/10 |

Remarks - Results

Very faint erythema was noted at two test sites 24 hours after challenge. The irritation cleared from both affected sites by 48 hours.

CONCLUSION

There was no evidence of reactions indicative of skin sensitisation to the test substance under the conditions of the test.

TEST FACILITY

PSL (1996d)

B.5. Genotoxicity – bacteria

TEST SUBSTANCE

Notified chemical (10% in DMSO)

METHOD

Similar to OECD TG 471 Bacterial Reverse Mutation Test.

Species/Strain

Plate incorporation procedure

S. typhimurium: TA1535, TA1537, TA98, TA100

E. coli: WP2uvrA

Metabolic Activation System
Vehicle

Aroclor 1254-induced rat liver (S9 homogenate)

DMSO

| | |
|-------------------|---|
| Remarks - Method | 0.1 mL/plate of test substance was used, with and without metabolic activation. Prepared mixtures were poured across triplicate plates, the plates incubated (2 days) and then the revertant colonies counted. Negative control: DMSO vehicle (with and without activation) Positive control: Dexon (TA98, TA100, TA1537); 2-aminofluorene (TA100), sodium azide (TA1535), 2-aminoanthracene (WP2uvrA) and 1-methylmethane-sulfonate (WP2uvrA) (all with and without activation). |
| RESULTS | |
| Remarks - Results | In no case was there a ≥ 2 -fold increase in the mean number of revertants of tester strains. |
| CONCLUSION | The test substance was not mutagenic to bacteria under the conditions of the test. |
| TEST FACILITY | NAMSA (2000) |

B.6. Comedogenicity

| | |
|-------------------|--|
| TEST SUBSTANCE | Notified chemical (10% in solution) |
| METHOD | In-house |
| Species/Strain | Rabbit/New Zealand albino |
| Number of Animals | 1 male, 2 female |
| Remarks - Method | The test substance was liberally applied to the right ear of each animal, once daily for 3 weeks (21 applications). The left ear served as a control. Irritation was evaluated based on scores determined prior to each application and on Day 22. The animals were euthanized and the ears removed and examined histologically. |
| RESULTS | The tissues were given a comedogenic score of zero, indicating no increase in visible follicular hyperkeratosis. |
| CONCLUSION | The test substance was non-comedogenic. |
| TEST FACILITY | PSL (1997) |

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