

# Active.Lite® Toxicology Data

**Code:** 22040  
**INCI Name:** Saccharomyces/ Grape Ferment Extract  
**CAS #:** 84929-27-1  
**EINECS #:** 284-511-6

Name of Study	Type of Study	Results
<b>Acute Daphnia Assay</b>	<i>in-vitro</i>	After 48 hours, the EC50 value for Active.Lite® was determined to be 952.6 mg/L. The conditions of OECD guideline 202 for the validity of the test were adhered to: The immobility of controls in purified drinking water (dilution water) did not exceed 10%. According to the EU Directive 93/67/EEC, this product is not classified and therefore not harmful to aquatic organisms. .
<b>Dermal &amp; Ocular Irritation Tests</b>	<i>in-vitro</i>	Both the dermal and ocular assays reveal that Active.Lite® is non-irritating and should not cause any of the aforementioned conditions.
<b>AMES Test</b>	<i>in-vitro</i>	The results of the Bacterial Reverse Mutation Assay indicate that under the conditions of this assay, that Active.Lite® was considered to be Non-Mutagenicto Salmonella typhimurium testerstrains TA98, TA100, TA1537, TA1535 and Escherichia coli WP2uvrA.



## OECD 202 Acute *Daphnia* Assay

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**Tradename:** Active.Lite®

**Code:** 22040

**CAS #:** 84929-27-1

**Test Request Form #:** 773

**Lot #:** NC140120-A

**Sponsor:** Active Concepts, LLC; 107 Technology Drive Lincolnton, NC 28092

**Study Director:** Erica Segura

**Principle Investigator:** Meghan Darley

**Test Performed:**

OECD 202

*Daphnia* spp. Acute Immobilization Test

### Introduction

The purpose of the present study is to determine the toxicity of **Active.Lite®** by exposing *Daphnia* spp. to the test substance for 48 hours and measuring the immobilization rate against the control. The present study defines an organism as being immobilized when it does not move for 15 seconds after the test vessel is gently shaken.

OECD Guideline 202 on "*Daphnia* spp., Acute Immobilization Test and Reproduction Test", adopted in 1984, included two parts: Part I – the 24 hour EC<sub>50</sub> acute immobilization test and Part II – the reproduction test (at least 14 days). Revision of the reproduction test resulted in the adoption and publication of Test Guideline 211 on "*Daphnia magna* Reproduction Test" in September 1998. Consequently, the new version of Guideline 202 is restricted to the acute immobilization test.

### Assay Principle

Young daphnids, aged less than 24 hours at the start of the test, are exposed to the test substance at a range of concentrations for a period of 48 hours. Immobilization is recorded at 24 hours and 48 hours and compared with control values. The results are analyzed in order to calculate the EC<sub>50</sub> at 48 hours. EC<sub>50</sub> is the concentration estimated to immobilize 50% of the daphnids within a stated exposure period. Immobilization refers to those animals that are not able to swim within 15 seconds after gentle agitation of the test vessel, even if they can still move their antennae.

The water solubility and vapor pressure of the test substance should be known. A reliable analytical method for the quantification of the substance in the test solutions with reported recovery efficiency and limit of determination should also be available.

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A reference substance may be tested for EC<sub>50</sub> as a means of assuring that the test conditions are reliable.

For this assay to be valid, the following performance criteria apply:

- In the control, not more than 10% of the daphnids should have been immobilized.
- The dissolved oxygen concentration at the end of the test should be at least 3 mg/L in control and test vessels.

## Materials

- Glass Test Tubes and/or Beakers
- Dissolved Oxygen Meter
- pH Meter
- Temperature Control Apparatus
- Total Organic Carbon (TOC) Analyzer
- Chemical Oxygen Demand (COD) Analyzer
- *Daphnia magna* Straus
  - Use organisms less than 24 hours old. Do not use first offspring of parents.
- Water
  - Use water suitable for culturing and testing *Daphnia* spp. It can be natural water (surface water or groundwater), dechlorinated tap water, or artificially prepared water (Table 1), but must satisfy the conditions listed in Table 2. Do not use Elendt M4 or M7 media or water containing chelating agents for testing metal-containing substances.

Substance	Concentration
Particulate Matter	<20 mg/L
Total Organic Carbon	<2 mg/L
Unionized Ammonia	<1 ug/L
Residual Chlorine	<10 ug/L
Total Organophosphorus Pesticides	<50 ng/L
Total Organochlorine Pesticides plus Polychlorinated Biphenyls	<50 ng/L
Total Organic Chlorine	<25 ng/L

**Table 1: Chemical Characteristics of Suitable Water**

Substance	Amount Added to 1 Liter Water	To prepare the reconstituted water, add the following volumes of stock solutions to 1 liter water
Calcium Chloride	11.76 grams	25 mL
Magnesium Sulfate	4.93 grams	25 mL
Sodium Bicarbonate	2.59 grams	25 mL
Potassium Chloride	0.23 grams	25 mL

**Table 2: Examples of Suitable Reconstituted Test Water**

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## Methods

### Test Conditions

- Test Method
  - Test is performed under a static, semi-static, or flow-through condition. If test substance is unstable, a semi-static or flow-through test is recommended.
- Exposure Period
  - 48 hours
- Test Volume
  - At least 2 milliliters
- Number of Test Organisms
  - At least 20 organisms for each test concentration and the control.
- Test Concentration
  - Adopt a concentration range of at least 5 concentrations, with the highest concentration inducing 100% immobilization and no effect at the lowest concentration.
- Culture Method
  - Illumination: The photoperiod is set to 16 hours light and 8 hours dark
  - Temperature: The temperature is between 18°C to 22°C
  - Dissolved Oxygen Concentration: Must be kept at 3mg/L or higher
  - Feeding: Do not feed test organisms

### Observation

- Observe mobility of the organisms at least twice (i.e., at 24 and 48 hours after exposure).
- The organisms are considered immobilized when they do not move for 15 seconds after test vessel is gently shaken.

### Measurement of Test Substance Concentrations

- At the beginning and end of exposure, measure test substance concentrations at the lowest and highest test concentration groups.
  - For volatile or adsorptive substances, additional measurements are recommended at 24 hours intervals during exposure period.

### Test Condition Measurements

- Measure dissolved oxygen in the control and at the highest test concentration at the beginning and end of the exposure period.
- Measure pH in the control and at the highest test concentration at the beginning and end of the exposure period.
- Water temperature should be measured at the beginning and end of the exposure period.

## Data and Reporting

### I. Data

- a. Data should be summarized in tabular form, showing for each treatment group and control, the number of daphnids used, and immobilization at each observation. The percentages immobilized at 24 and 48 hours are plotted against test concentrations. Data are analyzed by appropriate statistical methods (e.g. probit analysis, etc.) to calculate the slopes of the curves and the EC<sub>50</sub> with 95% confidence limits ( $p = 0.95$ ).
- b. Where the standard methods of calculating the EC<sub>50</sub> are not applicable to the data obtained, the highest concentration causing no immobility and the lowest concentration producing 100% immobility should be used as an approximation for the EC<sub>50</sub> (this being considered the geometric mean of these two concentrations).

### II. Test Report

- a. The test report must include the following:
  - i. Test substance:
    1. Physical nature and relevant physical-chemical properties
    2. Chemical identification data, including purity
  - ii. Test species:
    1. Source and species of *Daphnia*, supplier of source (if known), and the culture conditions (including source, kind and amount of food, feeding frequency)
  - iii. Test conditions:
    1. Description of test vessels: type and volume of vessels, volume of solution, number of daphnids per test vessel, number of test vessels (replicates) per concentration
    2. Methods of preparation of stock and test solutions including the use of any solvent or dispersants, concentrations used
    3. Details of dilution water: source and water quality characteristics (pH, hardness, Ca/Mg ratio, Na/K ratio, alkalinity, conductivity, etc); composition of reconstituted water if used
    4. Incubation conditions: temperature, light intensity and periodicity, dissolved oxygen, pH, etc.
  - iv. Results:
    1. The nominal test concentrations and the result of all analyses to determine the concentration of the test substance in the test vessels; the recovery efficiency of the method and the limit of determination should also be reported
    2. All physical-chemical measurements of temperature, pH and dissolved oxygen made during the test
    3. The EC<sub>50</sub> at 48 hours for immobilization with confidence intervals and graphs of the fitted model used for calculation, the slopes of the dose-response curves and their standard error; statistical procedures used for determination of EC<sub>50</sub>.



## OECD 202 Acute *Daphnia* Assay

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### Results

#### General Information:

<b>Name of new chemical substance</b>	Active.Lite®		
<b>INCI Nomenclature</b>	<i>Saccharomyces/Grape Ferment Extract</i>		
<b>CAS number</b>	N/A		
<b>Structural or rational formula</b> (if neither is available, summarize its formulation method)	Biotechnology/Botanical: <i>Saccharomyces cerevisiae &amp; Vitis vinifera</i>		
<b>Molecular weight</b>	425 Daltons		
<b>Purity of the new chemical substance used for the test (%)</b>	100%		
<b>Lot number of the new chemical substance used for the test</b>	NC140120-A		
<b>Names and contents of impurities</b>	n/a		
<b>Solubility in water</b>	100%		
<b>Properties at room temperature</b>	Clear to Slightly Hazy Liquid		
<b>Stability</b>	Heat Stable up to 70°C		
<b>Solubility in solvents, etc.</b>	<b>Solvent</b>	<b>Solubility</b>	<b>Stability in solvent</b>
	n/a	n/a	n/a

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### Test Materials and Methods:

Items		Contents	
<b>Test Organisms</b>	Species	<i>Daphnia magna</i>	
	Source	Carolina Biological Supply Company	
	Reference substance (EC <sub>50</sub> )	Potassium dichromate (0.94 mg/L)	
<b>Culture</b>	Kind of Medium	Elendt Medium M4	
	Conditions (Temperature/Photoperiod)	20°C/16 Hour Light-8 Hour Dark	
<b>Test Conditions</b>	Test Vessel	Glass	
	Material Water	Kind	Elendt Medium M4
		Hardness	250 mg/L
		pH	7.4
	Date of Exposure	03/03/2014	
	Test Concentrations	200, 90.9, 41.3, 18.8, 8.5 mg/L	
	Number of organisms	120	
	Number of Replicates	Exposure Group	4
		Control Group	4
	Test Solution Volume	2 mL	
	Vehicle	Use or Not	N/A
		Kind	N/A
		Concentration	N/A
		Number of Replicates	N/A
	Culture Method (Static, Semi-Static, Flow-Through)	Static	
	Water Temperature	20°C ± 2°C	
	Dissolved Oxygen Concentration (DO)	3 mg/L	
Photoperiod	16 Hour Light-8 Hour Dark		
Statistical Method	Probit Analysis		

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## OECD 202 Acute *Daphnia* Assay

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### Test Results:

Items		Contents
Toxicity Value	48hr EC50	952.6 mg/L
Exposure Concentrations Used for Calculation	Nominal Values	200, 90.9, 41.3, 18.8, 8.5 mg/L
Remarks		Not harmful to aquatic organisms

### Discussion

After 48 hours, the EC50 value for **Active.Lite**<sup>®</sup> was determined to be 952.6 mg/L. The conditions of OECD guideline 202 for the validity of the test were adhered to: The immobility of controls in purified drinking water (dilution water) did not exceed 10%. According to the EU Directive 93/67/EEC, this product is not classified and therefore not harmful to aquatic organisms.

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# Dermal and Ocular Irritation Tests

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**Tradename:** Active.Lite<sup>®</sup>

**Code:** 22040

**CAS #:** 84929-27-1

**Test Request Form #:** 727

**Lot #:** NC140120-A

**Sponsor:** Active Concepts, LLC; 107 Technology Drive Lincolnton, NC 28092

**Study Director:** Erica Segura

**Principle Investigator:** Meghan Darley

**Test Performed:**

In Vitro EpiDerm<sup>™</sup> Dermal Irritation Test (EPI-200-SIT)

EpiOcular<sup>™</sup> Eye Irritation Test (OCL-200-EIT)

## **SUMMARY**

*In vitro* dermal and ocular irritation studies were conducted to evaluate whether **Active.Lite<sup>®</sup>** would induce dermal or ocular irritation in the EpiDerm<sup>™</sup> and EpiOcular<sup>™</sup> model assays.

The product was tested according to the manufacture's protocol. The test article solution was found to be a **non-irritant**. Reconstructed human epidermis and cornea epithelial model were incubated in growth media overnight to allow for tissue equilibration after shipping from MatTek Corporation, Ashland, MA. Test substances were applied to the tissue inserts and incubated for 60 minutes for liquid and solid substances in the EpiDerm<sup>™</sup> assay and 30 minutes for liquid substances and 90 minutes for solid substances in the EpiOcular<sup>™</sup> assay at 37 °C, 5% CO<sub>2</sub>, and 95% relative humidity (RH). Tissue inserts were thoroughly washed and transferred to fresh plates with growth media. After post substance dosing incubation is complete, the cell viability test begins. Cell viability is measured by dehydrogenase conversion of MTT [(3-4,5-dimethyl thiazole 2-yl)], present in the cell mitochondria, into blue formazan salt that is measured after extraction from the tissue. The irritation potential of the test chemical is dictated by the reduction in tissue viability of exposed tissues compared to the negative control.

Under the conditions of this assay, the test article was considered to be **non-irritating**. The negative and positive controls performed as anticipated.

## **I. Introduction**

### **A. Purpose**

*In vitro* dermal and ocular irritation studies were conducted to evaluate whether a test article would induce dermal or ocular irritation in the EpiDerm<sup>™</sup> and EpiOcular<sup>™</sup> model assays. MatTek Corporation's reconstructed human epidermal and human ocular models are becoming a standard in determining the irritancy potential of test substances. They are able to discriminate between irritants and non-irritants. The EpiDerm<sup>™</sup> assay has accuracy for the prediction of UN GHS R38 skin irritating and no-label (non-skin irritating) test substances. The EpiOcular<sup>™</sup> assay can differentiate chemicals that have been classified as R36 or R41 from the EU classifications based on Dangerous Substances Directive (DSD) or between the UN GHS Cat 1 and Cat 2 classifications.

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# Dermal and Ocular Irritation Tests

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## **II. Materials**

- A. Incubation Conditions:** 37° C at 5% CO<sub>2</sub> and 95% relative humidity
- B. Equipment:** Forma humidified incubator, ESCO biosafety laminar flow hood, Synergy HT Microplate reader; Pipettes
- C. Media/Buffers:** DMEM based medium; DPBS; sterile deionized H<sub>2</sub>O
- D. Preparation:** Pre-incubate (37° C) tissue inserts in assay medium; Place assay medium and MTT diluent at 4° C, MTT concentrate at -20° C, and record lot numbers of kit components
- E. Tissue Culture Plates:** Falcon flat bottom 96-well, 24-well, 12-well, and 6-well tissue culture plates
- F. Reagents:** MTT (1.0mg/mL); Extraction Solution (Isopropanol); SDS (5%); Methyl Acetate
- G. Other:** Nylon Mesh Circles (EPI-MESH); Cotton tip swabs; 1mL tuberculin syringes; Ted Pella micro-spatula; 220mL specimen containers; sterile disposable pipette tips; Parafilm

## **III. Test Assay**

### **A. Test System**

The reconstructed human epidermal model, EpiDerm™, and cornea epithelial model, EpiOcular™, consist of normal human-derived epidermal keratinocytes which have been cultured to form a multilayer, highly differentiated model of the human epidermis and cornea epithelium. These models consist of organized basal, spinous, and granular layers, and the EpiDerm™ systems also contains a multilayer stratum corneum containing intercellular lamellar lipid layers that the EpiOcular™ system is lacking. Both the EpiDerm™ and EpiOcular™ tissues are cultured on specially prepared cell culture inserts.

### **B. Negative Control**

Sterile DPBS and sterile deionized water are used as negative controls for the EpiDerm™ and EpiOcular™ assays, respectfully.

### **C. Positive Control**

Known dermal and eye irritants, 5% SDS solution and Methyl Acetate, were used as positive controls for the EpiDerm™ and EpiOcular™ assays, respectfully.

### **D. Data Interpretation Procedure**

#### **a. EpiDerm™**

An irritant is predicted if the mean relative tissue viability of the 3 tissues exposed to the test substance is reduced by 50% of the mean viability of the negative controls and a non-irritant's viability is > 50%.

#### **b. EpiOcular™**

An irritant is predicted if the mean relative tissue viability of the 2 tissues exposed to the test substance is reduced by 60% of the mean viability of the negative controls and a non-irritant's viability is > 40%.

## **IV. Method**

### **A. Tissue Conditioning**

Upon MatTek kit arrival at Active Concepts, LLC the tissue inserts are removed from their shipping medium and transferred into fresh media and tissue culture plates and incubated at 37° C at 5% CO<sub>2</sub> and 95% relative humidity for 60 minutes. After those 60 minutes the inserts are transferred into fresh media and tissue culture plates and incubated at 37° C at 5% CO<sub>2</sub> and 95% relative humidity for an additional 18 to 21 hours.



# Dermal and Ocular Irritation Tests

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## B. Test Substance Exposure

### a. EpiDerm™

30µL (liquid) or 25mg (solid) of the undiluted test substance is applied to 3 tissue inserts and allowed to incubate for 60 minutes in a humidified incubator (37 °C, 5% CO<sub>2</sub>, 95% RH).

### b. EpiOcular™

Each tissue is dosed with 20µL DPBS prior to test substance dosing. 50µL (liquid) or 50mg (solid) of the undiluted test substance is applied to 2 tissue inserts and allowed to incubate for 90 minutes in a humidified incubator (37 °C, 5% CO<sub>2</sub>, 95% RH).

## C. Tissue Washing and Post Incubation

### a. EpiDerm™

All tissue inserts are washed with DPBS, dried with cotton tipped swab, and transferred to fresh media and culture plates. After 24 hours the inserts are again transferred into fresh media and culture plates for an additional 18 to 20 hours.

### b. EpiOcular™

Tissue inserts are washed with DPBS and immediately transferred into 5mL of assay medium for 12 to 14 minutes. After this soak the inserts are transferred into fresh media and tissue culture plates for 120 minutes for liquid substances and 18 hours for solid substances.

## D. MTT Assay

Tissue inserts are transferred into 300µL MTT media in pre-filled plates and incubated for 3 hours at 37 °C, 5% CO<sub>2</sub>, and 95% RH. Inserts are then removed from the MTT medium and placed in 2mL of the extraction solution. The plate is sealed and incubated at room temperature in the dark for 24 hours. After extraction is complete the tissue inserts are pierced with forceps and 2 x 200µL aliquots of the blue formazan solution is transferred into a 96 well plate for Optical Density reading. The spectrophotometer reads the 96-well plate using a wavelength of 570 nm.

## V. Acceptance Criterion

### A. Negative Control

The results of this assay are acceptable if the mean negative control Optical Density (OD<sub>570</sub>) is ≥ 1.0 and ≤ 2.5 (EpiDerm™) or ≥ 1.0 and ≤ 2.3 (EpiOcular™).

### B. Positive Control

#### a. EpiDerm™

The assay meets the acceptance criterion if the mean viability of positive control tissues expressed as a % of the negative control is ≤ 20%.

#### b. EpiOcular™

The assay meets the acceptance criterion if the mean viability of positive control tissues is < 60% of control viability.

### C. Standard Deviation

Since each irritancy potential is predicted from the mean viability of 3 tissues for EpiDerm™ and 2 tissues for EpiOcular™, the variability of the replicates should be < 18% for EpiDerm™ and < 20% EpiOcular™.



# Dermal and Ocular Irritation Tests

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## VI. Results

### A. Tissue Characteristics

The tissue inserts included in the MatTek EpiDerm™ and EpiOcular™ assay kits were in good condition, intact, and viable.

### B. Tissue Viability Assay

The results are summarized in Figures 1 and 2. In no case was the tissue viability  $\leq 50\%$  for EpiDerm™ or  $\leq 60\%$  for EpiOcular™ in the presence of the test substance. The negative control mean exhibited acceptable relative tissue viability while the positive control exhibited substantial loss of tissue viability and cell death.

### C. Test Validity

The data obtained from this study met criteria for a valid assay.

## VII. Conclusion

Under the conditions of this assay, the test article substance was considered to be **non-irritating**. The negative and positive controls performed as anticipated.

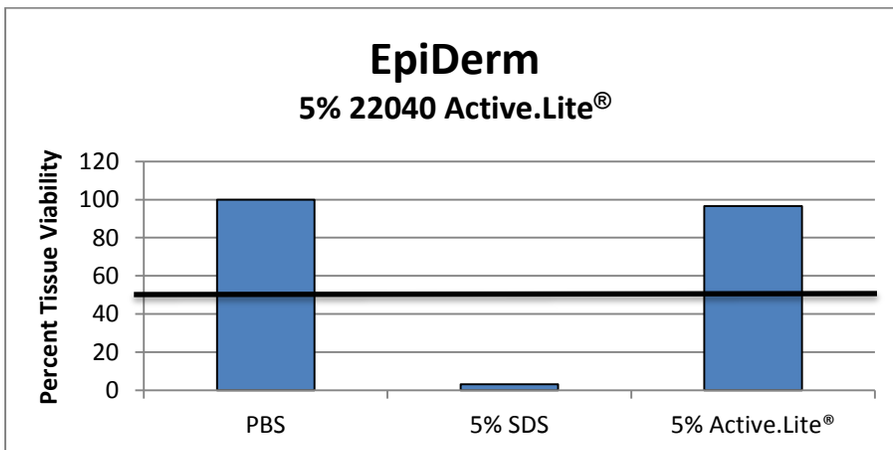


Figure 1: EpiDerm tissue viability

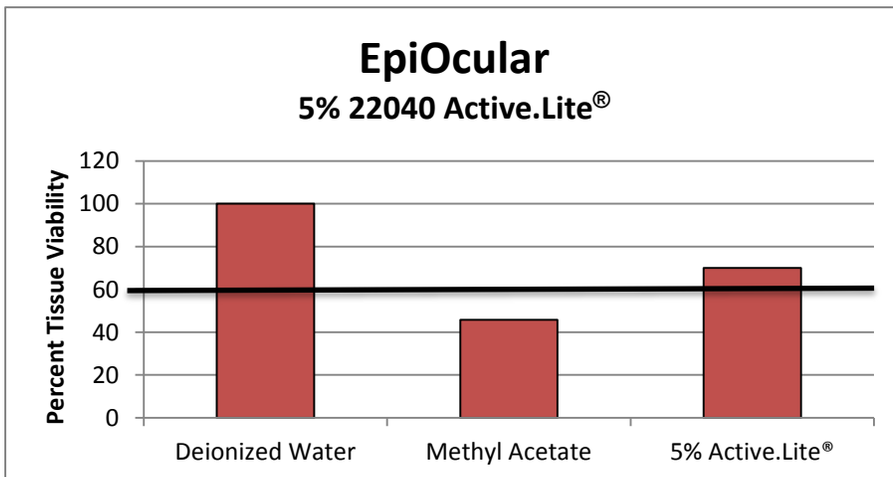


Figure 2: EpiOcular tissue viability

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# Bacterial Reverse Mutation Test

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**Test Article:** Active.Lite®  
**Code Number:** 22040  
**CAS #:** 84929-27-1

**Sponsor:**  
Active Concepts, LLC  
107 Technology Drive  
Lincolnton, NC 28092

**Study Director:** Erica Segura  
**Principle Investigator:** Monica Beltran

**Test Performed:**  
Genotoxicity: Bacterial Reverse Mutation Test  
**Test Request Number:** 715

**Reference:**  
OECD471/ISO10993.Part 3

## SUMMARY

A *Salmonella typhimurium*/*Escherichia coli* reverse mutation standard plate incorporation study described by Ames *et al.* (1975) was conducted to evaluate whether a test article solution **Active.Lite®** would cause mutagenic changes in the average number of revertants for histidine-dependent *Salmonella typhimurium* strains TA98, TA100, TA1537, TA1535 and tryptophan-dependent *Escherichia coli* strain WP2uvrA in the presence and absence of Aroclor-induced rat liver S9. This study was conducted to satisfy, in part, the Genotoxicity requirement of the International Organization for Standardization: Biological Evaluation of Medical Devices, Part 3: Tests for Genotoxicity, Carcinogenicity and Reproductive Toxicity.

The stock test article was tested at eight doses levels along with appropriate vehicle control and positive controls with overnight cultures of tester strains. The test article solution was found to be noninhibitory to growth of tester strain TA98, TA100, TA1537, TA1535 and WP2uvrA after Spot Inhibition Screen.

Separate tubes containing 2 ml of molten top agar at 45°C supplemented with histidine-biotin solution for the *Salmonella typhimurium* strains and supplemented with tryptophan for *Escherichia coli* strain were inoculated with 100 µl of tester strains, 100 µl of vehicle or test article dilution were added and 500 µl aliquot of S9 homogenate, simulating metabolic activation, was added when necessary. After vortexing, the mixture was poured across the Minimal Glucose Agar (GMA) plates. Parallel testing was also conducted with positive control correspond to each strain, replacing the test article aliquot with 50µl aliquot of appropriate positive control. After the overlay had solidified, the plates were inverted and incubated for 48 hours at 37°C. The mean numbers of revertants of the test plates were compared to the mean number of revertants of the negative control plates for each of the strains tested. The means obtained for the positive controls were used as points of reference.

Under the conditions of this assay, the test article solution was considered to be Non-Mutagenic to *Salmonella typhimurium* tester strains TA98, TA100, TA1537, TA1535 and *Escherichia coli* tester strain WP2uvrA. The negative and positive controls performed as anticipated. The results of this study should be evaluated in conjunction with other required tests as listed in ISO 100993, Part 3: Tests for Genotoxicity, Carcinogenicity, and Reproductive Toxicology.

## I. Introduction

### **A. Purpose**

A *Salmonella typhimurium*/*Escherichia coli* reverse mutation standard plate incorporation study was conducted to evaluate whether a test article solution would cause mutagenic changes in the average number of revertants for *Salmonella typhimurium* tester strains TA98, TA100, TA1537, TA1535 and *Escherichia coli* WP2uvrA in the presence and absences of the S9 metabolic activation. Bacterial reverse mutation tests have been widely used as rapid screening procedures for the determination of mutagenic and potential carcinogenic hazards.

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# Bacterial Reverse Mutation Test

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## II. Materials

- A. **Storage Conditions:** Room temperature (23-25C).
- B. **Vehicle:** Sterile DI Water.
- C. **Preparation:** Eight different doses level were prepared immediately before use with sterile DI water.
- D. **Solubility/Stability:** 100% Soluble and Stable.
- E. **Toxicity:** No significant inhibition was observed.

## III. Test System

### **A. Test System**

Each *Salmonella typhimurium* and *Escherichia coli* tester strain contains a specific deep rough mutation (*rfa*), the deletion of *uvrB* gene and the deletion in the *uvrA* gene that increase their ability to detect mutagens, respectively. These genetically altered *Salmonella typhimurium* strains (TA98, TA100, TA1537 and TA1535) and *Escherichia coli* strain (WP2*uvrA*) cannot grow in the absence of histidine and tryptophan, respectively. When placed in a histidine-tryptophan free medium, only those cells which mutate spontaneously back to their wild type states are able to form colonies. The spontaneous mutation rate (or reversion rate) for any one strain is relatively constant, but if a mutagen is added to the test system, the mutation rate is significantly increased.

<u>Tester strain</u>	<u>Mutations/Genotypic Relevance</u>
TA98	hisD3052, Dgal chlD bio <i>uvrB rfa</i> pKM101
TA100	hisG46, Dgal chlD BIO <i>uvrB rfa</i> pKM101
TA1537	hisC3076, <i>rfa</i> , Dgal chlD bio <i>uvrB</i>
TA 1535	hisG46, Dgal chlD bio <i>uvrB rfa</i>
WP2 <i>uvrA</i>	trpE, <i>uvrA</i>

<i>rfa</i>	=	causes partial loss of the lip polysaccharide wall which increases permeability of the cell to large molecules.
<i>uvrB</i>	=	deficient DNA excision-repair system (i.e., ultraviolet sensitivity)
pKM101	=	plasmid confers ampicillin resistance (R-factor) and enhances sensitivity to mutagens.
<i>uvrA</i>	=	All possible transitions and transversions, small deletions.

### **B. Metabolic Activation**

Aroclor induced rat liver (S9) homogenate was used as metabolic activation. The S9 homogenate is prepared from male Sprague Dawley rats. Material is supplied by MOLTOX, Molecular Toxicology, Inc.

### **C. Preparation of Tester strains**

Cultures of *Salmonella typhimurium* TA98, TA100, TA1537, TA1535 and *Escherichia coli* WP2*uvrA* were inoculated to individual flasks containing Oxoid broth No.2. The inoculated broth cultures were incubated at 37°C in an incubator shaker operating at 140-150 rpm for 12-16 hours.

### **D. Negative Control**

Sterile DI water (vehicle without test material) was tested with each tester strain to determine the spontaneous reversion rate. Each strain was tested with and without S9 activation. These data represented a base rate to which the number of revertants colonies that developed in each test plate were compared to determine whether the test material had significant mutagenic properties.

### **E. Positive Control**

A known mutagen for each strain was used as a positive control to demonstrate that tester strains were sensitive to mutation to the wild type state. The positive controls are tested with and without the presence of S9 homogenate.

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## F. Titer of the Strain Cultures:

Fresh cultures of bacteria were grown up to the late exponential or early stationary phase of growth; to confirm this, serial dilutions from each strain were conducted, indicating that the initial population was in the range of 1 to  $2 \times 10^9$ /ml.

## IV. Method

### A. Standard Plate Incorporation Assay:

Separate tubes containing 2 ml of molten top agar supplemented with histidine-biotin solution for the *Salmonella typhimurium* and tryptophan for *Escherichia coli* were inoculated with 100  $\mu$ l of culture for each strain and 100  $\mu$ l of testing solution or vehicle without test material. A 500  $\mu$ l aliquot of S9 homogenate, simulating metabolic activation, was added when necessary. The mixture was poured across Minimal Glucose Agar plates labeled with strain number and S9 activation (+/-). When plating the positive controls, the test article aliquot was replaced by 50  $\mu$ l aliquot of appropriate positive control. The test was conducted per duplicate. The plates were incubated for 37°C for 2 days. Following the incubation period, the revertant colonies on each plate were recorded. The mean number of revertants was determined. The mean numbers of revertants of the test plates were compared to the mean number of revertants of the negative control of each strain used.

## V. Evaluation

For the test solution to be evaluated as a test failure or "potential mutagen" there must have been a 2-fold or greater increase in the number of mean revertants over the means obtained from the negative control for any or all strains. Each positive control mean must have exhibited at least a 3-fold increase over the respective negative control mean of the *Salmonella* tester strain used.

## VI. Results and Discussion

### A. Solubility:

Water was used as a solvent. Solutions from the test article were made from 0.015 to 50mg/ml.

### B. Dose levels tested:

The maximum dose tested was 5000  $\mu$ g per plate. The dose levels tested were 1.5, 5.0, 15, 50, 150, 500, 1500 and 5000  $\mu$ g per plate.

### C. Titer (Organisms/ml):

$5 \times 10^8$  UFC/ml plate count indicates that the initial population was in the range of 1 to  $2 \times 10^9$  UFC/ml.

### C. Standard Plate Incorporation Assay

In no case was there a 2-fold or greater increase in the mean number of revertant testing strains TA98, TA100, TA1537, TA1535 and WP2uvrA in the presence of the test solution compared with the mean of vehicle control value. The positive controls mean exhibited at least a 3-fold increase over the respective mean of the *Salmonella typhimurium* and *Escherichia coli* tester strains used. The results are summarized in Appendix 2.

## VII. Conclusion

All criteria for a valid study were met as described in the protocol. The results of the Bacterial Reverse Mutation Assay indicate that under the conditions of this assay, the test article solution was considered to be Non-Mutagenic to *Salmonella typhimurium* tester strains TA98, TA100, TA1537, TA1535 and *Escherichia coli* WP2uvrA. The negative and positive controls performed as anticipated. The results of this study should be evaluated in conjunction with other required tests as listed in ISO 100993, Part 3: Tests for Genotoxicity, Carcinogenicity, and Reproductive Toxicology.



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## Appendix 2:

### Bacterial Mutation Assay Plate Incorporation Assay Results

	Concentration µg per Plate	<b>TA98</b>		
		Revertants per plate (CFU)		Mean
Test Solution w/ S9	5000	69	66	68
	1500	60	81	71
	500	98	62	80
	150	43	81	62
	50	54	76	65
	15	58	88	73
	5.0	98	64	81
	1.5	55	82	69
Test Solution w/o S9	5000	96	78	87
	1500	42	74	58
	500	73	78	76
	150	86	62	74
	50	75	43	59
	15	73	60	67
	5.0	63	68	66
	1.5	67	73	70
DI Water w/S9		53	60	57
DI Water w/o S9		79	84	82
2-aminoanthracen w/ S9		390	340	365
2-nitrofluorene w/o S9		150	165	158
Historical Count Positive w/S9		<b>43-1893</b>		
Historical Count Positive w/o S9		<b>39-1871</b>		
Historical Count Negative w/S9		<b>4-69</b>		
Historical Count Negative w/o S9		<b>3-59</b>		

\*CFU = Colony Forming Units

\*Mean = Average of duplicate plates

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	Concentration µg per Plate	TA100		
		Revertants per plate (CFU)		Mean
Test Solution w/ S9	5000	132	228	180
	1500	144	164	154
	500	108	152	130
	150	84	160	122
	50	118	118	118
	15	176	188	182
	5.0	148	140	144
	1.5	100	180	140
Test Solution w/o S9	5000	232	248	240
	1500	392	352	372
	500	212	208	210
	150	128	124	126
	50	156	200	178
	15	156	160	158
	5.0	156	140	148
	1.5	96	104	100
DI Water w/S9		136	144	140
DI Water w/o S9		1060	180	143
2-aminoanthracen w/ S9		810	812	811
Sodium azide w/o S9		1000	640	820
Historical Count Positive w/S9		<b>224-3206</b>		
Historical Count Positive w/o S9		<b>226-1837</b>		
Historical Count Negative w/S9		<b>55-268</b>		
Historical Count Negative w/o S9		<b>47-250</b>		

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	Concentration µg per Plate	<i>TA1537</i>		
		Revertants per plate (CFU)		Mean
Test Solution w/ S9	5000	7	10	9
	1500	11	4	8
	500	10	8	9
	150	7	11	9
	50	8	4	6
	15	5	8	7
	5.0	8	3	6
	1.5	8	6	7
Test Solution w/o S9	5000	26	15	21
	1500	11	9	10
	500	6	13	10
	150	13	6	10
	50	7	8	8
	15	13	7	10
	5.0	14	7	11
	1.5	8	13	11
DI Water w/S9		192	184	188
DI Water w/o S9		132	156	144
2-aminoanthracen w/ S9		50	60	60
2-aminoacridine w/o S9		520	792	656
Historical Count Positive w/S9		<b>13-1934</b>		
Historical Count Positive w/o S9		<b>17-4814</b>		
Historical Count Negative w/S9		<b>0-41</b>		
Historical Count Negative w/o S9		<b>0-29</b>		

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	Concentration µg per Plate	<b>TA1535</b>		
		Revertants per plate (CFU)		Mean
Test Solution w/ S9	5000	11	19	15
	1500	18	17	18
	500	11	19	15
	150	15	12	14
	50	16	14	15
	15	7	9	8
	5.0	16	13	15
	1.5	7	20	14
Test Solution w/o S9	5000	11	10	11
	1500	17	12	15
	500	12	8	10
	150	12	10	11
	50	7	12	10
	15	7	12	10
	5.0	12	13	13
	1.5	11	8	10
DI Water w/S9		13	18	16
DI Water w/o S9		10	16	13
2-aminoanthracen w/ S9		170	260	220
Sodium azide w/o S9		544	480	512
Historical Count Positive w/S9		<b>22-1216</b>		
Historical Count Positive w/o S9		<b>47-1409</b>		
Historical Count Negative w/S9		<b>1-50</b>		
Historical Count Negative w/o S9		<b>1-45</b>		

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	Concentration µg per Plate	<i>WP2uvrA</i>		
		Revertants per plate (CFU)		Mean
Test Solution w/ S9	5000	44	60	52
	1500	62	70	66
	500	68	69	69
	150	41	50	46
	50	56	38	47
	15	56	75	66
	5.0	75	74	75
	1.5	52	69	51
Test Solution w/o S9	5000	35	60	48
	1500	65	66	66
	500	47	64	56
	150	63	47	55
	50	71	51	61
	15	74	70	72
	5.0	73	56	65
	1.5	66	76	71
DI Water w/S9		33	66	50
DI Water w/o S9		68	71	70
2-aminoanthracen w/ S9		590	680	640
Methylmethanesulfonate w/o S9		250	211	231
Historical Count Positive w/S9		<b>44-1118</b>		
Historical Count Positive w/o S9		<b>42-1796</b>		
Historical Count Negative w/S9		<b>8-80</b>		
Historical Count Negative w/o S9		<b>8-84</b>		

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