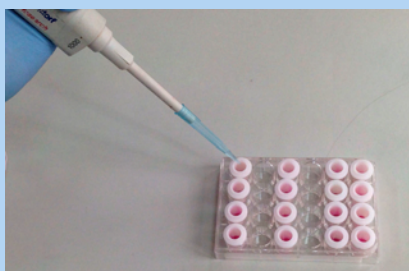
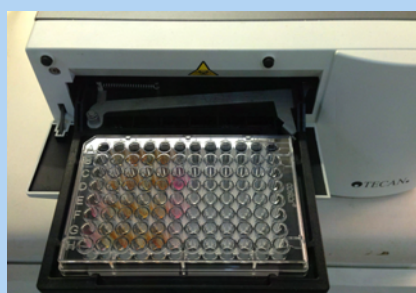




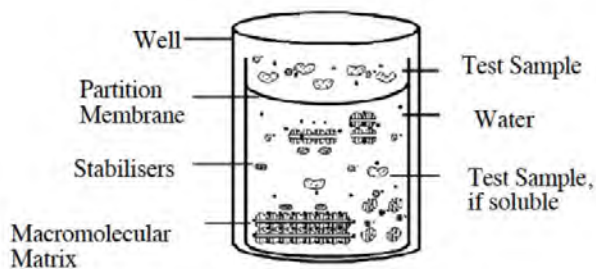
## STEPS



1. SAMPLE PREPARATION



2. PLATE READER



3. REACTION CELL CROSS-SECTION

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## Ocular Irritation

### Determination of Irritancy Potential

This test replaces the Draize test, previously conducted on the Rabbit.

### Supportable Claims

- Non Animal Safety Testing.
- Low Ocular Irritation.
- Accepted as OECD as alternative to animal testing.

### Principle

This in-vitro test is based on the principle that chemical compounds will promote measurable changes in target biomolecules and macromolecular structures. The proprietary Irritaction assay is a standardized and quantitative in vitro test which utilizes changes of relevant macromolecules to predict the acute ocular irritancy of chemicals and chemical formulations.

A macromolecular matrix, composed of proteins, glycoproteins, lipids and low molecular weight components, is included in each well.

### Plate Reader

After removal of the membrane and its support, turbidity of the reacted solutions is measured via a Plate Reader. More irritant substances produce greater turbidity and thus higher optical density.

### Calculation of Comparative Irritancy

The irritancy potential of a test sample is expressed as an Irritaction Draize Equivalent (IDE) score. This score is defined by comparing the changes in optical density (OD405) produced by the test material to a standard curve that is constructed by measuring the increase in OD405 produced by a set of Calibration substances.

Irritaction Draize Equivalent (IDE)	Predicted Ocular Irritancy Classification	UN GHS/EU CLP Classification
0.0 - 12.5	Minimal Irritant	Non-irritant [No Category]
12.6 - 30.0	Mild Irritant	Irritant [Category 1/Category 2]
30.1 - 51.0	Moderate Irritant	
51.1 - 80.0	Severe Irritant	

### Sample Preparation

The sample is serial diluted according to relevant in-use concentration. Aliquots of these samples are mixed with reagents and positive and negative controls. All of these solutions are set up in wells which have been fitted with a partition membrane which represents the membrane of the eye.

### Reporting References

1. Ocular Irritaction®: In Vitro Method for Testing Ocular Irritancy Bufo M., Ulmer R. SOFW Journal 5-2008
2. In vitro alternatives for ocular irritation. Curren R. D., Harbell J. W. Environ Health Perspect. Vol 106



Dermatest



Irritation - Ocular or Skin Irritation Test Checklist for Sample Submission

1. If the sample is less than pH 2 or greater than pH 9 then it is unsuitable for the test !

3. Is the sample highly coloured in solution? If so, please advise the dye/s used as we will need to adjust the wavelength for determination.

4. Is the sample oil based or water insoluble? If yes, we will need to solubilise it and will discuss this with you prior to running the test.

5. Please advise if the sample contains any of the following...

- Table with 2 columns: Substance name and 'No' response.

Company.....

Contact.....

Phone / fax/ email .....

- Checklist items with checkboxes: Please provide at least 20mL/gm of product, Clearly identify your product name and reference number, Any special instructions for pre-dilution or preparation prior to testing.

Please forward the sample/s to... Eurofins Dermatest Pty Ltd P.O. Box 1022 Rockdale NSW Australia 2216 ...or by Courier to 20 King St Rockdale