

### PRODUCT SPECIFICATION

# instaCELL Cytotoxicity Assay Kit

CatN°: SF020-01

Lot#:

Expiry Date: 01.01.2024

CX-220729

#### **PRODUCT DEFINITION**

Test kit to assess the cytotoxicity of chemicals and leachables by their application to cultures of mammalian cells and the subsequent determination of cell viability.

### **QUALITY SPECIFICATION OF THE CELLS**

	Batch Quality Control	Specification Limits
Cell Count	1.07E+07	9.00E+06<>1.20E+07
Homogenity (cell count)	98%	≥ 90%
Viability (after thawing)	96%	≥ 90%
Proliferative Capacity	98%	≥ 70%
Debris/Cell Ratio	0.3	≤ 1.0
Aggregation	1.2	≤ 2.0
Sterility (bacteria, yeast, fungi)	passed	negative after 7 days
Sterility (mycoplasma)	passed	negative by PCR
Morphology	passed	unalterd to reference
Cytotoxicity Assay (max signal)	16940 RFU	15000 RFU < result > 25000 RFU
Cytotoxicity Assay (min signal)	1104 RFU	1000 RFU < result > 4000 RFU
Cytotoxicity Assay (IC50)	Sodium Selenite: 2.3E-05M	1.0E-05 M < x < 2.0E-04 M
Cytotoxicity Assay (Z')	0.94	> 0.5

## KIT CONTENT

	Lot#	Storage
Recovery Buffer A	91-220628NR01	-20°C
Assay Buffer A	91-220701NR01	-20°C
Assay Medium A	91-220630NR01	-20°C
Cytotoxic Control	91-220701NR02	-20°C
Resazurin Solution	91-220727MD01	-20°C
96-well Assay Plate	I181673G	RT
Assay Ready L-929 Cells	92-220328NR01	< -140°C

Sterility was analyzed by microscopic/visual control after seven days according to sterility testing. Functionality of the content was tested by performing the assay with all listed batches.

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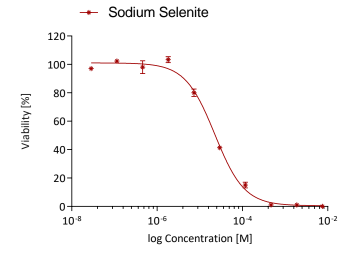
#### **MORPHOLOGY**





Morphology of cells 24 hours after seeding

### CYTOTOXICITY ASSAY



Dose response of the reference compound Sodium Selenite (Cytotoxic Control) performed according to the assay protocol.

#### **METHODS**

Cell Viability Parameters:

Viability parameters (viable cell count after thawing, grade of aggregation, percentage of debris) were determined from a pooled sample (SOP-2015-02). Briefly, assay ready cells were thawed in a water bath.  $100~\mu l$  of each sample were pooled, diluted 1:1000 in CASY Ton buffer and measured (3 replica) in a CASY TT automatic cell counter. Vial to vial variation was determined in a plate-based viability assay.

**Proliferative Capacity:** 

Proliferative Capacity compares mean growth rate (TO - T72 hours) of all sample vials with mean growth rate of exponentially growing culture. Freshly thawed cells from the assay ready cell samples were seed in a 96-well plate (3 replica each) according their specific 3 day seeding density. After 72 hours of cultivation, the proliferation of the cells was determined by addition of a metabolic cell dye (Resazurin) (SOP-2017-03).

**Sterility Testing:** 

Assay ready cells were seed in two specific bacteria growth brothes (Tryptic Soy Broth for aerob and Thioglycollate broth for anaerob conditions) and cultivated over a course of 14 days. After day 1, 4, 7 and 14 the cultures were analyzed microscopically for cell growth, cell morphology, and incidences of contamination (bacteria, yeast, or fungi). For mycoplasma testing from a three days old, sub-confluent culture 500  $\mu$ l of the supernatant was taken and analyzed by PCR using a Mycoplasma detection kit (Minerva). Assay was performed according to the manufacturer protocol (SOP-2015-06).

Cytotoxicity Assay:

The cells were seeded at 7E+04 c/well in a 96-well plate and treated with the reference substance Sodium Selenite for 24h at 37°C and 5% CO2. After the incubation phase 20µl of a 400µM Resazurin solution was added to the cells and after 4h the viability was determined by fluorescence measurement with a Tecan Safire2. Based on the dose-dependent viability the IC50 of each reference substance was calculated using GraphPad Prism.

### **LIMITED USE**

The product is provided under the terms of a limited use license provided with the kit. By breaking the sealed bag, the user is explicitly accepting the terms for limited use.