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Deutsche
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**Cytotoxicity Test to DIN EN ISO 10993-5
SOP 09-001**

2014-03-27

TESTREPORT

Identification of the test laboratory: SN 16673.1

Delivery date: 2014-03-20

Product: ProSkin[®]

Customer: Pro Seating BV

Test method: Cytotoxicity of eluates according to the
DIN EN ISO 10993-5:2009-10
Biological evaluation of medical devices
Part 5: tests for cytotoxicity: in vitro
SOP 09-001

Test time period: 2014-03-25 until 2014-03-27

Test conditions: Examining climate: 18°C / 41% rel. humidity
Incubation: 24 hours
The samples were checked in the delivery state.

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Description of the method

Extraction conditions: 25 cm² material into 10 ml MEM + 9 % serum +1 % antibiotic solution at 37°C for 24 h = **extraction medium**

cell culture FI-cells are derived from the human amnion. The stock cultures were carried out into 250 ml culture flasks (Greiner GmbH). The cells were trypsinised all 4 days. Only cells up to 100 passages were used.

Trypsinised cells were seeded in tissue culture plates.

The culture medium consists of MEM (Minimum Essential Medium) supplemented with 9 % calf serum, 1 % antibiotic solution (Penicilline G, Streptomycin sulfate, Neomycin) and L-glutamine.

Exposition After 24hours of cultivation the cells were available as monolayer. A medium change with extraction medium was accomplished. Therefore the culture medium was decanted and the extraction medium carefully pipetted into the wells (0.1 ml per well).

An incubation for 24h is following.

Measuring principle Vital cells incorporate the dye neutral red. Destroyed cells cannot incorporate the dye and remain unstained. The intensity of colour of the elution solution can be measured with a photometer.

Measurement At the end of the incubation time the microtiterplate will be washed with PBS (Phosphate Buffered Saline). Culture medium containing the dye neutral red (50µg/ml) was given to the cells. After an incubation time of 3 hours the microtiterplate was washed again to remove the spare dye. With a special elution solution (1% acetic acid in 50% ethyle alcohol) the dye was solved out of the cells. After 1 hour of elution the photometric measurement was conducted.

Controls As a negative control culture medium without a test solution was established.

To verify the sensitivity of the test system a positive control (1.5mg/ml Sodiumdodecylsulfate) in culture medium was exposed in the cell culture system.

Evaluation The optical density of 12 parallel tests was determined and used for statistical evaluation.

Results

Figure 1: box plot of the cellvitality

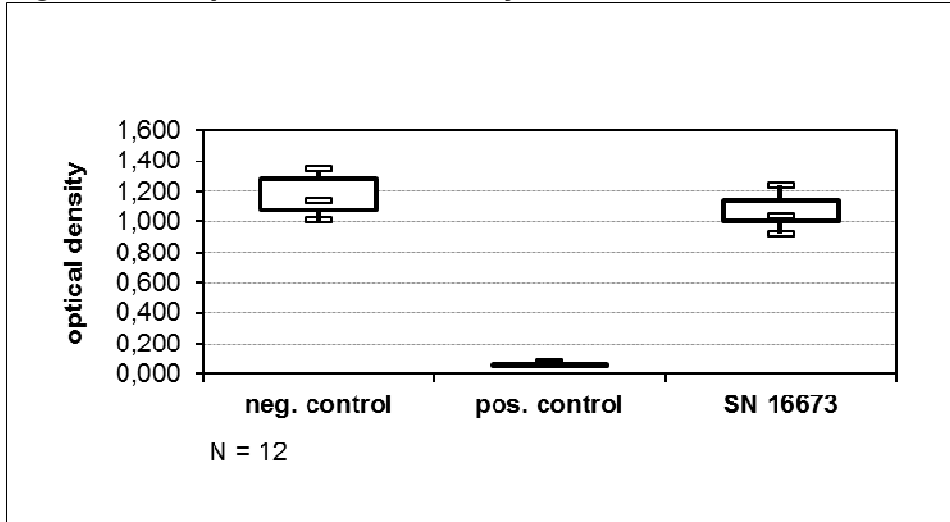


Table 1: Descriptive statistics (cellvitality)

	N	Mean	cell vitality (%)	minimum	maximum	Std. Deviation	p*
Negative control	9	1,172	100,00	1,005	1,340	0,127	-
Positive control	9	0,064	5,46	0,057	0,072	0,005	-
SN 16673	12	1,061	90,49	0,921	1,239	0,100	0,9562

*U test (Man Whitney) vs. Control

Archiving: The raw data with respect to this test and a copy of the report will be stored in the archive of HygCen.

Information: The test results exclusively refer to the samples described above. Account of extracts of this test report is only possible by written approval from HygCen.

Prof. Dr. med. H.-P. Werner
Manager of scientific-technical affairs

Dipl. Umweltwiss. J. Köhnlein
Vice department manager

Annex of testreport SN 16673 of 2014-03-27



Fig. 2: Material ProSkin ®

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2014-03-27

ProSkin®

Judgement

After testing the Cytotoxicity of the material "ProSkin®" according to the DIN EN ISO 10993-5:2009-10 -- test report of 2014-03-27 (Testreport SN 16673.1)-- I give the following statement:

An evaluation of the scope of biological testing was carried out as per EN ISO 10993-1:2010-04.

The intended use of the product, declared by the producer, involves contact with intact skin for a period of less than 24 hours. To evaluate the biocompatibility of the product, a cytotoxicity test as per EN ISO 10993-5:2009-10 was therefore considered sufficient.

Any knowledge to be gained from further biocompatibility testing with this product would not justify the unnecessarily high level of harm to experimental animals involved. As per EN ISO 10993-1:2010-04, chapter 4.6 and chapter 6.2.1 8), such tests will therefore not be performed in these cases.

The type and scope of the tests performed complies with the specifications as per EN ISO 10993-1:2010-04.

From the tested material only minimal cytotoxic compounds were extracted at 37°C. The extract of the test material reduced the cell growth to 90,49% of control. This is statistically not significant (testreport SN 16673, Fig. 1 and Tab. 1).

Using the test material as mentioned before described by the manufacturer no cytotoxic effects should be expected.

A handwritten signature in blue ink, consisting of several overlapping, fluid strokes that form a stylized, somewhat abstract shape.

Prof. Dr. med. H.-P. Werner