

## **ANTIGENES GmbH**

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Art.-No. ZR106020 20 applications IVD in vitro diagnostics

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# Manual

PLEASE READ CAREFULLY

# **SemenHos**

(Seminal Hypo-Osmotic test)

# Distribution:



# Professional Use Only

# **Application**

This SemenHos test is used to test the vitality of sperm cells. The hypo-osmotic swelling is based on the semi-permeability of the intact cell membrane and their ability of active water transport, in order not to burst. In sperms with intact membranes the flagellum swells up within 5 min. This change remains stable up to 30 min.

## **Principle**

In this hypo-osmotic swelling test swelling of cells only occurs in vital cells with an intact membrane by using hypotonic solution.

# Storage and stability

1 2-8°C. Sterile sampling. Contains no antibiotic.

24 months from date of manufacture. After opening use within 7 days.

# Content

SemenHos solution 20 x 900 µl

## **Necessary utensils**

- Coverslips (18 x 18 mm)
- Gloves
- Contrasting phase microscope
- Native ejaculate or washed sperm (105-110 µl)
- Slides
- Paper towels
- Pipettes and tips (10-100 μl)
- Water bath or heating cabinet (37°C)

#### **Preparation of SemenHos solution**

Preheat the SemenHos solution to 37°C

#### **Procedure**

- 1. Occasionally native ejaculate without additions includes hypo-osmotic sperm forms. Transfer 5-10 µI liquefied semen without air bubbles to a slide and cover it with cover slips. Microscope at 400x magnification. This is the zero-value.
- 2. Examine the percentage of sperms with swollen flagellum by observing 100 sperms, calculated in duplicate. Note this value note\* (a<sub>0</sub>%).
- **3.** Add 100 μl ejaculate without air bubbles to 900 μl preheated SemenHos solution and mix
- 4. Incubate this mixture for 10 min at 37°C.
- **5.** Transfer 10 µl of the mixture to a slide and place and cover it with a cover slip.
- 6. Microscope at 200x or 400x magnification.
- 7. Repeat twice step 2 to 6.

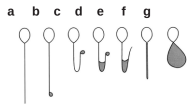


Fig.1: Schematic representation of typical morphological changes of human spermatozoa after exposure with hypo-osmotic solution (extract WHO 2010).

No change (a). Different tail changes (b-g). The swelling in the tail region is indicated in gray.

#### **Evaluation**

# Percentage of vitality of spermatozoa

Calculate the difference between the percentage of sperms with swollen flagellum before and after incubation with the SemenHos solution.

#### Example:

#### Before incubation:

1. Count 2/100

2. Count 3/100

Mean value: 2.5 /100= 2.5%

## After incubation:

1. Count 88/100

2. Count 97/100

Mean value: 92.5/100= 92.5%

Result: 92.5% - 2.5%= 90%

# 90 % of the sperms are vital

The SemenHos test is regarded as normal, if after incubation more than 60% of the sperms show a swollen flagellum. The sample is not normal when the result is less than 50% (WHO 2010).

## **Safety information / Precautions**

(Please read also safety data sheets)

- All semen samples should be considered potentially infectious. Handle with all samples like HIV or hepatitis infected material.
- When working with samples and reagents wear always protective clothing (gloves, gowns, eye / face protection).
- All ingredients of reagents are classified as nontoxic

#### References

- Drevius L, Eriksson H, (1966) Osmotic swelling of mammalian spermatozoa, Experimental Cell Research, 42: 136-56
- Jeyendran RS, et al, (1984) Development of an assay to assess the functional integrity of the human sperm membrane and its relationship to the other sperm characteristics, *Journal of Reproduction and Fertility*, 70: 219-28
- WHO Press, (2010) Laboratory manual for the examination and processing of human semen, 5<sup>Th</sup> edition
- Zaneveld LJD, (1984) Journal of Reproduction and Fertility, 70:219-228. 

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consult instructions for use

in vitro diagnostics

Temperature limitation

Lot number

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