



Test report No. 10719hd

EVALUATION OF FUNGICIDAL OR YEASTICIDAL ACTIVITY IN THE MEDICAL AREA (EN 13624)

Name of the product: CHEMISEPT MED

Batch number: 196010519

Date of test report: 24.07.2019

Client, representative:

Chemi-Pharm Ltd.

Tänassilma tee 11

Tänassilma küla

Saku vald, 76406

ESTONIA

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EVALUATION OF FUNGICIDAL OR YEASTICIDAL ACTIVITY IN THE MEDICAL AREA
(EN 13624)

Name of the product: CHEMISEPT MED
Batch number: 196010519
Order number: 19023
Manufacturer: Chemi-Pharm Ltd.
Client, representative: Chemi-Pharm Ltd., Tännassilma tee 11, Tännassilma küla, Saku vald,
76406, ESTONIA
Maris Millner, +372-51-77-090
Date of delivery: 03.07.2019
Test material conditions: No specific features, sample in the manufacturers tare
Storage conditions: In room temperature, dark
Active substance – conc.: Ethyl alcohol 72.5 (w/w) %; isopropyl alcohol (w/w) 7.5%
Appearance of the product: Transparent liquid
Test concentration: Ready to use
Contact time: 60s
Interfering substance: 3.0 g/l bovine albumin = clean conditions
Rinsing liquid: Tryptone 1 g/l + NaCl 9 g/l
Neutralizer: -
Test organisms: *Aspergillus brasiliensis* ATCC 16404
Testing method: EVS-EN 13624:2013
Quantitative suspension test for the evaluation of fungicidal or
yeastocidal activity in the medical area.
Testing date: 16.07.2019 – 18.07.2019
Results: look appendix 1-2




Allar Laaneleht
Chief specialist
Date of test report: 24.07.2019

TEST RESULTS (fungicidal suspension test)

EVS-EN 13624:2013; Phase 2, step 1;
Membrane filtration method;
Rinsing liquid: Tryptone 1 g/l + NaCl 9 g/l;
Test organism: *Aspergillus brasiliensis* ATCC 16404;
Test temperature: +20° C; Incubation temperature: +30° C
Interfering substance: 3.0 g/l bovine albumin = clean conditions;
Nordic Tersus Laboratory LLC.; Date of test: 16.07.2019
Responsible person: Allar Laaneleht

Validation and controls

Clean conditions

Validation suspension N_{vo}			Experimental conditions (A)			Filtration control (B)			Method validation (C)		
V_{C1}	V_{C2}	\bar{x}	V_{C1}	V_{C2}	\bar{x}	V_{C1}	V_{C2}	\bar{x}	V_{C1}	V_{C2}	\bar{x}
61	74	67.5	33	39	36	42	30	36	32	34	33
$30 \leq \bar{x} N_{vo} \leq 160$? yes X; no <input type="checkbox"/>			$\bar{x} A$ is $\geq 0.5 \bar{x} N_{vo}$? yes X; no <input type="checkbox"/>			$\bar{x} B$ is $\geq 0.5 \bar{x} N_{vo}$? yes X; no <input type="checkbox"/>			$\bar{x} C$ is $\geq 0.5 \bar{x} N_{vo}$? yes X; no <input type="checkbox"/>		

Test suspension and test

Testsuspension:	N	V_{C1}	V_{C2}	$\bar{x}_{wm} = 1.79 \times 10^8$; $\log N = 8.25$ $N_0 = N/100$; $\log N_0 = 6.25$ $6.17 \leq \log N_0 \leq 6.70$; yes X; no <input type="checkbox"/>
N and N_0	10^{-6}	194	164	
	10^{-7}	21	15	

Experimental results

Concentration of the product	Dilution step	V_{C1}	V_{C2}	N_a (= \bar{x} *10)	log N_a	logR	Contact time	Conditions
Ready to use	-	<14	<14	<140	<2.15	>4.10	60 s	Clean

Explanations:

V_C = count per ml (one plate or more)
 \bar{x} = average of V_{C1} and V_{C2} (1. + 2. duplicate)
 N = cfu/ml microbes in testsuspension
 N_0 = cfu/ml at the start of the contact time (t=0)
 N_{vo} = cfu/ml in the validation suspension (t=0)
 N_a = surviving microbes after the test
R = reduction factor (R= N_0/ N_a ; LogR=Log N_0 - Log N_a)

Appendix 2

Interpretation:

The ready to use product CHEMISEPT MED (batch no. 196010519) was tested according to the test method EVS-EN 13624:2013. The test was performed at 20 °C ± 1 °C, under clean conditions during contact time of 60s. The membrane filtration method was used for testing the product's effectiveness against the reference strain: *Aspergillus brasiliensis* ATCC 16404. Under clean conditions the tested product was effective against the reference strain within 60 s.

Conclusion:

The surviving count of the reference strain showed at least 4 lg reduction meaning that the ready to use product CHEMISEPT MED has a fungicidal effect under clean conditions within 60 s.



Allar Laaneleht
Chief specialist
24.07.2019