

# Compound Data Sheet

FM140/0

Grey

#### General description

Chlorobutyl compound with silicate filler and inorganic coloring system.

Unconventionally cured, free from MBT.

Universal application range: infusion, injection, lyophilization.

#### Physical properties

	Unit	Method	Target	Range	
		ISO 7619-1			
Hardness	°Shore A	(1 sec. Indentation)	46	± 5	
		Avg of 3 measurements			
Density	g/cm³	ISO 2781	1.323	± 0.025	
Ash	%	Internal Method(s):	45.5	± 2.0	
		Calc. 4h @ 700° C	45.5		
Compression Cot	0/	100.045	25	may	
Compression Set	%	ISO 815	35	max.	
Tensile Strength	N/mm <sup>2</sup>	ISO 37	3	min.	

#### Chemical properties

FM140(\*\*) meets the chemical requirements for Type I Closures specified in General Chapter 3.2.9. of the European Pharmacopoeia and specified in General Chapter <381> of the United States Pharmacopoeia.

Typical USP <381> / EP 3.2.9. data for FM140(\*\*) are presented in the table on page 2.

A typical UV spectrum of the USP <381> / EP 3.2.9. extract of FM140(\*\*) is presented in the figure on page 3.

# Biological properties

FM140(\*\*) is non-cytotoxic and meets the requirements of the Elution Test as described in General Chapter <87> of the United States Pharmacopoeia.

A typical USP <87> Elution Test Certificate is enclosed on page 4.

# **Pyrolysate**

An infrared spectrum of the pyrolysate of FM140(\*\*) is enclosed on page 5.

#### Compound statement

A statement about compound FM140(\*\*) in respect to natural rubber latex, nitrosamines, MCBT, Heavy metals, TSE/BSE and GMO is enclosed on page 6.

(\*\*) Note: FM140 refers to the type of compound, the extension "/0", "/1", ... refers to the colour of the said compound.

Differently coloured compounds might be used for testing throughout this document. It is generally accepted that the colour is irrelevant for the properties discussed in this document.

Prepared by:

**R&D** Laboratory

e: Plugust 16 2010

Reviewed by:

Material Development

: September 9, 2010

Approved by:

Quality Assurance

Date: 15 Sent loss

Valid from: August 16, 2010 (Replaces CDS of January 2, 2002) CDS FM140/0

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# Typical USP <381> / EP 3.2.9. chemical properties of FM140(\*\*)

Characteristic		Amount tested	Units	Limit	Typical Values	
Appearance of solution	Turbidity	Sol. S	NTU	Type I: ≤ 6.0 (*)		0.6
	Colour	Sol. S		See test procedure		pass
Acidity or alkalinity					<u>Blank</u>	
		Sol. S (20 ml)	ml 0.01M HCl	≤ 0.8		
			ml 0.01M NaOH	≤ 0.3	0.06	0.06
					EP	0.06
					USP	0.00
Absorbance		Sol. S	A <sub>max</sub> 220-360nm	Type I: ≤ 0.2		0.02
Reducing substances		Sol. S (20 ml)	ml 0.002M KMnO <sub>4</sub>	Type I: ≤ 3.0		0.4
Extractable heavy metals		Sol. S	ppm Pb	≤ 2	EP	<2
				<u> </u>	USP	<2
Extractable zinc		Sol. S	ppm Zn	≤ 5.0		0.31
Ammonium		Sol. S	ppm NH <sub>4</sub>	≤ 2		<2
Residue on evaporation (only for EP)		Sol. S (50 ml)	mg	Type I: ≤ 2.0		0.2
Volatile sulphides		20 cm <sup>2</sup>	mg S	≤ 0.02		<0.02

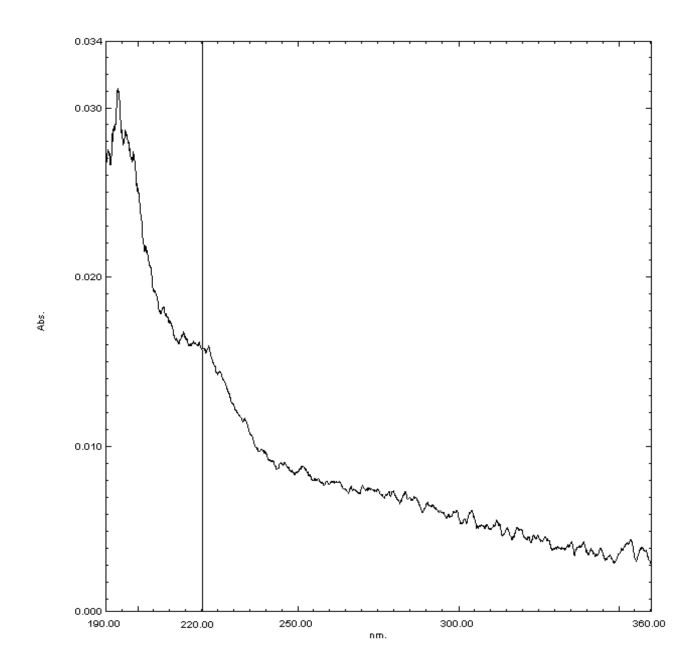
<sup>(\*)</sup> By definition corresponding with reference suspension II.

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# Typical UV-spectrum of USP <381> / EP 3.2.9. extract of FM140(\*\*)



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### Typical USP <87> Elution Test Certificate of FM140(



#### TEST RESULT REPORT N°:10-B0277-N1



Project Number: TE 10118

Study Number:

10-B0277-N1

Sponsor:

Helvoet Pharma Belgium NV Mrs. Nadia Nouri

Report Date:

18/02/2010

Contact:

Industrieterrein Kolmen 1519

Date Sample Arrival:

10/02/2010 15/02/2010

Address:

Study

3570 Alken, Belgium

Technical Initiation: **Technical Completion:**  18/02/2010

PO.Number:

PB1000518

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**Elution Test - ISO** Temp/Time 37°C/24 hours

Test Item	FM140/0 V9025 SAF1	Ratio	25cm²/20mL		
Lot	30194939	Vehicle	MEM-Complete		

REFERENCE: According to "ISO 10993-5, 2009: Biological Evaluation of Medical Devices- Part 5:Tests for In Vitro Cytotoxicity." and "USP 32-NF 27, 2009: <87> Biological reactivity test, in vitro." Toxikon Reference: SOP 3.1.2.3, rev. 08

PROCEDURE: The biological reactivity of a mammalian monolayer, L929 mouse fibroblast cell culture, in response to the test item extract was determined. The samples and control articles were autoclaved prior to the preparation of the extracts. Extracts were prepared at 37±1°C for 24 hours in a humidified atmosphere containing  $5\pm1\%$  carbon dioxide (static). Positive (natural rubber) and negative (silicone) control articles were prepared to verify the proper functioning of the test system. The maintenance medium on the cell cultures is replaced by the extracts of the test item or control article in triplicate and the cultures are subsequently incubated for 48 hours, at 37±1°C, in a humidified atmosphere containing 5±1% carbon dioxide. Biological reactivity was rated on the following scale: Grade 0 (No reactivity); Grade 1 (Slight reactivity), Grade 2 (Mild reactivity), Grade 3 (Moderate reactivity) and Grade 4 (Severe reactivity). The test item is considered non-cytotoxic if none of the cultures exposed to the test item shows greater than mild reactivity (Grade 2).

OPINION AND INTERPRETATION: No reactivity (Grade 0) was exhibited by the cell cultures exposed to the test item at the 48 hours observation. Severe reactivity (Grade 4) was observed for the positive control article. The negative control article showed no signs of reactivity (Grade 0).

OPINION AND INTERPRETATION: Based on the evaluation criteria mentioned above, the test item is considered non-cytotoxic.

RECORD STORAGE: All raw data generated in this study will be archived at Toxikon Europe, according to SOP 4.2.8.

AUTHORIZED PERSONNEL

ir. Peter Cornelis Study Director

Vanessa Ruymen Quality Assurance

The test results on the enclosed report are only referring to the tested articles. Partly reproduction of this report can only be allowed after written permission of Toxikon. Toxikon guarantees that all results are acquired by testing according to officially accepted scientific methodology.

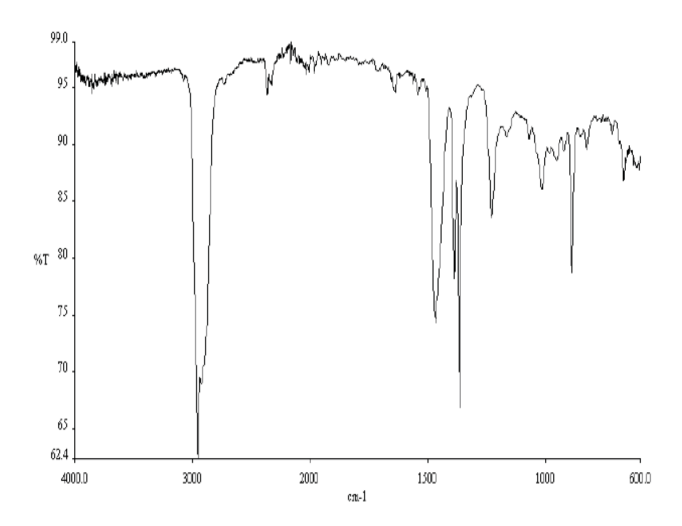
Toxikon Europe nv - Researchpark Haasrode 1724 - Romeinsestraat 12, B 3001 Leuven, Belgium - Tel. 32-16-40 04 84 - Fax 32-16-40 13 04 Fortis 230-0391575-06 - KBC 431-0597001-33 - BTW/TVA BE 0442,395,719 - H.R. Leuven 80.154 - www.toxikon.be

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# Typical infrared spectrum of a pyrolysate (4000-600 cm-1) of FM140(\*\*)



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### Compound statement for FM140(\*\*)

#### Natural rubber latex

Compound FM140(\*\*) is free from natural rubber and natural rubber latex.

#### **Nitrosamines**

Compound FM140(\*\*) does not use rubber chemicals that are associated with hazardous nitrosamines formation.

#### **MCBT**

Compound FM140(\*\*) does not contain 2-mercaptobenzothiazole (MCBT, also named MBT), or any of its derivatives.

#### **Heavy Metals**

- Compound FM140(\*\*) fulfils the European Community Guideline 94/62/EC for heavy metals in packaging materials.
- Compound FM140(\*\*) fulfils the CONEG regulation on reduction of toxics in Packaging Law.

Both directives state that packaging components should not contain more than 100 ppm of Lead (Pb), Cadmium (Cd), Mercury (Hg) and Hexavalent Chromium (VI) (Cr). Where the regulated metals are present at levels below the values stated above, they were not intentionally added during the manufacturing process.

#### TSE/BSE

Compound FM140(\*\*) does not contain material of animal origin and hence is not associated with TSE/BSE risks.

(TSE = Transmissible Spongiform Encephalopathy; BSE = Bovine Spongiform Encephalopathy)

#### **GMO**

Compound FM140(\*\*) does not contain ingredients made from GMO's (Genetically Modified Organisms).

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