Sealing Solutions



COMPOUND DATA SHEET FM27/0 Grey

General description

FM27/0 is a styrene butadiene compound. Conventionally cured, free from MBT.

Identification as per Ph.Eur. 3.2.9., method A

A typical ATR-FTIR spectrum of a clean, cut surface of FM27/0 is enclosed on page 2.

Physical properties

Physical properties	
Hardness	51 ± 5 Shore A
ISO 7619-1(1sec Indentation)	
Density	1.290 ± 0.025 g/cm ³
ISO 2781	
Ash content	43.7 ± 2.0 %
Internal method. Calc.4h@700℃	
Compression set	21 %
ISO 815, (typical value)	
Modulus 100	2.2 N/mm ²
Internal method. 100mm/min,ISO 37 Dumb-bell 2, (typical value)	
Modulus 300	6.0 N/mm ²
Internal method. 100mm/min,ISO 37 Dumb-bell 2, (typical value)	
Elongation @ Break	360 %
Internal method. 200mm/min, ISO 37 Dumb-bell 2, (typical value)	
Tensile Strength	9 N/mm²
Internal method. 200mm/min, ISO 37 Dumb-bell 2, (typical value)	
Water Vapor Transmission Rate	3.9 g/m ² .24h
Mocon, 38°C,100%RH, 1.2 mm thickness, (typical valu e)	J
Oxygen Transmission Rate	971 cc/m ² .24h
Mocon, 38℃, 50%RH, 100% O2, 1.2 mm thickness, (typ ical value)	

Compound ingredient declarations

Latex FM27/0 may contain natural rubber latex which may cause

allergic reactions in latex sensitive individuals. **Nitrosamines** FM27/0 is not made with chemicals that are associated with

nitrosamine formation as per the ASTM F1313-90 list.

Phthalates FM27/0 is not made using Di(2-EthylHexyl) Phthalate (DEHP)

or other phthalates. BSF/TSF

For the raw materials of FM27/0, certification confirming that such products are either of vegetable origin or are manufactured in severe process conditions for inactivation of prions as described in the EMA/410/01-rev.3 and in the European Pharmacopoeia 5.2.8. "Minimizing the risk of transmitting Animal Spongiform Encephalopathy Agents" is

available.

MBT FM27/0 is not made with 2-mercapto-benzothiazole (MCBT,

also named MBT), or any of its derivatives.

FM27/0 fulfills the EC Guideline 94/62/EC for heavy metals in **Heavy Metals** packaging materials and the CONEG regulation on reduction of toxics in Packaging Law: "packaging components shall not

contain more than 100 ppm of Pb, Cd, Hg, and Cr(VI)".

Biocompatibility

FM27/0 is non-cytotoxic and meets the requirements of the Elution Test as described in General Chapter <87>,"Biological Reactivity Tests, in vitro" of the USP. A typical USP<87> Elution Test certificate is enclosed on page 3.

Chemical properties

FM27/0 meets the chemical requirements for Type I closures specified in Ph.Eur. 3.2.9 and in USP <381>.

Typical results are given in the table below.

A typical UV spectrum is enclosed on page 2.

Characte	eristic	Limits		FM27/0
Appearance of solution	Turbidity	Type I: 6 NTU (*) Type II: 18 NTU (*)		2
	Color	Solution S is not more intensely colored than reference		Pass
		0.8 ml 0.01M HCl	EP	NA
Acidity or alkalinity			USP°	NA
(NA : Not applicat	ole)	0.3 ml 0.01M NaOH	EP	0.0
		0.3 IIII 0.0 IIVI NaOri	USP°	0.0
UV Absorbance (max 220-360 nm)		Type I: 0.2 Type II: 4.0		0.1
Reducing substances		Type I: 3.0 ml 0.01M $Na_2S_2O_3$ Type II: 7.0 ml 0.01M $Na_2S_2O_3$		0.9
Extractable heavy metals		2 ppm Pb ²⁺	EP USP	Pass Pass
Extractable zinc		5.0 ppm Zn ²⁺		0.5
Ammonium		2 ppm NH ₄ ⁺		Pass
Residue on evapo (only for EP)	oration	Type I: 2.0 mg Type II: 4.0 mg		0.2
Volatile sulphides		Any black stain on the paper is not more intense than that produced by a control solution		Pass

^(*) By definition corresponding with reference suspensions II and III (for Ph.Eur.) or B and C (for USP) respectively

Note: "FMxxx" refers to the type of compound, the extension "/x" refers to the color of the said compound. Differently colored compounds might be used for testing throughout this document. It is generally accepted that the color is inclevant for the properties discussed, except Ash content and Density which are different per color.

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(replaces CDS of October 21, 2010)

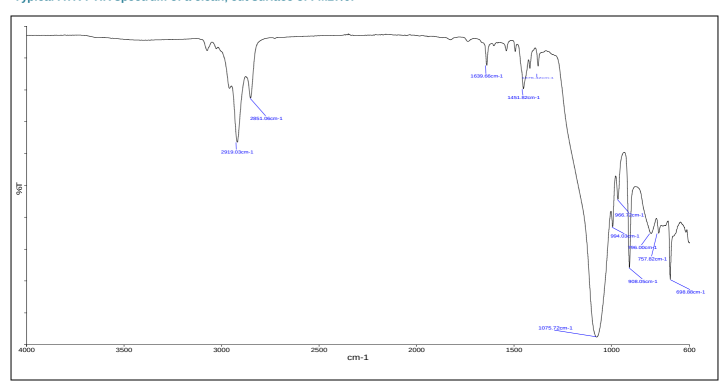
⁽⁹ corrected with blank

Note : Chemical testing is performed on non-post-treated standard test plates or products, prepared using representative process conditions.

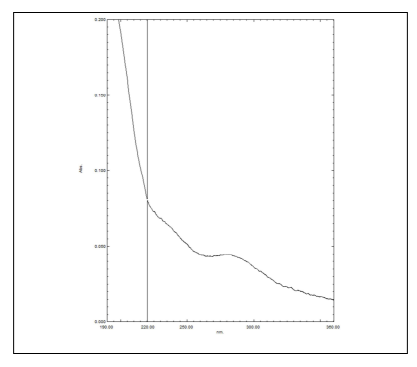


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Typical ATR-FTIR spectrum of a clean, cut surface of FM27/0:



Typical UV spectrum of the Solution S extract of FM27/0, measured as per the USP <381> / Ph.Eur. 3.2.9.:



Version: January 1, 2014 (replaces CDS of October 21, 2010)



COMPOUND DATA SHEET FM27/0 Grey

USP<87> Elution Test certificate for FM27/0:



TEST RESULT CERTIFICATE

08-B0508-N1 Study Number: Project Number: TE 08211 Helvoet Pharma Belgium NV 20/03/2008 Report Date: Sponsor:

Address

Industrieterrein Kolmen 1519 B-3570 Alken

Technical Initiation:

PO.Number:

Belgium PB0800805

Technical Completion:

17/03/2008 20/03/2008

Study	Elution Test - ISO	Temp/Time	37°C/24 hours
Test Item	P8568A FM27 ISAFNS 1/2 domed Not treated	Ratio	25 cm²/20 mL
Lot	Ch751915	Vehicle	MEM-Complete

REFERENCE: Based on "ISO 10993-5, 1999: Biological Evaluation of Medical Devices- Part 5:Tests for In Vitro Cytotoxicity." and "USP 30-NF 25, 2007: <87> Biological reactivity test, in vitro." Toxikon Reference: SOP 3.1.2.3, rev. 05

PROCEDURE: The biological reactivity of a mammalian monolayer, L929 mouse fibroblast cell culture, PROCEDURE: The biological reactivity of a mammalian monolayer, L929 mouse fibroblast cell culture, in response to the test item extract was determined. Extracts were prepared at 37±1°C for 24 hours in a humidified atmosphere containing 5±1% 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). cultures exposed to the test item shows greater than mild reactivity (Grade 2).

RESULTS: 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)

CONCLUSION: 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

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