



medical<sup>®</sup>  
leather

a healthy focus on feet

## Product Specific Information Document

<b>Product name:</b>	<b>Sheep fur lining</b>
Item code:	SH
Barcode:	N/A
CE Marking:	N/A
MDR Risk Classification:	Riskclass I
Product summary:	Sheep fur lining
Size / Contents product:	7-9 sqft
Packaging size:	hide
Packaging unit:	per hide
Color:	414 beige, 415 taupe, 417 black
Thickness:	1,5 - 2,0mm
Shorevalue:	N/A
Implementation:	non-perforated
Storage advise:	not in direct sunlight
Maintenance advise:	This leather can be cleaned with a mild, non alcoholic, cleaning substance
Sterilisation advise:	N/A
Manufacturing date:	N/A
Shelf life:	N/A

**Postbus 95, 5140 AB Waalwijk • Zanddonkweg 6, 5144 NX Waalwijk • The Netherlands**  
**Telefoon** +31 (0)416 37 69 87 • **Fax** +31(0)416 37 56 02 • **E-mail** info@medical-leather.nl •  
**Website** www.medical-leather.nl **Rabobank** 1308.95.377 • **BIC Code** RABONL2U • **IBAN nr.** NL63 RABO 0130 8953 77 •  
**KVK** 18124129 • **BTW nr.** NL8145 15 745 B01



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## DECLARATION OF CONFORMITY

The undersigned  
C.J. Maas – Leder LEFA BV. / Medical Leather  
In reference  
To the REACH Regulation 1907/2006/CE  
Concerning the Registration, Evaluation, Authorization  
And the restriction of chemical substances

### DECLARES THAT

- Our company purchases and sells foam sheets through the transformation of polymers, mineral fillers, pigments and chemicals. Therefore, as a user, our company is not bound to any registration
- According to the REACH Regulation polymers are exempted from registration (article 2) as well as most of the raw materials used.
- Through the raw material suppliers and sheet suppliers, we were able to ascertain compliance with the REACH legislation. In fact, the chemical manufacturers, where provided, have taken steps as for regular registration.
  - With regard to the presence of hazardous substances (SVHC) mentioned in **the last list published by ECHA on 25/06/2020** we declare that our products, after curing, as supplied by us, has no hint of SVHC in a concentration above the threshold limit of 0.1%..

Pieve del Cairo, 22/07/2020

REACH Responsible  
Pieter Maas

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Website [www.medical-leather.nl](http://www.medical-leather.nl) Rabobank 1308.95.377 • BIC Code RABONL2U • IBAN nr. NL63 RABO 0130 8953 77 • KVK 18124129 • BTW nr. NL8145 15 745 B01

**Expertise:** 1910787-02-30-01

Datum/ Date: 2019-12-05  
Dr. Anderie

**Orderer:** Medical Leather

Zanddonkweg 6  
5144 NX Waalwijk  
The Netherlands

**Order:** Examination of materials

Your letter/Fax/Email dated 2019-12-02

**48h**

Received: 2019-12-03

**Test item:** Lamb fur, Article 190007

**Test results (Test item as received):**

Description of procedure on the following pages.


Parameter	Unit	Result *	Material	Rating
Chromium VI	mg/kg	5.5	EP	2
Formaldehyde Leather	mg/kg	33	EP	0

**Rating:**

0 = Sample meets the requirements  
1 = Further test necessary  
2 = Sample failed the requirements  
- = not tested or material not available

**Remarks:**

n.n = not detectable (n.d.), MP = mixed materials, EP = single material test, SEP = see detailed results  
\* Explicit given results for mixed materials (MP) are the total content of the sample. For values with sign of inequality the content of each material of the mixed sample is below the given value



Dr. I. Anderie  
(Abteilungsleitung / Department Management)



Dr. K. Schulte  
(Institutsleitung / Managing Director)



Die Prüfergebnisse beziehen sich ausschließlich auf die Prüfgegenstände. Ohne schriftliche Genehmigung des PFI darf die Expertise nicht auszugsweise verwendet werden. Restliches Untersuchungsmaterial wird nach 10 Wochen vernichtet. In Einzelfällen kann eine Untervergabe an andere geeignete Prüflaboratorien erfolgen. Ein Verzeichnis der akkreditierten Prüfverfahren und unsere geltenden Allgemeinen Geschäftsbedingungen können auf unserer Internetseite eingesehen werden und werden Ihnen auf Anfrage gerne zugesandt. / Test results solely refer to the items tested. Extracts of the expertise may not be used without written permission by PFI. Remaining materials will be destroyed after 10 weeks. In individual cases, tests can be carried out by qualified subcontracting laboratories. A list of accredited test methods and our applying general terms and conditions are available on our website and will be sent to you on request.

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**Expertise: 1910787-02-30-01**

Datum/ Date: 2019-12-05  
Dr. Anderie

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**Examination Procedure:**

Test methods are part of the laboratory accreditation (D-PL-14150-01-00)

**Shortened form: Chromium VI**

**Determination of Chromium VI in leather**

DIN EN ISO 17075-2:2017-05 and BVL B 82.02-11:2008-10

**Shortened form: Formaldehyde Leather**

**Determination of Formaldehyde content in leather by High Performance Liquid Chromatography**

EN ISO 17226-1:2019

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