



5. July 2016 Version 3.0

Material Safety Data Sheet

Based on template version 5.0

Identification of the substance/mixture and of the company/undertaking

Product name: Biatain Silicone Ag

Product code: 39636, 39637, 39638, 39639, and 39640

Product information: Wound care product

Manufacturer: Coloplast A/S

Holtedam 1

DK-3050 Humlebaek

Denmark

Telephone +45 49111111 msds@coloplast.com

Hazards identification

This product consists primarily of polymer materials. The products pose no immediate hazard.

Composition/information on ingredients

This product is regulated as a medical device in European Economic Area (EEA). In other regions it may be regulated as a medical device, a cosmetic or not regulated.

This product does not contain substances classified as hazardous under EC Regulation No. 1272/2008/EC, Annex VI (EU) in concentrations above 0.1 % (w/w). Main ingredients and packaging materials are listed below.

Chemical name CAS no.

Wound dressing:

Polyurethane film 9009-54-5 Polyurethane foam 9009-54-5

Silver complex* -

USA

Coloplast Corp. 1601 West River Road N Minneapolis, MN 55411 Telephone: +1-800-533-0464 www.us.coloplast.com

Canada

Coloplast Canada Corporation 3300 Ridgeway Drive, Unit 12 Mississauga, Ont. L5L 5Z9 Telephone: +1-877-820-7008 www.coloplast.ca

Europe

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DK-3050 Humlebaek Telephone: +45 49 11 11 11

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Biatain Silicone Ag

Surfactant -

Silicone adhesive

Polyethylene film 9002-88-4

Packaging:

PE-coated paper

Polyester/ethylene vinyl acetate film

*: Content corresponds to a target of 0.95 mg silver per cm²

Disposal considerations

Dispose the device according to the recommended disposal technology at any approved facility. Under normal private use the product may be disposed of together with other household waste unless local regulations set special requirements. The disposal should always be in compliance with national, federal, state and local regulations. The product should not be discharged to the environment.

US

This product does not meet the criteria for hazardous waste as defined under the Resource Conversation and Recovery Act (RCRA) 40 CFR 261. Under normal private use the product may be disposed of together with other household waste per RCRA 40 RFT 261.4.B1.

European Union

Per The European Waste Catalogue (EWC), in accordance with EC Directive 75/422/ECC, the following Waste Code can be used: 18 01 04 00 wastes whose collection and disposal is not subject to special requirements in view of the prevention of infection (e.g. dressings, plaster casts, linen, disposable clothing, diapers). However, if the waste in view of the prevention of infection needs special requirements, other Waste Codes should be used. Under normal private use the product may be disposed of together with other household waste unless local regulations set special requirements.

Handling and storage

Handling: See instruction for use

Storage: Store until use as supplied and at room temperature un-

less other information is stated on the packaging or on the

leaflet.

Other information

This MSDS is supplied as an additional service to the customer. The product is a medical device, which is regulated under the Council Directive 93/42/ECC, Medical Device Directive. The product has been evaluated according to the requirements of medical devices. According to current knowledge this product is considered non-toxic. For further information please contact Coloplast A/S.

Signature Page for VV-0097605 v2.0

Approved by	DKPLYM Peter Lynge Madsen Senior Biosafety Specialist
	Technical / Specialist
	05-Jul-2016 11:48:04 GMT+0000

Signature Page for VV-0097605 v2.0

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