

Declaration of Conformity EU

Document ID: Created by: Approved by: Approval date:

Title: Mepilex Border Flex Lite

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We, Mölnlycke Health Care AB, Gamlestadsvägen 3C, Box 13080, SE-402 52 Göteborg, Sweden being exclusively responsible manufacturer for conformity of the device/s; declare that the devices listed in the attached schedule are in conformity with the provisions of the Council Directive 93/42/EEC of 14 June 1993, as amended by 2007/47/ EEC, concerning medical devices, the Medical Devices Act SFS 1993:584 and the Swedish Medical Product Agency regulations and guidelines: Medical Devices, LVFS 2003:11, as amended by LVFS 2009:18.

Trade name/ Product name:	Mepilex Border Flex Lite	
Product classification:	llb	
Sterility Status:	Sterile	
Measuring function:	Νο	
This declaration is supported by a conformity assessment procedure in accordance with		
Annex/es:	II	
Certificate number:	CE 01965	
Issued by:	BSI (2797)	
For non sterile, non-measuring Class I products, no certificate is issued by a Notified Body.		

Mölnlycke Health Care issues this declaration in recognition of all applicable harmonised standards.

Signed for and on behalf of Mölnlycke Health AB Care

Date: 2021-02-26

Function:

Regulatory Affairs Manager Compliance

Name:

Karin Darle Olsson

Signature:

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Title: Mepilex Border Flex Lite

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Product(s) covered by this declaration:

Product Reference:	Product Descriptor:	GMDN Code:
581011	Self-adherent soft silicone foam dressing	46854 Wound - nonadherent dressing, absorbent, sterile
581100	Self-adherent soft silicone foam dressing	46854 Wound - nonadherent dressing, absorbent, sterile
581200	Self-adherent soft silicone foam dressing	46854 Wound - nonadherent dressing, absorbent, sterile
581300	Self-adherent soft silicone foam dressing	46854 Wound - nonadherent dressing, absorbent, sterile
581500	Self-adherent soft silicone foam dressing	46854 Wound - nonadherent dressing, absorbent, sterile