

EU – Declaration of Conformity

We, the company,

Beiersdorf AG
SRN DE-MF-000014151

20245 Hamburg
Germany

hereby declare in our sole responsibility as legal manufacturer that the device(s)

48657-00000-46	EPL EL_FXN_BDG 4MX6CM 1 PCS EN
48657-00002-45	HP EL_FXN_BDG 4MX6CM 1 PCS ID
48657-00002-46	HP EL_FXN_BDG 4MX6CM 1 PCS ID
48658-00000-45	HP EL_FXN_BDG 4MX8CM 1 PCS DE_PT
48658-00001-45	HP EL_FXN_BDG 4MX8CM 1 PCS ID
48658-00002-46	EPL EL_FXN_BDG 4MX8CM 1 PCS EN
48658-00003-45	HP EL_FXN_BDG 4MX8CM 1 PCS DE_FR_NL

Product Identification:	Hansaplast Elastic Fixation Bandage
Basic-UDI-DI:	40058000000000000000000024CV
Intended Use:	For the fixation and protection of e.g. wound pads or wound dressings

is/are in compliance with Regulation (EU) 2017/745 concerning medical devices.

Classification

according to Annex VIII of the Regulation (EU) named above.

class I

Conformity Assessment Procedure

according to Article 52(7) and Annex IV of the Regulation (EU) named above.

Notified Body

not applicable for medical device class I.

This declaration is also valid for products which may carry additional labels to comply with local market and regulatory requirements (relabeling).

Hamburg (Germany), 28.08.2024
(date)



Martina Fraenkel
Regulatory Affairs Manager
(Signed for and on behalf of Beiersdorf AG)