

Beiersdorf	Beiersdorf international documentation system BEC.10245895.000.03 DoC Wound Spray Forward	Released: 12.05.21-... Page 1 of 2
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We,

Beiersdorf AG

Unnastr. 48
20253 Hamburg

are exclusively responsible for this declaration of conformity and hereby declare, that the following products
(article number refers to the smallest sales unit of the product)

48387-00000-45	HP WND_SPY PHMB 0.04% 15ML DE
48387-00001-45	HP WND_SPY PHMB 0.04% 15ML IT
48387-00002-45	CURI WND_SPY PHMB 0.04% 15ML ES
48387-00003-45	HP WND_SPY PHMB 0.04% 15ML ES
48388-00000-45	HP WND_SPY PHMB 0.04% 50ML DE
48388-00001-45	HP WND_SPY PHMB 0.04% 50ML DE_FR_IT
48388-00003-45	EPL WND_SPY PHMB 0.04% 50ML EN
48388-00004-45	EPL WND_SPY PHMB 0.04% 50ML FR
48388-00005-45	CURI WND_SPY PHMB 0.04% 50ML ES
48388-00006-45	HP WND_SPY PHMB 0.04% 50ML DE EUROHANGER
48388-00007-45	HP WND_SPY PHMB 0.04% 50ML DE_FR_NL
48388-00008-45	HP WND_SPY PHMB 0.04% 50ML ES
48388-00009-45	EPL WND_SPY PHMB 0.04% 50ML EN
48388-00011-45	HP WND_SPY PHMB 0.04% 50ML DE_FR_NL
48389-00000-45	HP WND_SPY PHMB 0.04% 100ML DE_FR_NL
48389-00002-45	HP WND_SPY PHMB 0.04% 100ML PT
48389-00003-45	HP WND_SPY PHMB 0.04% 100ML EL_IT
48389-00004-45	HP WND_SPY PHMB 0.04% 100ML AR_EN_MS
48389-00005-45	HP WND_SPY PHMB 0.04% 100ML INT
48389-00006-45	EPL WND_SPY PHMB 0.04% 100ML EN
48389-00009-45	EPL WND_SPY PHMB 0.04% 100ML EN
48389-00010-45	HP WND_SPY PHMB 0.04% 100ML DE_FR_NL
48389-00011-45	EPL WND_SPY PHMB 0.04% 100ML EN_FR
48390-00001-45	HP WND_SPY_KDS PHMB 0.04% 100ML DE_FR_NL
48390-00002-45	HP WND_SPY_KDS PHMB 0.04% 100ML EL_ES
48390-00002-47	HP WND_SPY_KDS PHMB 0.04% 100ML EL_ES_PT
48390-00003-45	HP WND_SPY_KDS PHMB 0.04% 100ML DE_FR_NL
48759-00000-45	HP WND_SPY PHMB 0.04% 250ML DE

within the

class IIb

comply with

Medical Device Directive 93/42/EEC, annex I.

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

Conformity assessment procedure follows

Medical Device Directive 93/42/EEC, annex II.

Notified Body for Conformity Assessment Procedure:

DEKRA Certification GmbH
Handwerkstraße 15
70565 STUTTGART
Country: Germany

Notified Body number: 0124

20253 Hamburg (Germany)
(place of issuance)

12.05.2021
(date)


Hedda Siebrecht
Senior Regulatory Affairs Manager

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valid until 26.05.2024