



湖北省潜江市江赫医用材料有限公司
HUBEI QIANJIANG KINGPHAR MEDICAL MATERIAL CO, LTD

Qianjiang, 2023-06-16

Declaration of Conformity

We

Hubei Qianjiang Kingphar Medical Material Co., Ltd.
Yuanguang Road, 433100, Qianjiang, P. R. China
SRN (Single Registration No): CN-MF-000009051

Manufacturer according to Medical Device Regulation (EU) 2017/745

hereby declare in our own responsibility
that the products

Askina® Mullkompressen

Non-sterile gauze compresses are ideally used for primary care of acute wounds, for the absorption of blood and exudate and for cleansing the skin's surface. For single use.

Basic UDI-DI: 694878811001PN
(article numbers see attachment I)

are in conformity with the requirements of the Medical Device Regulation (EU) 2017/745

Conformity Assessment Procedure

according to article 52 section 7
of the Regulation named above

Classification

according to annex VIII of the Regulation named above
Class I


Authorized Representative for the European Union (Article 11)

Shanghai International
Holding Corp. GmbH (Europe)
Eiffenstraße 8
20537 Hamburg
Germany

Valid until 2026-06-15

Name: Wang Zhongxiao

Function: President

Signature: 



湖北省潜江市江赫医用材料有限公司
HUBEI QIANJIANG KINGPHAR MEDICAL MATERIAL CO, LTD

Attachment I

Art. No.	Product name	Class
9031308	Askina® Mullkompressen	I
9031316	Askina® Mullkompressen	I
9031324	Askina® Mullkompressen	I
9033017	Askina® Mullkompressen	I
9033025	Askina® Mullkompressen	I
9033041	Askina® Mullkompressen	I