

**KONFORMITÄTSERKLÄRUNG / DECLARATION OF CONFORMITY**Name und Adresse der Firma /  
*Name and address of the company***Kulzer GmbH**  
Leipziger Straße 2, 63450 Hanau  
Deutschland / Germany

SRN: DE-MF-000007705

**Wir erklären in alleiniger Verantwortung, dass / We declare under our sole responsibility that**das Medizinprodukt / *the medical device***GLUMA Desensitizer**Bezeichnung, Typ oder Modell, Chargen- oder  
Seriennummer, ev. Herkunft und Stückzahl / *Name,  
type or model, batch or serial number, possibly  
sources and number of items*Artikelliste siehe Anhang / *List of Articles see Annex*EMDN-Nummer / *EMDN-Code*  
GMDN-Nummer / *GMDN code*  
UMDNS-Nummer / *UMDNS code*  
Basis-UDI-DI / *Basic UDI-DI*Q019008  
64794  
10-034  
++J0141103DES010104e5Jder Klasse / *of class*

IIa

nach Regel / *according to rule*8 nach Anhang VIII der Medizinprodukte-Verordnung, 2017/745 /  
*according to Annex VIII of Medical Device Regulation 2017/745***allen Anforderungen der Medizinprodukte-Verordnung 2017/745 entspricht, die anwendbar sind /  
*meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.***Angewandte harmonisierte Normen, nationale  
Normen oder andere normative Dokumente /  
*Applied harmonised standards, national standards  
or other normative documents*Angewandte Normen siehe Version 02 der Technischen  
Dokumentation von Produkt GLUMA Desensitizer / *Applied  
standards see Technical Documentation of Product GLUMA  
Desensitizer, Version 02*Konformitätsbewertungsverfahren nach /  
*Conformity assessment procedure acc. to*Medizinprodukte-Verordnung 2017/745 Anhang IX, Kapitel I,  
Abschnitt 2 und 3 and Kapitel III  
*Medical Device Regulation 2017/745 Annex IX, Chapter I, Section 2  
and 3 and Chapter III*Benannte Stelle / *Notified Body*TÜV Rheinland LGA Products GmbH  
Tillystrasse 2  
90431 Nürnberg / Germany

CE 0197

Registrierungsnr. / *Registration No.:*

HZ 1198082-1

Versionsnummer / *Version number*

02

Ersetzt Konformitätserklärung vom /  
*Replaces Declaration of Conformity from*

01.12.2022

Hanau, 25.11.2024

i.V.   
Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**Ort, Datum / *Place, date*Name und Funktion / *Name and function*Diese Konformitätserklärung ist gültig für 2 Jahre in Verbindung mit den Freigabe-Dokumenten für die jeweilige Charge der  
produzierten Medizinprodukte / *This statement of conformity is valid for 2 years in connection with the release documents for the  
respective batch of produced devices.*

**Artikelliste / List of Articles**  
**Anhang zur Konformitätserklärung / Annex to declaration of conformity**

das Medizinprodukt /  
*for the medical device*

**GLUMA Desensitizer**

Versionsnummer Artikelliste/  
*Version number article list*

**02**

Ersetzt Artikelliste vom /  
*Replaces article list from*

**01.12.2022**

Diese Artikelliste ist gültig für die  
Konformitätserklärung Version/ *This article list is  
valid for the declaration of conformity version*

**02**

| UDI-DI /<br><i>UDI-DI</i> | Artikelnummer /<br><i>Article number</i> | Name /<br><i>Name</i>                    |
|---------------------------|------------------------------------------|------------------------------------------|
| +J014658723540            | 65872354                                 | GLUMA DESENSITIZER 1 X 5 ML              |
| +J014660018540            | 66001854                                 | GLUMA DESENSITIZER SINGLE DOSE 40 X 1 ST |
| +J014660037640            | 66003764                                 | GLUMA DESENSITIZER 1 X 5 ML REGIO        |
| +J014660182210            | 66018221                                 | GLUMA DESENSITIZER VALUE PACK 3 X 5 ML   |

Hanau, 25.11.2024

Ort, Datum / *Place, date*

i.V.  
Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**



**Name und Funktion / *Name and function***

**OVERENSSTEMMELSESERKLÆRING / DECLARATION OF CONFORMITY**Virksomhedens navn og adresse /  
*Name and address of the company***Kulzer GmbH**  
Leipziger Straße 2, D-63450 Hanau  
Tyskland / Germany  
SRN: DE-MF-000007705**Vi erklærer hermed på eget ansvar, at / We declare under our sole responsibility that**  
det medicinske udstyr / *the medical device* **GLUMA Desensitizer**Betegnelse, type eller model, batch- eller  
serienummer samt eventuelt oprindelse og antal  
emner / *Name, type or model, batch or serial  
number, possibly sources and number of items*Produktlisten kan ses i bilaget / *List of Articles see Annex*EMDN-kode / *EMDN-Code*  
GMDN-kode / *GMDN code*  
UMDNS-kode / *UMDNS code*  
Grundlæggende UDI-DI / *Basic UDI-DI*Q019008  
64794  
10-034  
++J0141103DES010104e5Ji klasse / *of class*

IIa

i henhold til artikel / *according to rule*8 i bilag VIII i Europa-Parlamentets og Rådets forordning (EU)  
2017/745 om medicinsk udstyr / *according to Annex VIII of  
Medical Device Regulation 2017/745***lever op til alle de relevante bestemmelser i forordning (EU) 2017/745 om medicinsk udstyr. /  
*meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.***Anvendte harmoniserede standarder, nationale  
standarder eller andre normative dokumenter /  
*Applied harmonised standards, national standards  
or other normative documents*Andre anvendte standarder kan ses i det tekniske  
dokumentationsmateriale til produktet GLUMA Desensitizer,  
version 02  
*Further Applied standards see Technical Documentation of  
Product GLUMA Desensitizer, Version 02*Overensstemmelsesvurderingsprocedure iht. /  
*Conformity assessment procedure acc. to*Forordning (EU) 2017/745 om medicinsk udstyr, bilag IX, kapitel I,  
afsnit 2 og 3 samt kapitel III  
*Medical Device Regulation 2017/745 Annex IX, Chapter I, Section  
2 and 3 and Chapter III*Underrettet organ / *Notified Body*TÜV Rheinland LGA Products GmbH  
Tillystrasse 2  
D-90431 Nürnberg, Tyskland

CE 0197

Registreringsnummer / *Registration number:*

HZ 1198082-1

Versionsnummer / *Version number*

02

Erstatter overensstemmelseserklæring fra /  
*Replaces Declaration of Conformity from*

01.12.2022

Hanau, 25.11.2024

på vegne af Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**Sted, dato / *Place, date*Navn og stilling / *Name and function*Denne konformitetserklæring gælder i 2 år i forbindelse med frigivelsesdokumenterne for det aktuelle parti af produceret  
medicinsk udstyr. / *This statement of conformity is valid for 2 years in connection with the release documents for the respective  
batch of produced devices.*

**Artikelliste / List of Articles**  
**Bilag / Annex: Overensstemmelseserklæring / Declaration of Conformity**

Det medicinske udstyr /  
*The medical device*

**GLUMA Desensitizer**

Versionsnummer / *Version number*

**02**

Erstatter bilag fra /  
*Replaces Annex from*

**01.12.2022**

Denne artikelliste er gyldig i forbindelse med  
overensstemmelseserklæringen version /  
*This article list is valid for the declaration of  
conformity version*

**02**

| UDI-DI /<br><i>UDI-DI</i> | Artikelnummer /<br><i>Article number</i> | Name /<br><i>Name</i>                    |
|---------------------------|------------------------------------------|------------------------------------------|
| +J014658723540            | 65872354                                 | GLUMA DESENSITIZER 1 X 5 ML              |
| +J014660018540            | 66001854                                 | GLUMA DESENSITIZER SINGLE DOSE 40 X 1 ST |
| +J014660037640            | 66003764                                 | GLUMA DESENSITIZER 1 X 5 ML REGIO        |
| +J014660182210            | 66018221                                 | GLUMA DESENSITIZER VALUE PACK 3 X 5 ML   |

Hanau, 25.11.2024

på vegne af Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

  
Navn og stilling / *Name and function*

**DECLARACIÓN DE CONFORMIDAD / DECLARATION OF CONFORMITY**

Nombre y dirección de la empresa /  
*Name and address of the company*

**Kulzer GmbH**  
Leipziger Straße 2, 63450 Hanau  
Alemania / Germany  
SRN: DE-MF-000007705

**Declaramos bajo nuestra exclusiva responsabilidad que / We declare under our sole responsibility that**  
el producto sanitario / *the medical device*

**GLUMA Desensitizer**

Nombre, tipo o modelo, lote o número de serie,  
posiblemente fuentes y número de elementos /  
*Name, type or model, batch or serial number,*  
*possibly sources and number of items*

Lista de artículos en el Anexo / *List of Articles see Annex*

Código EMDN / *EMDN-Code*  
Código GMDN / *GMDN code*  
Código UMDNS / *UMDNS code*  
UDI-DI básico / *Basic UDI-DI*

Q019008  
64794  
10-034  
++J0141103DES010104e5J

de la clase / *of class*

Ila

de acuerdo con la norma / *according to rule*

8 de acuerdo con el Anexo VIII del Reglamento sobre productos  
sanitarios 2017/745 / *according to Annex VIII of Medical Device*  
*Regulation 2017/745*

**cumple todas las disposiciones del Reglamento sobre productos sanitarios 2017/745 que se le aplican. /**  
***meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.***

Normas armonizadas, normas nacionales u otros  
documentos normativos que se aplican / *Applied*  
*harmonised standards, national standards or other*  
*normative documents*

Para otras normas aplicadas consulte la documentación técnica del  
producto GLUMA Desensitizer, versión 02  
*Further Applied standards see Technical Documentation of*  
*Product GLUMA Desensitizer, Version 02*

Procedimiento de evaluación de la conformidad de  
acuerdo con /  
*Conformity assessment procedure acc. to*

Reglamento sobre productos sanitarios 2017/745 Anexo IX,  
Capítulo I, Secciones 2 y 3 y Capítulo III  
*Medical Device Regulation 2017/745 Annex IX, Chapter I, Section 2*  
*and 3 and Chapter III*

Organismo notificado / *Notified Body*

TÜV Rheinland LGA Products GmbH  
Tillystrasse 2  
90431 Nürnberg / Alemania

CE 0197

Número de registro / *Registration number:*

HZ 1198082-1

Número de versión / *Version number*

02

Sustituye a la declaración de conformidad del /  
*Replaces Declaration of Conformity from*

01.12.2022

Hanau, 25.11.2024

i.V. Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

Lugar, fecha / *Place, date*

Nombre y cargo / *Name and function*

La presente declaración de conformidad tendrá una validez de 2 años según la documentación emitida para el correspondiente  
lote de productos fabricados. / *This statement of conformity is valid for 2 years in connection with the release documents for the*  
*respective batch of produced devices.*

**Lista de artículos / *List of Articles***  
**Anexo / *Annex: Declaración de conformidad / Declaration of Conformity***

El producto sanitario /  
*The medical device*

Número de versión / *Version number*

Sustituye al Anexo del /  
*Replaces Annex from*

Esta lista de artículos es válida para la  
versión de la declaración de conformidad /  
*This article list is valid for the declaration of  
conformity version*

**GLUMA Desensitizer**

**02**

**01.12.2022**

**02**

| UDI-DI /<br><i>UDI-DI</i> | Artikelnummer /<br><i>Article number</i> | Name /<br><i>Name</i>                    |
|---------------------------|------------------------------------------|------------------------------------------|
| +J014658723540            | 65872354                                 | GLUMA DESENSITIZER 1 X 5 ML              |
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| +J014660037640            | 66003764                                 | GLUMA DESENSITIZER 1 X 5 ML REGIO        |
| +J014660182210            | 66018221                                 | GLUMA DESENSITIZER VALUE PACK 3 X 5 ML   |

Hanau, 25.11.2024

Lugar, fecha / *Place, date*

i.V. Dr. Matthias Hartmann

Head of Global Quality, Regulatory & Scientific Services

**Kulzer GmbH**



**Nombre y cargo / *Name and function***

**VAATIMUSTENMUKAISUUSVAKUUTUS / DECLARATION OF CONFORMITY**

Yhtiön nimi ja osoite /  
Name and address of the company

**Kulzer GmbH**  
Leipziger Straße 2, 63450 Hanau  
Saksa / Germany  
SRN: DE-MF-000007705

**Vakuutamme yksinomaisella vastuullamme, että / We declare under our sole responsibility that**

lääkinnällinen laite / the medical device

**GLUMA Desensitizer**

Laitteen nimi, tyyppi tai malli, erä- tai sarjanumero,  
mahdolliset lähteet ja lukumäärä / Name, type or  
model, batch or serial number, possibly sources and  
number of items

Artikkeliluettelo, ks. liite / List of Articles see Annex

EMDN-koodi / EMDN-Code  
GMDN-koodi / GMDN code  
UMDNS-koodi / UMDNS code  
Perus-UDI-DI / Basic UDI-DI

Q019008  
64794  
10-034  
++J0141103DES010104e5J

luokka / of class

IIa

säädös / according to rule

8 lääkitämisistä laitteista annetun asetuksen 2017/745 liitteen VIII  
mukaan / according to Annex VIII of Medical Device Regulation  
2017/745

**täyttää kaikki lääkitämisistä laitteista annetun asetuksen 2017/745 soveltuvat vaatimukset. /  
meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.**

Soveltuvat harmonisoidut standardit, kansalliset  
standardit tai muut säädökset / Applied harmonised  
standards, national standards or other normative  
documents

Muut sovellettavat standardit, ks. tekniset tiedot  
tuotteesta GLUMA Desensitizer, versio 02  
Further Applied standards see Technical Documentation of  
Product GLUMA Desensitizer, Version 02

Vaatimustenmukaisuuden arviointimenettelyn  
perusta /  
Conformity assessment procedure acc. to

Asetus lääkitämisistä laitteista 2017/745, liite IX, I luku, 2 ja  
3 kohta ja III luku  
Medical Device Regulation 2017/745 Annex IX, Chapter I,  
Section 2 and 3 and Chapter III

Ilmoitettu laitos / Notified Body

TÜV Rheinland LGA Products GmbH  
Tillystrasse 2  
90431 Nürnberg / Saksa

CE 0197

Rekisteröintinumero / Registration number:

HZ 1198082-1

Versionumero / Version number

02

Korvaa vaatimustenmukaisuusvakuutuksen /  
Replaces Declaration of Conformity from

01.12.2022

Hanau, 25.11.2024

i.V Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

Paikka, päiväys / Place, date

Nimi ja asema / Name and function

Tämä vaatimustenmukaisuusvakuutus on voimassa 2 vuotta tuotettujen laitteiden vastaavan erän julkaisuasiakirjojen kanssa. /  
This statement of conformity is valid for 2 years in connection with the release documents for the respective batch of produced  
devices.



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**Artikkeliluettelo / List of Articles**  
**Liite / Annex: Vaatimustenmukaisuusvakuutus / Declaration of Conformity**

|                                                                                                                                                               |                           |
|---------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------|
| Lääkinnällinen laite /<br><i>The medical device</i>                                                                                                           | <b>GLUMA Desensitizer</b> |
| Versionumero / <i>Version number</i>                                                                                                                          | <b>02</b>                 |
| Korvaa liitteen /<br><i>Replaces Annex from</i>                                                                                                               | <b>01.12.2022</b>         |
| Tämä artikkeliluettelo pätee<br>vaatimustenmukaisuusvakuutuksen versioon /<br><i>This article list is valid for the declaration of<br/>conformity version</i> | <b>02</b>                 |

| UDI-DI /<br><i>UDI-DI</i> | Artikelnummer /<br><i>Article number</i> | Name /<br><i>Name</i>                    |
|---------------------------|------------------------------------------|------------------------------------------|
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| +J014660018540            | 66001854                                 | GLUMA DESENSITIZER SINGLE DOSE 40 X 1 ST |
| +J014660037640            | 66003764                                 | GLUMA DESENSITIZER 1 X 5 ML REGIO        |
| +J014660182210            | 66018221                                 | GLUMA DESENSITIZER VALUE PACK 3 X 5 ML   |

Hanau, 25.11.2024

  
i.V. Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

Paikka, päiväys / *Place, date*

---

Nimi ja asema / *Name and function*

**DÉCLARATION DE CONFORMITÉ / DECLARATION OF CONFORMITY**

Nom et adresse de la société /  
*Name and address of the company*

**Kulzer GmbH**  
Leipziger Straße 2, 63450 Hanau  
Allemagne / Germany  
SRN: DE-MF-000007705

**Nous déclarons sous notre seule responsabilité que / We declare under our sole responsibility that**  
le dispositif médical / *the medical device* **GLUMA Desensitizer**

Nom, type ou modèle, numéro de lot ou de série,  
éventuellement sources et nombre d'articles /  
*Name, type or model, batch or serial number,*  
*possibly sources and number of items*

Liste des articles voir l'Annexe / *List of Articles see Annex*

Code EMDN / *EMDN-Code*  
Code GMDN / *GMDN code*  
code UMDNS / *UMDNS code*  
UDI-DI de base / *Basic UDI-DI*

Q019008  
64794  
10-034  
++J0141103DES010104e5J

de classe / *of class*

Ila

selon la règle / *according to rule*

8 conformément à l'Annexe VIII du Règlement des Dispositifs  
Médicaux 2017/745 / *according to Annex VIII of Medical Device*  
*Regulation 2017/745*

**répond à toutes les dispositions du Règlement des Dispositifs Médicaux 2017/745 qui lui sont applicables. / meets**  
***all the provisions of the Medical Device Regulation 2017/745 which apply to it.***

Application de normes harmonisées, de normes  
nationales ou d'autres documents normatifs /  
*Applied harmonised standards, national standards*  
*or other normative documents*

Autres normes appliquées voir Documentation technique du  
produit GLUMA Desensitizer, version 02  
*Further Applied standards see Technical Documentation of*  
*Product GLUMA Desensitizer, Version 02*

Procédure d'évaluation de la conformité selon /  
*Conformity assessment procedure acc. to*

Règlement relatif aux dispositifs médicaux 2017/745 Annexe IX,  
Chapitre I, Paragraphes 2 et 3 et Chapitre III  
*Medical Device Regulation 2017/745 Annex IX, Chapter I, Section 2*  
*and 3 and Chapter III*

Organisme notifié / *Notified Body*

TÜV Rheinland LGA Products GmbH  
Tillystrasse 2  
90431 Nürnberg / Allemagne

CE 0197

Numéro d'enregistrement / *Registration number:*

HZ 1198082-1

Numéro de version / *Version number*

02

Remplace la Déclaration de conformité de /  
*Replaces Declaration of Conformity from*

01.12.2022

Hanau, 25.11.2024

i.V. Dr. Matthias Hartmann   
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

Lieu, date / *Place, date*

Nom et fonction / *Name and function*

Cette déclaration de conformité est valable 2 ans en relation avec les documents de libération pour le lot respectif des  
dispositifs médicaux fabriqués. / *This statement of conformity is valid for 2 years in connection with the release documents for*  
*the respective batch of produced devices.*

**Déclaration de conformité / Declaration of Conformity**  
**Annexe / Annex : Liste des articles / List of Articles**

Le dispositif médical /  
The medical device

**GLUMA Desensitizer**

Numéro de version / Version number

**02**

Remplace l'annexe de /  
Replaces Annex from

**01.12.2022**

Cette liste d'articles est valable pour la  
déclaration de conformité, version / This  
article list is valid for the declaration of  
conformity version

**02**

| UDI-DI /<br>UDI-DI | Artikelnummer /<br>Article number | Name /<br>Name                           |
|--------------------|-----------------------------------|------------------------------------------|
| +J014658723540     | 65872354                          | GLUMA DESENSITIZER 1 X 5 ML              |
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| +J014660037640     | 66003764                          | GLUMA DESENSITIZER 1 X 5 ML REGIO        |
| +J014660182210     | 66018221                          | GLUMA DESENSITIZER VALUE PACK 3 X 5 ML   |

Hanau, 25.11.2024

Lieu, date / Place, date



i.V. Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

Nom et fonction / Name and function

**DICHIARAZIONE DI CONFORMITÀ / DECLARATION OF CONFORMITY**

Nome e indirizzo della società /  
*Name and address of the company*

**Kulzer GmbH**  
Leipziger Straße 2, 63450 Hanau  
Germania / Germany  
SRN: DE-MF-000007705

**Dichiariamo sotto la nostra esclusiva responsabilità che /  
*We declare under our sole responsibility that***

il dispositivo medico / *the medical device*

**GLUMA Desensitizer**

Nome, tipo o modello, numero di lotto o di serie,  
eventualmente fonti e numero di articoli / *Name, type  
or model, batch or serial number, possibly sources  
and number of items*

Elenco degli articoli vedi allegato / *List of Articles see Annex*

Codice EMDN / *EMDN-Code*  
Codice GMDN / *GMDN code*  
Codice UMDNS / *UMDNS code*  
UDI-DI di base / *Basic UDI-DI*

Q019008  
64794  
10-034  
++J0141103DES010104e5J

di classe / *of class*

Ila

secondo la norma / *according to rule*

8 secondo l'allegato VIII del regolamento sui dispositivi medici  
2017/745 / *according to Annex VIII of Medical Device Regulation  
2017/745*

**soddisfa tutte le disposizioni del regolamento sui dispositivi medici 2017/745 ad esso applicabili. /  
*meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.***

Norme armonizzate applicate, norme nazionali o altri  
documenti normativi / *Applied harmonised standards,  
national standards or other normative documents*

Ulteriori norme applicate vedi Documentazione tecnica di  
Prodotto GLUMA Desensitizer, Versione 02  
*Further Applied standards see Technical Documentation of  
Product GLUMA Desensitizer, Version 02*

Procedura di valutazione della conformità secondo il /  
*Conformity assessment procedure acc. to*

Regolamento sui dispositivi medici 2017/745 Allegato IX, Capitolo I,  
Paragrafi 2 e 3, e Capitolo III  
*Medical Device Regulation 2017/745 Annex IX, Chapter I, Section  
2 and 3 and Chapter III*

Organismo notificato / *Notified Body*

TÜV Rheinland LGA Products GmbH  
Tillystrasse 2  
90431 Norimberga / Germany

CE 0197

Numero di registrazione / *Registration number:*

HZ 1198082-1

Numero versione / *Version number*

02

Sostituisce la dichiarazione di conformità di /  
*Replaces Declaration of Conformity from*

01.12.2022

Hanau, 25.11.2024

  
i.V. Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

Luogo, data / *Place, date*

Nome e funzione / *Name and function*

La presente dichiarazione di conformità ha validità di 2 anni in relazione ai documenti di rilascio per il lotto corrispondente di dispositivi prodotti. / *This statement of conformity is valid for 2 years in connection with the release documents for the respective batch of produced devices.*



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**Elenco degli articoli / List of Articles**  
**Allegato / Annex: Dichiarazione di conformità / Declaration of Conformity**

Il dispositivo medico / **GLUMA Desensitizer**  
*The medical device*

Numero versione / *Version number* **02**

Sostituisce l'allegato da / **01.12.2022**  
*Replaces Annex from*

Questa lista di articoli è valida per la versione **02**  
della dichiarazione di conformità / *This article*  
*list is valid for the declaration of conformity*  
*version*

| <b>UDI-DI /<br/>UDI-DI</b> | <b>Artikelnummer /<br/>Article number</b> | <b>Name /<br/>Name</b>                   |
|----------------------------|-------------------------------------------|------------------------------------------|
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| +J014660037640             | 66003764                                  | GLUMA DESENSITIZER 1 X 5 ML REGIO        |
| +J014660182210             | 66018221                                  | GLUMA DESENSITIZER VALUE PACK 3 X 5 ML   |

Hanau, 25.11.2024

i.V. Dr. Matthias Hartmann   
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

Luogo, data / *Place, date*

Nome e funzione / *Name and function*

**VERKLARING VAN CONFORMITEIT / DECLARATION OF CONFORMITY**

Naam en adres van de onderneming /  
Name and address of the company

**Kulzer GmbH**  
Leipziger Straße 2, 63450 Hanau  
Duitsland / Germany  
SRN: DE-MF-000007705

**Wij verklaren geheel onder onze eigen verantwoordelijkheid dat /  
We declare under our sole responsibility that**

het medisch hulpmiddel / the medical device

**GLUMA Desensitizer**

Naam, type of model, batch of serienummer,  
mogelijke bronnen en aantal items / Name, type or  
model, batch or serial number, possibly sources and  
number of items

Voor lijst met artikelen, zie bijlage / List of Articles see Annex

EMDN-code / EMDN-Code  
GMDN-code / GMDN code  
UMDNS-code / UMDNS code  
Basis UDI-DI / Basic UDI-DI

Q019008  
64794  
10-034  
++J0141103DES010104e5J

van klasse / of class

IIa

in overeenstemming met regelgeving / according to  
rule

8 conform Bijlage VIII van de Verordening (EU) 2017/745  
betreffende medische hulpmiddelen / according to Annex VIII of  
Medical Device Regulation 2017/745

**voldoet aan alle voorschriften van de Verordening (EU) 2017/745 betreffende medische hulpmiddelen die erop van  
toepassing zijn. / meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.**

Toegepaste geharmoniseerde normen, nationale  
normen of andere normatieve documenten / Applied  
harmonised standards, national standards or other  
normative documents

Voor overige toegepaste normen, zie technische documenten van  
product GLUMA Desensitizer, versie 02  
Further Applied standards see Technical Documentation of Product  
GLUMA Desensitizer, Version 02

Conformiteitsbeoordelingsprocedure in  
overeenstemming met / Conformity assessment  
procedure acc. to

Verordening (EU) 2017/745 betreffende medische hulpmiddelen  
bijlage IX, hoofdstuk I, sectie 2 en 3 en hoofdstuk III  
Medical Device Regulation 2017/745 Annex IX, Chapter I, Section 2  
and 3 and Chapter III

Aangemelde instantie / Notified Body

TÜV Rheinland LGA Products GmbH  
Tillystrasse 2  
90431 Nürnberg / Duitsland

CE 0197

Registratienummer / Registration number:

HZ 1198082-1

Versienummer / Version number

02

Vervangt de verklaring van conformiteit van /  
Replaces Declaration of Conformity from

01.12.2022

Hanau, 25.11.2024

  
i.V. Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

Plaats, datum / Place, date

Naam en functie / Name and function

Deze conformiteitsverklaring is 2 jaar geldig in verband met de vrijgavedocumenten voor de respectieve partij van geproduceerde  
hulpmiddelen. / This statement of conformity is valid for 2 years in connection with the release documents for the respective batch  
of produced devices.



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**Lijst met artikelen / List of Articles**  
**Annex / Annex: Verklaring van conformiteit / Declaration of Conformity**

Het medisch hulpmiddel / **GLUMA Desensitizer**  
*The medical device*

Versienummer / *Version number* **02**

Vervangt de bijlage van / **01.12.2022**  
*Replaces Annex from*

Deze artikellijst is geldig voor de **02**  
conformiteitsverklaring, versie / *This article*  
*list is valid for the declaration of conformity*  
*version*

| <b>UDI-DI /<br/>UDI-DI</b> | <b>Artikelnummer /<br/>Article number</b> | <b>Name /<br/>Name</b>                   |
|----------------------------|-------------------------------------------|------------------------------------------|
| +J014658723540             | 65872354                                  | GLUMA DESENSITIZER 1 X 5 ML              |
| +J014660018540             | 66001854                                  | GLUMA DESENSITIZER SINGLE DOSE 40 X 1 ST |
| +J014660037640             | 66003764                                  | GLUMA DESENSITIZER 1 X 5 ML REGIO        |
| +J014660182210             | 66018221                                  | GLUMA DESENSITIZER VALUE PACK 3 X 5 ML   |

Hanau, 25.11.2024

i.V. Dr. Matthias Hartmann   
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

Plaats, datum / *Place, date*

Naam en functie / *Name and function*

**DEKLARACJA ZGODNOŚCI / DECLARATION OF CONFORMITY**

Nazwa i adres firmy /  
Name and address of the company

**Kulzer GmbH**  
Leipziger Straße 2, 63450 Hanau  
Niemcy / Germany  
SRN: DE-MF-000007705

**Niniejszym deklarujemy pod rygorem odpowiedzialności, że /  
We declare under our sole responsibility that**

wyrób medyczny / the medical device

**GLUMA Desensitizer**

Nazwa, typ lub model, numer partii lub serii, ewentualnie  
źródła i liczba elementów / Name, type or model, batch  
or serial number, possibly sources and number of items

Wykaz wyrobów znajduje się w załączniku / List of Articles see  
Annex

Kod wyrobu wg EMDN / EMDN-Code  
Kod wyrobu wg GMDN / GMDN code  
Kod wyrobu wg UMDNS / UMDNS code  
Kod Basic UDI-DI / Basic UDI-DI

Q019008  
64794  
10-034  
++J0141103DES010104e5J

klasy / of class

Ila

zgodnie z regułą / according to rule

8 zgodnie z załącznikiem VIII do Rozporządzenia 2017/745  
w sprawie wyrobów medycznych / according to Annex VIII of  
Medical Device Regulation 2017/745

**spełnia wszystkie przepisy Rozporządzenia 2017/745 w sprawie wyrobów medycznych, które go dotyczą. / meets all  
the provisions of the Medical Device Regulation 2017/745 which apply to it.**

Zastosowane normy zharmonizowane, normy krajowe  
lub inne dokumenty normatywne / Applied harmonised  
standards, national standards or other normative  
documents

Pozostałe stosowane normy znajdują się w dokumentacji  
technicznej produktu GLUMA Desensitizer, wersja 02  
Further Applied standards see Technical Documentation of  
Product GLUMA Desensitizer, Version 02

Procedura oceny zgodności wg. /  
Conformity assessment procedure acc. to

Rozporządzenie 2017/745 w sprawie wyrobów medycznych,  
załącznik IX, rozdział I, sekcja 2 i 3 oraz rozdział III  
Medical Device Regulation 2017/745 Annex IX, Chapter I, Section  
2 and 3 and Chapter III

Jednostka notyfikowana / Notified Body

TÜV Rheinland LGA Products GmbH  
Tillystrasse 2  
90431 Nürnberg/Niemcy

CE 0197

Numer rejestracyjny / Registration number:

HZ 1198082-1

Numer wersji / Version number

02

Zastępuje Deklarację zgodności z /  
Replaces Declaration of Conformity from

01.12.2022

25.11.2024  
Hanau,

i.V. Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

Miejscowość, data / Place, date

Imię i nazwisko, stanowisko / Name and function

Niniejsze deklaracja zgodności jest ważna przez 2 lata w połączeniu z dokumentami zwolnienia odpowiedniej partii  
wyprodukowanych wyrobów. / This statement of conformity is valid for 2 years in connection with the release documents for the  
respective batch of produced devices.



**Wykaz wyrobów / List of Articles**  
**Załącznik / Annex: Deklaracja zgodności / Declaration of Conformity**

Wyrób medyczny /  
The medical device

**GLUMA Desensitizer**

Numer wersji / Version number

**02**

Zastępuje załącznik z dnia /  
Replaces Annex from

**01.12.2022**

Poniższa lista artykułów obowiązuje dla  
następujących wersji deklaracji zgodności /  
This article list is valid for the declaration of  
conformity version

**02**

| <b>UDI-DI /<br/>UDI-DI</b> | <b>Artikelnummer /<br/>Article number</b> | <b>Name /<br/>Name</b>                   |
|----------------------------|-------------------------------------------|------------------------------------------|
| +J014658723540             | 65872354                                  | GLUMA DESENSITIZER 1 X 5 ML              |
| +J014660018540             | 66001854                                  | GLUMA DESENSITIZER SINGLE DOSE 40 X 1 ST |
| +J014660037640             | 66003764                                  | GLUMA DESENSITIZER 1 X 5 ML REGIO        |
| +J014660182210             | 66018221                                  | GLUMA DESENSITIZER VALUE PACK 3 X 5 ML   |

Hanau, 25.11.2024

i.V. Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

Miejscowość, data / Place, date

Imię i nazwisko, stanowisko / Name and function

**DECLARAÇÃO DE CONFORMIDADE / DECLARATION OF CONFORMITY**

Nome e morada da empresa /  
*Name and address of the company*

**Kulzer GmbH**  
Leipziger Straße 2, 63450 Hanau  
Alemanha / Germany  
SRN: DE-MF-000007705

**Declaramos, sob nossa exclusiva responsabilidade, que / We declare under our sole responsibility that**  
o dispositivo médico / *the medical device* **GLUMA Desensitizer**

Nome, tipo ou modelo, número de lote ou de série,  
possivelmente origem e quantidade de itens /  
*Name, type or model, batch or serial number,*  
*possibly sources and number of items*

Lista de artigos, ver Anexo / *List of Articles see Annex*

Código EMDN / *EMDN-Code*  
Código GMDN / *GMDN code*  
Código UMDNS / *UMDNS code*  
UDI-DI básico / *Basic UDI-DI*

Q019008  
64794  
10-034  
++J0141103DES010104e5J

da classe / *of class*

Ila

em conformidade com o regulamento / *according*  
*to rule*

8 em conformidade com o Anexo VIII do Regulamento 2017/745  
relativo aos Dispositivos Médicos / *according to Annex VIII of*  
*Medical Device Regulation 2017/745*

**cumpre todas as disposições aplicáveis do Regulamento 2017/745 relativo aos Dispositivos Médicos. /**  
***meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.***

Normas harmonizadas aplicadas, normas  
nacionais ou outros documentos normativos /  
*Applied harmonised standards, national standards*  
*or other normative documents*

Outras normas aplicadas, ver Documentação técnica do produto  
GLUMA Desensitizer, Versão 02  
*Further Applied standards see Technical Documentation of*  
*Product GLUMA Desensitizer, Version 02*

Procedimento de avaliação da conformidade de  
acordo com /  
*Conformity assessment procedure acc. to*

Anexo IX do Regulamento 2017/745 relativo aos Dispositivos  
Médicos, Capítulo I, secção 2 e 3 e Capítulo III  
*Medical Device Regulation 2017/745 Annex IX, Chapter I, Section 2*  
*and 3 and Chapter III*

Organismo notificado / *Notified Body*

TÜV Rheinland LGA Products GmbH  
Tillystrasse 2  
90431 Nürnberg / Alemanha

CE 0197

Número de registo / *Registration number:*

HZ 1198082-1

Número de versão / *Version number*

02

Substitui a Declaração de Conformidade de /  
*Replaces Declaration of Conformity from*

01.12.2022

Hanau, 25.11.2024

p.p. Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**



Local, data / *Place, date*

Nome e função / *Name and function*

A presente declaração de conformidade é válida durante 2 anos em associação aos documentos do respetivo lote de dispositivos produzidos. / *This statement of conformity is valid for 2 years in connection with the release documents for the respective batch of produced devices.*

**Lista de artigos / *List of Articles***  
**Anexo / *Annex: Declaração de Conformidade / Declaration of Conformity***

O dispositivo médico /  
*The medical device*

**GLUMA Desensitizer**

Número de versão / *Version number*

**02**

Substitui o Anexo de /  
*Replaces Annex from*

**01.12.2022**

A presente lista de artigos é válida para a  
versão da declaração de conformidade / *This  
article list is valid for the declaration of  
conformity version*

**02**

| <b>UDI-DI /<br/><i>UDI-DI</i></b> | <b>Artikelnummer /<br/><i>Article number</i></b> | <b>Name /<br/><i>Name</i></b>            |
|-----------------------------------|--------------------------------------------------|------------------------------------------|
| +J014658723540                    | 65872354                                         | GLUMA DESENSITIZER 1 X 5 ML              |
| +J014660018540                    | 66001854                                         | GLUMA DESENSITIZER SINGLE DOSE 40 X 1 ST |
| +J014660037640                    | 66003764                                         | GLUMA DESENSITIZER 1 X 5 ML REGIO        |
| +J014660182210                    | 66018221                                         | GLUMA DESENSITIZER VALUE PACK 3 X 5 ML   |

Hanau, 25.11.2024

p.p. Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**



Local, data / *Place, date*

Nome e função / *Name and function*

**DECLARAȚIE DE CONFORMITATE / DECLARATION OF CONFORMITY**

Numele și adresa companiei /  
*Name and address of the company*

**Kulzer GmbH**  
Leipziger Straße 2, 63450 Hanau  
Germania / Germany

SRN: DE-MF-000007705

**Declarăm pe propria răspundere că / We declare under our sole responsibility that**  
dispozitivul medical / *the medical device*

**GLUMA Desensitizer**

Nume, tip sau model, număr de lot sau de serie,  
eventual sursele și numărul de articole / *Name,  
type or model, batch or serial number, possibly  
sources and number of items*

Lista de articole, vezi Anexa / *List of Articles see Annex*

Cod EMDN / *EMDN-Code*  
Cod GMDN / *GMDN code*  
Cod UMDNS / *UMDNS code*  
UDI-DI de bază / *Basic UDI-DI*

Q019008  
64794  
10-034  
++J0141103DES010104e5J

din clasa / *of class*

IIa

în conformitate cu regula / *according to rule*

8 conform Anexei VIII la Regulamentul privind dispozitivele  
medicale 2017/745 / *according to Annex VIII of Medical Device  
Regulation 2017/745*

**respectă toate prevederile Regulamentului privind dispozitivele medicale 2017/745 corespunzător. /**  
***meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.***

Standarde armonizate, naționale aplicate sau alte  
documente normative / *Applied harmonised  
standards, national standards or other normative  
documents*

Alte standarde aplicate, vezi documentația tehnică a Produsului  
GLUMA Desensitizer, Versiunea 02  
*Further Applied standards see Technical Documentation of  
Product GLUMA Desensitizer, Version 02*

Procedură de evaluare a conformității în conf. cu /  
*Conformity assessment procedure acc. to*

Regulamentul privind dispozitivele medicale 2017/745, Anexa IX,  
Capitolul I, Secțiunile 2 și 3, și Capitolul III  
*Medical Device Regulation 2017/745 Annex IX, Chapter I,  
Section 2 and 3 and Chapter III*

Organism notificat / *Notified Body*

TÜV Rheinland LGA Products GmbH  
Tillystrasse 2  
90431 Nürnberg / Germany

CE 0197

Numărul de înregistrare / *Registration number:*

HZ 1198082-1

Număr versiune / *Version number*

02

Înlocuiește Declarația de conformitate din /  
*Replaces Declaration of Conformity from*

01.12.2022

Hanau, 25.11.2024

i.V.  
Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

Loc, dată / *Place, date*

Nume și funcție / *Name and function*

Prezenta declarație de conformitate este valabilă timp de 2 ani împreună cu documentele de autorizare pentru respectivul lot de  
dispozitive produse. / *This statement of conformity is valid for 2 years in connection with the release documents for the  
respective batch of produced devices.*

**Listă de articole / *List of Articles***  
**Anexă / *Annex: Declarație de conformitate / *Declaration of Conformity****

Dispozitivul medical /  
*The medical device*

**GLUMA Desensitizer**

Număr versiune / *Version number*

**02**

Înlocuiește Anexa de la /  
*Replaces Annex from*

**01.12.2022**

Această listă de articole este valabilă pentru  
declarația de conformitate versiunea / *This  
article list is valid for the declaration of  
conformity version*

**02**

| <b>UDI-DI /<br/><i>UDI-DI</i></b> | <b>Artikelnummer /<br/><i>Article number</i></b> | <b>Name /<br/><i>Name</i></b>            |
|-----------------------------------|--------------------------------------------------|------------------------------------------|
| +J014658723540                    | 65872354                                         | GLUMA DESENSITIZER 1 X 5 ML              |
| +J014660018540                    | 66001854                                         | GLUMA DESENSITIZER SINGLE DOSE 40 X 1 ST |
| +J014660037640                    | 66003764                                         | GLUMA DESENSITIZER 1 X 5 ML REGIO        |
| +J014660182210                    | 66018221                                         | GLUMA DESENSITIZER VALUE PACK 3 X 5 ML   |

Hanau, 25.11.2024

Loc, dată / *Place, date*



**i.V. Dr. Matthias Hartmann**  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

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**Nume și funcție / *Name and function***

**FÖRSÄKRAN OM ÖVERENSSTÄMMELSE / DECLARATION OF CONFORMITY**Företagets namn och adress /  
*Name and address of the company***Kulzer GmbH**  
Leipziger Straße 2, 63450 Hanau  
Tyskland / Germany

SRN: DE-MF-000007705

**Vi försäkrar på eget ansvar att / We declare under our sole responsibility that**den medicintekniska produkten / *the medical device***GLUMA Desensitizer**Namn, typ eller modell, batch eller serienummer,  
eventuella källor och antal artiklar / *Name, type or  
model, batch or serial number, possibly sources and  
number of items*Se bilaga för lista över artiklar / *List of Articles see Annex*EMDN-kod / *EMDN-Code*

Q019008

GMDN-kod / *GMDN code*

64794

UMDNS-kod / *UMDNS code*

10-034

Grundläggande UDI-DI / *Basic UDI-DI*

++J0141103DES010104e5J

i klass / *of class*

IIa

enligt paragraf / *according to rule*8 enligt bilaga VIII i förordningen om medicintekniska produkter  
2017/745 / *according to Annex VIII of Medical Device Regulation  
2017/745***uppfyller kraven i förordningen om medicintekniska produkter 2017/745 som gäller produkten. /  
*meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.***Tillämpade harmoniserade standarder, nationella  
standarder eller andra normerande dokument /  
*Applied harmonised standards, national standards  
or other normative documents*För ytterligare tillämpade standarder, se teknisk dokumentation för  
produkten GLUMA Desensitizer, version 02  
*Further Applied standards see Technical Documentation of  
Product GLUMA Desensitizer, Version 02*Förfarande för bedömning av överensstämmelse  
enl. /  
*Conformity assessment procedure acc. to*förordning om medicintekniska 2017/745 bilaga IX, kapitel I,  
avsnitt 2 och 3 och kapitel III  
*Medical Device Regulation 2017/745 Annex IX, Chapter I,  
Section 2 and 3 and Chapter III*Anmält organ / *Notified Body*TÜV Rheinland LGA Products GmbH  
Tillystrasse 2  
90431 Nürnberg/Tyskland

CE 0197

Registreringsnummer / *Registration number:*

HZ 1198082-1

Versionsnummer / *Version number*

02

Ersätter försäkran om överensstämmelse från /  
*Replaces Declaration of Conformity from*

01.12.2022

Hanau, 25.11.2024

i.V. Dr. Matthias Hartmann  
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**Ort, datum / *Place, date*Namn och funktion / *Name and function*Denna försäkran om överensstämmelse är giltig i 2 år tillsammans med dokumenten för frisläppande av respektive  
tillverkningsserie av medicintekniska produkter. / *This statement of conformity is valid for 2 years in connection with the release  
documents for the respective batch of produced devices..*



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**Lista över artiklar / List of Articles**  
**Bilaga / Annex: Försäkran om överensstämmelse / Declaration of Conformity**

Den medicintekniska produkten / **GLUMA Desensitizer**  
*The medical device*

Versionsnummer / *Version number* **02**

Ersätter bilaga från / **01.12.2022**  
*Replaces Annex from*

Denna artikellista gäller för förklaring av **02**  
överensstämmelse version / *This article list is*  
*valid for the declaration of conformity version*

| <b>UDI-DI /<br/>UDI-DI</b> | <b>Artikelnummer /<br/>Article number</b> | <b>Name /<br/>Name</b>                   |
|----------------------------|-------------------------------------------|------------------------------------------|
| +J014658723540             | 65872354                                  | GLUMA DESENSITIZER 1 X 5 ML              |
| +J014660018540             | 66001854                                  | GLUMA DESENSITIZER SINGLE DOSE 40 X 1 ST |
| +J014660037640             | 66003764                                  | GLUMA DESENSITIZER 1 X 5 ML REGIO        |
| +J014660182210             | 66018221                                  | GLUMA DESENSITIZER VALUE PACK 3 X 5 ML   |

Hanau, 25.11.2024

Ort, datum / *Place, date*

i.V. Dr. Matthias Hartmann   
Head of Global Quality, Regulatory & Scientific Services  
**Kulzer GmbH**

---

Namn och funktion / *Name and function*