

**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 thatdas Medizinprodukt / *the medical device***Charisma Bulk Flow One**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*Q010101
35870
16-724
++J0141103COMP010103i8Cder Klasse / *of class*

IIa

nach Regel / *according to rule*8-1, 19-3 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*EN ISO 4049 - Dentistry- Polymer-based restorative materials
Weitere angewandte Normen siehe Version 03 der Technischen
Dokumentation von Product Charisma Bulk Flow One / *Further
Applied standards see Technical Documentation of Product
Charisma Bulk Flow One, Version 03*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*

03

Ersetzt Konformitätserklärung vom /
Replaces Declaration of Conformity from

08.08.2024

Hanau, 21.10.2024

i.V.


Dr. Matthias Hartmann
Head of Global Quality, Regulatory & Scientific Services
Kulzer GmbHOrt, 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

Charisma Bulk Flow One

Versionsnummer Artikelliste /
Version number article list

04

Ersetzt Artikelliste vom /
Replaces article list from

08.08.2024

Diese Artikelliste ist gültig für die Konformitätserklärung Version / 03
This article list is valid for the declaration of conformity version

UDI-DI / <i>UDI-DI</i>	Artikelnummer / <i>Article number</i>	Name / <i>Name</i>
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Hanau, 21.10.2024

i.V. 

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

Name und Funktion / *Name and function*

Ort, Datum / *Place, date*

ДЕКЛАРАЦИЯ ЗА СЪОТВЕТСТВИЕ / DECLARATION OF CONFORMITY

Име и адрес на фирмата /
Name and address of the company

Kulzer GmbH
Leipziger Straße 2, 63450 Hanau
Германия / Germany

SRN: DE-MF-000007705

Декларираме на наша собствена отговорност, че / We declare under our sole responsibility that

медицинското изделие / the medical device

Charisma Bulk Flow One

Наименование, тип или модел, партиден или
сериен номер, евентуално произход и брой
елементи / Name, type or model, batch or serial
number, possibly sources and number of items

Списък с артикули, вижте Приложението /
List of Articles see Annex

Код по EMDN / EMDN-Code
Код по GMDN / GMDN code
Код по UMDNS / UMDNS code
Основна UDI-DI идентификация / Basic UDI-DI

Q010101
35870
16-724
++J0141103COMP010103i8C

от клас / of class

Ila

съгласно правило / according to rule

8-1, 19-3 съгласно Приложение VIII от Регламента за
медицинските изделия 2017/745 / according to Annex VIII of
Medical Device Regulation 2017/745

**отговаря на всички разпоредби на Регламента за медицинските изделия 2017/745, който се прилага за него. /
meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.**

Приложени хармонизирани стандарти, национални
стандарти или други нормативни документи /
Applied harmonised standards, national standards or
other normative documents

EN ISO 4049 - Dentistry- Polymer-based restorative materials
Други приложени стандарти, вижте техническата
документация на продукт Charisma Bulk Flow One Версия 03
Further Applied standards see Technical Documentation of
Product Charisma Bulk Flow One, Version 03

Процедура за оценка на съответствието съгласно /
Conformity assessment procedure acc. to

Регламента за медицинските изделия 2017/745 Приложение
IX, глава I, раздел 2 и 3 и глава III
Medical Device Regulation 2017/745 Annex IX, Chapter I, Section
2 and 3 and Chapter III

Нотифициран орган / Notified Body

TÜV Rheinland LGA Products GmbH
Tillystrasse 2
90431 Nürnberg / Германия
CE 0197

Регистрационен номер / Registration number:

HZ 1198082-1

Номер на версия / Version number

03

Заменя Декларация за съответствие от /
Replaces Declaration of Conformity from

08.08.2024



Ханау, 21.10.2024

от името на д-р Dr. Matthias Hartmann
Head of Global Quality, Regulatory & Scientific Services
Kulzer GmbH

Място, дата / Place, date

Име и длъжност / Name and function

Тази Декларация за съответствие е валидна за 2 години във връзка с публикуваните документи за съответната партида произведени
устройства / This statement of conformity is valid for 2 years in connection with the release documents for the respective batch of produced
devices.

Списък с артикули / *List of Articles*

Приложение / *Annex*: Декларация за съответствие / *Declaration of Conformity*

Медицинското изделие /
The medical device

Charisma Bulk Flow One

Номер на версия / *Version number*

04

Заменя Приложението от /
Replaces Annex from

08.08.2024

Този списък със статии е валиден във връзка с декларацията за съответствие, версия / *This article list is valid for the declaration of conformity version*

03

UDI-DI / <i>UDI-DI</i>	Artikelnummer / <i>Article number</i>	Name / <i>Name</i>
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Ханау, 21.10.2024


от името на д-р Dr. Matthias Hartmann
Head of Global Quality, Regulatory & Scientific Services
Kulzer GmbH

Място, дата / *Place, date*

Име и длъжност / *Name and function*

**PROHLÁŠENÍ O SHODĚ / DECLARATION OF CONFORMITY**

Název a adresa společnosti /
Name and address of the company

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

Prohlašujeme na svou výlučnou zodpovědnost, že / We declare under our sole responsibility that
zdravotnický prostředek / the medical device

Charisma Bulk Flow One

Název, typ nebo model, šarže nebo výrobní číslo,
případně zdroje a počet kusů / Name, type or
model, batch or serial number, possibly sources and
number of items

Seznam položek je uveden v příloze /
List of Articles see Annex

Kód EMDN / EMDN-Code
Kód GMDN / GMDN code
Kód UMDNS / UMDNS code
Základní UDI-DI / Basic UDI-DI

Q010101
35870
16-724
++J0141103COMP010103i8C

třídy / of class

IIa

podle pravidla / according to rule

8-1, 19-3 podle přílohy VIII k nařízení 2017/745 o zdravotnických
prostředcích / according to Annex VIII of Medical Device Regulation
2017/745

splňuje všechna ustanovení nařízení 2017/745 o zdravotnických prostředcích, která se ho týkají. /
meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.

Použité harmonizované normy, národní normy nebo
jiné normativní dokumenty / Applied harmonised
standards, national standards or other normative
documents

EN ISO 4049 - Dentistry- Polymer-based restorative materials
Další použité normy najdete v technické dokumentaci k
výrobku Charisma Bulk Flow One, verze 03
Further Applied standards see Technical Documentation of
Product Charisma Bulk Flow One, Version 03

Procedura posouzení shody podle /
Conformity assessment procedure acc. to

nařízení 2017/745 o zdravotnických prostředcích, příloha IX,
kapitola I, oddíl 2 a 3 a kapitola III

Medical Device Regulation 2017/745 Annex IX, Chapter I,
Section 2 and 3 and Chapter III

Notifikovaná osoba / Notified Body

TÜV Rheinland LGA Products GmbH
Tillystrasse 2
90431 Nürnberg / Německo

CE 0197

Registrační číslo / Registration number:

HZ 1198082-1

Číslo verze / Version number

03

Nahrazuje Prohlášení o shodě ze dne /
Replaces Declaration of Conformity from

08.08.2024

Hanau, 21.10.2024

i.V.

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

Místo, datum / Place, date

Jméno a funkce / Name and function

Toto prohlášení o shodě je platné po dobu 2 let ve spojení s příbalovými informacemi pro příslušnou šarži vyrobených
zdravotnických prostředků. / This statement of conformity is valid for 2 years in connection with the release documents for the
respective batch of produced devices.

Seznam položek / List of Articles

Příloha / Annex: Prohlášení o shodě / Declaration of Conformity

Zdravotnický prostředek /
The medical device

Charisma Bulk Flow One

Číslo verze / Version number

04

Nahrazuje přílohu ze dne /
Replaces Annex from

08.08.2024

Tento seznam zboží platí pro verzi
prohlášení o shodě / This article list is valid
for the declaration of conformity version:

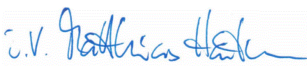
03

UDI-DI / UDI-DI	Artikelnummer / Article number	Name / Name
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Hanau,

21.10.2024

Místo, datum / Place, date



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

Jméno a funkce / Name and function

**VASTAVUSDEKLARATSIOON / DECLARATION OF CONFORMITY**

Ettevõtte nimi ja aadress /
Name and address of the company

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

Ettevõtte kinnitab oma ainuvastusel, et / We declare under our sole responsibility that
meditsiiniseade / *the medical device*

Charisma Bulk Flow One

Nimi, tüüp või mudel, partii või seerianumber,
võimalikud allikad ja osad / *Name, type or model,
batch or serial number, possibly sources and
number of items*

Artiklite loendit vt lisast / *List of Articles see Annex*

EMDN-kood / *EMDN-Code*
GMDN-i kood / *GMDN code*
UMDNS-i kood / *UMDNS code*
Põhi-UDI-DI / *Basic UDI-DI*

Q010101
35870
16-724
++J01411103COMP010103i8C

klassil / *of class*

Ila

kooskõlas reegluga / *according to rule*

8-1, 19-3 kooskõlas meditsiiniseadme määrusega 2017/745 lisaga
VIII / *according to Annex VIII of Medical Device Regulation
2017/745*

vastab kõigile meditsiiniseadme määruse 2017/745 sätetele, mis seadmele kohaldub. /
meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.

Kehtivad harmoniseeritud standardid, riiklikud
standardid ja muud normatiivdokumendid / *Applied
harmonised standards, national standards or other
normative documents*

EN ISO 4049 - Dentistry- Polymer-based restorative materials
Muid rakenduvaid standardeid vt tehnilistest
dokumentidest tootele Charisma Bulk Flow One, Version 03
*Further Applied standards see Technical Documentation of
Product Charisma Bulk Flow One, Version 03*

Vastavushindamismenetlus vastavalt /
Conformity assessment procedure acc. to

meditsiiniseadmete määruse 2017/745 IX. lisa, I. peatükk, 2. ja 3.
jaotis ning III. peatükk

*Medical Device Regulation 2017/745 Annex IX, Chapter I, Section
2 and 3 and Chapter III*

Teavitatud asutus / *Notified Body*

TÜV Rheinland LGA Products GmbH
Tillystrasse 2
90431 Nürnberg / Saksamaa

CE 0197

Registreerimisnumber / *Registration number:*

HZ 1198082-1

Versiooni number / *Version number*

03

Asendab vastavusdeklaratsiooni alates /
Replaces Declaration of Conformity from

08.08.2024

Hanau, 21.10.2024

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

Koht, kuupäev / *Place, date*

Nimi ja funktsioon / *Name and function*

See vastavusdeklaratsioon kehtib 2 aastat koos toodetud seadmete vastava partii väljastusdokumentidega / *This statement of
conformity is valid for 2 years in connection with the release documents for the respective batch of produced devices.*

Artiklite loend / List of Articles
lisa / Annex: Vastavusdeklaratsioon / Declaration of Conformity

Meditsiiniseade /
The medical device

Versioni number / Version number

Asendab lisa alates /
Replaces Annex from

See artiklite loend kehtib
vastavusdeklaratsiooni versiooni kohta / This
article list is valid for the declaration of
conformity version

Charisma Bulk Flow One

04

08.08.2024

03

UDI-DI / UDI-DI	Artikelnummer / Article number	Name / Name
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

21.10.2024
Hanau,

Koht, kuupäev / Place, date


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

Nimi ja funktsioon / 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*

Charisma Bulk Flow One

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*

Q010101
35870
16-724
++J0141103COMP010103i8C

de la clase / *of class*

Ila

de acuerdo con la norma / *according to rule*

8-1, 19-3 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

EN ISO 4049 - Dentistry– Polymer-based restorative materials
Para otras normas aplicadas consulte la documentación técnica del
producto Charisma Bulk Flow One, versión 03
Further Applied standards see Technical Documentation of
Product Charisma Bulk Flow One, Version 03

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

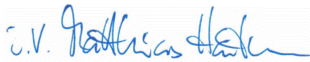
Número de versión / *Version number*

03

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

08.08.2024

Hanau, 21.10.2024


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*

Charisma Bulk Flow One

04

08.08.2024

03

UDI-DI / <i>UDI-DI</i>	Artikelnummer / <i>Article number</i>	Name / <i>Name</i>
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Hanau, 21.10.2024

Lugar, fecha / *Place, date*


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

Nombre y cargo / *Name and function*

**ΔΗΛΩΣΗ ΣΥΜΜΟΡΦΩΣΗΣ / DECLARATION OF CONFORMITY**Επωνυμία και διεύθυνση εταιρείας /
Name and address of the company**Kulzer GmbH**
Leipziger Straße 2, 63450 Hanau
Γερμανία / Germany
SRN: DE-MF-000007705**Δηλώνουμε με δική μας ευθύνη ότι / We declare under our sole responsibility that**

το ιατροτεχνολογικό προϊόν / the medical device

Charisma Bulk Flow OneΕπωνυμία, τύπος ή μοντέλο, παρτίδα ή αριθμός
σειράς, πιθανές πηγές και αριθμός ειδών / Name,
type or model, batch or serial number, possibly
sources and number of items

Κατάλογος ειδών Παράρτημα / List of Articles see Annex

Κωδικός EMDN / EMDN-Code
Κωδικός GMDN / GMDN code
Κωδικός UMDNS / UMDNS code
Βασικό UDI-DI / Basic UDI-DIQ010101
35870
16-724
++J0141103COMP010103i8C

κλάσης / of class

IIa

σύμφωνα με τον κανόνα / according to rule

8-1, 19-3 σύμφωνα με το Παράρτημα VIII του Κανονισμού
2017/745 για τα ιατροτεχνολογικά προϊόντα / according to Annex
VIII of Medical Device Regulation 2017/745**πληροί όλες τις ισχύουσες διατάξεις του Κανονισμού 2017/745 για τα ιατροτεχνολογικά προϊόντα. /
meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.**Εφαρμοζόμενα εναρμονισμένα πρότυπα, εθνικά
πρότυπα ή άλλα κανονιστικά έγγραφα / Applied
harmonised standards, national standards or other
normative documentsEN ISO 4049 - Dentistry- Polymer-based restorative materials
Για περαιτέρω εφαρμοζόμενα πρότυπα βλ. την τεχνική
τεκμηρίωση του
προϊόντος Charisma Bulk Flow One, έκδοση 03
Further Applied standards see Technical Documentation of
Product Charisma Bulk Flow One, Version 03Διαδικασία αξιολόγησης συμμόρφωσης σύμφωνα με
/ Conformity assessment procedure acc. toΚανονισμός 2017/745 για τα ιατροτεχνολογικά προϊόντα,
Παράρτημα IX, Κεφάλαιο I, Τμήμα 2 και 3, και Κεφάλαιο III
Medical Device Regulation 2017/745 Annex IX, Chapter I, Section
2 and 3 and Chapter III

Κοινοποιημένος οργανισμός / Notified Body

TÜV Rheinland LGA Products GmbH
Tillystrasse 2
90431 Nürnberg / Γερμανία

CE 0197

Αριθμός καταχώρησης / Registration number:

HZ 1198082-1

Αριθμός έκδοσης / Version number

03

Αντικαθιστά τη δήλωση συμμόρφωσης από /
Replaces Declaration of Conformity from

08.08.2024

Hanau, 21.10.2024

i.V. Dr. Matthias Hartmann
Head of Global Quality, Regulatory & Scientific Services
Kulzer GmbHΤόπος, ημερομηνία / Place,
date

Ονοματεπώνυμο και τίτλος / Name and function

Αυτή η δήλωση συμμόρφωσης ισχύει για 2 χρόνια σε σχέση με τα έγγραφα κυκλοφορίας για την αντίστοιχη παρτίδα των
παραγόμενων προϊόντων. / This statement of conformity is valid for 2 years in connection with the release documents for the
respective batch of produced devices.

Κατάλογος ειδών / *List of Articles*
Παράρτημα / *Annex: Δήλωση συμμόρφωσης / *Declaration of Conformity**

Το ιατροτεχνολογικό προϊόν /
The medical device

Charisma Bulk Flow One

Αριθμός έκδοσης / *Version number*

04

Αντικαθιστά το Παράρτημα από /
Replaces Annex from

08.08.2024

Αυτός ο κατάλογος προϊόντων ισχύει για την
έκδοση δήλωσης συμμόρφωσης / *This article
list is valid for the declaration of conformity
version*

03

UDI-DI / <i>UDI-DI</i>	Artikelnummer / <i>Article number</i>	Name / <i>Name</i>
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Hanau, 21.10.2024

Τόπος, ημερομηνία / *Place, date*



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

Ονοματεπώνυμο και τίτλος / *Name and function*

IZJAVA O USKLAĐENOSTI / DECLARATION OF CONFORMITY

Naziv i adresa tvrtke / Name and address of the company	Kulzer GmbH Leipziger Straße 2, 63450 Hanau Njemačka / Germany SRN: DE-MF-000007705
Izjavljujemo pod punom odgovornošću da / We declare under our sole responsibility that	
medicinski proizvod / the medical device	Charisma Bulk Flow One
Naziv, tip ili model, broj serije, po mogućnosti izvori i broj stavki / Name, type or model, batch or serial number, possibly sources and number of items	Popis artikala, pogledajte Dodatak / List of Articles see Annex
šifra EMDN / EMDN-Code	Q010101
šifra GMDN / GMDN code	35870
šifra UMDNS / UMDNS code	16-724
osnovni UDI-DI / Basic UDI-DI	++J0141103COMP010103i8C
klase / of class	Ila
u skladu s pravilom / according to rule	8-1, 19-3 u skladu s Dodatkom VIII Uredbe 2017/745 o medicinskim proizvodima / according to Annex VIII of Medical Device Regulation 2017/745
ispunjava sve odredbe Uredbe 2017/745 o medicinskim proizvodima koje se na njega odnose. / meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.	
Primijenjene usklađene norme, državne norme ili drugi normativni dokumenti / Applied harmonised standards, national standards or other normative documents	EN ISO 4049 - Dentistry– Polymer-based restorative materials Druge primijenjene norme, pogledajte Tehničku dokumentaciju za proizvod Charisma Bulk Flow One, verzija 03 Further Applied standards see Technical Documentation of Product Charisma Bulk Flow One, Version 03
Postupak procjene usklađenosti prema / Conformity assessment procedure acc. to	Prilog IX Uredbi 2017/745 o medicinskim proizvodima, Poglavlje I, Odjeljak 2 i 3 te Poglavlje III Medical Device Regulation 2017/745 Annex IX, Chapter I, Section 2 and 3 and Chapter III
Obaviješteno tijelo / Notified Body	TÜV Rheinland LGA Products GmbH Tillystrasse 2 90431 Nürnberg / Njemačka
	CE 0197
Registracijski broj / Registration number:	HZ 1198082-1
Broj verzije / Version number	03
Zamjenjuje Izjavu o usklađenosti od / Replaces Declaration of Conformity from	08.08.2024 
Hanau, 21.10.2024	i.V. Dr. Matthias Hartmann Head of Global Quality, Regulatory & Scientific Services Kulzer GmbH
Mjesto, datum / Place, date	Ime i funkcija / Name and function

Ova Izjava o usklađenosti valjana je 2 godine u odnosu na dokumente o izdanju za odgovarajuće serije proizvedenih umedicinskih proizvoda. / This statement of conformity is valid for 2 years in connection with the release documents for the respective batch of produced devices.

Popis artikala / List of Articles

Dodatak / Annex: Izjava o usklađenosti / Declaration of Conformity

Medicinski proizvod /
The medical device

Charisma Bulk Flow One

Broj verzije / Version number

04

Zamjenjuje Dodatak od /
Replaces Annex from

08.08.2024

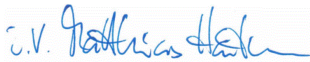
Ovaj popis artikala valjan je za verziju izjave
u sukladnosti / This article list is valid for the
declaration of conformity version

03

UDI-DI / UDI-DI	Artikelnummer / Article number	Name / Name
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Hanau,^{21.10.2024}

Mjesto, datum / Place, date



i.V. Dr. Matthias Hartmann

Head of Global Quality, Regulatory & Scientific Services

Kulzer GmbH

Ime i funkcija / Name and function

**MEGFELELŐSÉGI NYILATKOZAT / DECLARATION OF CONFORMITY**

A vállalat neve és címe /
Name and address of the company

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

Kizárólagos felelősségünkre kijelentjük, hogy / We declare under our sole responsibility that
az orvostechnikai eszköz / the medical device **Charisma Bulk Flow One**

Név, típus vagy modell, tétel vagy sorozatszám,
esetleg források és tételek száma / Name, type or
model, batch or serial number, possibly sources and
number of items

A cikkek listáját lásd a mellékletben / List of Articles see Annex

EMDN kód / EMDN-Code
GMDN kód / GMDN code
UMDNS kód / UMDNS code
Alapvető UDI-DI / Basic UDI-DI

Q010101
35870
16-724
++J0141103COMP010103i8C

osztálya / of class

Ila

a következő szabály szerint / according to rule

8-1, 19-3 az orvostechnikai eszközökről szóló 2017/745 rendelet VIII.
melléklete szerint / according to Annex VIII of Medical Device
Regulation 2017/745

**megfelel az orvostechnikai eszközökről szóló, 2017/745 rendelet valamennyi rá vonatkozó rendelkezésének. / meets
all the provisions of the Medical Device Regulation 2017/745 which apply to it.**

Alkalmazott harmonizált szabványok, nemzeti
szabványok vagy más normatív dokumentumok /
Applied harmonised standards, national standards
or other normative documents

EN ISO 4049 - Dentistry- Polymer-based restorative materials
További alkalmazott szabványokat lásd a műszaki dokumentációban,
termék: Charisma Bulk Flow One, 03. verzió
Further Applied standards see Technical Documentation of
Product Charisma Bulk Flow One, Version 03

Megfelelőségértékelési eljárás a következő szerint
/ Conformity assessment procedure acc. to

Az orvostechnikai eszközökről szóló, 2017/745 rendelet IX. függeléke,
az I. fejezet 2. és 3. szakasza, és a III. fejezet
Medical Device Regulation 2017/745 Annex IX, Chapter I, Section 2
and 3 and Chapter III

Bejelentett szervezet / Notified Body

TÜV Rheinland LGA Products GmbH
Tillystrasse 2
90431 Nürnberg / Németország

CE 0197

Regisztrációs szám / Registration number:

HZ 1198082-1

Verziószám / Version number

03

Felváltja a megfelelőségi nyilatkozatot ettől /
Replaces Declaration of Conformity from

08.08.2024

Hanau, 21.10.2024

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

Hely, dátum / Place, date

Név és funkció / Name and function

Ez a megfelelőségi nyilatkozat 2 évig érvényes a gyártott eszközök adott tételére vonatkozó kibocsátási dokumentumokkal együtt. / This statement of conformity is valid for 2 years in connection with the release documents for the respective batch of produced devices.

Cikkek listája / List of Articles
Melléklet / Annex: Megfelelőségi nyilatkozat / Declaration of Conformity

Az orvostechnikai eszköz /
The medical device

Charisma Bulk Flow One

Verziószám / Version number

04

Felváltja a mellékletet ettől /
Replaces Annex from

08.08.2024

Ez a tétellista a megfelelőségi nyilatkozat
következő verziójához érvényes / This article
list is valid for the declaration of conformity
version

03

UDI-DI / UDI-DI	Artikelnummer / Article number	Name / Name
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Hanau, 21.10.2024

Hely, dátum / Place, date


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

Név és funkció / Name and function

**ATITIKTIES DEKLARACIJA / DECLARATION OF CONFORMITY**

Bendrovės pavadinimas ir adresas /
Name and address of the company

Kulzer GmbH
Leipziger Straße 2, 63450 Hanau
Vokietija / Germany

SRN: DE-MF-000007705

Prisiimdami visą atsakomybę pareiškiame, kad / We declare under our sole responsibility that

medicinos prietaisas / *the medical device*

Charisma Bulk Flow One

Pavadinimas, tipas arba modelis, partija arba serijos
numeris, galimi šaltiniai ir elementų skaičius / *Name,
type or model, batch or serial number, possibly
sources and number of items*

Prekių sąrašo ieškokite Priede / *List of Articles see Annex*

EMDN kodas / *EMDN-Code*
GMDN kodas / *GMDN code*
UMDNS kodas / *UMDNS code*
Pagrindinis UDI-DI / *Basic UDI-DI*

Q010101
35870
16-724
++J0141103COMP010103i8C

klasės / *of class*

Ila

pagal taisyklę / *according to rule*

8-1, 19-3 Pagal Medicinos prietaisų reglamento 2017/745 VIII
priedą / *according to Annex VIII of Medical Device Regulation
2017/745*

**atitinka visas jam taikomas Medicinos prietaisų reglamento 2017/745 sąlygas. /
*meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.***

Taikomi harmonizuotieji standartai, nacionaliniai
standartai ar kiti normatyviniai dokumentai / *Applied
harmonised standards, national standards or other
normative documents*

EN ISO 4049 - Dentistry- Polymer-based restorative materials
atkuriamosiomis medžiagomis Kitus taikomus standartus žr.
produkto Charisma Bulk Flow One, techninėje dokumentacijoje, 03
versijoje
*Further Applied standards see Technical Documentation of
Product Charisma Bulk Flow One, Version 03*

Atitikties patvirtinimo procedūra pagal /
Conformity assessment procedure acc. to

Medicinos priemonių reglamento 2017/745 IX priedas, I skyrius, 2 ir
3 straipsniai bei III skyrius
*Medical Device Regulation 2017/745 Annex IX, Chapter I, Section
2 and 3 and Chapter III*

Notifikuotoji įstaiga / *Notified Body*

TÜV Rheinland LGA Products GmbH
Tillystrasse 2
90431 Nürnberg / Vokietija

CE 0197

Registracijos numeris / *Registration number:*

HZ 1198082-1

Versijos numeris / *Version number*

03

Pakeičia atitikties deklaraciją nuo /
Replaces Declaration of Conformity from

08.08.2024

Hanau, 21.10.2024

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

Vieta, data / *Place, date*

Vardas, pavardė ir pareigos / *Name and function*

Ši atitikties deklaracija galioja 2 metus kartu su atitinkamos pagamintų priemonių partijos pateikimo į rinką dokumentais. / *This statement of conformity is valid for 2 years in connection with the release documents for the respective batch of produced devices.*

Prekių sąrašas / List of Articles

Priedas / Annex: Atitikties deklaracija / Declaration of Conformity

Medicinos prietaisas /
The medical device

Charisma Bulk Flow One

Versijos numeris / Version number

04

Pakeičia Priedą nuo /
Replaces Annex from

08.08.2024

Šis straipsnių sąrašas tinka atitikties
deklaracijai, kurios versija yra / This article
list is valid for the declaration of conformity
version

03

UDI-DI / UDI-DI	Artikelnummer / Article number	Name / Name
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Hanau21.10.2024

Vieta, data / Place, date



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

Vardas, pavardė ir pareigos / Name and function

**ATBILSTĪBAS DEKLARĀCIJA / DECLARATION OF CONFORMITY**Uzņēmuma nosaukums un adrese /
*Name and address of the company***Kulzer GmbH**
Leipziger Straße 2, 63450 Hanau
Vācija / Germany

SRN: DE-MF-000007705

Mēs vienīgi uz savu atbildību deklarējam, ka / We declare under our sole responsibility that
medicīniskā ierīce / *the medical device***Charisma Bulk Flow One**Preču nosaukums, tips vai modelis, partijas vai
sērijas numurs, iespējamie avoti un skaits / *Name,*
type or model, batch or serial number, possibly
*sources and number of items*Preču sarakstu skatiet pielikumā / *List of Articles see Annex*EMDN kods / *EMDN-Code*
GMDN kods / *GMDN code*
UMDNS kods / *UMDNS code*
Pamata UDI-DI / *Basic UDI-DI*Q010101
35870
16-724
++J0141103COMP010103i8Cklase / *of class*

IIa

saskaņā ar noteikumu / *according to rule*8-1, 19-3 saskaņā ar Medicīnisko ierīču regulas 2017/745
VIII pielikumu / *according to Annex VIII of Medical Device*
*Regulation 2017/745***atbilst visiem tiem Medicīnisko ierīču regulas 2017/745 noteikumiem, kas uz to attiecas. /**
meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.Izmantotie saskaņotie standarti, nacionālie standarti
vai citi normatīvie dokumenti / *Applied harmonised*
standards, national standards or other normative
*documents*EN ISO 4049 - Dentistry- Polymer-based restorative materials
Citus izmantotos standartus skatiet tehniskajā dokumentācijā par
produktu Charisma Bulk Flow One, 03. versija
Further Applied standards see Technical Documentation of
*Product Charisma Bulk Flow One, Version 03*Atbilstības novērtēšanas procedūra saskaņā ar /
*Conformity assessment procedure acc. to*Medicīnisko ierīču regulas 2017/745 IX pielikumu, I nodaļu, 2. un
3. iedaļu un III nodaļu*Medical Device Regulation 2017/745 Annex IX, Chapter I, Section*
*2 and 3 and Chapter III*Pilnvarotā iestāde / *Notified Body*TÜV Rheinland LGA Products GmbH
Tillystrasse 2
90431 Nürnberg/Vācija

CE 0197

Reģistrācijas numurs / *Registration number:*

HZ 1198082-1

Versijas numurs / *Version number*

03

Aizstāj atbilstības deklarāciju, kas datēta ar /
Replaces Declaration of Conformity from

08.08.2024

Hānava 21.10.2024

v.i. Dr. Matthias Hartmann
Head of Global Quality, Regulatory & Scientific Services
Kulzer GmbHVieta, datums / *Place, date*Vārds, uzvārds un amats / *Name and function*

Šis paziņojums par atbilstību ir derīgs 2 gadus saistībā ar ražoto ierīču atbilstošās partijas izlaišanas dokumentiem. / *This statement of conformity is valid for 2 years in connection with the release documents for the respective batch of produced devices.*

Preču saraksts / List of Articles

Pielikums / Annex: Atbilstības deklarācija / Declaration of Conformity

Medicīniskā ierīce /
The medical device

Charisma Bulk Flow One

Versijas numurs / *Version number*

04

Aizstāj pielikumu, kas datēts ar /
Replaces Annex from

08.08.2024

Šis izstrādājumu saraksts ir derīgs šādai
atbilstības deklarācijas versijai / *This article
list is valid for the declaration of conformity
version*

03

UDI-DI / <i>UDI-DI</i>	Artikelnummer / <i>Article number</i>	Name / <i>Name</i>
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Hānava 21.10.2024

Vieta, datums / *Place, date*



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

Vārds, uzvārds un amats / *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

Charisma Bulk Flow OneNazwa, 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 itemsWykaz wyrobów znajduje się w załączniku / List of Articles see
AnnexKod wyrobu wg EMDN / EMDN-Code
Kod wyrobu wg GMDN / GMDN code
Kod wyrobu wg UMDNS / UMDNS code
Kod Basic UDI-DI / Basic UDI-DIQ010101
35870
16-724
++J0141103COMP010103i8C

klasy / of class

IIa

zgodnie z regułą / according to rule

8-1, 19-3 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
documentsEN ISO 4049 - Dentistry- Polymer-based restorative materials
Pozostałe stosowane normy znajdują się w dokumentacji
technicznej produktu Charisma Bulk Flow One, wersja 03
Further Applied standards see Technical Documentation of
Product Charisma Bulk Flow One, Version 03Procedura oceny zgodności wg. /
Conformity assessment procedure acc. toRozporządzenie 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

03

Zastępuje Deklarację zgodności z /
Replaces Declaration of Conformity from

08.08.2024

Hanau, 21.10.2024

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

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

Charisma Bulk Flow One

Numer wersji / Version number

04

Zastępuje załącznik z dnia /
Replaces Annex from

08.08.2024

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

03

UDI-DI / UDI-DI	Artikelnummer / Article number	Name / Name
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Hanau, 21.10.2024

Miejscowość, data / Place, date



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

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*

Charisma Bulk Flow One

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*

Q010101
35870
16-724
++J0141103COMP010103i8C

da classe / *of class*

Ila

em conformidade com o regulamento / *according to rule*

8-1, 19-3 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

EN ISO 4049 - Dentistry– Polymer-based restorative materials
Outras normas aplicadas, ver Documentação técnica do produto
Charisma Bulk Flow One, Versão 03
Further Applied standards see Technical Documentation of
Product Charisma Bulk Flow One, Version 03

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*

03

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

08.08.2024



Hanau, 21.10.2024

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

Charisma Bulk Flow One

Número de versão / *Version number*

04

Substitui o Anexo de /
Replaces Annex from

08.08.2024

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*

03

UDI-DI / <i>UDI-DI</i>	Artikelnummer / <i>Article number</i>	Name / <i>Name</i>
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Hanau, 21.10.2024

Local, data / *Place, date*



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

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*

Charisma Bulk Flow OneNume, 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*Q010101
35870
16-724
++J0141103COMP010103i8Cdin clasa / *of class*

IIa

în conformitate cu regula / *according to rule*8-1, 19-3 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*EN ISO 4049 - Dentistry- Polymer-based restorative materials
Alte standarde aplicate, vezi documentația tehnică a Produsului
Charisma Bulk Flow One, Versiunea 03
*Further Applied standards see Technical Documentation of
Product Charisma Bulk Flow One, Version 03*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*

03

Înlocuiește Declarația de conformitate din /
Replaces Declaration of Conformity from

08.08.2024

Hanau, 21.10.2024

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

Charisma Bulk Flow One

Număr versiune / *Version number*

04

Înlocuiește Anexa de la /
Replaces Annex from

08.08.2024

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

03

UDI-DI / <i>UDI-DI</i>	Artikelnummer / <i>Article number</i>	Name / <i>Name</i>
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Hanau, 21.10.2024

Loc, dată / *Place, date*


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

Nume și funcție / *Name and function*

**IZJAVA O SKLADNOSTI / DECLARATION OF CONFORMITY**Ime in naslov podjetja /
*Name and address of the company***Kulzer GmbH**
Leipziger Straße 2, 63450 Hanau
Nemčija / Germany

SRN: DE-MF-000007705

Z izključno odgovornostjo izjavljamo, da / We declare under our sole responsibility thatmedicinski pripomoček / *the medical device***Charisma Bulk Flow One**Ime, vrsta ali model, številka šarže ali serijska številka, po možnosti izvor in število izdelkov /
*Name, type or model, batch or serial number, possibly sources and number of items*Seznam artiklov je na voljo v Prilogi / *List of Articles see Annex*Koda EMDN / *EMDN-Code*
Koda GMDN / *GMDN code*
Koda UMDNS / *UMDNS code*
Osnovni UDI-DI / *Basic UDI-DI*Q010101
35870
16-724
++J0141103COMP010103i8Crazreda / *of class*

IIa

v skladu s členom / *according to rule*8-1, 19-3, v skladu s Prilogo VIII Uredbe o medicinskih pripomočkih 2017/745 / *according to Annex VIII of Medical Device Regulation 2017/745***izpolnjuje vse določbe Uredbe o medicinskih pripomočkih 2017/745, ki veljajo zanj. /
*meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.***Uveljavljeni usklajeni standardi, nacionalni standardi ali drugi normativni dokumenti / *Applied harmonised standards, national standards or other normative documents*EN ISO 4049 - Dentistry- Polymer-based restorative materials
Drugi uveljavljeni standardi so na voljo v Tehnični dokumentaciji izdelka Charisma Bulk Flow One, različica 03
*Further Applied standards see Technical Documentation of Product Charisma Bulk Flow One, Version 03*Postopek ugotavljanja skladnosti v skladu z /
Conformity assessment procedure acc. to

Uredbo o medicinskih pripomočkih 2017/745, Priloga IX, poglavje I, oddelka 2 in 3, poglavje III

*Medical Device Regulation 2017/745 Annex IX, Chapter I, Section 2 and 3 and Chapter III*Priglašeni organ / *Notified Body*TÜV Rheinland LGA Products GmbH
Tillystrasse 2
90431 Nürnberg/Nemčija

CE 0197

Registrska številka / *Registration number*

HZ 1198082-1

Številka različice / *Version number*

03

Nadomešča Izjavo o skladnosti z dne /
Replaces Declaration of Conformity from

08.08.2024

Hanau, 21.10.2024


zastopnica Dr. Matthias Hartmann
Head of Global Quality, Regulatory & Scientific Services
Kulzer GmbHKraj, datum / *Place, date*Ime in položaj / *Name and function*Ta izjava o skladnosti je veljavna 2 leti v povezavi z dokumenti o izdaji za zadevne serije proizvedenih pripomočkov. / *This statement of conformity is valid for 2 years in connection with the release documents for the respective batch of produced devices.*

Seznam artiklov / List of Articles

Priloga / Annex: Izjava o skladnosti / Declaration of Conformity

Medicinski pripomoček /
The medical device

Charisma Bulk Flow One

Številka različice / Version number

04

Nadomešča Prilogo z dne /
Replaces Annex from

08.08.2024

Ta seznam izdelkov velja za naslednjo
različico izjave o skladnosti / This article list is
valid for the declaration of conformity version

03

UDI-DI / UDI-DI	Artikelnummer / Article number	Name / Name
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Hanau,^{21.10.2024}


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

Kraj, datum / Place, date

Ime in položaj / Name and function

**VYHLÁSENIE O ZHODE / DECLARATION OF CONFORMITY**

Názov a adresa spoločnosti /
Name and address of the company

Kulzer GmbH
Leipziger Straße 2, 63450 Hanau
Nemecko / Germany

SRN: DE-MF-000007705

Vyhlasujeme na svoju výlučnú zodpovednosť, že / We declare under our sole responsibility that
zdravotnícka pomôcka / the medical device

Charisma Bulk Flow One

Názov, typ alebo model, číslo šarže alebo sériové
číslo, prípadne zdroje a počet kusov / Name, type or
model, batch or serial number, possibly sources and
number of items

Zoznam položiek je uvedený v prílohe / List of Articles see Annex

Kód EMDN / EMDN-Code

Q010101

Kód GMDN / GMDN code

35870

Kód UMDNS / UMDNS code

16-724

Základné identifikačné číslo UDI-DI / Basic UDI-DI

++J0141103COMP010103i8C

triedy / of class

Ila

podľa pravidiel / according to rule

8-1, 19-3 podľa prílohy VIII k nariadeniu
2017/745 o zdravotníckych pomôckach / according to Annex VIII
of Medical Device Regulation 2017/745

spĺňa všetky ustanovenia nariadenia 2017/745 o zdravotníckych pomôckach, ktoré sa na ňu vzťahujú. /
meets all the provisions of the Medical Device Regulation 2017/745 which apply to it.

Použitie harmonizované normy, národné normy
alebo iné normatívne dokumenty / Applied
harmonised standards, national standards or other
normative documents

EN ISO 4049 - Dentistry- Polymer-based restorative materials
Ďalšie použité normy nájdete v technickej dokumentácii verzie 03k
produktu Charisma Bulk Flow One
Further Applied standards see Technical Documentation of
Product Charisma Bulk Flow One, Version 03

Postup posúdenia zhody podľa /
Conformity assessment procedure acc. to

prílohy IX k nariadeniu 2017/745 o zdravotníckych pomôckach,
kapitola I, časť 2 a 3 a kapitola III

Medical Device Regulation 2017/745 Annex IX, Chapter I, Section
2 and 3 and Chapter III

Notifikovaný orgán / Notified Body

TÜV Rheinland LGA Products GmbH
Tillystrasse 2
90431 Nürnberg / Nemecko

CE 0197

Registračné číslo / Registration number:

HZ 1198082-1

Číslo verzie / Version number

03

Nahrádza vyhlásenie o zhode z /
Replaces Declaration of Conformity from

08.08.2024

Hanau, 21.10.2024

i.V.


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

Miesto, dátum / Place, date

Meno a funkcia / Name and function

Toto vyhlásenie o zhode je platné 2 roky v súvislosti s dokumentmi o uvoľnení príslušnej šarže vyrobených pomôcok. /
This statement of conformity is valid for 2 years in connection with the release documents for the respective batch of produced
devices.

Zoznam položiek / List of Articles
Príloha / Annex: Vyhlásenie o zhode / Declaration of Conformity

Zdravotnícka pomôcka /
The medical device

Charisma Bulk Flow One

Číslo verzie / Version number

04

Nahrádza prílohu z /
Replaces Annex from

08.08.2024

Tento zoznam tovaru je platný pre vyhlásenie
o zhode, verzia / This article list is valid for
the declaration of conformity version

03

UDI-DI / UDI-DI	Artikelnummer / Article number	Name / Name
+J014660955200	66095520	CHARISMA BULK FLOW SYR 1 x 2G ONE ROW
+J014660953360	66095336	CHARISMA BULK FLOW ONE VALUE-KIT SYRINGE
+J014660947250	66094725	CHARISMA BULK FLOW SYR 1 X 2G ONE

Hanau, 21.10.2024

Miesto, dátum / Place, date



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

Meno a funkcia / Name and function

Abschlusszertifikat

Umschlag-ID: DE58ADD645A04D78BF67F809D7CF90EA		Status: Abgeschlossen
Betreff: Mit Docusign abschließen: Charisma Bulk Flow One_MDR_DoC_V03_2024-10-xx_Annex_V04_2024-10-xx.pdf		
Quellumschlag:		
Dokumentenseiten: 30	Signaturen: 30	Umschlagsteller:
Zertifikatsseiten: 1	Initialen: 0	Daniela Wienhold
Signatur mit Anleitung: Aktiviert		Leipziger Str. 2
Umschlag-ID-Stempel: Aktiviert		Hanau, Hessen 63450
Zeitzone: (UTC+01:00) Amsterdam, Berlin, Bern, Rom, Stockholm, Wien		Daniela.Wienhold@kulzer-dental.com
		IP-Adresse: 52.233.243.152

Eintragsverfolgung

Status: Original	Inhaber: Daniela Wienhold	Standort: DocuSign
18.10.2024 14:52:37	Daniela.Wienhold@kulzer-dental.com	

Unterzeichnerereignisse	Signatur	Zeitstempel
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Matthias Hartmann		Gesendet: 18.10.2024 14:56:57
matthias.hartmann@kulzer-dental.com		Eingesehen: 21.10.2024 09:04:23
Head of Global Quality, Regulatory & Scientific Services		Signiert: 21.10.2024 09:04:51
Sicherheitsstufe: E-Mail, Kontoauthentifizierung (keine)	Signaturübernahme: Hochgeladenes Signaturbild Mit IP-Adresse: 77.186.183.50	

Vereinbarung bezüglich elektronischer Unterlagen und Signaturen:
Nicht über DocuSign angeboten

Vor-Ort-Unterzeichner – Ereignisse	Signatur	Zeitstempel
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Bearbeiterversandereignisse	Status	Zeitstempel
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Beauftragtenzustellereignisse	Status	Zeitstempel
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Vermittlerversandereignisse	Status	Zeitstempel
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Zertifizierter Versand - Ereignisse	Status	Zeitstempel
-------------------------------------	--------	-------------

Kopienereignisse	Status	Zeitstempel
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Anette Stadtfeld	Kopiert	Gesendet: 18.10.2024 14:56:58
Anette.Stadtfeld@kulzer-dental.com		Eingesehen: 22.10.2024 08:33:55
Sicherheitsstufe: E-Mail, Kontoauthentifizierung (keine)		

Vereinbarung bezüglich elektronischer Unterlagen und Signaturen:
Nicht über DocuSign angeboten

Zeugen-Ereignisse	Signatur	Zeitstempel
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Notarereignisse	Signatur	Zeitstempel
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Umschlagereignisse – Überblick	Status	Zeitstempel
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Umschlag gesendet	Hash-codiert/verschlüsselt	18.10.2024 14:56:58
Zertifiziert zugestellt	Sicherheitsprüfung ausgeführt	21.10.2024 09:04:23
Signiervorgang abgeschlossen	Sicherheitsprüfung ausgeführt	21.10.2024 09:04:51
Abgeschlossen	Sicherheitsprüfung ausgeführt	21.10.2024 09:04:51

Zahlungen	Status	Zeitstempel
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