

## EU Declaration of Conformity

for a medical device class I (according to the EU regulation 2017/745, annex IV)

The manufacturer:

**Franz Mensch GmbH**  
**Werner-von-Siemens-Str. 2**  
**D - 86807 Buchloe**

declares under sole responsibility, that the class I medical device according to the classification rules of the EU regulation 2017/745, annex VIII

<b>Item REF</b>	<b>27066</b>
<b>Description</b>	<b>Nitrile gloves Extra Safe   powder-free</b>
<b>Brand</b>	<b>Hygostar</b>
<b>Version</b>	<b>Packing unit: Carton</b>
	<b>Color: Blue</b>
	<b>Size: 7/S</b>
	<b>Length: 24cm</b>

complies and meets all the provisions of the conformity assessment procedure of the EU regulation 2017/745 (annex I).

in accordance with confirmed records, test results or certificates, complies/comply with the requirements of the relevant harmonization legislation:

EN 455-1:2020  
EN 455-2:2015+A1:2011  
EN 455-3:2015  
EN 455-4:2009

The achieved performance levels of this medical device correspond to:

Signed for and on behalf of Franz Mensch GmbH,

Buchloe, 31.05.2021



Amanda Kreuzmann  
Head of Quality Management