

Fielmann

Zentrale



Name of the manufacturer: **Fielmann Group AG
Weidestraße 118a
22083 Hamburg**

Single Registration Number (SRN) **DE-MF-000006245**

We declare under our sole responsibility, pursuant to article 52 (7) MDR, the conformity with Regulation (EU) 2017/745 of the European Parliament and of the Council for the following product:

Basic UDI DI according to Annex VI MDR: **4066209plafuladu38**

Product name: **MF 031 FA PEPE**

Risk class: **1 (according to rules set out in Annex VIII MDR)**

Intended use: **Spectacle frames to be combined with corrective lenses to treat refractive errors**

The above-mentioned product is a medical device according to article 2 (1) of the Regulation. It fulfils the basic safety and performance requirements according to Annex I of the Regulation and complies with the state of the art and applicable harmonized standards.

Hamburg, 20.03.2024

Place, date

Valter Larentis
Qualified Person (Art. 15 MDR)

Fielmann
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