

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: **LN 009 MOD II FLEX CL**

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, 16.01.2024

Place, date

Valter Larentis
Qualified Person (Art. 15 MDR)

fielmann
Fielmann Group AG
Weidestraße 118a
22083 Hamburg