

**EC-Certificate****Directive 93/42/EEC, Annex II excluding (4)****Full Quality Assurance System**

Benannt durch/Designated by  
Zentralstelle der Länder  
für Gesundheitsschutz  
bei Arzneimitteln und  
Medizinprodukten  
www.zlg.de  
ZLG-BS-207.15.04

**Berlin Cert**

Prüf- und Zertifizierstelle für Medizinprodukte GmbH

hereby certifies that

**miha bodytec GmbH**

Siemensstr.1, 86368 Gersthofen, Germany

has implemented and uses a quality assurance system for the following scope of application:

**Development, production and final inspection of devices for electrical stimulation of muscles (see appendix)**

The audit in accordance with Annex II of MDD 93/42/EEC (report no. B-18-140-S-EZ) provided confirmation that the requirements of Annex II of MDD 93/42/EEC have been fulfilled. The Manufacturer has to be inspected periodically by the notified body according the requirements of Annex II, Article 5 of MDD 93/42/EEC. The manufacturer is allowed to use this certification in his process for the declaration of conformity.

The manufacturer is allowed to place the CE-mark on the above-mentioned products in combination with the identification No. **0633**.

issued on: **12.04.2021**valid from: **12.04.2021**valid to: **20.02.2024**

Prüf- und Zertifizierstelle  
**BERLIN**  
**CERT**  
AFNOR Group  
miha bodytec GmbH  
Dipl.-Ing. Martin Tettke  
Signature of authorized representative

