

Zeulenroda-Triebes, 08.05.2020

EC Declaration of Conformity

(According to Appendix VII of the Directive 93/42/EEC for medical devices)

We,

Bauerfeind AG
Triebeser Str. 16
D-07937 Zeulenroda-Triebes,

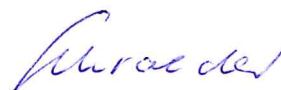
declare in exclusive accountability, that our products

VenoTrain® business, VenoTrain® clinic, VenoTrain® cocoon, VenoTrain® delight, VenoTrain® delight T, VenoTrain® discretion, VenoTrain® glider, VenoTrain® glider plus, VenoTrain® impuls, VenoTrain® look, VenoTrain® micro, VenoTrain® pure, VenoTrain® soft, VenoTrain® soft S, VenoTrain® ulcertec

and their spares, in case there exist any, comply with the relevant regulations of the Directive 93/42/EEC for class 1 medical devices.



Ines Exner
Authorized MD representative
Bauerfeind AG



Petra Schroeder
Quality and Regulatory Affairs
Technical Documentation

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