



MEDES limited
AUTHORIZED REPRESENTATIVE

CERTIFICATE OF CONFORMANCE

Acting as Regulatory Authorized Representative in Europe for **Perflex Ltd., from Israel**, we hereby declare that:

The following products listed hereunder or in the "Products Schedule" have been registered with the European Competent Authority in the UK (MHRA) on 5 of May 2009 Ref. No. CA010953 As Class I.

Products group

A Denture Base Polymer

By submitting the information to the MHRA, the legal requirements for registration have been met. **Perflex Ltd.** complies with the requirements of the Medical Device Directive 93/42 EEC amended by 2007/47 and the above products can be CE marked according to the above directive.

Product Schedule

Sr. No.	Product Name/Family
1	Perflex FN - Flexi Nylon
2	Perflex TC - T-Crystal
3	Perflex BS - BioSens
4	Perflex AC - Acetal
5	Perflex AF - Acry Free
6	Perflex TF - Thermofix
7	Perflex PPP - Perflex Pure Permanent
8	Perflex EC - Easy Clasp

For Medes Ltd.

Benny Arazy
General Manager
Medes Ltd.

Date: 27.11.14



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