

EC Declaration of Conformity

Kinetik Medical Devices Ltd, herewith declare that the products

Fully Automatic Blood Pressure Monitor - WBP1

meets the provisions of Directive 93/42/EEC which apply to them.

The medical devices has been assigned to class IIa according to Annex IX of the Directive 93/42/EEC. It bears the mark



The product concerned has been designed and manufactured under a quality management system according to Annex II of Directive 93/42/EEC.

Compliance of the designated product with the Directive 93/42/EEC has been assessed and certified by the Notified Body

TÜV Rheinland LGA Products GmbH Tillystraße 2, 90431, Nürnberg, Germany

Certificate No.: HD 60147729 0001 Issue date: 2020-08-24 Expiry date: 2024-05-26

following the procedure relating to the EC Declaration of Conformity set out in Annex II of Directive 93/42/EEC.

This Declaration of conformity is valid in connection with the release document fo the respective batch of produced devices.

Place / Date:	Surrey / November 3, 2020
	JRGrover
Signed: Name:	James Grover
Position:	Sales Director