



CEC
International

CEC International s.a., Hrabľová 18, 821 06 Bratislava, Slovakia
Notified Body No. 2265

EC CERTIFICATE

No. 2015-MDD/QS-010

Issued in compliance with the Council Directive 93/42/EEC as amended by 2007/47/EC, which is implemented by the Slovak Government Decree No. 582/2008 Coll. as amended by 215/2013 Coll., certifies that the medical device of Class IIa

Blood Bank Centrifuge
Model KBM 70 Plus

manufactured by company

REMI ELEKTROTECHNIK LTD.

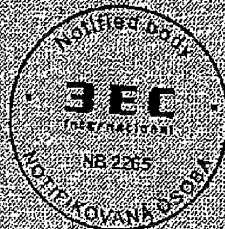
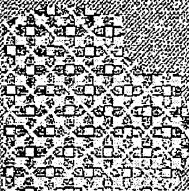
"B" Building, Survey No. 65/1, Vally, Vasai (East), Thane - 401208, Maharashtra, India

is manufactured under conditions fulfilling the quality system requirements of Annex II, excluding (4), of the Directive 93/42/EEC as amended by 2007/47/EC.

The Notified Body No. 2265 has performed an audit of the above device quality system. The full quality assurance system has been assessed and found that it meets the requirements above. The quality system is subject to continuous surveillance according to Annex I, Sections 3.3, and 5, of the Directive 93/42/EEC as amended by 2007/47/EC. The detailed description of the system, requirements and measures applied by the manufacturer are presented in the Audit Report No. 310109, and the Final protocol No. 310109/2011 that is enclosed to this certificate.

This certificate is issued under the following conditions:

It applies only to the quality system maintained in the manufacture of the above referenced model of medical device and it does not substitute the design or type examination procedures, if requested. The certificate remains valid until the manufacturing conditions or the quality system are changed but until May 15th, 2016 at the latest. The certificate validity is conditional upon positive results of regular surveillance audits and fulfilment of relevant legal and other requirements by manufacturer.



Dr. Katarína Šimová
Responsible to act on behalf of NB 2265

REMI ELEKTROTECHNIK LIMITED

In Bratislava, on May 15th, 2015