

EC CERTIFICATE

Certificate No 1959/MDD

Production Quality Assurance System Approval Certificate

On the basis of our assessment carried out according to Annex V, section 3 and considering the Annex VII, section 5 of the Directive 93/42/EEC and its revised version, we hereby certify that:

KORPO SRL

16121 GENOVA (GE) - VIA XX SETTEMBRE 3/28 (ITA) - Italy

manages in the factories of:

16121 GENOVA (GE) - VIA XX SETTEMBRE 3/28 (ITA) - Italy

a quality assurance system ensuring the conformity of the following products:

Adipose tissue blender
Type ref. ADIPOPIMER
Trade mark KORPO

with the relevant essential requirements of the aforementioned directive (concerning the manufacturing stage relevant to the reaching and the keeping of the sterile conditions) and it is subject to surveillance as specified in section 4 of Annex V.

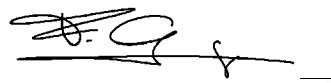

Reference to IMQ files Nos:

DM17-0008198-01; DM17-0019151.

This Approval Certificate is issued by IMQ S.p.A. as Notified Body for the Directive 93/42/EEC and its revised version.

Notified Body notified to European Commission under number: 0051.

Date: 2017-09-07
Updated: 2018-05-03
Substitution Date: 2017-09-07
Expiry Date: 2023-05-02


IMQ 

 **IMQ** 
ISTITUTO ITALIANO DEL MARCHIO DI QUALITA'

IMQ S.p.A. - I-20138 Milano
Via Quintiliano 43
tel. + 39 0250731
www.imq.it

This Approval Certificate is subjected to the provisions laid down in the "Rules for managing the EC Certification of Medical Devices on the basis of the Directive 93/42/EEC".

This is a translation of the Italian text, which prevails in case of doubts