



FOR AFR 012 / 02 Version 5.0

2.3 EC Declaration of conformity

We herewith declare on our sole responsibility, that the design, production, and packaging of the described product is compliant with the specific requirements of the Directive 93/42/EEC concerning medical devices, that the product has been classified according to the rules of classification of the Annex IX of the Directive 93/42/EEC and satisfied all requirements of the Annex II (without Section 4) of the Directive 93/42/EEC.

Product terralin PAA

Item code 126203

Manufacturer BIOXAL SA – Route des Varennes - 71100 CHALON-SUR-SAONE –

France

Notified Body DQS Medizinprodukte GmbH

August-Schanz-Str. 21

D-60433 Frankfurt am Main GERMANY

Notified Body EC 0297

Class of the medical device

(Directive 93/42/EEC, Annex IX, Rule 15)

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Product group Disinfectant, medical devices

Product category (EN ISO 15225) Hospital hardware

Issued certificates EN ISO 9001 _ Cert. Reg. No. 368588 QM15

EN ISO 13485_ Cert. Reg. No. 368588 MP2016

Annex II _ Cert. Reg. No. 368588 MR2

Standards applied Applied standards are listed in Sec. 2.4 of the technical

documentation.

I, the undersigned, declare that BIOXAL SA, bears the sole responsibility for issuing this Declaration.

Position of the responsible person

General Manager

Name of the responsible person

Sylvain LEMAIRE

Location

Chalon-sur-Saône

Date of issue

14/12/2022

Sianature

This Declaration is valid until an updated version has been issued, but not longer than 2024-05-26. Localization of the technical documentation: Bioxal SA, Regulatory Affairs office.