

TO WHOM IT MAY CONCERN

Whereas ANTONIO MATACHANA, S.A who are established and reputable manufacturers of Infection Control equipments, with headquarters in Calle Almogávares 174-176, 08018 Barcelona (Spain),

Do hereby certify that the company Echipamed PLUS ERL, with business address at Valea Trandafirilor, 24B, of.80 – MD-2001 Chisinau, Republic of Moldova.

They are fully authorised to perform after sales activities (quote, sell spare parts) as well carry out maintenance activities in Moldova.

This sole agency agreement will be valid until 10th of January 2023.

And in proof of this, we hereunder sign the present certificate in the city of Barcelona, on the 10th of January 2022.

ANTONIO MATACHANA, S.A.



Ramón Torres

Technical Manager Europe & USA

000 matachana
Servicio Técnico Internacional
Service & Support
Raurell, 21-29 Nave 4
Poligono Industrial Camí Rei
08860-Castelldefels (Barcelona)



Certificate of Approval

This is to certify that the Management System of:

Antonio Matachana, S.A.

Almogavares, 174-176, 08018 Barcelona, Spain

has been approved by Lloyd's Register to the following standards:

ISO 9001:2015

Approval number(s): ISO 9001 – 0036999

This certificate is valid only in association with the certificate schedule bearing the same number on which the locations applicable to this approval are listed.

The scope of this approval is applicable to:

Design and manufacture of sterilizers, disinfection devices and treatment equipment for medical devices, food, associated research, industry, medical waste and pharmaceutical laboratories. Distribution of communities, surgical, sterilisation, disinfection and mortuary equipment. Technical and validation service at customer's premises (sterilizers, washer-disinfectors and sealing machines).

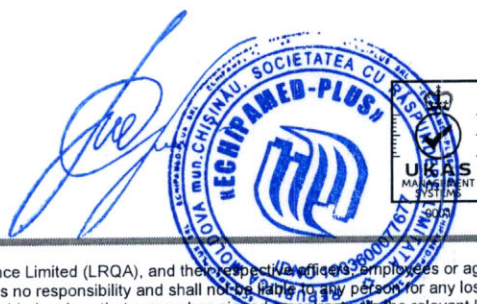


Daniel Oliva Marcilio de Souza

Area Operations Manager - South Europe

Issued by: Lloyd's Register Quality Assurance España, S.L.U.

for and on behalf of: Lloyd's Register Quality Assurance Limited



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Certificate

Standard **ISO 14001:2015**

Certificate Registr. No. 3.00.13062

TÜV Rheinland Ibérica Inspection, Certification & Testing S.A.
certifies:

Certificate Owner:

●●● matachana

ANTONIO MATACHANA, S.A.
Copèrnic, 8
E - 08860 Castelldefels (Barcelona)

Scope:

Design and manufacture of sterilizers, disinfection devices and treatment equipment for medical devices, food, associated research, industry, medical waste and pharmaceutical laboratories.

An audit was performed, Report No. 3.00.13062.
Proof has been furnished that the requirements according to ISO 14001:2015 are fulfilled.

The due date for all future audits is 10-02 (mm - dd).

Validity:

The certificate is valid from 2019-11-15 until 2022-11-14.
First certification 2013.

2019-11-04

TÜV Rheinland Ibérica Inspection, Certification & Testing S.A.
Garrotxa, 10-12 - E-08820 El Prat de Llobregat

DECLARACIÓN DE CONFORMIDAD CE

EC DECLARATION OF CONFORMITY / DECLARATION DE CONFORMITÉ CE

La empresa: ANTONIO MATACHANA, S.A.
The company: Almogàvers, 174
La société: E-08018 – Barcelona

Declara bajo su única responsabilidad que el equipo,
Declares that, in its own responsibility, the equipment
Déclare sous sa propre responsabilité, que l'appareil,

Marca: Matachana **Tipo:** Esterilizador de vapor.
Brand / Marque: Matachana *Type / Tipe:* Steam sterilizer / Stérilisateur à la vapeur d'eau.

Serie: S1000 **Modelos:** S1004 V-1, S1006 V-1, S1008 V-1, S1010 V-1, S1012 V-1,
Series / Série Models / Modèles: S1004 E-1, S1006 E-1, S1008 E-1, S1010 E-1, S1012 E-1,
 S1004 V-2, S1006 V-2, S1008 V-2, S1010 V-2, S1012 V-2,
 S1004 E-2, S1006 E-2, S1008 E-2, S1010 E-2, S1012 E-2

siendo productos en el ámbito de la Directiva Europea 93/42/CEE por la que se regulan los
productos sanitarios y clasificado como clase IIb según la regla 15 del Anexo IX,
Being devices within the scope of the regulation of the European Directive 93/42/EEC concerning Medical
Devices (MDD), and classified as Class IIb according to rule 15 of Annex IX therein,
étant dispositifs dedans le champ d'application de la Directive Européenne 93/42/CEE relative aux dispositifs
médicaux, et classifié comme Classe IIb selon la règle 15 de l'Annexe IX,

se hallan en conformidad con:

conforms to / sont conformes à:

- **Directiva 93/42/CEE relativa a los Productos Sanitarios, Anexo I.**
- *Directive 93/42/EEC concerning Medical Devices, Annex I.*
- *Directive 93/42/CEE relative aux dispositifs médicaux, Annexe I.*

Evaluación de la conformidad según:

Conformity assessment according to: / Évaluation de la conformité selon:

- **Directiva 93/42/CEE relativa a los Productos Sanitarios, Anexo II excepto II.4**
- *Directive 93/42/EEC concerning Medical Devices, Annex II without II.4*
- *Directive 93/42/CEE relative aux dispositifs médicaux, Annexe II à l'exception du II.4*

Con número de registro HD 60029459 0001 del Organismo Notificado Europeo:

Under the Registration number
Avec registre numéro

from the European Notified Body number:
de l'Organisme Notifié Européen :

TÜV Rheinland LGA Products GmbH
An Grauen Stein
D-51105 Köln



Además de la Directiva de Productos Sanitarios 93/42/CEE los productos mencionados o sus
componentes cumplen con los requisitos esenciales de las siguientes Directivas que son relevantes:
Additionally to the Medical Device Directive 93/42/EEC the a. m. devices or the included components fulfil the
essential requirements of the following relevant Directives:
En addition à la Directive relative aux Dispositifs Médicaux 93/42/CEE les dispositifs mentionnés si dessus les
exigences essentielles des suivantes Directives relevantes:

EMC 2004/108/EC, LVD 2006/95/EC, MD 2006/42/EC, PED 97/23/EC

Estando disponibles las correspondientes Declaraciones o Certificados de Conformidad emitidos por los fabricantes.

Respective Declarations or Certificates of Conformity issued by the manufacturers are available.

Étant disponibles les respectives Déclarations ou Certifications de Conformité livrées par les fabricants

Nombre:

Sagué

Name / Nom:

Jose Carlos López

Director Industrial

ANTONIO MATACHANA, S.A. Ramón Calvera

Calle Hierro, 20-22
 E-08038 Barcelona - SPAIN
 Tel. +34.93 223 26 28
 Fax + 34.93 223 33 61

Cargo:

Title / Position:

Director de Calidad

Lugar y fecha:

Barcelona, 14 de mayo de 2010

Place and date / lieu et date:

