

EC CERTIFICATE

for the Quality Assurance System



according the Directive 93/42/EEC, Annex II excluding section (4)

As a Notified Body of the European Union, DEKRA Certification GmbH certifies, that the company
E.M.S. Electro Medical Systems S.A.

Chemin de la Vuarpillière 31, 1260 Nyon, Switzerland

Certified location:

Chemin de la Vuarpillière 31, 1260 Nyon, Switzerland
E.M.S. Electro Medical Systems S.A., Rte. de Champ-Colin 2, 1260 Nyon, Switzerland
E.M.S. Electro Medical Systems S.A., Rte. de Champ-Colin 18, 1260 Nyon, Switzerland
E.M.S. Foncine S.A.S., Rte. de Pontarlier 32, 39460 Foncine-le-Haut, France

applies a quality assurance system according to the Directive 93/42/EEC Annex II for the medical devices listed in the annex. The approval is based on the result of the re-certification audit report no. 50081-Z7-00, the decision dated 2020-06-15 and is only valid in connection with the successful performance of the annual surveillance audits.

This certificate is valid from 2020-06-15 to 2024-05-26

Registration No.: 50081-16-09



Ruth Delbeck-Bayer
DEKRA Certification GmbH Stuttgart; 2020-06-15
Notified Body ID-number: 0124

DEKRA Certification GmbH * Handwerkstraße 15 * D-70565 Stuttgart * www.dekra-certification.de



Benannt durch/Designated by
Zentralstelle der Länder
für Gesundheitsschutz
bei Arzneimitteln und
Medizinprodukten
www.zlg.de
ZLG-BS-295.10.02

Annex to the EC Certificate No. 50081-16-09

Valid from 2020-06-15 to 2024-05-26

Revision status of the annex: 0 dated 2020-06-15

Devices/device categories included in the certificate:

Class I s:

For the products listed below, review of the Quality Assurance System refers exclusively to aspects of manufacture concerned with securing and maintaining sterile conditions.

IntraCorporal Lithotripters systems
- Stone Catcher

Class II a:

IntraCorporal Lithotripters systems
- Swiss LITHOPUMP
- Probes (pneumatic, ultrasound and combined)
- Probes Sets (pneumatic, ultrasound and combined)

Piezo-ceramic systems for dentistry - Piezon
- PIEZON product group
- PIEZON Surgery product group
- PIEZON Handpieces, PIEZON instruments

Air abrasion / air polisher devices for dental prophylaxis
- AIRFLOW product group
- PERIOFLOW nozzles
- AIRFLOW, PERIOFLOW Handpieces

Air abrasion / air polisher devices for dental prophylaxis with Piezo-ceramic systems for dentistry
- AIRFLOW / PIEZON product group

Surgical HO_YAG laser systems
- SmartFibers product group

Class II b:

IntraCorporal Lithotripters systems
- LITHOCLAST product group
- LITHOCLAST handpieces (pneumatic, ultrasound and combined)
- Handpiece Sets (pneumatic, ultrasound and combined)

ExtraCorporal shock wave therapy systems
- DOLORCLAST product group
- DOLORCLAST Handpieces, DOLORCLAST applicators

Surgical HO_YAG laser-system
- Swiss LASERCLAST product group


Ruth Delbeck-Bayer



DEKRA Certification GmbH, Stuttgart, 2020-06-15

Notified Body ID-number: 0124

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Signature Manifest

Document Number: CER-0002

Revision: D

Title: EC certificate for the Quality Assurance System according the Directive 93/42/EEC, Annex II excluding section (4)

All dates and times are in UTC+01:00.

CER-0002 EC certificate

Author and reviewers

Name/Signature	Title	Date	Meaning/Reason
Timothée Deblock (TDE)		29 Jun 2020, 03:46:27 PM	Approved

Manufacturer's Declaration

in relation to Regulation (EU) 2023/607 amending Regulations (EU) 2017/745 and (EU) 2017/746 as regards the transitional provisions for certain medical devices and in vitro diagnostic medical devices, in particular with respect to

- the validity of certificates issued under Council Directive 90/385/EEC on Active Implantable Medical Devices (AIMDD) or Council Directive 93/42/EEC on Medical Devices (MDD) (Directive Certificates) *and/or*¹
- the compliance of the devices and us as their manufacturer with the conditions for the continued placing on the market and putting into service

Manufacturer name	E.M.S. Electro Medical Systems S.A.
Manufacturer address and contact details	Chemin de la Vuarpillière 31, 1260 Nyon, Switzerland +41 22 994 47 00
Single Registration Number (SRN) (if available)	CH-MF-000026136

Authorised Representative name (if applicable)	E.M.S. Electro Medical Systems FRANCE SARL
Authorised Representative address and contact details	32, Route de Pontarlier 39460 Foncine-Le-Haut France +33 3 84 51 90 01
Single Registration Number (SRN) (if available)	FR-AR-000011266

Notified body name (if applicable)	DEKRA Certification GmbH
Notified body number (if applicable)	0124
Directive Certificate number(s) to which this confirmation is made (if applicable)	50081-16-09

¹ The first condition is not applicable in case of devices for which the conformity assessment procedure pursuant to MDD did not require the involvement of a notified body, for which the declaration of conformity was drawn up prior to 26 May 2021 and for which the conformity assessment procedure pursuant to this Regulation requires the involvement of a notified body.

Original expiry date as indicated on the Directive Certificate prior to the extension of the validity (if applicable)	2024.05.26
End date of extended validity/transition period	2028.12.31

We, as the manufacturer declare under our sole responsibility:

- for the above listed **Directive Certificate** (or see attached schedule, if multiple certificates) the conditions for the legal extension of validity as required in Article 120.2 of the MDR are met *and/or*²
- the listed **device(s)** in the attached schedule and we as their manufacturer are in compliance with the conditions listed in Article 120.3c of the MDR for continued placing on the market and putting into service,

namely by fulfilling the following conditions:

➤ **Directive Certificate(s)** as listed above or in the attached schedule

- Directive Certificate(s) covering the listed device(s) was/were issued after 25 May 2017, was/were valid on 26 May 2021 and have not been withdrawn afterwards.

Choose applicable statements:

Expired *before* 20 March 2023:

- Before the original date of expiry as indicated on the Directive Certificate(s), we and the notified body have signed written agreement(s) in accordance with Section 4.3, second subparagraph of Annex VII to this Regulation for the conformity assessment(s) in respect of the device(s) covered by the expired certificate(s) or in respect of a device(s) intended to substitute that/those device(s), or
- A Competent Authority has granted a derogation from the applicable conformity assessment procedure in accordance with Article 59(1) MDR (may be provided upon request), or
- A Competent Authority has required the manufacturer, in accordance with Article 97(1) MDR, to carry out the applicable conformity assessment procedure (may be provided upon request)

Choose one of the following statements only if a derogation per Article 59(1) or a requirement per Article 97(1) has been granted by a Competent Authority:

- Formal application(s) to the notified body in accordance with Section 4.3, first subparagraph of Annex VII MDR for conformity assessment has/have been made or will be made/submitted by us to a notified body no later than 26 May 2024 for the device(s) listed in the attached schedule or its/their substitute(s) and signed written agreement(s) is/will be in place in accordance with Section 4.3, second subparagraph of Annex VII MDR before 26 September 2024.

² The first condition is not applicable in case of devices for which the conformity assessment procedure pursuant to MDD did not require the involvement of a notified body, for which the declaration of conformity was drawn up prior to 26 May 2021 and for which the conformity assessment procedure pursuant to this Regulation requires the involvement of a notified body

We do not intent to lodge an application for conformity assessment by 26 May 2024, therefore the transition period will end on 26 May 2024.

Expired/expires *after* 20 March 2023:

Choose one applicable statement:

Formal application(s) to the notified body in accordance with Section 4.3, first subparagraph of Annex VII MDR for conformity assessment has/have been made or will be made/submitted by us to a notified body no later than 26 May 2024 for the device(s) listed in the attached schedule or its/their substitute(s) and signed written agreement(s) is/will be in place in accordance with Section 4.3, second subparagraph of Annex VII MDR before 26 September 2024.

We do not intent to lodge an application for conformity assessment by 26 May 2024, therefore the transition period will end on 26 May 2024.

➤ **Upclassified devices (case of AIRFLOW Powders)**

In case of devices for which the conformity assessment procedure pursuant to MDD did not require the involvement of a notified body, for which the declaration of conformity was drawn up prior to 26 May 2021 and for which the conformity assessment procedure pursuant to this Regulation requires the involvement of a notified body:

Choose one applicable statement:

Formal application(s) to the notified body in accordance with Section 4.3, first subparagraph of Annex VII MDR for conformity assessment has/have been made or will be made/submitted by us to a notified body no later than 26 May 2024 for the device(s) listed in the attached schedule or its/their substitutes and signed written agreement(s) is/will be in place in accordance with Section 4.3, second subparagraph of Annex VII MDR before 26 September 2024.

We do not intent to lodge an application for conformity assessment by 26 May 2024, therefore the transition period will end on 26 May 2024.

➤ **Quality Management System (QMS)**

Choose one applicable statement:

A QMS in accordance with Article 10(9) MDR will be put in place by no later than 26 May 2024.

A QMS in accordance with Article 10(9) MDR is in place.

A notified body has issued the attached certificate for the MDR-compliant QMS.

➤ **Device(s) as listed in the attached schedule**

- The device(s) continue to comply with the AIMDD or MDD.
- There are no significant changes in the design and intended purpose.
- The device(s) do not present an unacceptable risk to health or safety of patients, users or other persons, or to other aspects of the protection of public health.

Signed for and on behalf of the manufacturer: .

Full Company Name: E.M.S. Electro Medical Systems S.A.

Location & Date: Nyon, 29.04.2024

Signature, Print Name, Title:



Timothée Deblock
Head of Quality

Contact Details (at least email): ra-pcn@ems-ch.com

Schedule of Devices

The above Manufacturer's Declaration is valid for the following devices:

Identification of the device(s) ³ (e.g., device name, family/group name, device model or catalogue number)	Directive Certificate number(s) to which this confirmation is made (if applicable)	Original expiry date as indicated on the Directive Certificate (s) prior to the extension of the validity (if applicable)	Notified Body name and number that issued the Directive Certificate (if applicable)	Notified Body name and number where the MDR application was lodged/contract signed (if applicable)	End date of extended validity / transition period	Substitute Device(s) (if applicable)
BUDIDI Group name						
07613353001KQ LithoClast TRILOGY	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353008L6 LithoClast Master	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353013KX LithoClast 2	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353040L2 LITHO Handpieces	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353011KT LITHO Probes	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353048LJ LASER Fibers	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A

³ for devices with AIMDD/MDD certificate(s) the identification should be as in the certificate, and only if the certificate has a generic scope it should be as defined above)

Identification of the device(s) ³ (e.g., device name, family/group name, device model or catalogue number)	Directive Certificate number(s) to which this confirmation is made (if applicable)	Original expiry date as indicated on the Directive Certificate (s) prior to the extension of the validity (if applicable)	Notified Body name and number that issued the Directive Certificate (if applicable)	Notified Body name and number where the MDR application was lodged/contract signed (if applicable)	End date of extended validity / transition period	Substitute Device(s) (if applicable)
BUDIDI Group name						
07613353039LH Stone Catcher	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353051L7 Handpieces Suction Tubes	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353049LL Probes Suction Tubes	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353019LB LithoPump	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353002KS AIRFLOW Prophylaxis Master and AIRFLOW One	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353030KX PIEZON 250	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353028LC PIEZON Kits	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353033L5 AIRFLOW Handy 3.0	50081-16-09	2024.05.26	DEKRA Certification GmbH	DEKRA Certification GmbH	2028.12.31	N/A

Identification of the device(s) ³ (e.g., device name, family/group name, device model or catalogue number)	Directive Certificate number(s) to which this confirmation is made (if applicable)	Original expiry date as indicated on the Directive Certificate (s) prior to the extension of the validity (if applicable)	Notified Body name and number that issued the Directive Certificate (if applicable)	Notified Body name and number where the MDR application was lodged/contract signed (if applicable)	End date of extended validity / transition period	Substitute Device(s) (if applicable)
BUDIDI Group name			0124	0124		
07613353021KW PIEZON Handpieces	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353016L5 PERIOFLOW Handpieces	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353035L9 AIRFLOW Handpieces	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353037LD PIEZON Instruments	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353036LB PERIOFLOW Nozzles	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353034L7 Radial Shock Wave	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353031KZ DolorClast Master	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A

Identification of the device(s) ³ (e.g., device name, family/group name, device model or catalogue number)	Directive Certificate number(s) to which this confirmation is made (if applicable)	Original expiry date as indicated on the Directive Certificate (s) prior to the extension of the validity (if applicable)	Notified Body name and number that issued the Directive Certificate (if applicable)	Notified Body name and number where the MDR application was lodged/contract signed (if applicable)	End date of extended validity / transition period	Substitute Device(s) (if applicable)
BUDIDI Group name						
07613353026L8 DolorClast Smart 20	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353046LE DOLOR Handpieces	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353020KU AIRFLOW Powders	N/A, product up-classified under Regulation EU 2017/745			DEKRA Certification GmbH 0124	2028.12.31	N/A
07613353012KV DOLOR Applicators	50081-16-09	2024.05.26	DEKRA Certification GmbH 0124	Product down-classified as Class I under Regulation EU 2017/745		N/A

DEKRA Certification GmbH – Handwerkstraße 15 – D-70565 Stuttgart

E.M.S. Electro Medical Systems S.A.
Mr. Timothée Deblock
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1260 Nyon
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Date 2020-12-17

Decision on the audit 1 / certification audit

Certification

Regulation (EU) 2017/746 Annex IX Chapter I

Dear Mr. Deblock

Based on audit report no. 50081-R1-00 it has been verified that your quality management system complies with the requirements.

Please note that a technical documentation must be positively verified before a certificate according to Regulation (EU) 2017/745 can be issued.

Notes

Basis of the continuing validity of the certificate is the regular performance of a yearly surveillance audit during the certificate's period of validity. The intention of the surveillance audit is the evaluation of the Quality management system's continued effectiveness according to the corresponding requirements.

Please note that your next audits are aligned with the already scheduled audits according to Directive 93/42EEC and EN ISO 13485:2016 and are to be carried out with the following periods:

audit 2 / 1st surveillance audit: between **2021-05-23 and 2021-08-23**
audit 3 / 2nd surveillance audit: between **2022-05-23 and 2022-08-23**

The prices for these audits will be included in a separate supplement to offer no. A20021254.

Yours sincerely

DEKRA Certification GmbH



Markus Kopf



Enclosures:

Audit Report No. 50081-R1-00 (sent electronically)
Invoice (sent electronically)

DEKRA Certification GmbH
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Geschäftsführer:
Dr. Rolf Krökel, Thomas Thees

DEKRA Certification GmbH – Handwerkstraße 15 – D-70565 Stuttgart

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Date 2024-05-02

Subject: Notified Body Confirmation Letter

Our reference: 50081-CoL-00, Rev.0

Confirmation of the status of a formal application, written agreement, and appropriate surveillance in the framework of Regulation (EU) 2023/607 amending Regulations (EU) 2017/745 as regards the transitional provisions for certain medical devices and in vitro diagnostic medical devices

Dear Mr. Deblock

This letter confirms that, DEKRA Certification GmbH, a Notified Body (NB) designated against Regulation (EU) 2017/745 (MDR) and identified by the number 0124 on NANDO, has received a formal application in accordance with Section 4.3, first subparagraph of Annex VII of MDR and has signed a written agreement in accordance with Section 4.3, second subparagraph of Annex VII of MDR with the following manufacturer:

E.M.S. Electro Medical Systems S.A.
Ch. de la Vuarpillière 31
1260 Nyon
Switzerland

SRN Number: CH-MF-000026136

The devices covered by the formal application and the written agreement mentioned above are identified in the Table provided in the Annex. This table identifies the devices for which an MDR application has been received, written agreement concluded and for which the NB is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive MDD.

In the case of devices covered by certificates issued under Directive 93/42/EEC (MDD) that expired after 26 May 2021 and before 20 March 2023, without having been withdrawn, this letter also confirms that the manufacturer signed the written agreement under MDR by the date of MDD certificate expiry; or provided evidence that a competent authority of a Member State had granted a derogation or exemption from the applicable conformity assessment

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medizinprodukte

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IBAN: DE76 6008 0000 0901 4949 00
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Ust.-ID-Nr. DE 811 976 119

Managing director:
Dr. Rolf Krökel

procedure in accordance with Article 59(1) of MDR or Article 97(1) of the MDR respectively, by the 20 Mar 2023 for the relevant devices.

The transition timelines that apply to the devices covered by this letter, subject to the manufacturer's continued compliance to the other conditions specified in Article 120.3c of MDR (as amended by (EU) 2023/607), are shown below:

- 26 May 2026 for Class III custom-made implantable devices
- 31 December 2027 for Class III devices and Class IIb implantable devices excluding Well-established technologies (WET - sutures, staples, dental fillings, dental braces, tooth crowns, screws, wedges, plates, wires, pins, clips and connectors)
- 31 December 2028 for other Class IIb devices, Class IIa, Class I devices placed on the market in sterile condition or have a measuring function
- 31 December 2028 for devices not requiring the involvement of a notified body under MDD but requiring it under MDR (e.g., class I devices that qualify as re-usable surgical instruments)

On behalf of the Notified Body,

Stephanie Donner
2024-05-02

Enclosures:

Confirmation Letter Annex

Annex to Notified Body Confirmation Letter 50954-CoL-00, Rev.0

Devices covered by this letter and for which the Notified body DEKRA Certification GmbH is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<p>07613353001KQ LithoClast TRILOGY</p>	<p>Class IIb excluding Class IIb implantable non-WET</p>	<p>N/A</p>	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15 Annex Rev. 0, dated: 2020-06-15 Notified Body: DEKRA (0124)</p>
<p>07613353008L6 LithoClast Master</p>	<p>Class IIb excluding Class IIb implantable non-WET</p>	<p>N/A</p>	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15 Annex Rev. 0, dated: 2020-06-15 Notified Body: DEKRA (0124)</p>
<p>07613353013KX LithoClast 2</p>	<p>Class IIb excluding Class IIb implantable non-WET</p>	<p>N/A</p>	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15 Annex Rev. 0, dated: 2020-06-15 Notified Body: DEKRA (0124)</p>
<p>07613353040L2 LITHO Handpieces</p>	<p>Class IIb excluding Class IIb implantable non-WET</p>	<p>N/A</p>	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15 Annex Rev. 0, dated: 2020-06-15 Notified Body: DEKRA (0124)</p>
<p>07613353011KT LITHO Probes</p>	<p>Class IIa</p>	<p>N/A</p>	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15 Annex Rev. 0, dated: 2020-06-15 Notified Body:</p>

			DEKRA (0124)
07613353048LJ LASER Fibers	Class IIa	N/A	Certificate number: Certificate 50081-16-09; dated: 2020-06-15 Annex Rev. 0, dated: 2020-06-15 Notified Body: DEKRA (0124)
07613353039LH Stone Catcher	Class I devices placed on the market in sterile condition	N/A	Certificate number: Certificate 50081-16-09; dated: 2020-06-15 Annex Rev. 0, dated: 2020-06-15 Notified Body: DEKRA (0124)
07613353051L7 Handpieces Suction Tubes	Class I devices placed on the market in sterile condition	N/A	Certificate number: Certificate 50081-16-09; dated: 2020-06-15 Annex Rev. 0, dated: 2020-06-15 Notified Body: DEKRA (0124)
07613353002KS AIRFLOW Prophylaxis Master and AIRFLOW One	Class IIa	N/A	Certificate number: Certificate 50081-16-09; dated: 2020-06-15 Annex Rev. 0, dated: 2020-06-15 Notified Body: DEKRA (0124)
07613353030KX PIEZON 250	Class IIa	N/A	Certificate number: Certificate 50081-16-09; dated: 2020-06-15 Annex Rev. 0, dated: 2020-06-15 Notified Body: DEKRA (0124)
07613353028LC PIEZON Kits	Class IIa	N/A	Certificate number: Certificate 50081-16-09; dated: 2020-06-15 Annex Rev. 0, dated: 2020-06-15 Notified Body: DEKRA (0124)

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<p>07613353033L5 AIRFLOW Handy 3.0</p>	Class IIa	N/A	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15</p> <p>Annex Rev. 0, dated: 2020-06-15</p> <p>Notified Body: DEKRA (0124)</p>
<p>07613353021KW PIEZON Handpieces</p>	Class IIa	N/A	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15</p> <p>Annex Rev. 0, dated: 2020-06-15</p> <p>Notified Body: DEKRA (0124)</p>
<p>07613353016L5 PERIOFLOW Handpieces</p>	Class IIa	N/A	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15</p> <p>Annex Rev. 0, dated: 2020-06-15</p> <p>Notified Body: DEKRA (0124)</p>
<p>07613353035L9 AIRFLOW Handpieces</p>	Class IIa	N/A	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15</p> <p>Annex Rev. 0, dated: 2020-06-15</p> <p>Notified Body: DEKRA (0124)</p>
<p>07613353037LD PIEZON Instruments</p>	Class IIa	N/A	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15</p> <p>Annex Rev. 0, dated: 2020-06-15</p> <p>Notified Body: DEKRA (0124)</p>

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<p>07613353036LB PERIOFLOW Nozzles</p>	Class IIa	N/A	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15</p> <p>Annex Rev. 0, dated: 2020-06-15</p> <p>Notified Body: DEKRA (0124)</p>
<p>07613353034L7 Radial Shock Wave</p>	Class IIa	N/A	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15</p> <p>Annex Rev. 0, dated: 2020-06-15</p> <p>Notified Body: DEKRA (0124)</p>
<p>07613353031KZ DolorClast Master</p>	Class IIa	N/A	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15</p> <p>Annex Rev. 0, dated: 2020-06-15</p> <p>Notified Body: DEKRA (0124)</p>
<p>07613353026L8 DolorClast Smart 20</p>	Class IIa	N/A	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15</p> <p>Annex Rev. 0, dated: 2020-06-15</p> <p>Notified Body: DEKRA (0124)</p>
<p>07613353046LE DOLOR Handpieces</p>	Class IIa	N/A	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15</p> <p>Annex Rev. 0, dated: 2020-06-15</p> <p>Notified Body: DEKRA (0124)</p>

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<p align="center">07613353037LD PIEZON Instruments</p>	Class IIa	N/A	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15</p> <p>Annex Rev. 0, dated: 2020-06-15</p> <p>Notified Body: DEKRA (0124)</p>
<p align="center">07613353049LL Probes Suction Tubes</p>	Class IIa	N/A	<p>Certificate number: Certificate 50081-16-09; dated: 2020-06-15</p> <p>Annex Rev. 0, dated: 2020-06-15</p> <p>Notified Body: DEKRA (0124)</p>
<p align="center">07613353019LB LithoPump</p>	Class IIa	N/A	<p>Certificate number: 50081-16-09</p> <p>Notified Body: DEKRA (0124)</p>

CERTIFICATE



EN ISO 13485:2016 + AC:2018 + A11:2021

DEKRA Certification GmbH hereby certifies that the organization

E.M.S. Electro Medical Systems S.A.

Scope of certification:

Design, manufacturing, distribution and servicing of medical devices and accessories for dentistry, urology and orthopedics

Certified location:

Chemin de la Vuarpillière 31, 1260 Nyon, Switzerland
(further locations see annex)

has established and maintains a quality management system according to the above mentioned standard.
The conformity was adduced with audit report no. 50081-R3-00.

Certificate registration no.: 50081-21-00

Certificate valid from: 2024-04-23

Validity of previous certificate: 2024-04-22

Certificate valid to: 2026-06-14



K. Leicht

Karin Leicht
DEKRA Certification GmbH, Stuttgart, 2024-04-23



Deutsche
Akkreditierungsstelle
D-ZM-16029-08-00

Annex to the Certificate No. 50081-21-00

valid from 2024-04-23 to 2026-06-14

The following locations/companies belong to the certificate above:

	Headquarters	Certified location	Scope of certification
	E.M.S. Electro Medical Systems S.A.	Chemin de la Vuarpillière 31 1260 Nyon Switzerland	see page 1
	at the following locations/at the companies at the following locations		Scopes of certification
1.		Rte. de Champ-Colin 2 1260 Nyon Switzerland	Manufacturing of medical devices and accessories for dentistry, urology and orthopedics.
2.		Rte. de Champ-Colin 18 1260 Nyon Switzerland	Manufacturing and logistic activities of medical devices and accessories for dentistry, urology and orthopedics.
3.		Rte. de Pontarlier 32 39460 Foncine-le-Haut France	Manufacturing of medical devices and accessories for dentistry, urology and orthopedics



K. Leicht

Karin Leicht

DEKRA Certification GmbH, Stuttgart, 2024-04-23