ActiMaris AG Sandgrube 29 ■ 9050 Appenzell SWITZERLAND

Fon: +41 71 505 75 25 Fax: +41 71 505 75 24

www.actimaris.com info@actimaris.com



# **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/or1
- 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	ActiMaris AG
Manufacturer address and contact details	Sandgrube 29, 9050 Appenzell, SWITZERLAND info@actimaris.com +41 71 505 75 25
Single Registration Number (SRN) (if available)	CH-MF-000035551

Authorised Representative name (if applicable)	Adriatic Health Factory d.o.o.
Authorised Representative address and contact details	Nikole Tesle 17, 44000 Sisak, CROATIA info@ahf.hr +385 44 523 550
Single Registration Number (SRN) (if available)	HR-AR-000029035

Notified body name (if applicable)	SIQ Ljubljana
Notified body number (if applicable)	CE 1304

 $<sup>^{</sup>m 1}$  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.

### ActiMaris AG Sandgrube 29 ■ 9050 Appenzell SWITZERLAND

Fon: +41 71 505 75 25 Fax: +41 71 505 75 24



www.actimaris.com info@actimaris.com

Directive Certificate number(s) to which this confirmation is made (if applicable)	7036 □ See	attached schedu <b>l</b> e
Original expiry date as indicated on the Directive Certificate prior to the extension of the validity (if applicable)	26 May 2024 □ See	e attached schedule
End date of extended validity/transition period	31 December 2028	attached schedu <b>l</b> e

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/or2
- 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 attac	ched schedule	Э
--	-----------	----------------	-----------	-------------	-----------	---------------	---

ective Certificate(s) as listed above of ill 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 <i>before</i> 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
$\square$ 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
$\square$ 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)

Doc-ID: 3934-02-FO\_ Declaration-Letters-General - Release: Date / Visum: 2022-10-28 /

 $<sup>^{2}</sup>$  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

ActiMaris AG Sandgrube 29 9050 Appenzell **SWITZERLAND** 

Fon: +41 71 505 75 25 Fax: +41 71 505 75 24



ActiMaris AG

www.actimaris.com info@actimaris.com

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 subpara-
graph 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.

## ⊠ 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

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:

ActiMaris AG Sandgrube 29 ■ 9050 Appenzell SWITZERLAND

Fon: +41 71 505 75 25 Fax: +41 71 505 75 24



www.actimaris.com info@actimaris.com

 $\square$  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:

ActiMaris AG

Appenzell, 14.5.2023

ActiMaris AG
Sandgrube 29
9050 Appenzell
Switzerland
www.actimaris.com

Dani Marincic

CEO

dani.marincic@actimaris.com +41 71 505 75 25



ActiMaris AG



info@actimaris.com www.actimaris.com

natural medical solutions

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

Schedule of Devices

Identification of the device(s) <sup>3</sup> (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)
ActiMaris Wound Gel 20 g REF: 30220	7063	26 May 2024	SIQ / 1304	SIQ / 1304	31 December 2028	N/A
ActiMaris Wound Gel 50 g REF: 30250	2063	26 May 2024	SIQ / 1304	SIQ / 1304	31 December 2028	N/A
ActiMaris SENSITIVE Wound Irrigation Solution 50 ml	7063	26 May 2024	SIQ / 1304	SIQ / 1304	31 December 2028	N/A
ActiMaris SENSITIVE Wound Irrigation Solution 300 ml	7063	26 May 2024	SIQ / 1304	SIQ / 1304	31 December 2028	N/A
ActiMaris SENSITIVE Wound Irrigation Solution 1000 ml	7063	26 May 2024	SIQ / 1304	SIQ / 1304	31 December 2028	N/A
ActiMaris FORTE Wound Irrigation Solution 50 mI REF: 30355	7063	26 May 2024	SIQ / 1304	SIQ / 1304	31 December 2028	N/A
ActiMaris FORTE Wound Irrigation Solution 300 mI REF: 30350	7063	26 May 2024	SIQ / 1304	SIQ / 1304	31 December 2028	N/A
ActiMaris FORTE Wound Irrigation Solution 1000 ml REF: 31050	7063	26 May 2024	SIQ / 1304	SIQ / 1304	31 December 2028	N/A

<sup>&</sup>lt;sup>3</sup> 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)