

# EU Technical Documentation Assessment Certificate

Certificate no.  
13560GB450240322

Final Assessment Report no.  
13560AU08F

Effective date  
2024-03-22

Expiry date  
2029-03-21

This is to certify that  
**Medical devices listed on the following pages**

Manufactured by

**Luciole Medical AG**

Baslerstr. 30, 8048 Zürich, Switzerland

SRN: CH-MF-000015856

Have been assessed and found to comply with respect to

**Technical Documentation Assessment as described in Annex IX,  
Chapter II of Regulation (EU) 2017/745 on Medical Devices**

Any applicable limitations for certain medical devices are included in the following list or recorded in the final assessment report. This certificate assumes that DNV MEDCERT has to be informed about any changes of the assessed device. Changes need further approval by DNV MEDCERT.

For conditions or for limitations to the validity refer to the relevant final assessment report. Examinations and tests performed, e. g. reference to relevant common specifications, harmonised standards, test reports and audit report(s), are recorded in the relevant reports. For placing on the market of the medical devices covered by this certificate, an additional EU Quality Management System Certificate according to Annex IX Chapter I of Regulation (EU) 2017/745 is required.

Place and date  
Hamburg, 2024-03-22

For the issuing office  
DNV MEDCERT GmbH – Notified Body 0482  
Pilatuspool 2, 20355 Hamburg, Germany



Benannt durch/Designated by  
Zentralstelle der Länder  
für Gesundheitsschutz  
bei Arzneimitteln und  
Medizinprodukten  
www.zlg.de  
BS-MDR-096

The certificate is only valid when provided entirely with all of its pages. To verify the validity of this certificate, contact Medcert-Info@dnv.com

  
Markus Bianchi  
Director Certification Body



Certificate no.: [13560GB450240322](#)  
Place and date: [Hamburg, 2024-03-22](#)

**Authorised representative**

Veranex Germany GmbH, Landsbergerstr. 302, 80687 München, Germany  
SRN: DE-AR-000005578





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### Products covered by this certificate

<b>Class</b>	III
<b>Category</b>	MDN 1201
<b>Basic UDI-DI</b>	7649989882RS2.0UV
<b>Intended purpose</b>	Probe for monitoring brain parameters
<b>Model (REF#)</b>	<b>Medical device name</b>
20020	RheoSens 2.0

