



**CE DECLARATION OF CONFORMITY according to European medical device regulation 2017/745**  
**DECLARATION CE DE CONFORMITE selon le règlement européen 2017/745**

MANUFACTURER FABRICANT	<b>BioSerenity</b> ICM-iPEPS 47, Boulevard de l'Hôpital 75013 Paris France
SRN NUMBER NUMERO SRN	FR-MF-000000497
PRODUCT DESIGNATION DÉSIGNATION DU PRODUIT	Neuronaute® IceCap Neuronaute® CareCap
PRODUCT REFERENCE REFERENCE PRODUIT	1001-01010-EU 1001-01001-UN 1001-01002-UN 1001-01003-UN
Basic UDI-DI NUMBER NUMERO IUD-ID de base	361522WEMUelectrodsV0BC
INTENDED USE INDICATION D'UTILISATION	Neuronaute enables the acquisition, recording, storage, transmission, and display of 21-lead electroencephalogram (EEG) in order to analyse potential neurological disorders. Neuronaute permet l'acquisition, l'enregistrement, le stockage, la transmission et l'affichage d'un électroencéphalogramme (EEG) à 21 dérivations afin d'analyser les troubles neurologiques potentiels.
EMDN CODE CODE EMDN	N01010299 – EEG electrodes - other
CLASSIFICATION	Class I - Rule 1

We hereby declare that the above-mentioned products meet the requirements of the European medical device regulation 2017/745. All supporting documentation is retained at the premises of the manufacturer. This declaration of conformity is issued under the sole responsibility of BioSerenity.

Nous certifions que les produits mentionnés ci-dessus sont conformes au règlement 2017/745/CEE. Les preuves de conformité sont maintenues dans les locaux du fabricant. Cette déclaration de conformité est délivrée sous la seule responsabilité de BioSerenity.

PLACE  
A Paris

DATE OF ISSUE  
DATE January 20<sup>th</sup>, 2022  
20 Janvier 2022

SIGNATURE

NAME / NOM  
POSITION / TITRE Mélanie RENAUD SAMIRI  
Quality and Regulatory Affairs Director  
Directeur Qualité et Affaires Réglementaires



## REFERENCE OF APPLIED REGULATORY STANDARDS REFERENCE DES NORMES RÉGLEMENTAIRES APPLIQUÉES

Standard number Numéro du standard	Standard title Titre du standard
<b>ISO 10993-1:2018</b>	Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk management process (ISO 10993-1:2009)
<b>NF EN ISO 10993-2:</b>	Biological evaluation of medical devices — Part 2 : Animal Welfare Requirements
<b>EN ISO 10993-5:2009</b>	Biological evaluation of medical devices — Part 5: Tests for in vitro cytotoxicity
<b>ISO 10993-10:2010</b>	Biological evaluation of medical devices — Part 10: Tests for irritation and skin sensitization
<b>EN ISO 10993-12:2021</b>	Biological evaluation of medical devices — Part 12: Sample preparation and reference materials
<b>EN ISO 10993-17:2009</b>	Biological evaluation of medical devices — Part 17: Establishment of allowable limits for leachable substances (ISO 10993-17:2002)
<b>EN ISO 10993-18:2020</b>	Biological evaluation of medical devices — Part 18: Chemical characterization of materials
<b>ISO/TS 10993-19:2020</b>	Biological evaluation of medical devices — Part 19: Physico-chemical, morphological and topographical characterization of materials
<b>EN ISO 10993-23:2021</b>	Biological evaluation of medical devices — Part 23: Tests for irritation
<b>EN ISO 13485:2016/AC:2018</b>	Medical devices — Quality management systems — Requirements for regulatory purposes
<b>EN ISO 14971:2019</b>	Medical devices — Application of risk management to medical devices
<b>EN ISO 15223-1:2021</b>	Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied — Part 1: General requirements
<b>ISO 20417 :2021</b>	Medical devices — Information to be supplied by the manufacturer
<b>ISO/TR 24971:2020</b>	Medical devices — Guidance on the application of ISO 14971
<b>EN 60601-1:2006 + A1:2013/AC:2014</b>	Medical electrical equipment - Part 1-1: General requirements for safety - Collateral standard: Safety requirements for medical electrical systems
<b>IEC 60601-1-2:2014+AMD1:2020</b>	Medical electrical equipment — Part 1-2: General requirements for basic safety and essential performance — Collateral Standard: Electromagnetic disturbances — Requirements and tests IEC 60601-1-2:2014
<b>IEC 60601-1-6:2010 +AMD1:2013 +AMD2:2020</b>	Medical electrical equipment - Part 1-6: General requirements for basic safety and essential performance - Collateral standard: Usability
<b>EN 60601-1-11:2015/AMD 1:2020</b>	Medical electrical equipment - Part 1-11: General requirements for basic safety and essential performance - Collateral standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment
<b>EN IEC 80601-2-26:2020</b>	Medical electrical equipment - Part 2-26: Particular requirements for the safety of electroencephalographs
<b>IEC 80601-2-49:2020</b>	Medical electrical equipment - Part 2-49: Particular requirements for the safety of multifunction patient monitoring equipment
<b>EN 62366-1:2015+A1:2020</b>	Medical devices - Application of usability engineering to medical devices
<b>EN 62366-2:2016</b>	Guidance on the application of usability engineering to medical devices



Applied Regulation Règlement appliqué	Regulation title Titre du règlement
REG (UE) 2017/745	REGULATION (EU) 2017/745 OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL of 5 April 2017 on medical devices, amending Directive 2001/83/EC, Regulation (EC) No 178/2002 and Regulation (EC) No 1223/2009 and repealing Council Directives 90/385/EEC and 93/42/EEC
Rect REG (UE) 2017/745	Corrigendum to Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices, amending Directive 2001/83/EC, Regulation (EC) No 178/2002 and Regulation (EC) No 1223/2009 and repealing Council Directives 90/385/EEC and 93/42/EEC
Rect REG (UE) 2017/745 (2)	Corrigendum to Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices, amending Directive 2001/83/EC, Regulation (EC) No 178/2002 and Regulation (EC) No 1223/2009 and repealing Council Directives 90/385/EEC and 93/42/EEC
REG (UE) 2020/561	REGULATION (EU) 2020/561 OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL of 23 April 2020 amending Regulation (EU) 2017/745 on medical devices, as regards the dates of application of certain of its provisions
REG (UE) 1907/2006/CE	C1 REGULATION (EC) No 1907/2006 OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL of 18 December 2006 concerning the Registration, Evaluation, Authorisation and Restriction of Chemicals (REACH), establishing a European Chemicals Agency, amending Directive 1999/45/EC and repealing Council Regulation (EEC) No 793/93 and Commission Regulation (EC) No 1488/94 as well as Council Directive 76/769/EEC and Commission Directives 91/155/EEC, 93/67/EEC, 93/105/EC and 2000/21/EC
DÉCISION (UE) 2020/437	COMMISSION IMPLEMENTING DECISION (EU) 2020/437 of 24 March 2020 on the harmonised standards for medical devices drafted in support of Council Directive 93/42/EEC
DECISION (EU) 2021/610	COMMISSION IMPLEMENTING DECISION (EU) 2021/610 of 14 April 2021 amending Implementing Decision (EU) 2020/437 as regards harmonised standards on medical vehicles and their equipment, anaesthetic and respiratory equipment, biological evaluation of medical devices, packaging for terminally sterilised medical devices, sterilisation of health care products, clinical investigation of medical devices for human subjects, non-active surgical implants, medical devices utilising animal tissues and their derivatives, electroacoustics and medical electrical equipment
DECISION (EU) 2021/1182	COMMISSION IMPLEMENTING DECISION (EU) 2021/1182 of 16 July 2021 on the harmonised standards for medical devices drafted in support of Regulation (EU) 2017/745 of the European Parliament and of the Council
REC 2013/172/UE	COMMISSION RECOMMENDATION of 5 April 2013 on a common framework for a unique device identification system of medical devices in the Union
REG (EU) 2012/19/EU	WEEE - DIRECTIVE 2012/19/EU OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL of 4 July 2012 on waste electrical and electronic equipment (WEEE)
REG (EU) 2011/65/EU	DIRECTIVE 2011/65/EU OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment
REG (EU) 2012/207	COMMISSION REGULATION (EU) No 207/2012 of 9 March 2012 on electronic instructions for use of medical devices



**REFERENCE OF APPLIED COMMON SPECIFICATIONS**  
**REFERENCE DES SPECIFICATIONS COMMUNES APPLIQUÉES**

**No common specification applicable.**

**Pas de spécifications communes appliquées.**

**List of accessories included in the Neuronaute IceCap and Neuronaute CareCap**  
**Liste des accessoires inclus dans le Neuronaute IceCap et Neuronaute CareCap**

<b>Accessories</b> <b>Accessoires</b>	<b>Reference</b> <b>Référence</b>
IceAdapter	1001-15012-EU / 1001-15020-UN
IceBox PCB	1001-15019-EU
eEEG Comfort + - Foam Electrodes	1013-06001-EU
Care ECG Cable	1001-15011-EU
Care REF/EOG	1001-15010-EU
DB25 Cable	3000-0001