

## EC Declaration of Conformity

To whom it may concern,

Samsø, Denmark, May 2024

We,

**Exam Vision ApS, Industrivej 11, Tranebjerg, 8305 Samsø, Denmark**

hereby declare in our capacity as the product manufacturer, that the Light System Control Unit, Xtend, which is intended to be used

*as a visual aid, illuminating the visual field for the Healthcare Professional while performing medical procedures,*

complies with the requirements of the

*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 with amendments.*

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|--|---|
| <b>Actor ID/SRN:</b>   | DK-MF-000032397   |
| <b>Basic UDI-DI:</b>   | 574400023EVLight5F  |
| <b>Product Name/Trade name:</b>  | Xtend   |
| <b>Intended Purpose:</b>   | The Light System is intended to be used as a visual aid, illuminating the visual field for the Healthcare Professional while performing medical procedures.   |
| <b>Risk Class (MDR):</b>   | Class I, rule I.  |
| <b>Conformity Assessment Route:</b>  | Annex I General Safety and Performance Requirements<br>Annex II Technical Documentation<br>Annex III Technical Documentation on Post-Market Surveillance      |
| <b>Other relevant legislation, applicable harmonised standards, and normative documents:</b> | RoHS & REACH<br>IEC 60601-1:2005 + A1:2012 + A2:2020<br>IEC 60601-1-2:2014 + A1:2020<br>IEC 60601-1-6 + A1:2+13 + A2:2020<br>IEC 62471:2006<br>ISO 14971:2019 |

Issue date: 1 May 2024



Signature:



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**Ole Anker Aagaard/Head of Legal**