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**EC Declaration of Conformity**

**No.: REG-003791**

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**Product Name**    Ambu® aView™ 2 Advance

**REF**                405011000

**Identification  
of the Device**    Category:            Medical Device Accessory  
                          Type:                Displaying Unit  
                          Software ver.:    v1.0.0

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**Identification:**    All products manufactured after issue date.

**We, the manufacturer, hereby declare that the device(s) covered by the present declaration is in conformity with the requirements specified in the relevant Union legislation:**

**Council directive 93/42/EEC**, Annex I and Annex VII enforced in Danish law.

Device classification: Class I, non-sterile according to Annex IX, rule 12

**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 (RoHS recast) as is specified in Article 4.

RoHS category of device: Category 8 – Medical devices

## Radio Equipment Directive 2014/53/EU

Article 3.1(a) Health & Safety	EN 62311: 2008
Article 3.1(b) EMC	EN 301 489-1 v2.1.1 EN 301 489-17 v3.1.1 EN 55032:2015 + AC:2016, Class A EN 61000-3-2:2014 EN 61000-3-3:2013 EN 61000-4-2:2009 EN 61000-4-3:2006 + A1:2008 + A2:2010 EN 61000-4-4:2012 EN 61000-4-5:2014 + A1:2017 EN 61000-4-6:2014 + AC:2015 EN 61000-4-11:2004 + A1:2017
Article 3.2 Spectrum	EN 300 328 v2.1.1 EN 301 893 v2.1.1

**Signed for and on behalf of Ambu A/S, Denmark**

01 May 2020



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Kristine Rasmussen, Head of Regulatory Affairs Displaying Units

**First Issued: 01 May 2020**

**EC Declaration of Conformity – Annex I: GMDN Code**

**No.: REG-003791**

**Product Name: Ambu® aView™ 2 Advance**

The Ambu® aView™ 2 Advance are covered by the following GMDN Code:

**GMDN Code:**56654

**Term:** Endoscopic video image display monitor

