

# EU DECLARATION OF CONFORMITY

Manufacturer: **WSAUD A/S**  
Nymoellevej 6  
DK-3540 Lyngø  
Denmark

Brand: **WIDEX**

Product Family: **WIDEX CROS DEVICES**

Type of Device: **CROS audio streaming device**

Basic UDI-DI: **5714880-WSA-62-10-3S**

GMDN Code: **59460 Contralateral hearing unit**

Product Identification: **See below**

We declare under our sole responsibility that above products are in conformity with the following Regulations and Directives:

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**REGULATION (EU) 2017/745 (MDR) OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL**

Conformity assessment procedure: Annex IX of Regulation (EU) 2017/745

Classification of device: **Class I** (according to Annex VIII Rule 13 to Regulation (EU) 2017/745)

The products meet all applicable standards and the general safety and performance requirements of the Regulation (EU) 2017/745 Annex I. Applicable standards are listed in the respective technical documentation.

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**Council Directive 2011/65/EU (RoHS) as amended by Dir. 2017/2102/EC (RoHS2)**

Relevant Harmonized Standards: EN50531, EN62321

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**Council Directive 2014/53/EU (RED)**

Relevant Harmonized Standards: EN 62479, EN 301 489-1, EN 301 489-3, EN 300 330

Standard versions valid on the date when this DoC is issued.

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Product Identification	Type of Device
CROS-FA	CROS audio streaming device

This Declaration of Conformity includes all hearing aid components and spare parts of the products listed above.

Place and valid from date: Lyngø, October 05, 2021

Name: Hans-Otto Bindeballe  
Global Regulatory Affairs Manager

Signature: 

This declaration will be renewed on any significant change of product, product range, standards and laws.