

## EC – Declaration of Conformity

**Manufacturer:** GN Hearing A/S  
Lautrupbjerg 7  
DK-2750 Ballerup  
Denmark

**Conformity Assessment Procedure:** Annex VII of Medical Device Directive  
(MDD) 93/42/EEC  
Annex III of Radio and  
Telecommunications Terminal Equipment  
(R&TTE) 1999/5/EC

**Identification of the Device:** **Category:** Accessory  
**Type:** Assistive Listening Device  
**Brand:** GN Hearing  
**Model:** Multi Mic  
SM-2P

**GMDN code:** 57886

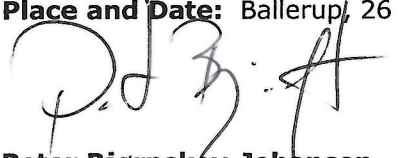
**Revision:** Report No. 296

**Classification of the Device (MDD):** Class I, Rules 1 and 12, MDD 93/42/EEC

**Applied standards and normative standards:**  
MDD: ISO 10993-5:2009, ISO 10993-10:2010, ISO 14971:2012  
R&TTE: EN/(IEC) 60950-1:2006, EN/(IEC) 62479:2010 (Health & Safety)  
EN 301 489-17 2.2.1 (EMC), EN 300-328 V1.9.1 (Spectrum)

***We, the manufacturer hereby declare that the above-mentioned devices comply with the relevant provisions of the EU Council Directive 93/42/EEC (MDD) Annex I - Essential Requirements and the essential requirements and other relevant requirements of the R&TTE Directive 1999/5/EC and their relevant transpositions into national laws of the Member States in which the above mentioned medical devices are distributed.***

**Place and Date:** Ballerup, 26 February 2016

  
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