

EC Declaration of Conformity

(Following the provisions of the R&TTE Directive 1999/5/EC)

We,

**Widex A/S
Nymoellevej 6
DK-3540 Lyngø
Denmark**

Declare under our sole responsibility that the following **EQUIPMENT**

| | |
|-------------|--|
| Model(s) | C4-CIC, C3-CIC, C2-CIC, C4-CIC-TR, C3-CIC-TR, C2-CIC-TR |
| Description | Hearing aid |


Is manufactured in accordance with the requirements of 1999/5/EC Annex III and that the product is in conformity with the essential requirements of the following **DIRECTIVE**

Council Directive 1999/5/EC (R&TTE)

The product conforms to the following **STANDARDS AND SPECIFICATIONS** applying versions valid on the date when this DoC is issued

| | |
|--------------|---|
| EN 60950-1 | : Safety of Information Technology Equipment |
| EN 60601-1-2 | : Electromagnetic Compatibility, Medical electrical equipment |
| EN 301 489-1 | : Electromagnetic Compatibility. Common technical requirements |
| EN 301 489-3 | : Electromagnetic Compatibility, Short range devices 9 kHz – 40 GHz |
| EN 300 330-2 | : EMC and Radio Spectrum Matters 9kHz – 30 MHz |
| EN 50371 | : Human exposure standard for electromagnetic fields 10 MHz – 300 GHz |

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Place of issue: Lyngø



Jan Tøpholm
CEO