

# EU Declaration of Conformity

Manufacturer:

Nitzbon AG

SRN: DE-MF-000009971

The manufacturer declares under its sole responsibility that the medical device:

Product name:

WP 5 Hole Grip World & Grip Set, Item no. 1623610

Intended purpose:

Occupational therapy, work simulation training, matching job profiles with skill profiles, and conducting occupational therapy and labour market-related performance analyses.

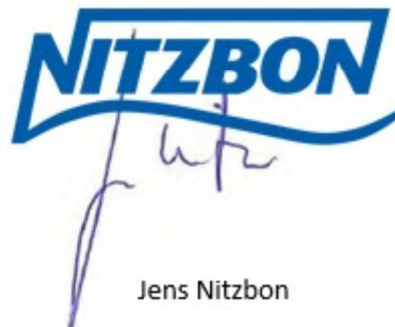
Basic UDI-DI: 4260101481623610VX

is in conformity with Regulation (EU) 2017/745 on medical devices.

The device is classified as Class I in accordance with Annex VIII, Rule 1 of Regulation (EU) 2017/745.

This declaration is valid until 31 December 2027 or until significant changes require revision of this declaration.

Hamburg, 1 January 2026  
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The logo consists of the word "NITZBON" in a bold, blue, sans-serif font, enclosed within a blue rectangular border that has a slight wave at the bottom. Below the logo is a handwritten signature in blue ink, which appears to be "Jens Nitzbon".

Jens Nitzbon

## Annex to the Declaration of Conformity

Supplement for legacy products pursuant to Regulation (EU) 2017/745 (MDR, Art. 120)

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### 1. Scope

This annex supplements the existing Declaration of Conformity dated January 1, 2026, for the following products that were placed on the market in accordance with Directive 93/42/EEC (MDD) and are subject to the transitional provisions of the MDR.

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### 2. Affected products

| Product Name                    | UDI-DI        | Basis UDI-DI       | Valid until |
|---------------------------------|---------------|--------------------|-------------|
| WP 5 Hole Grip World & Grip Set | 4260101482044 | 4260101481623610VX | 12/31/2027  |

*Note: The product is a Class I device and has complied with the UDI requirements of the MDR since July 2021.*

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### 3. Legal Basis

- The products were placed on the market in accordance with MDD 93/42/EEC.
  - They are subject to the transitional provisions of the MDR 2017/745 (Art. 120) and may continue to be used until the specified date.
  - The UDI-DI and Basic UDI-DI comply with the requirements of the MDR (Annex VI).
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### 4. Declaration

The manufacturer confirms that the above-mentioned products comply with the transitional provisions of the MDR and that the UDI requirements as well as post-market surveillance, vigilance, and market surveillance for legacy devices are met.

Hamburg, April 8, 2026

Jens Nitzbon PRRC

