



# EU DECLARATION OF CONFORMITY

**Manufacturer** HoverTech International  
4482 Innovation Way  
Allentown, PA 18109

**SRN** US-MF-000008435  

EC	REP
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**Authorized Representative Name** CEpartner4U

**Authorized Representative Address** Esdoornlaan 13, 3951DB Maarn, The Netherlands

**Statement** This EU declaration of conformity is issued under the sole responsibility of the manufacturer.  
The device(s) covered by present declaration is/are in conformity with EU Regulation 2017/745 on medical devices.

**Basic UDI-DI** 081629901HMSM

**Intended purpose** The HoverMatt® Air Transfer System is used to assist caregivers with patient transfers, positioning (including boosting and turning), and proning.

**Product / device name** HoverMatt  
HoverMatt SPU  
HoverSling  
HoverMatt PROS

**Risk class of the device** Class 1, rule 1

**Place** Allentown, PA USA

**Date of issue** January 30, 2025

**Name and function** Susan Pavelko /Quality Manager

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Signature, on behalf of HoverTech International