



260320/ha-RS-1

## EU DECLARATION OF CONFORMITY

**Manufacturer:** Oy Klippan Ab  
Santaradantie 8  
FI-01370 Vantaa  
SRN: FI-MF-000052644

**Manufactured for:** High Adventure Oy  
Pulttitie 14  
00880 HELSINKI

**Basic UDI-DI:** 641764002816AX

**Name of Device:** HARRT Pro RS – Restraint Strap  
**Product code:** HA-RS

**Intended Use:** For use in conjunction with basket stretchers

**Classification:** Class 1, according to annex VIII, of MDR 2017/745

### Conformity Claim

The device/accessory covered by this declaration is in conformity with Regulation (EU) 2017/745.

- EN ISO 14971:2019 + A11:2021
- EN ISO 15223-1

We hereby declare that this EU Declaration of Conformity is issued under the sole responsibility of the manufacturer, and the medical device specified above meet the provision of the Regulation (EU) MDR 2017/745 for medical devices.

Oy Klippan Ab or representative will make available upon request all applicable technical documentation to allow assessment of conformity of its products

Vantaa 20.3.2026

Oy Klippan Ab

Hans Bäckström  
CEO



### OY KLIPPAN AB

Santaradantie 8  
FI-01370 VANTAA

tel. +358 (0) 9 836 243 0  
e-mail: [info@klippan.fi](mailto:info@klippan.fi)  
[www.klippan.fi](http://www.klippan.fi)

Y-tunnus: 0113210-1  
VAT No: FI01132101

CERTIFIED  
ISO 9001  
ISO 14001

