

## EU DECLARATION OF CONFORMITY

DOC NAME					PAGES	
EU Declaration of Conformity MDR - Navina Smart app for iOS/Android					1(3)	
DOC TYPE	DOC NO	VERSION	ENCLOSURE	DATE	STATE	
DC	10105	E	None	2023-12-19	Approved	
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We,

Wellspect HealthCare  
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Sweden

being the Manufacturer, registered in the European Union under Single Registration Number (SRN) No. SE-MF-000026271, of the Navina Mobile Application (App), including the products listed in the Annex I to this document, with the following characteristics:

- device class I, as determined by Rule 11, according to Regulation (EU) 2017/745, Annex VIII
- intended for (optional) use with Navina Smart System 'Transanal irrigation' device
- GMDN code: 61114
- EMDN category G/ code(s):
  - o G99 Gastrointestinal devices - other
- Basic UDIDI/Global Model Number: 733338724103CZ

Declare under our sole responsibility that the product(s) conform to the requirements of: Regulation (EU) 2017/745 of the European Parliament and of the Council, on medical devices, and meet(s) the relevant General Safety and Performance Requirements of Annex I.

All devices are designed, manufactured, tested, and released for sale in accordance with the technical documentation according to Annex II and III of regulation (EU) 2017/745 and the applicable standards.

This declaration of Conformity is approved and signed as dated on first page.

Wellspect HealthCare, Mölndal, Sweden

  
TONI JØRGENSEN  
VICE PRESIDENT QUALITY ASSURANCE & REGULATORY AFFAIRS

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DC	10105	E	2023-12-19	2 (3)

**ANNEX I**

<b>Article (model) No.</b>	<b>Product Name, Description</b>
75750	Navina Smart app for iOS
75751	Navina Smart app for Android

DOC TYPE	DOC NO	VERSION	DATE	PAGES
DC	10105	E	2023-12-19	3 (3)

## REVISION HISTORY

Document Version	Change note/Description
E	New Basic UDI-DI assigned due to an error in previous Basic UDI-DI identifier.