

MANUFACTURER'S DECLARATION: MEDICAL DEVICES REGULATION 2017/745

As this component is not a complete "Medical Device" under the definition provided in the Regulation but a part with no function on its own, a Declaration of Conformity or CE mark is not appropriate until the component is incorporated. The following Declaration is issued in order to aid the installation of the component below into a finished appliance, for use by, or sale to, an enduser.

The following declares the status of these products with regard to Union Regulation 2017/745:

Product Description	Product Name	Part Numbers
VR2 Dual Attendant Module	VR2 Dual Module	D50872

The object of the declaration described above is in conformity with the relevant Union harmonisation legislation to the extent possible as a component to be integrated into a complete product.

APPLICABLE STANDARDS

The following standards (and those called by them) have been used in order to assess a presumption of conformity with the essential requirements of the above regulation as far as the component allows:

Medical devices — Application of risk management to medical devices
Medical device software — Software life-cycle processes
Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process
Electrically powered wheelchairs, scooters and their chargers. Requirements and test methods
Wheelchairs – Part 9: Climatic tests for electric wheelchairs
Wheelchairs Part 14: Power and control systems for electrically powered wheelchairs and scooters Requirements and test methods
Wheelchairs Part 21: Requirements and test methods for electromagnetic compatibility of electrically powered wheelchairs and scooters, and battery chargers

Nigsl D Mills 14 June 2023

Nigel Mills, Senior Manager, Engineering

Signed at, for and on behalf of:

Penny & Giles Controls Ltd.,

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