

EU Quality Management System Certificate

Certificate no.:
C538821

Initial certification date:
27 January 2025

Valid Until:
27 January 2030

This is to certify that the quality system of

Charder Electronic Co., Ltd.

No. 103, Guozhong Rd., Dali Dist., Taichung City 41262, Taiwan

SRN: TW-MF-000013673

For design, production, and final product inspection/testing of:

Scale, Body Composition Analyzer, Height Measurement

Has been assessed and found to comply with respect to:

The conformity assessment procedure described in Annex IX, (Chapter I & III) of Regulation (EU) 2017/745 on Medical Devices

Place and date:
Høvik, 27 January 2025

For the issuing office:
DNV Product Assurance AS – Notified Body 2460
Veritasveien 1, 1363 Høvik, Norway



Sholeh Gheissar

Sholeh Gheissar
Management Representative

Jurisdiction

Application of Regulation 2017/745 on medical devices, adopted as “Forordning (EU) 2017/745 om medisinsk utstyr (MDR)” by the Norwegian Ministry of Health and Care Services.

Certificate history:

Revision	Description	Report No.	Issue Date
0.0	Original Certificate	2694021	27 January 2025

Products covered by this Certificate:

Product Description	Product Name	Class*
Body Composition Analyzer	MBF6000, MBF6010, MA601, MA801	IIa
Bed Scale	MS6000, MS6001, MS7800, M-999, MS6080	Im
Chair Scale	MS5410, MS5440, MS5460, MS5461, MS5810, MS5811, M-200, M-230, M-250, MS5470, MS5480, MS5880	
Infant Scale	MS21NEOV, MS2400, MS3500, MS4200, MS4400I, MS5900, M-400, M-400BT, M-410, M-410BT, MS5980	
Lift Scale	MHS2500I, MHS2510I, MHS2600I, MHS2610I, M-600, M-605, MHS2700, MHS2710	
Stand-on Floor Scale	MS2504, MS3200, MS3400-1, MS3450, MS3910, MS4202L, MS4640, MS4900, MS4910, MS4970, MS4971, MS5750, MS5751, MS6110, MS6111, M-110, M-125, M-420, M-420BT, M-430, M-430BT, M-510, M-545, M-550, MS2580, MS3980, MS4980, MS4203	
Wheel Chair Scale	MS2350, MS3830, M-610, M-650, M-615, MS3880	
Height Measurement	HM80M, HM80P, HM101M, HM110M, HM200PW, HM200P, HM201M, HM202P, HM230M, HM80D, HM80DT, HM200D, HM201D, HM100D, HM130D, HM210D, HM200U, HM250U	Im

* Class III and class IIb devices referred to in the second subparagraph of Article 52(4): Technical documentation assessment is covered by a separate EU Technical Documentation Assessment Certificate No.: NA

The complete list of devices is filed with the Notified Body

Sites covered by this certificate

Site Name	Address
Charder Electronic Co., Ltd.	No. 103, Guozhong Rd., Dali Dist., Taichung City 41262, Taiwan

EU Representative

Obelis s.a., Boulevard Général Wahis, 53 1030 Brussels, Belgium SRN: BE-AR-000000106

Terms and conditions

The certificate is subject to the following terms and conditions:

- Any producer (see 2001/95/EC for a precise definition) is liable for damage caused by a defect in his product(s), in accordance with directive 85/374/EEC, as amended, concerning liability of defective products.
- The certificate is only valid for the products and/or manufacturing premises listed above.
- The Manufacturer shall fulfil the obligations arising out of the quality system as approved and uphold it so that it remains adequate and efficient.
- The Manufacturer shall inform the Notified Body of any intended updating of the quality system and the Notified Body will assess the changes and decide if the certificate remains valid.
- Periodical audits will be held, in order to verify that the Manufacturer maintains and applies the quality system. Notified Body reserves the right, on a spot basis or based on suspicion, to pay unannounced visits.
- For the class III devices and IIb devices falling under Article 52 (4) covered this certificate is dependent on the continued validity of the EU Technical Documentation Assessment Certificate, covering the devices.

The following may render this Certificate invalid:

- Changes in the quality system affecting production.
- Periodical audits not held within the allowed time window

Specific conditions - Class I devices, Systems and Procedure Packs:

- For class I device being placed on the market in a sterile condition, Class I devices with a measurement function and class I devices being reusable surgical instruments covered by this certificate the audit by the notified body of the quality management system was limited to the aspects required under article 52(7) of the regulation.
- For system and procedure packs being placed on the market in a sterile condition, covered by this certificate the audit by the notified body of the quality management system was limited to the aspects required under article 22(3) of the regulation.
- For Custom Made Class III implantable device the certification only relates to the Quality management system. Technical documentation assessment and issuance of EU Technical Documentation Assessment Certificate does not apply.