

# EU Quality Management System Certificate

Regulation (EU) 2017/745, Annex IX Chapter I and III

## MDR 784266 R000

**Manufacturer:** Cardinal Health 200, LLC

**Address:**

3651 Birchwood Drive  
Waukegan  
Illinois  
60085  
USA

**Single Registration Number:** US-MF-000006765

**EU Authorised Representative:** Cardinal Health Ireland Manufacturing Limited

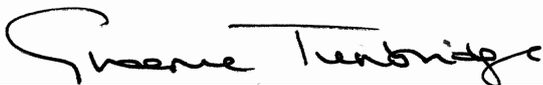
**Address:**

Tullamore Business & Technology Park  
Tullamore  
County Offaly  
R35 H903  
Ireland

**Scope:** See attached **Device Schedule**

On the basis of our examination of the quality system in accordance with Regulation (EU) 2017/745, Annex IX Chapter I and III, the quality system meets the requirements of the Regulation. For the placing on the market of Class III devices, and Class IIb implantable devices that are not considered well-established technologies as specified in Article 52(4) an additional Annex IX Chapter II certificate is required.

For and on behalf of BSI, a Notified Body for the above Regulation (Notified Body Number 2797):



Graeme Tunbridge, Senior Vice President Global Regulatory & Quality

First Issue Date: **2024-02-27**

Current Issue Date: **2025-02-11**

Starting Validity Date: **2025-02-11**

Expiry Date: **2029-02-26**

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### Device Schedule: Class III and Class IIb devices

Class IIb	Intended purpose
Sterile polyurethane dressing	Intended to act as a primary or secondary layer in wound dressings, to serve as a barrier between the skin and/or wound and the environment, and to absorb exudate.
Sterile nasogastric intestinal tubes	Intended for use in an established gastrostomy tract as a replacement tube for the administration of nutrition, fluids, and medications to the stomach of a patient that is physically unable to manage nutritional intake through normal mastication and deglutition.
Sterile alginate dressing	Intended for use to absorb exudates and protect the wound from contamination.

### Device Schedule: Class IIa, Custom-made and other devices

Device(s)	Risk Classification
Enteral feeding pump	Class IIa
Sterile and non-sterile enteral feeding pump tubing sets	Class IIa
Sterile and non-sterile gastrointestinal feeding/aspiration tubes	Class IIa
Sterile and non-sterile cannulas and tips for aspiration and irrigation	Class IIa
Sterile powder free surgical gloves	Class IIa
Sterile Hypodermic syringe needles, with safety systems	Class IIa
Sterile Surgical Drainage Systems	Class IIa
Pneumatic compression equipment	Class IIa
Digital thermometers	Class IIa

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### Certificate History

*(References to applicable Common Specifications, Harmonized Standards complied with, and the relevant test and audit reports that support any of the below certificate changes may be requested from Certificate.Verification@bsigroup.com)*

Date	Reference Number	Action
2024-02-27	3833425	Issued
2024-05-07	30122257	Amended - Change to the intended purpose of the Sterile Polyurethane dressing to align with TD. Specification on sterility status to align with device schedule. Supplemented - Addition of sterile nasogastric tubes devices to the MDR device schedule.
2024-11-19	30204608	Amended - Change to the Gastrointestinal feeding/aspiration tubes device group name to make it more general. Supplemented - Addition of Cannulas and tips for aspiration and irrigation, powder free surgical gloves, Nasogastric intestinal tubes, Hypodermic syringe needles with safety systems devices and Surgical drainage systems to the MDR device schedule.
Current	30312203	Supplemented - Addition of Alginate dressing, Pneumatic compression equipment and Digital thermometers to the MDR device schedule.

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