Changes to legislation: There are currently no known outstanding effects for the The Medical Devices (Northern Ireland Protocol) Regulations 2021. (See end of Document for details)

STATUTORY INSTRUMENTS

2021 No. 905

MEDICAL DEVICES CONSUMER PROTECTION

The Medical Devices (Northern Ireland Protocol) Regulations 2021

Made - - - - 26th July 2021
Coming into force 27th July 2021

THE MEDICAL DEVICES (NORTHERN IRELAND PROTOCOL) REGULATIONS 2021

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Preliminary

- 1. Citation and commencement
- 2. Extent and application
- 3. Interpretation
- 4. Scope

PART 2

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- 5. Reprocessing of single-use devices
- 6. Requirement on health institutions to provide information relating to implanted devices
- 7. Registration of custom-made devices
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PART 3

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- 11. Ethical review of clinical investigations
- 12. Prior authorisation of clinical investigations by the Secretary of State

Status: Point in time view as at 27/07/2021.

Changes to legislation: There are currently no known outstanding effects for the The Medical Devices (Northern Ireland Protocol) Regulations 2021. (See end of Document for details)

- 13. Arbitration following the refusal of a clinical investigation application
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- 18. Notified bodies
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- 20. Language requirements
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- 23. Offence of breaching certain provisions
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- 27. Investigatory powers under the Consumer Rights Act 2015
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- 29. Amendments to the Medical Devices Regulations 2002
- 30. Amendment of regulation 2 (interpretation)
- 31. Amendment of regulation 2A (medical devices which are qualifying Northern Ireland goods)
- 32. Revocation and transitional provision
- 33. Amendment of regulation 10A (UK(NI) indication: general medical devices)
- 34. Amendment of regulation 19 (registration of persons placing general medical devices on the market)
- 35. Amendment of regulation 21B (registration of persons placing active implantable medical devices on the market)
- 36. Amendment of regulation 24A (UK(NI) indication: active implantable medical devices)
- 37. Amendment of regulation 36A (UK(NI) indication: in vitro diagnostic medical devices)

Status: Point in time view as at 27/07/2021.

Changes to legislation: There are currently no known outstanding effects for the The Medical Devices (Northern Ireland Protocol) Regulations 2021. (See end of Document for details)

PART 9

Amendments of other secondary legislation

- 38. Amendment of the Blood Safety and Quality Regulations 2005
- 39. Amendment of the Human Tissue (Quality and Safety for Human Application) Regulations 2007
- 40. Amendment of the Legislative and Regulatory Reform (Regulatory Functions) Order 2007
- 41. Amendment of the Medicines (Products for Human Use) (Fees) Regulations 2016
- 42. Amendment of the Economic Growth (Regulatory Functions) Order 2017
- 43. Amendment of the Market Surveillance (Northern Ireland) Regulations 2021
 Signature

SCHEDULE 1 — Fees for clinical investigations

SCHEDULE 2 — Fees in connection with the designation of notified bodies

SCHEDULE 3 — Provisions breach of which is an offence under regulation 23

Explanatory Note

Status:

Point in time view as at 27/07/2021.

Changes to legislation:

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