#### STATUTORY INSTRUMENTS

# 2021 No. 905

# The Medical Devices (Northern Ireland Protocol) Regulations 2021

#### PART 7

# Amendment of primary legislation

### **Investigatory powers under the Consumer Rights Act 2015**

- **27.**—(1) Schedule 5 to the Consumer Rights Act 2015(1)(investigatory powers etc) is amended in accordance with this regulation.
- (2) In paragraph 8 (interpretation of Schedule 5), after the definition of "online interface order" insert—
  - ""Regulation (EU) 2017/745 on medical devices" means Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices, amending Directive 2001/83/EC, Regulation (EC) No 178/2002 and Regulation (EC) No 1223/2009 and repealing Council Directives 90/385/EEC and 93/42/EEC."
- (3) In paragraph 10 (enforcer's legislation: duties and powers mentioned in paragraph 9(1)(a)) at the end insert "regulation 26 of the Medical Devices (Northern Ireland Protocol) Regulations 2021".
- (4) In paragraph 19 (exercise of powers in Part 4), in sub-paragraph (7A)(a)(2), for the words from "a breach of" to the end substitute—

#### "a breach of-

- (i) the Medical Devices Regulations 2002 (S.I. 2002/618),
- (ii) regulations made under section 15(1) of the Medicines and Medical Devices Act 2021,
- (iii) the Medical Devices (Northern Ireland Protocol) Regulations 2021, or
- (iv) Regulation (EU) 2017/745 on medical devices, and".
  - (5) In paragraph 30A(3) (power to decommission or switch off fixed medical devices)—
    - (a) in sub-paragraph (1), for the words from "pursuant to" to the end substitute— "pursuant to—
      - (a) the duty in regulation 61(1A) or (1B) of the Medical Devices Regulations 2002 (S.I. 2002/618),
      - (b) a duty in regulations made under section 15(1) of the Medicines and Medical Devices Act 2021, or
      - (c) the duty in regulation 26 of the Medical Devices (Northern Ireland Protocol) Regulations 2021.";

<sup>(1) 2015</sup> c.15.

<sup>(2)</sup> Sub-paragraph (7A) was inserted by section 41(2)(c) of the Medicines and Medical Devices Act 2021 (c.3).

<sup>(3)</sup> Paragraph 30A was inserted by section 41(2)(d) of the Medicines and Medical Devices Act 2021 (c.3).

- (b) in sub-paragraph (2), for "medical device to which the Medical Devices Regulations 2002 apply" substitute "relevant medical device";
- (c) after sub-paragraph (2), insert—
  - "(3) In sub-paragraph (2), "relevant medical device" means—
    - (a) where a domestic enforcer is acting pursuant to a duty mentioned in subparagraph (1)(a) or (b), any medical device to which the Medical Devices Regulations 2002 apply;
    - (b) where a domestic enforcer is acting pursuant to the duty mentioned in subparagraph (1)(c), any medical device to which Regulation (EU) 2017/745 on medical devices applies."

## **Commencement Information**

II Reg. 27 in force at 27.7.2021, see reg. 1(2)

Changes to legislation:
There are currently no known outstanding effects for the The Medical Devices (Northern Ireland Protocol) Regulations 2021, Section 27.