

## STATUTORY INSTRUMENTS

# 2021 No. 905

## The Medical Devices (Northern Ireland Protocol) Regulations 2021

### PART 8

#### Amendment of the Medical Devices Regulations 2002

##### Amendments to the Medical Devices Regulations 2002

**29.** The Medical Devices Regulations 2002(1) are amended in accordance with this Part.

##### Commencement Information

**11** Reg. 29 in force at 27.7.2021, see [reg. 1\(2\)](#)

##### Amendment of regulation 2 (interpretation)

**30.** In regulation 2, in paragraph (1)—

(a) for the definition of “medical device” substitute—

““medical device” has the meaning given in Article 2(1) of Regulation (EU) 2017/745 and includes devices intended to administer a medicinal product or which incorporate as an integral part a substance which, if used separately, would be a medicinal product and which is liable to act upon the body with action ancillary to that of the device;”;

(b) after the definition of “Regulation 722/2012” insert—

““Regulation (EU) 2017/745” 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](#)”.

##### Commencement Information

**12** Reg. 30 in force at 27.7.2021, see [reg. 1\(2\)](#)

##### Amendment of regulation 2A (medical devices which are qualifying Northern Ireland goods)

**31.** In regulation 2A—

(a) in paragraph (1)(a), after “Northern Ireland” insert “or of Regulation (EU) 2017/745”;

(1) [S.I. 2002/618](#), as amended by [S.I. 2003/1400](#), [2003/1697](#), [2005/2759](#), [2909](#), [2007/400](#), [803](#), [2008/2936](#), [2009/383](#), [2010/557](#), [2012/1426](#), [2013/525](#), [2327](#), [2017/207](#), [2019/791](#), [1385](#) and [2020/1478](#).

- (b) in paragraph (2)—
- (i) the words from ““qualifying Northern Ireland good”” to the end become sub-paragraph (a); and
  - (ii) after that sub-paragraph insert—
    - “(b) “Regulation (EU) 2017/745” 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](#).”.

**Commencement Information**

**I3** Reg. 31 in force at 27.7.2021, see [reg. 1\(2\)](#)

**Revocation and transitional provision**

**32.** After regulation 3 (scope), insert—

**“Revocation, transitional and saving provisions in respect of Regulation (EU) 2017/745**

- 3ZA.**—(1) Subject to paragraph (2)—
- (a) Parts 2 and 3 only apply in Northern Ireland for the purposes of regulating qualifying devices.
  - (b) Parts 5 to 7 only apply in Northern Ireland for the purposes of regulating qualifying devices and devices within the scope of Part 4.
- (2) The following provisions continue to apply in Northern Ireland in accordance with this paragraph whether or not the device to which they apply is referred to in paragraph (1)—
- (a) for the purposes of registration of medical devices and persons placing medical devices on the market in Northern Ireland—
    - (i) regulation 19 (registration of persons placing general medical devices on the market),
    - (ii) regulation 21B (registration of persons placing active implantable medical devices on the market), and
    - (iii) regulation 53 (fees in connection with the registration of devices and changes to registration details),
 apply until the date which is 6 months after the date of publication of the notice referred to in Article 34(3) of Regulation (EU) 2017/745.
  - (b) Parts 5 to 7 apply for purposes related to the designation of conformity assessment bodies for the purposes of a UK mutual recognition agreement.
- (3) For the purposes of paragraph (1), a device is a qualifying device if, by virtue of Article 120 of Regulation (EU) 2017/745—
- (a) it may be placed on the market, put into service or made available in Northern Ireland in accordance with the requirements of Directive 93/42 or Directive 90/385, rather than Regulation (EU) 2017/745; and

- (b) it is placed on the market, put into service or made available in Northern Ireland in accordance with, and subject to the requirements of and the arrangements set out in, Parts 2, 3 and 5 to 7.”

**Commencement Information**

**I4** Reg. 32 in force at 27.7.2021, see [reg. 1\(2\)](#)

**Amendment of regulation 10A (UK(NI) indication: general medical devices)**

**33.** In regulation 10A, after paragraph (3), insert—

“(3A) The UK(NI) indication may be less than 5mm high provided that it is the same height as the CE marking that it accompanies.”

**Commencement Information**

**I5** Reg. 33 in force at 27.7.2021, see [reg. 1\(2\)](#)

**Amendment of regulation 19 (registration of persons placing general medical devices on the market)**

**34.** In regulation 19—

- (a) omit paragraph (1)(a)(ii);
- (b) in paragraph (1)(b) for “and custom-made devices” substitute “that are not custom-made devices”;
- (c) omit paragraph (2)(c);
- (d) omit paragraph (5); and
- (e) omit paragraph (6).

**Commencement Information**

**I6** Reg. 34 in force at 27.7.2021, see [reg. 1\(2\)](#)

**Amendment of regulation 21B (registration of persons placing active implantable medical devices on the market)**

**35.** In regulation 21B—

- (a) omit paragraph (1)(a)(ii);
- (b) omit paragraph (1)(b);
- (c) omit paragraph (2)(c);
- (d) omit paragraph (5); and
- (e) omit paragraph (6).

**Commencement Information**

**I7** Reg. 35 in force at 27.7.2021, see [reg. 1\(2\)](#)

**Amendment of regulation 24A (UK(NI) indication: active implantable medical devices)**

36. In regulation 24A, after paragraph (3), insert—

“(3A) The UK(NI) indication may be less than 5mm high provided that it is the same height as the CE marking that it accompanies.”

**Commencement Information**

**18** Reg. 36 in force at 27.7.2021, see [reg. 1\(2\)](#)

**Amendment of regulation 36A (UK(NI) indication: in vitro diagnostic medical devices)**

37. In regulation 36A, after paragraph (3), insert—

“(3A) The UK(NI) indication may be less than 5mm high provided that it is the same height as the CE marking that it accompanies.”

**Commencement Information**

**19** Reg. 37 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, PART 8.