

Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on in vitro diagnostic medical devices and repealing Directive 98/79/EC and Commission Decision 2010/227/EU (Text with EEA relevance) (revoked)

## CHAPTER VII

### POST-MARKET SURVEILLANCE, VIGILANCE AND MARKET SURVEILLANCE

#### *Section 1*

#### ***Post-market surveillance***

#### *Article 81*

#### **Periodic safety update report**

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#### **Textual Amendments applied to the whole legislation**

- F1** Regulation revoked (31.12.2020) by [The Medical Devices Regulations 2002 \(S.I. 2002/618\)](#), **reg. 4P** (as inserted by [S.I. 2019/791](#), regs. 1(1), 3(7) (as amended by [S.I. 2020/1478](#), regs. 1(3), Sch. 2 paras. 2, 9(m)); 2020 c. 1, Sch. 5 para. 1(1))

**Status:**

This version of this provision no longer has effect.

**Changes to legislation:**

There are currently no known outstanding effects for the Regulation (EU) 2017/746 of the European Parliament and of the Council, Article 81.