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 2

Vigilance

Article 82

Reporting of serious incidents and field safety corrective actions

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This version of this provision no longer has effect.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 82.