SCHEDULES

SCHEDULE 2

Amendment of the Human Medicines (Amendment etc.) (EU Exit) Regulations 2019

PART 4

Amendment of Part 5 (amendment of Part 5 (marketing authorisations))

68. For regulation 91 (omission of regulation 94A (offences relating to Commission Regulation 2016/161)) substitute—

"Amendment of regulation 94A (offences relating to Commission Regulation 2016/161)

- 91. In regulation 94A—
 - (a) for paragraph (1) substitute—
 - "(1) A person who is—
 - (a) the holder of a UKMA(NI), UKMA(UK) or parallel import licence, or
 - (b) a parallel distributor,
 - is guilty of an offence if the holder fails to comply with a requirement or obligation contained in a provision of Commission Regulation 2016/161 listed in paragraph (2).";
 - (b) for paragraph (3) substitute—
 - "(3) In this regulation "parallel distributor" means a person who imports into Northern Ireland from an EEA state a product which has been granted a marketing authorisation under Regulation (EC) No 726/2004 and in relation to which that person is not the holder of a UKMA(NI), UKMA(UK), Article 126a authorisation, COR(NI), COR(UK), THR(NI) or THR(UK)."."

Commencement Information

I1 Sch. 2 para. 68 in force at 31.12.2020 immediately before IP completion day, see reg. 1

Changes to legislation:
There are currently no known outstanding effects for the The Human Medicines (Amendment etc.) (EU Exit) Regulations 2020, Paragraph 68.