
DRAFT STATUTORY INSTRUMENTS

2019 No.

**The Human Medicines (Amendment
etc.) (EU Exit) Regulations 2019**

PART 10

Amendment of Part 10 (exceptions to requirement for marketing authorisations etc)

Amendment of regulation 168 (use of non-prescription medicines in the course of a business)

135. In regulation 168, in paragraph (8)—

(a) in sub-paragraph (a), for “EEA State” substitute “approved country for import”; and

(b) for sub-paragraph (b) substitute—

“(b) imported from an approved country for import—

(i) it is manufactured or assembled in that country by a person who is the holder of an authorisation in that country in relation to its manufacture or assembly, and

(ii) it is imported by the holder of a wholesale dealer’s licence under Part 3 that includes the import of a medicinal product from such a country.”.