
STATUTORY INSTRUMENTS

2022 No. 352

The Human Medicines (Amendments Relating to the Early Access to Medicines Scheme) Regulations 2022

Amendment of Schedule 4

- 14.**—(1) Schedule 4 (standard provisions of licences under Part 3) is amended as follows.
- (2) After paragraph 14A(1), insert—
- “**14B.** A licence holder may only manufacture or assemble EAMS medicinal products if and to the extent that such products are required for the Early Access to Medicines Scheme.”.
- (3) In paragraph 22(2)—
- (a) in sub-paragraph (5), after “special medicinal products” insert “or EAMS medicinal products”;
- (b) in sub-paragraph (7), after “special medicinal product” insert “or EAMS medicinal product”;
- (c) in sub-paragraph (8) after “special medicinal product” insert “or EAMS medicinal product”; and
- (d) in sub-paragraph (9), after “special medicinal product” insert “or EAMS medicinal product”.
- (4) After paragraph 23A(3), insert—
- “**23B.** A licence holder may only import EAMS medicinal products if and to the extent that such products are required for the Early Access to Medicines Scheme.”.
- (5) After paragraph 33, insert—
- “**33A.** A licence holder may only import EAMS medicinal products if and to the extent that such products are required for the Early Access to Medicines Scheme.”.
- (6) In paragraph 37, after “special medicinal products” insert “or EAMS medicinal products”.
- (7) In paragraph 39, after “special medicinal product” insert “or EAMS medicinal product”.
- (8) In paragraph 40, after “special medicinal product” insert “or EAMS medicinal product”.
- (9) In paragraph 41, after “special medicinal product” insert “or EAMS medicinal product”.

(1) Inserted by [S.I. 2019/775](#).
(2) Amended by [S.I. 2019/775](#).
(3) Inserted by [S.I. 2019/775](#).