STATUTORY INSTRUMENTS

## 1977 No. 675

## **MEDICINES**

The Medicines (Standard Provisions for Licences and Certificates) Amendment Regulations 1977

Made	6th April 1977
Laid before Parliament	21st April 1977
Coming into Operation	12th May 1977

## THE MEDICINES (STANDARD PROVISIONS FOR LICENCES AND CERTIFICATES) AMENDMENT REGULATIONS 1977

- 1. Citation, interpretation and commencement
- 2. Amendment of regulation 2(1) of the principal regulations
- 3. Additional regulations
- 4. Additional schedules Signature

Schedules added to the principal regulations

SCHEDULE 4 -

PART I — Standard provisions for manufacturer's licences and manufacturer's licences of right relating to vaccines

- 1. The licence holder shall ensure that the premises on which...
- 2. The licence holder shall provide separate premises or separate parts...
- 3. The licences holder shall ensure that any procedure which, in...
- 4. The licence holder shall ensure that no person who has...
- 5. Before an animal is used in the production of vaccine...
- 6. (1) The licence holder shall ensure that animals used in...
- The licence holder shall provide a special room capable of...
- 8. Without prejudice to any other requirements to keep records, where...
  - PART II Standard provisions for manufacturer's licences and manufacturer's licences of right relating to smallpox vaccine
- 1. (1) The licence holder shall ensure that animals used in...
- 2. Where it is necessary for an animal which has been...
  - PART III Standard provisions for manufacturer's licences and manufacturer's licences of right relating to BCG vaccine

- 1. The licence holder shall provide separate premises or separate parts...
- 2. The licence holder shall ensure that any procedure which involves...
- 3. The licence holder shall ensure that all media, glassware and...
- 4. The licence holder shall not permit animals to be in...
- 5. (1) The licence holder shall arrange for all persons engaged...
- 6. The licence holder shall ensure that no person who has...
  - PART IV Standard provisions for manufacturer's licences and manufacturer's licences of right relating to toxins
- 1. The licence holder shall ensure that the premises on which...
- 2. The licence holder shall provide separate premises or separate parts...
- 3. The licence holder shall ensure that any procedure which in... PART V — Standard provisions for manufacturer's licences and manufacturer's
  - licences of right relating to sera
- 1. The licence holder shall ensure that the premises on which...
- 2. The licence holder shall ensure that blood used in the...
- 3. The licence holder shall ensure that an adequate system of...
- 4. Before an animal is used in the production of any...
- 5. The licence holder shall notify the licensing authority if any...
- 6. The licence holder shall notify the licensing authority if any...
- 7. The licence holder shall ensure that laboratories in which any...
- 8. The licence holder shall provide such number of sterilizers as...
- 9. Without prejudice to any other requirements to keep records, the...

SCHEDULE 5 —

PART I — Standard provisions for product licences including product licences of right relating to medicinal products to which regulation 5 of these regulations applies

- 1. In this Part of this Schedule "expiry date" means the...
- 2. The licence holder shall, within 28 days of any request...
- 3. Until the expiry date and for six months thereafter the...
- 4. Where the licence holder has supplied the licensing authority with...
- 5. Where the licence holder has been informed by the licensing...
- 6. Unless and to the extent that the licensing authority otherwise...
- 7. (1) Unless the licensing authority otherwise direct in writing, the...
- 8. The licence holder shall not sell, supply, import or export...
- 9. The provisions of this Schedule shall not have effect until... PART II — Tests for sterility
- 1. In this Part of this Schedule— "batch" means a homogeneous...
- 1. In this Fait of this Schedule— batch means a homogeneous.
- 2. (1) The test for bacterial sterility shall be applied to...
- 3. (1) The quantity of the medicinal product required for the...
- 4. (1) Subject to the provisions of sub-paragraph (2) below, the...
- 5. In the case of a medicinal product which is itself...
- 6. Where more than one test is to be performed on...
- 7. (1) The tests for sterility shall be made on a...
- 8. (1) The sample shall be applied to the media selected...
- 9. The tubes or vessels of media to which the sample...
- 10. (1) If at the examination at the end of the...
- 11. (1) Notwithstanding the provisions of paragraph 10 above, where the... PART III — Test for abnormal toxicity
- 1. (1) The amount of medicinal product as is specified in...
- 2. Where, by virtue of the nature of any substance used,... PART IV — Test for pyrogens
- 1. In this Part of this Schedule— "maximum temperature", in relation...

- 2. A test for pyrogenic substances (hereinafter referred to as a...
- 3. A rabbit shall not be used in a pyrogen test—...
- 4. (1) The measurement of the temperatures of the rabbits in...
- 5. A pyrogen test shall be conducted in a quiet room...
- 6. The rabbits used in the pyrogen test shall not be...
- 7. (1) A pyrogen test shall be made by—
- 8. The medicinal product shall be regarded as having satisfied the...

Explanatory Note