

(iv) By retail pharmacists or other authorized retail distributors and by institutions or qualified persons in the duly authorized exercise of therapeutic or scientific functions, or

(v) As "special stocks".

(y) "Territory" means any part of a State which is treated as a separate entity for the application of the system of import certificates and export authorizations provided for in article 31. This definition shall not apply to the term "territory" as used in articles 42 and 46.

2. For the purposes of this Convention a drug shall be regarded as "consumed" when it has been supplied to any person or enterprise for retail distribution, medical use or scientific research; and "consumption" shall be construed accordingly.

ARTICLE 2

Substances under control

1. Except as to measures of control which are limited to specified drugs, the drugs in Schedule I are subject to all measures of control applicable to drugs under this Convention and in particular to those prescribed in articles 4 (c), 19, 20, 21, 29, 30, 31, 32, 33, 34 and 37.

2. The drugs in Schedule II are subject to the same measures of control as drugs in Schedule I with the exception of the measures prescribed in article 30, paragraphs 2 and 5, in respect of the retail trade.

3. Preparations other than those in Schedule III are subject to the same measures of control as the drugs which they contain, but estimates (article 19) and statistics (article 20) distinct from those dealing with these drugs shall not be required in the case of such preparations, and article 29, paragraph 2 (c) and article 30, paragraph 1 (b) (ii) need not apply.

4. Preparations in Schedule III are subject to the same measures of control as preparations containing drugs in Schedule II except that article 31, paragraphs 1 (b) and 4 to 15 need not apply, and that for the purpose of estimates (article 19) and statistics (article 20) the information required shall be restricted to the quantities of drugs used in the manufacture of such preparations.

5. The drugs in Schedule IV shall also be included in Schedule I and subject to all measures of control applicable to drugs in the latter schedule, and in addition thereto:

(a) A Party shall adopt any special measures of control which in its opinion are necessary having regard to the particularly dangerous properties of a drug so included; and

(b) A Party shall, if in its opinion the prevailing conditions in its country render it the most appropriate means of protecting the public health and welfare, prohibit the production, manufacture, export and import of, trade in, possession or use of any such drug except for amounts which may be necessary for medical and scientific research only, including clinical trials therewith to be conducted under or subject to the direct supervision and control of the Party.