Medicated plasters

DEFINITION
Medicated plasters are flexible preparations containing one or more active substances. They are intended to be applied to the skin. They are designed to maintain the active substance(s) in close contact with the skin such that these may be absorbed slowly, or act as protective or keratolytic agents.

Medicated plasters consist of an adhesive basis, which may be coloured, containing one or more active substances, spread as a uniform layer on an appropriate support made of natural or synthetic material. It is not irritant or sensitising to the skin. The adhesive layer is covered by a suitable protective liner, which is removed before applying the plaster to the skin. When removed, the protective liner does not detach the preparation from the outer, supporting layer.

Medicated plasters are presented in a range of sizes directly to the skin. When removed, the protective liner does not detach the preparation from the outer, supporting layer.

TESTS
Dissolution. A suitable test may be required to demonstrate the appropriate release of the active substance(s), for example one of the tests described in Dissolution test for transdermal patches (2.9.4).

01/2005:1154

STICKS

Styli

Additional requirements for sticks may be found, where appropriate, in other general monographs, for example Nasal preparations (0676).

DEFINITION
Sticks are solid preparations intended for local application. They are rod-shaped or conical preparations consisting of one or more active substances alone or which are dissolved or dispersed in a suitable basis which may dissolve or melt at body temperature.

Urethral sticks and sticks for insertion into wounds are sterile.

PRODUCTION
In the manufacture, packaging, storage and distribution of sticks, suitable means are taken to ensure their microbial quality; recommendations on this aspect are provided in the text on Microbiological quality of pharmaceutical preparations (5.1.4).

Urethral sticks and other sterile sticks are prepared using materials and methods designed to ensure sterility and to avoid the introduction of contaminants and the growth of micro-organisms; recommendations on this aspect are provided in the text on Methods of preparation of sterile products (5.1.1).

In the manufacture of sticks means are taken to ensure that the preparation complies with a test for mass uniformity or, where appropriate, a test for uniformity of content.

TESTS
Sterility (2.6.1). Urethral sticks and sticks for insertion into wounds comply with the test for sterility.

LABELLING
The label states:
- the quantity of active substance(s) per stick,
- for urethral sticks and sticks to be inserted into wounds that they are sterile.

01/2005:0478

TABLETS

Compressi

The requirements of this monograph do not necessarily apply to preparations that are presented as tablets intended for use other than by oral administration. Requirements for such preparations may be found, where appropriate, in other general monographs; for example Rectal preparations (1145), Vaginal preparations (1164) and Oromucosal preparations (1807). This monograph does not apply to lozenges, lyophilisates, oral pastes and oral gums. Where justified and authorised, the requirements of this monograph do not apply to tablets for veterinary use.

DEFINITION
Tablets are solid preparations each containing a single dose of one or more active substances and usually obtained by compressing uniform volumes of particles. Tablets are intended for oral administration. Some are swallowed whole, some after being chewed, some are dissolved or dispersed in water before being administered and some are retained in the mouth where the active substance is liberated. The particles consist of one or more active substances with or without excipients such as diluents, binders, disintegrating agents, glidants, lubricants, substances capable of modifying the behaviour of the preparation in the digestive tract, colouring matter authorised by the competent authority and flavouring substances.

Tablets are usually right, circular solid cylinders, the end surfaces of which are flat or convex and the edges of which may be bevelled. They may have lines or break-marks and may bear a symbol or other markings. Tablets may be coated. Where applicable, containers for tablets comply with the requirements for Materials used for the manufacture of containers (3.1 and subsections) and Containers (3.2 and subsections).

Several categories of tablets for oral use may be distinguished:
- uncoated tablets,
- coated tablets,
- effervescent tablets,
- soluble tablets,
- dispersible tablets,
- oродispersible tablets,
- gastro-resistant tablets,
- modified-release tablets.

PRODUCTION
Tablets are usually prepared by compressing uniform volumes of particles or particle aggregates produced by granulation methods. In the manufacture of tablets, means are taken to ensure that they possess a suitable mechanical strength to avoid crumbling or breaking on handling or subsequent processing. This may be demonstrated by examining the Friability of uncoated tablets (2.9.7) and the Resistance to crushing of tablets (2.9.8). Chewable tablets are prepared to ensure that they are easily crushed by chewing.
For tablets for which subdivision is authorised, it is demonstrated to the satisfaction of the competent authority that the subdivided parts comply either with test A for Uniformity of content of single-dose preparations (2.9.6) or with the test for Uniformity of mass of single-dose preparations (2.9.5), as appropriate.

In the manufacture, packaging, storage and distribution of tablets, suitable means are taken to ensure their microbiological quality; recommendations on this aspect are provided in the text on Microbiological quality of pharmaceutical preparations (5.1.4).

**TESTS**

**Uniformity of content** (2.9.6). Unless otherwise prescribed or justified and authorised, tablets with a content of active substance less than 2 mg or less than 2 per cent of the total mass comply with test A for uniformity of content of single-dose preparations. If the preparation has more than one active substance, the requirement applies only to those substances which correspond to the above conditions.

Unless otherwise justified and authorised, coated tablets other than film-coated tablets comply with test A for uniformity of content of single-dose preparations irrespective of their content of active substance(s).

**Uniformity of mass** (2.9.5). Uncoated tablets and, unless otherwise justified and authorised, film-coated tablets comply with the test for uniformity of mass of single-dose preparations. If the test for uniformity of content is prescribed or justified and authorised for all the active substances, the test for uniformity of mass is not required.

**Dissolution.** A suitable test may be carried out to demonstrate the appropriate release of the active substance(s), for example one of the tests described in Dissolution test for solid dosage forms (2.9.3).

Where a dissolution test is prescribed, a disintegration test may not be required.

**Uncoated tablets**

**DEFINITION**

Uncoated tablets include single-layer tablets resulting from a single compression of particles and multi-layer tablets consisting of concentric or parallel layers obtained by successive compression of particles of different composition. The excipients used are not specifically intended to modify the release of the active substance in the digestive fluids.

Uncoated tablets conform to the general definition of tablets. A broken section, when examined under a lens, shows either a relatively uniform texture (single-layer tablets) or a stratified texture (multi-layer tablets) but no signs of coating.

**TESTS**

**Disintegration.** Uncoated tablets comply with the test for disintegration of tablets and capsules (2.9.1). Use water R as the liquid. Add a disc to each tube. Operate the apparatus for 15 min, unless otherwise justified and authorised, and examine the state of the tablets. If the tablets fail to comply because of adherence to the discs, repeat the test on a further 6 tablets omitting the discs. The tablets comply with the test if all 6 have disintegrated.

Chewable tablets are not required to comply with the test.

**Coated tablets**

**DEFINITION**

Coated tablets are tablets covered with one or more layers of mixtures of various substances such as natural or synthetic resins, gums, gelatin, inactive and insoluble fillers, sugars, plasticisers, polyols, waxes, colouring matter authorised by the competent authority and sometimes flavouring substances and active substances. The substances used as coatings are usually applied as a solution or suspension in conditions in which evaporation of the vehicle occurs. When the coating is a very thin polymeric coating, the tablets are known as film-coated tablets.

Coated tablets have a smooth surface which is often coloured and may be polished; a broken section, when examined under a lens, shows a core surrounded by one or more continuous layers with a different texture.

**PRODUCTION**

Where justified, uniformity of mass or uniformity of content of coated tablets other than film-coated tablets may be ensured by control of the cores.

**TESTS**

**Disintegration.** Coated tablets other than film-coated tablets comply with the test for disintegration of tablets and capsules (2.9.1). Use water R as the liquid. Add a disc to each tube. Operate the apparatus for 30 min, unless otherwise justified and authorised, and examine the state of the tablets. If any of the tablets has not disintegrated, repeat the test on a further 6 tablets, replacing water R with 0.1 M hydrochloric acid. The tablets comply with the test if all 6 have disintegrated in the acid medium.

Film-coated tablets comply with the disintegration test prescribed above except that the apparatus is operated for 30 min, unless otherwise justified and authorised.

If coated tablets or film-coated tablets fail to comply because of adherence to the discs, repeat the test on a further 6 tablets omitting the discs. The tablets comply with the test if all 6 have disintegrated.

Chewable coated tablets are not required to comply with the test.

**Effervescent tablets**

**DEFINITION**

Effervescent tablets are uncoated tablets generally containing acid substances and carbonates or hydrogen carbonates which react rapidly in the presence of water to release carbon dioxide. They are intended to be dissolved or dispersed in water before administration.

**TESTS**

**Disintegration.** Place 1 tablet in a beaker containing 200 ml of water R at 15-25 °C; numerous bubbles of gas are evolved. When the evolution of gas around the tablet or its fragments ceases the tablet has disintegrated, being either dissolved or dispersed in the water so that no agglomerates of particles remain. Repeat the operation on 5 other tablets. The tablets comply with the test if each of the 6 tablets used disintegrates in the manner prescribed within 5 min, unless otherwise justified and authorised.
Soluble tablets

DEFINITION
Soluble tablets are uncoated or film-coated tablets. They are intended to be dissolved in water before administration. The solution produced may be slightly opalescent due to the added excipients used in the manufacture of the tablets.

TESTS
Disintegration. Soluble tablets disintegrate within 3 min when examined by the test for disintegration of tablets and capsules (2.9.1), but using water R at 15-25 °C.

Dispersible tablets

DEFINITION
Dispersible tablets are uncoated or film-coated tablets intended to be dispersed in water before administration giving a homogeneous dispersion.

TESTS
Disintegration. Dispersible tablets disintegrate within 3 min when examined by the test for disintegration of tablets and capsules (2.9.1), but using water R at 15-25 °C.

Finness of dispersion. Place 2 tablets in 100 ml of water R and stir until completely dispersed. A smooth dispersion is produced, which passes through a sieve screen with a nominal mesh aperture of 710 µm.

Orodispersible tablets

DEFINITION
Orodispersible tablets are uncoated tablets intended to be placed in the mouth where they disperse rapidly before being swallowed.

TESTS
Disintegration. Orodispersible tablets disintegrate within 3 min when examined by the test for disintegration of tablets and capsules (2.9.1).

Modified-release tablets

DEFINITION
Modified-release tablets are coated or uncoated tablets which contain special excipients or which are prepared by special procedures, or both, designed to modify the rate, the place or the time at which the active substance(s) are released. Modified-release tablets include prolonged-release tablets, delayed-release tablets and pulsatile-release tablets.

PRODUCTION
A suitable test is carried out to demonstrate the appropriate release of the active substance(s).

Gastro-resistant tablets

DEFINITION
Gastro-resistant tablets are delayed-release tablets that are intended to resist the gastric fluid and to release their active substance(s) in the intestinal fluid. Usually they are prepared from granules or particles already covered with a gastro-resistant coating or in certain cases by covering tablets with a gastro-resistant coating (enteric-coated tablets). Tablets covered with a gastro-resistant coating conform to the definition of coated tablets.

PRODUCTION
For tablets prepared from granules or particles already covered with a gastro-resistant coating, a suitable test is carried out to demonstrate the appropriate release of the active substance(s).

TESTS
Disintegration. For tablets covered with a gastro-resistant coating carry out the test for disintegration (2.9.1) with the following modifications. Use 0.1 M hydrochloric acid as the liquid. Operate the apparatus for 2 h, or other such time as may be justified and authorised, without the discs and examine the state of the tablets. The time of resistance to the acid medium varies according to the formulation of the tablets to be examined. It is typically 2 h to 3 h but even with authorised deviations is not less than 1 h. No tablet shows signs of either disintegration (apart from fragments of coating) or cracks that would allow the escape of the contents. Replace the acid by phosphate buffer solution pH 6.8 R and add a disc to each tube. Operate the apparatus for 60 min and examine the state of the tablets. If the tablets fail to comply because of adherence to the discs, repeat the test on a further 6 tablets omitting the discs. The tablets comply with the test if all 6 have disintegrated.

Dissolution. For tablets prepared from granules or particles already covered with a gastro-resistant coating, a suitable test is carried out to demonstrate the appropriate release of the active substance(s), for example the test described in Dissolution test for solid dosage forms (2.9.3).

Tablets for use in the mouth

DEFINITION
Tablets for use in the mouth are usually uncoated tablets. They are formulated to effect a slow release and local action of the active substance(s) or the release and absorption of the active substance or substances at a defined part of the mouth. They comply with the requirements of the monograph on Orodispersal preprations (1807).

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TAMPONS, MEDICATED

Tamponae medicatae

Additional requirements for medicated tampons may be found, where appropriate, in other general monographs, for example Rectal preparations (1145), Vaginal preparations (1164) and Ear preparations (0652).

DEFINITION
Medicated tampons are solid, single-dose preparations intended to be inserted into the body cavities for a limited period of time. They consist of a suitable material such as cellulose, collagen or silicone impregnated with one or more active substances.

PRODUCTION
In the manufacture, packaging, storage and distribution of medicated tampons, suitable means are taken to ensure their microbial quality; recommendations on this aspect are provided in the text on Microbiological quality of pharmaceutical preparations (5.1.4).

LABELLING
The label states the quantity of active substance(s) per tampon.