Section 3: Pharmaceutical form

SmPC training presentation

Note: for full information refer to the European Commission’s Guideline on summary of product characteristics (SmPC)

SmPC Advisory Group
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I. General objectives of section 3

This section should state the pharmaceutical form according to the European Pharmacopoeia and a visual description of the product.
II.1 Term for pharmaceutical form

Should be described by a **full standard term of European Pharmacopoeia using SINGULAR FORM**

Term used in Section 1 should be the same as used in this section

If a **patient friendly term** (formerly **short term**) of the European Pharmacopoeia is used on **small immediate packaging material**, the **patient friendly term** should be **added in brackets in this section**

http://www.edqm.eu/StandardTerms/ (subscription only access)
Examples full standard term

A full term of European Pharmacopoeia using singular form

Prolonged-release tablet.

Solution for intraperitoneal use.

Powder for concentrate for solution for infusion.
### Examples short term

<table>
<thead>
<tr>
<th>A full term of European Pharmacopoeia using singular form</th>
<th>The patient friendly (formerly short) term should be added in brackets in this section</th>
</tr>
</thead>
<tbody>
<tr>
<td>Film-coated tablet (tablet).</td>
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<tr>
<td>Eye drops, suspension (eye drops).</td>
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<tr>
<td>Concentrate for solution for infusion (sterile concentrate).</td>
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</tbody>
</table>
II.2 Description of pharmaceutical form

A **VISUAL DESCRIPTION** of the appearance of the product such as

- **COLOUR**
- **markings**
- **SIZE**

For **tablets designed with a score line**, information should be given on whether or not the tablet can be divided into equal halves.

Information on **pH** and **osmolarity**, as appropriate.

**Appearance before reconstitution** should be stated in this section. **Appearance of the product after reconstitution** should be stated in sections 4.2 & 6.6.
A **VISUAL DESCRIPTION** of the appearance of the product

**COLOUR**  **markings**  **SIZE**

7.0 mm, round, normal convex, white film-coated tablets debossed “OZ 2.5” on one side and “G” on the other side.
Example-pH and osmolarity

A **VISUAL DESCRIPTION** of the appearance of the product

**COLOUR**

Clear, colourless to pale yellow solution, with a pH of 6.0-7.5 and an osmolality of 260 – 320mOsm/kg.
Example–score line

For tablets designed with a score line, information should be given on whether or not reproducible dividing of the tablets has been shown.

The scoreline is only to facilitate breaking for ease of swallowing and not to divide into equal doses.

The tablet can be divided into equal halves.

(SmPC Guideline)
Example-appearance before reconstitution

**Appearance before reconstitution** should be stated in section 3. Appearance after reconstitution should be stated in section 4.2 and 6.6.

Active substance X 250 IU powder and solvent for solution for injection

**Section 3**
White to off-white cake/powder. Clear, colourless solvent.

**Section 4.2**
For reconstitution instructions prior to administration, see section 6.6.

**Section 6.6**
The solution will be clear or slightly opalescent and colourless. The solution is to be discarded if visible particulate matter or discolouration is observed.
Thank you for consulting this training presentation

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Please note the presentation includes examples that may have been modified to best illustrate the related principle