

TECHNICAL DATA SHEET

002540-TDS-ENG-2026

| IVERMECTINA USO HUMANO (EUR. PH.) | | |
|--|--|------------------------------|
| DESCRIPTION DCI: IVERMECTIN | | DESCRIPTION DOE: IVERMECTINA |
| CAS Nº: 70288-86-7 | EC Nº: 274-536-0 | AEMPS CODE: 7778A |
| MOL. WEIGHT: 1736,16 | MOL. FORMULA: C ₉₅ H ₁₄₆ O ₂₈ | ARTICLE CODE: 002540 |

| ATTRIBUTES | SHOULD BE |
|--|---|
| Appearance | White or yellowish-white, crystalline powder, slightly hygroscopic |
| Solubility | Practically insoluble in water, freely soluble in methylene chloride, soluble in ethanol (96 %) |
| Identification A | Complies |
| Identification B | Complies |
| Appearance of solution | Clear and not more intensely coloured than ref. sol. BY7 |
| Specific optical rotation | -20 / -17 |
| Related substances | |
| Impurity with a relative retention of 1.3 to 1.5 | = < 2.5 % |
| Any other impurity | = < 1 % |
| Total impurities | = < 5 % |
| Ethanol and formamide | |
| Ethanol | = < 5.0 % |
| Formamide | = < 3.0 % |
| Water | = < 1.0 % |
| Sulfated ash | = < 0.1 % |
| Assay | |
| Ivermectin (H2B1a + H2B1b) | 95.0 - 102.0 % |
| Ratio H2B1a/(H2B1a + H2B1b) | = > 90.0 % |

COMPLIES WITH

European Pharmacopoeia 12.2

STORAGE

Keep the container tightly closed in a dry and cool place.

REMARKS

Ivermectin is subjected to the requirements of the ICH Q3D "Elemental Impurities" guideline and the requirements of guides EMA/CHMP/ICH/82260/2006.

Absence of N-nitrosamines impurities has been ensured after a risk evaluation according to ICH Q9, ICH M7 and in accordance with guidelines EMA/428592/2019 Rev 2 and EMA/189634/2019.

Certificates of residual solvents, allergens, non-GMO and BSE-TSE, among others, are available upon request.

All methods of analysis are validated by official pharmacopoeias or are validated by internal methods of the manufacturer, which can be obtained at specific request. The above information does not exempt from the obligation to identify the

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product before use.