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Dolasetron Mesylate Compounded Oral Solution

DEFINITION

Dolasetron Mesylate Compounded Oral Solution contains NLT 90.0% and NMT 110.0% of the labeled amount of dolasetron mesylate $(C_{10}H_{20}N_2O_3 \cdot CH_aO_3S)$.

Prepare Dolasetron Mesylate Compounded Oral Solution 10 mg/mL as follows (see <u>Pharmaceutical Compounding—Nonsterile Preparations</u> (795)).

Dolasetron Mesylate powder	1 g
Vehicle for Oral Solution (regular or sugar-free), NF, a sufficient quantity to make	100 mL

Add *Dolasetron Mesylate powder* and 15 mL of *Vehicle* to a mortar, and mix. Add the *Vehicle* in small portions almost to volume, and mix thoroughly after each addition. Transfer the contents of the mortar, stepwise and quantitatively, to a calibrated bottle. Add enough *Vehicle* to bring to final volume, and mix.

ASSAY

Proceduri

Solution A: 0.05 M ammonium acetate adjusted with diluted ammonium hydroxide to a pH of 7.5

Mobile phase: Acetonitrile and Solution A (24:76). Filter, and degas.

Diluent: Acetonitrile and water (24:76)

Standard stock solution: 500 µg/mL of USP Dolasetron Mesylate RS in Diluent

Standard solution: 10 µg/mL of USP Dolasetron Mesylate RS from Standard stock solution in Mobile phase

 $\textbf{Sample solution:} \ 10\ \mu\text{g/mL of dolase} tron\ mesylate\ prepared\ from\ Oral\ Solution\ and\ \textit{Diluent}.\ Shake\ each\ sample\ thoroughly\ by\ hand\ for\ 15\ s,$

centrifuge at 1000 rpm for 2 min, and use the supernatant.

Chromatographic system

(See Chromatography (621), System Suitability.)

Mode: LC

Detector: UV 280 nm

Column: 4.6-mm × 15-cm; 3-µm packing L10

Column temperature: 30° Flow rate: 0.8 mL/min Injection volume: 5 µL System suitability

Sample: Standard solution

[Note—The retention time for dolasetron mesylate is about 6.9 min.]

Suitability requirements

Relative standard deviation: NMT 1.4% for replicate injections

Analysis

Samples: Standard solution and Sample solution

 $Calculate \ the \ percentage \ of \ the \ labeled \ amount \ of \ dolase tron \ mesylate \ (C_{19}H_{20}N_2O_3 \cdot CH_4O_3S) \ in \ the \ portion \ of \ Oral \ Solution \ taken:$

Result =
$$(r_{II}/r_{S}) \times (C_{S}/C_{II}) \times 100$$

 r_{ij} = peak response from the Sample solution

 r_{o} = peak response from the Standard solution

 C_S = concentration of <u>USP Dolasetron Mesylate RS</u> in the *Standard solution* (µg/mL)

 C_{μ} = nominal concentration of dolasetron mesylate in the Sample solution (µg/mL)

Acceptance criteria: 90.0%-110.0%

SPECIFIC TESTS

• PH (791): 3.6-4.6

ADDITIONAL REQUIREMENTS

- PACKAGING AND STORAGE: Package in tight, light-resistant containers. Store at controlled room temperature, or in a refrigerator.
- BEYOND-USE DATE: NMT 90 days after the date on which it was compounded when stored at controlled room temperature, or in a refrigerator
- LABELING: Label it to state the Beyond-Use Date.
- USP Reference Standards (11)

USP Dolasetron Mesylate RS

Auxiliary Information - Please check for your question in the FAQs before contacting USP.

Topic/Question	Contact	Expert Committee
DOLASETRON MESYLATE COMPOUNDED ORAL SOLUTION	Brian Serumaga Science Program Manager	CMP2020 Compounding 2020

Chromatographic Database Information: Chromatographic Database

Most Recently Appeared In:

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