Vancomycin Capsules

Details for the public consultation of this monograph are as follows:

<table>
<thead>
<tr>
<th>EAG ABS</th>
<th>Antibiotics</th>
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</table>
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| Deadline for Comment | 31st December 2021 |
| Target Publication (subject to change) | BP 2023 |
| Notes: | Revised Monograph  
If limits are too restrictive, please provide batch/stability data to demonstrate that an increase is required.  
Production: Statement replaced with Dissolution test.  
Identification: Test A, concordant retention time in Vancomycin B replaced with concordant UV spectra and retention time in Related substance and Vancomycin B.  
Dissolution: New test added.  
Water: Tests removed.  
Impurities: New section added. |

Action and use

Glycopeptide antibacterial.

DEFINITION

Vancomycin Capsules contain Vancomycin Hydrochloride.

The capsules comply with the requirements stated under Capsules and with the following requirements.

IDENTIFICATION

A. In the test for Related substances and Vancomycin B, record the UV spectrum of the principal peak in the chromatograms obtained with solutions (1) and (2) with a diode array detector in the range of 210 to 400 nm:

the UV spectrum of the principal peak in the chromatogram obtained with solution (1) is similar to that of the peak in the chromatogram obtained with solution (2);
the retention time of the principal peak in the chromatogram obtained with solution (1) is similar to that of the peak in the chromatogram obtained with solution (2).

B. The contents of the capsules yield reaction A characteristic of chlorides, Appendix VI.

TESTS

Dissolution

Comply with the requirements in the dissolution test for tablets and capsules, Appendix XII B1.

TEST CONDITIONS

(a) Use Apparatus 1, rotating the basket at 100 revolutions per minute.
(b) Use 900 mL of water, at a temperature of 37°, as the dissolution medium.

PROCEDURE

After 45 minutes withdraw a sample of the medium and filter. Dilute to 100 IU/mL with water, if necessary, and carry out the microbiological assay of antibiotics, Appendix XIV A.

LIMITS

The amount of vancomycin released is not less than 85% (Q) of the stated amount.

Related substances and Vancomycin B

Carry out the method for liquid chromatography, Appendix III D, using the following solutions and the normalisation procedure.

(1) Shake a quantity of the capsule contents containing 0.1 g of Vancomycin Hydrochloride with 20 mL of water and dilute to 25 mL.
(2) 0.4% w/v of vancomycin for system suitability EPCRS.
(3) Expose 4 mg of vancomycin for system suitability EPCRS to 80-100% relative humidity at 42 ± 2° for at least 7 days. Allow to cool. Add 1 mL of water and dissolve the sample with the aid of ultrasound (in situ generation of impurities B, D, E, G, and L).
(4) Dilute 1 volume of solution (2) to 100 volumes with 0.1% v/v acetic acid. Dilute 1 volume of this solution to 10 volumes with the same solvent.

CHROMATOGRAPHIC CONDITIONS

(a) Use a stainless steel column (15 cm × 2.1 mm) packed with end-capped, charged-surface, ethylene-bridged, octadecylsilyl silica gel for chromatography (hybrid material) (1.7 µm) (Acquity CSH C18 is suitable).
(b) Use gradient elution and the mobile phase described below.
(c) Use a flow rate of 0.3 mL per minute.
(d) Use a column temperature of 40°.
(e) Use a detection wavelength of 280 nm.
(f) Use an autosampler temperature of 5°.
(g) Inject 20 µL of each solution.

MOBILE PHASE

Mobile phase A 3 volumes of acetonitrile, 4 volumes of methanol and 93 volumes of a 0.7% w/v solution of tris(hydroxymethyl)aminomethane previously adjusted to pH 8.1 with a 20% v/v solution of glacial acetic acid.
Mobile phase B 10 volumes of acetonitrile, 40 volumes of methanol and 50 volumes of a 0.7% w/v solution of tris(hydroxymethyl)aminomethane previously adjusted to pH 8.1 with a 20% v/v solution of glacial acetic acid.

<table>
<thead>
<tr>
<th>Time (Minutes)</th>
<th>Mobile phase A (% v/v)</th>
<th>Mobile phase B (% v/v)</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-7</td>
<td>88</td>
<td>12</td>
<td>isocratic</td>
</tr>
<tr>
<td>7-21</td>
<td>88→75</td>
<td>12→25</td>
<td>linear gradient</td>
</tr>
<tr>
<td>21-35</td>
<td>75→25</td>
<td>25→75</td>
<td>linear gradient</td>
</tr>
<tr>
<td>35-37</td>
<td>25</td>
<td>75</td>
<td>isocratic</td>
</tr>
<tr>
<td>37-38</td>
<td>25→88</td>
<td>75→12</td>
<td>linear gradient</td>
</tr>
<tr>
<td>38-45</td>
<td>88</td>
<td>12</td>
<td>re-equilibration</td>
</tr>
</tbody>
</table>

When the chromatograms are recorded under the prescribed conditions the retention times relative to vancomycin B (retention time, about 19 minutes) are: impurity E, about 0.37; impurity L, about 0.66; impurity B, about 0.70; impurity A, about 0.76; impurity F, about 0.82; impurity G, about 0.90; impurity H, about 0.94; impurity M, about 1.11; impurity I, about 1.14; impurity J, about 1.20; impurity D, about 1.24; impurity K, about 1.50 and impurity C, about 1.86.

SYSTEM SUITABILITY

The test is not valid unless, in the chromatogram obtained with solution (3):

the resolution between the peaks due to impurity L and impurity B is between 1.5 to 5.0;

the resolution between the peaks due to impurity G and impurity H is between 1.5 to 4.0.

If the resolution between the peaks due to impurities L and B is greater than 5.0, adjust the pH of solution A to a lower value. If the resolution between the peaks due to impurities G and H is greater than 4.0, adjust the pH of solution A to a higher value.

LIMITS

Identify any peaks in the chromatogram obtained with solution (1) corresponding to vancomycin impurities A, C, F, H, I, J, K and M using the chromatogram obtained with solution (2) and any peaks corresponding to vancomycin impurities B, D, E, G, and L using the chromatogram obtained with solution (3).

In the chromatogram obtained with solution (1), integrate all peaks present with an area greater than the area of the principal peak in the chromatogram obtained with solution (4) to determine the total peak area. Calculate the percentage content of each of the components and impurities by normalisation:

the content of Vancomycin B is not less than 91.0%;

the area of any peaks due to impurities A or H are not greater than 3.0%;

the sum of the areas of any peaks due to impurities B and E is not greater than 2.0%;

the area of any peak due to impurity J is not greater than 1.6%;

the area of any peaks due to impurities D, F or M are not greater than 1.5%;

the area of any peaks due to impurities G, I or K are not greater than 1.2%;
the area of any peak due to impurity C is not greater than 1.0%;

the area of any other secondary peak is not greater than 0.8%;

the sum of the areas of all the secondary peaks is not greater than 9.0%.

**ASSAY**

Dissolve a quantity of the mixed contents of 20 capsules containing 1,000,000 IU of vancomycin as completely as possible in sufficient water to produce 1000 mL. Dilute 10 mL to 100 mL and carry out the *microbiological assay of antibiotics*, Appendix XIV A. The precision of the assay is such that the fiducial limits of error are not less than 95% and not more than 105% of the estimated potency.

The upper fiducial limit of error is not less than 90.0% and the lower fiducial limit of error is not more than 110.0% of the stated number of IU.

**LABELLING**

The label states (1) the total number of IU (Units) and (2) the number of IU (Units) per mg in each capsule.

**IMPURITIES**

The impurities limited by the requirements of this monograph include those listed under Vancomycin Hydrochloride.