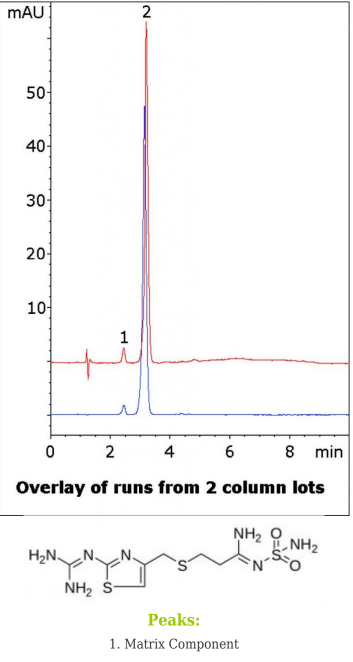
MICROS

Famotidine Tablet Analyzed with HPLC – AppNote

API Separation from Matrix Component

This Method for Analysis of Famotidine Tablets is easy to perform and produces a Symmetrical Peak Shape for the API. This compound has numerous amines which can be problematic in terms of Peak Shape with conventional Columns. Separation from a component from the tablet extract matrix is obtained as well, illustrating specificity of the Method.

Reproducibility is shown by the overlay of runs from two different Column lots.



2. Famotidine

Method Conditions



Column: Cogent Diamond Hydride™, 4µm, 100Å

Catalog No.: 70000-7.5P

Dimensions: 4.6 x 75mm

Mobile Phase:

A: DI Water with 0.1% Trifluoroacetic Acid (TFA) v/v

B: Acetonitrile with 0.1% Trifluoroacetic Acid (TFA) v/v

Gradient:

Time (minutes)	%B
0	95
2	95
6	50
7	95

Post Time: 3 minutes Injection vol.: 1µL Flow rate: 1.0mL / minute Detection: UV @ 265nm

Sample Preparation: 10mg strength Famotidine tablet was ground and added to a 25mL volumetric flask. A portion of 50:50 Solvent A / Solvent B diluent was added and the flask was sonicated 10 minutes. It was then diluted to mark and filtered with a 0.45µm Nylon Syringe Filter (MicroSolv Tech Corp.).

to: 0.9 minutes



Attachment

No 221 Famotidine Tablet Analyzed with HPLC pdf 0.4 Mb Download File

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