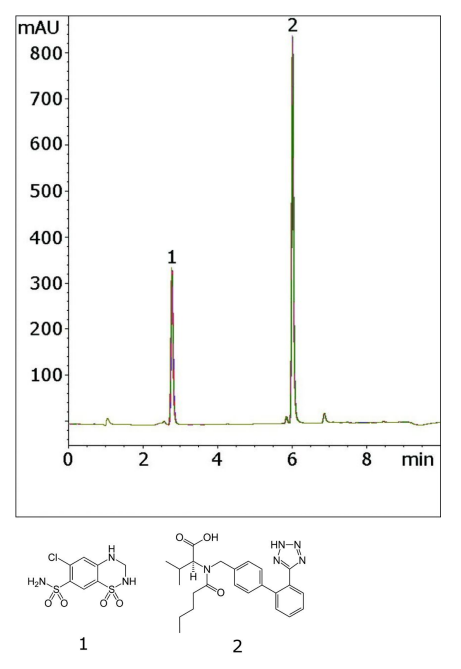
# MICROS

## Valsartan and Hydrochlorothiazide Analyzed with HPLC – AppNote

## **Improved Gradient Method with Faster Equilibration**

The USP Assay Method for Valsartan in combination with Hydrochlorothiazide features a 27 minute gradient with a 13 minute re-equilibration for a total run time of 40 minutes. In this method, the run time was a quarter of the USP method, and the Column equilibrates much faster when gradients are used. This demonstrates a substantial time and solvent savings for the analytical laboratory.

Five Chromatograms are overlaid below which shows the Robustness and Precision of this Method.





## 1. Hydrochlorothiazide (HCT)

2. Valsartan

## **Method Conditions**

Column: Cogent Bidentate C18™, 4µm, 100Å

Catalog No.: 40018-75P

Dimensions: 4.6 x 75mm

### Mobile Phase:

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

B: Acetonitrile / 0.1% Trifluoroacetic Acid (TFA)

### Gradient:

| Time (minutes) | %B |
|----------------|----|
| 0              | 10 |
| 8              | 90 |
| 9              | 10 |

Post Time: 1 minute

Injection vol.: 10µL

Flow rate: 1.0ml / minute

Detection: UV @ 265nm

#### Sample Preparation:

Stock Solution: A Diovan® HCT brand tablet containing 160mg Valsartan and 25 mg Hydrochlorothiazide was ground and added to a 50 mL volumetric flask. The flask was diluted to mark with 50:50 Solvent A / Solvent B mixture and sonicated. A portion was then filtered with a 0.45µm Nylon Syringe Filter AQ<sup>™</sup> Brand (MicroSolv Tech Corp.).

*Working Solution:* 100µL of the stock solution was diluted with 900µL of a 50:50 Solvent A / Solvent B mixture.

**to**: 1 minute

Comparison of This Method and the USP MethodBidentate C18Ordinary C18Total Run Time10 MinutesColumn Volume Equilibration1 ColumnSolvent Usage per Run10ml



Attachment

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