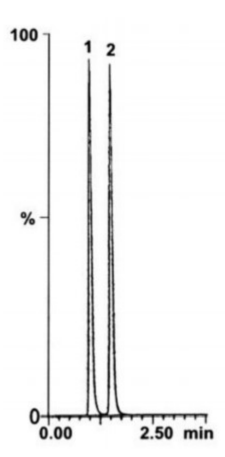
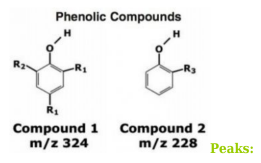


## Substituted Phenols Analyzed with LCMS - AppNote

## **Separation by Functional Groups**

Two proprietary compounds, which are precursors for a catalyst or Prodrugs are analyzed using a C18 Column under Normal Phase conditions. Separation of the two compounds is extremely reproducible (%RSD 0.1) and is very easy. The amount of moisture in your Mobile Phase is not an issue with this Method.





- 1. Compound 1
- 2. Compound 2

## **Method Conditions**

Column: Cogent Bidentate C18<sup>™</sup>, 4μm, 100Å

Catalog No.: 40018-75P Dimensions: 4.6 x 75mm

Mobile Phase: 95% Hexane / 5% Ethyl Acetate



Injection vol.:  $1\mu L$ 

Flow rate: 1mL / minute

Detection: Mass Spectrometer - Atmospheric Pressure Chemical Ionization in positive mode: APCI+ Single Ion

Monitoring

Sample Preparation: 1mg / mL of proprietary compound. 1 (m/z 324) and 2 (m/z 228) dissolved in the Mobile

Phase.

**Notes:** Because silanols on the Silica surface are substituted with Si-H, Water is not retained by the Stationary Phase, so drying of all the Solvents is not essential and analyses are very reproducible.



## Attachment

No 23 Substituted Phenols Analyzed with HPLC pdf 0.2 Mb Download File

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