Preparative Scale Chromatography of a Commercially-Available Sweetener using Hydrophilic Interaction Chromatography

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• Jo-Ann M. Jablonski
• Christopher J. Hudalla
• Kenneth J. Fountain

• Waters Corporation

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Abstract

The analysis and isolation of polar compounds from natural products is often challenging because of the difficulty in retaining hydrophilic compounds on a reversed-phase column. Hydrophilic Interaction Chromatography (HILIC), is an orthogonal chromatographic separation technique which separates hydrophilic compounds by their interaction with a polar stationary phase. In this application note, we demonstrate the utility of the BEH Amide particle for the analysis and isolation of a stevia-related compound in a commercially-available sweetener.

Benefits

BEH Amide columns are specifically designed to enhance the retention of polar compounds by HILIC, making analysis, scaling, and isolation of these molecules easy.

- The isolation of polar compounds on BEH Amide columns is easier because hydrophilic components are retained longer than on reversed-phase packings allowing separation to occur.
- The BEH Amide bonded phase is more compatible with slightly higher aqueous content in mobile phases and sample diluents than unbonded HILIC stationary phases, making the BEH Amide column a more viable alternative for the purification of hydrophilic compounds such as sugars and peptides.
- Improved mass loading of polar compounds on BEH Amide columns reduces the number of injections required to isolate the product, promoting process efficiency.
- Reproducible chromatograms at the analytical and preparative scales reduce ambiguity in the identification of target compounds and demonstrate BEH Amide column scalability.

Introduction

Although reversed-phase is usually the chromatographic mode of choice for many samples, extremely polar compounds are often difficult to analyze because they do not retain on non-polar columns. Hydrophilic Interaction Chromatography (HILIC) is a complementary technique that utilizes a polar stationary phase to bind hydrophilic compounds. Analytes are separated based on a unique combination of liquid-liquid partitioning, adsorption, ionic interaction and hydrophobic retention mechanisms. Compounds elute from the column as the gradient transitions from low aqueous to high aqueous mobile-phase composition. The BEH Amide column, with a trifunctionally-bonded amide phase, was first introduced in 2009 with 1.7 \( \mu \)m particles for the analysis of polar compounds using the ACQUITY UPLC System. Demand for a column capable of analyzing and isolating compounds such as saccharides, synthetic sugars, glycopeptides, and polar compounds from natural products has driven the development of a larger 5 \( \mu \)m particle for use in preparative HPLC applications. In this application note, we demonstrate the utility of the BEH Amide particle for the analysis and isolation of a stevia-related compound in a commercially-available sweetener.

Experimental

LC Conditions
System: Waters 2525 Binary Gradient Module, 2767 Sample Manager, Column Fluidics Organizer, 2996 Photodiode Array Detector, ZQ 2000 Mass Spectrometer, and 2420 Mass Detector

Columns: XBridge BEH Amide, 5 μm, 4.6 x 150 mm, part number 186006595

XBridge BEH Amide, 5 μm, 19 x 150 mm, part number 186006605

Column Temp.: 25 °C

Mobile Phase A: 80/20 acetonitrile/water with 0.1% ammonium hydroxide

Mobile Phase B: 30/70 acetonitrile/water with 0.1% ammonium hydroxide

Weak Needle Wash: 75/25 acetonitrile/water

Strong Needle Wash: 20/80 acetonitrile/water

Seal Wash: 50/50 acetonitrile/water

Sample Diluent: 50/50 acetonitrile/water

Flow Rate: Reported in figures

Gradient: Reported in figures

Injection Volume: Reported in figures

Sample Preparation

Analytical Scale
1.2 mg of commercially-available stevia sweetener was dissolved in 1 mL 50/50 acetonitrile/water. The sample mixture was vortexed and filtered through a 13 mm, 0.45 μm GHP syringe filter, part number WAT200516.

**Preparative Scale**

832 mg of commercially-available stevia sweetener were dissolved in 20 mL 50/50 acetonitrile/water. The sample mixture was vortexed and filtered through a 25 mm, 0.45 μm GHP syringe filter, part number WAT200514.

**Results and Discussion**

**Analytical HPLC**

The analysis and isolation of polar compounds from natural products is often challenging because of the difficulty in retaining hydrophilic compounds on a reversed-phase column. Hydrophilic Interaction Chromatography (HILIC), is an orthogonal chromatographic separation technique which separates hydrophilic compounds by their interaction with a polar stationary phase. Liquid-liquid partitioning, adsorption, ion exchange, and hydrogen bonding mechanisms all contribute to the retention of the sample. Analytes are eluted from the column by increasing the polarity of the mobile phase. The selectivity and retentivity of compounds on different stationary phases is dependent upon the specific properties of the column packing. As shown in Figure 1, the selectivity and retentivity of the analytes are different for each of the three HILIC stationary phases when the column dimensions and the chromatographic method are held constant. The BEH Amide column shows the most retention for the various types of compounds and a different selectivity compared to the other two columns.

Figure 1. Here is a chromatographic representation of the differences in retentivity and selectivity of the 3 HILIC stationary phases. The Atlantis HILIC Silica provides more retention and a different selectivity compared to the BEH HILIC (due to its higher surface polarity). The Amide column exhibits the most retention for various types of compounds and a different selectivity compared to the other two columns. Conditions: Isocratic, 12 mM ammonium formate (pH 3) with 90% acetonitrile; UV at 254 nm. Compounds: (1) acenaphthene, (2) thymine, (3) 5-fluoroorotic acid, (4) adenine, (5) cytosine.

Although polar compound retention is the primary reason for employing HILIC chromatography, the sample diluent also plays a role in retention, influencing sample solubility and peak shape. Traditional unbonded HILIC stationary phases usually require diluents and mobile phases with high organic concentration which limit the solubility of polar compounds at the high sample concentrations used in prep chromatography. Small amounts of water, even 10-20%, make the injection solvent incompatible with initial HILIC conditions on unbonded phases. Since the BEH Amide bonded phase tolerates mobile phases and injection solvents which are higher in aqueous content, polar compounds like sugars and peptides can be solubilized at concentrations amenable to preparative chromatography. In addition, because of the higher organic content employed in HILIC in general, the mass spectrometer response is usually enhanced for ionizable compounds and the fraction drying time is reduced.

Stevia rebaudiana Bertoni is an herb that grows as a small shrub in the mountainous regions of Paraguay and Brazil. The natural sweetness of stevia is attributed to stevioside and rebaudioside, two polar glycosides isolated from the leaves of the stevia plant. Stevioside and rebaudioside are about 300 times sweeter than
sugar, yet they are calorie- and carbohydrate-free. These properties alone help to promote Stevia’s acceptance for use in food products.

As shown in Figure 2, the analysis of a commercially-available stevia sweetener on a 5 μm, 4.6 x 150 mm BEH Amide column indicates the presence of a minor component eluting prior to the main peak. Upon further investigation, it was determined that the stevia-related compound is actually the minor component in the formulation. To better understand the composition of a compound mixture, scientists often isolate minor components for identification. Repetitive injections of the sample are usually required to effectively isolate an appropriate amount for subsequent analyses.

Figure 2. Analysis of a commercially-available stevia sweetener on a BEH Amide, 4.6 x 150 mm, 5 μm column.

A loading study was performed at the small scale to determine the sample mass that could be loaded on the column without compromising the resolution. Improved peak resolution leads to higher mass capacity and better purity for isolated compounds. Figure 3 illustrates the results of a loading study performed on the analytical column with the maximum injection volume of 50 μL, equivalent to 2.1 mg of sample on column.

Figure 3. ELS detection of stevia sweetener loaded in increasing amounts on a BEH Amide, 4.6 x 150 mm, 5 μm column.

**Preparative HPLC**

A sample mass of 2.1 mg on the analytical column scales to about 36 mg on a 19 mm preparative column, but excellent resolution between the minor component and the larger main peak suggested that an even higher load on the 19 mm preparative column would be acceptable.

Without further investigation, the separation method was scaled to the 19 x 150 mm column. As shown in Figure 4, 2 mL of the sample mix, equivalent to about 83 mg on column, gives excellent resolution between the minor component and the main peak. The 3 mL injection volume with a load of about 125 mg also indicates that this larger mass capacity can be successfully applied to the column with reasonable resolution between the minor component of interest and the main peak. While a load of 166 mg can be injected on the preparative column, the loss of resolution between the peaks of interest can reduce the purity of the isolated peaks. The mass capacity on the BEH Amide column in this study is comparable to the estimated mass capacity on a reversed-phase column with the same dimensions.

Figure 4. ELS detection of stevia sweetener loaded in increasing amounts on a BEH Amide, 19 x 150 mm, 5 μm column. Good resolution between the minor component and the main peak permit higher sample loading.

Scaling separations requires matching column chemistry as well as appropriately scaled gradients. A properly-scaled separation gives the same chromatographic profile at increased mass capacity as the chromatographic profile at the smaller scale. Figure 5 illustrates the BEH Amide column scalability by comparing the chromatography using the small-scale gradient at maximum load with the large-scale chromatography used for the isolation. Note the baseline resolution between the minor component eluting at about 4 minutes and the main peak.

Figure 5. Stevia sweetener at the maximum load on a 4.6 x 150 mm column (top) compared with the preparative run at the maximum load on a 19 x 150 mm column. The chromatographic profile is the same at both scales.
Conclusion

- The isolation of polar compounds on BEH Amide columns is easier because hydrophilic components are retained longer than on reversed-phase packings allowing separation to occur.
- The BEH Amide bonded phase is more compatible with slightly higher aqueous content in mobile phases and sample diluents than unbonded HILIC stationary phases, making the BEH Amide column a more viable alternative for the purification of hydrophilic compounds such as sugars and peptides.
- Improved mass loading of polar compounds on BEH Amide columns reduces the number of injections required to isolate the product, promoting process efficiency.
- Reproducible chromatograms at the analytical and preparative scales reduce ambiguity in the identification of target compounds and demonstrate BEH Amide column scalability.

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