The Separation of Δ⁸-THC, Δ⁹-THC, and Their Enantiomers by UPC² Using Trefoil Chiral Columns

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APPLICATION BENEFITS

■ The stereoisomers of Δ⁸ and Δ⁹-THC can be fully resolved more quickly using convergence chromatography on the Waters® UPC² System than by using HPLC.

■ The three Waters Trefoil™ chiral stationary phases exhibit orthogonal selectivity for the stereoisomers of THC. Providing an advantage when resolving the stereoisomers from matrix interferences.

■ The ACQUITY UPC² separation is repeatable and can be used for qualitative and quantitative analysis of the THC stereoisomers from natural and synthetic cannabis products.

INTRODUCTION

Cannabinoids are chemicals found in Cannabis sativa L (Cannabis) that have been shown to exhibit various pharmacokinetic activities. Specifically, Tetrahydrocannabinol (THC) is considered the key psychoactive cannabinoid present in cannabis.¹ Depending on how it is derived, THC exists in many isomeric forms, four are predominant, and include (+)trans-Δ⁸-THC, (−)trans-Δ⁸-THC, (+)trans-Δ⁹-THC, and (−)trans-Δ⁹-THC (structures in Figure 1). The main plant-derived stereoisomer is (−)trans-Δ⁹-THC, while many synthetic preparations of THC produce the more stable Δ⁸-THC isomer, or a mixture of positional and stereo isomers. Finally, cannabidiol (another major active cannabinoid) isomerizes under acidic conditions to form mixtures that contain THC isomers and other cannabinoid related compounds.²,³ Just as with other pharmaceutically active ingredients, the stereochemistry of THC affects pharmacological activity.

Analytical procedures capable of separating the positional and stereoisomers of THC are required in many applications.¹ For active chiral pharmaceutical compounds, such as THC, the FDA requires that stereoisomeric composition be quantified.⁴ This will be important as the use of cannabis for medicinal purposes becomes legal in many states. Currently, however, cannabis and many of the constituents derived from cannabis (including THC) are classified as Schedule I illicit drugs. Therefore, the separation of THC stereoisomers can be useful not only for pharmaceutical stereoisomeric composition, but also for analysis of seized drugs in forensics, where achiral analysis alone does not provide a comprehensive sample profile.⁵ Where legalized, many consumables are being infused with cannabis derived products. Monitoring the stability of products that contain THC or CBD is important, in case new mixtures of positional and stereoisomers form that result in changes in potency, pharmacological activity, or toxicity.
To that end, the separation of the $\Delta^8$-THC and $\Delta^9$-THC positional isomers, as well as their respective stereoisomers, was investigated using Waters ACQUITY UPC² technology and Trefoil chiral columns. Once an optimized UPC² method was established, calibration and repeatability was performed for each enantiomer. The method was used to determine the THC composition of a commercially available CBD product, before and after undergoing acidic conversion to THC.

![Figure 1. Structures of THC positional and stereoisomers.](image)

**EXPERIMENTAL**

Federally exempt standards were obtained from Cerilliant including: $(\pm)\Delta^8$-THC at 0.1 mg/ml in heptane, $(\pm)\Delta^9$-THC at 0.1 mg/ml in heptane, $(\pm)\Delta^8$-THC at 1 mg/ml in methanol, and $(\pm)\Delta^9$-THC at 1 mg/ml in methanol. For initial method development, a mixture of all four isomers was used. The mixed sample was a 50:50 mix of the $(\pm)\Delta^8$-THC and the $(\pm)\Delta^9$-THC standards resulting in 0.05 mg/ml solution (0.0125 mg/ml each isomer). The $(\pm)\Delta^8$-THC and $(\pm)\Delta^9$-THC samples were diluted to 0.1 mg/ml in ethanol and were used to determine peak order. Serial dilutions of the $(\pm)\Delta^8$-THC and the $(\pm)\Delta^9$-THC standards were used to create calibration curves for the four stereoisomers.

A commercially available CBD oil extract was obtained and used as an example product. Three aliquots of the CBD oil samples were made up using the following conditions:

- **Sample 1:** 15.65 mg of CBD Oil in 3 mL 200-proof ethanol, heated overnight at 55 °C
- **Sample 2:** 15.26 mg of CBD Oil in 3 mL 0.1 M HCl in 200-proof ethanol, heated overnight at 55 °C
- **Sample 3:** 15.40 mg of CBD Oil in 3 mL 200-proof ethanol, room temperature

All three samples were filtered and diluted 1:10 in ethanol before injecting on the UPC²

**UPC² screening conditions**

| System: | ACQUITY UPC² System with an ACQUITY PDA Detector |
| Columns: | ACQUITY UPC² Trefoil AMY1 Column, 2.5 µm, 3 mm x 150 mm |
| | ACQUITY UPC² Trefoil CEL1 Column, 2.5 µm, 3 mm x 150 mm |
| | ACQUITY UPC² Trefoil CEL2 Column, 2.5 µm, 3 mm x 150 mm |
| Mobile phase A: | Carbon Dioxide |
| Mobile phase B: | 200-proof ethanol |
| Gradient: | 2 to 20% B over 5 minutes |
| Column temperature: | 50 °C |
| Injection volume: | 1 µL |
| Flow rate: | 2 mL/min |
| ABPR: | 2000 psi |
| PDA absorbance: | 228 nm, compensation reference: 500–600 nm |
| Optimized conditions: | noted on figures |
RESULTS AND DISCUSSION

In normal phase chromatography, and SFC in particular, the first step in method development involves column screening. In this case, the columns used in the screening were amylose (AMY1) and cellulose (CEL1 and CEL2) based chiral stationary phases, which are commonly used because of their wide range of applicability. In particular, the three Trefoil columns provide orthogonal selectivity, which is optimal for successful chiral method development. Much of the work involving cannabis quality control and post-extraction processing uses ethanol as the preferred solvent in order to avoid toxicity from solvent contamination in any final product or consumable. Consequently, ethanol was chosen as the sample diluent and co-solvent for method development. As is typical in column screening, a generic gradient from 2% to 20% ethanol was utilized (Figure 2). It was observed, that all three columns showed good separation of the four THC isomers. As a result, all three separations could be optimized for qualitative or quantitative analysis of the THC isomers. The AMY1 column was chosen for method development because it showed high resolution and more retention of these compounds, resulting in more room for optimization of the separation than with the two cellulose (CEL) columns.

![Figure 2. Screening of the four isomers of THC on three Trefoil chiral columns using a 2–20% gradient over 5 minutes.](image)

Optimization of the mobile phase conditions usually involves either focusing the gradient or determining isocratic conditions. Isocratic methods are ideal because they are easy to develop and no equilibration is required between runs, improving productivity. Using retention times and gradient slope, and compensating for system and column volume delay, the co-solvent percentage at elution was determined for the separation on the AMY1 column.
With a gradient delay of 0.34 min, gradient slope of 3.6%/min, and 2% starting percentage, the co-solvent percentage at elution of the first peak at 4.15 was calculated using the following equation:

\[
\text{%Co-solvent at elution} = (\text{retention time} - \text{gradient delay}) \times \text{gradient slope} + \text{starting %}
\]

\[
\text{%Co-solvent at elution} = (4.15 - 0.34 \text{ min}) \times 3.6\%/\text{min} + 2\%
\]

\[
\text{% Co-solvent at elution} = 15.7\%
\]

In SFC, usually the best starting point for optimization is 5% below the calculated percentage. Therefore, the initial isocratic method was run at 10% ethanol. Figure 3 shows separation of the THC isomers at 10% and 15% ethanol conditions. At 15%, the separation of the positional and stereo isomers of THC is achievable in less than 3 minutes. Comparatively, the liquid chromatography separation of these compounds does not result in baseline resolution of all four enantiomers, even with a relatively long 23 minute run time.³

In this case, the separation was going to be used to analyze THC in a complex matrix containing cannabidiol (CBD). As a result, the methods were investigated to ensure separation of the THC isomers from CBD. The 10% method showed acceptable resolution from the CBD and matrix, and was subsequently used for calibration and repeatability. Figure 4 shows calibration curves at concentrations of 0.003125, 0.00625, 0.0125, 0.025, and 0.05 mg/mL for all four THC isomers on the AMY1 column at 10% ethanol conditions. The 0.025 mg/mL standard was used to determine repeatability. The calibration curves are linear, with \( R^2 \) values greater than 0.998, and the peak areas show good reproducibility, with area count RSD values less than 2% (n=7).
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Depending on the strain of cannabis used in the production of CBD oil, the resulting product can contain varying amounts of other cannabinoids. THC content is of particular interest for purposes of product quality control. In this case, the product was a CBD oil extract that was marketed as “CBD only,” indicating there is no THC present in the untreated sample. Once the method was calibrated, and shown to be reproducible, the separation was applied to the analysis of the CBD oil extract. Three samples of the product were analyzed; a heated sample, a sample exposed to heat and acidic conditions, and a third control sample that was left untreated. Comparative analysis of the three samples against the THC standards can be seen in Figure 5.

Figure 4. Calibration curves for the four THC isomers on the AMY1 column at 10% isocratic conditions. The calibration curve concentrations were 0.003125, 0.00625, 0.0125, 0.025, and 0.05 mg/mL, and the injection volume was 1 µL.

Figure 5. UPC2 chromatograms showing (1) separation of the THC isomer standards, (2) analysis of CBD oil after being heated in an oven overnight at 55 °C, (3) analysis of CBD oil after being exposed to acidic conditions and heating overnight at 55 °C, and (4) analysis of untreated CBD oil (control). THC isomers that were identified in the CBD samples are indicated. The separations were achieved at 2 mL/min and 10% ethanol on the AMY1 column using 1 µL injections.
Analysis of the CBD oil samples showed a very small amount of THC in the initial product (control sample). The heat treated sample showed no significant change in THC content. The sample exposed to acidic conditions contained a significant amount of (−)Δ⁹-THC, calculated to be 1.16mg or approximately 7.6% of the initial 15.26 mg sample. There was also a very small detectable amount of (−)Δ⁸-THC in the sample exposed to acidic conditions. The samples were incubated for one overnight period (approximately 16 hrs), given more time the conversion would likely be even more pronounced.³

CONCLUSIONS

■ The separation of (±)Δ⁸-THC and (±)Δ⁹-THC and their enantiomers can be accomplished in less than 3 minutes on the Trefoil AMY1 column at 15% ethanol mobile phase conditions using the Waters ACQUITY UPC² system.

■ All three Waters Trefoil chiral stationary phases separate the THC isomers, while also exhibiting different elution order between the amylose (AMY) and cellulose (CEL) based columns. This is advantageous when resolving the stereoisomers from matrix interferences.

■ The ACQUITY UPC² separation of the four THC isomers on the Trefoil AMY1 column showed excellent repeatability and the calibration curves showed good linearity. As a result, the methodology can be used for qualitative and quantitative analysis of the Δ⁸-THC and Δ⁹-THC stereoisomers in natural and synthetic cannabinoid products.

References


