

Summary

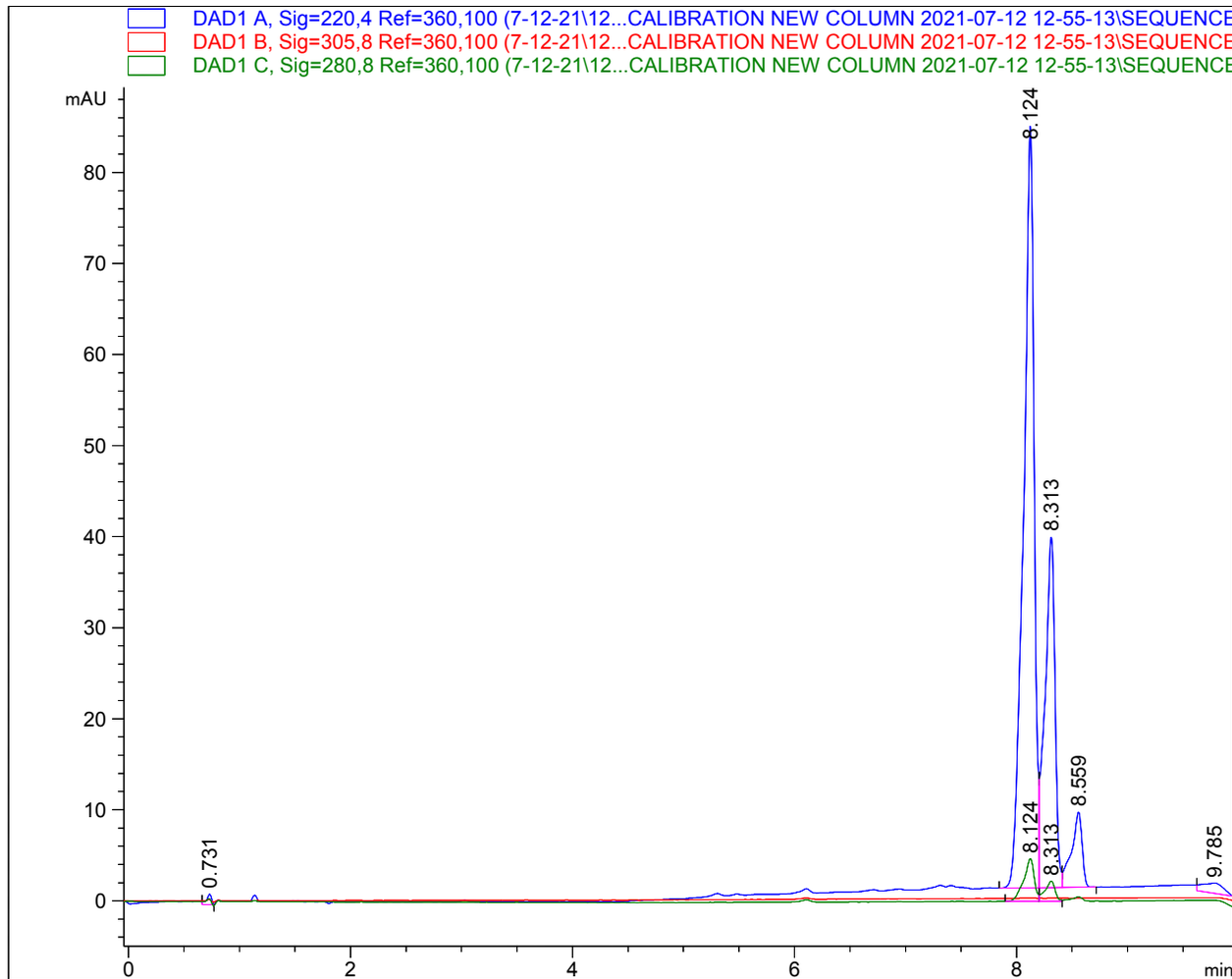
Bearly Legal Hemp has partnered with a leading cGMP certified manufacturer of rare cannabinoids, whom recently accessed Hexahydrocannabinol (HHC) at commercial scale. The manufacturer's HHC concentrate contains three different stereoisomers of HHC and less than 0.2% other cannabinoids. There is currently no Certified Reference Standard commercially available for HHC. The manufacturer is working with various third-party cannabis testing labs including KCA Labs and Green Scientific Labs to purify and isolate the various HHCs and have them turned into Certified Reference Materials for the industry. Once a created Certified Reference Standard, the manufacturer will have exclusive testing rights on HHC for a period of 6 months. We expect to have the standard certified within the next 30 to 45 days.

We have attached the manufacturer's in-house HPLC chromatogram which shows no other cannabinoids detected other than HHC and trace levels of Delta-8 THC. There are three versions of HHC present in our product each of which corresponds to an individual peak on the chromatogram. A COA from Gobi shows that our HHC is truly Farm Bill Compliant, meaning that this product is below 0.3% Total THC. The technician at Gobi also noticed unidentified peaks which they do not have a standard for.

We have a third-party NMR which confirms that the structure of what we made is HHC. We have also sent a sample to KCA labs to verify the identity of these unknown peaks and their Mass Spec data confirms that the sample is HHC. The manufacturer is also working with toxicology and product safety labs to verify the safety of HHC using the latest *in vitro* techniques.

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=====
Acq. Operator : Seq. Line : 3
Acq. Instrument : Instrument 1 Location : Vial 10
Injection Date : 7/12/2021 1:23:54 PM Inj : 1
Inj Volume : 5.0 µl
Acq. Method : C:\CHEM32\1\DATA\7-12-21\12_16_20 CALIBRATION NEW COLUMN 2021-07-12 12-55-13\
1100 DAD HIGH THROUGHPUT (NO CBT).M
Last changed : 7/12/2021 1:03:17 PM
(modified after loading)
Analysis Method : C:\CHEM32\1\METHODS\PERCENT AREA METHOD.M
Last changed : 7/6/2021 7:24:35 PM
(modified after loading)
Method Info : First Runs Shutdown



=====
Area Percent Report
=====

Sorted By : Signal
Calib. Data Modified : 7/6/2021 7:24:35 PM
Multiplier: : 1.0000
Dilution: : 1.0000
Use Multiplier & Dilution Factor with ISTDs

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Signal 1: DAD1 A, Sig=220,4 Ref=360,100

Peak #	RetTime [min]	Type	Width [min]	Area [mAU*s]	Area %	Name
1	2.402		0.0000	0.00000	0.0000	CBDV
2	4.467		0.0000	0.00000	0.0000	CBD
3	4.718		0.0000	0.00000	0.0000	CBG
4	5.470		0.0000	0.00000	0.0000	CBDA
5	7.134		0.0000	0.00000	0.0000	D9 THC
6	7.863		0.0000	0.00000	0.0000	CBN
7	10.329		0.0000	0.00000	0.0000	D8 THC

Totals : 0.00000 0.0000

Uncalibrated Peaks:

Peak #	RetTime [min]	Type	Width [min]	Area [mAU*s]	Area %	Name
1	0.731	VB	0.0485	3.60868	0.4125	?
2	8.124	BV	0.0936	556.16852	63.5785	?
3	8.313	VV	0.0920	244.60983	27.9626	?
4	8.559	VB	0.0962	55.31372	6.3232	?
5	9.785	VBA	0.1801	15.07346	1.7231	?

Uncalib. totals : 874.77422 100.0000

Signal 2: DAD1 B, Sig=305,8 Ref=360,100

Uncalibrated Peaks:

Signal 3: DAD1 C, Sig=280,8 Ref=360,100

Uncalibrated Peaks:

Peak #	RetTime [min]	Type	Width [min]	Area [mAU*s]	Area %	Name
1	8.124	BV	0.0933	31.06334	68.8874	?
2	8.313	VV	0.0919	14.02956	31.1126	?

Uncalib. totals : 45.09291 100.0000

1 Warnings or Errors :

Warning : Calibrated compound(s) not found

*** End of Report ***

Analytical Report - Certificate of Analysis

Manifest: 2107090002
Sample Id: 1A-GHEMP-2107090002-0001
Sample Name: D10 ML
Sample Type: Concentrate
Client Id: CID-00120

Test Performed: Hemp Lab
Report No: P-2107090002-V1
Receive Date: 2021-07-09
Test Date: 2021-07-09
Report Date: 2021-07-09
Sample Condition: Good
Method Reference: GH-OP-06

Scope

The content of sixteen cannabinoids was determined by an in-house developed method for solvent extraction followed by High Performance Liquid Chromatography with Diode Array Detection.

Cannabinoids	Percent	mg/gram
CBDV	ND	ND
CBDA	ND	ND
CBGA	ND	ND
CBG	ND	ND
CBD	ND	ND
THCV	ND	ND
CBN	0.13	1.26
Δ9-THC	ND	ND
CBC	ND	ND
THCA	ND	ND
CBDVA	ND	ND
THCVA	ND	ND
CBNA	ND	ND
Δ8-THC	0.12	1.19
CBL	ND	ND
CBCA	ND	ND

ND - not detected; T - trace; ULOQ - limit of quantitation

	Percent	mg/gram
Total Δ9-THC	0.00	0.00
Total CBD	0.00	0.00
Total CBG	0.00	0.00
Total Cannabinoids	0.24	2.45

Total Δ9-THC = Δ9-THC + (THCA x 0.877)
 Total CBD = CBD + (CBDA x 0.877)
 Total CBG = CBG + (CBGA x 0.877)

Laboratory Comments:

Presence of two large peaks at 6.043 min and 6.249 min that are unidentified by our standards.



Jerry Hogan - Director of Operations

2021-07-09

Date

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Gobi Hemp
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 • Wheat Ridge CO 80033 •
 • ISO/IEC 17025:2017 Accredited •
 • (303) 955-4934 •



HHC

Sample ID: 2105KCA0856.2132

Cultivar: N/A

Matrix: Concentrates & Extracts

Type: Distillate

Sample Size:

Received: 05/17/2021

Completed: 05/27/2021

Batch#:



Summary

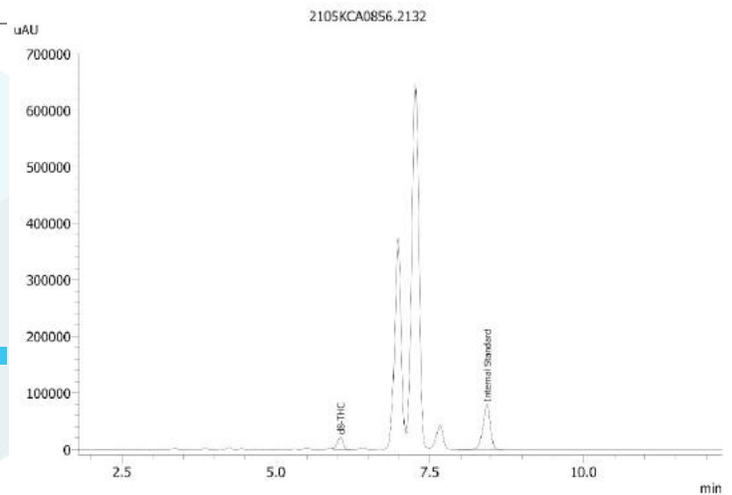
Test	Date Tested	Result
Cannabinoids	05/27/2021	Complete

Cannabinoids by HPLC-PDA

Complete

ND Total THC	ND Total CBD	1.5359% Total Cannabinoids	NT Not Tested Moisture Content	Not Tested Foreign Matter
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Analyte	LOD %	LOQ %	Result %	Result mg/g
CBC	0.0095	0.0284	ND	ND
CBCA	0.0181	0.0543	ND	ND
CBCV	0.0060	0.0180	ND	ND
CBD	0.0081	0.0242	ND	ND
CBDA	0.0043	0.0130	ND	ND
CBDV	0.0061	0.0182	ND	ND
CBDVA	0.0021	0.0063	ND	ND
CBG	0.0057	0.0172	ND	ND
CBGA	0.0049	0.0147	ND	ND
CBL	0.0112	0.0335	ND	ND
CBLA	0.0124	0.0371	ND	ND
CBN	0.0056	0.0169	ND	ND
CBNA	0.0060	0.0181	ND	ND
Δ8-THC	0.0104	0.0312	1.5359	15.359
Δ9-THC	0.0076	0.0227	ND	ND
THCA	0.0084	0.0251	ND	ND
THCV	0.0069	0.0206	ND	ND
THCVA	0.0062	0.0186	ND	ND
Total THC			ND	ND
Total CBD			ND	ND
Total			1.5359	15.359



Total THC = THCA * 0.877 + Δ9-THC

Total CBD = CBDA * 0.877 + CBD

LOD = Limit of Detection

LOQ = Limit of Quantitation

ND = None Detected

For plant material, the reported result is based on a sample weight with the applicable moisture content for that sample.



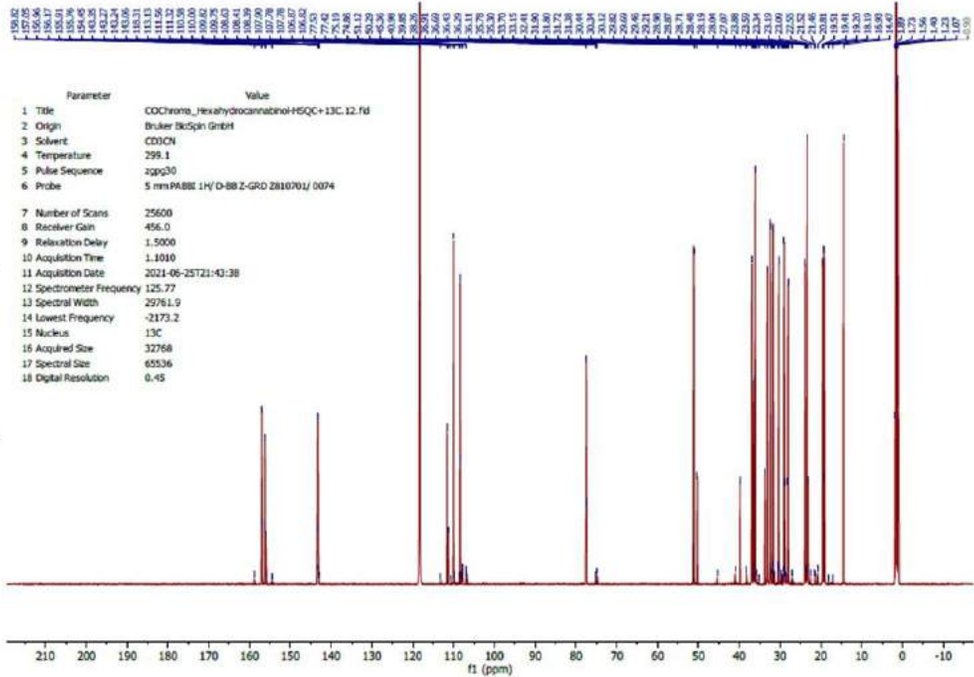
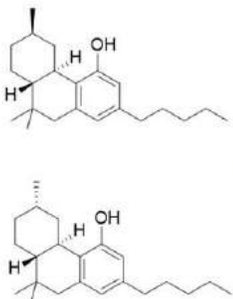
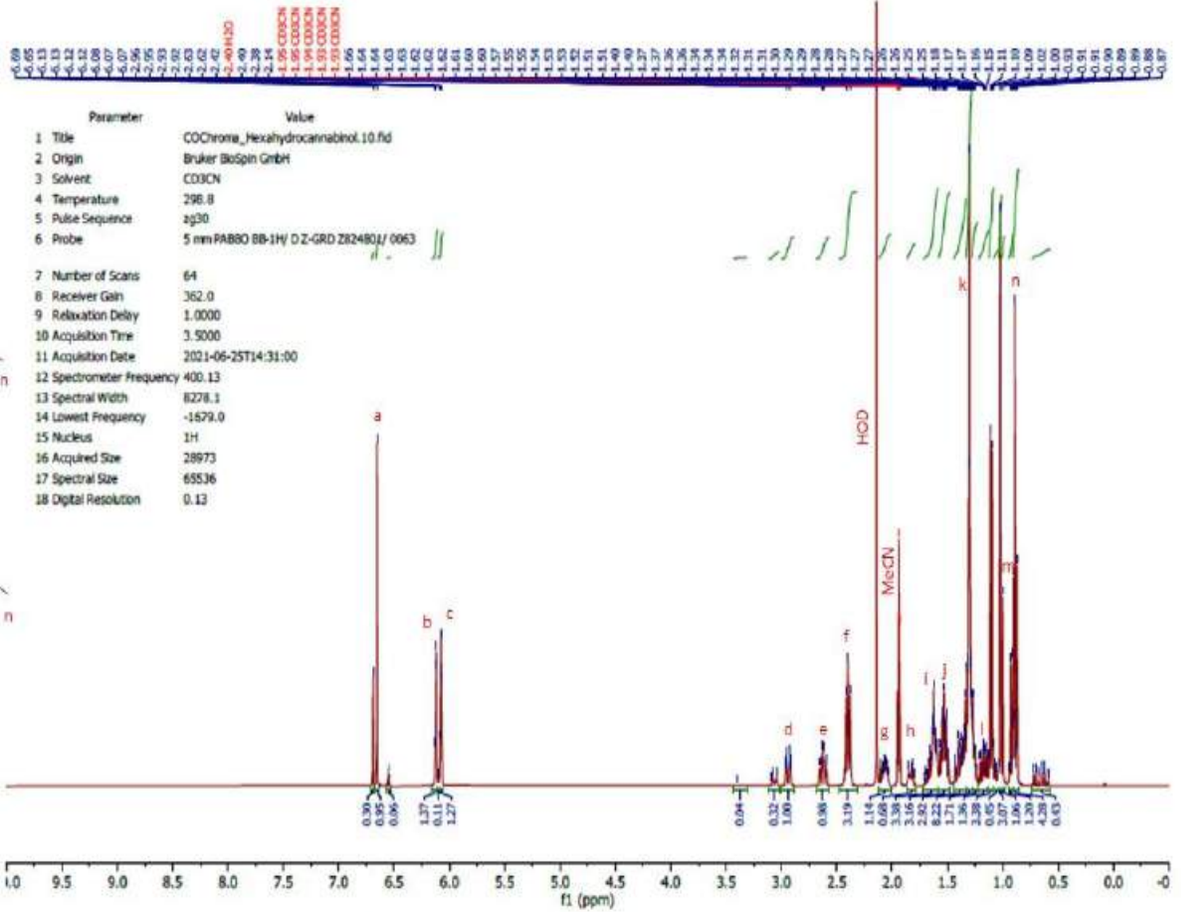
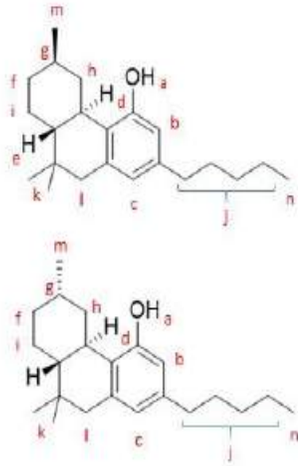
Wes Rogers

Wes Rogers
 Principal Scientist
 05/27/2021



ISO/IEC 17025:2017 Accredited
 Accreditation #108651

1H NMR and 13C NMR for HHC

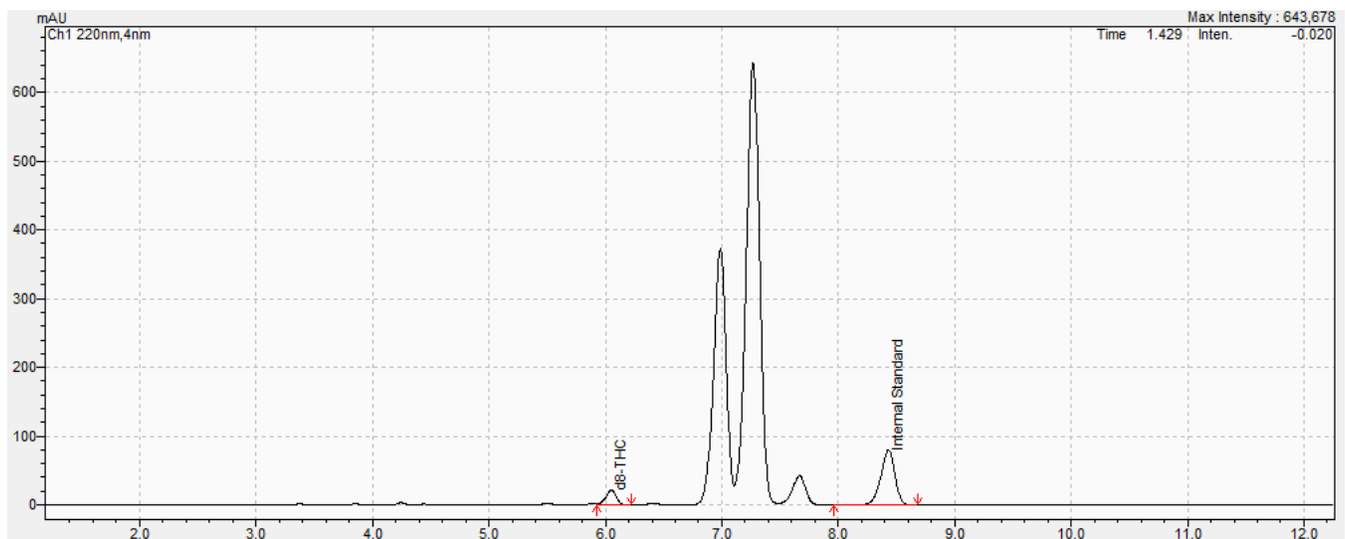


Report 28 May 2021

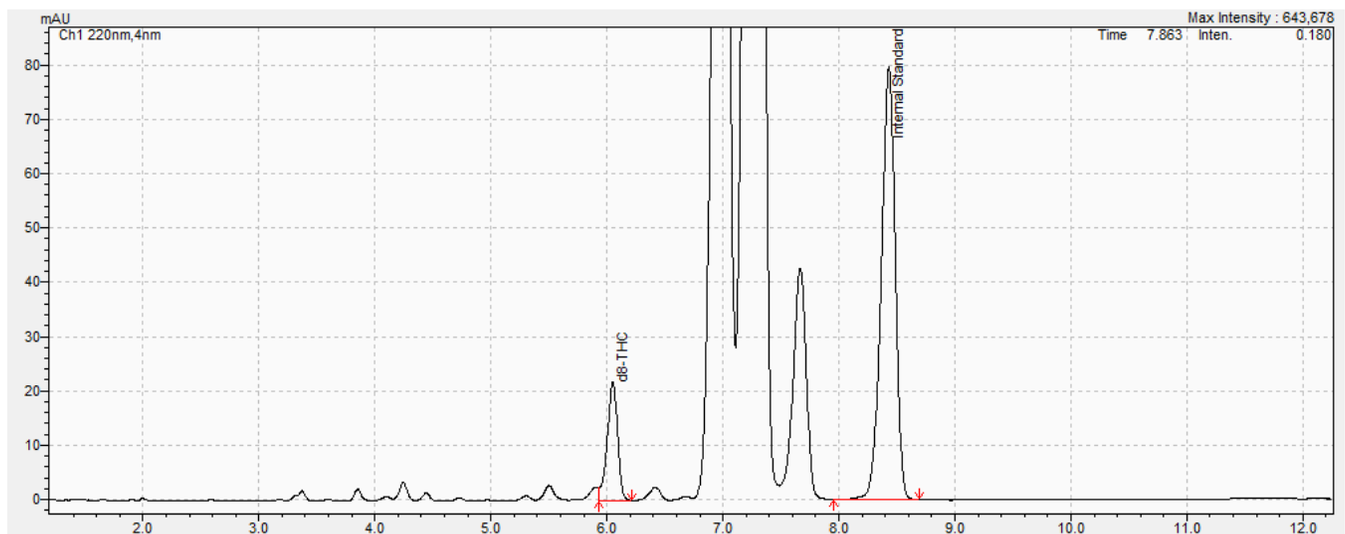
HPLC-PDA Data Analysis

Samples were initially analyzed by HPLC-PDA to identify and quantify cannabinoids in the mixture of 18 standard cannabinoids plus additional cannabinoids (*i.e.*, CBND, CBT, CBE, Δ^10 -THC, $\Delta^{6a,10a}$ -THC etc.). Cannabinoids were identified by comparison of their relative retention times (RRT) and the UV-spectral data. Samples were analyzed twice – once at a concentrated level to detect and quantify minor cannabinoids and once after dilution to detect and quantify major cannabinoids).

Sample ...2132

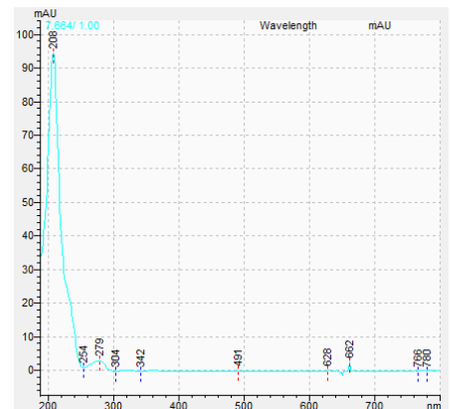
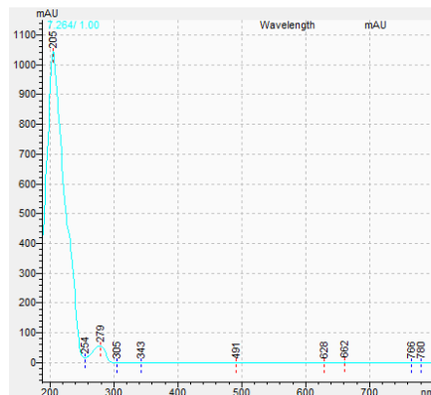
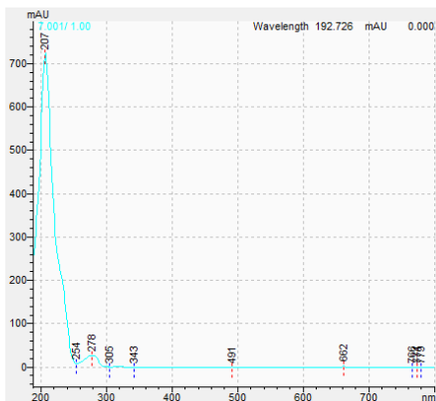
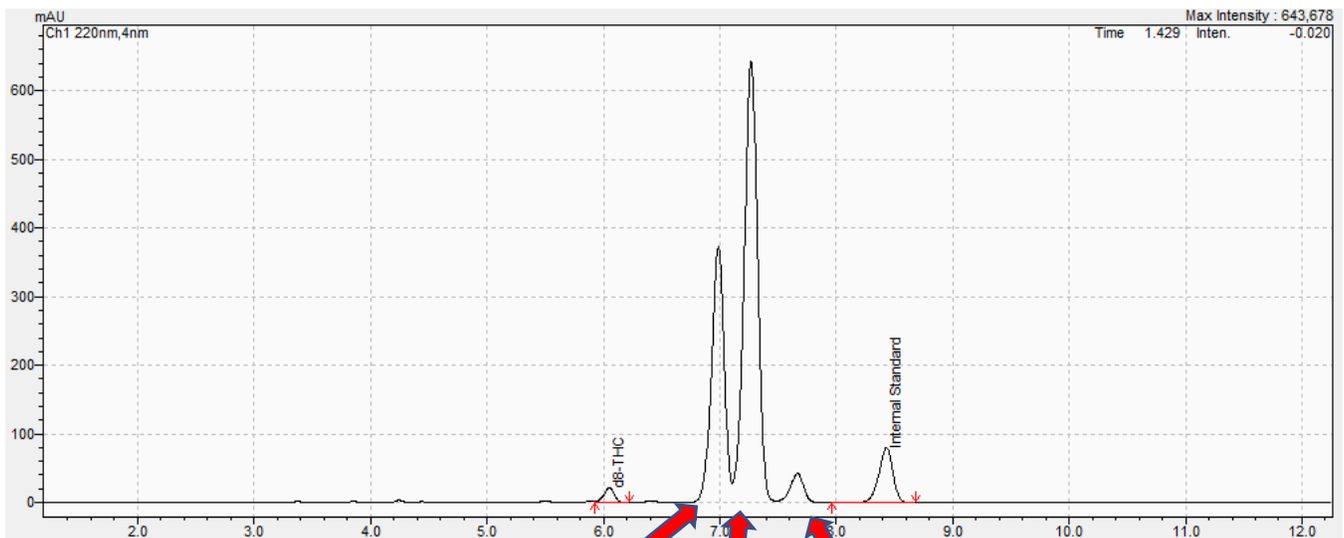


Sample ...2132 (Zoomed In)



Δ^8 -THC was the only cannabinoid identified from the reference standards analyzed by this method. The spectra of unknown peaks are displayed below. At least three unidentified substances eluted after Δ^8 -THC indicating that they are more lipophilic than Δ^8 -THC. If the molar absorptivities of these substances are similar to that of Δ^8 -THC, then at least two of them are much more abundant than that of Δ^8 -THC and the third at a retention time of about 7.7 min is about twice the abundance of Δ^8 -THC.

Origins of PDA Spectral Data



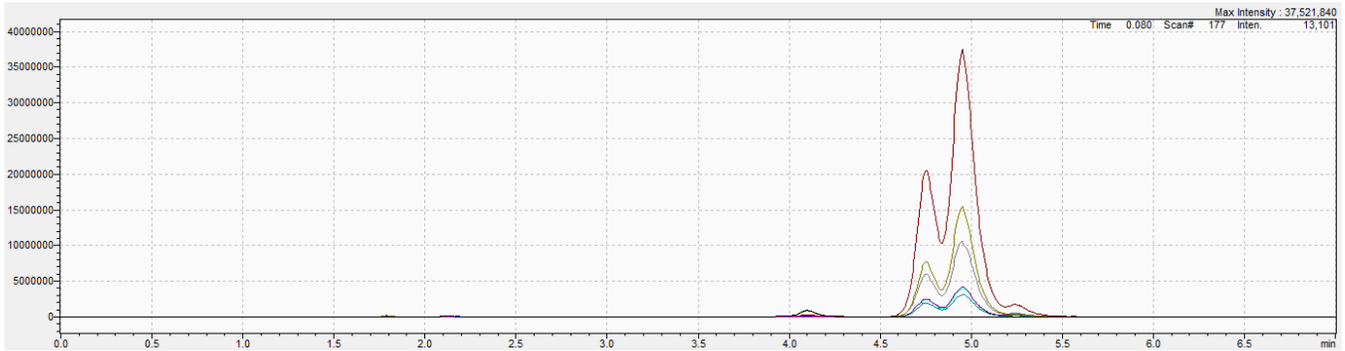
The PDA spectral data for these peaks appear similar to those of other cannabinoids such as Δ^8 -THC, Δ^9 -THC, and CBD without extended conjugation.

Since the major components of the sample were not identified by HPLC-PDA, we subjected the sample to LC-MS/MS analysis as shown on the following pages.

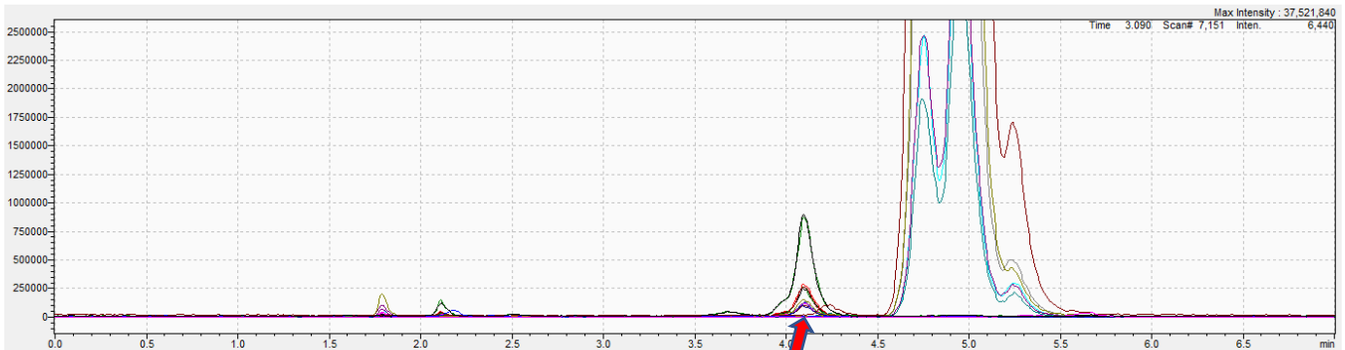
LC-MS/MS Data Analysis

Full Scan Data

Sample ...2132



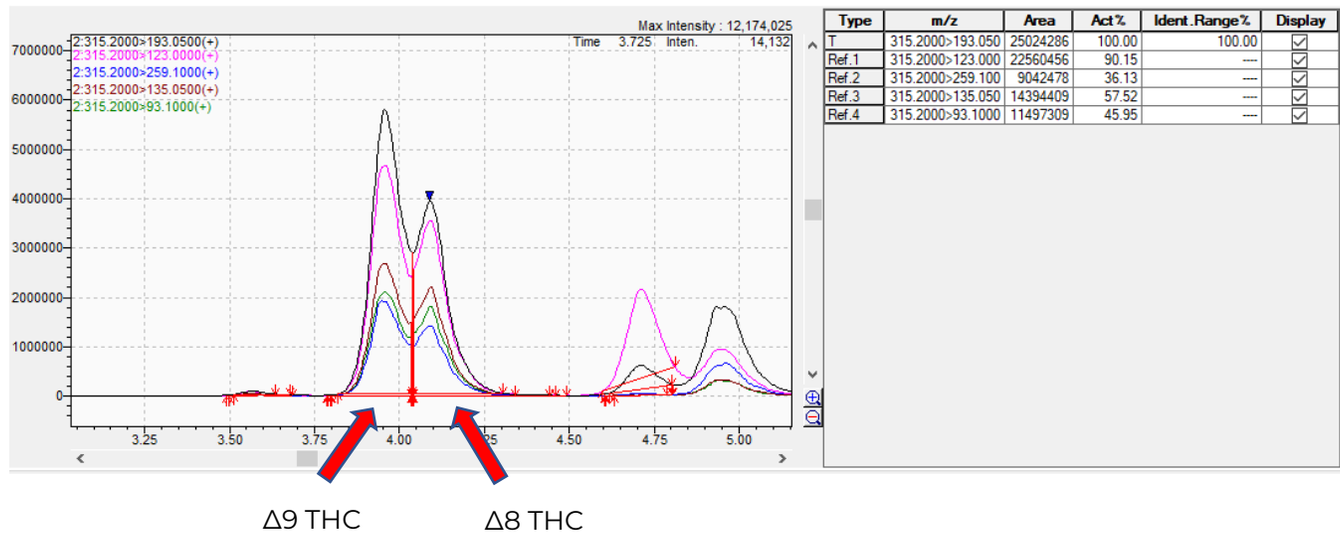
Sample ...2132 (Zoomed In)



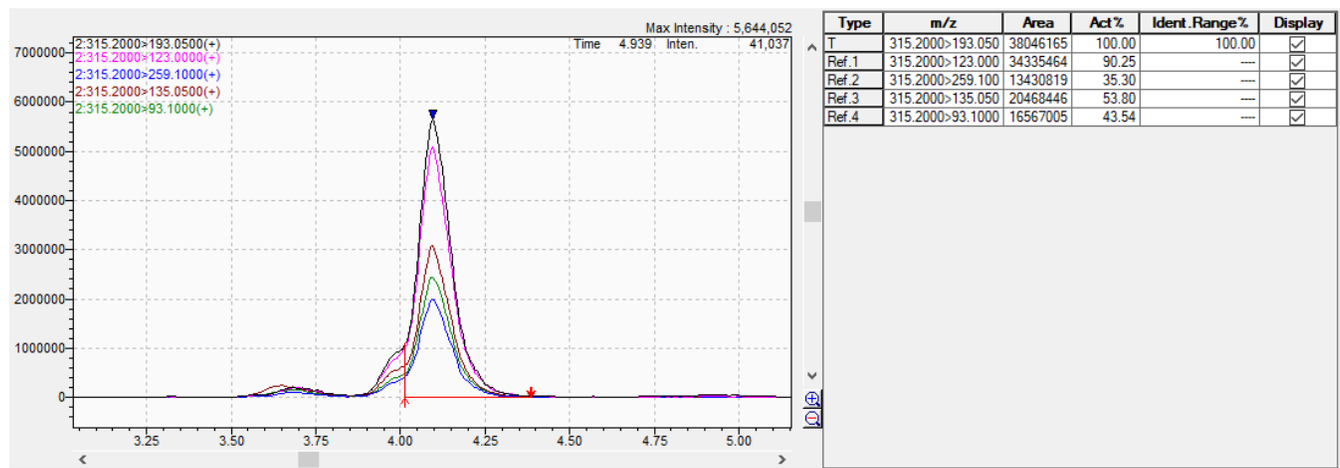
$\Delta 8$ THC

Data were then acquired in MRM (multiple reaction monitoring mode). The MRM transitions are listed to the right of the chromatogram. The “Act%” is the relative abundance (RA) of each MRM transition compared to the primary transition (calculated by area counts). To identify a compound, the relative abundances (RA) of the MRM transitions of the peak are compared to those of a reference standard.

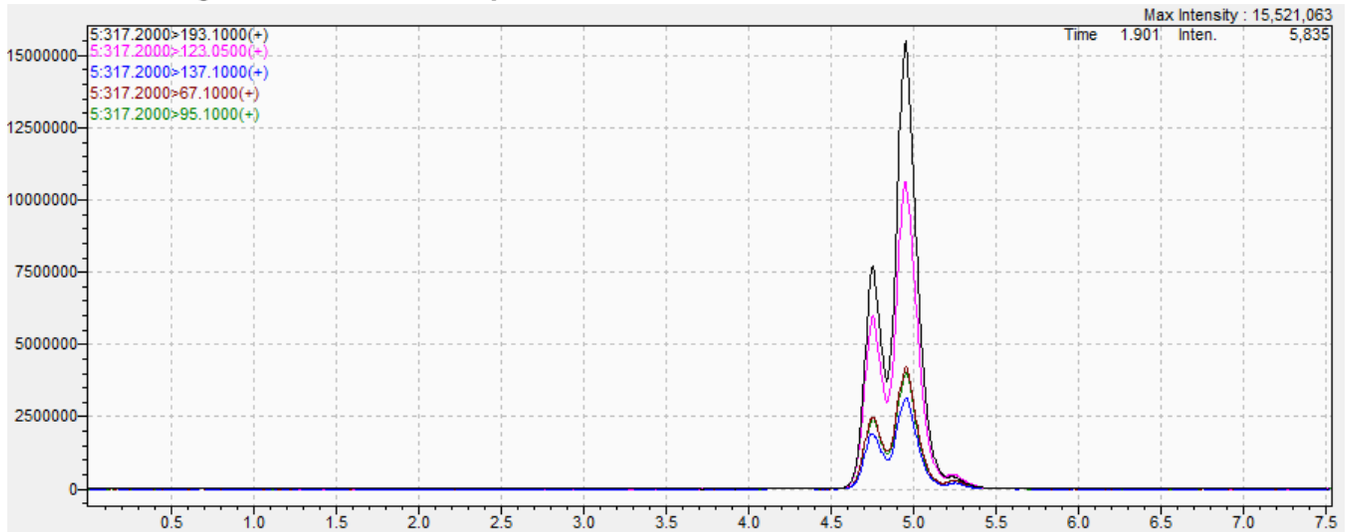
Calibration Mix containing cannabinoid standards.



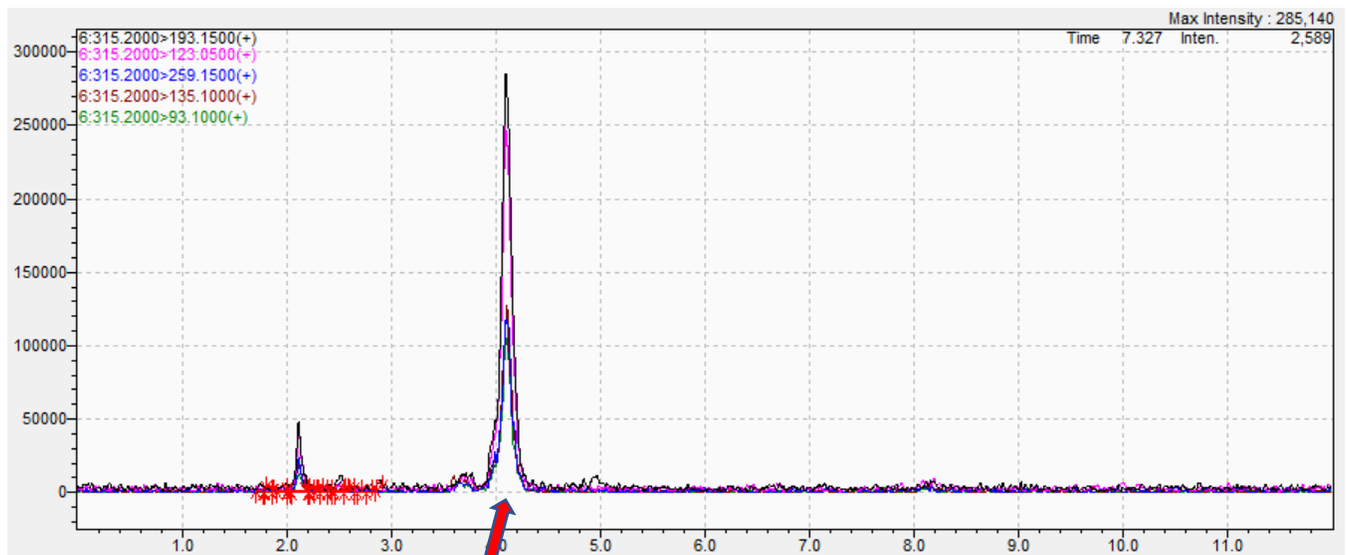
Sample ...2132



The presence of Δ^B-THC was confirmed by the ions and ion area ratios of the peak at 4.10 minutes in sample #2132. The peak just to the left of the peak for Δ^B-THC has ions characteristic of Δ⁹-THC but we do not believe that identification criteria were met for this identification.

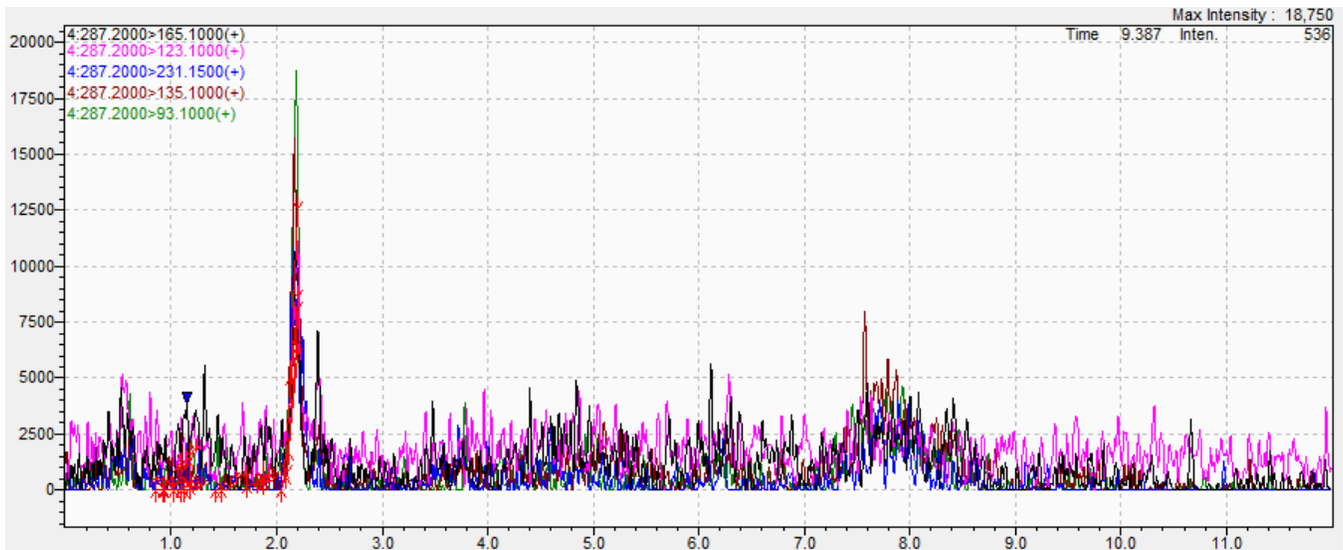
Ion chromatogram for m/z 317.2 in positive ionization mode

These peaks have masses and fragmentation patterns similar to those of CBG but the retention times are incorrect for CBG. The apparent molecular mass of the three substances eluting between 4.5 and 5.3 minutes is 316. This mass is identical to that of various isomers of hexahydrocannabinol.

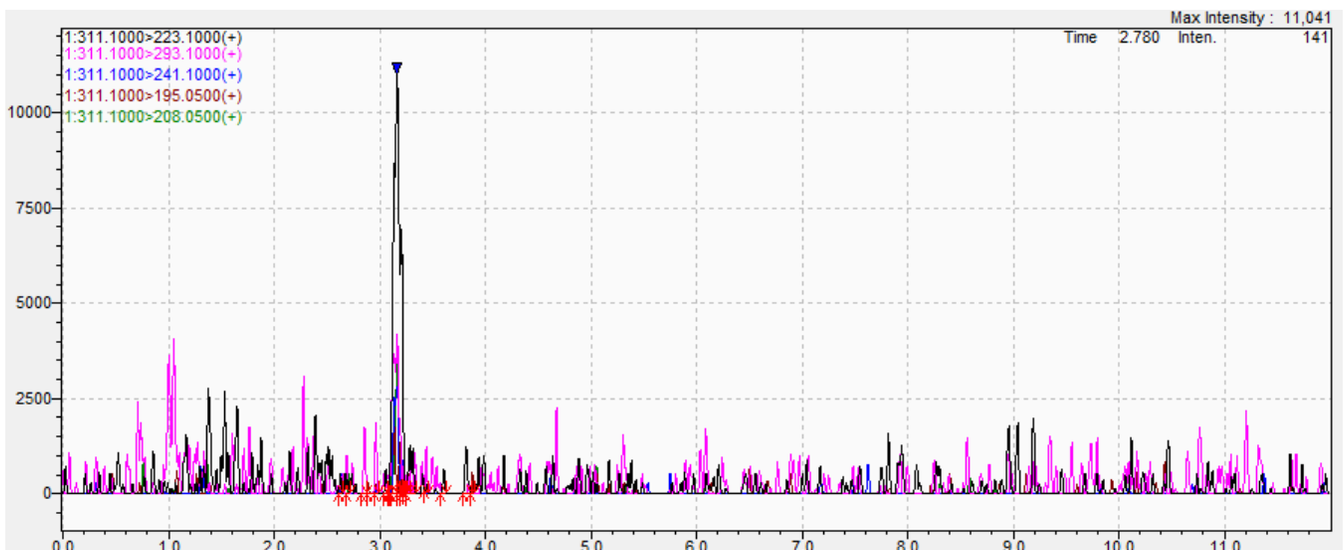
Ion chromatogram for m/z 315.2 in positive ionization mode

Δ^8 -THC

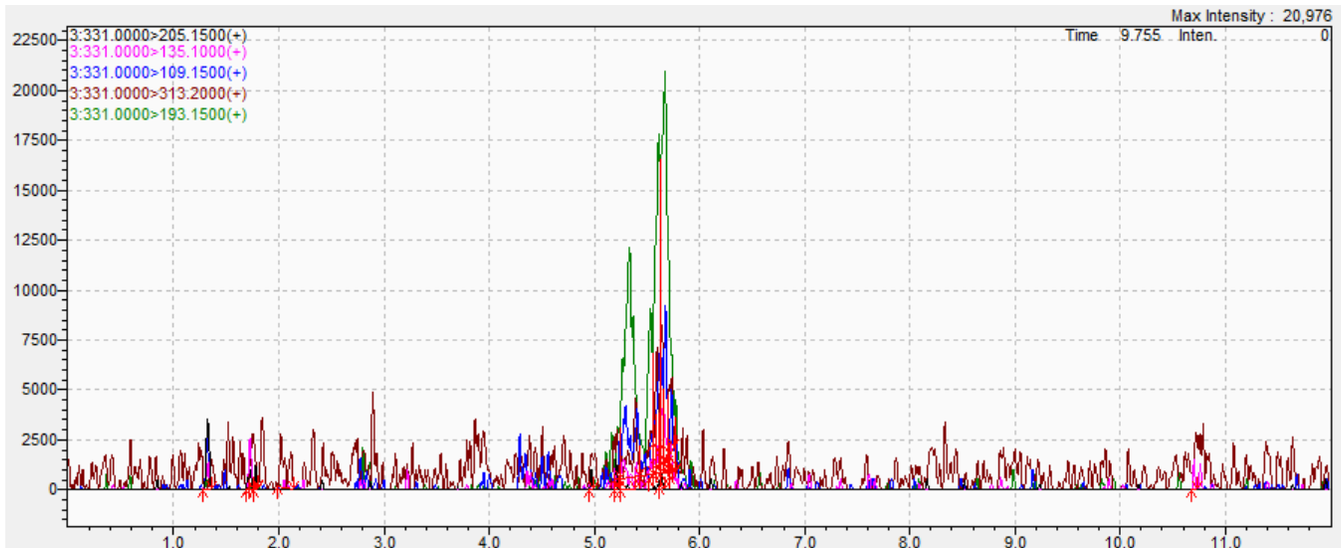
The identification of Δ^8 -THC was confirmed.

Ion chromatogram for m/z 287.2 in positive ionization mode

Nothing was identified in this ion chromatogram. The ions are of low intensity and probably do not represent substantial products.

Ion chromatogram for m/z 311.1 in positive ionization mode

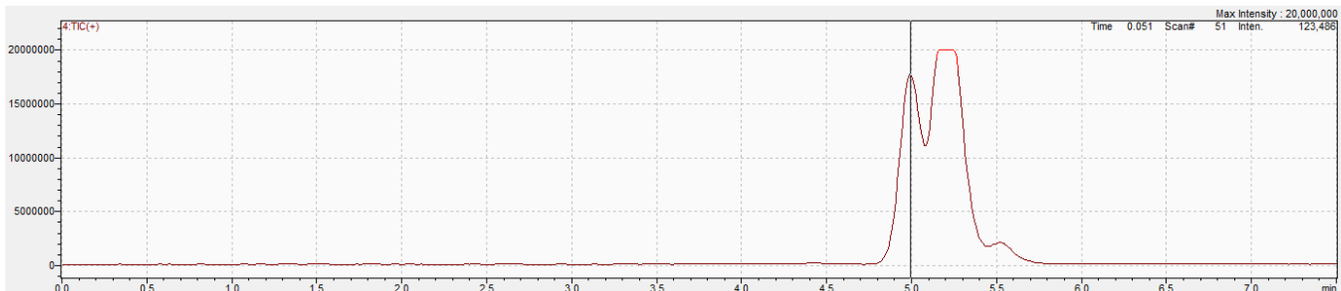
Nothing was identified in this ion chromatogram. The ions are of low intensity and probably do not represent substantial products.

Ion chromatogram for m/z 331.0 in positive ionization mode

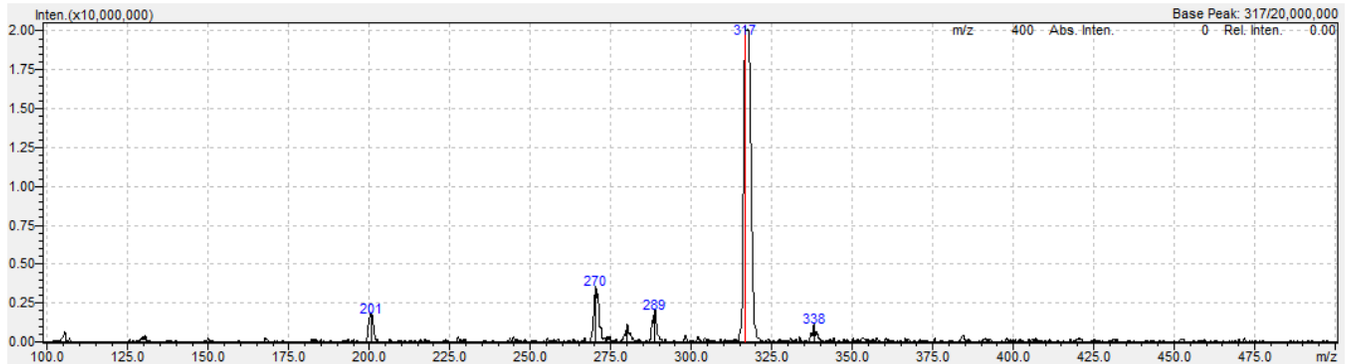
Nothing was identified in this ion chromatogram. The ions are of low intensity and probably do not represent substantial products.

The selected ion monitoring data for the ion at m/z 317 were collected and are displayed in the following ion chromatograms. The vertical line in the ion chromatogram indicates the time at which the data displayed in the second window were obtained.

The Selected Ion Monitoring (SIM) results for m/z 317 in positive ionization mode are shown below. The data at 5.0 minutes were selected for analysis.



The prominent peak at 5.0 min is characterized by m/z 317 and the full scan spectrum is characterized by ions at m/z 289, 270, and 201.

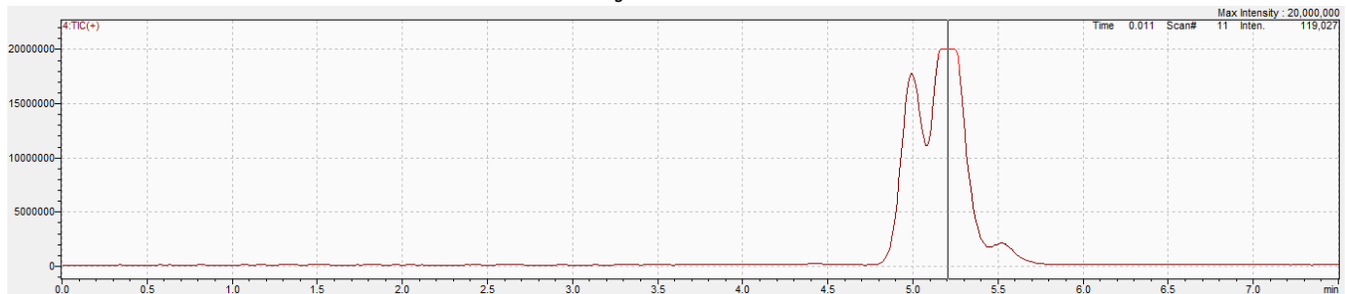


The product ion scan for m/z 317 at 5.0 min is characterized by ions at m/z 193 and 123.

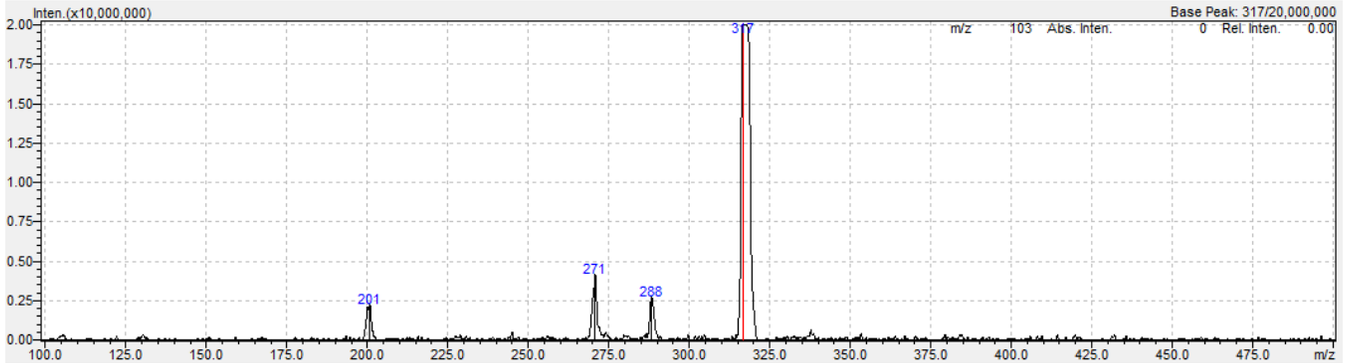


SIM of m/z 317

The Selected Ion Monitoring (SIM) results for m/z 317 in positive ionization mode are shown below. The data at 5.2 minutes were selected for analysis.



The prominent peak at 5.2 min is characterized by m/z 317 and the full scan spectrum is characterized by ions at m/z 2898 271, and 201.

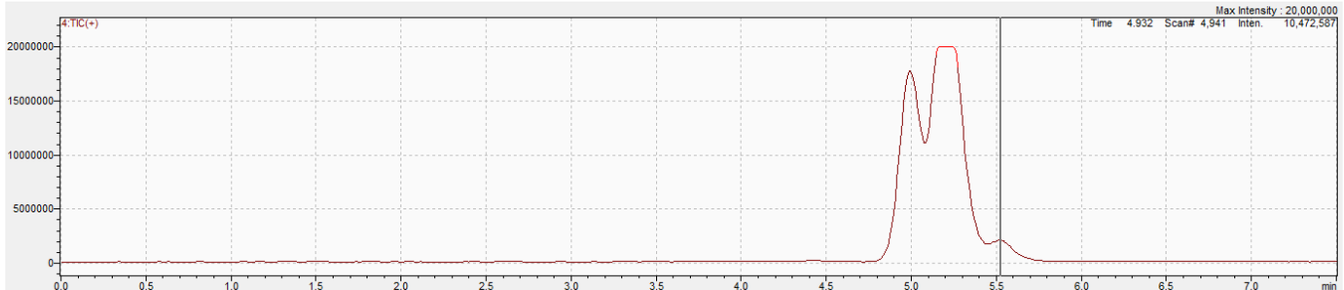


The product ion scan for m/z 317 at 5.2 min is characterized by fragment ions at m/z 194 and 123.

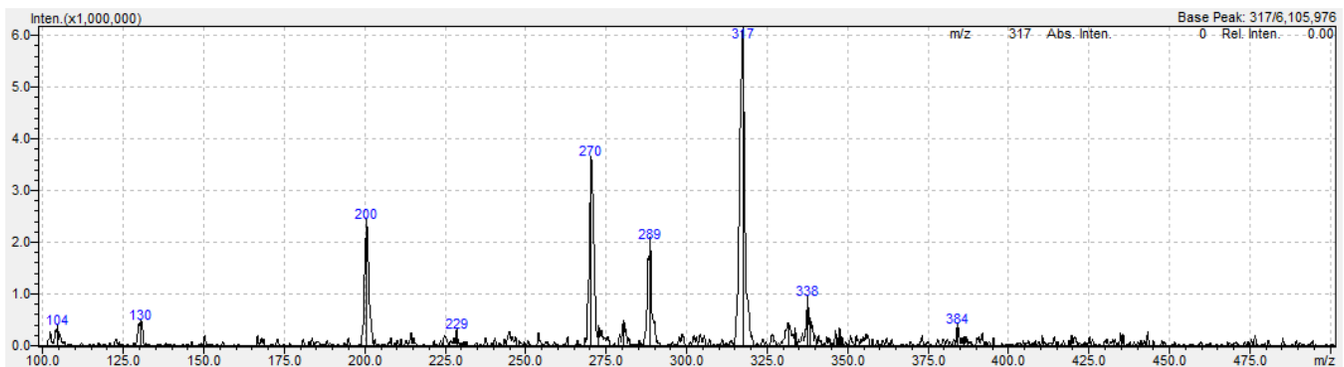


SIM of m/z 317

The Selected Ion Monitoring (SIM) results for m/z 317 in positive ionization mode are shown below. The data at 5.5 minutes were selected for analysis.



The prominent peak at 5.5 min is characterized by m/z 317 and the full scan spectrum is characterized by ions at m/z 289, 270, and 200.



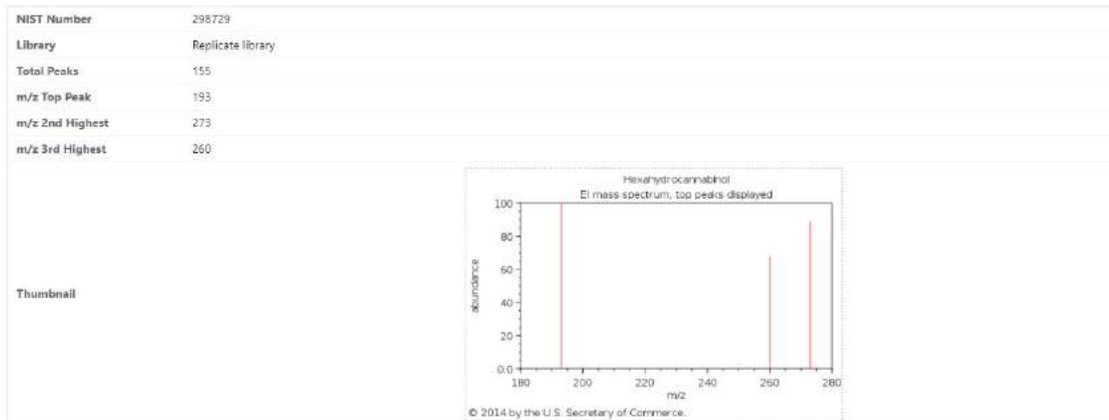
The product ion scan for m/z 317 at 5.5 min is characterized by fragment ions at m/z 194 and 124.



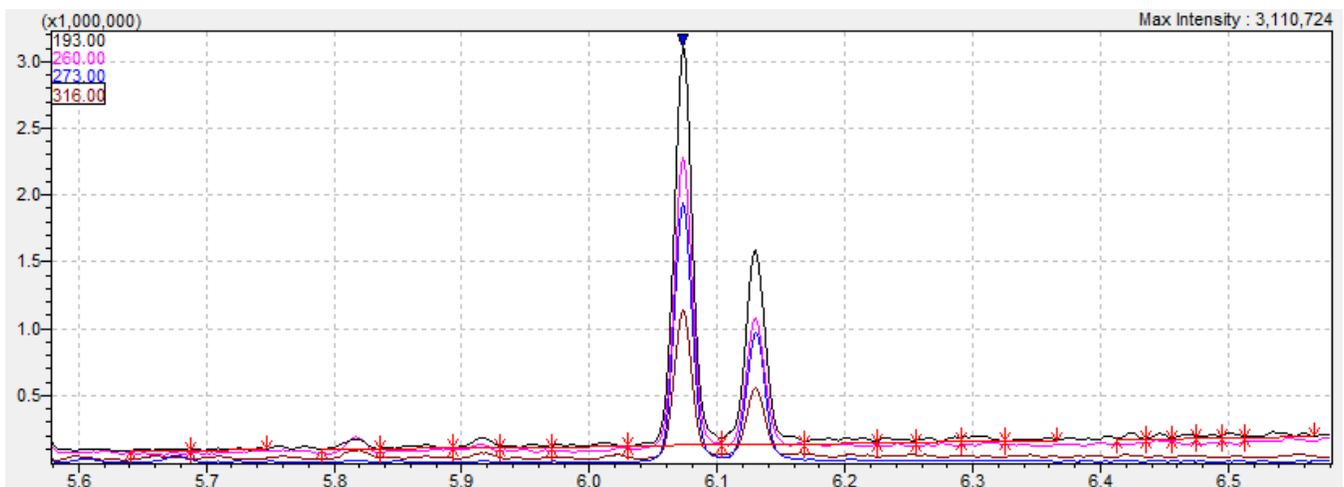
GC-MS/MS Analysis of Sample #2132

The sample was subjected to GC-MS/MS analysis to obtain evidence for the presence of hexahydrocannabinol diastereomers and confirm the identity of the Δ^8 -THC.

The GC/MS spectrum of hexahydrocannabinol was obtained from the NIST library and is shown below. Prominent ions are reported at m/z 193 (base peak), m/z 273, and m/z 260.

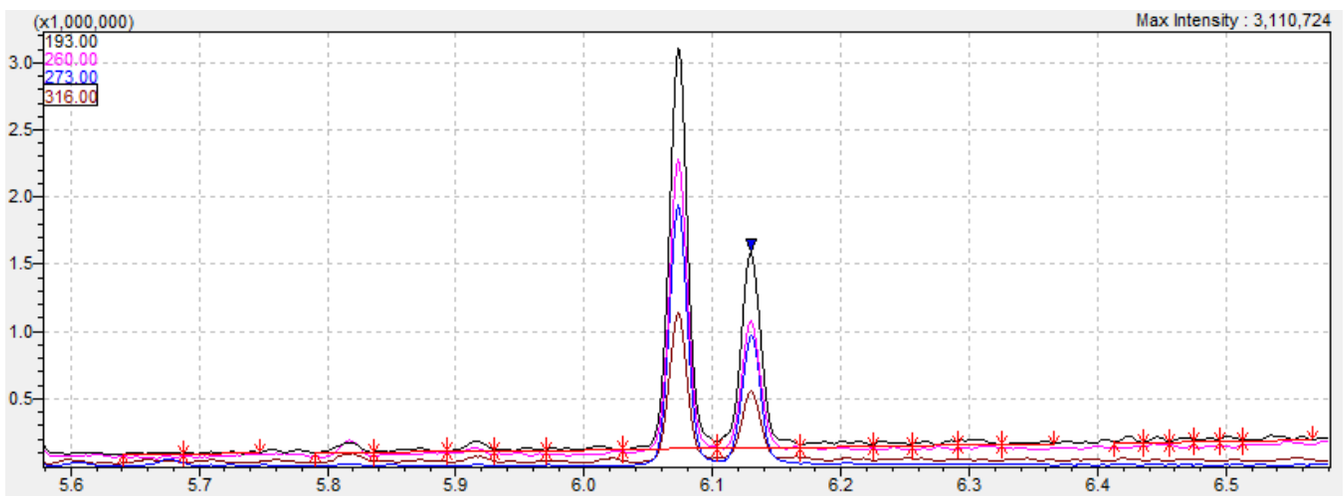


Those ions characteristic of hexahydrocannabinol were monitored during the GC/MS analysis of Sample 2123. The selected ion chromatogram is shown below and the relative abundances of the ions characteristic of hexahydrocannabinol from the 1st prominent peak at about 6.08 min are shown in the table below.



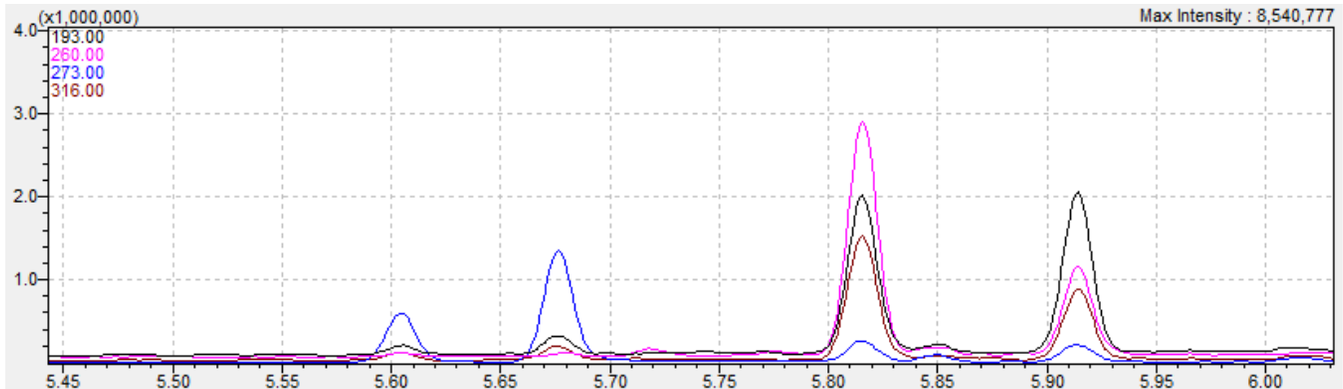
Type	m/z	Intensity	Set%	Act. %	Ref. Band
Target	193.00	3107990	100.00	100.00	---
Ref. Ion1	260.00	2249844	0.00	72.39	30
Ref. Ion2	273.00	2044301	0.00	65.78	30
Ref. Ion3	316.00	846728	0.00	27.24	30
Ref. Ion4					
Ref. Ion5					

The selected ion chromatogram is shown below and the relative abundances of the ions characteristic of hexahydrocannabinol from the 2nd prominent peak at about 6.13 min are shown in the table below.

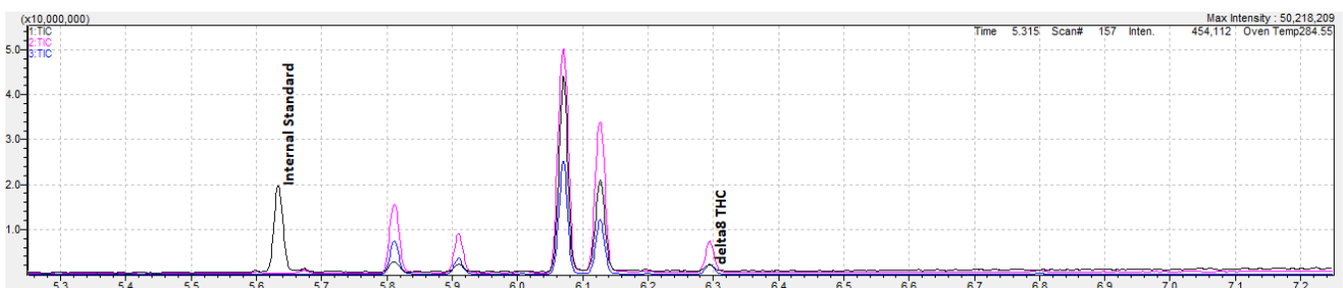


Type	m/z	Intensity	Set%	Act. %	Ref. Band
Target	193.00	1472511	100.00	100.00	---
Ref. Ion1	260.00	994232	0.00	67.52	30
Ref. Ion2	273.00	955394	0.00	64.88	30
Ref. Ion3	316.00	518360	0.00	35.20	30
Ref. Ion4					
Ref. Ion5					

The selected ion chromatograms of the ions characteristic of hexahydrocannabinol from the less abundant peaks before 6.0 min are shown below.

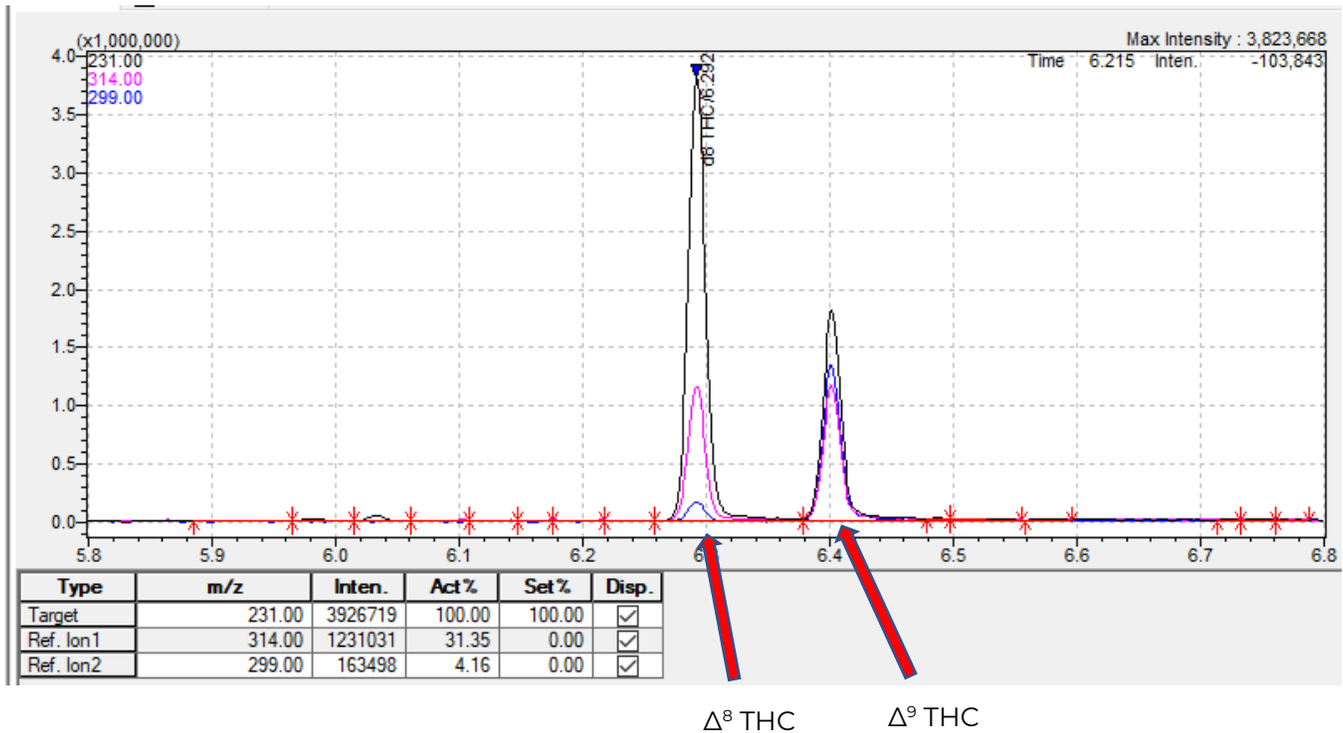


These results indicate with reasonable assurance that at least two and possibly more isomers of hexahydrocannabinol are present in the sample submitted for analysis. Proof of identity would require a reference standard.



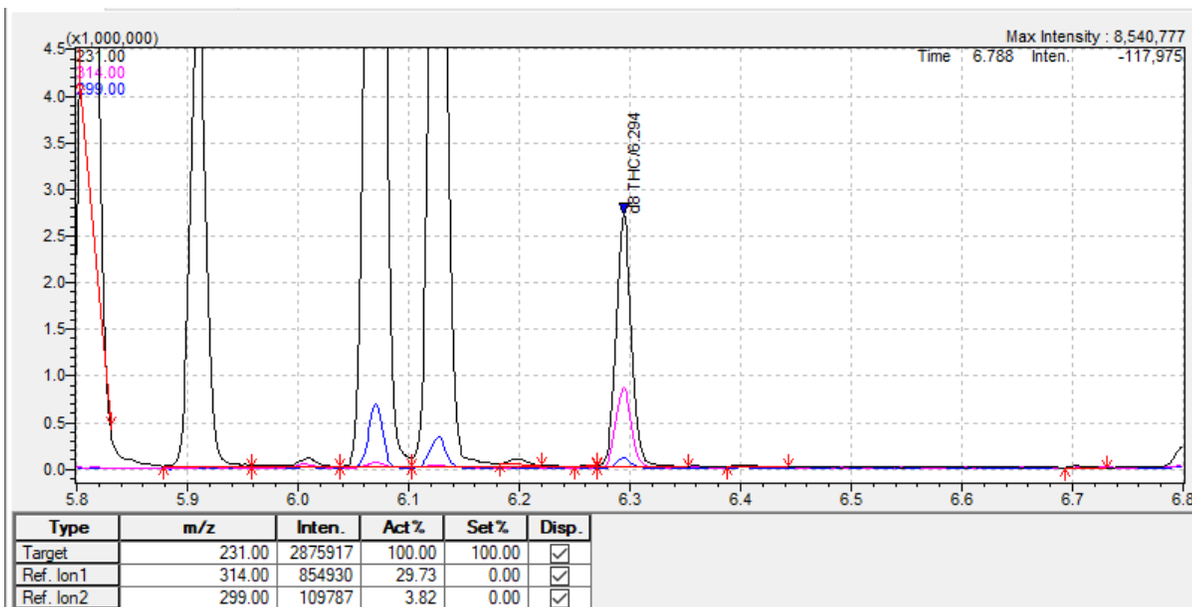
The peak at 6.3 minutes was identified as Δ^8 -THC. See the following page for confirmation of the identity of Δ^8 -THC.

The following is a SIM analysis of a Standards Mix showing ions and retention times for Δ^8 -THC and Δ^9 -THC. The table below the ion chromatogram reports the relative abundances of the most abundant ion for Δ^8 -THC at m/z 231, the molecular ion at m/z 314, and a fragment ion at m/z 299.



The following is a SIM analysis of Sample 2132 showing ions and retention times for Δ^8 -THC and Δ^9 -THC. The table below the ion chromatogram reports the relative abundances of the most abundant ion for Δ^8 -THC at m/z 231, the molecular ion at m/z 314, and a major fragment ion at m/z 299.

SIM analysis of Sample 2132.





232 North Plaza Drive
Nicholasville, KY 40356

The presence of Δ^8 -THC in sample #2132 was confirmed by comparison of the retention time and relative abundances of the qualifier ions with those of a certified reference standard.

Conclusions

- Sample #2132 contains a small amount of D⁸-THC based on HPLC-PDA, LC-MS, and GC-MS data.
- Sample #2132 contains components characterized by greater lipophilicity than D⁸-THC based on longer HPLC and LC retention times. The components eluting after D⁸-THC are characterized by a pseudomolecular ion at m/z 317 based on positive ionization electrospray LC-MS analysis indicating an apparent molecular weight of 316, consistent with that of the hexahydrocannabinol isomers. Analysis of sample #2132 by GC/MS analysis indicated the presence of at least two substances with an apparent molecular mass of 316 and characterized by fragment ions at m/z 193, m/z 260, and m/z 273. These fragment ions are consistent with reported fragment ions of hexahydrocannabinol.
- Confirmation of the presence of hexahydrocannabinol in sample #2132 requires availability of a reference standard.
- Sample #2132 contains no other detectable cannabinoids from our collection of more than forty reference standards.