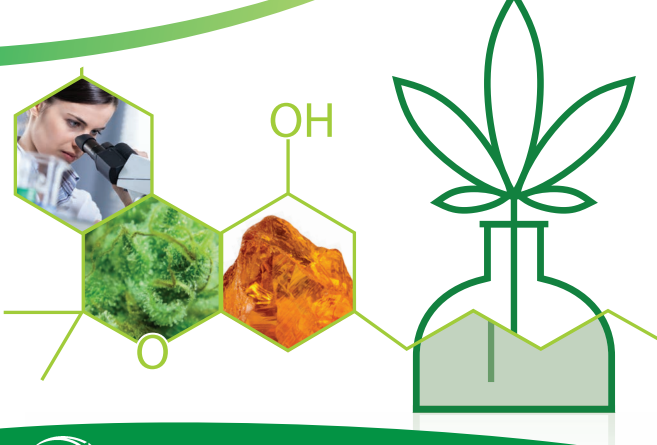


Growing Analytical Solutions for Cannabis Testing

INNOVATIVE PRODUCTS AND EXPERTISE FOR ACCURATE AND RELIABLE RESULTS



RESTEK

Pure Chromatography

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1 (of 102) >2015

Restek has been helping cannabis labs establish innovative, cost-effective analytical solutions from the very beginning, and we will continue to help you manage your everchanging analytical challenges every step of the way.

Technical Expertise &

By Breaking Boundaries in Our Industry, We Help You Succeed in Yours

We get it. Your market is quickly changing and you need a chromatography partner that understands that. Whether you are part of a well-established safety and potency lab or starting a new lab, Restek has the products and expertise you need for successful cannabis analyses. Being an employee-owned and independent chromatography company, every employee at Restek has a vested interest in your success. We design the best solutions for your lab, regardless of the instrumentation and techniques used. In this brochure, you will find innovative LC and GC products and methodologies designed to fit your toughest analytical problems.

We've been in your shoes. That's why we understand your challenges and focus on solving them. Using our expertise to develop innovative products that help chromatographers has always been, and continues to be, Restek's top priority. We strive to develop industry-leading technologies that fit the needs of today's analysts. When setting up a laboratory for cannabis testing, we realize that you need dependable products that deliver high quality data without considerable capital investment. We know you need to work with a company that understands the challenges of your market and supports you with tailored solutions and superior customer service.

Rxi® GC COLUMNS





The chemists at Restek have combined their analytical expertise and wide range of polymer chemistries to provide a solution for straightforward analysis of terpenes and residual solvents on a single Rxi® column platform, streamlining workflows for busy labs. Rxi® columns deliver more accurate, reliable results than any other fused silica column on the market. To ensure the highest level of performance, all Rxi® capillary columns for the cannabis industry are manufactured and individually tested to meet stringent requirements for exceptional inertness, low bleed, and unsurpassed column-to-column reproducibility.

Sky® GC INLET LINERS

True Blue Performance—State-of-the-Art Deactivation With a 100% Satisfaction Guarantee



Whether you're determining cannabinoids, residual solvents, pesticides, or terpenes by GC, the inertness of your inlet is crucial for the success of your analyses. Sky* inlet liners from Restek use a comprehensive, state-of-the-art deactivation and are the only blue liners on the market—making them an easy-to-recognize solution to common inlet problems. The innovative deactivation used for Sky* liners results in exceptional inertness for a wide range of analyte chemistries. In addition to improved data quality, you'll benefit from fewer liner changes and less downtime for maintenance.



www.restek.com/cannabis

Product Innovation



Raptor™ LC COLUMNS

Maximize Analytical Performance and Minimize Your Capital Investment



Raptor™ LC columns combine the speed of a superficially porous particle (SPP or "core-shell") with the separation power of optimized USLC phase chemistry. These columns are ideal for cannabis testing because they quickly separate your target compounds, providing higher sample throughput. Raptor™ LC columns maximize your instrument performance so you won't need to buy expensive UHPLC equipment or extend your capital investment when the sample volume increases. Build a solid analytical foundation on any instrument with fast, rugged Raptor™ LC columns.

Q-sep® SAMPLE PREP SUPPLIES

Everything You Need for Fast, Simple Sample Prep



Cannabis products present a broad array of challenging matrices, from foods, to plant materials, to concentrates. For pesticides analysis, a fast, easy cleanup method is required to remove the matrix background for accurate, reliable results. Restek's versatile line of Q-sep™ QuEChERS extraction and cleanup salts allows for the development of quick, easy, and affordable sample preparation methods without capital investment in extraction equipment. The friendly experts at Restek are always willing to help with method development questions, too.

CERTIFIED REFERENCE MATERIALS (CRMs)

Get Results You can Trust With World-Class CRMs Produced in ISO-Accredited Labs

In order to achieve accurate results, samples must be quantified using certified reference materials. Restek has the widest offering of cannabinoid standards in the industry, and we are continually expanding our product line in order to meet the evolving needs of the cannabis industry. Restek's certified reference materials are manufactured and QC tested under our ISO Guide 34 and ISO/ IEC 17025 accreditations, helping ensure confidence in results and compliance with changing regulations.



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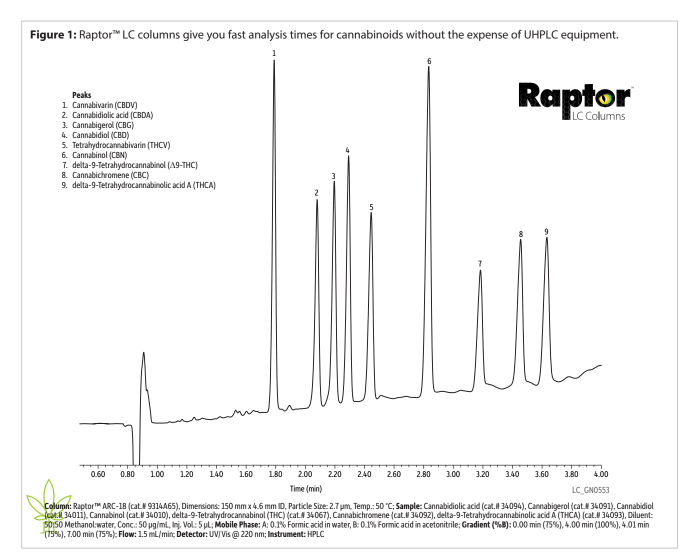
Applications

PRODUCT POTENCY TESTING

Our High-Throughput LC and GC Cannabinoids Methods Produce Results Quickly Without the Cost of New Equipment

When setting up a lab, often you just can't invest in the latest instrumentation, but you still need to get results fast. We understand that. That's why Restek has developed both LC and GC methods for cannabinoids that let you report potency results quickly. For LC, we created a fast analysis that can be performed on any LC instrument. By utilizing Raptor™ column technology, as shown in Figure 1, we developed a 3.7 minute analysis (7 minutes total cycle time) that is compatible with any HPLC instrument—so you get UHPLC speed on your existing equipment without the capital investment. Also, we specifically chose an easy-to-make mobile phase that can be directly

transferred to LC-MS, if you ever need to move to MS due to regulation changes. For labs using GC equipment, you can analyze cannabinoids in just minutes using an Rxi®-35Sil MS column and the instrument conditions shown in Figure 2. We also offer a similar 35-type stationary phase on metal MXT® tubing for labs using SRI GC instruments. Why did we focus on fast cannabinoid analyses? Potency testing is the cornerstone of your lab. Building a fast method means your productivity increases and you can analyze more samples per day on the same instrument, delaying the need for expensive capital investments in new equipment.



TECH TIP

Using syringe filters is an economical way to remove particulate matter that could clog your column. Visit **www.restek.com/filters** to access our solvent/syringe filter compatibility guide and quickly find the best filter for your method.





Figure 2: Determine critical cannabinoids in minutes by GC using an Rxi®-35Sil MS column. Conc. tR (sec) (µg/mL) 50 100 Phencyclidine (IS) Cannabidivarin 115.2 100 Tetrahydrocannabivarin 130.3 100 138.3 Cannabichromene Cannabidiol 100 150.0 Δ8-Tetrahydrocannabinol 152.1 7. $\Delta 9$ -Tetrahydrocannabinol 100 156.1 8. Cannabigerol 100 159.5 9. Cannabinol 100 165.6 10. Prazepam (IS) 192.4 Column: Rxi®-35Sil MS, 15 m, 0.25 mm ID, 0.25 µm (cat.# 13820), Sample: Phencyclidine (cat.# 34027); Cannabichromene (CBC) (cat.# 34092); Cannabinoids standard (cat.# 34014); delta-8-Tetrahydrocannabinol (THC) (cat.# 34090): Cannabigerol (CBG) (cat.# 34091); Prazepam (cat.# 34055); Injection: Inj. Vol.: 1 µL split (split ratio 20:1); Liner: Sky® 4 mm Precision® liner w/wool (cat.# 23305.5); Inj. Temp.: 250 °C; **Oven:** Oven Temp.: 190 °C (hold 0.1min) to 330 °C at 35 °C/min (hold 0.9 min); Carrier Gas: H2, constant flow; Flow Rate: 2.5 mL/min; Detector: FID @ 350 °C; Constant Column + Constant Make-up: 50 mL/min; Make-up Gas Type: N2; Hydrogen flow: 40 mL/min; Air flow: 450 mL/min; Data Rate: 20 Hz; Instrument: Agilent/HP6890 GC; Notes: Cannabidivarin and tetrahydrocannabivarin standards were obtained from BOC Sciences. 100 180 200 80 120 140 160 Time (sec) GC_FS0549

POTENCY TESTING PRODUCTS

Raptor™ ARC-18 LC Columns (USP L1)

Properties:

- Well-balanced retention profile.
- Sterically protected and acid-resistant to resist harsh, low-pH mobile phases.
- Ideal for use with sensitive detectors like mass spec.

Description	cat.#
2.7 µm Columns 150 mm, 4.6 mm ID	9314A65

For guard cartridges, visit our website at www.restek.com

Rxi®-35Sil MS Columns (fused silica)

(midpolarity Crossbond® phase)

- Provides superior separation for cannabinoids.
- Very low-bleed phase for GC-MS analysis.
- Extended temperature range: 50 °C to 340/360 °C.

Description	temp. limits	qty.	cat.#
15 m, 0.25 mm ID, 0.25 μm	50 to 340/360 °C	ea.	13820

Sky® 4.0 mm ID Precision® Inlet Liner w/Wool

For Agilent GCs equipped with split/splitless inlets

ID x OD x L	qty.	cat.#	
Precision, Sky Technology, Borosilicate Gla	ss with Quartz Wool		
4.0 mm x 6.3 mm x 78.5 mm	ea.	23305.1	
4.0 mm x 6.3 mm x 78.5 mm	5-pk.	23305.5	
4.0 mm x 6.3 mm x 78.5 mm	25-pk.	23305.25	

Patent pending

Medical Marijuana Singles

Concentration is $\mu g/mL$. Volume is 1 mL/ampul.

CAS#	Solvent	Conc.	cat.#
20675-51-8	PTM	1,000	34092
13956-29-1	PTM	1,000	34011
1244-58-2	ACN	1,000	34094
25654-31-3	PTM	1,000	34091
521-35-7	PTM	1,000	34010
5957-75-5	PTM	1,000	34090
1972-08-3	М	1,000	34067
23978-85-0	PTM	1,000	34093
1693-74-9	PTM	2,000	30112
104874-50-2	М	100	34068
	20675-51-8 13956-29-1 1244-58-2 25654-31-3 521-35-7 5957-75-5 1972-08-3 23978-85-0 1693-74-9	20675-51-8 PTM 13956-29-1 PTM 1244-58-2 ACN 25654-31-3 PTM 521-35-7 PTM 5957-75-5 PTM 1972-08-3 M 23978-85-0 PTM 1693-74-9 PTM	20675-51-8 PTM 1,000 13956-29-1 PTM 1,000 1244-58-2 ACN 1,000 25654-31-3 PTM 1,000 521-35-7 PTM 1,000 5957-75-5 PTM 1,000 1972-08-3 M 1,000 23978-85-0 PTM 1,000 1693-74-9 PTM 2,000

M = methanol; PTM = purge-and-trap grade methanol; ACN = acetonitrile

Cannabinoids Standard (3 components)

Cannabidiol (13956-29-1) Cannabinol (521-35-7)

delta-9-Tetrahydrocannabinol (Δ ⁹-THC) (1972-08-3)

 $1,\!000\,\mu g/mL$ each in P&T methanol, $1\,mL/ampul$ cat.# 34014 (ea.)

Quantity discounts not available.

Phencyclidine

Phencyclidine (956-90-1)

 $1,000 \, \mu g/mL$ in P&T methanol, $1 \, mL/ampul$ cat.# 34027 (ea.)

Prazepam

Prazepam (2955-38-6)

 $1,000 \mu g/mL$ in P&T methanol, 1 mL/ampul cat.# 34055 (ea.)



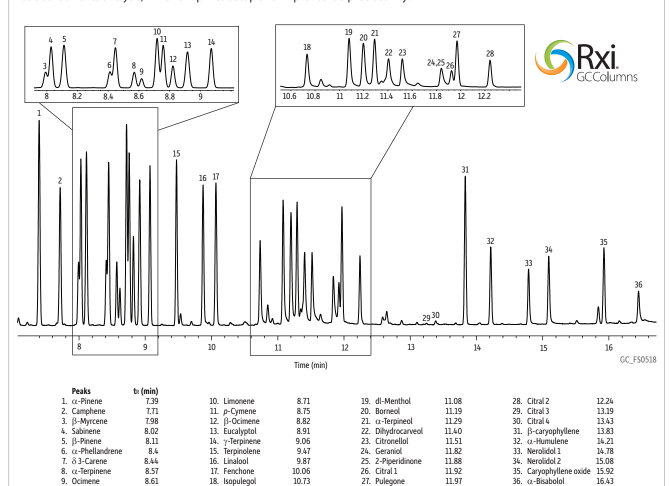
TERPENE PROFILING

Reduce Capital Investments—Analyze Terpenes by GC on the Same Setup Used for Residual Solvents

Cannabis has a complex terpene profile, which is theorized to increase its therapeutic effects. Terpene profiling is used for both product quality testing and strain identification. These complex and sometimes problematic compounds are challenging to analyze, but the experts at Restek have developed

GC methodology for terpene profiling that fits easily into required laboratory workflows. To keep things simple, the GC terpene profile analysis in Figure 3 can be performed on the same instrument and column that we recommend for residual solvent testing (see page 8).

Figure 3: Comprehensive terpene analysis by headspace GC-FID can be done on the same instrument and GC column as residual solvents analysis, which simplifies setup and improves lab productivity.



Column: Rxi® -624Sil MS, 30 m, 0.25 mm ID, 1.40 µm (cat.# 13868); Sample:Terpenes mix; Diluent: Isopropyl alcohol; Conc.: 200 ng/µL (0.02% wt/vol). The sample was prepared by placing 10 µL into the headspace vial.; Injection: headspace-loop split (split ratio 10:1); Liner: Sky® 1.0 mm ID straight inlet liner (cat.# 23333.1); Headspace-Loop; Inj. Port Temp.: 250 °C; Instrument: Tekmar HT-3; Inj. Time: 1.0 min; Transfer Line Temp.: 160 °C; Valve Oven Temp.: 160 °C; Needle Temp.: 140 °C; Sample Temp.: 140 °C; Sample Equil. Time: 30.0 min; Vial Pressure: 20 psi; Loop Pressure: 15 psi; Oven: Oven Temp.: 60 °C (hold 0.10 min) to 300 °C at 12.50 °C/min (hold 3.0 min); Carrier Gas: He, constant flow; Linear Velocity: 33 cm/sec; Detector: FID @ 320 °C; Make-up Gas Flow Rate: 45 mL/min; Make-up Gas Type: Nz; Hydrogen flow: 40 mL/min; Air flow: 450 mL/min; Data Rate: 20 Hz; Instrument Agilent/HP6890 GC

TECH TIP

For full method details on headspace GC analysis of terpenes, visit www.restek.com/cannabis_terpenes





TERPENE TESTING PRODUCTS

Rxi®-624Sil MS Columns (fused silica)

(midpolarity Crossbond® phase)

- Low-bleed, high-thermal stability column—maximum temperatures up to 320 °C.
- Inert—excellent peak shape for a wide range of compounds.
- Selective—G43 phase highly selective for volatile organics and residual solvents, great choice for USP<467>.
- Manufactured for column-to-column reproducibility—well-suited for validated methods.

Description	temp. limits	qty.	cat.#
30 m, 0.25 mm ID, 1.40 μm	-20 to 300/320 °C	ea.	13868



Sky® 1.0 mm ID Straight Inlet Liner

for Agilent GCs equipped with split/splitless inlets

qty.	cat.#
ea.	23333.1
5-pk.	23333.5
25-pk.	23333.25
	ea. 5-pk.

* 100% SATISFACTION GUARANTEE: If your Sky® inlet liner does not perform to your expectations for any reason, simply contact Restek® Technical Service or your local Restek® representative and provide a sample chromatogram showing the problem. If our GC experts are not able to quickly and completely resolve the issue to your satisfaction, you will be given an account credit or replacement product (same cat.#) along with instructions for returning any unopened product. (Do not return product prior to receiving authorization.) For additional details about Restek's return policy, visit www.restek.com/warranty



Headspace Crimp Vials (20 mm)

Description	Volume	Color	Dimensions	100-pk.	1,000-pk.
Headspace Vial, Flat Bottom	20 mL	Clear	23 x 75 mm	24685	24686

Vial-to-instrument compatibility are designated in instrument reference chart on the product web page.



Medical Cannabis Terpenes Standards

Medical Cannabis Terpenes Standard #1 (19 components)

(-)-alpha-Bisabolol (23089-26-1) Camphene (79-92-5) delta-3-Carene (13466-78-9) beta-Caryophyllene (87-44-5) Geraniol (106-24-1) (-)-Guaiol (489-86-1) alpha-Humulene (6753-98-6) p-Isopropyltoluene (p-cymene) (99-87-6) (-)-Isopulegol (89-79-2) d-Limonene (5989-27-5)

beta-Myrcene (123-35-3) Nerolidol (7212-44-4) Ocimene (13877-91-3) alpha-Pinene (80-56-8) (-)-beta-Pinene (18172-67-3) alpha-Terpinene (99-86-5) gamma-Terpinene (99-85-4) Terpinolene (586-62-9)

Linalool (78-70-6)

 $2,\!500\,\mu g/mL$ each in isopropanol, $1\,mL/ampul$ cat.# 34095 (ea.)

Did you **know**?

You'll save money ordering from Restek because we understand the need to control costs and build efficient workflows. We develop as many analyses as possible using the same columns and consumables, so you can minimize the number of products you need to stock.

Medical Cannabis Terpenes Standard #2 (2 components)

(-)-Caryophyllene oxide (1139-30-6) 1,8-Cineole (Eucalyptol) (470-82-6)

 $2,500 \, \mu g/mL$ each in isopropanol, $1 \, mL/ampul$ cat.# 34096 (ea.)

TECH TIP

Did you know that headspace analysis eliminates the possibility of column contamination from nonvolatile matrix components? This results in an extremely clean chromatogram, minimal instrument maintenance, and longer column lifetimes.



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RESIDUAL SOLVENT ANALYSIS

Improve Productivity—Keep Analyzing Samples Instead of Changing Columns Between Residual Solvent and Terpene Methods.

As the popularity of medical cannabis grows, so does concern over the safety of the drug products. Cannabis concentrates can contain residual solvents left over from manufacturing that can be harmful to human health. Because of this risk, many states will require residual solvent testing of cannabis concentrates. Due to their high volatility, residual solvents can

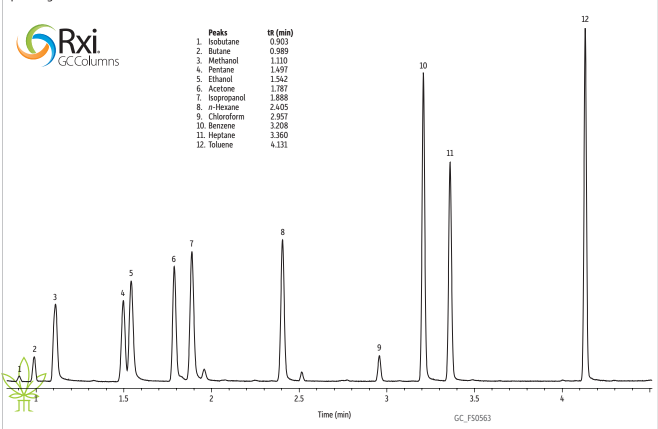
only be analyzed using GC techniques. The chemists at Restek have developed a quick and easy method that allows for residual solvent analysis (Figure 4) and terpene profiling (Figure 3) on the same column and instrument platform with minimal sample preparation (see page 6 for terpene profiling).

TECH TIP

For full method details on headspace GC analysis of residual solvents, visit **www.restek.com/cannabis_solvents**



Figure 4: Improve productivity and reduce downtime for column changes—this sensitive headspace GC-FID analysis of residual solvents can be accomplished on the same instrument and Rxi®-624Sil MS column that is used in Restek's terpenes profiling method.



Column: Rxi®-624Sil MS, 30 m, 0.25 mm ID, 1.40 µm (cat.# 13868); Sample: Residual solvent mix; Diluent: Dimethyl sulfoxide (DMSO); Conc.: 25 ppm (For the HS-FET technique, 10 µL of a 50 µg/mL standard was placed into a 20 mL headspace vial to represent a 25 ppm sample concentration, assuming a 20 mg sample weight); Injection: headspace-loop split (split ratio 10:1); Liner: Sky® 1.0 mm ID straight inlet liner (cat.# 23333.1); Headspace-Loop: Inj. Port Temp.: 250 °C; Instrument: Tekmar HT3; Inj. Time: 1.0 min; Transfer Line: Temp.: 160 °C; Valve Oven Temp.: 160 °C; Nedel Temp.: 140 °C; Sample Temp.: 140 °C; Platen temp equil. time: 1.0 min; Sample Equil. Time: 30.0 min; Vial Pressure: 20 psi; Pressurize Time: 5.0 min; Loop Pressure: 15 psi; Loop Fill Time: 2.0 min; Oven Temp.: 35 °C (hold 1.5 min) to 300 °C at 30 °C/min (hold 2.0 min); Carriere Gas: He, constant flow; Linear Velocity: 80 cm/sec; Detector: FID @ 320 °C; Make-up Gas Flow Rate: 45 mL/min; Make-up Gas Type: N2; Hydrogen flow: 40 mL/min; Air flow: 450 mL/min; Data Rate: 20 Hz; Instrument: Agilent/IHP6890 GC; Notes: The butane used for standard preparation was a mixture of butane and isobutane in an unknown ratio. The concentrations should be considered approximate, but do not exceed 50 ppm for any component.

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RESIDUAL SOLVENT TESTING PRODUCTS

Rxi®-624Sil MS Columns (fused silica)

(midpolarity Crossbond® phase)

- Low-bleed, high-thermal stability column—maximum temperatures up to 320 °C.
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- Selective—G43 phase highly selective for volatile organics and residual solvents, great choice for USP<467>.
- Manufactured for column-to-column reproducibility—well-suited for validated methods.

Description	temp. limits	qty.	cat.#
30 m, 0.25 mm ID, 1.40 μm	-20 to 300/320 °C	ea.	13868



Sky® 1.0 mm ID Straight Inlet Liner

for Agilent GCs equipped with split/splitless inlets

ID x OD x L	qty.	cat.#
Straight, Sky Technology, Borosilicate Glass		
1.0 mm x 6.3 mm x 78.5 mm	ea.	23333.1
1.0 mm x 6.3 mm x 78.5 mm	5-pk.	23333.5
1.0 mm x 6.3 mm x 78.5 mm	25-pk.	23333.25

RESTEK

^{* 100%} SATISFACTION GUARANTEE: If your Sky® inlet liner does not perform to your expectations for any reason, simply contact Restek® Technical Service or your local Restek® representative and provide a sample chromatogram showing the problem. If our GC experts are not able to quickly and completely resolve the issue to your satisfaction, you will be given an account credit or replacement product (same cat.#) along with instructions for returning any unopened product. (Do not return product prior to receiving authorization.) For additional details about Restek's return policy, visit www.restek.com/warranty



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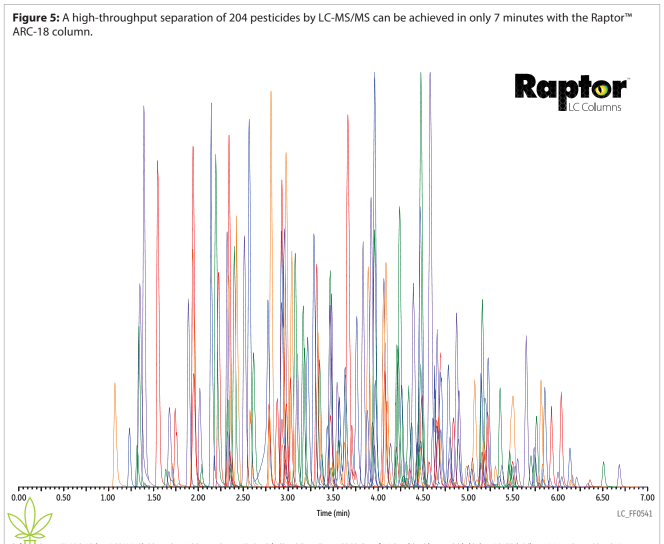
www.restek.com/cannabis

PESTICIDE ANALYSIS

Ensure Product Safety With Fast, Selective Multiresidue Pesticide Analysis

In addition to residual solvents, cannabis products can contain residues of pesticides that were applied to cannabis plants during growth in order to control agricultural pests. These pesticides can be analyzed by LC-MS/MS, GC-MS/MS, and GC-MS. Regardless of the technique used, lists of target compounds can be extensive, so column selectivity is an important factor in achieving good separations. Both Raptor™

ARC-18 LC columns (Figure 5) and Rxi®-5ms GC columns (Figure 6) provide the selectivity needed for accurate and reliable multiresidue pesticides analysis. Removing matrix interferences while also recovering the analytes of interest is also crucial for a successful pesticide analysis using either LC or GC, and Restek's Q-sep® QuEChERS products allow for fast, easy, adaptable cleanup of a wide variety of matrices.



Column: Raptor ARC-18 (cat.# 9314A12), Dimensions: 100 mm x 2.1 mm ID, Particle Size: 2.7 µm, Temp.: 50 °C; Sample: LC multiresidue pesticide kit (cat.# 31971), Diluent: Water, Conc.: 20 ng/mL, Inj. Vol.: 5 µL; Mobile Phase: A: Water + 2 mM ammonium formate + 0.2% formic acid, B: Methanol + 2 mM ammonium formate + 0.2% formic acid; Gradient (%B): 0.00 min (5%), 2.00 min (5%), 4.00 min (75%), 6.00 min (100%), 7.01 min (5%), 9.50 min (5%); Flow: 0.4 mL/min; Max Pressure: 525 bar; Detector: Waters Xevo TQ-S, Ion Source: Waters Zspray ES, Ion Mode: ESI+, Mode: MRM, Instrument: Waters ACQUITY UPLC® I-Class; Notes: When combining a large number of compounds with different chemical functionalities, mix stability can be an issue. In formulating our LC multi-residue pesticide standard kit (cat.# 31971), we extensively studied the 204 compounds involved, then grouped them into as few mixes as possible while still ensuring maximum long-term stability and reliability. Several of these compounds are isomeric and separation of the isomers accounts for 216 peaks in the chromatogram compound list. For quantitative analysis, we recommend analyzing each mix separately to ensure accurate results for every compound.

Figure 6: Peak List

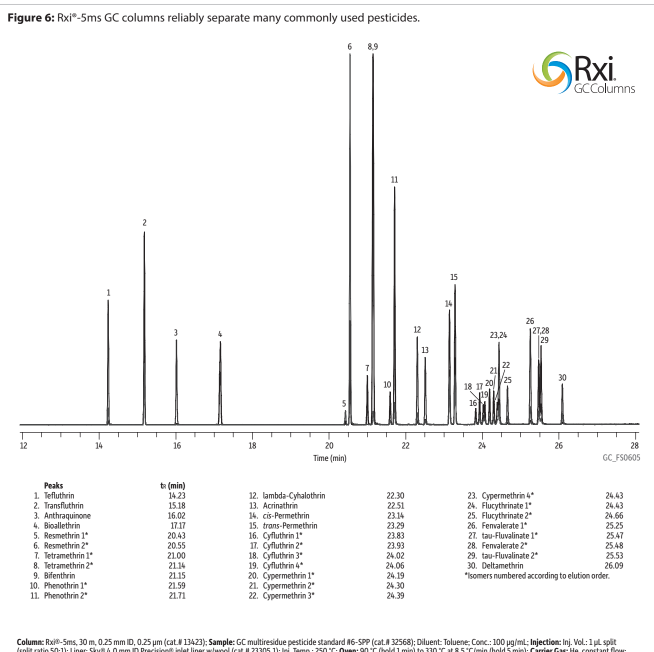
		Draguraar	Duaduat	Duaduat			Duagungan	Duaduat	Dundust			Duagunga	. Duadust	Duaduat
Peaks	t _R (min)	Precursor Ion	lon 1	lon 2	Peaks	tr (min)	Precursor Ion	lon 1	lon 2	Peaks	tr (min)	Precursor	lon 1	lon 2
Cyromazine	1.07	167.0	85.0	108.1	75. Isocarbophos	3.48	291.1	121.1	231.1	149. Bromuconazole isor		376.0	158.9	70.1
2. Methamidophos	1.23	142.0	93.9	124.9	76. Isoproturon	3.48	207.0	72.0	47.0	150. Flubendiamide	4.89	683.0	408.0	274.0
Formetanate HCl	1.32	222.0	165.0	46.0	77. Pyrimethanil	3.48	200.0	107.0	82.0	151. Carfentrazone ethy		412.0	346.0	266.0
4. Aminocarb	1.34	209.0	137.0	152.0	78. Desmedipham	3.55	318.0	182.0	154.0	152. Diclobutrazol	4.91	328.0	70.0	59.1
 Pymetrozine Acephate 	1.35 1.40	218.0 184.1	105.0 143.0	79.0 125.1	79. Metalaxyl 80. Spiroxamine isomer 1	3.56 3.57	280.1 298.0	220.1 144.0	192.1 100.0	153. Kresoxim-methyl 154. Tebuconazole	4.92 4.98	314.1 308.0	206.0 70.1	116.0 125.0
7. Propamocarb	1.40	189.1	102.0	144.0	81. Phenmedipham	3.63	301.0	168.0	136.0	155. Penconazole	5.00	284.0	70.1	159.0
8. Omethoate	1.55	214.1	125.1	183.1	82. Spiroxamine isomer 2	3.63	298.0	144.0	100.0	156. Spinosyn A	5.04	732.6	142.0	98.1
Aldicarb sulfoxide	1.64	207.0	89.0	132.0	83. Chlorantraniliprole	3.66	483.9	286.0	453.0	157. Prothioconazole	5.05	344.0	326.0	189.0
10. Dinotefuran	1.64	203.0	129.0	157.0	84. Cycluron	3.68	199.0	89.1	69.2	158. Alanycarb	5.06	400.0	238.2	254.1
11. Butoxycarboxim	1.67 1.68	223.0 271.1	106.0 125.9	166.0 224.9	85. Prometryn	3.71 3.76	242.0 242.1	158.0 186.1	200.1 91.0	159. Zoxamide 160. Famoxadone	5.08 5.10	336.0 392.2	187.1 331.1	159.0 238.0
 Nitenpyram Aldicarb sulfone 	1.08	240.0	148.0	86.0	86. Terbutryn 87. Linuron	3.83	242.1	160.1	182.0	161. Prochloraz	5.10	376.0	308.0	70.1
14. Carbendazim	1.74	192.1	160.1	132.1	88. Fenobucarb	3.84	208.0	94.9	152.0	162. Triflumuron	5.15	359.0	156.1	139.1
15. Oxamyl	1.78	237.0	72.0	90.0	89. Diethofencarb	3.88	268.0	226.0	124.0	163. Benalaxyl	5.16	326.1	148.0	91.0
16. Flonicamid	1.89	230.0	203.1	174.1	90. Ethofumesate	3.89	287.1	121.1	259.1	164. Hexaconazole	5.16	314.0	70.1	159.0
17. Methomyl	1.91	163.0	106.0	88.0	91. Azoxystrobin	3.92	404.1	372.0	329.0	165. Hydramethylnon	5.17	495.1	323.2	151.1
 Thiabendazole Thiamethoxam 	1.94 1.94	202.0 292.0	175.0 211.0	131.0 181.0	92. Ethriprole 93. Fenamidone	3.94 3.96	396.9 312.1	350.9 236.1	255.2 92.0	166. Metconazole 167. Propiconazole	5.19	320.1	70.0	125.0
20. Mexacarbate	1.94	222.9	151.1	166.1	94. Methiocarb	3.96	226.0	121.0	169.0	isomer 1 & 2	5.19	342.0	159.0	69.0
21. Monocrotophos	2.02	224.1	127.1	98.1	95. Siduron	3.96	233.0	93.8	137.0	168. Clofentezine	5.22	303.0	138.0	102.0
22. Fuberidazole	2.04	185.0	157.0	156.0	96. Fludioxonil	3.97	249.1	229.1	158.1	169. Pyraclostrobin	5.23	388.1	163.0	193.9
23. Dicrotophos	2.14	238.0	112.0	193.0	97. Furalaxyl	3.97	302.1	270.1	242.2	170. Bitertanol	5.27	338.1	269.2	70.1
24. Imidacloprid	2.19	256.1	175.1	209.1	98. Halofenozide	3.99	331.1	104.9	275.0	171. Benzoximate	5.29	364.0	199.1	105.0
25. Clothianidin 26. Trichlorfon	2.22 2.32	250.0 257.0	169.0 109.0	132.0 79.0	99. Acibenzolar-S-methyl 100. Boscalid	4.06 4.06	210.9 342.9	91.0 307.0	135.9 139.9	172. Spinosyn D 173. Thiobencarb	5.31 5.31	746.5 257.9	142.0 125.1	98.1 100.1
27. 3-Hydroxycarbofuran	2.33	238.0	181.0	163.0	101. Dimethomorph isomer:		388.1	300.9	165.0	174. Diniconazole	5.35	326.1	70.2	159.0
28. Fenuron	2.33	165.0	71.9	45.9	102. Nuarimol	4.08	315.0	252.0	81.1	175. Pencycuron	5.36	329.1	125.0	218.0
29. Dimethoate	2.34	230.1	125.0	199.0	103. Mandipropamid	4.09	412.3	328.2	356.2	176. Spinetoram	5.38	748.5	142.2	98.1
30. Vamidothion	2.34	288.0	146.0	118.0	104. Flutolanil	4.10	324.1	262.1	65.0	177. Hexaflumuron	5.46	461.0	158.0	141.0
31. Dioxacarb	2.35	224.1	123.1	167.1	105. Promecarb	4.10	208.1	151.0	109.0	178. Indoxacarb	5.46	528.0	203.0	218.0
 Mevinphos isomer 1 Acetamiprid 	2.36 2.40	225.1 223.0	127.1 126.0	193.1 56.1	106. Paclobutrazol 107. Thiofanox	4.14 4.19	294.1 219.1	125.1 172.9	70.2 129.0	179. Ipconazole isomer 1 180. Triflumizole	5.46 5.49	334.2 346.0	70.0 277.9	125.1 60.0
34. Ethirimol	2.43	210.1	140.0	98.0	108. Cyproconazole isomer 1		292.2	125.1	70.2	181. Difenoconazole	3.43	340.0	211.5	00.0
35. Cymoxanil	2.46	199.0	128.0	111.0	109. Mepronil	4.21	270.1	119.0	91.0	isomer 1 & 2	5.50	406.0	251.1	111.1
36. Pirimicarb	2.51	239.1	72.0	182.1	110. Bupirimate	4.22	317.0	166.0	108.0	182. Trifloxystrobin	5.50	409.0	186.0	145.0
37. Thiacloprid	2.56	253.0	126.0	90.1	111. Dimethomorph isomer		388.1	300.9	165.0	183. Novaluron	5.53	493.0	158.0	141.0
38. Mevinphos isomer 2	2.58	225.1	127.1	193.1	112. Myclobutanil	4.26	289.1	70.2	125.1 268.1	184. Ipconazole isomer 2	5.56	334.2	70.0	125.1
39. Mesotrione 40. Butocarboxim	2.62 2.68	340.1 213.0	228.1 156.0	104.0 116.0	113. Clethodim isomer 1 114. Methoxyfenozide	4.28 4.30	360.0 369.1	164.0 149.1	313.2	185. Emamectin benzoate B1b	5.57	872.4	158.2	126.1
41. Aldicarb	2.71	213.1	89.1	116.1	115. Chloroxuron	4.31	291.1	164.1	111.0	186. Clethodim isomer 2		360.0	164.0	268.1
42. Oxadixyl	2.77	279.0	219.0	132.0	116. Cyprodinil	4.32	226.0	93.0	108.0	187. Buprofezin	5.70	306.1	201.0	57.4
43. Carbetamide	2.79	237.0	118.0	192.0	117. Triadimefon	4.34	294.1	197.2	69.3	188. Teflubenzuron	5.74	380.9	158.0	140.9
44. Tricyclazole	2.79	190.0	163.0	136.0	118. Bifenazate	4.35	301.1	198.0	170.0	189. Emamectin	F 7F	006 5	150.1	1061
45. Simetryn 46. Thiophanate-methyl	2.81 2.88	214.0 343.0	124.0 151.0	95.9 93.0	 Triadimenol Cyproconazole isomer 2 	4.35 2 4.38	296.1 292.2	99.1 125.1	70.2 70.2	benzoate B1a 190. Benfuracarb	5.75 5.76	886.5 411.1	158.1 195.0	126.1 190.0
47. Bendiocarb	2.93	224.1	109.0	167.0	121. Mefenacet	4.39	299.0	148.0	120.0	191. Fluazinam	5.78	464.8	373.0	338.1
48. Prometon	2.93	226.0	184.3	86.3	122. Mepanipyrim	4.40	224.1	106.0	77.0	192. Metaflumizone	5.79	507.0	287.2	267.1
49. Secbumeton	2.93	226.2	100.2	170.2	123. Iprovalicarb isomer 1	4.44	321.1	119.1	203.1	193. Furathiocarb	5.82	383.2	194.9	252.0
50. Thidiazuron	2.93	221.0	101.9	93.9	124. Fluquinconazole	4.45	376.0	348.8	306.9	194. Lufenuron	5.83	511.2	158.0	141.0
51. Propoxur 52. Metribuzin	2.95 2.96	210.0 215.0	111.0 131.0	168.0 89.0	125. Fenhexamid 126. Bromuconazole isomer	4.46	302.1 376.0	97.2 158.9	55.3 70.1	195. Temephos 196. Tebufenpyrad	5.83 5.86	467.1 334.0	125.0 117.0	418.9 145.0
53. Terbumeton	2.96	226.1	114.1	170.1	127. Fluoxastrobin	4.47	459.0	427.0	188.0	197. Pyriproxifen	5.91	322.1	96.0	227.1
54. Carbofuran	2.98	222.1	123.0	165.1	128. Iprovalicarb isomer 2	4.47	321.1	119.1	203.1	198. Piperonyl butoxide	5.93	356.3	176.9	119.0
55. Imazalil	2.98	297.0	159.0	69.0	129. Butafenacil	4.48	492.0	180.0	331.0	199. Hexythiazox	6.01	353.0	228.1	168.1
56. Sulfentrazone	3.03	387.0	307.0	145.8	130. Tetraconazole	4.48	372.0	159.0	70.1	200. Quinoxyfen	6.04	308.0	197.0	161.9
57. Pyracarbolid 58. Tebuthiuron	3.04 3.08	218.1 229.0	125.1 172.0	97.1 116.0	131. Flufenacet 132. Triticonazole	4.49 4.52	364.0 318.1	152.1 70.1	194.1 124.9	201. Flufenoxuron 202. Amitraz	6.05 6.14	489.1 294.0	158.0 163.0	141.0 122.0
59. Carbaryl	3.08	202.0	145.0	127.0	133. Cyazofamid	4.52 4.57	325.0	10.1	261.0	202. Affiliaz 203. Propargite	6.14	368.2	175.0	231.1
60. Carboxin	3.10	236.0	143.0	87.0	134. Spirotetramat	4.58	374.2	330.3	302.2	204. Etoxazole	6.16	360.2	304.2	177.2
61. Monolinuron	3.17	215.0	126.0	99.0	135. Diflubenzuron	4.63	311.1	141.0	158.1	205. Spiromesifen	6.20	371.1	273.1	255.1
62. Fluometuron	3.18	233.2	72.2	46.4	136. Epoxiconazole	4.66	330.0	121.0	101.0	206. Chlorfluazuron	6.21	539.8	382.9	158.0
63. Ethiofencarb	3.20	226.1	107.0	164.0	137. Etaconazole isomer 1 138. Fenbuconazole	4.66	328.1	205.0	159.0	207. Spirodiclofen	6.33	411.1	313.0	71.2
64. Ametryn 65. Chlortoluron	3.21 3.29	228.1 213.0	186.1 72.0	68.1 46.0	138. Fenbuconazole 139. Fenarimol	4.67 4.68	337.0 331.0	125.0 268.0	70.1 81.0	208. Fenpyroximate 209. Abamectin B1b	6.36 6.48	422.2 876.6	366.1 553.4	138.1 291.0
66. Metobromuron	3.32	259.1	170.0	148.1	140. Etaconazole isomer 2	4.70	328.1	205.0	159.0	210. Pyridaben	6.51	365.1	147.1	309.1
67. Methoprotryne	3.33	272.2	170.2	198.2	141. Fipronil	4.70	437.0	367.9	290.0	211. Eprinomectin	6.53	914.6	186.0	154.0
68. Propham	3.33	180.0	138.0	120.1	142. Flusilazole	4.78	316.0	247.0	165.0	212. Abamectin B1a	6.61	890.5	305.2	567.3
69. Flutriafol	3.35	302.1	123.1	70.2	143. Picoxystrobin	4.79	368.0	145.1	205.1	213. Fenazaquin	6.69	307.2	161.0	57.2
70. Isoprocarb 71. Fenpropimorph	3.37	194.1	95.1 1/-7.1	137.1 57.2	144. Fenoxycarb 145. Neburon	4.80	302.1	116.1 88.0	88.0 57.0	214. Doramectin 215. Moxidectin	6.82	916.6	331.2	593.4 528.4
71. Fenpropimorph 72. Methabenzthiazuron	3.44 3.46	304.2 222.0	147.1 165.0	57.2 150.0	145. Rotenone	4.80 4.84	275.0 395.0	213.1	57.0 192.1	216. Ivermectin	6.82 7.01	640.5 892.6	498.3 569.4	528.4 551.4
73. Diuron	3.47	233.0	72.1	46.3	147. Tebufenozide	4.87	353.1	133.0	297.1	210. IVERINCEUM	1.01	JJL.U	303.4	JJ1.T
74. Forchlorfenuron	3.47	248.1	129.0	93.0	148. Dimoxystrobin	4.88	327.1	116.1	205.2					

TECH TIP

Using syringe filters is an economical way to remove particulate matter that could clog your column. Visit **www.restek.com/filters** to access our solvent/syringe filter compatibility guide and quickly find the best filter for your method.

www.restek.com/cannabis



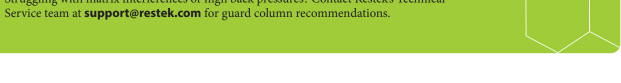


Column: Rxi®-5ms, 30 m, 0.25 mm ID, 0.25 μm (cat.# 13423); Sample: GC multiresidue pesticide standard #6-SPP (cat.# 32568); Diluent: Toluene; Conc.: 100 μg/mL; Injection: Inj. Vol.: 1 μL split (split ratio 50:1); Liner: Sky® 4.0 mm ID Precision® inlet liner w/wool (cat.# 23305.1); Inj. Temp.: 250 °C; Oven: 90 °C (hold 1 min) to 330 °C at 8.5 °C/min (hold 5 min); Carrier Gas: He, constant flow; Flow Rate: 1.4 mL/min; Detector: MS; Mode: Scan; Start Time: 5 min; Scan Range: 55-550 amu; Scan Rate: 7 scans/sec; Transfer Line Temp.: 290 °C; Analyzer Type: Quadrupole; Source Temp.: 325 °C; Electron Energy: 70 eV; Solvent Delay Time: 5 min; lonization Mode: EI; Instrument: Thermo Scientific TSQ 8000 Triple Quadrupole GC-MS; Notes: Bioallethrin isomers are only slightly resolved with this method, so they are treated as one peak. Chromatogram is reconstructed from select ions.



TECH TIP

Struggling with matrix interferences or high back pressures? Contact Restek's Technical



PESTICIDE ANALYSIS PRODUCTS

Raptor™ ARC-18 LC Columns (USP L1)



Properties:

- Well-balanced retention profile.
- Sterically protected and acid-resistant to resist harsh, low-pH mobile phases.
- · Ideal for use with sensitive detectors like mass spec.

Description	cat.#
2.7 μm Columns 100 mm, 2.1 mm ID	9314A12

For guard cartridges, visit our website at www.restek.com



Q-sep® QuEChERS Extraction Salts

Fast, Simple Sample Prep for Multiresidue Pesticide Analysis

- Salt packets eliminate the need for a second empty tube to transfer salts.
- Go green by using packets with reusable tubes.
- · Convenient and easy to use.

Description	Material	Methods	qty.	cat.#
Q-sep Kit	6 g MgSO4, 1.5 g NaOAc with 50 mL Centrifuge Tube	AOAC 2007.01	50 packets & 50 tubes	26237

NaOAc—sodium acetate

For LC Analysis

Q-sep® QuEChERS dSPE Tubes for Extract Cleanup

Fast, Simple Sample Prep for Multiresidue Pesticide Analysis

Packaged in foil subpacks of 10 for enhanced protection and storage stability.

Multiple sorbents are used to extract different types of interferences.

- MgSO4 removes excess water
- · PSA removes sugars, fatty acids, organic acids, and anthocyanine pigments
- C18 removes nonpolar interferences

Description	Methods	qty.	cat.#
2 mL Micro-Centrifuge Tubes for dSPE (cleanup of 1 mL extract)			
150 mg MgSO ₄ , 50 mg PSA, 50 mg C18	AOAC 2007.01	100-pk.	26125
PSA—primary and secondary amine			

Rxi®-5ms Columns (fused silica)

(low-polarity phase; Crossbond® diphenyl dimethyl polysiloxane)

- General-purpose columns for semivolatiles, phenols, amines, residual solvents, drugs of abuse, pesticides, PCB congeners (e.g., Aroclor mixes), solvent impurities.
- · Most inert column on the market.
- Tested and guaranteed for ultra-low bleed; improved signalto-noise ratio for better sensitivity and mass spectral integrity.
- Equivalent to USP G27 and G36 phases.

Description	temp. limits	qty.	cat.#	
30 m, 0.25 mm ID, 0.25 μm	-60 to 330/350 °C	ea.	13423	

OuEChERS Performance Standards Kit

- · Kit contains organochlorine, organonitrogen, organophosphorus, and carbamate pesticides commonly used on fruits and vegetables.
- Ideal for initial method evaluations and ongoing method performance validations.
- · Analytes are divided into three ampuls based on compatibility for maximum stability and shelf life.*
- Precise formulations improve data quality and operational efficiency; spend more time running samples and less time sourcing and preparing standards.

Contains 1 mL each of these mixtures. 31153: QuEChERS Performance Standard A 31154: QuEChERS Performance Standard B 31155: OuEChERS Performance Standard C



300 µg/mL each in acetonitrile/acetic acid (99.9:0.1), 1 mL/ampul. Blend equal volumes of all three ampuls for a 100 μ g/mL final solution.

cat.# 31152 (kit)

*When combining compounds with different functionalities, chemical stability can be an issue. The analytes in this kit are separated into three mixes to ensure maximum long-term storage stability. For analysis, a fresh working standard should be prepared by combining the three kit mixes in a 1:1:1 ratio to prepare a 100 $\mu g/mL$ working standard solution. Once blended, Restek does not recommend storing working standards or subsequent dilutions for future use.

For GC Analysis

Pesticide Residue Cleanup SPE Cartridges

- Convenient, multiple adsorbent beds in a single cartridge.
- For use in multiresidue pesticide analysis to remove matrix interferences.
- Excellent for cleanup of dietary supplement extracts.

SPE Cartridge	qty.	cat.#
6 mL Combo SPE Cartridge Packed with 500 mg CarboPrep 90/500 mg PSA, Polyethylene Frits	30-pk.	26194

PSA-primary and secondary amine





www.restek.com/cannabis visit



PESTICIDE ANALYSIS PRODUCTS (CONT.)

LC Multiresidue Pesticide Kit

- Accurately detect and quantify pesticides of global food safety concern in a wide range of fruits, vegetables, and other commodities by LC-MS/MS.
- Full kit contains 204 compounds of interest, covering many LC-determined pesticides listed by government agencies; individual ampuls also sold separately.
- Formulated and grouped for maximum long-term stability* and well-balanced chromatographic performance, even for early eluting compounds.
- Quantitatively tested to confirm composition; detailed support documentation provided.
- Optimized multiresidue pesticide method is offered free of charge; downloadable XLS file includes conditions and transition tables.
- Certified reference material (CRM) manufactured and QC-tested in Restek's ISO-accredited labs satisfies your ISO requirements.

Cat.# 31972: LC Multiresidue Pesticide Standard #1 (13 components)

Organophosphorus Compounds Acephate (30560-19-1) Carbaryl (Sevin) (63-25-2) Dicrotophos (141-66-2) Dimethoate (60-51-5) Dimethomorph (110488-70-5) Isocarbophos (24353-61-5) Methamidophos (10265-92-6) Mevinphos (7786-34-7) Monocrotophos (6923-22-4) Omethoate (1113-02-6) Temephos (Abate) (3383-96-8) Trichlorfon (Dylox) (52-68-6) Vamidothion (Vamidoate) (2275-23-2)

Cat.# 31973: LC Multiresidue Pesticide Standard #2 (16 components)

Carbamate/Uron Compounds Alanycarb (83130-01-2) Aldicarb (116-06-3) Aldicarb sulfone (1646-88-4) Aldicarb sulfoxide (1646-87-3) Benfuracarb (82560-54-1) Butocarboxim (34681-10-2) Butoxycarboxim (34681-23-7) Ethiofencarb (29973-13-5) Furathiocarb (65907-30-4) Methabenzthiazuron (18691-97-9) Methiocarb (2032-65-7) Methomyl (16752-77-5) Oxamyl (23135-22-0) Tebuthiuron (34014-18-1) Thidiazuron (51707-55-2) Thiophanate-methyl

Cat.# 31974: LC Multiresidue Pesticide Standard #3 (38 components)

(23564-05-8)

Carbamate/Uron Compounds
Bendiocarb (22781-23-3)
Bifenarate (14987T-41-8)
Carbofutan (1563-66-2)
Chlorfuazurón (71422-67-8)
Chlorfuazurón (71422-67-8)
Chlorfuigurón (15545-48-9)
Cyclurón (2163-69-1)
Diethofencarb (87130-20-9)
Diflubenzurón (35367-38-5)
Dioxacarb (6988-21-2)

Diuron (330-54-1) Fenobucarb (BPMC) (3766-81-2) Fenoxycarb (79127-80-3) Fenuron (101-42-8) Flufenoxuron (101463-69-8) Fluometuron (2164-17-2) Forchlorfenuron (68157-60-8) Hexaflumuron (86479-06-3) 3-Hydroxycarbofuran (16655-82-6) Indoxacarb (173584-44-6) Iprovalicarb (140923-17-7) Isoprocarb (2631-40-5) Isoproturon (34123-59-6) Linuron (330-55-2) Lufenuron (103055-07-8) Metobromuron (3060-89-7) Monolinuron (1746-81-2) Neburon (555-37-3) Novaluron (116714-46-6) Pirimicarb (23103-98-2) Promecarb (2631-37-0) Propham (122-42-9) Propoxur (Baygon) (114-26-1) Pyraclostrobin (175013-18-0) Siduron (1982-49-6) Teflubenzuron (83121-18-0) Thiobencarb (28249-77-6) Triflumuron (64628-44-0)

Cat.# 31975: LC Multiresidue Pesticide Standard #4 (63 components)

Organonitrogen Compounds Abamectin (71751-41-2) Acetamiprid (135410-20-7) Ametryn (834-12-8) Amitraz (33089-61-1) Azoxystrobin (131860-33-8) Benalaxyl (71626-11-4) Benzoximate (29104-30-1) Boscalid (188425-85-6) Butafenacil (134605-64-4) Carbetamide (16118-49-3) Carfentrazone-ethyl (128639-02-1) Chlorantraniliprole (500008-45-7) Clofentezine (74115-24-5) Cymoxanil (57966-95-7) Cyprodinil (121552-61-2) Cyromazine (66215-27-8) Dimoxystrobin (149961-52-4) Dinotefuran (165252-70-0) Doramectin (117704-25-3) Eprinomectin (123997-26-2)

Famoxadon (131807-57-3) Fenazaquin (120928-09-8) Fenhexamid (126833-17-8) Fenpyroximate (111812-58-9) Flonicamid (158062-67-0) Fluazinam** (79622-59-6) Fludioxonil (131341-86-1) Fluoxastrobin (361377-29-9) Flutolanil (66332-96-5) Furalaxyl (57646-30-7) Halofenozide (112226-61-6) Imazalil (35554-44-0) Imidacloprid (138261-41-3) Ivermectin (70288-86-7) Kresoxim-methyl (143390-89-0) Mandipropamid (374726-62-2) Mepanipyrim (110235-47-7) Mepronil (55814-41-0) Metaflumizone (139968-49-3) Metalaxyl (57837-19-1) Methoxyfenozide (161050-58-4) Moxidectin (113507-06-5) Myclobutanil (88671-89-0) Nitenpyram (120738-89-8) Oxadixyl (77732-09-3) Picoxystrobin (117428-22-5) Piperonyl butoxide (51-03-6) Prochloraz (67747-09-5) Prometon (1610-18-0) Pymetrozine (123312-89-0) Pyracarbolid (24691-76-7) Pyrimethanil (53112-28-0) Pyriproxyfen (95737-68-1) Quinoxyfen (124495-18-7) Rotenone (83-79-4) Secbumeton (26259-45-0) Spiroxamine (118134-30-8) Tebufenozide (112410-23-8) Tebufenpyrad (119168-77-3) Terbumeton (33693-04-8) Triadimefon (43121-43-3) Trifloxystrobin (141517-21-7) Zoxamide (156052-68-5)

Cat.# 31976: LC Multiresidue Pesticide Standard #5 (30 components)

Organonitrogen Compounds Acibenzolar-S-methyl (135158-54-2) Bupirimate (41483-43-6) Buprofezin (69327-76-0) Carboxin (5234-68-4) Clethodim (99129-21-2) Clothianidin (210880-92-5) Cyazofamid (120116-88-3)

Ethiprole (181587-01-9) Ethofumesate (26225-79-6) Fenamidone (161326-34-7) Fipronil (120068-37-3) Flubendimide (272451-65-7) Flufenacet (Fluthiamide) (142459-58-3) Hexythiazox (78587-05-0) Mefenacet (73250-68-7) Mesotrione (104206-82-8) Methoprotryne (841-06-5) Metribuzin (21087-64-9) Prometryne (7287-19-6) Propargite (2312-35-8) Prothioconazole (178928-70-6) Pyridaben (96489-71-3) Simetryn (1014-70-6) Sulfentrazone (122836-35-5) Terbutryn (886-50-0) Thiabendazole (148-79-8) Thiacloprid (111988-49-9) Thiamethoxam (153719-23-4) Thiofanox (39196-18-4) Tricyclazole (Beam) (41814-78-2)

Cat.# 31977: LC Multiresidue Pesticide Standard #6 (28 components)

Organonitrogen Compounds Baycor (Bitertanol) (55179-31-2) Bromuconazole (116255-48-2) Cyproconazole (113096-99-4) Diclobutrazol (75736-33-3) Difenoconazole (119446-68-3) Diniconazole (83657-24-3) Epoxiconazole (133855-98-8) Etaconazole (60207-93-4) Ethirimol (23947-60-6) Etoxazole (153233-91-1) Fenarimol (60168-88-9) Fenbuconazole (114369-43-6) Fluquinconazole (136426-54-5) Flusilazole (85509-19-9) Flutriafol (76674-21-0) Fuberidazole (3878-19-1) Hexaconazole (79983-71-4) Ipconazole (125225-28-7) Metconazole (125116-23-6) Nuarimol (63284-71-9) Paclobutrazol (76738-62-0) Penconazole (66246-88-6) Propiconazole (Tilt) (60207-90-1)

Tebuconazole (107534-96-3) Tetraconazole (112281-77-3) Triadimenol (55219-65-3) Triflumizole (68694-11-1) Triticonazole (131983-72-7)

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Reference Materials

Cat.# 31978: LC Multiresidue Pesticide Standard #7 (7 components)

Organonitrogen Compounds
Emamectin-benzoate
(155569-91-8)
Fenpropimorph (67564-91-4)
Spirodiclofen (148477-71-8)
Spinosad (168316-95-8)
Spirotetramat (203313-25-1)
Spinetoram (J&L) (187166-40-1)
Spiromesifen (283594-90-1)

Cat.# 31979: LC Multiresidue Pesticide Standard #8 Organonitrogen Compounds

Organonitrogen Compounds Hydramethylnon (67485-29-4)

Cat.# 31980: LC Multiresidue Pesticide Standard #9 (7 components)

Carbamate/Uron Compounds Aminocarb (2032-59-9) Desmedipham (13684-56-5) Formetanate HCL (23422-53-9) Mexacarbate (Zectran) (315-18-4) Monceren (Pencycuron) (66063-05-6) Phenmedipham (13684-63-4) Propamocarb free base (24579-73-5)

Cat.# 31981: LC Multiresidue Pesticide Standard #10 Carbamate/Uron Compounds

Carbendazim (10605-21-7)



Contains 1 mL each of these mixtures. cat.# 31971 (kit)

Quantity discounts not available.

* NOTE: When combining a large number of compounds with different chemical functionalities, mix stability can be an issue. In formulating these standards, we extensively studied the 204 compounds involved, then grouped them into as few mixes as possible while still ensuring maximum long-term stability and reliability. For quantitative analysis, we recommend analyzing each mix separately to ensure accurate results for every compound.

** NOTE: In this standard, fluazinam should only be used for qualitative analysis. A single-component standard (cat.# 31982) is available for quantitative analysis



GC Multiresidue Pesticide Kit

- Accurately identify and quantify pesticide residues by GC-MS/MS in fruits, vegetables, botanicals, and herbals like tea, ginseng, ginger, Echinacea, and dietary supplements.
- Comprehensive 203-compound kit covers food safety lists by the FDA, USDA, and other global governmental agencies; individual ampuls also sold separately.
- Formulated and grouped for maximum long-term stability* and well-balanced chromatographic performance, even for early eluting compounds.
- Quantitatively tested to confirm composition; detailed support documentation provided.
- Certified reference material (CRM) manufactured and QC-tested in Restek's ISO-accredited labs satisfies your ISO requirements.



Reference Materials

Cat.# 32563: GC Multiresidue Pesticide Standard #1 (16 components)

Organophosphorus Compounds Azinphos ethyl (2642-71-9) Azinphos-methyl (86-50-0) Chlorpyrifos (2921-88-2) Chlorpyrifos methyl (5598-13-0) Diazinon (333-41-5) EPN (2104-64-5) Fenitrothion (122-14-5) Isazophos (42509-80-8) Phosalone (2310-17-0) Phosmet (732-11-6) Pirimiphos ethyl (23505-41-1) Pirimiphos methyl (29232-93-7) Pyraclofos (77458-01-6) Pyrazophos (13457-18-6) Pyridaphenthion (119-12-0) Quinalphos (13593-03-8)

Cat.# 32564: GC Multiresidue Pesticide Standard #2

(40 components) Organochlorine Compounds Aldrin (309-00-2) alpha-BHC (319-84-6) beta-BHC (319-85-7) delta-BHC (319-86-8) gamma-BHC (Lindané) (58-89-9)Chlorbenside (103-17-3) cis-Chlordane (5103-71-9) trans-Chlordane (5103-74-2) Chlorfenson (Ovex) (80-33-1) Chloroneb (2675-77-6) 2,4'-DDD (53-19-0) 4,4'-DDD (72-54-8) 2.4'-DDE (3424-82-6) 4,4'-DDE (72-55-9) 2,4'-DDT (789-02-6) 4.4'-DDT (50-29-3) 4,4'-Dichlorobenzophenone (90-98-2) Dieldrin (60-57-1) Endosulfan I (959-98-8) Endosulfan II (33213-65-9) Endosulfan ether (3369-52-6) Endosulfan sulfate (1031-07-8) Endrin (72-20-8) Endrin aldehyde (7421-93-4) Endrin ketone (53494-70-5) Ethylan (Perthane) (72-56-0) Fenson (80-38-6) Heptachlor (76-44-8) Heptachlor epoxide (Isomer B) (1024-57-3)Hexachlorobenzene (118-74-1)

Isodrin (465-73-6)

2,4'-Methoxychlor (30667-99-3) 4,4'-Methoxychlor olefin (2132-70-9) Mirex (2385-85-5) cis-Nonachlor (5103-73-1) trans-Nonachlor (39765-80-5) Pentachloroanisole (1825-21-4) Pentachlorobenzene (608-93-5) Pentachlorothioanisole (1825-19-0) Tetradifon (116-29-0)

Cat.# 32565: GC Multiresidue Pesticide Standard #3

(25 components) Organonitrogen Compounds Benfluralin (1861-40-1) Biphenyl (92-52-4) Chlorothalonil (1897-45-6) Dichlofluanid (1085-98-9) Dichloran (99-30-9) 3,4-Dichloroaniline (95-76-1) 2,6-Dichlorobenzonitrile (Dichlobenil) (1194-65-6) Diphenylamine (122-39-4) Ethalfluralin (55283-68-6) Fluchloralin (33245-39-5) Isopropalin (33820-53-0) Nitralin (4726-14-1) Nitrofen (1836-75-5) Oxyfluorfen (42874-03-3) Pendimethalin (40487-42-1) Pentachloroaniline (527-20-8) Pentachlorobenzonitrile (20925-85-3) Pentachloronitrobenzene (Quintozene) (82-68-8) Prodiamine (29091-21-2) Profluralin (26399-36-0) 2,3,5,6-Tetrachloroaniline (3481-20-7) Tetrachloronitrobenzene (Tecnazene) (117-18-0) THPI (Tetrahydrophthalimide) (1469-48-3) Tolylfluanid (731-27-1) Trifluralin (1582-09-8)

Cat.# 32566: GC Multiresidue Pesticide Standard #4

(28 components) Organonitrogen Compounds Acetochlor (34256-82-1) Alachlor (15972-60-8) Allidochlor (93-71-0) Clomazone (Command) (81777-89-1) Cycloate (1134-23-2) Diallate (cis and trans)

(2303-16-4)Dimethachlor (50563-36-5) Diphenamid (957-51-7) Fenpropathrin (39515-41-8) Fluquinconazole (136426-54-5) Flutolanil (66332-96-5) Linuron (330-55-2) Metazachlor (67129-08-2) Methoxychlor (72-43-5) Metolachlor (51218-45-2) N-(2,4-Dimethylphenyl) formamide (60397-77-5) Norflurazon (27314-13-2) Oxadiazon (19666-30-9) Pebulate (1114-71-2) Pretilachlor (51218-49-6) Prochloraz (67747-09-5) Propachlor (1918-16-7) Propanil (709-98-8) Propisochlor (86763-47-5) Propyzamide (23950-58-5) Pyridaben (96489-71-3) Tebufenpyrad (119168-77-3) Triallate (2303-17-5)

Cat.# 32567: GC Multiresidue Pesticide Standard #5 (34 components)

Organonitrogen Compounds Atrazine (1912-24-9) Bupirimate (41483-43-6) Captafol (2425-06-1) Captan (133-06-2) Chlorfenapyr (122453-73-0) Cyprodinil (121552-61-2) Etofenprox (80844-07-1) Etridiazole (2593-15-9) Fenarimol (60168-88-9) Fipronil (120068-37-3) Fludioxonil (131341-86-1) Fluridone (Sonar) (59756-60-4) Flusilazole (85509-19-9) Flutriafol (76674-21-0) Folpet (133-07-3) Hexazinone (Velpar) (51235-04-2) Iprodione (36734-19-7) Lenacil (2164-08-1) MGK-264 (113-48-4) Myclobutanil (88671-89-0) Paclobutrazol (76738-62-0) Penconazole (66246-88-6) Procymidone (32809-16-8) Propargite (2312-35-8) Pyrimethanil (53112-28-0) Pyriproxyfen (95737-68-1) Tebuconazole (107534-96-3) Terbacil (5902-51-2)

Triadimefon (43121-43-3) Triadimenol (55219-65-3) Tricyclazole (Beam) (41814-78-2) Triflumizole (68694-11-1) Vinclozolin (50471-44-8)

Cat.# 32568: GC Multiresidue Pesticide Standard #6 (18 components)

Synthetic Pyrethroid Compounds Acrinathrin (101007-06-1) Anthraguinone (84-65-1) Bifenthrin (82657-04-3) Bioallethrin (584-79-2) Cyfluthrin (68359-37-5) lambda-Cyhalothrin (91465-08-6) Cypermethrin (52315-07-8) Deltamethrin (52918-63-5) Fenvalerate (51630-58-1) Flucythrinate (70124-77-5) tau-Fluvalinate (102851-06-9) cis-Permethrin (61949-76-6) trans-Permethrin (61949-77-7) Phenothrin (cis & trans) (26002-80-2) Resmethrin (10453-86-8) Tefluthrin (79538-32-2) Tetramethrin (7696-12-0) Transfluthrin (118712-89-3)

Cat.# 32569: GC Multiresidue Pesticide Standard #7 (10 components)

Herbicide Methyl Esters Acequinocyl (57960-19-7) Bromopropylate (18181-80-1) Carfentrazone ethyl (128639-02-1) Chlorobenzilate (510-15-6) Chlorpropham (101-21-3) Chlozolinate (84332-86-5) DCPA methyl ester (Chlorthal-dimethyl) (1861-32-1) Fluazifop-p-butyl (79241-46-6) Metalaxyl (57837-19-1) 2-Phenylphenol (90-43-7)

Cat.# 32570: GC Multiresidue Pesticide Standard #8 (24 components)

Organophosphorus Compounds Bromfenvinfos-methyl (13104-21-7) Bromfenvinphos (33399-00-7) Bromophos ethyl (4824-78-6) Bromophos methyl (2104-96-3) Carbophenothion (786-19-6) Chlorfenvinphos (470-90-6) Chlorthiophos (60238-56-4) Coumaphos (56-72-4) Edifenphos (17109-49-8) Ethion (563-12-2) Fenamiphos (22224-92-6) Fenchlorphos (Ronnel) (299-84-3) Fenthion (55-38-9) Iodofenphos (18181-70-9) Leptophos (21609-90-5) Malathion (121-75-5) Methacrifos (62610-77-9) Profenofos (41198-08-7) Prothiofos (34643-46-4) Sulfotepp (3689-24-5) Sulprofos (35400-43-2) Terbufos (13071-79-9) Tetrachlorvinfos (22248-79-9) Tolclofos-methyl (57018-04-9)

Cat.# 32571: GC Multiresidue Pesticide Standard #9 (8 components)

Organophosphorus Compounds Disulfoton (298-04-4) Fonofos (944-22-9) Methyl parathion (298-00-0) Mevinphos (7786-34-7) Parathion (Ethyl parathion) (56-38-2) Phorate (298-02-2) Piperonyl butoxide (51-03-6) Triazophos (24017-47-8)



Contains 1 mL each of these mixtures. cat.# 32562 (kit)

* NOTE: When combining a large number of compounds with different chemical functionalities, mix stability can be an issue. In formulating these standards, we extensively studied the 203 compounds involved, then grouped them into as few mixes as possible while still ensuring maximum long-term stability and reliability. For quantitative analysis, we recommend analyzing each mix separately to ensure accurate results for every compound.

Terbuthylazine (5915-41-3)

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Growing Analytical Solutions for Medical Cannabis Labs

PRODUCTS AND EXPERTISE FOR **ACCURATE, RELIABLE RESULTS**

EVERY TIME

Explore our products and resources and grow your medical cannabis testing business with trusted analytical solutions from Restek. We're proud to have helped medical cannabis labs establish sound analytical practices from the beginning, and we will continue to be there for you every step of the way as the testing landscape changes. Our commitment to medical marijuana analysis extends beyond a trusted, well-rounded product line, so if you're struggling with any part of your cannabinoid analyses, contact our in-house experts for assistance.

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Lit. Cat.# FFBR2073A-UNV



Technical Article

High-Quality Analysis of Pesticides in Cannabis

Using QuEChERS, Cartridge SPE Cleanup, and GCxGC-TOFMS

By Jack Cochran, Julie Kowalski, Sharon Lupo, Michelle Misselwitz, and Amanda Rigdon

- Quickly and effectively extract medical marijuana samples for pesticide analysis.
- Cartridge SPE cleanup of dirty extracts improves GC inlet and column lifetimes.
- Selective GC columns increase accuracy of pesticide determinations for complex samples.

Over 20 states in the U.S. have legalized the use of recreational or medical cannabis because of therapeutic benefits for ailments such as cancer, multiple sclerosis, and ALS. Dosing methods include smoking or vaporizing and baked goods. Unlike other prescribed medicines regulated by U.S. FDA, marijuana is a Schedule 1 drug and is illegal on the federal level. As a result, medical cannabis patients have no safety assurances for their medication, which could contain harmful levels of pesticide residues. Currently, medical marijuana pesticide residue analysis methods are poorly defined and challenging to develop due to matrix complexity and a long list of potential target analytes.

In order to address matrix complexity, we combined a simple QuEChERS extraction approach with cartridge SPE (cSPE) cleanup, followed by GCxGC-TOFMS. Acceptable recoveries were obtained for most pesticides, and incurred pesticide residues were detected in some of the illicit marijuana samples used for method development.

QuEChERS Extraction Saves Time and Reduces Hazardous Solvent Use

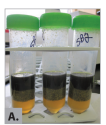
Trace residue extraction procedures from dry materials like medical cannabis typically involve large amounts of solvent, long extraction times, and tedious concentration steps similar to the Soxhlet procedure or multiresidue methods from the Pesticide Analytical Manual. QuEChERS, with its simple 10 mL acetonitrile shake extraction and extract partitioning with salts and centrifugation, offers time savings, glassware use reduction, and lower solvent consumption.

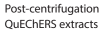
Water was added to finely ground, dry cannabis samples to increase QuEChERS extraction efficiency, especially for more polar pesticides. A vortex mixer was used to shake the solvent and sample for at least 30 minutes prior to extract partitioning. When finished, it was easy to transfer the supernatant from the QuEChERS extraction tube for subsequent cSPE cleanup prior to analysis with GC or LC (Figure 1).

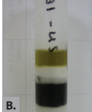
Cartridge SPE Cleanup Improves **GC Inlet Uptime**

Injecting chlorophyll-laden extracts into a GC gives reduced recoveries for less volatile pesticides, and results in degradation of sensitive pesticides like DDT and Dicofol (Table I). SPE cleanup with a 500 mg graphitized carbon black/500 mg PSA cartridge removes chlorophyll and traps fatty acids that interfere with qualitative pesticide identification and bias quantification. cSPE has increased sorbent capacity over dispersive SPE for thorough cleanup of complex extracts.

Figure 1: A quick and easy QuEChERS extraction, combined with cSPE, effectively prepared extracts for pesticide residue analysis from highly complex marijuana samples.







QuEChERS extracts loaded on SPE cartridge



Final extract



RESTEK Pure Chromatography

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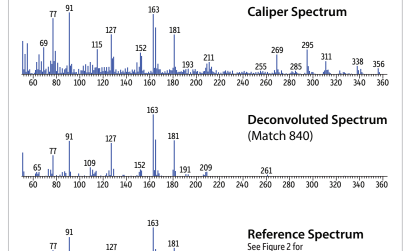
Orthogonal GC Columns Increase Separation Power for More Accurate Pesticide Results

GCxGC is a powerful multidimensional approach that gives two independent separations in one instrumental analysis. An Rxi®-5Sil MS and Rtx®-200 column combination distributes pesticides broadly in both dimensions, providing a highly orthogonal GCxGC system. More important though is separating pesticides from potential isobaric matrix interferences, as seen in the surface plot for the insecticide cypermethrin (Figure 2). Cypermethrin gas chromatographs as four isomers, and all would have experienced qualitative interference and quantitative bias from peaks in the foreground of the surface plot had only 1-dimensional GC been used. With GCxGC-TOFMS, cypermethrin was unequivocally identified in a marijuana sample at a low ppm level (Figure 3).

Summary

QuEChERS and cSPE produced usable extracts from highly complex cannabis samples for high-quality pesticide residue analysis. The multidimensional separation power of GCxGC-TOFMS was then used to correctly identify and quantify pesticides in these complex extracts.

Figure 3: Positive mass spectral identification of incurred cypermethrin in illicit marijuana.



Acknowledgment: Randy Hoffman, a Police Evidence Technician at The Pennsylvania State University (PSU), supplied the seized marijuana samples while overseeing their handling. Frank Dorman at PSU assisted with QuEChERS extractions.

209

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Initially published in Restek® Advantage.

115

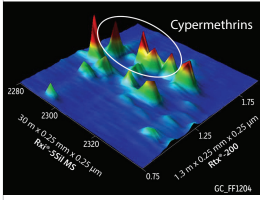
120 140 160 180 200 220

Table I: Pesticide recoveries for a QuEChERS extract of cannabis give higher results when cSPE is used for cleanup. Dicofol and DDT are degraded in the inlet for the dirtier extract, yielding high DDD results.

Pesticide	Classification	With cSPE Cleanup (%)	Without cSPE Cleanup (%)
4,4'-DDD	Organochlorine	83	230
4,4'-DDT	Organochlorine	77	9
Bifenthrin	Pyrethroid	86	89
Dicofol	Organochlorine	84	ND
Azinphos methyl	Organophosphorus	79	53
trans-Permethrin	Organochlorine	68	17
Pyraclostrobin	Strobilurin	73	19
Fluvalinate	Pyrethroid	72	23
Difenoconazole	Triazole	67	21
Deltamethrin	Pyrethroid	68	20
Azoxystrobin	Strobilurin	72	27

ND = no peak detected

Figure 2: GCxGC-TOFMS and orthogonal Rxi®-5Sil MS and Rtx®-200 columns allow incurred cypermethrins in a marijuana extract to be separated from interferences (m/z 163 quantification ion).



Peaks	RT 1 (sec.)	RT 2 (sec.)
 Cypermethrin 1 	2292	1.50
2. Cypermethrin 2	2304	1.54
Cypermethrin 3	2310	1.53
4. Cypermethrin 4	2313	1.58

Column: Rxi \circ -5Sil MS 30 m, 0.25 mm ID, 0.25 μ m (cat.# 13623), Rtx \circ -200 1.3 m, 0.25 mm ID, 0.25 μ m (cat.# 15124); Sample: Diluent: Toluene; Injection: Inj. Vol.: 1 μ L splitless (hold 1 min); Liner: Sky \circ ν m single taper w/wool (cat.# 23303.1); Inj. Temp: 250 °C; Purge Flow: 40 mL/min; Oven: Oven: Oven Temp: Rxi \circ -5Sil MS: 80 °C (hold 1 min) to 310 °C at 5 °C/min, Rtx \circ -200: 85 °C hold 1 min) to 315 °C at 5 °C/min; Carrier Gas: He, corrected constant flow (2 mL/min); Modulation: Modulator Temp. Offset: 20 °C; Second Dimension Separation Time: 3 sec.; Hot Pulse Time: 0.9 sec.; Cool Time between Stages: 0.6 sec.; Instrument: LECO Pegasus 4D GCxGC-TOFMS; For complete conditions, visit www.restek.com and enter GC. FF1204 in the search.



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instrument conditions.

260 280

GC FF1206

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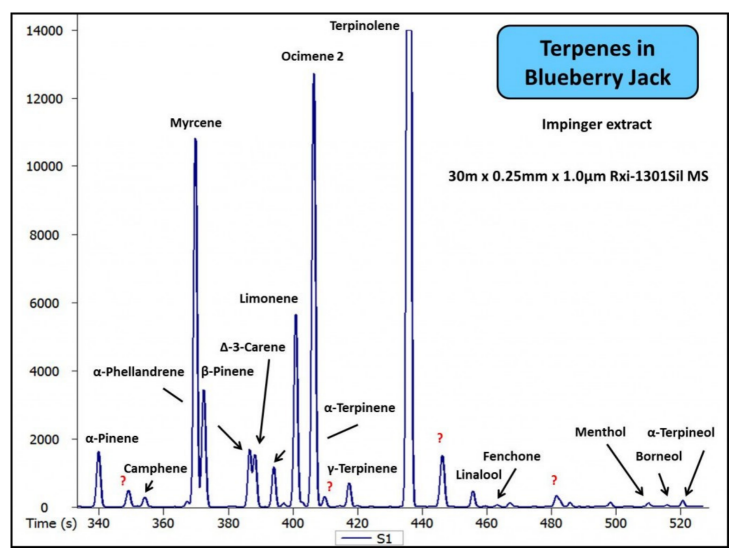
« Early Eluting Terpenes – GC – Medical Cannabis Keep up with ChromaBLOGraphy — new subscription option added »

Terpenes in Blueberry Jack Medical Cannabis – GC – More Identified

March 26th, 2014 by Jack Cochran

Based on acquisition of new terpene standards I was able to better profile the Blueberry Jack medical cannabis impinger sample on the beta-version 30m x 0.25mm x 1.0µm Rxi-1301Sil MS GC column. Check it out...

I'm looking for suggestions on terpene identification for the ones marked by "?" in the chromatogram below. Help, please!



This entry was posted on Wednesday, March 26th, 2014 at 10:37 pm and is filed under New GC Columns, Medical Marijuana. You can follow any responses to this entry through the RSS 2.0 feed. You can leave a response, or trackback from your own site.

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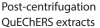
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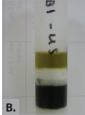
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QuEChERS extracts loaded on SPE cartridge



Final extract



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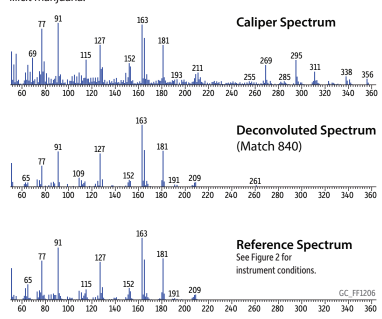
Orthogonal GC Columns Increase Separation Power for More Accurate Pesticide Results

GCxGC is a powerful multidimensional approach that gives two independent separations in one instrumental analysis. An Rxi®-5Sil MS and Rtx®-200 column combination distributes pesticides broadly in both dimensions, providing a highly orthogonal GCxGC system. More important though is separating pesticides from potential isobaric matrix interferences, as seen in the surface plot for the insecticide cypermethrin (Figure 2). Cypermethrin gas chromatographs as four isomers, and all would have experienced qualitative interference and quantitative bias from peaks in the foreground of the surface plot had only 1-dimensional GC been used. With GCxGC-TOFMS, cypermethrin was unequivocally identified in a marijuana sample at a low ppm level (Figure 3).

Summary

QuEChERS and cSPE produced usable extracts from highly complex cannabis samples for high-quality pesticide residue analysis. The multidimensional separation power of GCxGC-TOFMS was then used to correctly identify and quantify pesticides in these complex extracts.

Figure 3: Positive mass spectral identification of incurred cypermethrin in illicit marijuana.



Acknowledgment: Randy Hoffman, a Police Evidence Technician at The Pennsylvania State University (PSU), supplied the seized marijuana samples while overseeing their handling. Frank Dorman at PSU assisted with QuEChERS extractions.

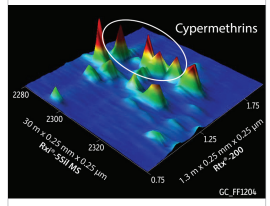
Initially published in Restek® Advantage.

Table 1: Pesticide recoveries for a QuEChERS extract of cannabis give higher results when cSPE is used for cleanup. Dicofol and DDT are degraded in the inlet for the dirtier extract, yielding high DDD results.

Pesticide	Classification	With cSPE Cleanup (%)	Without cSPE Cleanup (%)
4,4'-DDD	Organochlorine	83	230
4,4'-DDT	Organochlorine	77	9
Bifenthrin	Pyrethroid	86	89
Dicofol	Organochlorine	84	ND
Azinphos methyl	Organophosphorus	79	53
trans-Permethrin	Organochlorine	68	17
Pyraclostrobin	Strobilurin	73	19
Fluvalinate	Pyrethroid	72	23
Difenoconazole	Triazole	67	21
Deltamethrin	Pyrethroid	68	20
Azoxystrobin	Strobilurin	72	27

ND = no peak detected

Figure 2: GCxGC-TOFMS and orthogonal Rxi®-5Sil MS and Rtx®-200 columns allow incurred cypermethrins in a marijuana extract to be separated from interferences (m/z 163 quantification ion).



Peaks		RT 1 (sec.)	RT 2 (sec.
1.	Cypermethrin 1	2292	1.50
2.	Cypermethrin 2	2304	1.54
3.	Cypermethrin 3	2310	1.53
4.	Cypermethrin 4	2313	1.58

 $\begin{array}{l} \textbf{Column:} \ Rxi@-5Sil\ MS\ 30\ m, 0.25\ mm\ ID, 0.25\ \mum\ (cat.\#\ 13623), \ Rtx@-200\ 1.3\ m, 0.25\ mm\ ID, 0.25\ \mum\ (cat.\#\ 136124); \ \textbf{Sample:} \ Diluent:\ Toluene;\ \textbf{Injection:} \ lnj.\ Vol.:\ 1\ \muL\ splitless\ (hold\ 1\ min);\ Liner:\ Sky@\ 4m\ msingle\ taper\ w/wool\ (cat.\#\ 23303.1);\ lnj.\ Temp:\ 250°C;\ Purge\ Flow:\ 40\ mL/min;\ \textbf{Oven:}\ Oven:\ Oven:\ Temp:\ Rxi@-5Sil\ MS:\ 80°C\ (hold\ 1\ min)\ to\ 310°C\ at\ 5°C/min;\ Carrier\ Gas:\ He,\ corrected\ constant\ flow\ (2\ mL/min);\ \textbf{Modulation:}\ Modulator:\ Temp.\ 0ffset:\ 20°C;\ Second\ Dimension\ Separation\ Time:\ 3\ sec.;\ Hot\ Pulse\ Time:\ 0.9\ sec.;\ Cool\ Time\ between\ Stages:\ 0.6\ sec.;\ Instrument:\ LECO\ Pegasus\ 4D\ GCxGC-ToFMS;\ For\ complete\ conditions,\ visit\ www.restek.com\ and\ enter\ GC_F1204\ in\ the\ search. \end{array}$



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Technical Article

Don't Overestimate Cannabidiol During Medical Cannabis Potency Testing by Gas Chromatography

By Jack Cochran

Accurate potency testing of medical cannabis with gas chromatography (GC) depends principally on choosing a column with the right selectivity; otherwise, coelutions between cannabinoids of interest may cause error in potency measurements. Cannabidiol is one of the chief cannabinoids with pharmacological value and provides relief against nausea, anxiety, and inflammation. Potency testing for medical marijuana is often done using "5-type" GC columns since they are commonly available in most labs. However, on 5-type columns cannabidiol can coelute with cannabichromene, a compound that likely also has medical value and is increasingly becoming part of potency testing. To identify and report both of these compounds accurately, a GC column with a different stationary phase is needed.

Proper Column Choice Results in More Accurate Potency Data

As shown in Figure 1, cannabinoids are aromatic compounds, meaning they will likely be better separated on a column that contains aromatics in the stationary phase because these stationary phases are more selective for aromatic-containing analytes. A fully non-aromatic stationary phase, like a "1-type" (100% dimethyl polysiloxane) column is not appropriate for this analysis since cannabichromene (CBC) and cannabidiol (CBD) will coelute completely. While 5-type columns (5% phenyl) contain some aromatic component, they generally also produce coelutions for cannabichromene and cannabidiol, depending on the conditions used. At best, CBC and CBD can be only partially resolved on 15 m 5% phenyl columns. Much better separations are obtained on higher phenyl-content phases, such as Rxi*-35Sil MS (35% phenyl type) and Rxi*-17Sil MS (50% phenyl type) columns, as they offer excellent selectivity for aromatic cannabinoids. Not only do both columns resolve cannabichromene and cannabidiol, the chromatograms in Figures 2 and 3 demonstrate that they also separate delta-8-tetrahydrocannabinol (d8-THC), delta-9-tetrahydrocannabinol (d9-THC), cannabigerol (CBG), and cannabinol (CBN). Although both columns perform well, the Rxi*-35Sil MS column is recommended because of the slightly faster analysis time and greater space overall between the peaks of interest.

While stationary phase selectivity is the most important factor in choosing a GC column for cannabinoid analysis, there are some additional aspects of this work that will benefit labs doing medical marijuana potency testing. First, cost savings were achieved by using a 15 m column. When a column with the proper selectivity is used, a 15 m column easily provides the separating power needed for this analysis at about half the cost of a 30 m column. Also, the 0.25 mm x 0.25 µm format has good sample loading capacity and is robust, especially when a proper split injection is used with a Sky* Precision* split liner with wool. Finally, hydrogen carrier gas was used here instead of helium. Using hydrogen provides a faster analysis, increasing sample throughput. Hydrogen carrier gas is a convenient way to speed up run times, increase productivity, and reduce the cost and availability concerns associated with using helium carrier gas.



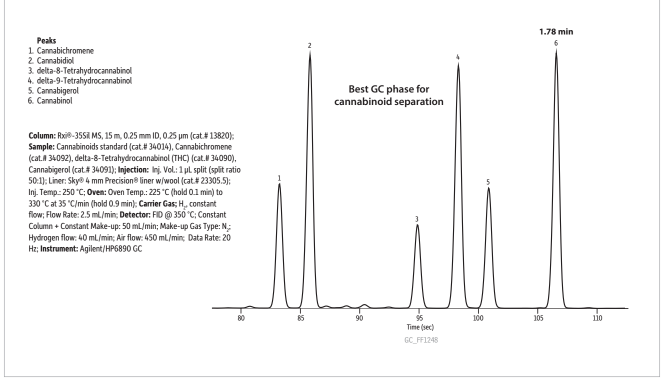
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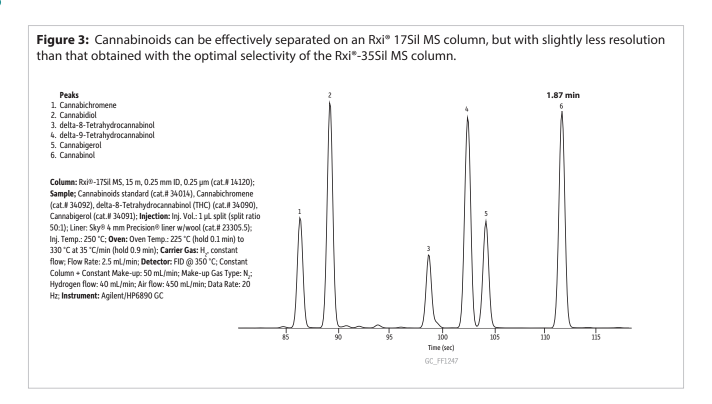


Figure 1: Since cannabinoids are aromatic compounds, a GC column that contains aromatics in the stationary phase will provide much better separations than a column with a non-aromatic phase.

Figure 2: The Rxi®-35Sil MS column provides both the best separation and the fastest analysis time, making it the ideal GC column choice for medical cannabis potency testing.

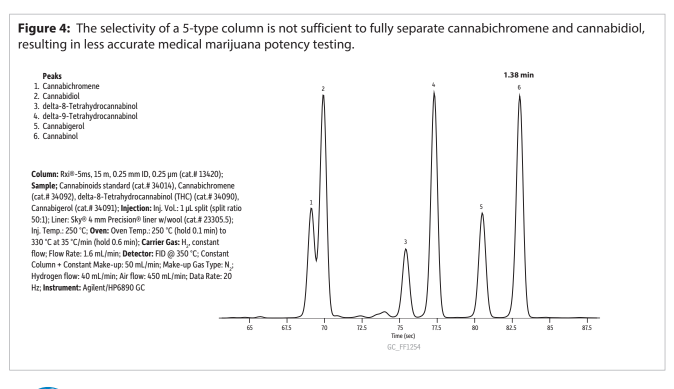


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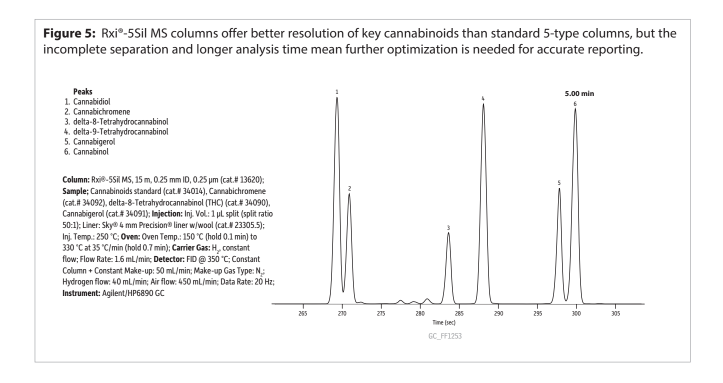


Adjusting Conditions for 5-Type Columns

While using an Rxi $^{\circ}$ -35Sil MS column provides the best selectivity and speed for cannabinoid analysis, cannabidiol potency can be determined in medical cannabis using a 5-type column under certain conditions. If you already have a 5-type column for this work, you can vary the GC conditions, especially carrier flow and oven temperature program, and still separate cannabichromene and cannabidiol, just not as quickly or easily as with the Rxi $^{\circ}$ -35Sil MS column. Figures 4 and 5 show this analysis on Rxi $^{\circ}$ -5ms and Rxi $^{\circ}$ -5Sil MS columns, respectively. Again, the 0.25 mm x 0.25 µm format was used here because it offers better efficiency than wider bore columns (e.g., 0.32 mm and 0.53 mm IDs), which may not separate cannabichromene and cannabidiol under any operational conditions.







Note that even though these are both 5-type columns, the elution order of cannabichromene and cannabidiol changed. This is due to two things. The first is that Rxi*-5ms and Rxi*-5Sil MS columns differ slightly in selectivity for certain compounds; even though they are both considered 5-type columns, they contain different stationary phases that retain some compounds differently. The second reason is that the GC oven programs are different for the columns, which means that the compounds are eluting at different temperatures. You may be able to further optimize the separation of cannabichromene and cannabidiol on a 5-type column, but the selectivity and faster analysis that can be obtained using a high-phenyl content Rxi*-35Sil MS column make it ideal for potency determinations in medical cannabis.

To sum things up, proper column choice is essential for accurate and robust cannabis potency testing. Using the right column not only gives you more confidence in your potency values, but it also saves you time and money. Switching to hydrogen carrier gas can reduce your costs even further, while increasing sample throughput.

Visit www.restek.com/medical-cannabis for Restek® GC and LC columns, accessories, reference standards, and other products and resources for medical marijuana analysis.



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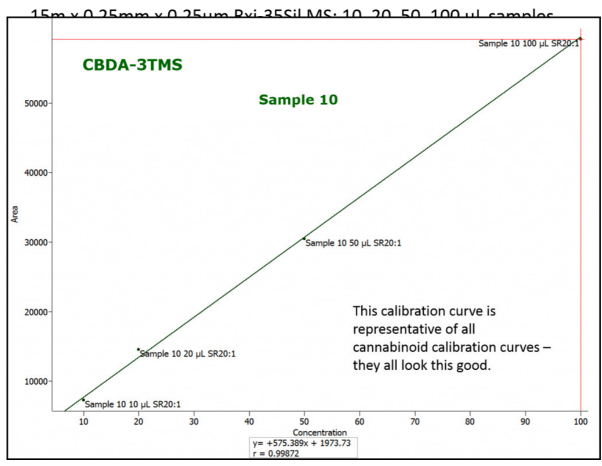
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Sample 10 Dilution Derivatization Linearity

Compound	Mol Wt	$t_{\rm R}$ sec	t _R min	r
CBDV-2TMS	430	94.6	1.58	0.9994
CBD-2TMS	458	121.0	2.02	0.9995
THCV-1TMS	358	128.2	2.14	0.9991
CBC-1TMS	386	148.2	2.47	0.9989
CBG-2TMS	460	149.9	2.50	0.9993
Δ ⁸ -THC-1TMS	386	156.4	2.61	0.9993
Δ ⁹ -THC-1TMS	386	160.7	2.68	0.9990
CBDA-3TMS	574	164.8	2.75	0.9987
CBN-1TMS	382	178.7	2.98	0.9988
Δ ⁹ -THCA-2TMS	502	212.3	3.54	0.9992



In addition to verifying that the derivatization reaction goes to completion in the presence of plant matrix, we also verified the procedure using several different samples which were generated at the same time as the sample shown in the figure above. Our preliminary work is still looking good, which is exciting, but what about all of the other matrices cannabis chemists have to work with? Well, we're planning on moving the work forward into edible matrices next, so stay tuned for an update!

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Foods, Flavors & Fragrances Applications

A Preliminary FET Headspace GC-FID Method for Comprehensive Terpene Profiling in Cannabis

By Amanda Rigdon, Corby Hilliard, and Jack Cochran

Abstract

This application note describes an FET headspace GC-FID method that was developed in hops for the analysis of terpenes in cannabis. Good chromatographic separation allowed quantification of critical compounds across the volatility range, including α -pinene, β -myrcene, α -humulene, β -caryophyllene, and caryophyllene oxide.

Introduction

In addition to cannabinoids, cannabis contains a suite of compounds known as terpenes. Terpenes are not only responsible for the characteristic aromas of cannabis strains, but they also are suspected to contribute to the therapeutic properties of cannabis. By themselves, terpenes have anti-inflammatory and anti-microbial properties, and they also reportedly contribute to an "entourage effect" with cannabinoids, modulating and/or enhancing their activity [1,2].

Because terpenes may contribute to the therapeutic effects of cannabis, there is a growing demand for analytical methods that profile terpenes in marijuana samples. In addition to analyzing terpenes for therapeutic purposes, terpenes can also be used as differentiators among cannabis strains and terpene profiles can be used for strain identification.

While relatively few terpenes have been studied for therapeutic purposes, cannabis strains can contain dozens of terpenes in varying levels. Of these, the primary compounds of interest include α -pinene, β -myrcene, α -humulene, and β -caryophyllene [2,3]. Accurately profiling these analytes and other emerging terpenes of interest depends heavily on separating them from potentially interfering compounds. When an interfering terpene, or other compound, coelutes with a terpene of interest, quantification will be compromised and, since many terpenes have the same molecular weight and share fragment ions, mass spectrometry cannot be relied upon to distinguish a terpene of interest from a coeluting interference terpene. The only way to accurately identify and quantify terpenes is to ensure that the terpenes of interest are chromatographically separated from all interfering compounds. GC is an excellent technique for accomplishing this.

Here we present a headspace gas chromatography–flame ionization detection (GC-FID) method for a comprehensive set of 38 terpenes found in cannabis. Since cannabis is illegal in Pennsylvania where this work was done, we developed the method using hops as a model system since they are related to cannabis and contain a similar suite of terpenes [2,3,4]. The headspace method presented here utilizes full evaporation technique (FET) sample preparation because cannabis product matrices are extremely varied and plant material will not dissolve in solvent. FET involves the use of a very small sample amount (10–50 mg), which effectively creates a single phase gas system in the headspace vial at equilibrium, making it ideal for this application [5,6,7]. Figure 1 illustrates the basic principle of headspace gas chromatography using FET. To achieve chromatographic separation, a 30 m x 0.25 mm x 1.4 μ m Rxi®-624Sil MS column was used. This column was chosen based on several factors. First, and most importantly, the cyano-based stationary phase of the Rxi®-624Sil MS has excellent selectivity for terpenes, making it ideal to effect a good separation for a large suite of these compounds. Second, in addition to its excellent selectivity for terpenes, the maximum temperature of this column is 320 °C, which allows for elution of some of the less volatile terpenes and matrix compounds that may be present in the headspace sample. Third, this GC column phase is also well-suited for residual solvent analysis, potentially minimizing the number of columns and instruments required by labs to test cannabis.



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Experimental

Sample Preparation

Pelletized hops from three strains (UK East Kent Golding, Citra, and Cascade) were purchased from HopUnion. The pelletized hops were first ground to a fine powder using an IKA® mill. Because the hops were already ground and pelletized, very little grinding was necessary. For cannabis plant material, it is recommended that samples be frozen prior to grinding or that grinding occur under liquid nitrogen. This keeps the samples cold during the grinding process, reducing loss of the more volatile terpenes such as α-pinene. 10 mg samples of each strain were then placed in headspace vials (Figure 2). An incubation temperature of 140 °C was used to ensure volatilization of all terpenes and terpenoids in the sample. This temperature was chosen because it is also sufficient to melt samples of cannabis concentrates. An incubation time of 30 minutes was used to ensure the establishment of equilibrium during incubation, which is required for reproducible, quantitative results.

Gas Chromatographic Conditions

Samples were analyzed on an Agilent® 6890 gas chromatograph equipped with a Tekmar® HT-3 headspace autosampler. A 30 m x 0.25 mm x 1.4 µm Rxi®-624Sil MS column was installed based on its selectivity for terpenes and because it could also be used for analysis of residual solvents in cannabis concentrates. A 1 mm straight Sky® inlet liner was used to limit the volume in the GC inlet. For headspace instruments, reducing the inlet volume increases efficiency by reducing band broadening during sample introduction. Greater efficiency maximizes peak separation, which is essential for this analysis. Complete chromatographic conditions are presented in Figure 4.

Figure 1: Setup and Basic Principle of FET Headspace Injection Coupled With GC-FID Analysis

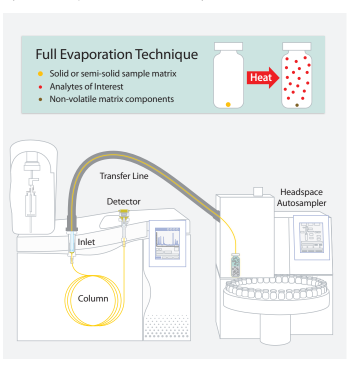


Figure 2: Grinding samples maximizes and normalizes surface area from sample to sample, increasing sensitivity and reproducibility.





Quantification

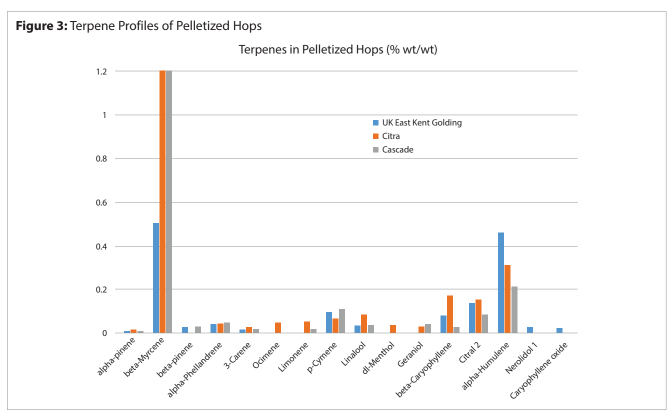
To aid in peak identification, a multi-component terpene standard was prepared with each compound at approximately 0.02% wt/vol. $10~\mu$ L of this standard solution was injected into a capped headspace vial and analyzed by FET headspace GC-FID. Standards were analyzed under the same conditions as the samples in order to eliminate the potential for discrimination across the volatility range (e.g., more volatile terpenes may show higher responses than less volatile terpenes). Since any discrimination effect would be the same in both the sample and standard, analytes were quantified based on their relative response factor compared to the standard as shown in Equation 1. This normalizes the values between sample and standard, ensuring accurate quantification across the full range of volatility for terpenes. Note that while the relative response factor technique improves accuracy, the semi-quantitative preparation of the standard and lack of well-characterized certified reference materials for terpenes limits the overall quantitative accuracy that can be obtained for this analysis. Additionally, the lack of pure, neat standards available to prepare a more concentrated standard resulted in a standard well below the level of many of the terpenes detected in this work. For accurate quantification, a calibration curve encompassing the expected concentration range of all analytes is required. The data presented in this article should be considered semi-quantitative.

Equation 1: Sample Concentration Calculation
$$Given: \frac{Standard\ Area}{Standard\ Concentration} = \frac{Sample\ Area}{Sample\ Concentration}$$

$$Sample\ Concentration = \frac{(Sample\ Area\times Standard\ Concentration)}{Standard\ Area}$$

Results and Discussion

The purpose of this study was to develop an FET headspace GC-FID method for the analysis of terpenes in cannabis using hops as a model system. The terpenes found in our samples matched well with literature descriptions of the terpenes present in hops [4]. High levels of terpenes were found across the volatility range, indicating that the FET headspace GC-FID technique was appropriate and that analysis of the standard adequately normalized any discrimination between the more and less volatile terpenes (Figure 3). Due to the starting concentration of some of the commercially available terpene standards, the maximum concentration at which the mixed terpene standard used for quantification could be prepared was 0.02% wt/vol, which is significantly lower than the concentration of some of the more prevalent terpenes in hops and cannabis. The use of a more concentrated standard solution is recommended to improve quantification of the higher concentrations found in these samples.

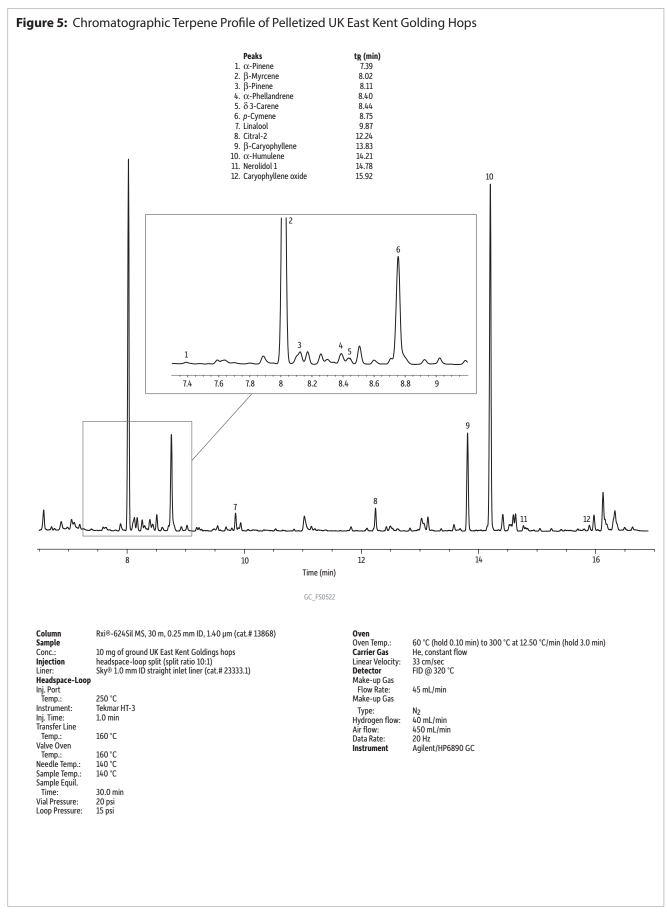


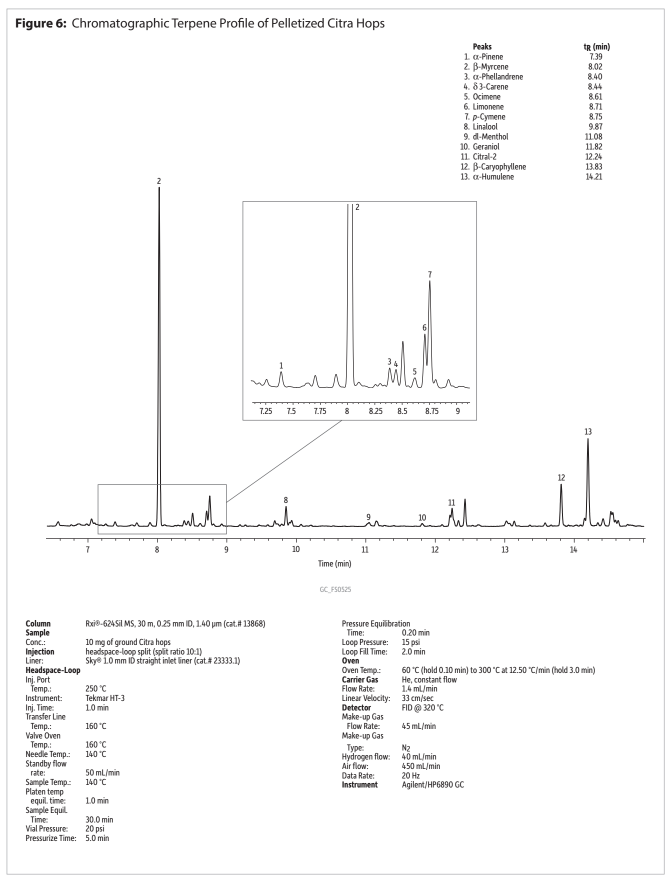
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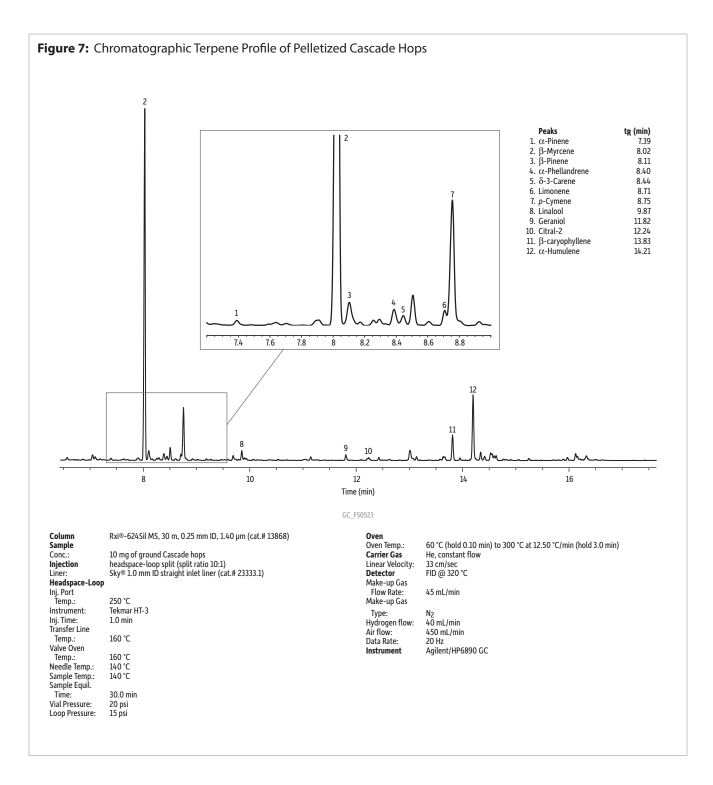
Figures 4–7 show individual chromatograms for the standard and each sample profiled for terpenes. Note that α -pinene, β -myrcene, α -humulene, β -caryophyllene, and caryophyllene oxide are well separated from interferences. For complex matrices, such as hops and marijuana, excellent chromatographic efficiency and selectivity are required to separate terpenes from one another and from other volatile matrix components in order to obtain accurate quantification. The selectivity of the Rxi®-624Sil MS column used here provided good separation of most terpenes and the small bore configuration (0.25 mm internal diameter) improved column efficiency, ultimately resulting in greater resolution between closely eluting terpenes than would be obtained using a wider bore column.

Figure 4: A 0.02% wt/vol multi-component terpenes standard analyzed on an Rxi®-624Sil MS column (30 m x 0.25 mm x 1.4 µm) demonstrates that this column provides the selectivity and efficiency needed to separate key terpenes using a simple FET headspace GC-FID method. **Peaks** te (min) Peaks te (min) **Peaks** te (min) **Peaks** te (min) 1. α-Pinene 7.39 10. Limonene 8.71 19. dl-Menthol 28. Citral 2 12.24 Camphene 7.71 11. p-Cymene 8.75 20. Borneol 11.19 29. Citral 3 13.19 β-Myrcene 798 12. β-Ocimene 8 82 21. α -Terpineol 11 29 30. Citral 4 13 43 31. β -caryophyllene Sabinene 8.02 13. Eucalyptol 8 91 22. Dihydrocarveol 11 40 13.83 14. Y-Terpinene 5. B-Pinene 8.11 9.06 23. Citronellol 11.51 32. α-Humulene 14.21 α-Phellandrene 15. Terpinolene 9.47 24. Geraniol 11.82 33. Nerolidol 1 14.78 8.4 7. δ 3-Carene 8.44 16. Linalool 9.87 25. 2-Piperidinone 34. Nerolidol 2 15.08 17. Fenchone 10.06 26. Citral 1 35. Caryophyllene oxide 15.92 8. α-Terpinene 11.92 9. Ocimene 8.61 18. Isopulegol 10.73 27. Pulegone 11.97 36. α -Bisabolol 16.43 18 28 10.6 11.6 10.8 11 11.2 11 4 11.8 12 8.2 31 17 9 10 13 14 16 11 12 Time (min) GC FS0518 Rxi® -624Sil MS, 30 m, 0.25 mm ID, 1.40 µm (cat.# 13868) Sample Equil. Column 30.0 min Sample Isopropyl alcohol 200 ng/µL (0.02% wt/vol). The sample was prepared by placing 10 µL Diluent: Vial Pressure: Loop Pressure: Conc.: into the headspace vial. headspace-loop split (split ratio 10:1) Sky® 1.0 mm ID straight inlet liner (cat.# 23333.1) 60 °C (hold 0.10 min) to 300 °C at 12.50 °C/min (hold 3.0 min) Injection Oven Temp.: Carrier Gas He, constant flow Liner: Headspace-Loop Linear Velocity: FID @ 320 °C Inj. Port Temp.: Instrument: 250 °C Detector Tekmar HT-3 Make-up Gas Inj. Time: 1.0 min Flow Rate: 45 mL/min Transfer Line Make-up Gas 160°C Temp.: Type: N₂ 40 mL/min Valve Oven Hydrogen flow: 160°C Air flow: Temp.: 450 mL/min Needle Temp.: Data Rate Sample Temp.: Agilent/HP6890 GC RESTÈK









While many cyano-based columns are commercially available, the Rxi®-624Sil MS column is recommended for terpene analysis because, in addition to offering optimized selectivity, the stationary phase is stabilized with silarylene, which significantly increases the operational temperature range of the column and improves its robustness. This is important for terpene analysis because some of the less-volatile terpenes require relatively high elution temperatures that would tax non-silarylene cyano stationary phases, resulting in shorter column lifetimes.

Although the Rxi®-624Sil MS column performs exceptionally well for the analysis of terpenes and residual solvents, it is too retentive for cannabinoids. In fact, cannabinoids do not elute from the Rxi®-624Sil MS column even at its 320 °C maximum

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temperature. Injection of cannabinoids on this column can potentially result in reduced column lifetime, selectivity changes, or baseline disturbances due to cannabinoids "bleeding" off of the stationary phase over time. Since both cannabinoids and terpenes will be present in cannabis samples, the sample preparation method must minimize the introduction of cannabinoids onto the analytical column. The full evaporation technique headspace sampling approach used here is ideal for terpene profiling because it introduces the volatile terpenes onto the GC column while eliminating the introduction of less volatile cannabinoids and nonvolatile matrix components into the system. This results in longer column lifetime and reduced inlet maintenance. Headspace sampling in general is simple to perform and requires no extraction or cleanup. While other methods exist that could remove cannabinoids from the sample while leaving the terpenes behind, these sample preparation methods are more time- and labor-intensive, and the increased amount of sample handling could result in loss of some of the more volatile terpenes, such as α -pinene. Grinding samples under dry ice is an additional measure that could be taken to minimize the loss of more volatile terpenes as it reduces the heat generated during the grinding process.

Conclusion

An FET headspace GC-FID method was used to analyze a comprehensive suite of terpenes in hops that are also found in cannabis samples. Compounds of interest across the volatility range were chromatographically separated and quantified. This method utilizes straightforward FET sample preparation, which minimizes manual labor and sample handling time. In addition, because it prevents nonvolatile material from entering the GC system, using the FET approach can increase column lifetime and reduce inlet maintenance. This technique, column, and instrument setup can also be used to analyze residual solvents in cannabis concentrates, eliminating the need for additional capital investment for different instrumentation and/or columns.

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Foods, Flavors & Fragrances Applications

A Fast, Simple FET Headspace GC-FID Technique for Determining Residual Solvents in Cannabis Concentrates

By Corby Hilliard; Amanda Rigdon; William Schroeder*, Ph.D.; Christi Schroeder*, Ph.D.; and Theo Flood*

*Cal-Green Solutions

Abstract

Due to rapid growth in the medical cannabis industry, demand is increasing for analysis of residual solvents in cannabis concentrates in order to protect consumer safety. This application note details a simple, fast test for common residual solvents using full evaporation technique headspace GC-FID and an Rxi° -624Sil MS column.

Introduction

As the popularity of cannabis concentrates increases, consumer safety concerns are resulting in the establishment of new regulations to control the level of residual solvents in commercial cannabis concentrates. The State of Colorado, for example, published allowable concentrations of certain residual solvents in Rule R 712. This is because, although cannabis concentrates can be produced in numerous ways, one of the most common means of extracting therapeutic compounds, like tetrahydrocannabinol (THC), cannabidiol (CBD), and terpenes, from cannabis is through extraction with an organic solvent, such as butane. After the cannabinoids and terpenes are extracted from the plant material, the organic solvent is allowed to evaporate and then is purged off using heat and/or vacuum. These extraction solvents can be difficult to purge completely, so the finished product needs to be tested to ensure that residual solvents are only present at or below safe levels. For consumer safety, especially with medicinal products, accurate and comprehensive analysis of residual solvents is necessary for concentrates and extracts.

Since residual solvents are extremely volatile, they cannot be analyzed by HPLC and lend themselves nicely to GC analysis. One of the most common and reliable ways to quantify residual solvents is through headspace gas chromatography–flame ionization detection (GC-FID). Headspace injection works by driving volatile compounds of interest from the sample into a gas phase in the headspace of the vial above the sample. An aliquot is then withdrawn from the headspace of the vial and analyzed by GC-FID in order to determine the volatile components of the sample. One approach for headspace GC-FID that is particularly useful for analyzing cannabis concentrates is the full evaporation technique (FET). FET sample preparation involves the use of a very small sample amount (e.g., 20–50 mg), which effectively creates a single-phase gas system in the headspace vial at equilibrium [1]. FET is ideal for difficult and varied matrices like cannabis concentrates because it eliminates matrix interferences that can cause inaccurate quantification, and it also has the advantages of little to no manual sample handling and a very small sample size. Additionally, high sensitivity can be achieved through the creation of a single-phase system in the headspace vial. Figure 1 illustrates the basic principle of headspace GC using the full evaporation technique.

The work described here demonstrates the viability of FET headspace injection and GC-FID analysis of residual solvents in cannabis concentrates. The method is simple to implement, quick to run, and does not require expensive dynamic headspace equipment or mass spectrometric detectors. While the methodology presented here is suitable for residual solvents in cannabis concentrates, it is not applicable for finished tinctures in alcohol. Finished alcohol tinctures contain large amounts of alcohol which will severely interfere with quantification of other residual solvents in the sample. Therefore, an alternate approach is required for alcohol tinctures. This technique also may be applicable for oil or glycerin tinctures; however, it has not been evaluated for that use.



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Experimental

Headspace and GC Method Optimization

An Rxi*-624Sil MS column was selected for this work as it is designed specifically for volatiles analysis and is widely used for the analysis of residual solvents in pharmaceutical products. Final FET headspace injector and GC-FID operating conditions are presented in Figure 3. Initially, modeled conditions for analyzing the specific compounds of interest were generated using Restek's *EZGC*TM chromatogram modeler. The method from the modeler was then optimized to account for headspace analysis employing a headspace instrument with a transfer line.

The following parameters were optimized for this method:

Linear velocity: Linear velocity was increased to 80 cm/sec to allow for fast sample transfer through the headspace instrument transfer line. Fast sample transfer minimizes band broadening, which maximizes efficiency, resolution, and sensitivity. The original GC oven program generated by the EZGC™ chromatogram modeler was translated using the EZGC™ method translator to give a new oven program optimized for the new carrier flow. Method translation is required when changing flow rates in order to keep elution temperatures constant. Changes in

Full Evaporation Technique

Solid or semi-solid sample matrix
Analytes of Interest
Non-volatile matrix components

Transfer Line

Detector

Headspace
Autosampler

elution temperatures between the original and the translated method will sometimes result in drastically different separations or even coelutions, especially on highly selective phases like the Rxi*-624Sil MS column.

- *GC inlet liner choice*: The liner used for this work was a 1 mm straight Sky* inlet liner (cat.# 23333.1). The use of a small internal diameter liner minimizes band broadening by reducing the overall volume of the inlet, again resulting in higher efficiency, resolution, and sensitivity.
- Split ratio: A split ratio of 10:1 was used for this work. Although maximum sensitivity is required due to very low expected levels of target analytes, using a split ratio of at least 10:1 ensures high sample velocity through the GC inlet, which minimizes band broadening, increasing resolution without compromising sensitivity. Sharper peaks are taller peaks, so any loss in sensitivity is mitigated through an increase in signal-to-noise ratio.
- Equilibration temperature: Samples were equilibrated at 140 °C to encourage complete melting of waxy concentrates. By melting the extracts, the ratio of surface area to volume is maximized, ensuring 100% transfer of the analytes of interest into the headspace. The use of a larger sample size will compromise this ratio; therefore, sample sizes should be kept as small as possible to ensure accurate quantification (20 mg is recommended for this application). Representative concentrates are shown in Figure 2. Small samples (20–25 mg) of each concentrate type were placed in a capped headspace vial and incubated for 30 minutes at 140 °C. All concentrates melted completely at the 140 °C incubation temperature, forming a thin film at the bottom of the headspace vial.
- Equilibration time: The equilibration time for this method was 30 minutes. This allows enough time for waxy concentrates to melt completely and ensures equilibrium is reached in the headspace vial. Equilibrium is required for accurate and reproducible quantification.
- Oven program: The oven program was optimized for speed for this application. In samples that contain terpenes, it is recommended that the oven ramp be extended to 320 °C and the isothermal hold time be extended to 5 minutes in order to ensure complete elution of any terpenes that may be present in the sample.



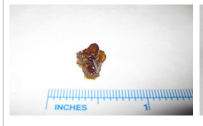
Figure 2: Cannabis concentrate samples are solid before FET incubation (left) and then melt completely into a thin liquid layer after a 30-minute incubation at 140 °C (right).

Crumble - Melting point = ~115 °C





Shatter - Melting point = 108 °C





Taffy - Melting point = 102 °C





Photos and melting point data courtesy Cal-Green Solutions

Table I: Commodity and Calibration Standard Curve Equivalency Levels

Concentration in Commodity (ppm)	Amount in 20 mg Sample (µg)	Concentration in 10 µL Standard (µg/mL)
500	10	1,000
250	5	500
100	2	200
50	1	100
25	0.5	50
10	0.2	20
5	0.1	10

Calibration Curve Preparation

When preparing standards for FET headspace GC-FID, it is necessary to calculate the total mass of analyte that will be present in a representative sample, since the equilibrium state results in a single-phase system. For example, a 20 mg sample containing a residual solvent at 50 ppm contains 1 µg of that residual solvent. Therefore, the 50 ppm point in the calibration curve should contain 1 µg of each compound of interest. Since FET headspace GC-FID depends on the establishment of a single phase system, very small volumes are required for standards. The volume used for standards in this application was 10 µL, which was placed directly into a capped headspace vial by injecting it through the vial septum with a clean syringe. Table I presents the 7-point calibration curve standards and their corresponding concentrations in commodity samples.

Standards were prepared in dimethyl sulfoxide (DMSO), which is a less-volatile, later-eluting solvent that does not interfere with the residual solvents of interest. Because FET establishes a single-phase system in the headspace vial without partitioning, it is not necessary to matrix-match standards and samples, which simplifies standard preparation for varied matrices.

The calibration curve was prepared by first making a 1,000 μ g/mL stock solution for dilution. The stock solution was prepared as follows:

- Prepare a 5,000 μ g/mL stock solution of butane by bubbling butane standard through DMSO on a balance in a fume hood. The butane used for this work was a mixture of butane and isobutane
- Prepare a 1,000 μ g/mL stock solution by adding 2 mL of 5,000 μ g/mL butane stock to a 10 mL volumetric flask, adding ~4 mL DMSO, and then volumetrically adding each neat solvent to the flask using a syringe. Volumes required for the 1,000 μ g/mL stock standard were adjusted to account for the density of each solvent as shown in Table II.
- After the addition of neat solvents, fill the flask to the line with DMSO and mix by gently inverting the flask three times and rotating to swirl the contents between inversions.

Table II: Density-Adjusted Volumes Used to Prepare 10 mL of the 1,000 μg/mL Stock Solution

Compound	Density (g/mL)	Volume Required (µL)
Butane	measured gravimetrically	2,000
Chloroform	1.48	6.7
Isobutane	NA	2,000
Acetone	0.79	12.6
Methanol	0.79	12.6
Ethanol	0.79	12.7
IPA	0.79	12.7
Benzene	0.88	11.4
Toluene	0.87	11.5
Pentane	0.63	16.0
Hexane	0.65	15.3
Heptane	0.68	14.7

The 1,000 μ g/mL stock solution prepared using Table II was used as the highest calibration standard. All other calibration points were prepared in 5 mL volumetric flasks with separate dilutions of the 1,000 μ g/mL stock solution. Serial dilution was not used for this work in order to minimize time-consuming syringe rinsing during calibration curve preparation. Because the compounds used here are volatile, work needed to be completed as quickly as possible to prepare the calibration standards. In addition, volumetric flasks were kept capped to minimize evaporative loss. Table III details the preparation of the calibration curve standards.

Table III: Calibration Curve Preparation

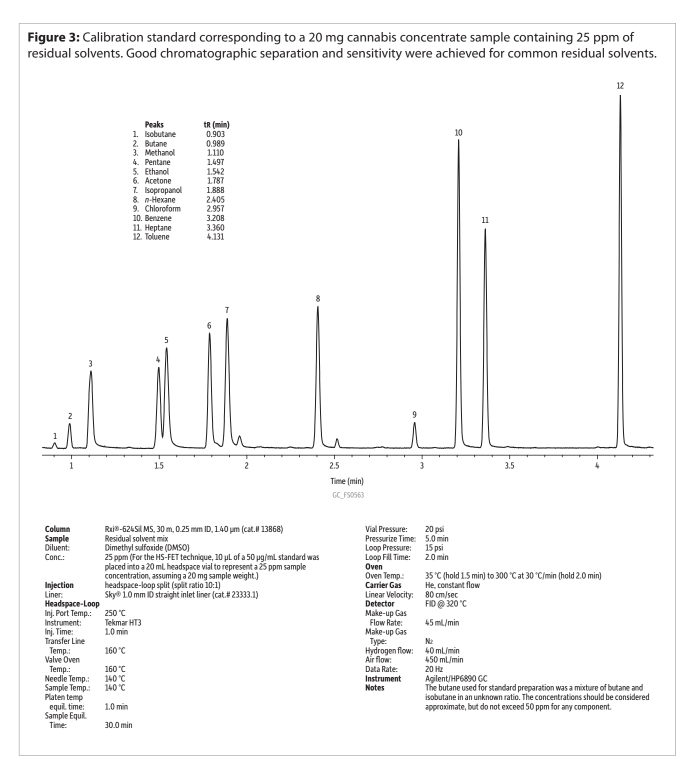
Calibration Level (ppm in Commodity)	Volume of 1,000 µg/mL Stock Solution (mL)	Final Volume (mL)	Final Calibration Standard Concentration (µg/mL)
500	5	5	1,000
250	2.5	5	500
100	1	5	200
50	0.5	5	100
25	0.25	5	50
10	0.1	5	20
5	0.05	5	10

After preparation, all calibration standards were divided into 2.5 mL aliquots and stored in the refrigerator at 5 °C. Since DMSO freezes under refrigeration, calibration standards were allowed to thaw completely prior to use. By aliquoting the calibration standards into separate vials, freeze/thaw cycles were reduced for the entire volume of the calibration solution, allowing for longer storage life of calibration and stock solutions. If desired, calibration standards may be split into aliquots smaller than 2.5 mL to further reduce freeze/thaw cycles. This can be accomplished by pipetting aliquots into gas-tight vials using a glass pipet and immediately capping the vials.

Results and Discussion

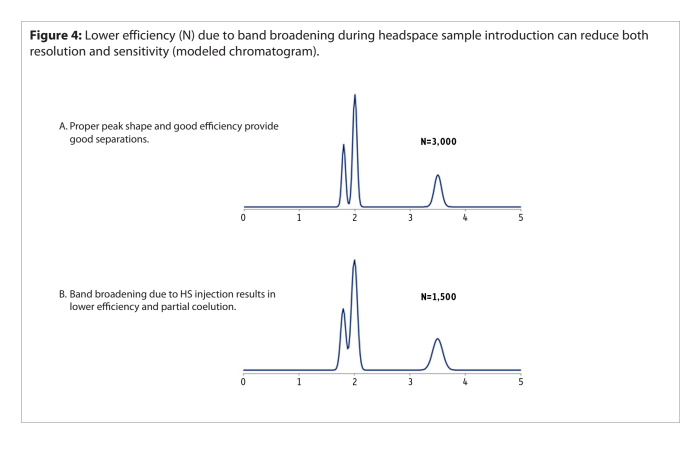
Good chromatographic peak shape, separation, and sensitivity were achieved for all analytes of interest. Figure 3 shows the 25 ppm calibration standard. Use of the Restek® Rxi®-624Sil MS column allowed for the separation of the wide variety of solvents that may be present in cannabis concentrates in a short analysis time, while retaining and resolving highly volatile butane isomers. This column was selected for the FET headspace GC-FID method because it was designed specifically for volatiles analysis and is widely used for the analysis of residual solvents in pharmaceutical products. Additionally, the column's unique selectivity also resolves dozens of terpenes [2]. This allows cannabis terpene profiling to be done without changing columns or injection technique, which decreases downtime between methods and improves lab productivity.





In addition to using a highly efficient, selective Rxi®-624Sil MS column, it is critical to optimize several GC parameters for headspace analyses in order to prevent band broadening. Early-eluting compounds such as isobutane and butane do not focus on the head of the analytical column, so band broadening through the headspace system and injection port can reduce efficiency, severely impacting sensitivity and resolution for these compounds (Figure 4). As detailed in the Experimental section, band broadening was controlled by using a fast linear velocity, narrow bore inlet liner, and a 10:1 split ratio. This approach speeds up sample transfer and ensures good chromatographic peak shape and response.

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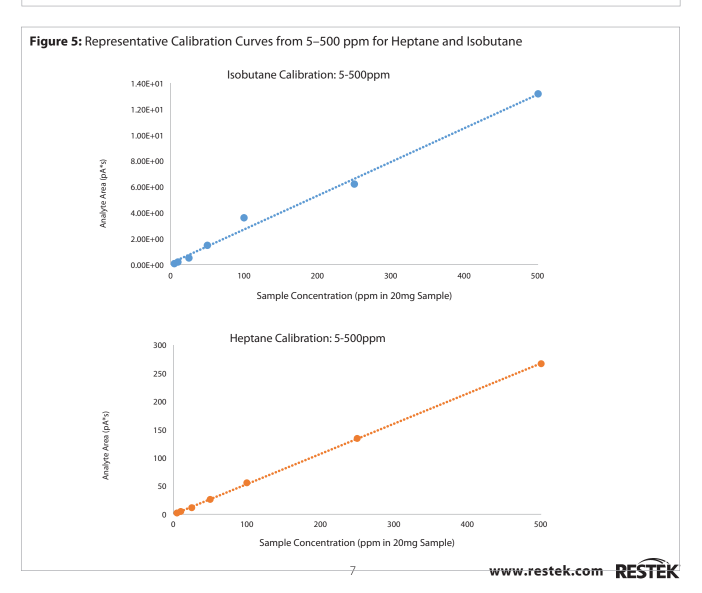


Analysis of calibration standards resulted in good sensitivity and linear responses for all analytes of interest. Table IV shows the signal-to-noise ratios at 10 ppm and 50 ppm (current Colorado regulatory cutoff values), as well as the correlation coefficients (r values) and coefficients of determination (r² values) for all analytes. All compounds exhibited adequate signal-to-noise ratios (> 10:1) at their respective Colorado state regulatory limits. Signal-to-noise ratios were > 10:1 for all compounds at 10 ppm, with the exception of isobutane. The Colorado cutoff for isobutane was 50 ppm at the time of this study; however, prior to publication, Colorado changed the limits and solvents of interest for residual solvent testing. This method will be suitable for the new regulations as well as the older ones.

Figure 5 shows plots of the most linear (heptane) and least linear (isobutane) calibration curves. All calibration curves exhibited acceptable linearity without the use of an internal standard. The use of an internal standard may improve linearity and reproducibility, if desired.

Table IV: Using full evaporation technique sample introduction for headspace GC-FID resulted in good sensitivity and linearity for all residual solvents as shown by peak response and correlation data for the calibration standards.

Compound	S:N 10 ppm	S:N 50 ppm	r	r ²
Isobutane	5.30	30.7	0.996	0.992
Butane	18.8	119	0.997	0.994
Methanol	48.1	189	0.999	0.999
Pentane	19.0	50.0	0.998	0.995
Ethanol	45.2	88.1	0.999	0.998
Acetone	49.9	97.0	0.999	0.999
Isopropanol	56.4	107	0.998	0.996
Hexane	45.6	109	0.999	0.998
Chloroform	11.5	22.5	0.999	0.998
Benzene	150	293	0.999	0.998
Heptane	88.4	193	1.00	1.00
Toluene	166	317	0.999	0.998



Conclusion

By combining a selective Rxi*-624Sil MS GC column with the FET headspace GC-FID technique, excellent sensitivity and linearity were achieved for residual solvent compounds applicable to cannabis concentrates. The use of FET headspace GC-FID should allow quantification without the use of matrix-matched standards by creating a single non-partitioning phase system in the headspace vial. This technique also has the added benefit of needing very little sample and is applicable for the analysis of other volatile compounds, such as terpenes, in cannabis products.

References

[1] B. Kolb, L. Ettre, Static Headspace-Gas Chromatography: Theory and Practice, John Wiley & Sons, Hoboken, NJ, 2006.

[2] J. Cochran, Terpenes in Medical Cannabis, ChromaBLOGraphy, Restek Corporation, 2014 http://blog.restek.com/?p=11451 (accessed July 18, 2014).



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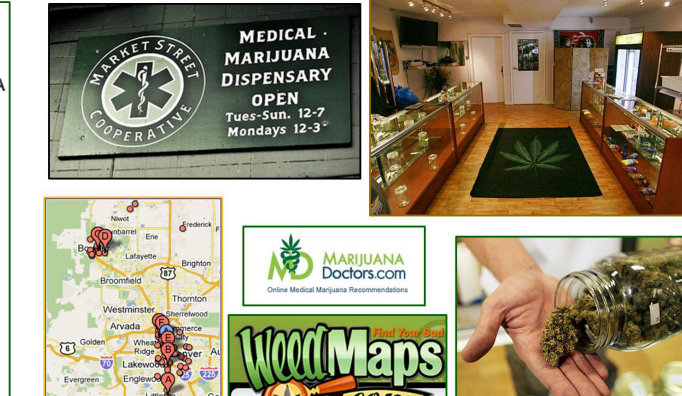
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High Quality Analysis of Pesticides in Marijuana for Food and Medicine using QuEChERS, Cartridge SPE, GCxGC-TOFMS, and LC-MS/MS

Jack Cochran, Julie Kowalski, Sharon Lupo, Michelle Misselwitz, Amanda Rigdon, Jason Thomas, Restek Corporation Frank Dorman, Jessica Westland, Amanda Leffler, The Pennsylvania State University

- We Over 15 states in the USA have medical marijuana laws.
- * Therapeutic benefits include pain relief, nausea control, appetite stimulation, and muscle relaxation.
- Marijuana is illegal on the federal level so patients have no assurances on medicine safety, including for pesticide residues.
- We used the QuEChERS sample preparation approach for extracting pesticides from marijuana.
 - But dispersive SPE did not have the cleanup capacity for GCxGC work.
 - ★ Instead, we employed cartridge SPE for cleanup for GCxGC.
- **W** GCxGC-TOFMS and LC-MS/MS were used for pesticide determinations in cleaned up QuEChERS extracts.
 - * The selectivity of advanced techniques was needed due to sample extract complexity, even after dilution/cleanup.
 - LC-MS/MS was necessary for abamectin because it does not gas chromatograph.







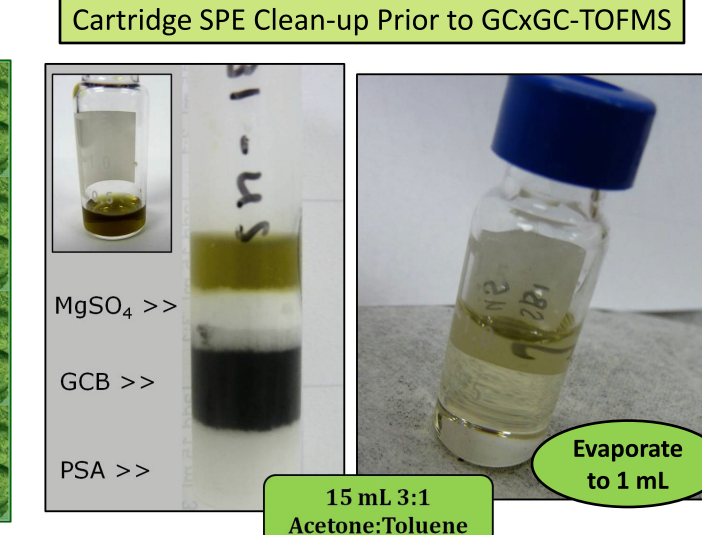




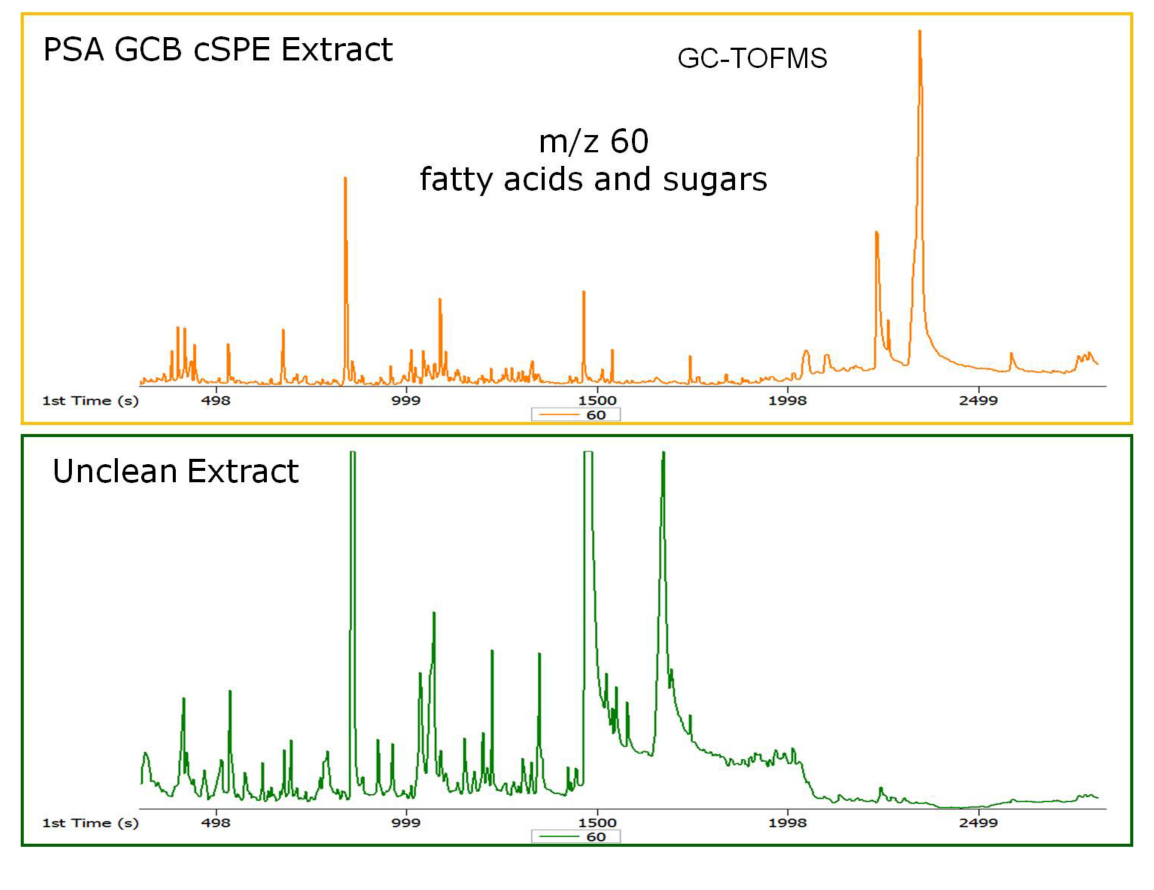


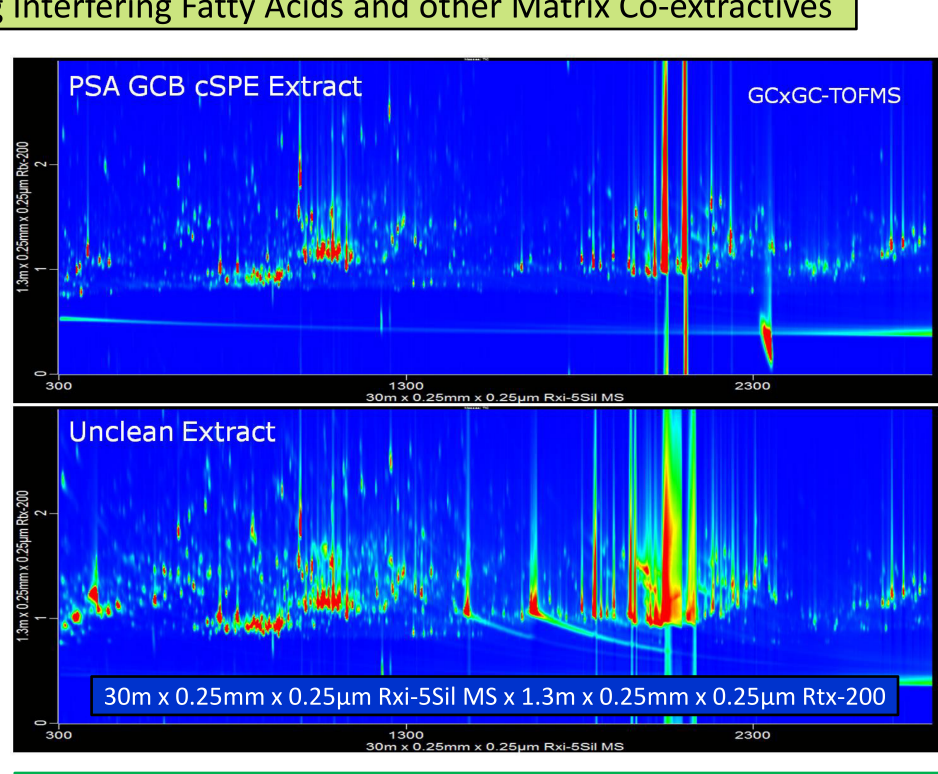






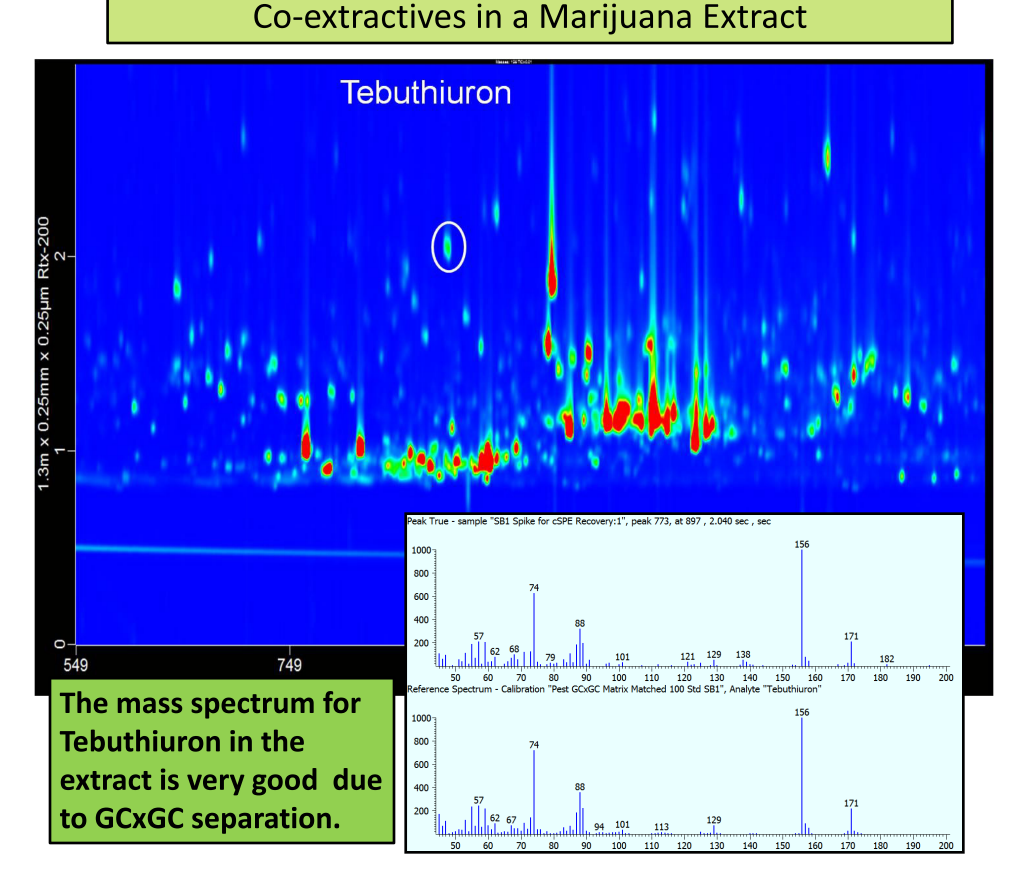
High Capacity Cartridge SPE Produces a Cleaner Extract by Removing Interfering Fatty Acids and other Matrix Co-extractives



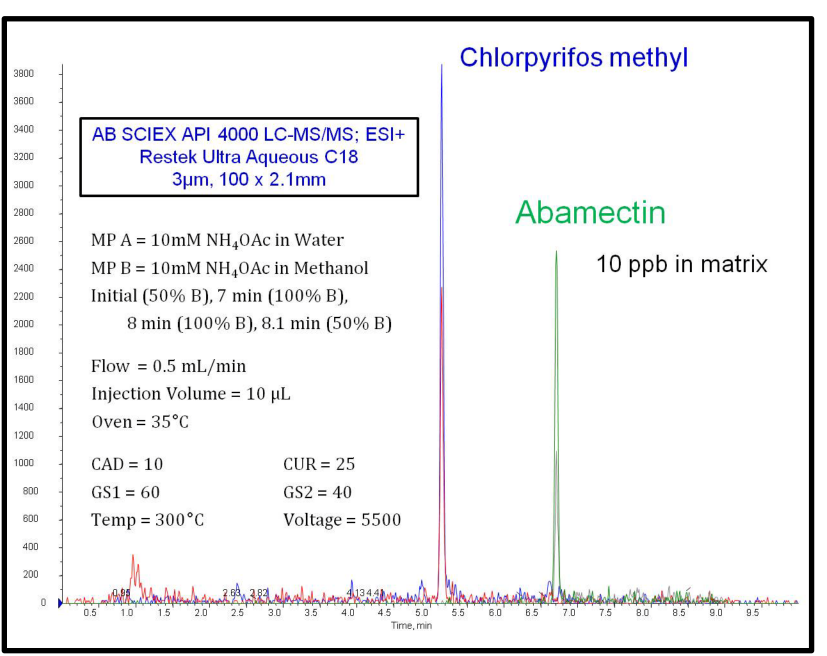


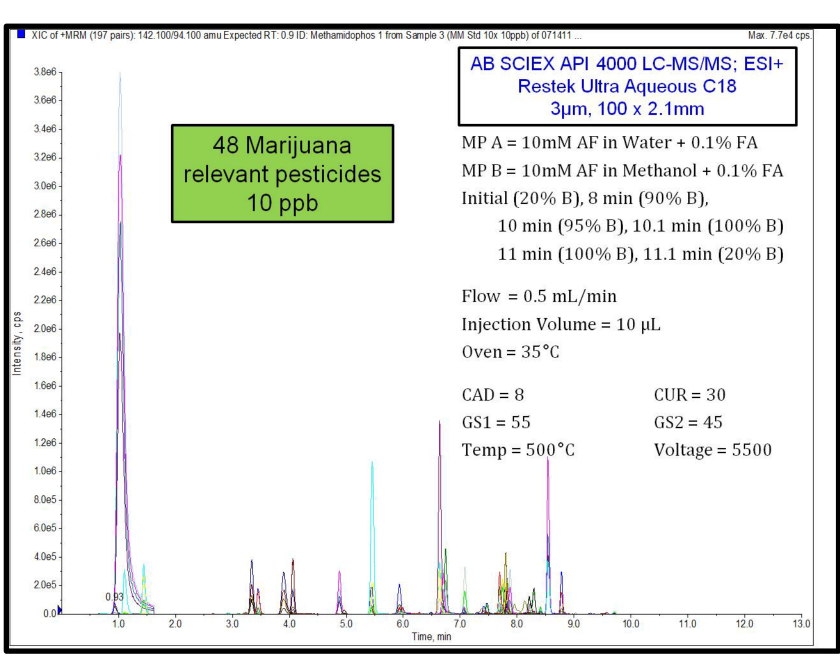
Splitless injection, 250°C, 1µL, valve 60 sec (4mm single taper Sky liner with wool) Primary oven: 80°C (1 min), 5°C/min to 310°C; Secondary oven: +5°C offset He, corrected constant flow 2 mL/min; Modulation time: 3 sec LECO Pegasus GC-TOFMS, El 70 eV, Source temp 225°C, 45 to 550 u, 100 spectra/sec

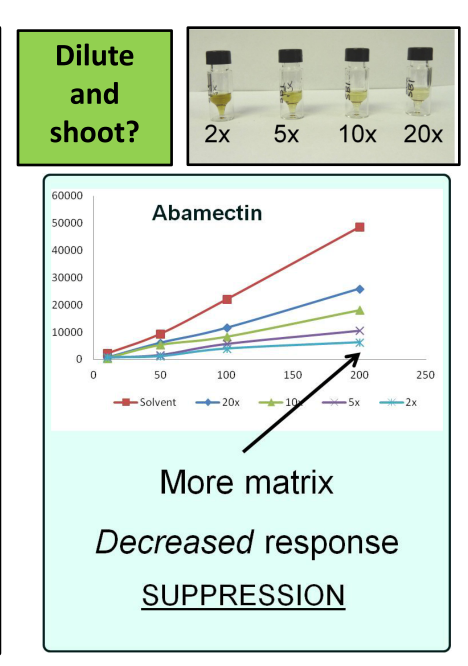
GCxGC Separates Pesticides from Remaining Matrix

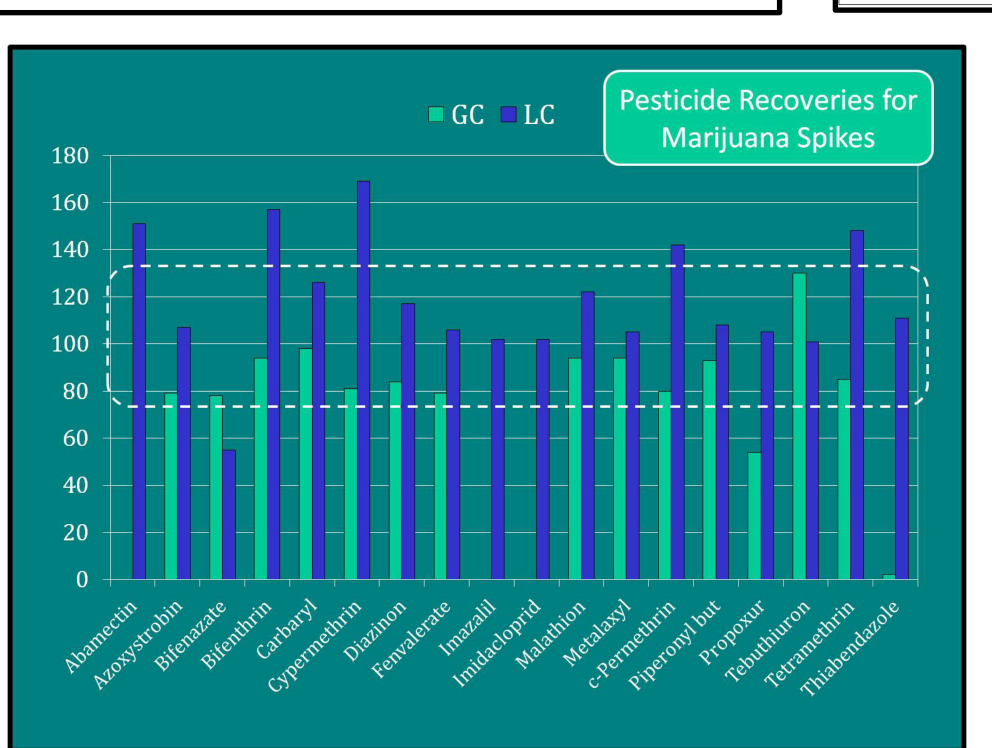


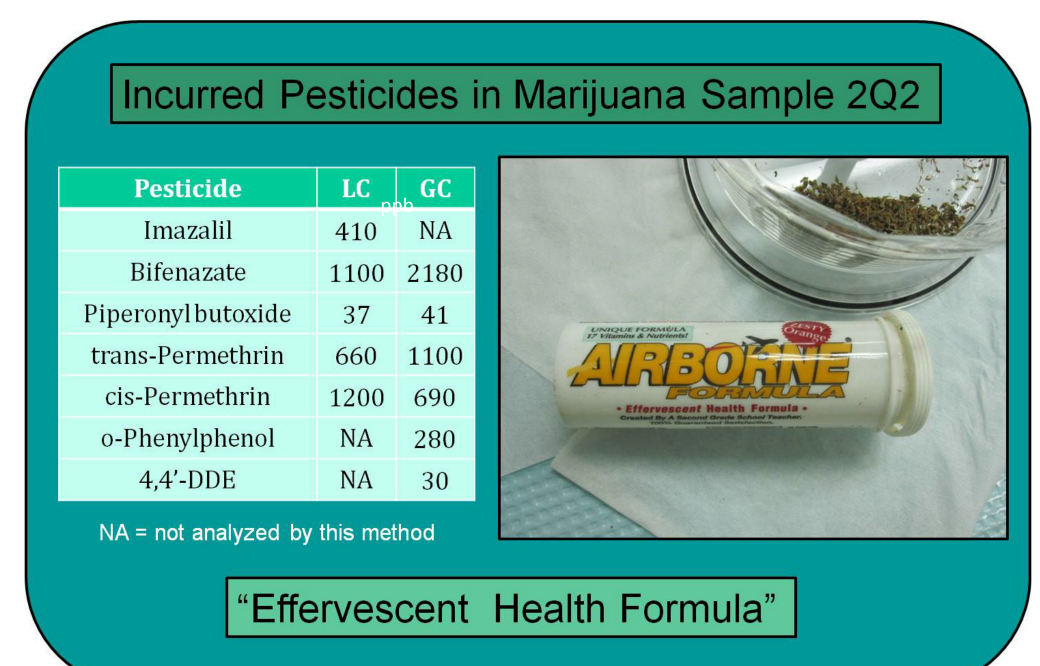
LC-MS/MS of Marijuana Pesticides – Abamectin Required a Single Analyte Method Approach



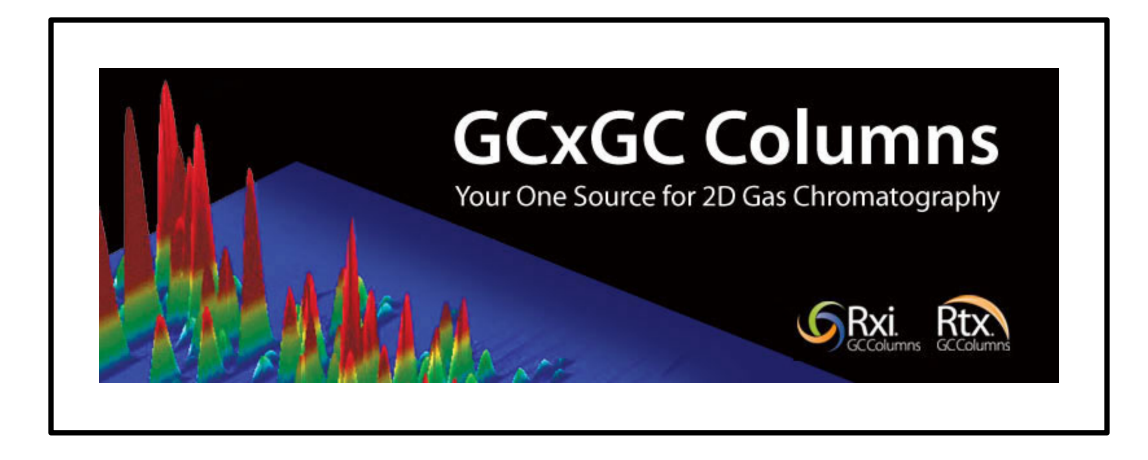


















Technical Article

High-Quality Analysis of Pesticides in Cannabis

Using QuEChERS, Cartridge SPE Cleanup, and GCxGC-TOFMS

By Jack Cochran, Julie Kowalski, Sharon Lupo, Michelle Misselwitz, and Amanda Rigdon

- Quickly and effectively extract medical marijuana samples for pesticide analysis.
- Cartridge SPE cleanup of dirty extracts improves GC inlet and column lifetimes.
- Selective GC columns increase accuracy of pesticide determinations for complex samples.

Over 20 states in the U.S. have legalized the use of recreational or medical cannabis because of therapeutic benefits for ailments such as cancer, multiple sclerosis, and ALS. Dosing methods include smoking or vaporizing and baked goods. Unlike other prescribed medicines regulated by U.S. FDA, marijuana is a Schedule 1 drug and is illegal on the federal level. As a result, medical cannabis patients have no safety assurances for their medication, which could contain harmful levels of pesticide residues. Currently, medical marijuana pesticide residue analysis methods are poorly defined and challenging to develop due to matrix complexity and a long list of potential target analytes.

In order to address matrix complexity, we combined a simple QuEChERS extraction approach with cartridge SPE (cSPE) cleanup, followed by GCxGC-TOFMS. Acceptable recoveries were obtained for most pesticides, and incurred pesticide residues were detected in some of the illicit marijuana samples used for method development.

QuEChERS Extraction Saves Time and Reduces Hazardous Solvent Use

Trace residue extraction procedures from dry materials like medical cannabis typically involve large amounts of solvent, long extraction times, and tedious concentration steps similar to the Soxhlet procedure or multiresidue methods from the Pesticide Analytical Manual. QuEChERS, with its simple 10 mL acetonitrile shake extraction and extract partitioning with salts and centrifugation, offers time savings, glassware use reduction, and lower solvent consumption.

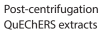
Water was added to finely ground, dry cannabis samples to increase QuEChERS extraction efficiency, especially for more polar pesticides. A vortex mixer was used to shake the solvent and sample for at least 30 minutes prior to extract partitioning. When finished, it was easy to transfer the supernatant from the QuEChERS extraction tube for subsequent cSPE cleanup prior to analysis with GC or LC (Figure 1).

Cartridge SPE Cleanup Improves **GC Inlet Uptime**

Injecting chlorophyll-laden extracts into a GC gives reduced recoveries for less volatile pesticides, and results in degradation of sensitive pesticides like DDT and Dicofol (Table I). SPE cleanup with a 500 mg graphitized carbon black/500 mg PSA cartridge removes chlorophyll and traps fatty acids that interfere with qualitative pesticide identification and bias quantification. cSPE has increased sorbent capacity over dispersive SPE for thorough cleanup of complex extracts.

Figure 1: A quick and easy QuEChERS extraction, combined with cSPE, effectively prepared extracts for pesticide residue analysis from highly complex marijuana samples.







OuEChERS extracts loaded on SPE cartridge



Final extract



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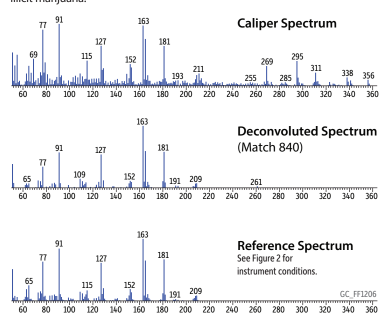
Orthogonal GC Columns Increase Separation Power for More Accurate Pesticide Results

GCxGC is a powerful multidimensional approach that gives two independent separations in one instrumental analysis. An Rxi®-5Sil MS and Rtx®-200 column combination distributes pesticides broadly in both dimensions, providing a highly orthogonal GCxGC system. More important though is separating pesticides from potential isobaric matrix interferences, as seen in the surface plot for the insecticide cypermethrin (Figure 2). Cypermethrin gas chromatographs as four isomers, and all would have experienced qualitative interference and quantitative bias from peaks in the foreground of the surface plot had only 1-dimensional GC been used. With GCxGC-TOFMS, cypermethrin was unequivocally identified in a marijuana sample at a low ppm level (Figure 3).

Summary

QuEChERS and cSPE produced usable extracts from highly complex cannabis samples for high-quality pesticide residue analysis. The multidimensional separation power of GCxGC-TOFMS was then used to correctly identify and quantify pesticides in these complex extracts.

Figure 3: Positive mass spectral identification of incurred cypermethrin in illicit marijuana.



Acknowledgment: Randy Hoffman, a Police Evidence Technician at The Pennsylvania State University (PSU), supplied the seized marijuana samples while overseeing their handling. Frank Dorman at PSU assisted with QuEChERS extractions.

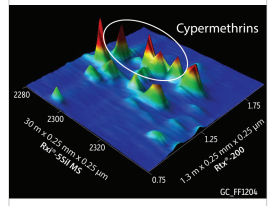
Initially published in Restek® Advantage.

Table 1: Pesticide recoveries for a QuEChERS extract of cannabis give higher results when cSPE is used for cleanup. Dicofol and DDT are degraded in the inlet for the dirtier extract, yielding high DDD results.

Pesticide	Classification	With cSPE Cleanup (%)	Without cSPE Cleanup (%)
4,4'-DDD	Organochlorine	83	230
4,4'-DDT	Organochlorine	77	9
Bifenthrin	Pyrethroid	86	89
Dicofol	Organochlorine	84	ND
Azinphos methyl	Organophosphorus	79	53
trans-Permethrin	Organochlorine	68	17
Pyraclostrobin	Strobilurin	73	19
Fluvalinate	Pyrethroid	72	23
Difenoconazole	Triazole	67	21
Deltamethrin	Pyrethroid	68	20
Azoxystrobin	Strobilurin	72	27

ND = no peak detected

Figure 2: GCxGC-TOFMS and orthogonal Rxi®-5Sil MS and Rtx®-200 columns allow incurred cypermethrins in a marijuana extract to be separated from interferences (m/z 163 quantification ion).



eaks		RT 1 (sec.)	RT 2 (sec.)
1.	Cypermethrin 1	2292	1.50
2.	Cypermethrin 2	2304	1.54
3.	Cypermethrin 3	2310	1.53
4.	Cypermethrin 4	2313	1.58

Column: Rxi®-55il MS 30 m, 0.25 mm ID, 0.25 µm (cat.# 13623), Rtx®-200 1.3 m, 0.25 mm ID, 0.25 µm (cat.# 15124); Sample: Diluent: Toluene; Injection: Inj. Vol.: 1 µL splitless (hold 1 min); Liner: Sky® 4 mm single taper w/wool (cat.# 23303.1); Inj. Temp: 250 °C; Purge Flow: 40 mL/min; Oven: Oven Temp: Rxi®-55il MS: 80 °C (hold 1 min) to 310 °C at 5 °C/min, Rtx®-200: 85 °C (hold 1 min) to 315 °C at 5 °C/min; Carrier Gas: He, corrected constant flow (2 mL/min); Modulation: Modulator Temp. Offset: 20 °C; Second Dimension Separation Time: 3 sec.; Hot Pulse Time: 0.9 sec.; Cool Time between Stages: 0.6 sec.; Instrument: LECO Pegsaus 4D GCxGC-TOFMS; For complete conditions, visit www.restek.com and enter GC FF1204 in the search.



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Reliably Confirm Cannabinoids by GC-MS

Using a 12m x 0.20mm ID 0.33µm Rxi®-5ms Column

by Kristi Sellers, Clinical/Forensic Innovations Chemist

- · Baseline resolution for all major metabolites.
- Ultra-low bleed at 300°C, for accurate data.
 Bake column at 340°C, to remove derivatization by-products and prolong column life.

Marijuana is one of the most abused substances in the United States. Its common abuse stems from its widespread availability and because it is inexpensive, compared to other abused substances such as cocaine and heroin. Marijuana use typically is determined by screening for its major metabolite in urine, 11-nor-9-carboxy- Δ^9 -tetrahydrocannabinol (Δ^9 -carboxy-THC), using an immunoassay. When screening results are positive, gas chromatography/mass spectrometry (GC/MS) is employed for confirmation. Marijuana use also can be determined by analyzing other sample matrices, such as blood, hair, oral fluid, or body tissues but, again, positive results require GC/MS confirmation.

GC/MS confirmation methods require sample clean-up and derivatization of target analytes, and call for a capillary GC column that can produce reliable identification and quantification results. Δ_9 -carboxy-THC is the primary target in GC/MS confirmation analysis, but other marijuana metabolites present in samples include cannabinol, cannabidiol, 11-hydroxy- Δ_9 -tetrahydrocannabinol (Δ_9 -hydroxy-THC), Δ_9 -tetrahydrocannabinol (Δ_9 -THC), and Δ_8 -tetrahydrocannabinol (Δ_9 -THC). Further, a guard column typically is recommended for this analysis, to prevent non-volatile residue in the sample matrix from contaminating the analytical column. The guard column should have an internal diameter approximately equal to that of the analytical column, to minimize changes in flow rate.

For the analysis we show in this article, we used MTBSTFA (N-methyl-N-(tert-butyldimethylsilyl)-trifluoroacetamide) to derivatize the target compounds. The analytical column we chose is our new $12m \times 0.20mm$ ID $\times 0.33\mu m$ RxiTM-5ms column (5% diphenyl / 95% dimethylpolysiloxane stationary phase). The small internal diameter makes this column compatible for use with mass spectrometers, because the column can be operated using a 1.0mL/min. flow rate. The short length produces analysis times of less than 15 minutes for the major metabolite, $\Delta 9$ -carboxy-THC, which elutes last. Because the target compounds have relatively high molecular weights (310-358 amu, underivatized — see Figure 1), the GC oven must be programmed to a relatively high temperature, 300°C, to keep analysis time short.

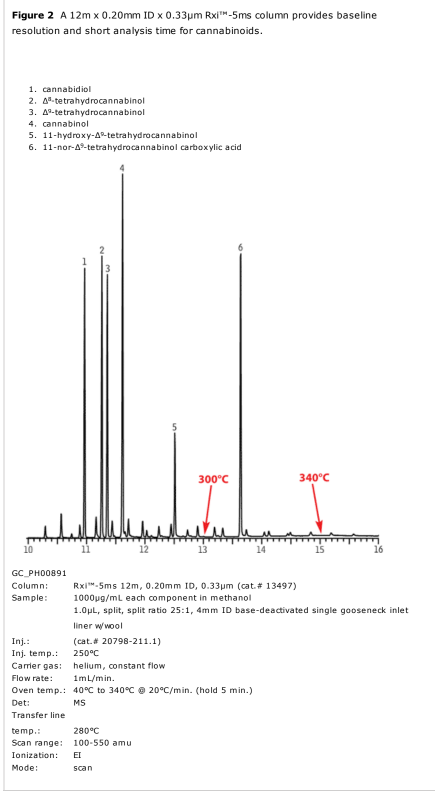
The column and conditions we used ensure baseline resolution for all of the metabolites in Figure 2. Figure 2 also shows that the ultra-low bleed exhibited by the Rxi[™]-5ms column does not interfere with the analysis. The GC oven must heated to an even higher temperature between samples, 340°C, to bake sample matrix interferences and derivatization by-products from the system. Derivatization by-products can be seen in the baseline in Figure 2.

The results of this analysis demonstrate that a $12m \times 0.20mm$ ID $\times 0.33\mu m$ RxiTM-5ms column has the selectivity and inertness needed to provide baseline resolution, suitably short analysis times, and no interference from bleed at high temperature. We highly recommend it for this analysis.

Figure 1 Cannabinoids have relatively high molecular weights, so high temperatures must be used in their analysis.

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- 1. Smith, F. and J. Siegel Handbook of Forensic Drug Analysis Elsevier Academic Press, 2005, pp. 98-151.
- 2. Clouette, R., M. Jacob, P. Koteel, and M. Spain Journal of Analytical Toxicology 17 (1): 1-4 (Jan./Feb. 1993).

RELATED SEARCHES

marijuana, cannabinoid metabolites, Rxi-5ms, THC



Technical Article

Don't Overestimate Cannabidiol During Medical Cannabis Potency Testing by Gas Chromatography

By Jack Cochran

Accurate potency testing of medical cannabis with gas chromatography (GC) depends principally on choosing a column with the right selectivity; otherwise, coelutions between cannabinoids of interest may cause error in potency measurements. Cannabidiol is one of the chief cannabinoids with pharmacological value and provides relief against nausea, anxiety, and inflammation. Potency testing for medical marijuana is often done using "5-type" GC columns since they are commonly available in most labs. However, on 5-type columns cannabidiol can coelute with cannabichromene, a compound that likely also has medical value and is increasingly becoming part of potency testing. To identify and report both of these compounds accurately, a GC column with a different stationary phase is needed.

Proper Column Choice Results in More Accurate Potency Data

As shown in Figure 1, cannabinoids are aromatic compounds, meaning they will likely be better separated on a column that contains aromatics in the stationary phase because these stationary phases are more selective for aromatic-containing analytes. A fully non-aromatic stationary phase, like a "1-type" (100% dimethyl polysiloxane) column is not appropriate for this analysis since cannabichromene (CBC) and cannabidiol (CBD) will coelute completely. While 5-type columns (5% phenyl) contain some aromatic component, they generally also produce coelutions for cannabichromene and cannabidiol, depending on the conditions used. At best, CBC and CBD can be only partially resolved on 15 m 5% phenyl columns. Much better separations are obtained on higher phenyl-content phases, such as Rxi*-35Sil MS (35% phenyl type) and Rxi*-17Sil MS (50% phenyl type) columns, as they offer excellent selectivity for aromatic cannabinoids. Not only do both columns resolve cannabichromene and cannabidiol, the chromatograms in Figures 2 and 3 demonstrate that they also separate delta-8-tetrahydrocannabinol (d8-THC), delta-9-tetrahydrocannabinol (d9-THC), cannabigerol (CBG), and cannabinol (CBN). Although both columns perform well, the Rxi*-35Sil MS column is recommended because of the slightly faster analysis time and greater space overall between the peaks of interest.

While stationary phase selectivity is the most important factor in choosing a GC column for cannabinoid analysis, there are some additional aspects of this work that will benefit labs doing medical marijuana potency testing. First, cost savings were achieved by using a 15 m column. When a column with the proper selectivity is used, a 15 m column easily provides the separating power needed for this analysis at about half the cost of a 30 m column. Also, the 0.25 mm x 0.25 μ m format has good sample loading capacity and is robust, especially when a proper split injection is used with a Sky* Precision* split liner with wool. Finally, hydrogen carrier gas was used here instead of helium. Using hydrogen provides a faster analysis, increasing sample throughput. Hydrogen carrier gas is a convenient way to speed up run times, increase productivity, and reduce the cost and availability concerns associated with using helium carrier gas.

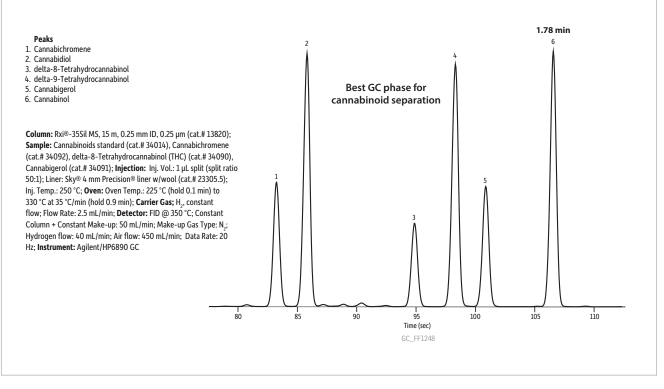


Pure Chromatography

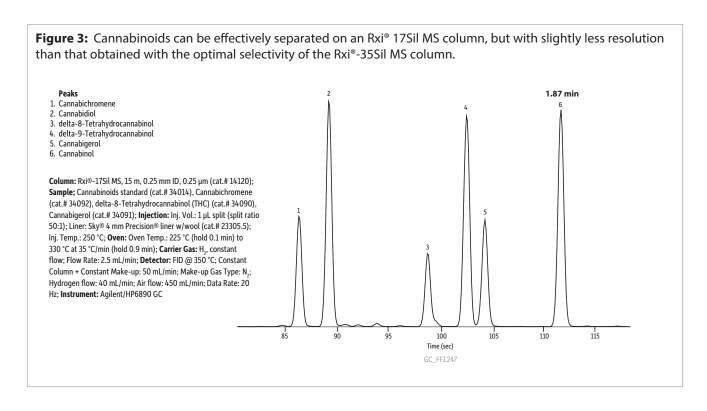
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Figure 1: Since cannabinoids are aromatic compounds, a GC column that contains aromatics in the stationary phase will provide much better separations than a column with a non-aromatic phase.

Figure 2: The Rxi®-35Sil MS column provides both the best separation and the fastest analysis time, making it the ideal GC column choice for medical cannabis potency testing.

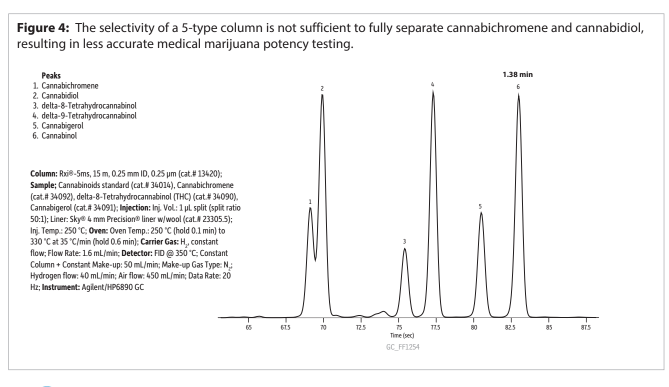


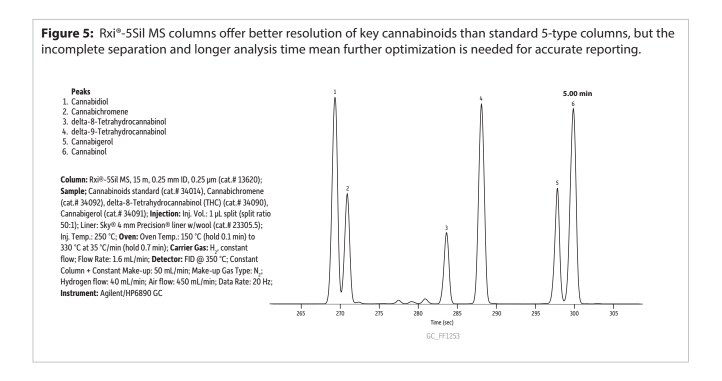
2



Adjusting Conditions for 5-Type Columns

While using an Rxi $^{\circ}$ -35Sil MS column provides the best selectivity and speed for cannabinoid analysis, cannabidiol potency can be determined in medical cannabis using a 5-type column under certain conditions. If you already have a 5-type column for this work, you can vary the GC conditions, especially carrier flow and oven temperature program, and still separate cannabichromene and cannabidiol, just not as quickly or easily as with the Rxi $^{\circ}$ -35Sil MS column. Figures 4 and 5 show this analysis on Rxi $^{\circ}$ -5ms and Rxi $^{\circ}$ -5Sil MS columns, respectively. Again, the 0.25 mm x 0.25 µm format was used here because it offers better efficiency than wider bore columns (e.g., 0.32 mm and 0.53 mm IDs), which may not separate cannabichromene and cannabidiol under any operational conditions.





Note that even though these are both 5-type columns, the elution order of cannabichromene and cannabidiol changed. This is due to two things. The first is that Rxi*-5ms and Rxi*-5Sil MS columns differ slightly in selectivity for certain compounds; even though they are both considered 5-type columns, they contain different stationary phases that retain some compounds differently. The second reason is that the GC oven programs are different for the columns, which means that the compounds are eluting at different temperatures. You may be able to further optimize the separation of cannabichromene and cannabidiol on a 5-type column, but the selectivity and faster analysis that can be obtained using a high-phenyl content Rxi*-35Sil MS column make it ideal for potency determinations in medical cannabis.

To sum things up, proper column choice is essential for accurate and robust cannabis potency testing. Using the right column not only gives you more confidence in your potency values, but it also saves you time and money. Switching to hydrogen carrier gas can reduce your costs even further, while increasing sample throughput.

Visit www.restek.com/medical-cannabis for Restek® GC and LC columns, accessories, reference standards, and other products and resources for medical marijuana analysis.



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Lit. Cat.# FFAR1954-UNV

High Quality Analysis of Pesticides in Marijuana for Medicine using QuEChERS, Cartridge SPE Cleanup, and GCxGC-TOFMS

Jack Cochran, Julie Kowalski, Sharon Lupo, Michelle Misselwitz, Amanda Rigdon

Restek Corporation, Bellefonte, PA, USA

Frank Dorman

The Pennsylvania State University, University Park, PA, USA

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Booming Business

Growers Dispensaries Edibles Labs





¥ Testing Potency

■ Safety



Cannabidiol, Δ^9 -THC, cannabinol, etc. Pesticides Microbiological Bacteria, mold, fungus, yeast





















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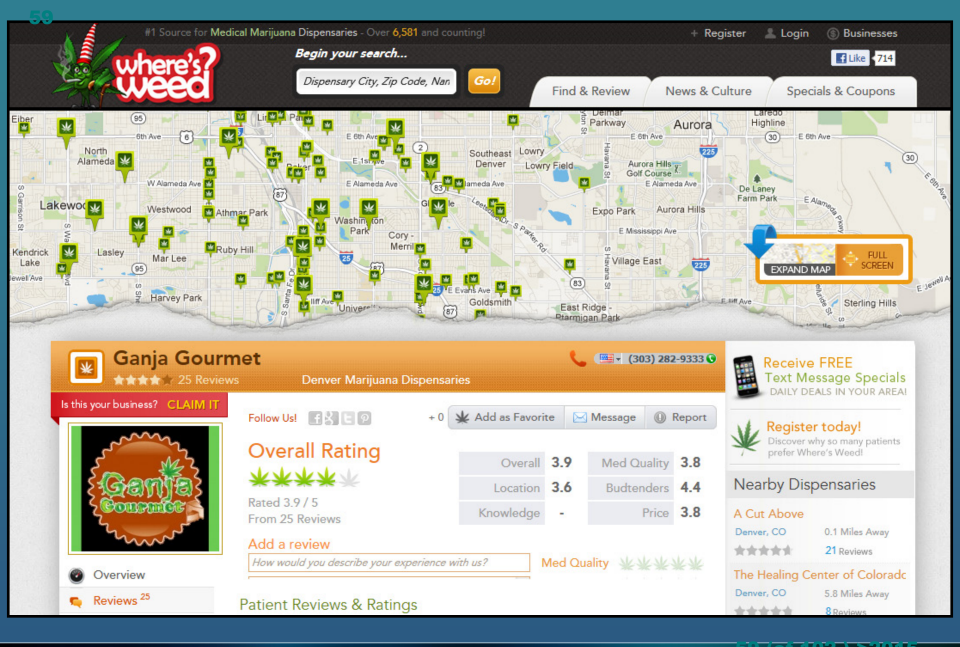




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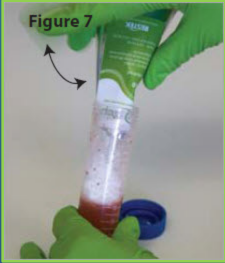


























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2 g ground marijuana - 50 mL centrifuge tube 10 mL MeCN 10 mL H₂O Shake to wet Soak one hour Add spikes and internal standards Vortex 30 min Add QuEChERS EN salts **Shake 1 min** Centrifuge 5 min at 3000g Remove extract for cleanup and analysis **GCxGC-TOFMS and LC-MS/MS**







MgSO₄ >>

GCB >>

PSA >>



Acetone:Toluene

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- 30m x 0.25mm x 0.25µm Rxi-5Sil MS
 - 5% phenyl (silphenylene) / 95% dimethyl
- Corrected constant flow He at 2.0 mL/min
- 80°C (1min), 5°C/min to 310°C
- Thermal modulation, 3 sec
- 1.3m x 0.25mm x 0.25μm Rtx-200
 - Trifluoropropylmethyl, selectivity for pesticides
 - +5° temp offset from primary column







LECO Pegasus[®] TOFMS for Pesticides

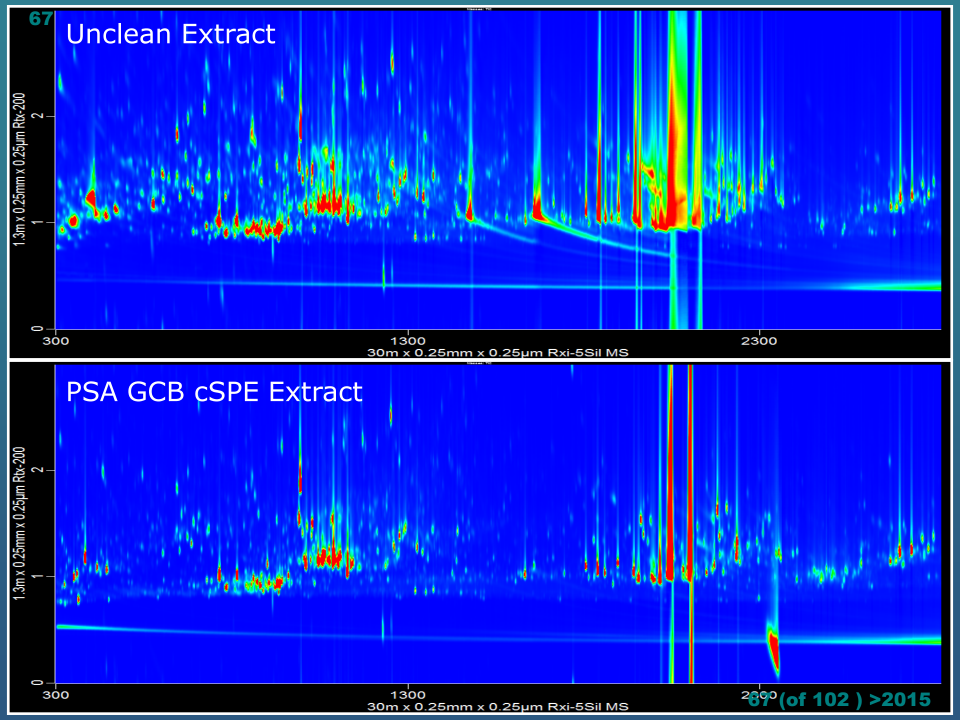
- Source temperature: 225°C
- Electron ionization: 70 eV
- Stored mass range: 45 to 550 u
- Acquisition rate: 100 spectra/sec

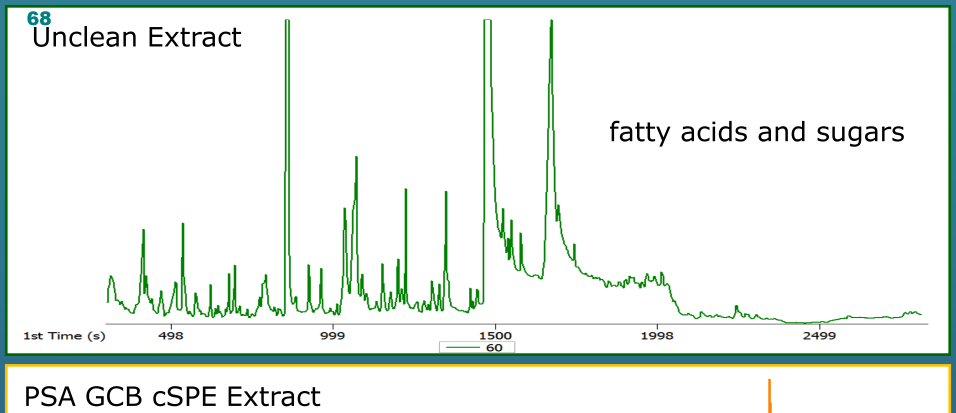


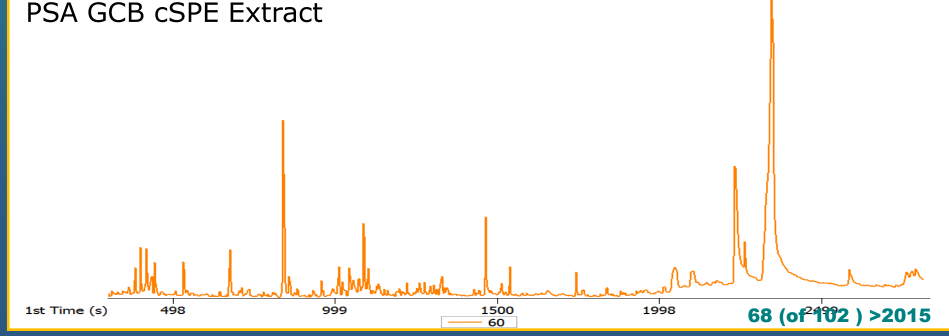
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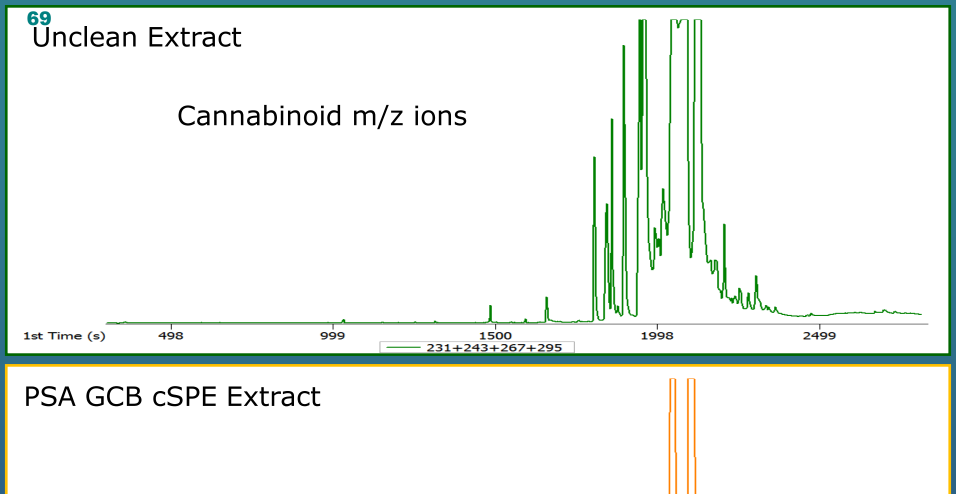


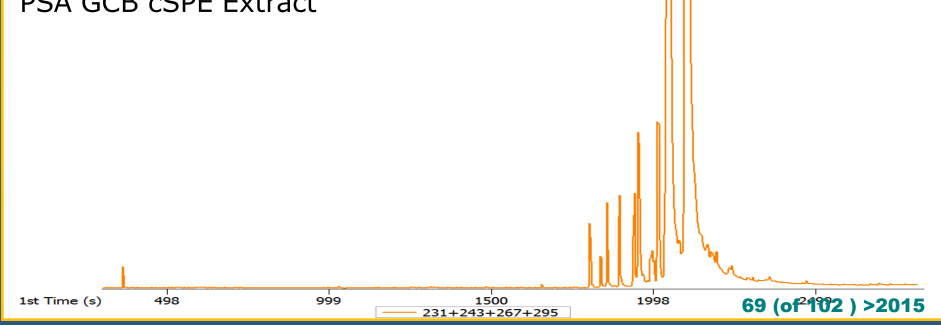












Pesticide Recoveries for Marijuana Spikes Later Eluting Compounds – Clean versus Unclean

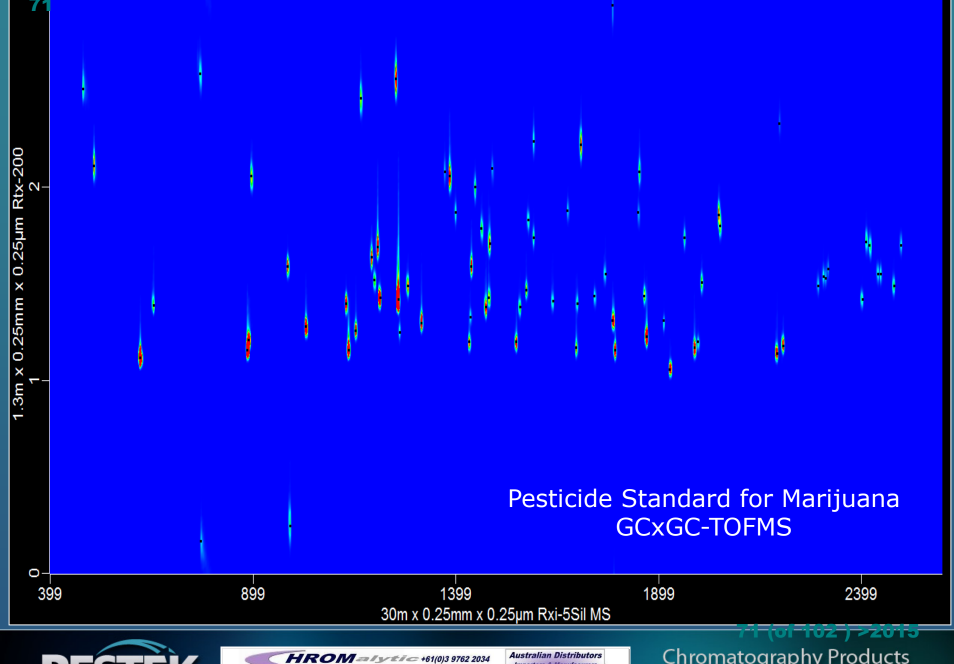




Pesticide	Classification	SB3 cSPE	SB3 No cSPE
4,4'-DDD	Organochlorine	83	230
4,4'-DDT	Organochlorine	77	9
Bifenthrin	Pyrethroid	86	89
Dicofol	Organochlorine	84	ND
Azinphos methyl	Organophosphorus	79	53
trans-Permethrin	Organochlorine	68	17
Pyraclostrobin	Strobilurin	73	19
Fluvalinate	Pyrethroid	72	23
Difenoconazole	Triazole	67	21
Deltamethrin	Pyrethroid 68		20
Azoxystrobin	Strobilurin	72	27

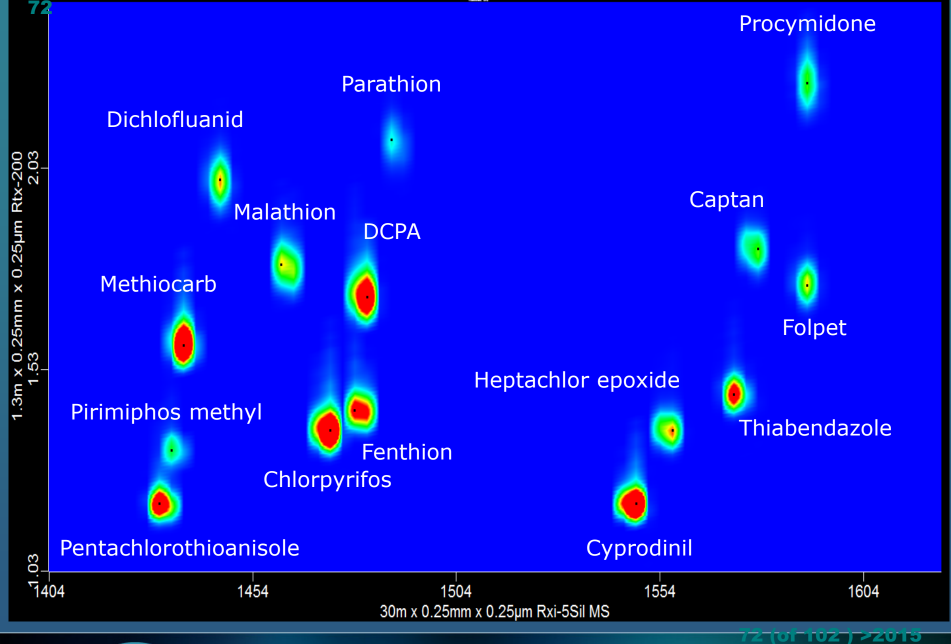
















Pesticide Recoveries for Marijuana Spikes

Pesticide	Classification	SB1 cSPE	SB3 Q + cSPE	S3 Q + cSPE
o-Phenylphenol	Unclassified	91	83	97
Tebuthiuron	Organonitrogen	100	104	94
Hexachlorobenzene	Organochlorine	73	44	71
Chlorothalonil	Organochlorine	77	80	81
Anthracene	QC STD	108	105	119
Diazinon	Organophosphorus	86	89	102
Carbaryl	Carbamate	91	103	100
Metalaxyl	Organonitrogen	93	96	90
Malathion	Organophosphorus	98	106	104
Chlorpyrifos	Organophosphorus	87	92	93
Captan	Organochlorine	71	80	91
Endosulfan I	Organochlorine	87	86	102





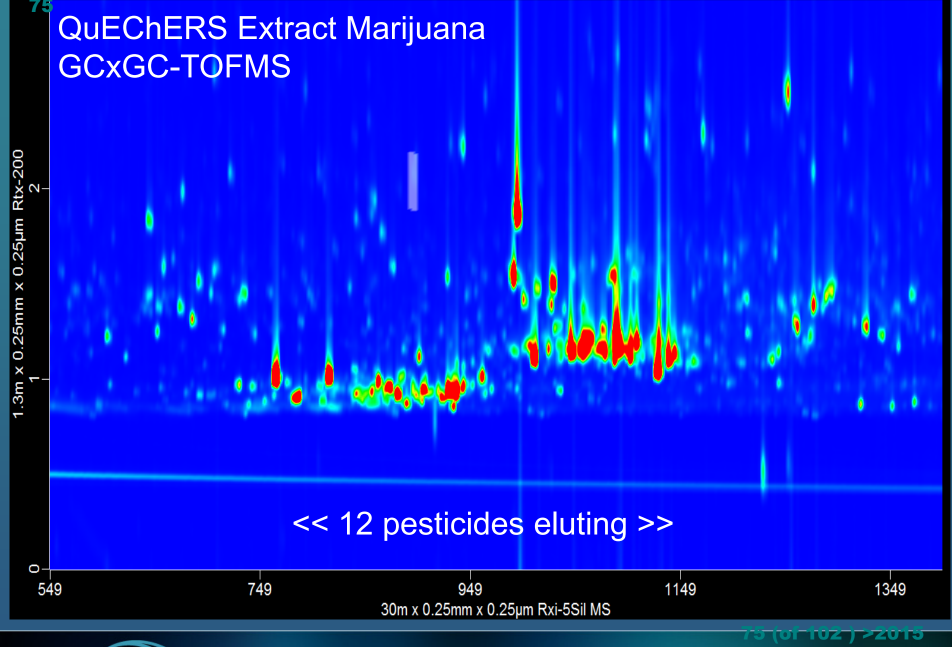
Pesticide Recoveries for Marijuana Spikes

Pesticide	Classification	SB1 cSPE	SB3 Q + cSPE	S3 Q + cSPE
Imazalil	Organonitrogen	83	77	91
Endosulfan II	Organochlorine	86	80	113
Endosulfan sulfate	Organochlorine	82	88	105
4,4'-DDT	Organochlorine	83	77	99
Bifenthrin	Pyrethroid	82	86	96
Dicofol	Organochlorine	40	84	73
Azinphos methyl	Organophosphorus	92	79	97
cis-Permethrin	Pyrethroid	72	64	91
trans-Permethrin	Pyrethroid	52	68	90
Cypermethrin	Pyrethroid	I	I	89
Deltamethrin	Pyrethroid	77	68	99

I = incurred pesticide

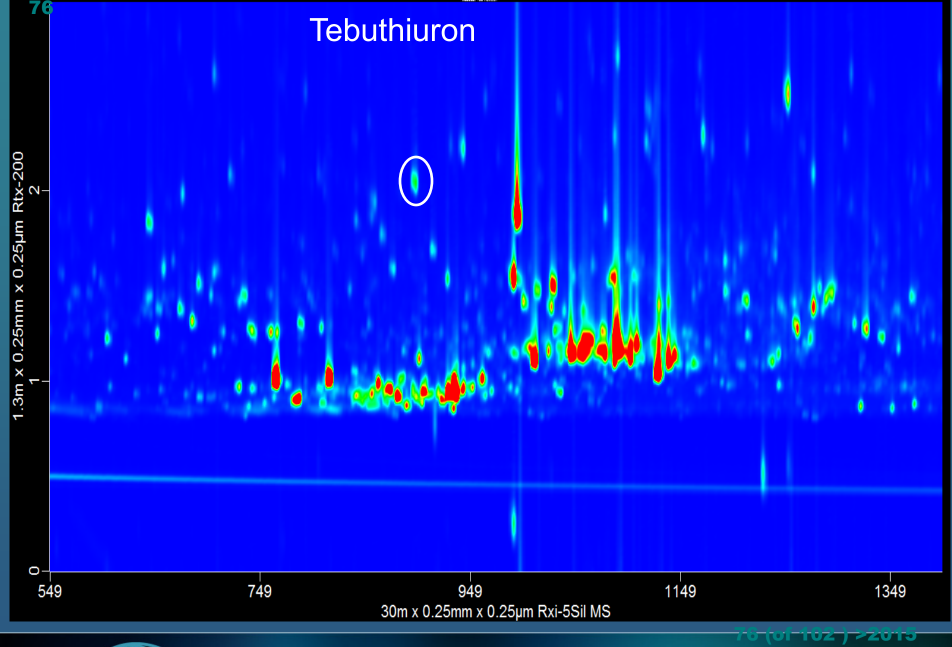






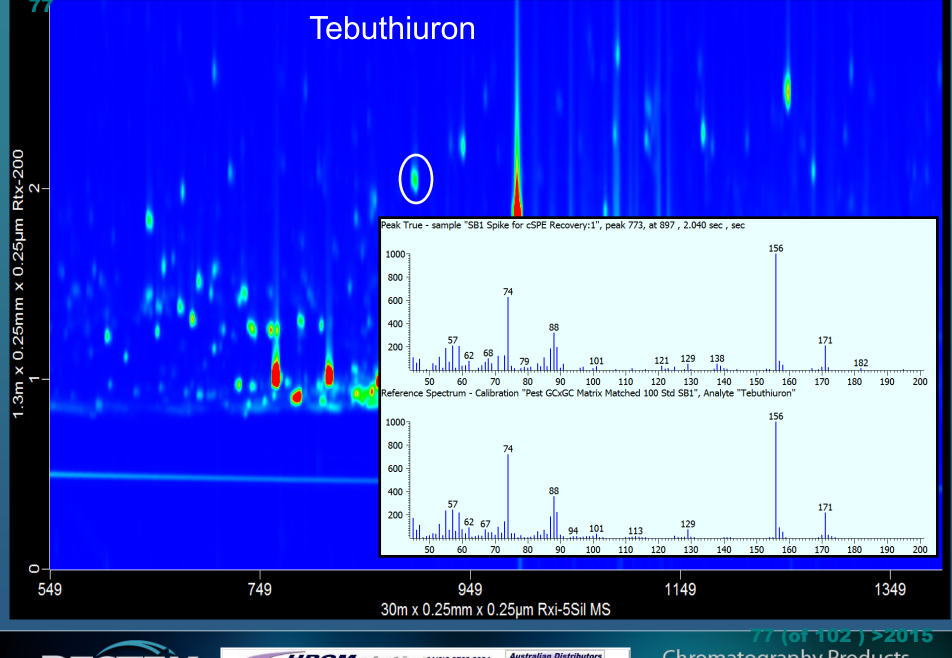
















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Hexachlorobenzene

Imazalil

OH 🖳	ÇI
	CI
	CI
	ĊI

SZ	Pesticide	ppb
	o-Phenylphenol	190
	Chlorothalonil	330
	Metalaxyl	400

CI	O—CH ₂	CI
		CI
$0 \rightarrow 0$	\times	

SB	Pesticide Pesticide	ppb
	o-Phenylphenol	58
	Chlorothalonil	29
	Cypermethrin	2200

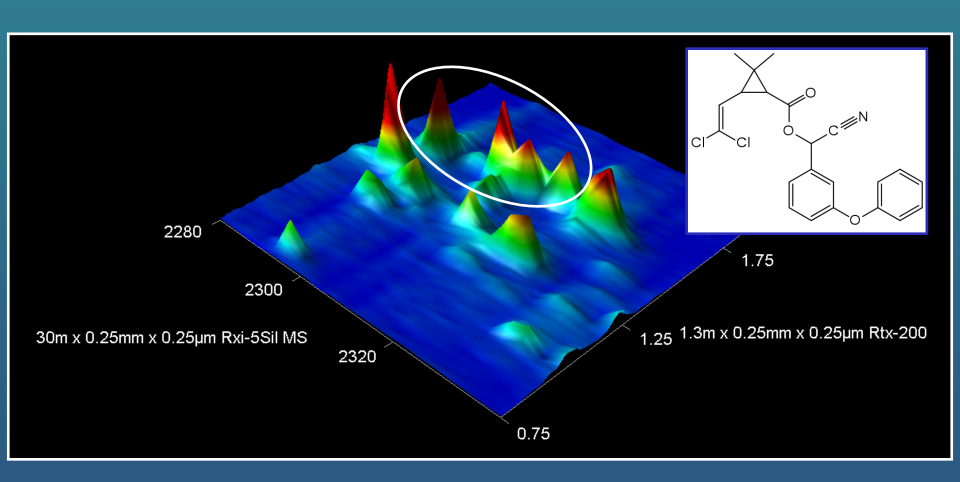




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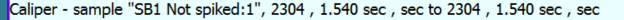
Incurred Cypermethrins in Marijuana m/z 163 – Quantification Ion

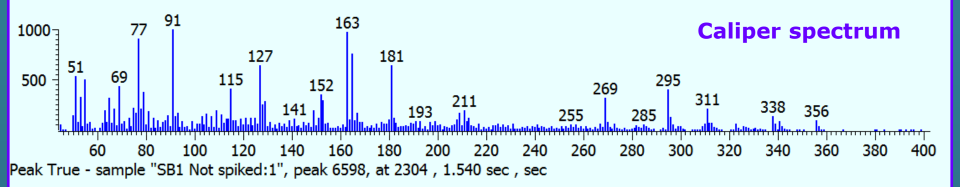


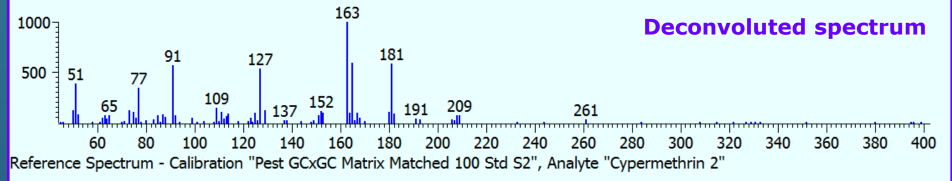


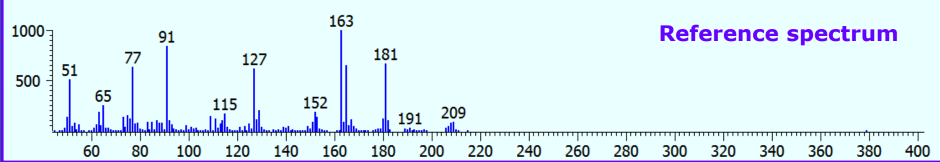


Incurred Cypermethrin in Illicit Marijuana – QuEChERS GCxGC-TOFMS













Incurred Pesticides in Marijuana Sample 2Q2

Pesticide	LC	GC
Imazalil	410	NA
Bifenazate	1100	2180
Piperonyl butoxide	37	41
trans-Permethrin	660	1100
cis-Permethrin	1200	690
o-Phenylphenol	NA	280
4,4'-DDE	NA	30



NA = not analyzed by this method

"Effervescent Health Formula"



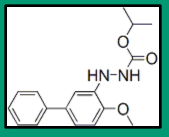


Bifenazate















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Summary

- QuEChERS is a viable extraction approach for cannabis, but cartridge SPE cleanup necessary
- GCxGC-TOFMS was very helpful in pesticide determinations in cannabis
 - Sample extracts are VERY complex
 - Detectability boost through thermal modulation process







Search

« Some answers on the Chromatography challenge posted before: Does a GC-capillary column produce different retention times when installed in the opposite direction?

LC/MS/MS Analysis of Synthetic Cannabinoid Metabolites in Urine – The Saga Begins »

First QuEChERS Extraction of Marijuana with GCxGC-TOFMS Analysis, dudes...

April 20th, 2011 by Jack Cochran

My Restek colleagues Julie Kowalski, Michelle Misselwitz, and Amanda Rigdon, along with Professor Frank Dorman from The Pennsylvania State University (PSU), report here what we believe is the first QuEChERS extraction of marijuana, with subsequent analysis using GCxGC-TOFMS. We were assisted in this task by Randy Hoffman, a Police Officer Specialist/Evidence Technician at PSU, who very kindly donated the samples confiscated from some students who probably should have had their minds on class, not grass.

Our interest in this topic is mainly about medicine, since at least 15 states (Pennsylvania is not one of them) and Washington DC have enacted laws to legalize medical marijuana. When you fill your prescription, how do you know your remedy is active (potency, or cannabinoid content), pesticide-free, and without bacteria or mold or fungus? Well, you probably don't, but eventually FDA might get involved and we'll need good, robust analytical methods, especially for pesticide analysis. We think that you might be able to do one extraction for both potency and pesticide determinations and we're high on QuEChERS, so we went for it.

First, the potency work, or cannabinoids determination. Although you don't need GCxGC for the BIG THREE (cannabidiol, Δ^9 -THC, cannabinol; by the way, Restek has a reference material containing these compounds...), we used it to illustrate one of the benefits of that technique, the structured chromatogram. In the first figure below, the GCxGC contour plot (or chromatogram), you can see that compound classes position themselves in certain areas. This helps identification, and makes discovery of new compounds within classes a bit easier (e.g. perhaps there are undiscovered cannabinoids out there with medicinal benefits). Zooming in, we can see the terpenoid classes, which are thought to have therapeutic effects. Finally, you can see the cannabinoids, including cannabidiol, one of much interest given that "it has been shown to relieve convulsion, inflammation, anxiety, and nausea, as well as inhibit cancer cell growth" (http://en.wikipedia.org/wiki/Cannabidiol).

We quantified cannabidiol (CBD), Δ^9 -THC (THC), and cannabinol (CBN) for 4 marijuana samples using QuEChERS and GCxGC-TOFMS and the results are presented in the table below. Since the samples had been stored in an evidence locker for over a year in some cases, the CBN content is relatively high versus fresh marijuana. CBN increases as THC degrades. The THC content is in line with what is typically reported for higher grade illicit marijuana.

Stay tuned for a report on pesticide analysis of marijuana using QuEChERS and GCxGC-TOFMS. As you might imagine, the extracts are extremely complex, similar to what we saw in our dietary supplements work.





Marijuana for grinding prior to QuEChERS extractions.





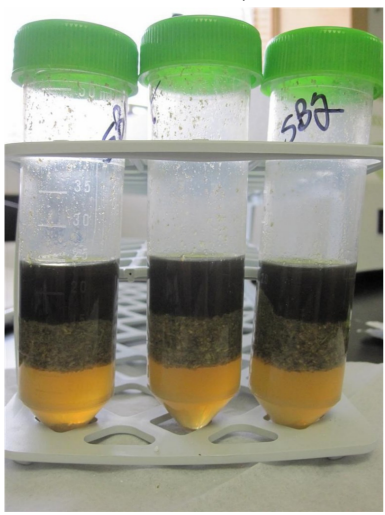
Professor Frank Dorman at Penn State University grinds the goods.





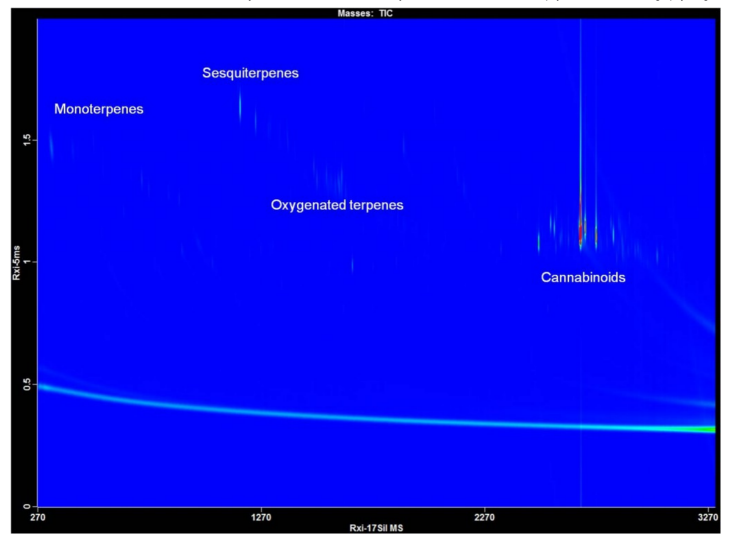
Weighing the marijuana into the QuEChERS extraction tubes. It is full of static!





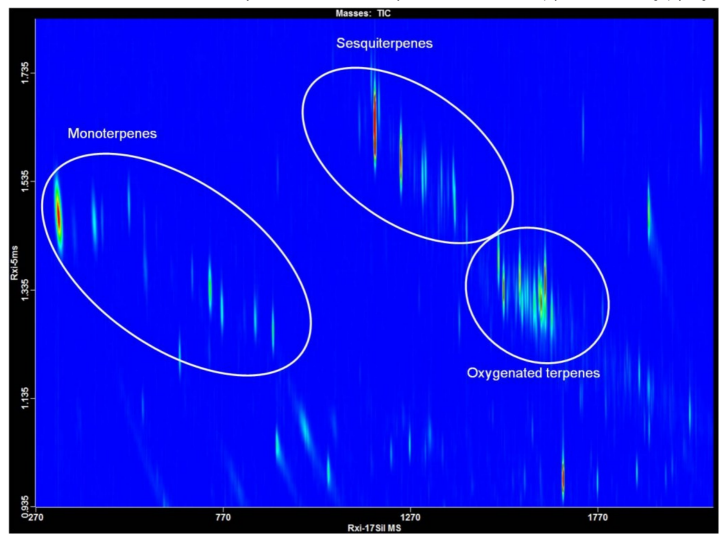
The first QuEChERS extracts of marijuana. They are almost black, and are very complex.





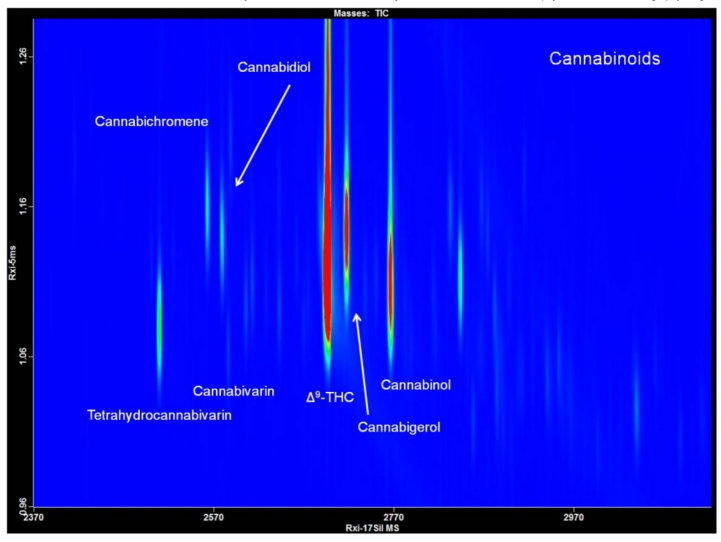
GCxGC-TOFMS contour plot of QuEChERS marijuana extract showing "structured chromatogram", where compound classes elute in certain regions. Rxi-17Sil MS x Rxi-5ms column combination.





Zooming in on the GCxGC terpenoid region for QuEChERS extracts of marijuana.





The cannabinoid region of the GCxGC chromatogram of a QuEChERS extract of marijuana.

Sample	CBD	THC	CBN
S1	0.029	12	1.1
S2	0.016	4.3	1.3
S3	0.034	9.0	1.3
SB9	0.15	10	1.7

Cannibinoid results in percent for samples of marijuana analyzed by QuEChERS and GCxGC-TOFMS.

This entry was posted on Wednesday, April 20th, 2011 at 11:14 pm and is filed under <u>GC/MS</u>, <u>QuEChERS</u>, <u>GCxGC</u>, <u>Medical Marijuana</u>. You can follow any responses to this entry through the <u>RSS 2.0</u> feed. You can <u>leave a response</u>, or <u>trackback</u> from your own site.

6 Responses to "First QuEChERS Extraction of Marijuana with GCxGC-TOFMS Analysis, dudes..."





First QuEChERS Extraction of Marijuana with GCxGC-TOFMS Analysis, dudes... « ChromaBLOGraphy: Restek's Chromatography Blog

Great work! I have been doing a Quechers extraction of cannabinoids for edible food products as well as a Quechers extraction of cannabis flowers for pesticide residue testing for over a year. Our laboratory tests medical cannabis samples in California. We would love to see restek offer standards for CBG, THCV, CBC, as well as THCA and CBDA (THCA and CBDA are the major cannabinoid constituents of raw cannabis flowers but are degraded into THC and CBD on a GC column).

I would love to share some data with your group if you are interested.

Thanks, Josh Wurzer Laboratory Director SC Laboratories Inc.



Greetings Josh!

Thanks so much for your kind comments and for letting me know of your use of QuEChERS. I'll forward your reference materials request to our standards group. I do know they are interested in expanding the line, but some of those neat compounds are SO expensive. I'm just getting ready to post on our pesticide results for the illicit marijuana we extracted. We found numerous pesticides, so the work turned out to be quite interesting. The samples are unbelievably complex, and needed a multidimensional technique, in this case, GCxGC, for the quantitative effort.

Regards,

Jack

3. Blake Meinert says:
May 19, 2011 at 8:45 pm

Hello Jack,

I have had substantial experience analyzing for chlorinated pesticides in soil and water, but I had not considered their use in marijuana crops. What pesticides did you primarily see? Is there one primary pesticide of choice with growers that you know of? This is interesting.

Thanks, Blake

4. <u>Jack Cochran</u> says: May 23, 2011 at 2:47 pm

Hi Blake:

The pesticides we saw on our small sample size were o-phenylphenol, hexachlorobenzene, metalaxyl, chlorothalonil, imazalil, and cypermethrin. Interestingly, all are fungicides except for the insecticide, cypermethrin. Mold/fungus apparently is a big problem for marijuana that is being dried/stored, so maybe this finding isn't surprising.

With the small sample size we had, and the fact that all of our samples were illicit marijuana, I'm not sure if there is a "primary pesticide of choice". At least with the medical marijuana, it may be that bifenazate (Floramite) and abamectin (Avid) are the "pesticides of choice" to control spider mites in indoor grow operations. But since this doesn't seem to be a well regulated area yet, I'm not sure if we know what to expect as regards pesticide use.

Jack

5. <u>High Quality Analysis of Pesticides in Marijuana using QuEChERS, cartridge SPE cleanup, and GCxGC-TOFMS « ChromaBLOGraphy</u> says:

May 22, 2011 at 3:28 am

- [...] we reported on what we believe is the first application of QuEChERS for marijuana, using it for potency analysis with GCxGC-TOFMS. Ultimately, the plan was to determine pesticides [...]
- 6. *The Bard Hits the Bong? « ChromaBLOGraphy* says: July 2, 2011 at 9:47 pm
 - [...] cannabis to (1) develop methods for possibly fingerprinting marijuana types, (2) characterize marijuana potency, and (3) analyze for pesticides in marijuana with GCxGC-TOFMS and [...]

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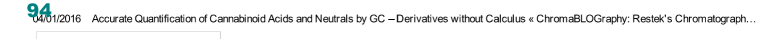
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Accurate Quantification of Cannabinoid Acids and Neutrals by GC – Derivatives without Calculus

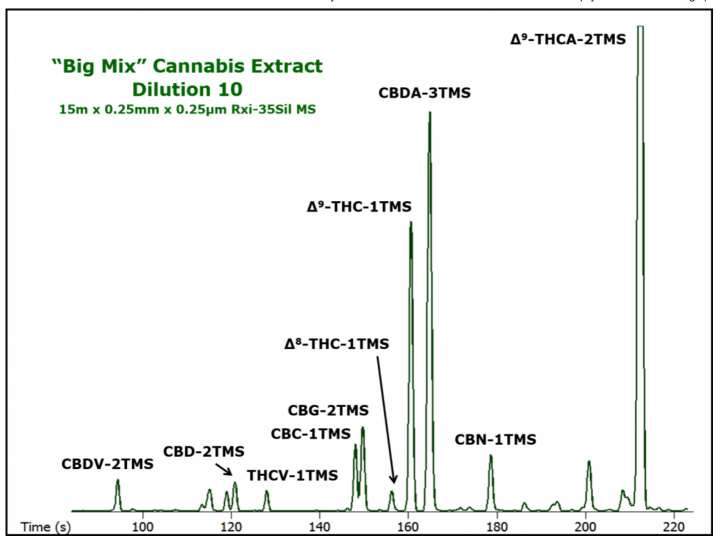
September 9th, 2015 by Amanda Rigdon

Derivatization is a widely-used technique for GC sample preparation across many industries and in widely varied matrices from soil to plastics to blood that is used to make polar and active compounds more amenable to good GC analysis. If you're careful about testing your derivatization procedure during method development, you can be confident that you'll have a reproducible method that can vastly improve the quality of your GC results. While derivatization does require some extra sample handling, the procedure I developed for cannabis plant matrix is very straightforward and easy to perform:

Derivatization Procedure for Cannabis and Hemp Plant Matrices:

- Place 100μL of plant extract into a <u>1mL Micro-Vial</u>
- Evaporate to dryness at 50°C under a gentle stream of nitrogen
- Add 50μL ethyl acetate and 50μL BSTFA + 1% TMCS
- Incubate at 70°C for 30 minutes
- Cool and dilute with ethyl acetate if desired

In my <u>last blog</u>, I introduced the concept of derivatization for use in cannabis or hemp cannabinoid testing. Once acidic cannabinoids are derivatized, they no longer break down in the GC inlet and can be quantified separately from the neutral cannabinoids. I demonstrated this through derivatization of high-level solvent standards, but work with solvent standards is a far cry from matrix work, which means the procedure needed to be tested in matrix. To kick off the matrix test, I spiked an extract with the most common cannabinoids of interest, derivatized it using the procedure listed above, and my colleague, Jack Cochran, analyzed it via GC-FID with our <u>Rxi-35Sil MS</u> GC column. We can see that we have a beautiful chromatogram with all of the derivatized cannabinoids separated, and very little matrix interference.



In addition to confirming that all derivatization sites are indeed derivatized by analyzing the standards with GC-MS (this is shown in my last blog), we also tested derivatization efficiency using a cannabis extract previously generated at Penn State University with the help of Professor Frank Dorman and a Police Officer Specialist. Because derivatization is a chemical reaction, the derivatization reagent gets used up during the derivatization reaction. Because plant matrix contains many other derivatizable compounds like sugars and sterols, these other compounds may compete for the derivatizing reagent, possibly resulting in the reagent getting used up before all of our analytes of interest can be derivatized.

So how can we be sure our derivatization is going to completion in the presence of matrix? There are a couple things we can do, the first of which is really simple. We can see in our procedure that we use a hefty $50\mu L$ of derivatizing reagent per $100\mu L$ of cannabis extract. We know that our extract contains a lot less than 50mg of plant matrix, not all of which is derivatizable. This means that by adding 50mg of BSTFA per $100\mu L$ of sample, we can be confident that we have a significant excess of derivatizing reagent as compared to derivatizable groups in our sample. Excess derivatizing reagent means that it will never be completely used up, ensuring the reaction will go to completion no matter what.

A more quantitative way to test derivatization efficiency in a matrix where you can't get blanks is to evaluate analyte linearity with differing amounts of matrix. For example, if you derivatize four THCA-containing samples prepared using $10, 20, 50, \text{ and } 100\mu\text{L}$ of cannabis extract and plot the area of THCA versus sample amount, you should end up with a straight line if your derivatization is going to completion. If it's not, then you'll likely see THCA area fall off for the samples containing more matrix since the derivatization reagent is being used up before all the analyte in the higher matrix level sample is derivatized. To test our procedure, we did just that. We can see that our linearity looks beautiful for all of the cannabinoids, indicating the derivatization does indeed go to completion.

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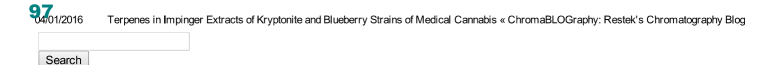
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« <u>Contents inside your baseplate trap</u> <u>Something to think about before you buy antimicrobial soaps</u> »

Terpenes in Impinger Extracts of Kryptonite and Blueberry Strains of Medical Cannabis

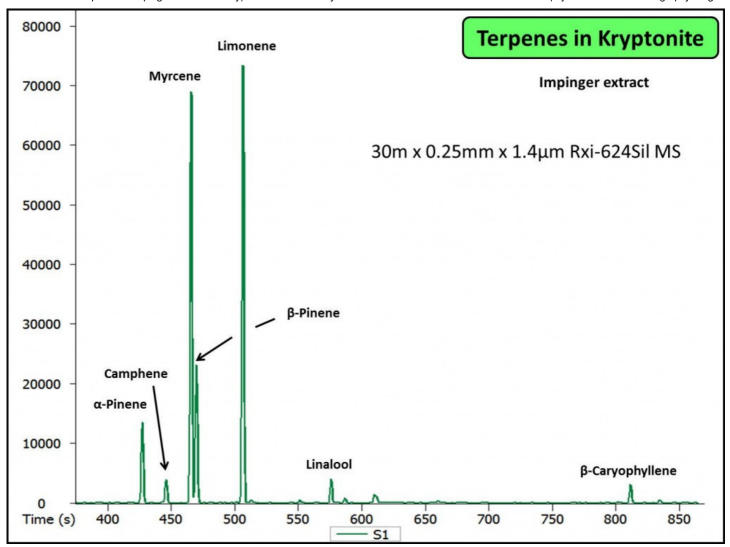
March 17th, 2014 by Jack Cochran

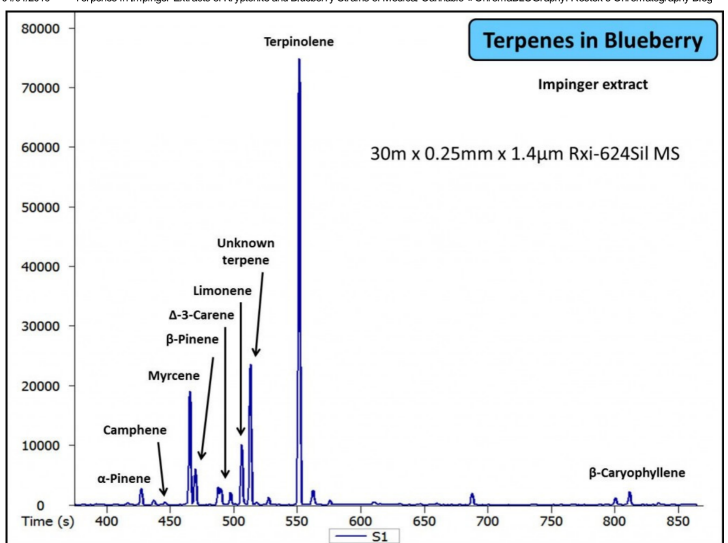
As noted in my earlier post, <u>Terpenes in Medical Cannabis</u>, terpenes are an important class of aroma compounds that may contribute to the medicinal benefits of cannabis, via the so-called "<u>entourage effect</u>". I profiled some of the terpenes listed as important for medical cannabis using our 30m x 0.25mm x 1.40µm Rxi-624Sil MS, achieving a promising separation on a standard I put together. Shown below are some impinger extracts provided by <u>SRI Instruments</u> for Kryptonite and Blueberry strains of medical cannabis. Importantly, these extracts do NOT contain any cannabinoids, which would elute late, if at all, from the thick-film 624Sil MS column, nor do they contain chlorophyll, another compound that plays havoc with GC inlet liners and stationary phases. Part of the beauty of headspace extraction techniques for terpenes is leaving the involatile material behind, and in this case, compressed air was used to sweep the terpenes from the cannabis to a vial containing methanol for trapping the terpenes.

As you look at the chromatograms below, it is important to note that this is ONLY qualitative work at this point and that different headspace methods (e.g. purge-and-trap, static headspace, SPME, etc.) could yield much different chromatograms. Solvent extraction or steam distillation, would likely be even more different, including resulting in more intense peaks for later eluting (less volatile) terpenes. The point of this work is to show initial efforts to characterize chromatographic elution order for some medical marijuana terpenes and analyze the first "real world" samples to show how terpene profiles for different medicines can be dissimilar.

Take a look back on the <u>GCxGC-TOFMS</u> work that shows very nice multidimensional separations of terpenes, sesquiterpenes, and oxygenated terpenes in a solvent extract for cannabis.

Restek continues to support the medical cannabis analysis community with GC and LC columns, accessories, and reference materials. Check out our <u>Medical Marijuana web page</u>.



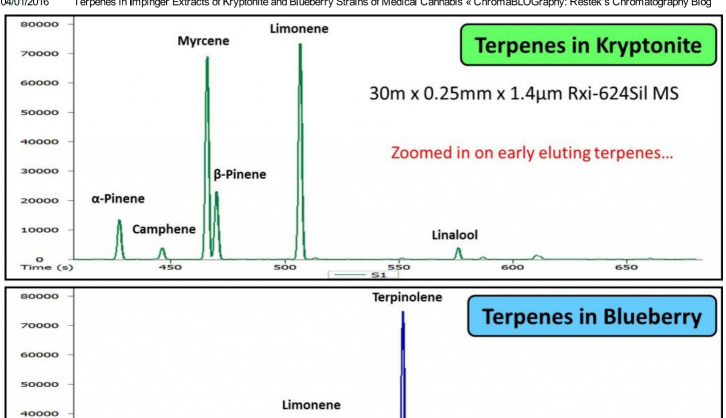




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Unknown

terpene

y-Terpinene

Linalool

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Δ-3-Carene

β-Pinene

Myrcene

Camphene

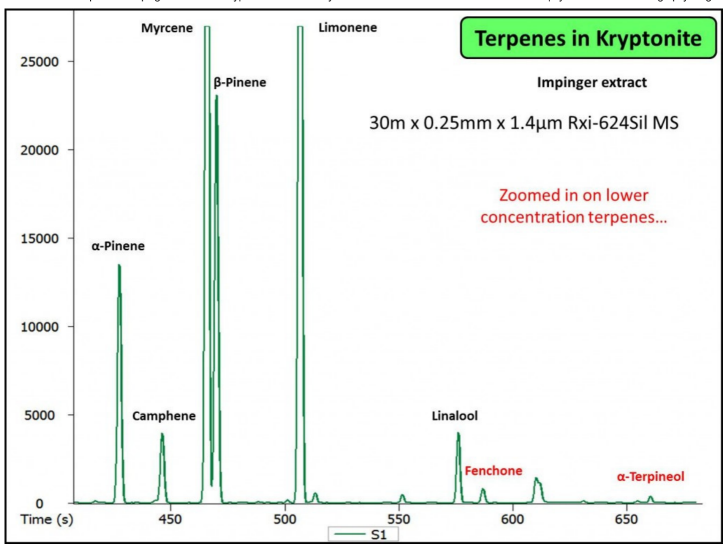
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α-Pinene

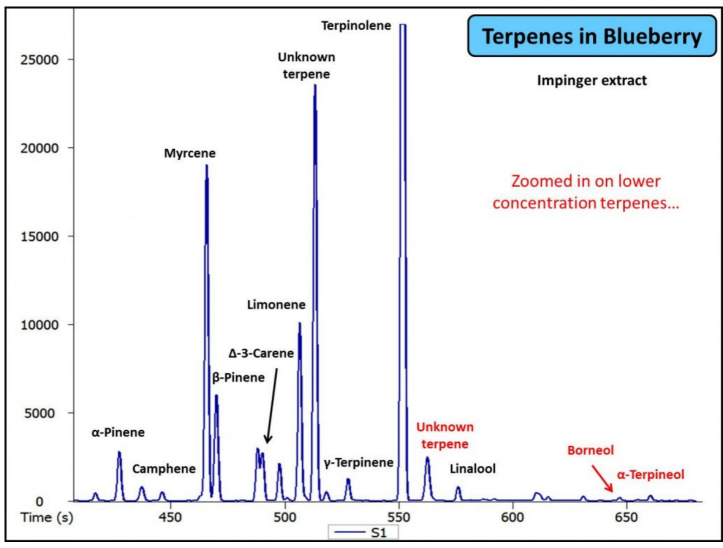
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This entry was posted on Monday, March 17th, 2014 at 6:32 pm and is filed under QuEChERS, GCxGC, Medical Marijuana. You can follow any responses to this entry through the RSS 2.0 feed. You can leave a response, or trackback from your own site.

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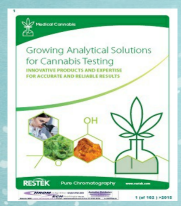
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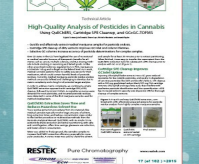
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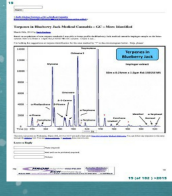






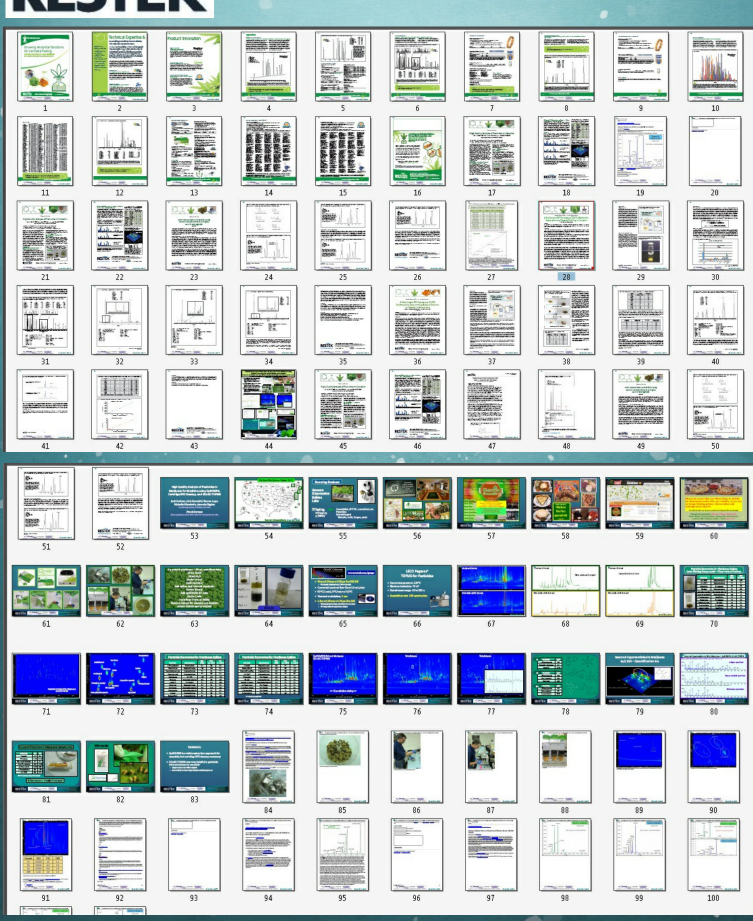






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- 4 High Quality Analysis of Pesticides in Marijuana for Food and Medicine using Quechers, Cartridge SPE, GCxGC-TOFMS and LC-MS-MS_MJrafa2011_med_mj_1-1
- 5 Don't Overestimate Cannabidiol During Medical Cannabis Potency Testing by GC_FFAR1954-UNV_1-4
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- 8 Accurate Quantification of Cannabinoid Acids and Neutrals by GC Derivatices without Calculus Blog
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- 10 Terpenes in Blueberry Jack Medical Cannabis GC More Identified

See SRI GCs-Cannabis - for some h'ware related Custom GCs and accessories

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