Practical Steps in GC Troubleshooting

Techniques, Tips, and Tricks

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What went wrong and how to fix it...

- Common problems
- **Troubleshooting tools**
- Troubleshooting examples



"Everything was just fine and then this happened!

How do I go about TROUBLESHOOTING?"



"Everything was just fine and then this happened!"

Logic = Something changed (slowly or sudden) = Something is different

Track Events – log book

- -Changed column, liner, septum, syringe, etc.
- -Injected samples, other method, etc.

-Did maintenance, cut column, inlet flush, etc.



Logical Troubleshooting

Troubleshooting Starts with Isolating the problem

- •There are 5 basic areas from where the problem arises
- INJECTOR
 FLOW
 COLUMN
 DETECTOR
 ELECTRONICS

•But of course it can always be some COMBINATION

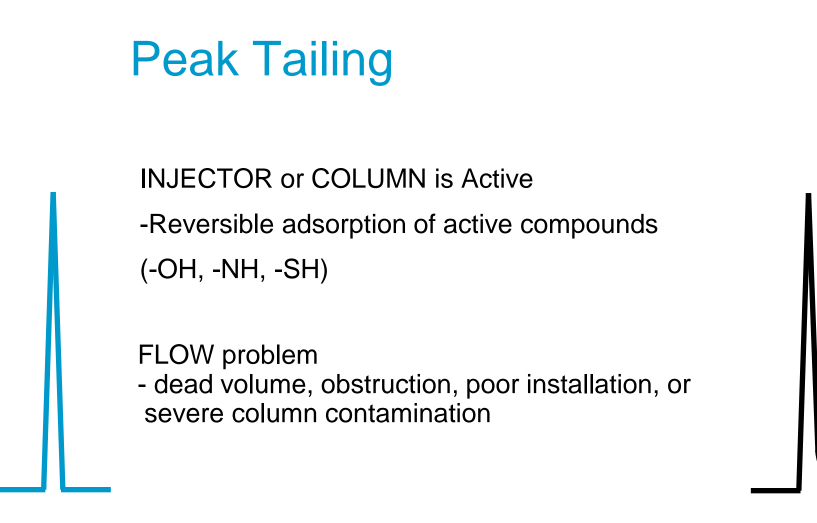
Knowing what can & <u>can't</u> cause the symptom is the key



Typical Problems of Optimized Methods becoming Unoptimized...and the Reason Why.

- Peak Tailing Flow Path or Activity
- Bonus Peaks In Sample or Back Flash (Carry Over)
- Split Peaks Injector Problems, Mixed Solvent
- No Peaks Wasn't Introduced, Wasn't Detected
- Response Changes Activity, Injector Discrimination, Detector Problem
- Peak Fronting Overload or Solubility Mismatch, Injector Problems
- Shifting Retention Leaks, Column Aging, Contamination or Damage
- Loss of Resolution Separation Decreasing, Peak Broadening
- Baseline Disturbances Column Bleed, Contamination, Electronics
- Noisy or Spiking Baseline Electronics or Contaminated Detector
- Quantitation Problems Activity, Injector or Detector Problems



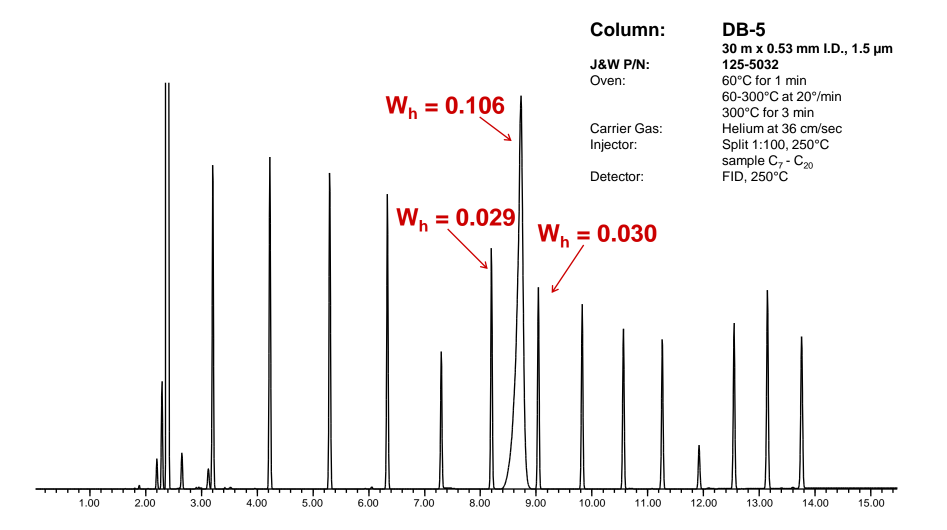


Miscellaneous - overloading of PLOT columns, co-elution, polarity mismatch between phase, solute or solvent, and some compounds always tail

*Tip = Inject a light hydrocarbon, should not tail unless flow path problem.



Bonus Peaks





Bonus Peaks or Ghost Peaks

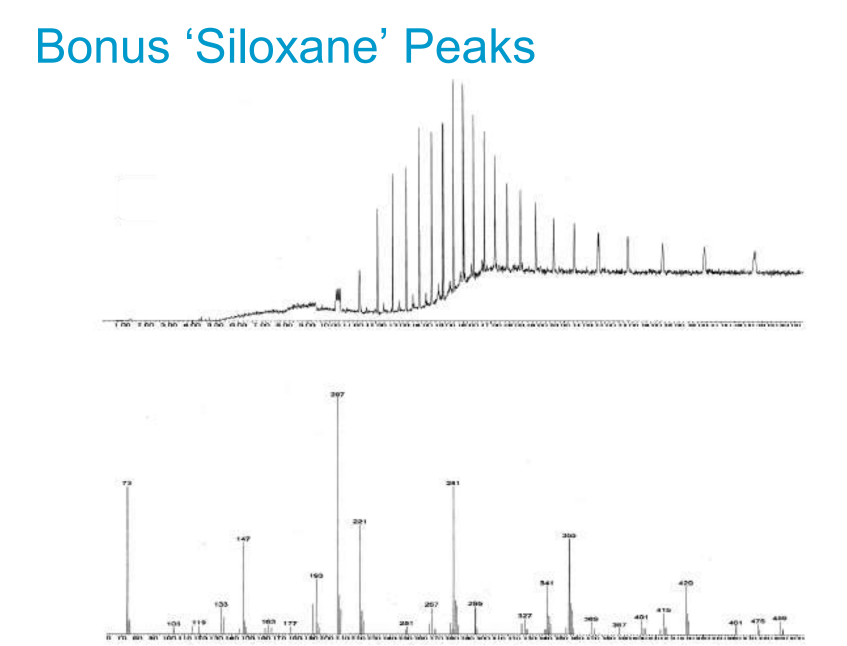


Contamination in INJECTOR, COLUMN or FLOW (carrier gas)

- -Carry-over from a backflash or previous sample
- -Bad tank of gas or traps have expired
- -Septum bleed

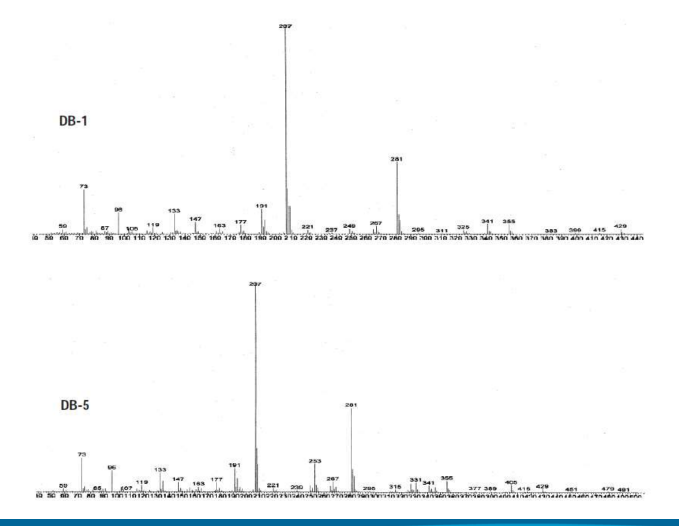
*TIP = Run a blank run...it should be blank!





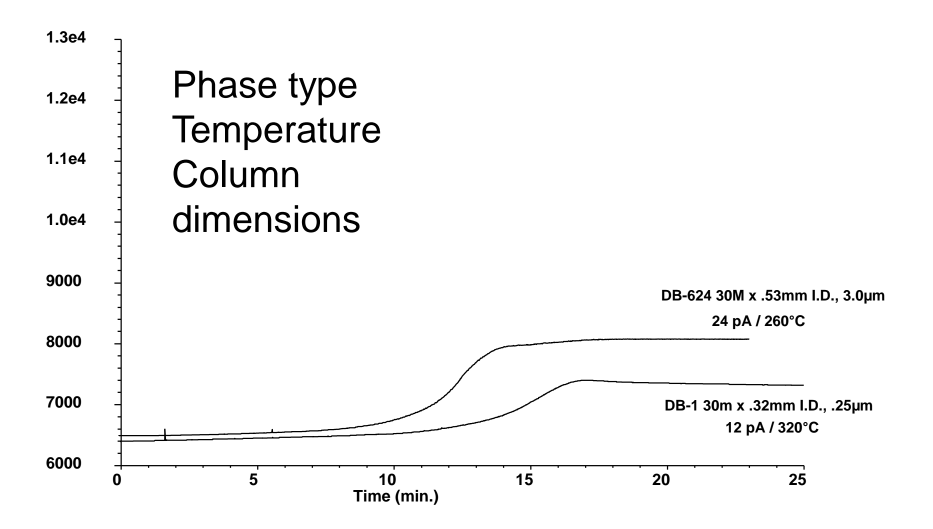


GC Column Bleed Ions





Column Bleed is Influenced by:





INJECTOR (poor sample introduction)
-Injecting the sample twice (some how?)
-Mixed sample solvent (polarity difference)
-Sample in syringe needle (manual inject)

Split Peaks

INJECTOR (activity)

-Breakdown (not really a split peak, 2 peaks)

-Sample degradation in injector

VOLATILITY

High boilers dropping out on Cold Spots

- -Transfer line temps
- -Unions or fittings not tracking column temp



No Peaks

DETECTOR (not on or not operational)

INJECTOR (not working)

-Plugged syringe/plunger not moving

-Wrong injector (or detector)

-Huge leak (older systems)

-No carrier gas flow

NOT the COLUMN Unless...

-Broken column or No column



Peak Response

All Change in Size

INJECTOR

-Leaky syringe

-Split ratio set incorrectly

-Wrong purge activation time

-Septum purge flow too high

-Injector temperature too low*

DETECTOR (response problem)

- -Settings or flows changed
- -Electronics failing

*Tip = Ask is it all of them or some of them, if all then injector or detector



Peak Response

Some Change in Size

INJECTOR or COLUMN is active/contaminated

-Irreversible adsorption of active compounds (-OH, -NH, -SH)

- -Decomposition of sample
- -Temperature Change Discrimination
- -Evaporation from sample

*Tip = If only some change, then ask which ones? If active compounds then activity. If tracks volatility then cold spots or inlet discrimination.



Peak Fronting

Shark Fin Shaped or Just Slight

COLUMN (contaminated)

-Overload (More pronounced with large solute and phase polarity differences)

INJECTOR

-Column installation

-Compound very soluble in injection solvent (need retention gap)

-Mixed sample solvent

OTHER

- -Co-elution
- -Breakdown



Retention Time Shift

4.75

INJECTOR

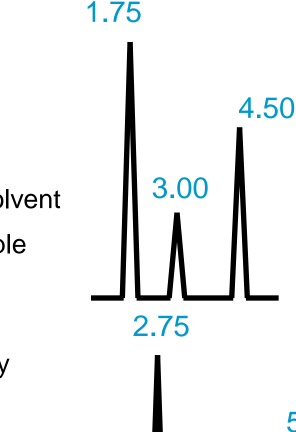
- -Leak in the septum
- -Change in injection solvent
- -Large change in sample concentration

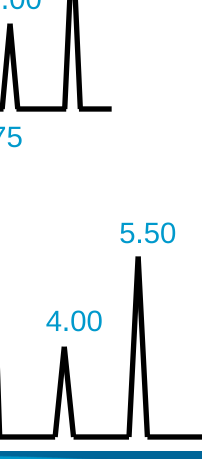
FLOW

-Change in gas velocity

COLUMN

- -Contamination
- -Damaged stationary phase
- -Loss of stationary phase
- -Change in temperature

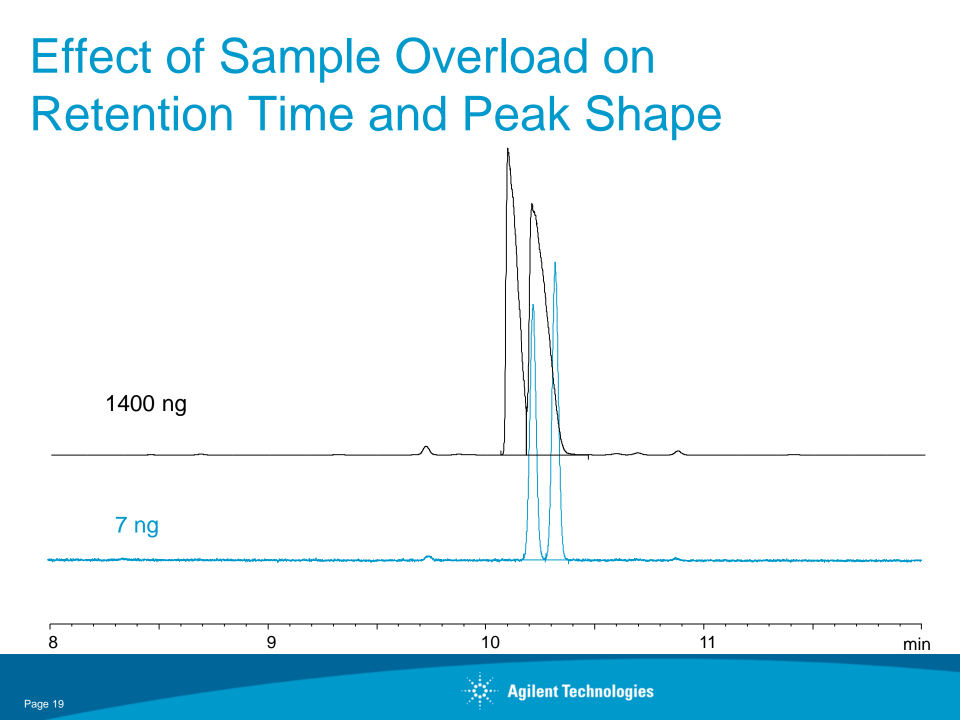




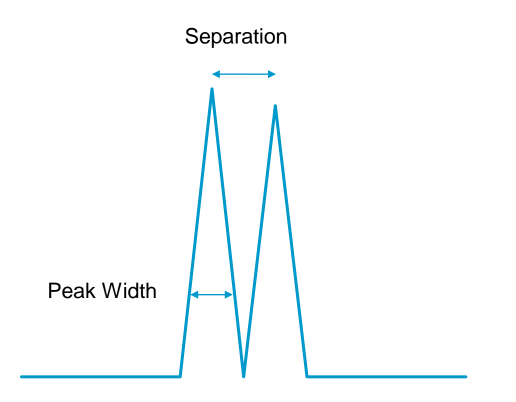


2.00

3.25



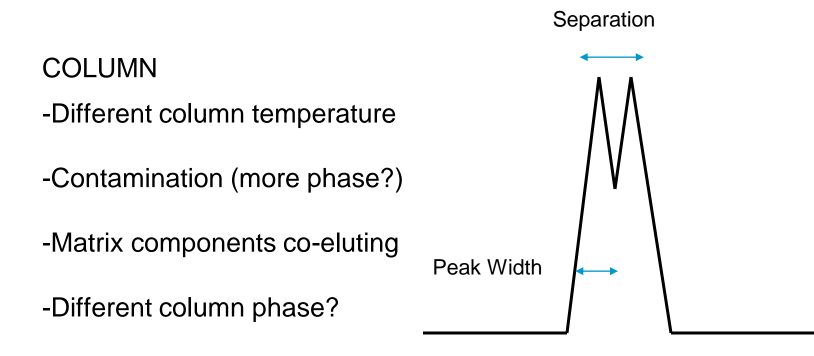
Loss of Resolution



Resolution is a function of separation and peak width

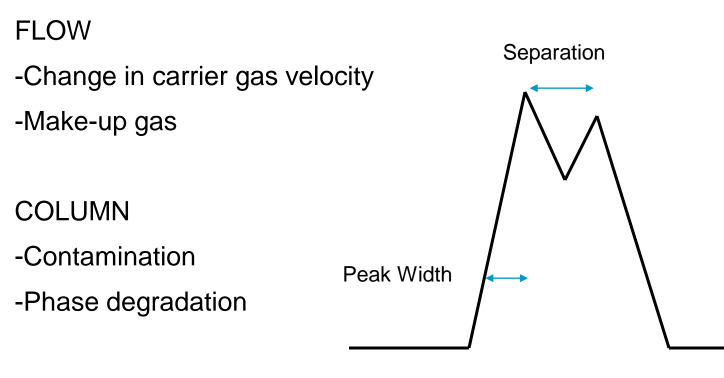


Loss of Resolution - Separation Decrease





Loss of Resolution - Peak Broadening



INJECTOR (efficiency)

-Settings, Liner, Installation, etc.



Baseline Disturbances



COLUMN or DETECTOR

-Not fully conditioned or stabilized (electronics)

-Contamination

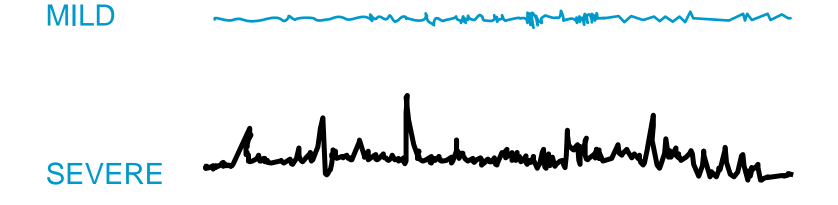
FLOW

-Changes in carrier and/or detector gas flows

-Valves switching, leaks



Noisy Baseline



- FLOW
- -Contaminated gas
- -Incorrect detector settings
- COLUMN
- -Bleed if at high temperature
- -In detector flame (poor installation)

DETECTOR

-Air leak - ECD, TCD

-Electronics malfunction

Spiking Baseline

DETECTOR

-Particles entering the detector

-Random: poor connection

-Regular: nearby "cycling" equipment (electronics)



Quantitation Problems

DETECTOR

-Poor stability (electronics) or Baseline disturbances (contamination)

-Outside detector's linear range or wrong settings

Activity (adsorption) in INJECTOR or COLUMN 392538403

OTHER

INJECTOR

-Technique, settings, conditions

-Syringe worn

-Co-elution

- -Matrix effects
 - -Sample evaporation leaky vials
 - -Sample decomposition



What is NOT caused by a Column???

Peaks!!

Any reproducible, sharp 'chromatographed' peak!

Siloxanes

Degradation product peaks: Endrin Aldehyde, Endrin Ketone, DDE, DDP....

Carryover of sample compounds

Splitting of peaks



Troubleshooting "Tools"

Bleed Profile: *baseline problems*

Inject a non-retained peak: peak shape problems

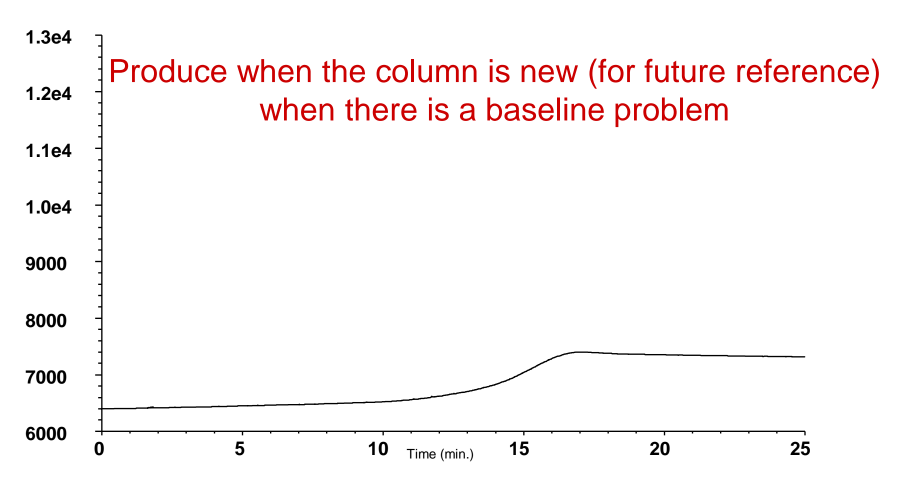
Test mix: *all problems*

Isolate the components: *all problems*

Condensation Test: *baseline problems* Jumper Tube Test: *baseline problems*

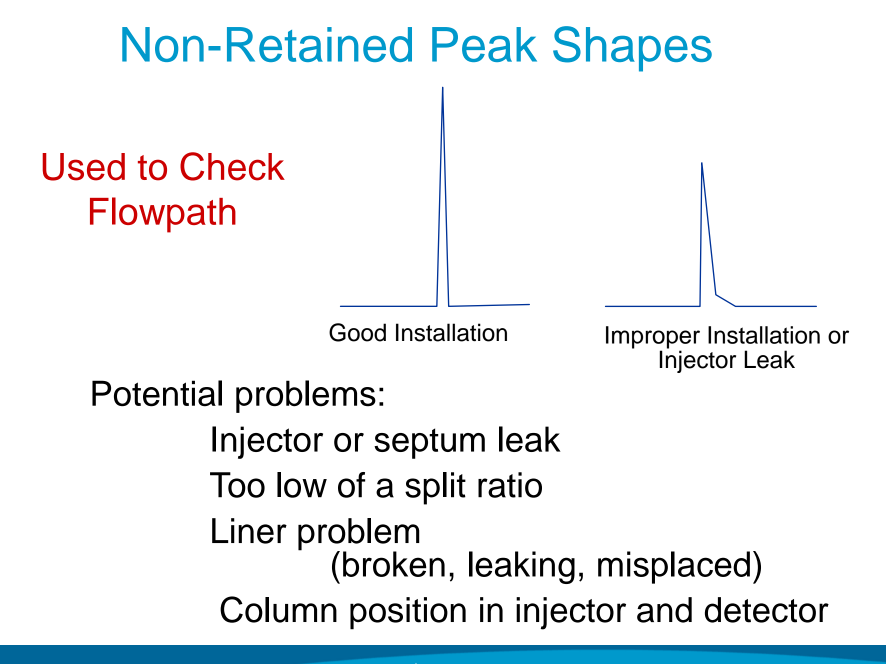


Generating a Bleed Profile



*DB-1 30m x .32mm I.D., .25μm Temperature program // 40°C, hold 1 min // 20°/min to 320°C, hold 10 min.

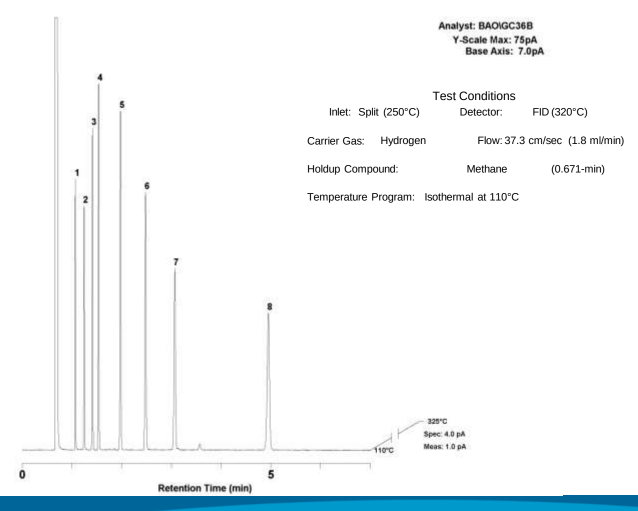








Used to determine how "good" the column is or if the problem is related to the chemical properties of the analytes.





Test Mixture Components

| <u>Compounds</u> | |
|------------------|--|
| Hydrocarbons | |
| | |
| Alcohols | |
| FAME's, PAH's | |
| Acids | |
| Bases | |

Purpose Efficiency

Retention

Activity

Retention

Acidic Character

Basic Character



Own Test Mixture

- More specific to your application
- Selective detectors
- Concentrations specific to your application
- Use same instrument conditions
- Easiest to simply inject a calibration standard
- Store for future measure of column performance



Isolate the Components

Simplify the system:

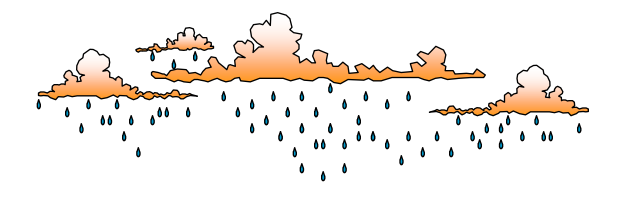
example - Direct injection instead of P&T sample introduction

Put in a known good column

Move column to a different GC, inlet or detector



Condensation Test



- Used* to isolate the cause of:
 - Erratic baselines
 - Ghost peaks or carryover

*Use when problems are worse after periods of GC non-use



Condensation Test

Procedure

Leave GC at 40-50°C for > 8 hours

Blank run

Repeat a blank run immediately after the first blank run is complete

Compare the two blank runs



Condensation Test

Results

First blank run is worse:

- Contaminants (from injector, lines, traps or carrier gas) carried into the column

Blank runs the same: contaminants are not strongly focused on the front of the column



Purpose

Helps to locate the source of contamination or noise

Isolates GC components



Isolate the Detector

Remove column from the detector

Cap detector and turn on

Blank run



Isolation of Detector - Results

Detector OK



Detector is the problem



Isolate the Injector

Connect the injector and detector

- 1-2 meters deactivated fused silica tubing

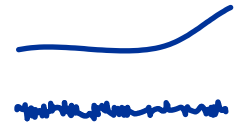
Turn on carrier gas

Blank run



Isolate the Injector - Results

Injector OK



Injector, lines or carrier gas contaminated



Isolate the Column

Reinstall the column

Setup as before

Blank Run



Isolate the Column - Results

Problem returns: It's the column

Problem gone: Previous leak, solid debris, or installation problem



And Now Let's do Some

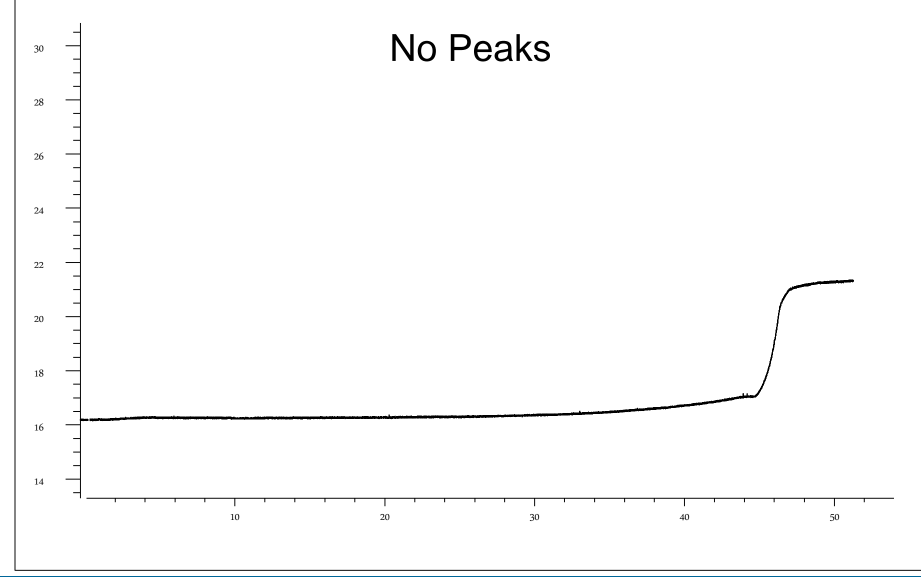




Troubleshooting-Example #1



A Real Troubleshooting Example





Logical Steps Taken to Find Peaks (most of our problems are leaks and plugs)

Is the flame Lit?

- → put glass piece over FID outlet----Answer in this case, Water condenses
- look at output in instrument guage-- is the digital value greater than 0.0?
 Answer in this case is approximately 16.2 pico amps

Is there flow through the column?

disconnect column from detector and measure flow with bubble solution or meter
 Answer in this case was YES THERE IS FLOW

Assess the observations

- Flame is lit and we have flow from end of column
- Hypothesis: Sample not getting on column-syringe plugged?

Take syringe out and make injection manually on a dry paper towel

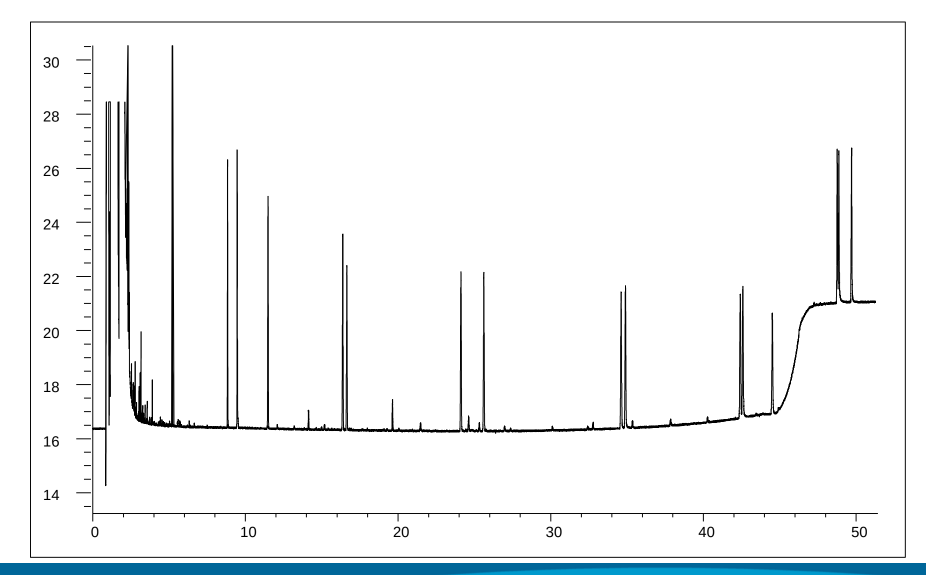
Answer – towel stayes dry (Syringe was clogged with septum)

Pull plunger out top, add solvent and replace plunger will usually dislodge septum particle (should hear a little pop) If you can't dislodge plug, Replace syringe

Reassemble the Injector & Re-inject



Peaks !!



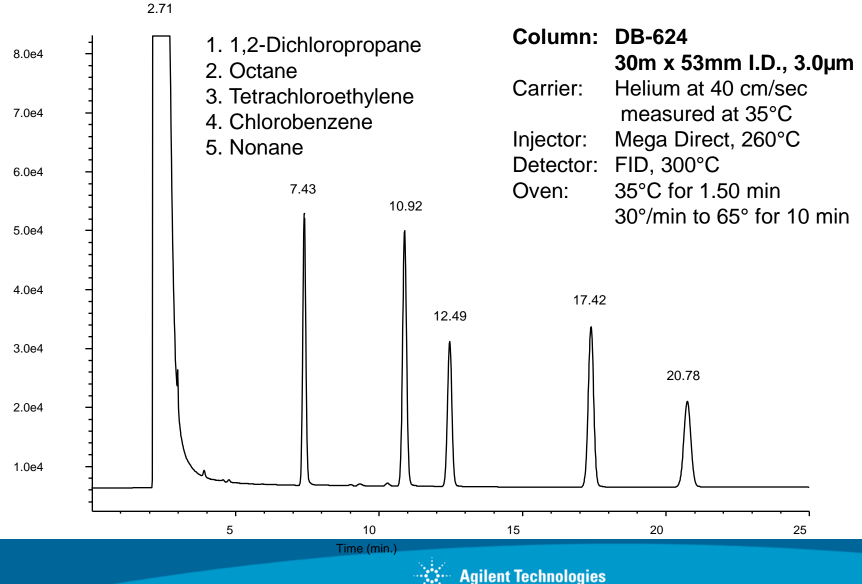


Troubleshooting-Example #2

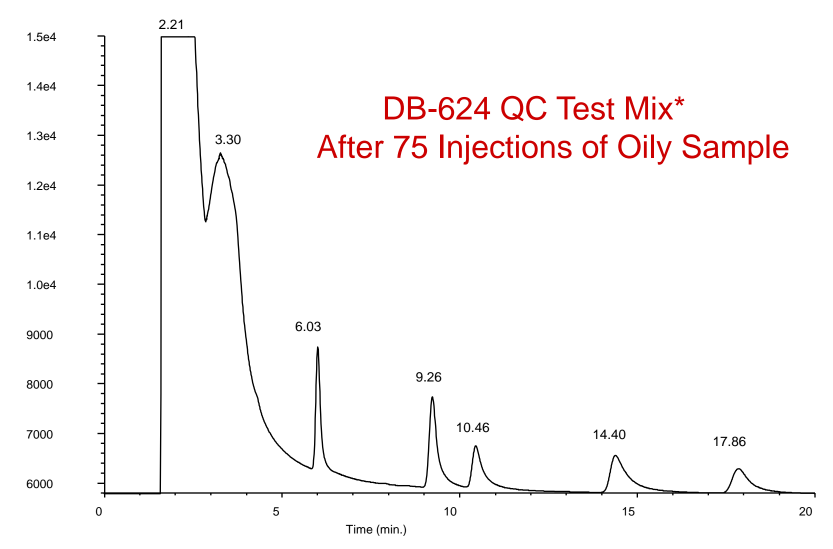


DB-624 COLUMN

QC Test Mix

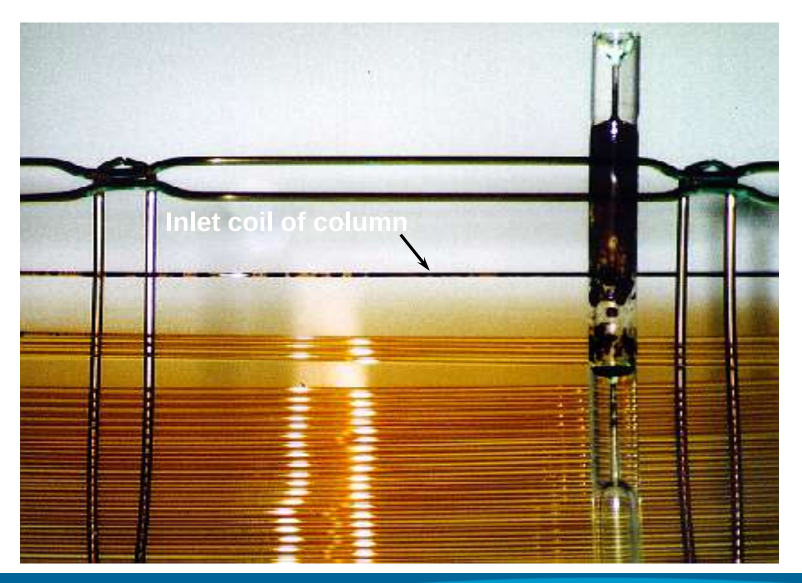


Example of Column Contamination



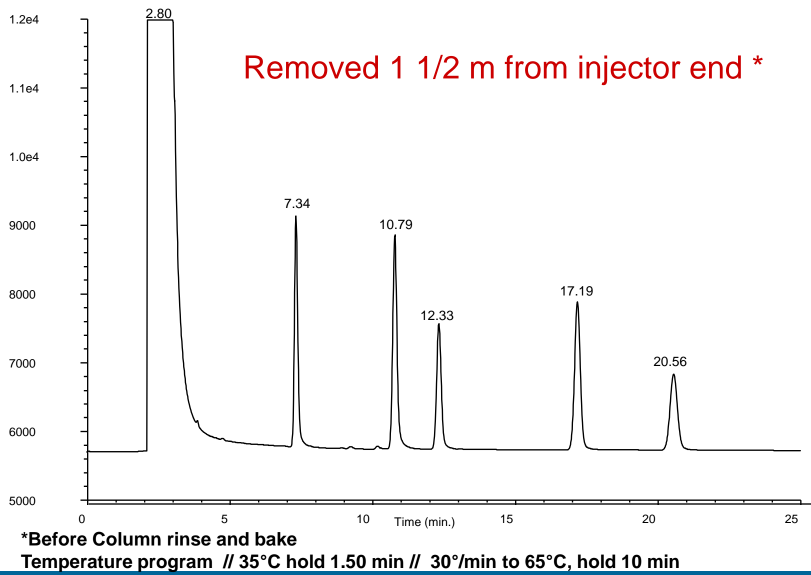


Column and Liner Contamination





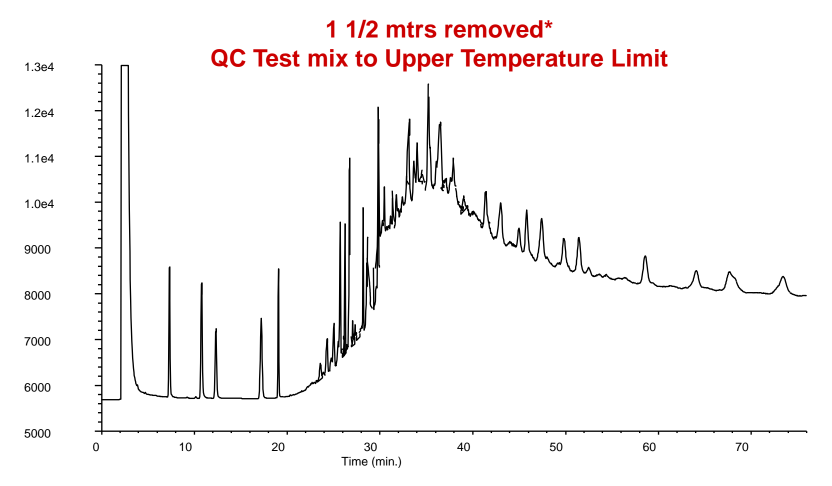
Example of Column Contamination



Looks Fixed Doesn't it?



Example of Column Contamination

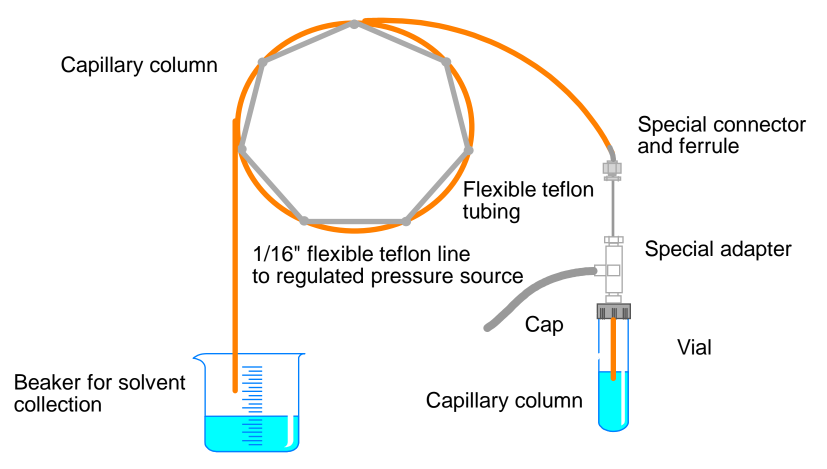


*Before Column rinse and bake. Temperature program // 35°C, hold 1.50 min // 30°/min to 65°C, hold 15 min // 20°/min to 260°, hold 50 min



Backflush Column

Rinse with 10ml each: Methanol, Methylene Chloride, Hexane



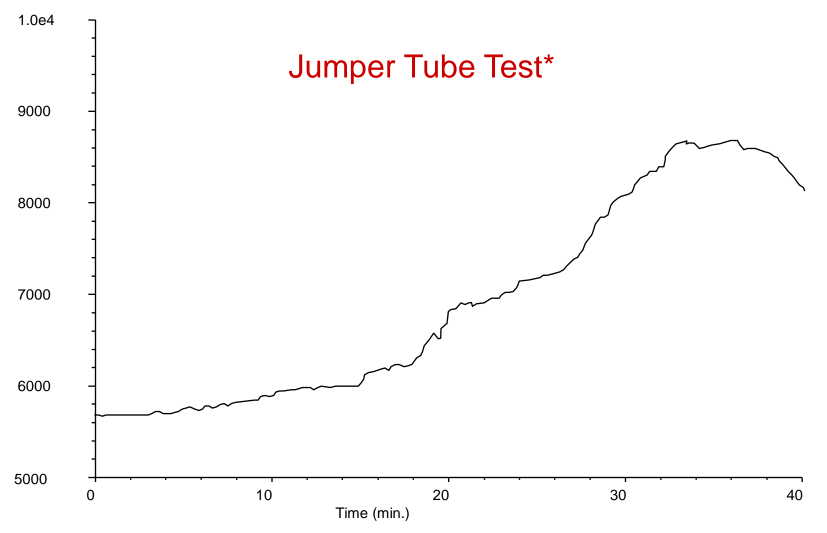


Used to Isolate Source of Contamination

- . Cap off the detector and establish normal gas flows and temperature.
- . Plot the baseline using a temperature program. If flat.....
- . Connect 1 meter of deactivated tubing between the injector and detector
- . Plot the baseline using a temperature program. If flat.....
- . Install the column.
- . Plot the baseline using a temperature program.

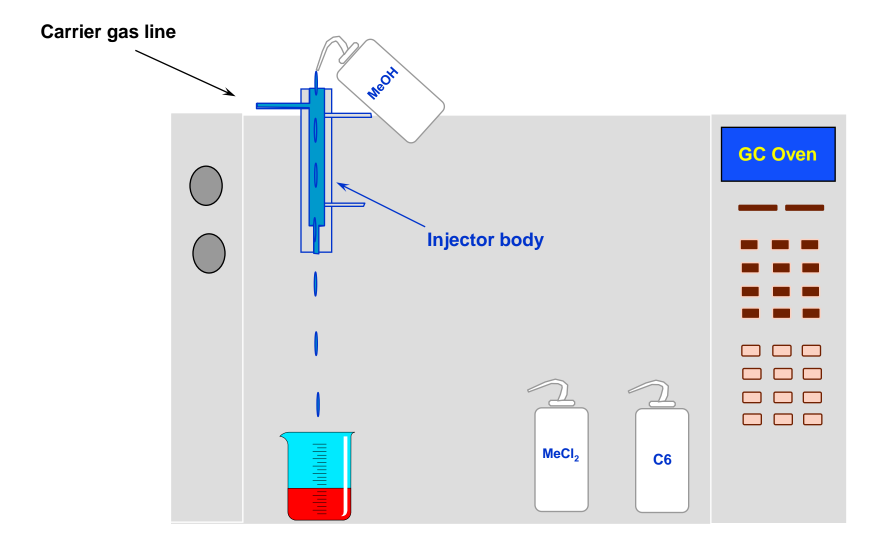


Contaminated Inlet





Rinsing Injector





Troubleshooting Tips

1. Isolate the problem.

(Blank Run, Inject Un-retained Compound, Jumper Tube Test)

2. Change only one variable at a time.

3. Compare before/after chromatograms.

(Peak shape, response, retention, baseline rise, background, look for trends, etc.)

4. Utilize Technical Support.





Complete system = Carrier Gas + Injector + Column + Detector + Data System

Multiple cause and effect

Do not change too many variables at once



TECHNICAL SUPPORT

1-800-227-9770, #3, #3, #1

E-mail: gc-column-support@agilent.com



