

# Installation, Care and Maintenance of Capillary Gas Chromatography Columns

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Application Engineer  
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# Installation, Care and Maintenance of Capillary Gas Chromatography Columns

**or....**

**"It's not what your column can do for you, but what you can do for your column"**



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# Column Installation

**"Getting off to a good start"**



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# Column Installation Procedure

- **Install the column**
- **Leak and installation check**
- **Column conditioning**
- **Setting linear velocity or flow rate**
- **Bleed profile**
- **Test mix**



# Column Installation

**What type of ferrule should I use?**

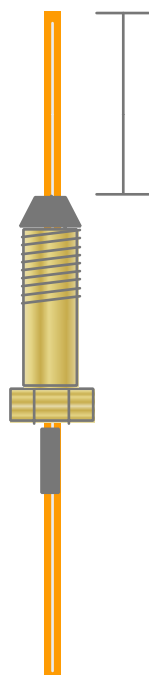
- **Graphite**
- **Graphite/Vespel**



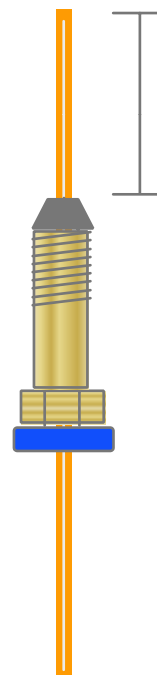
# Column Installation

## Measuring the right distance

**White out**



**Septa**



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# Cutting The Column

**Gently scribe through the polyimide coating.  
Do not attempt to cut the glass.**

**Recommended tools:**

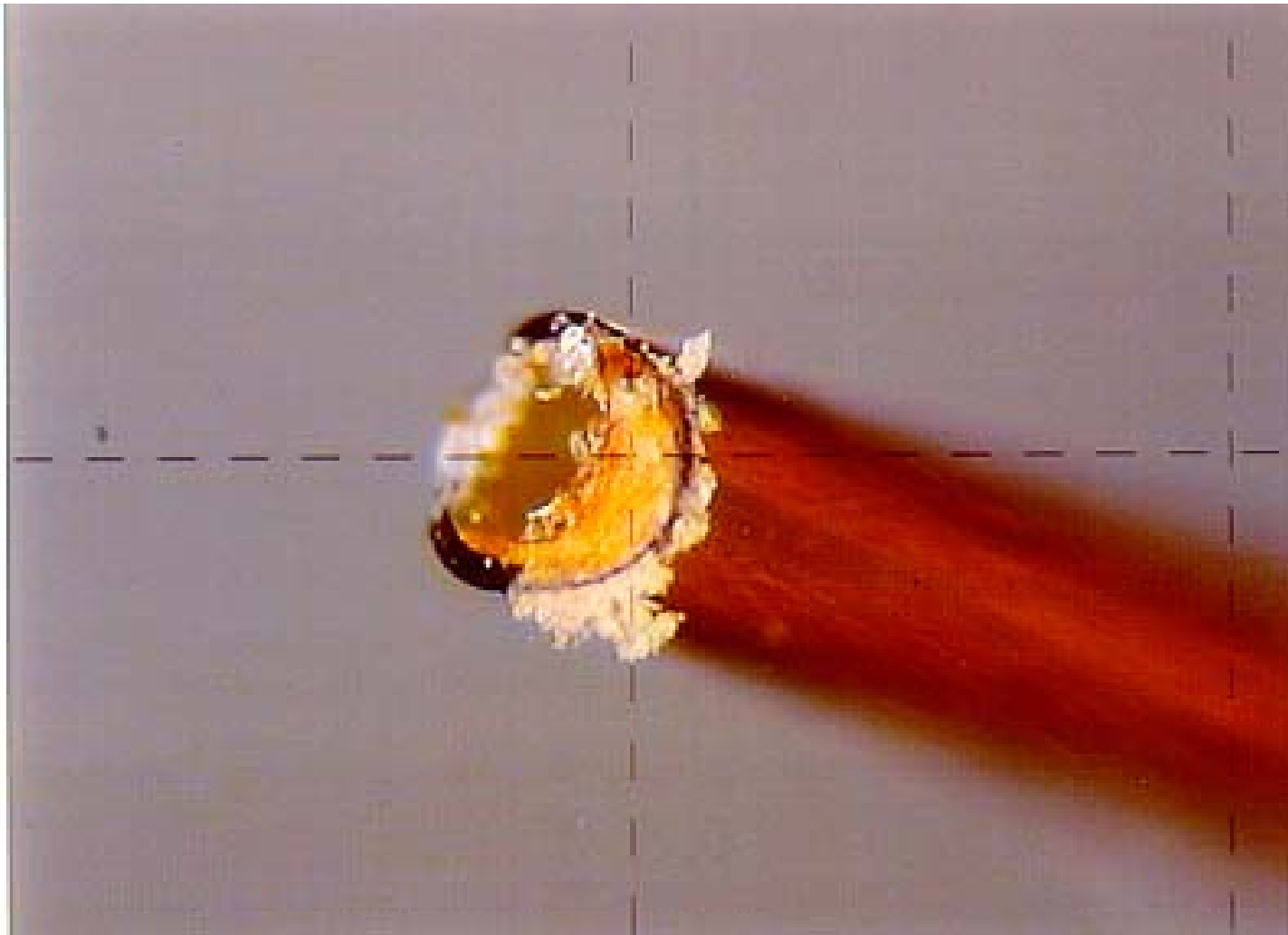
**Diamond or carbide tipped pencil; or sapphire  
cleaving tool, ceramic wafer  
Ocular**

**Do not use:**

**Scissors, file, etc.**



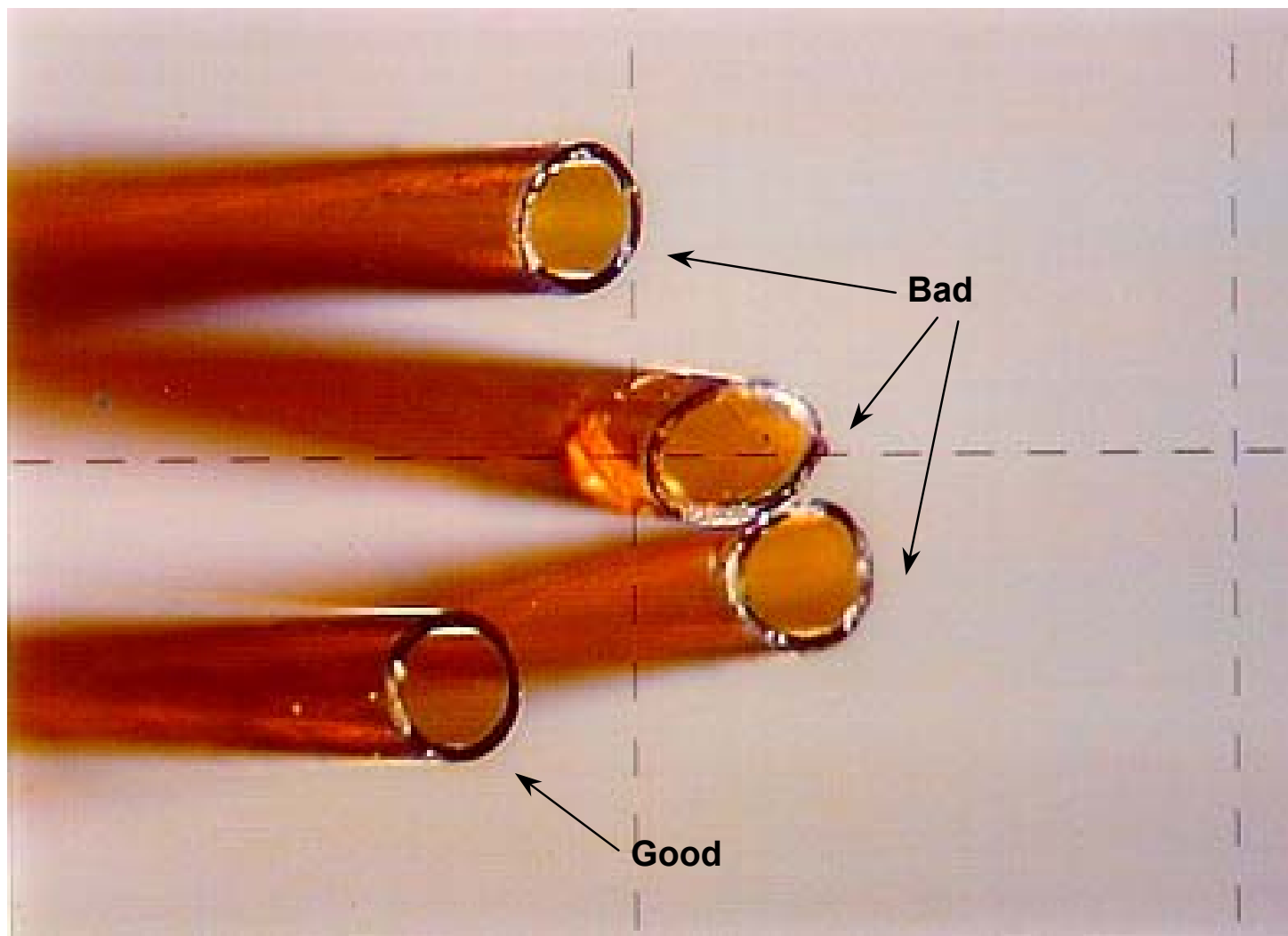
## Example of a Bad Cut



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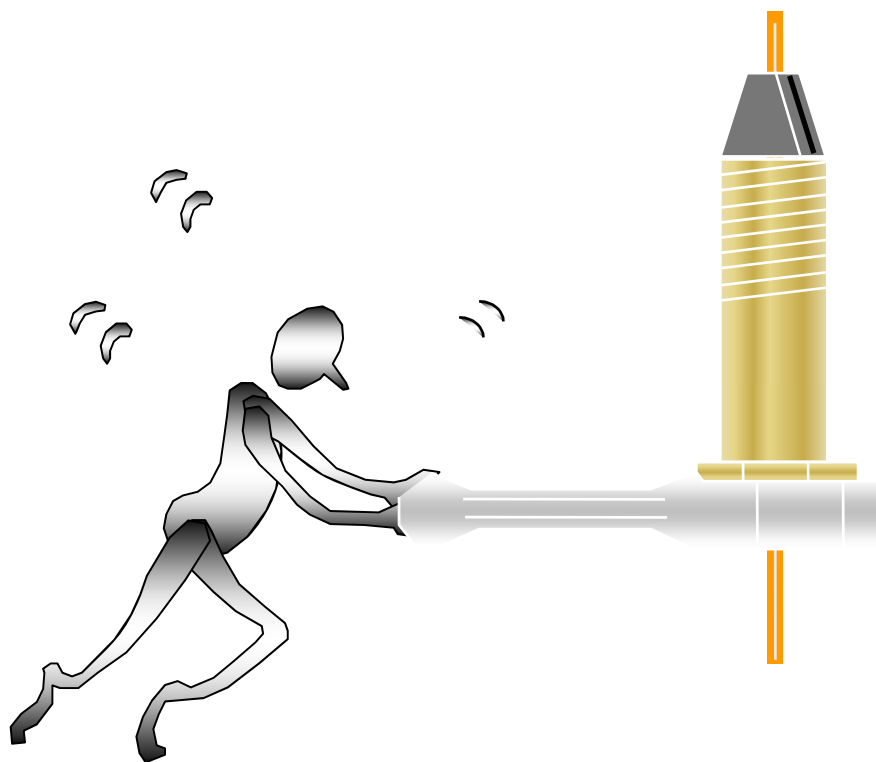


## Examples of Column Cuts



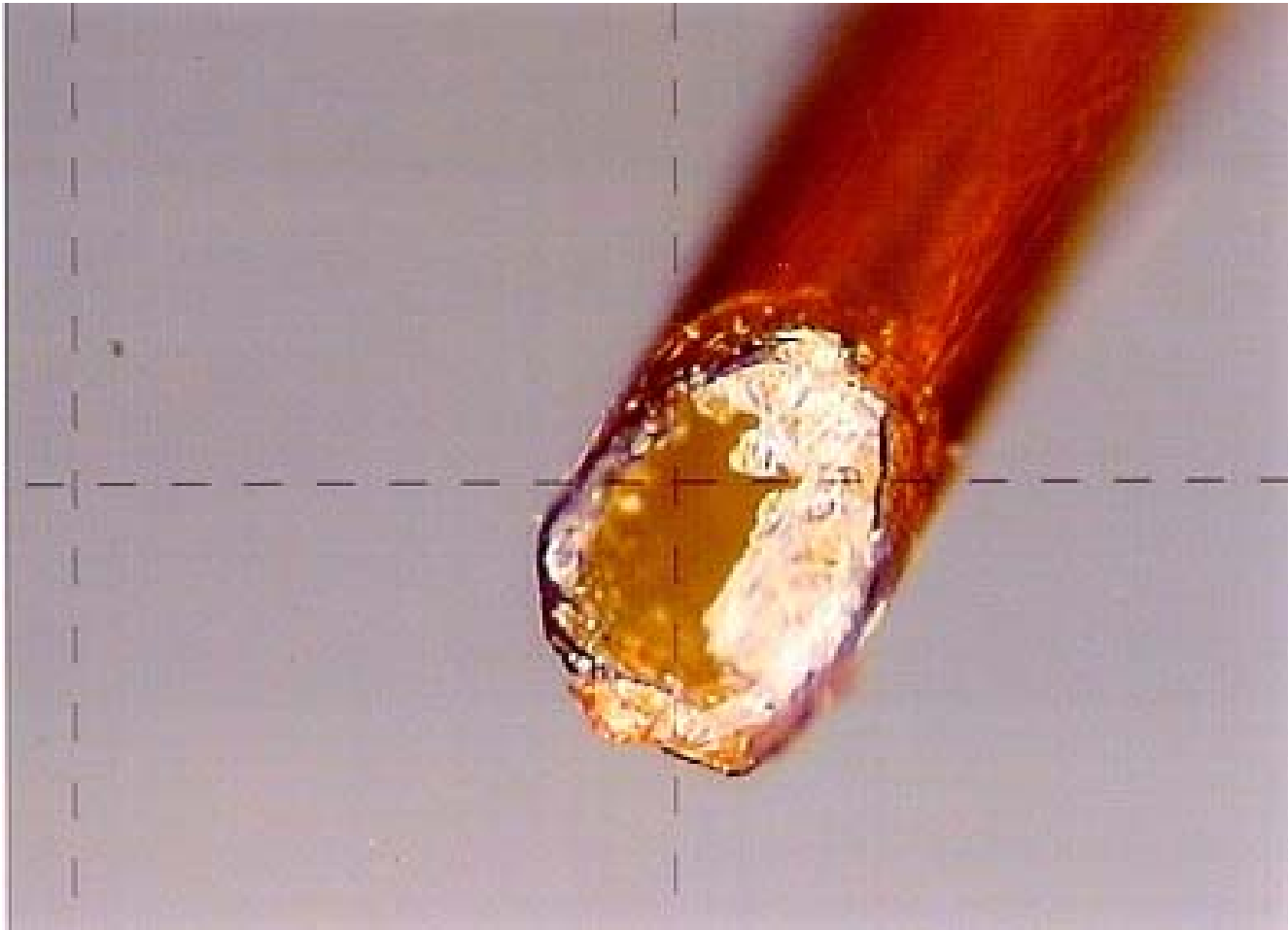
# Column Installation

**How tight is tight?**



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# Overtightened Ferrule



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# Column Installation

## Leak Check

**DO NOT USE SNOOP**

**Electronic leak detector**

**IPA/Water**

**Inject a non-retained peak**



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# Leak and Installation Check

Inject a non-retained compound vs DB-1

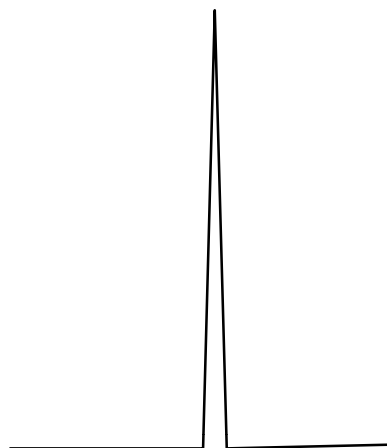
Detector	Compound
FID	Methane or Butane
ECD	$\text{MeCl}_2$ (headspace or diluted)
NPD	$\text{CH}_3\text{CN}$ -acetonitrile (headspace or diluted)
TCD	Air
MS	Air or Butane

The peak should be sharp and symmetrical

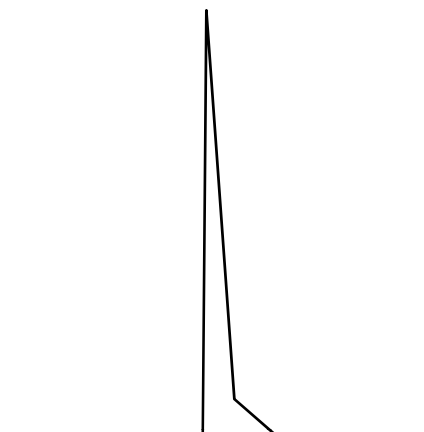


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## Non-Retained Peak Shapes



Good Installation



Improper Installation or  
Injector Leak

- Check for:**
- Too low of a split ratio**
  - Injector or septum leak**
  - Liner problem:**  
(broken, leaking, misplaced)
  - Column position in injector and detector**



## Calculating Linear Velocity

Inject a non-retained compound and obtain the retention time:

$$\bar{\mu} = \frac{L}{t_0}$$

$\bar{\mu}$  = Average linear velocity (cm/sec)

$L$  = Column length (cm)

$t_0$  = Retention time (sec)

$\bar{\mu}$  is *dependent* on column temperature

$\bar{\mu}$  is *independent* of column diameter

He 35-40 cm/sec

H<sub>2</sub> 45-60 cm/sec



## Calculating Flow Rate

**Inject a non-retained compound and obtain the retention time:**

$$\bar{F} = \frac{\pi r^2 L}{t_o}$$

$\bar{F}$  = Flow rate (mL/min)

$r$  = Column radius (cm)

$L$  = Column length (cm)

$t_o$  = Retention time (sec)

$\bar{F}$  is dependent on column temperature

**Measuring flow with a flow meter is often inaccurate**





# Column Conditioning

**System must be leak free before conditioning column**

**Heat the column to the lower of:**

**Isothermal maximum temperature OR**

**20° to 30°C above highest operation temperature**

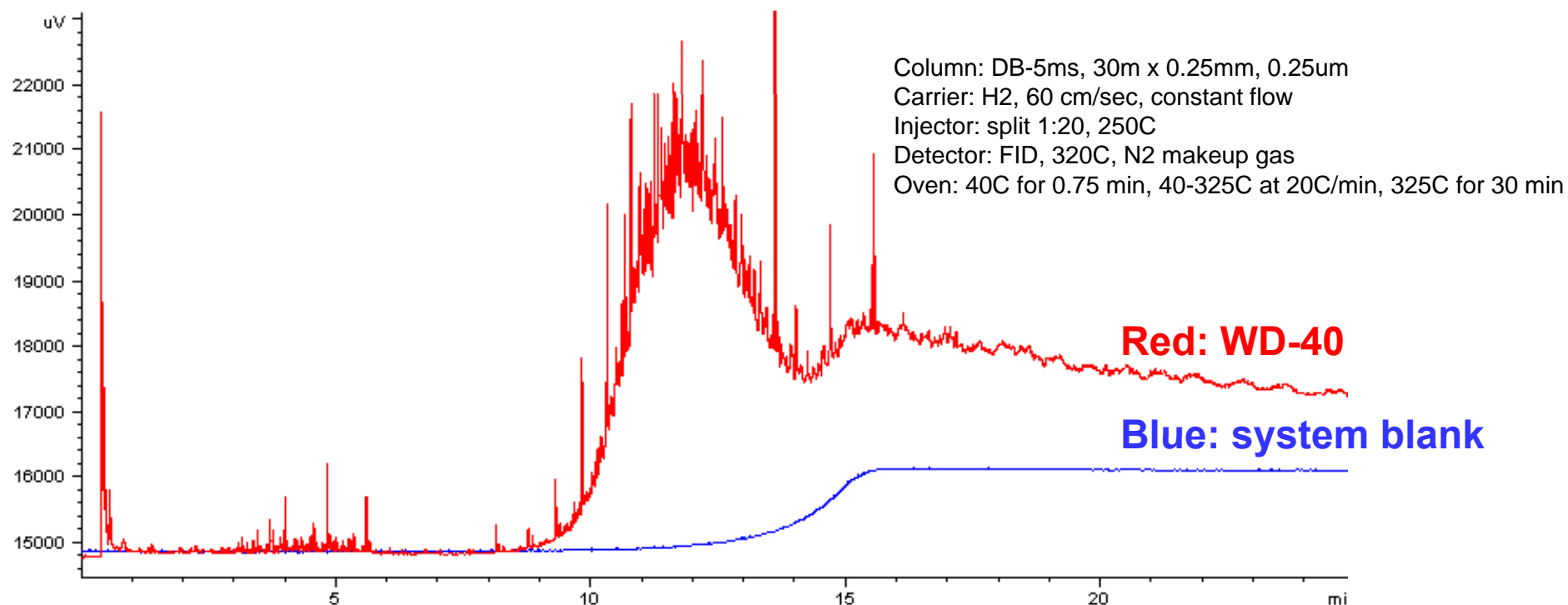
**Temperature programming is not necessary**

**Stop conditioning when the stable baseline is obtained:**

**1 to 2 hours in most cases**



# Contamination of System by Residue on Fingers During Column Installation



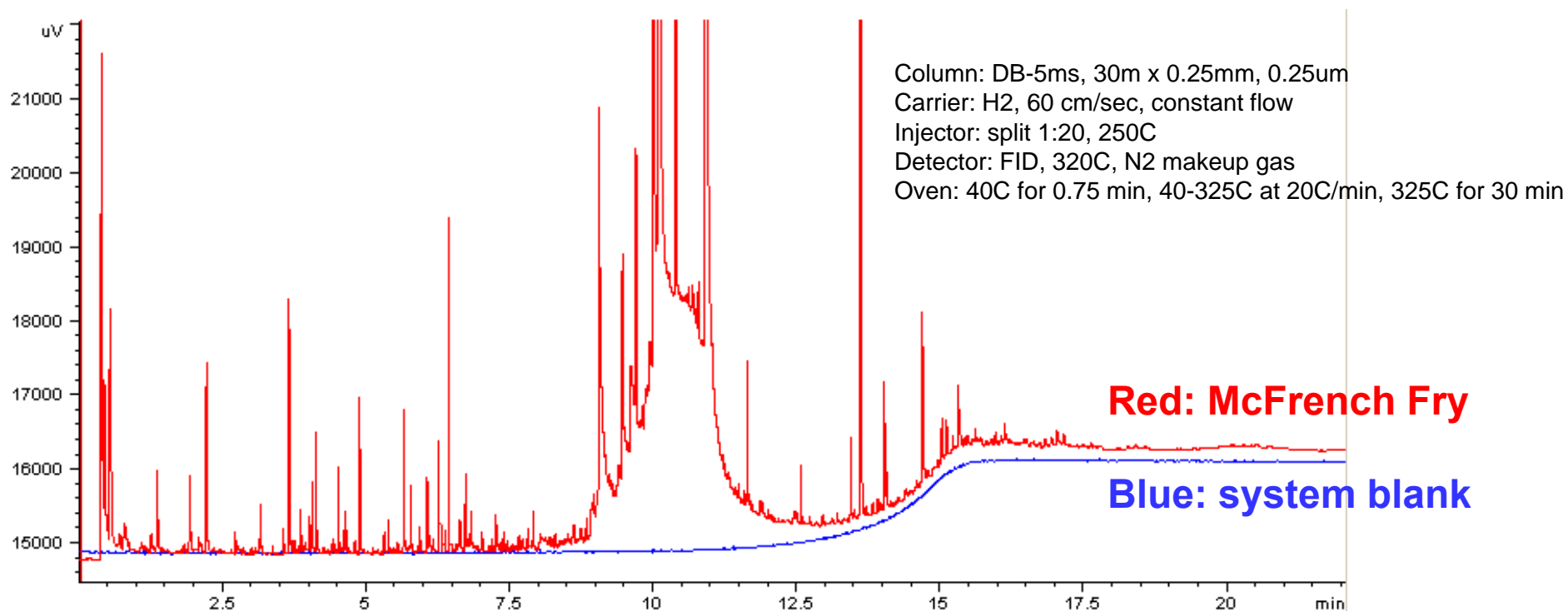
## Procedure:

- (1) One very small drop of liquid placed on one fingertip.
- (2) Fingertip was wiped with paper towel to remove as much of the offending material as possible.
- (3) Lightly touched the part of the column sticking up above the ferrule.
- (4) Installed column into injector.
- (5) Set oven temperature to 40C.
- (6) Started oven temperature program as soon as oven reached 40C.



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# Contamination from French Fry Grease



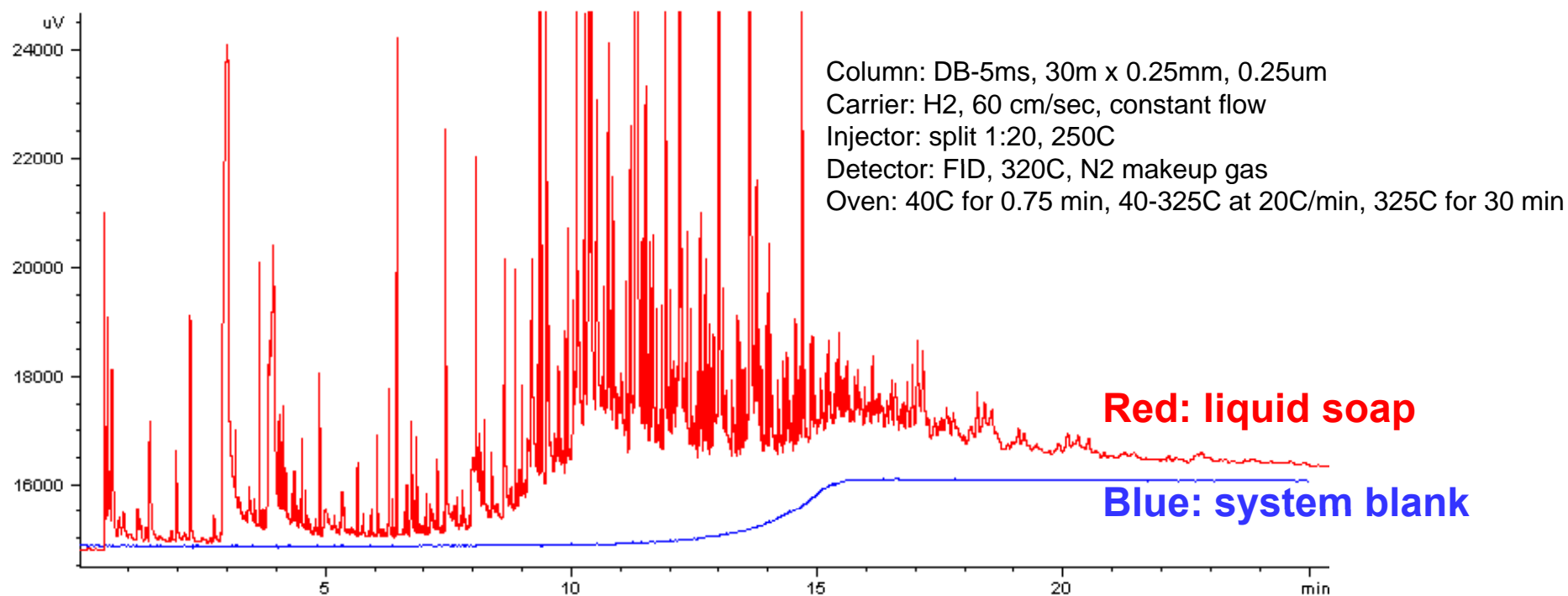
## Procedure:

- (1) Held french fry for 5 seconds.
- (2) Fingertip was wiped with paper towel to remove as much of the offending material as possible.
- (3) Lightly touched the part of the column sticking up above the ferrule.
- (4) Installed column into injector.
- (5) Set oven temperature to 40C.
- (6) Started oven temperature program as soon as oven reached 40C.



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# Contamination from Liquid Soap



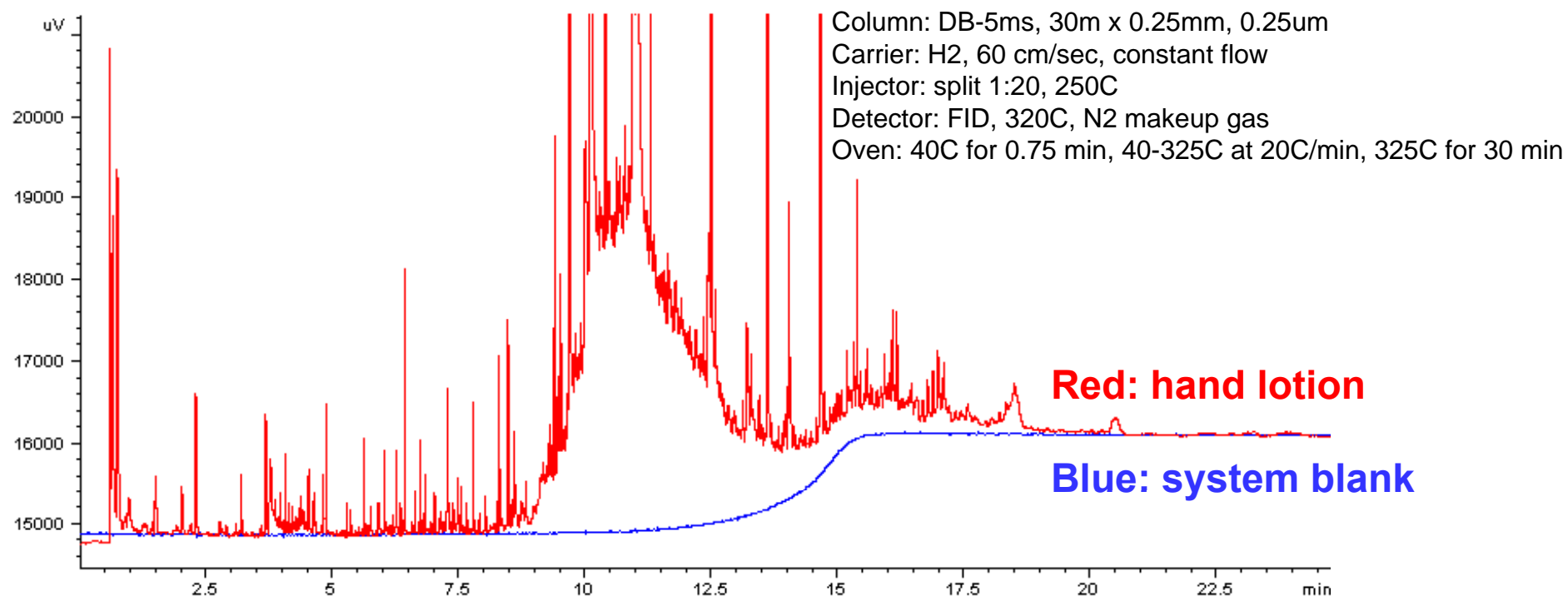
## Procedure:

- (1) One very small drop of liquid placed on one fingertip.
- (2) Fingertip was wiped with paper towel to remove as much of the offending material as possible.
- (3) Lightly touched the part of the column sticking up above the ferrule.
- (4) Installed column into injector.
- (5) Set oven temperature to 40C.
- (6) Started oven temperature program as soon as oven reached 40C.



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# Contamination from Hand Lotion



## Procedure:

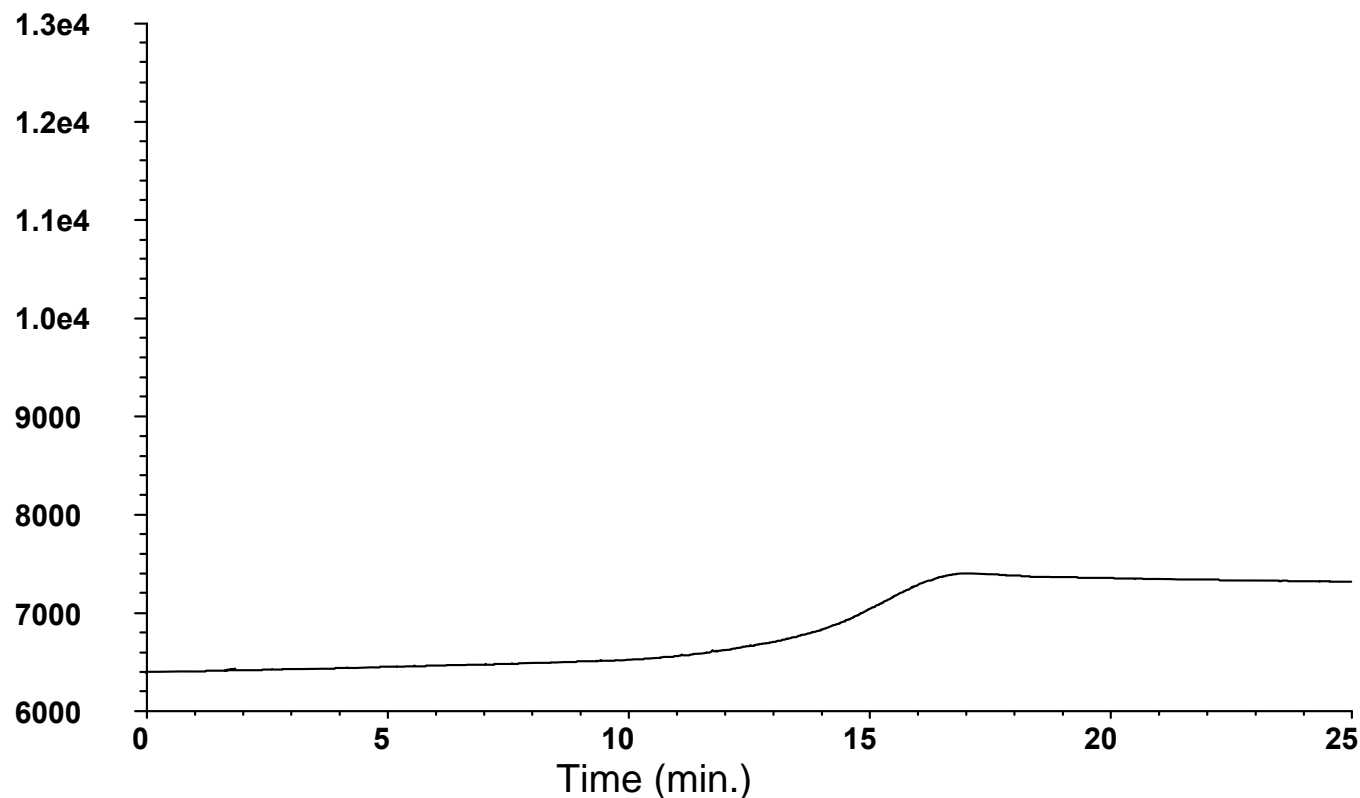
- (1) One very small drop of liquid placed on one fingertip.
- (2) Fingertip was wiped with paper towel to remove as much of the offending material as possible.
- (3) Lightly touched the part of the column sticking up above the ferrule.
- (4) Installed column into injector.
- (5) Set oven temperature to 40C.
- (6) Started oven temperature program as soon as oven reached 40C.



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# Generating a Bleed Profile

Temperature program the column without an injection\*



\*DB-1 30m x .32mm I.D., .25 $\mu$ m

Temperature program // 40°C, hold 1 min // 20°/min to 320°C, hold 10 min.



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# Test Mixes

Used to determine how "good" the column is



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# Column Performance Summary

PART NO: 1225032  
 COLUMN I.D. NO.: 3303121  
 LIQUID PHASE: DB-5  
 FILM THICKNESS: 0.25 µm  
 COLUMN DIMENSIONS:  
 30 m X 0.252 mm  
 TEMPERATURE LIMITS:  
 -60° C TO 325° C 350° C PROGRAM)

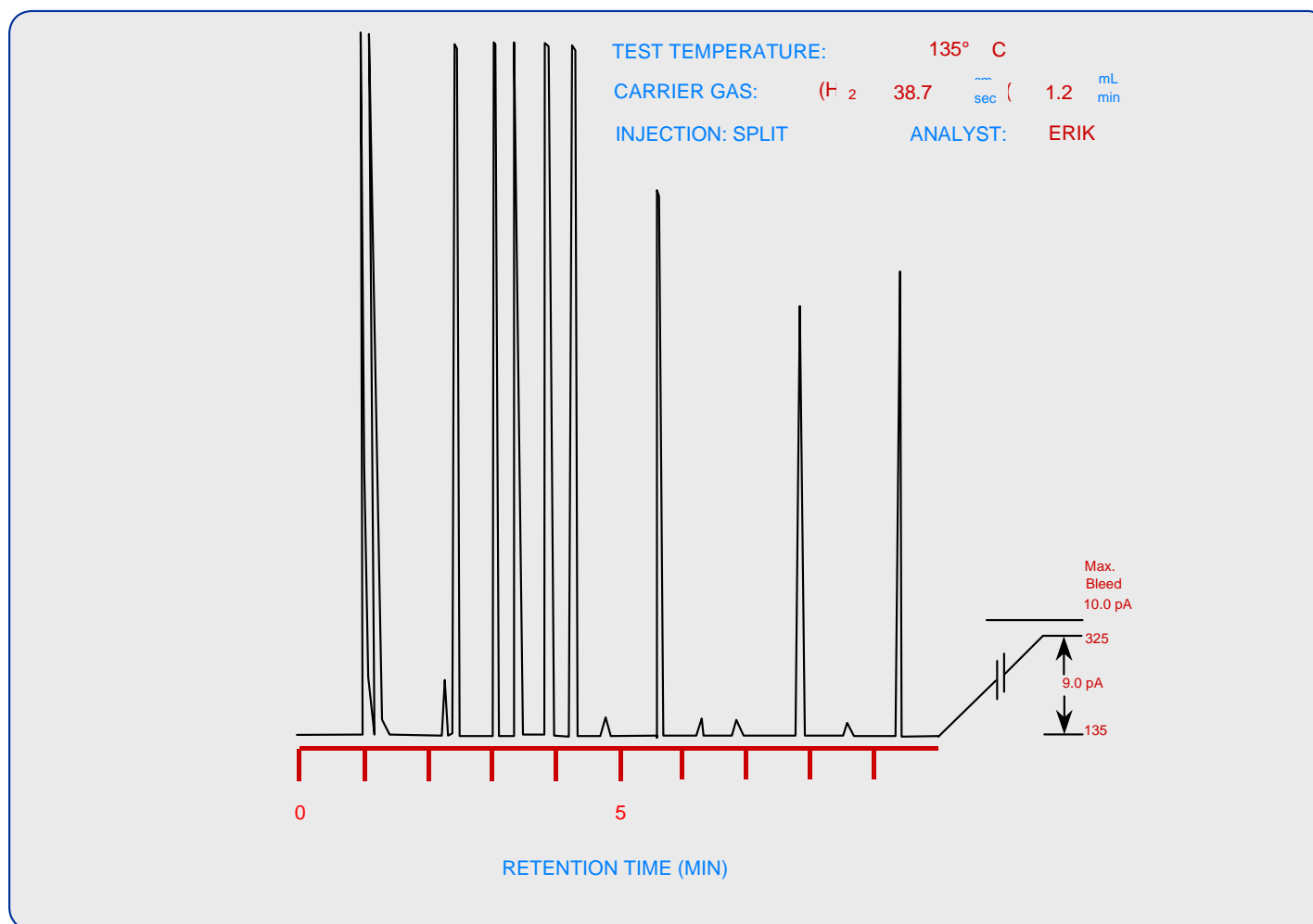
THEORETICAL PLATES/METER: MIN SPEC ACTUAL  
 PENTADECANE 3900 4389  
 COATING EFFICIENCY:  
 PENTADECANE 90.0 95.5  
 RETENTION INDEX: MIN SPEC MAX SPEC ACTUAL  
 1-UNDECANOL 1371.04 1372.04 1371.43  
 ACENAPHTHYLENE 1459.34 1460.34 1459.53  
 PEAK HEIGHT RATIO:  
 4-CHLOROPHENOL/  
 METHYL NONANOATE 0.83  
 4-PROPYLANILINE/  
 METHYL NONANOATE 1.14

COMPOUND IDENTIFICATION	RETENTION TIME (R)	PARTITION RATIO (k)	PEAK WIDTH (W 1/2)
1,6-HEXANEDIOL	2.51	0.9	0.019
4-CHLOROPHENOL	2.95	1.3	0.022
METHYL NONANOATE	3.21	1.5	0.022
4-PROPYLANILINE	3.81	1.9	0.026
TRIDECANE	4.20	2.2	0.027
1-UNDECANOL	5.52	3.3	0.036
ACENAPHTHYLENE	8.00	5.2	0.053
PENTADECANE	9.58	6.4	0.062
Approximately 5-10 ng on column			
o	1.29		





# Chromatographic Performance



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# Test Mixture Components

## Compounds

Hydrocarbons

FAME's, PAH's

Alcohols

Acids

Bases

## Purpose

Efficiency

Retention

Retention

Activity

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**



# An Ounce of Prevention.....



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# Common Causes of Column Performance Degradation

- **Physical damage to the polyimide coating**
- **Thermal damage**
- **Oxidation (O<sub>2</sub> damage)**
- **Chemical damage by samples**
- **Contamination**

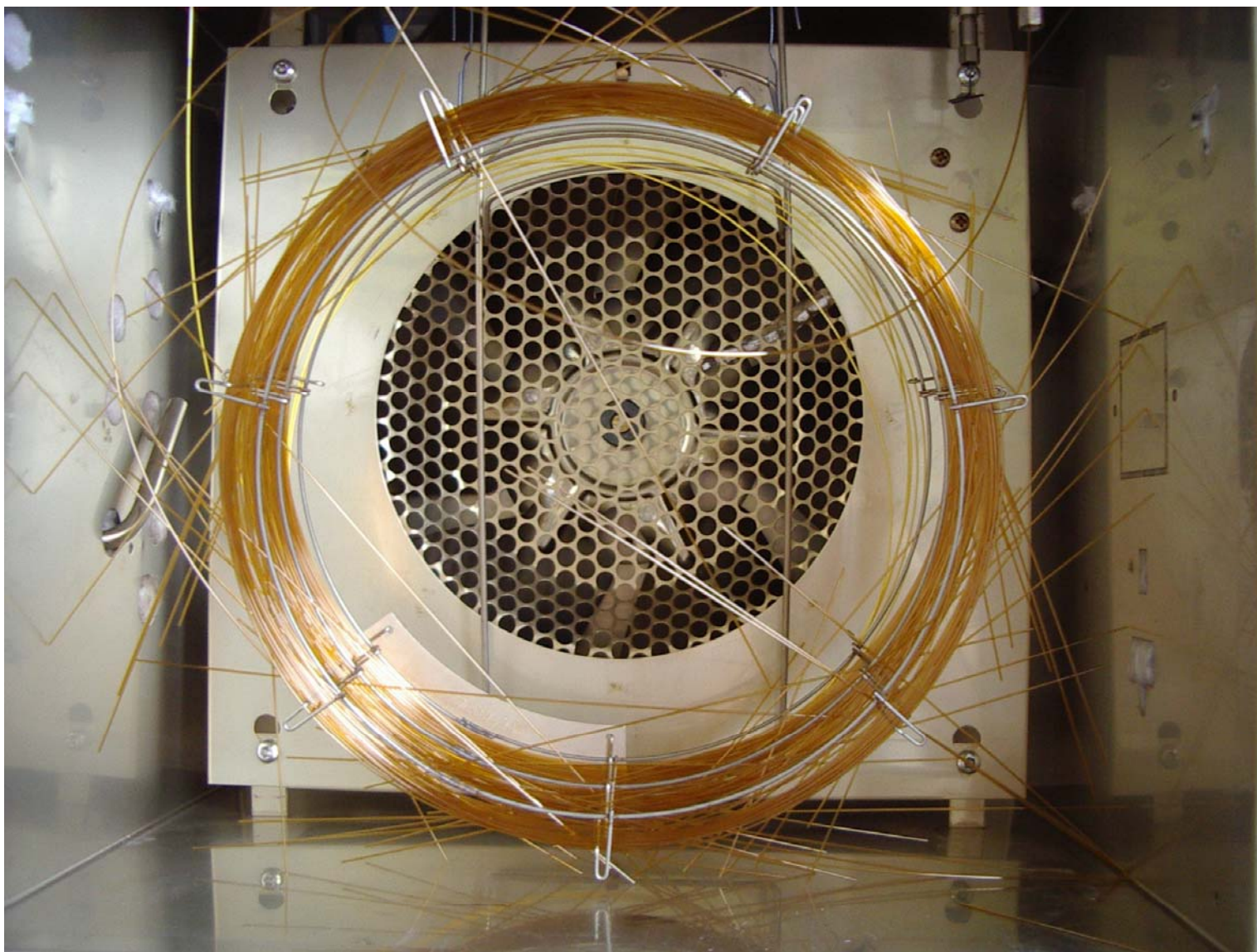


# Physical Damage to The Polyimide Coating

- **Smaller diameter tubing is more flexible than larger diameter tubing.**
- **Avoid scratches and abrasions**
- **Immediate breakage does not always occur upon physical damage**



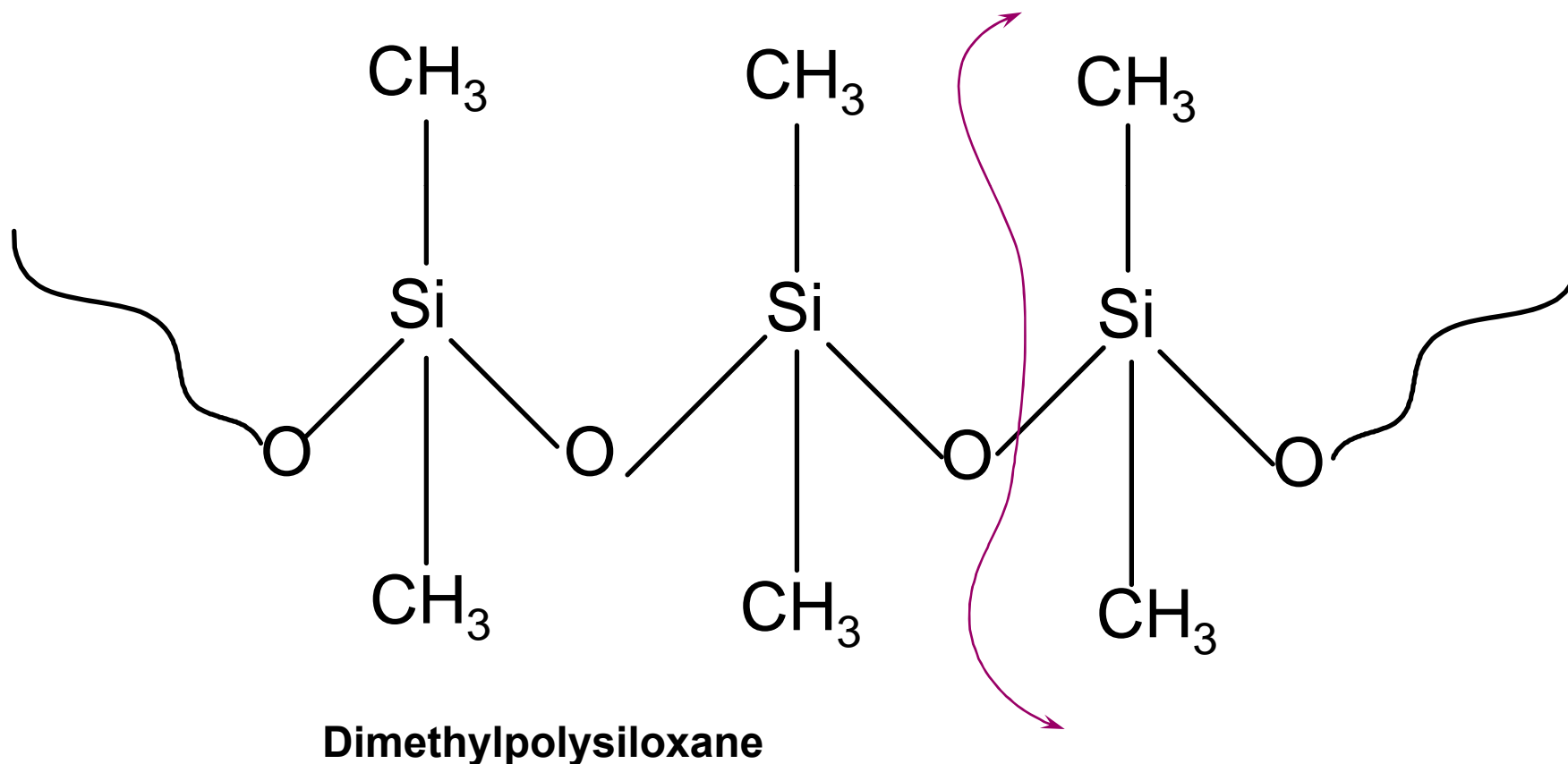
**NOT what you want your column to look like!**



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# Thermal Damage

Degradation of the stationary phase is increased at higher temperatures. Breakage along the polymer backbone.





# Thermal Damage

## What To Do If It Happens

- **Disconnect column from detector**
- **“Bake out” overnight at isothermal limit**
- **Remove 10-15 cm from column end**



# Thermal Damage

- **Rapid degradation of the stationary phase caused by excessively high temperatures**

**Isothermal limit = Indefinite time**

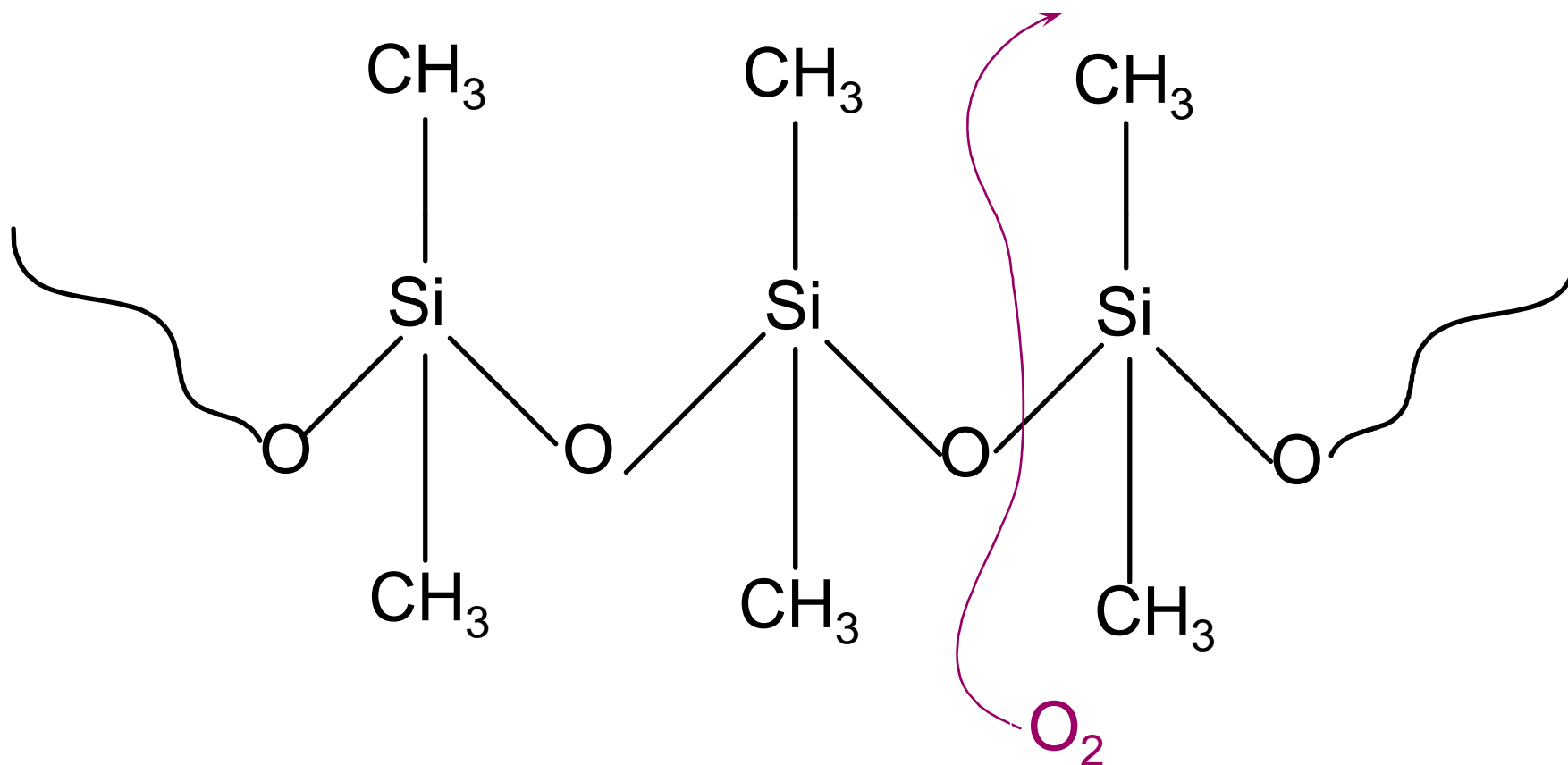
**Programmed limit = 5-10 minutes**

- **Temporary "column failure" below lower temperature limit**



## Oxidation (O<sub>2</sub> Damage)

Oxygen in the carrier gas rapidly degrades the stationary phase. The damage is accelerated at higher temperatures. Damage along the polymer backbone is irreversible.



Dimethylpolysiloxane



# Oxygen Damage

- **Causes rapid damage to the column**
- **Usually results in irreversible column damage**

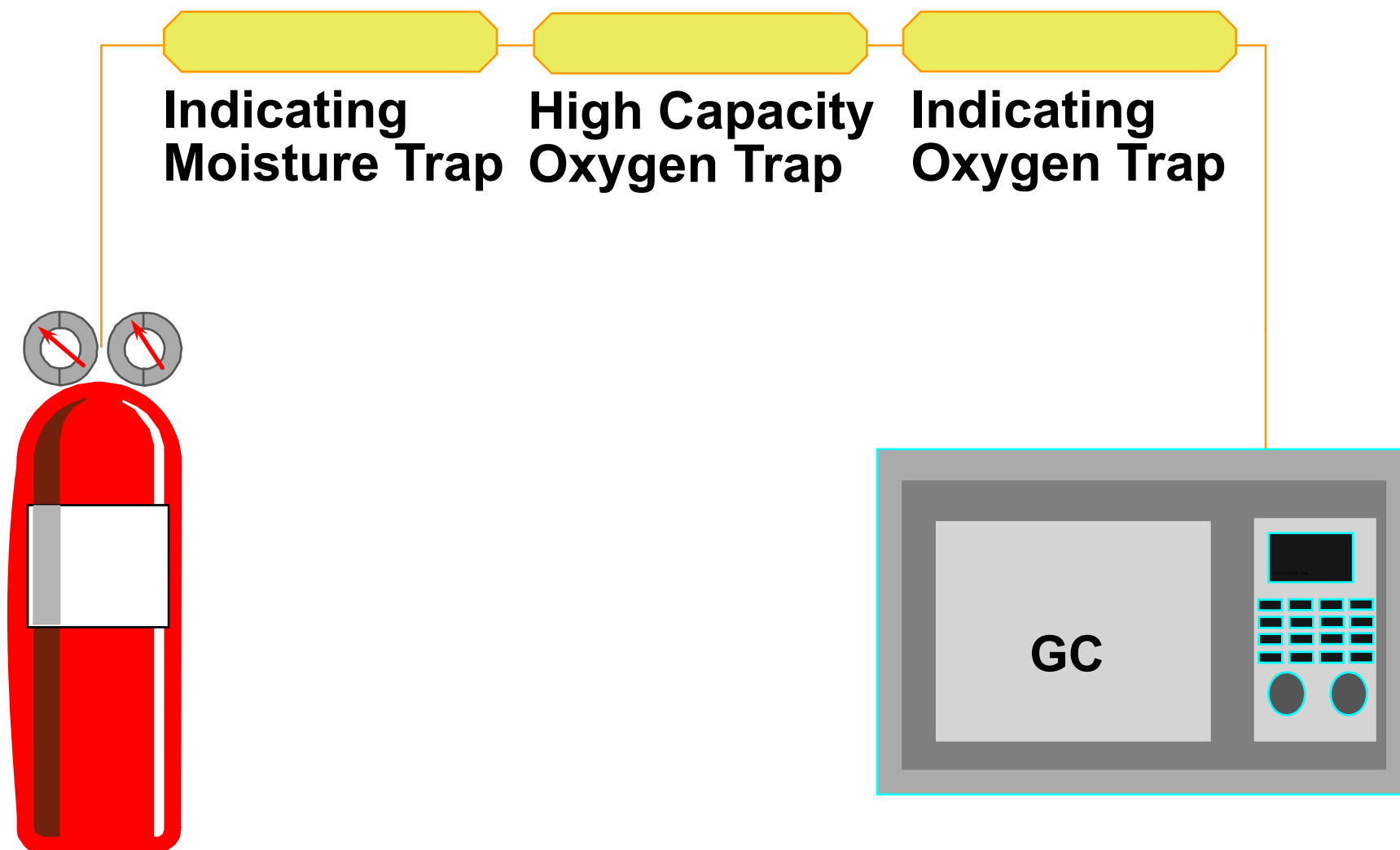


# How to Prevent Column Damage by Oxygen

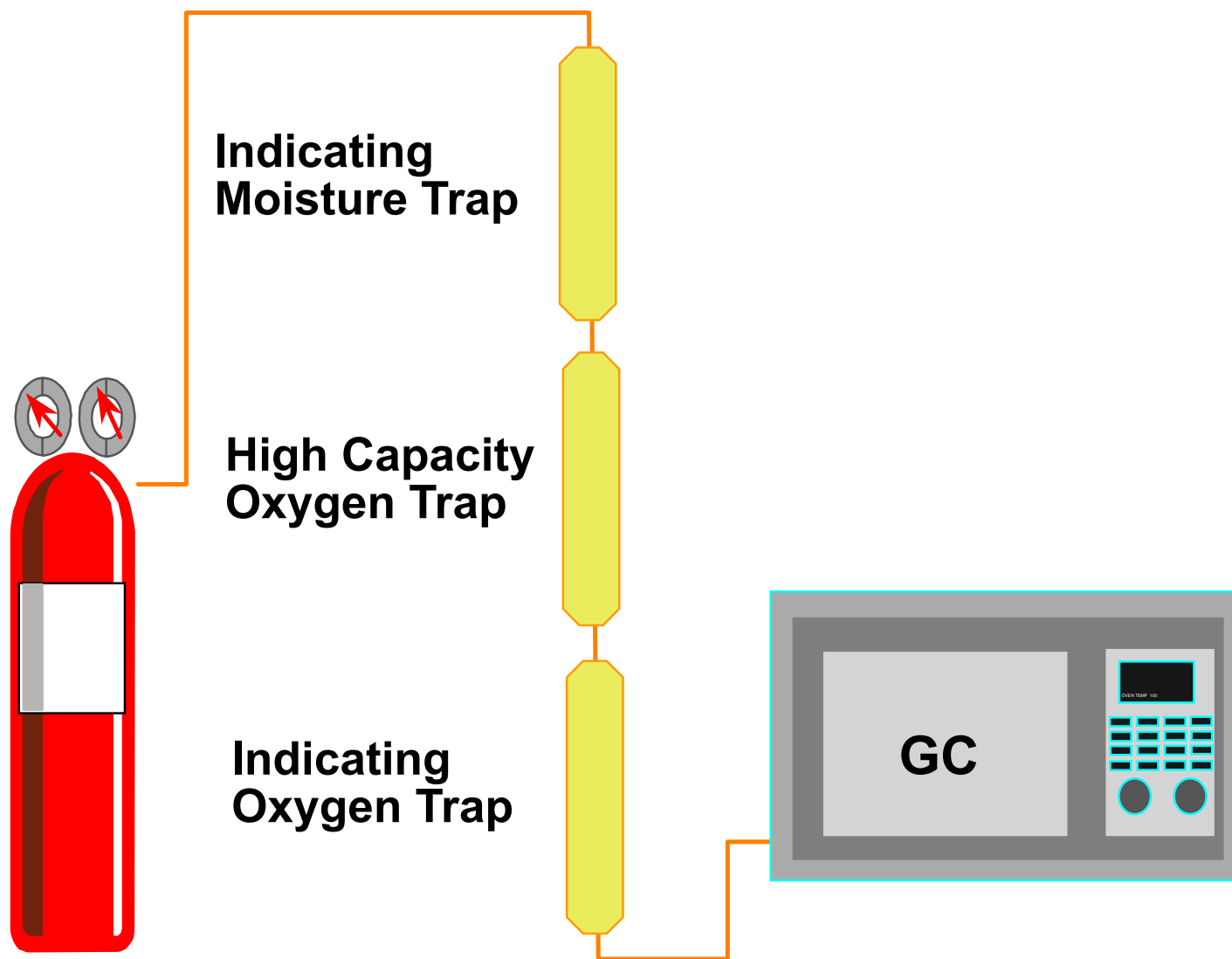
- **High quality carrier gas (4 nine's or greater)**
- **Leak free injector and carrier lines**
  - Change septa**
  - Maintain gas regulator fittings**
- **Appropriate impurity traps**



# Configurations for Carrier Gas Purifiers



# Configurations for Carrier Gas Purifiers



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# Chemical Damage

**Bonded and cross-linked columns have excellent chemical resistance except for inorganic acids and bases**



**Chemical damage will be evident by excessive bleed, lack of inertness or loss of resolution/retention**





# Chemical Damage

## What To Do If It Happens

- Remove 1/2 - 1 meter from the front of the columns
- Severe cases may require removal of up to 5 meters

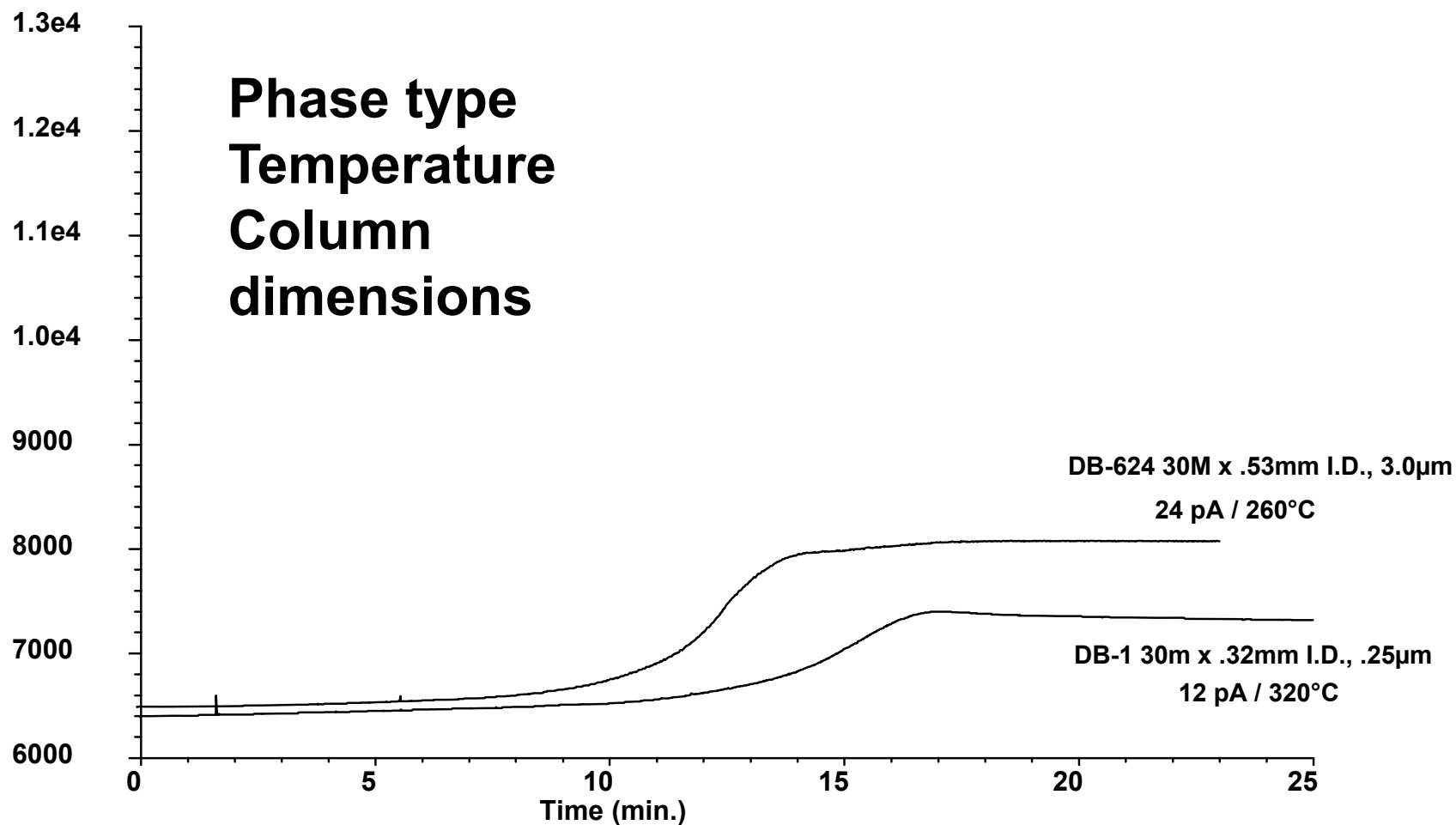


# What is Normal Column Bleed

Normal background signal generated by the elution of normal degradation products of the column stationary phase

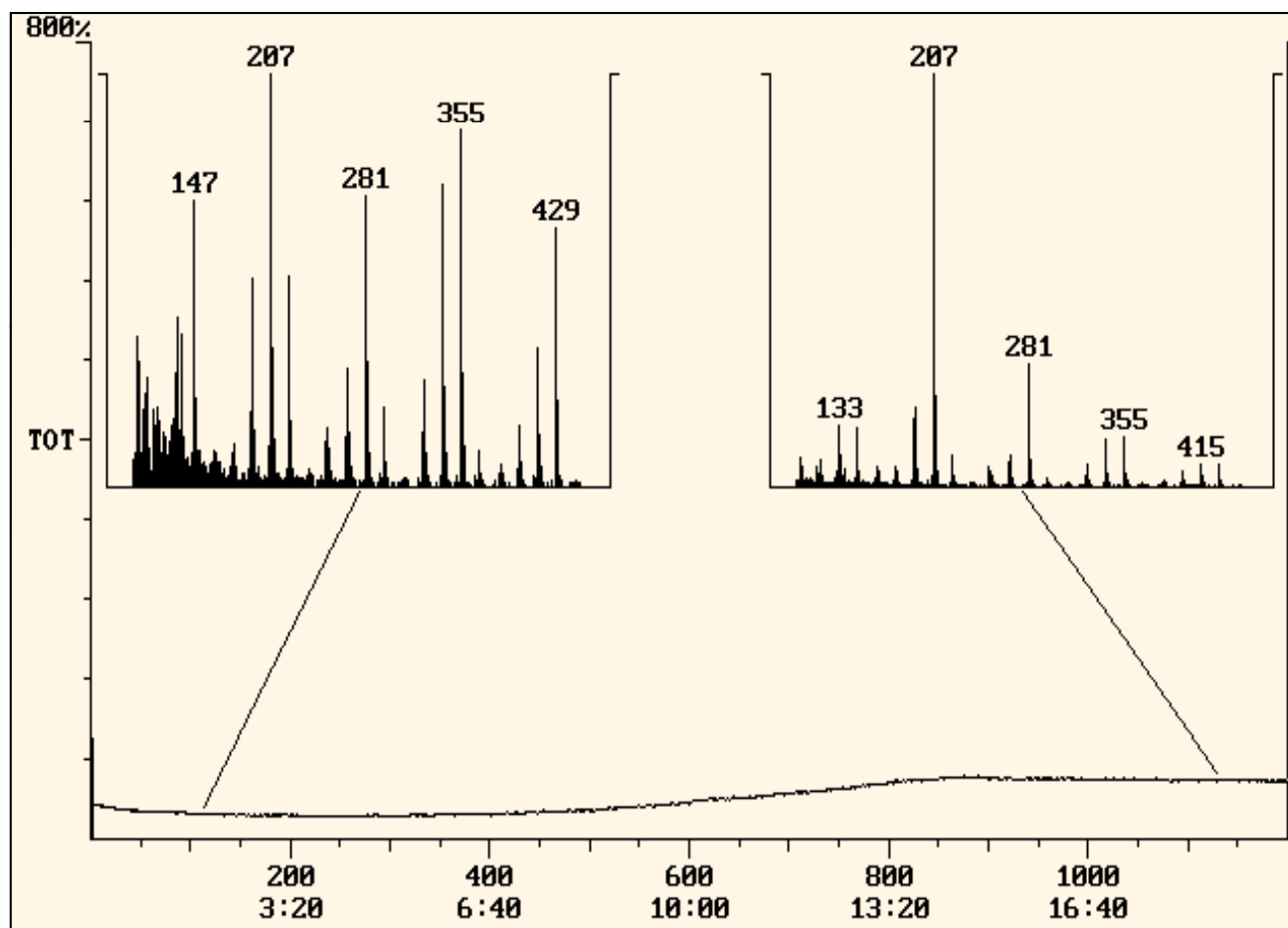


# Column Bleed is Influenced by:



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# Mass Spectrum of Phenylmethylpolysiloxane Column Bleed (Normal Background)



Mass spectral library search is not always accurate



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## What is a Bleed Problem?

### IT IS:

An abnormal elevated baseline at high temperature

### IT IS NOT:

A high baseline at low temperature

Wandering or drifting baseline at any temperature

Discrete peaks



# Column Contamination

- **Fouling of GC and column by contaminants**
- **Mimics nearly every chromatographic problems**



# Symptoms of Contamination

- **Poor peak shape**
- **Loss of separation (resolution)**
- **Changes in retention**
- **Reduced peak size**
- **Baseline disturbances (semi-volatiles only)**



# Typical Samples That Contain a Large Amount of Residues

**Biological (Blood, Urine, Tissue, Plants)**

**Soils**

**Foods**

**Waste Water**

**Sludges**

**All samples contain residues!! (even standards!)**



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# Other Sources of Contamination

- **Septum and ferrule particles**
- **Gas and trap impurities**
- **Unknown sources (vials, syringes, etc.)**



## Non-Volatile Residues

Any portion of the sample that does not elute from the column or remains in the injector.

## Semi-Volatile Residues

Any portion of the sample that elutes from the column after the current chromatographic run.

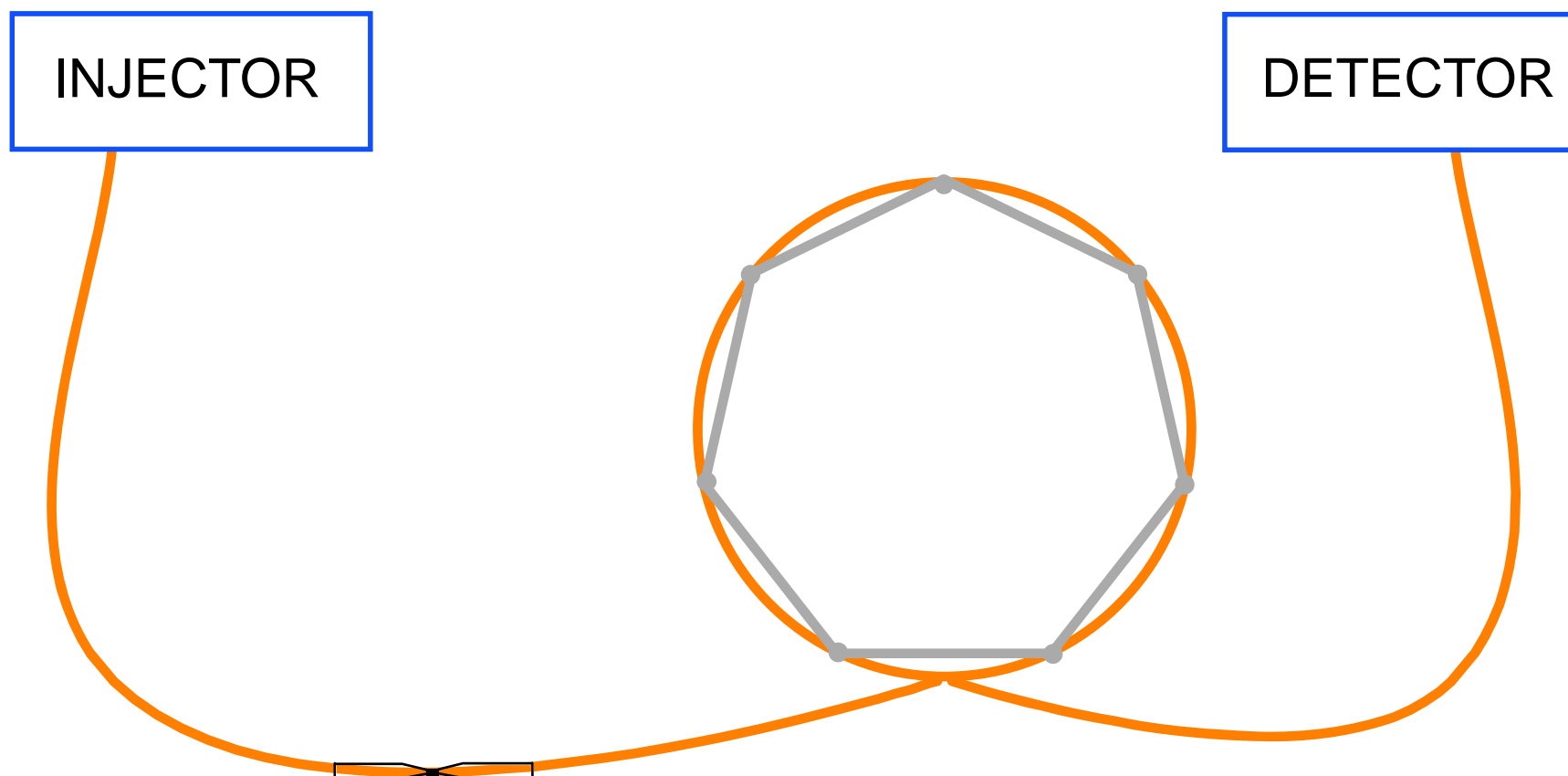


# Methods to Minimize Non-Volatile Residue Problems

- **Sample cleanup**
- **Packed injection port liners**
- **Guard columns**



# Guard Column or Retention Gap



**The guard column is 3 - 5 meters of deactivated fused silica tubing with the same diameter as the analytical column. It is connected with a zero dead volume union.**



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# Non-Volatile Contamination

## What To Do If It Happens

- Do not “bake out” the column
- Front End Maintenance
  - clean or change the injector liner
  - clean the injector
  - cut off 1/2 -1 meter of the front of the column
- Turn the column around
- Solvent rinse the column
- Cut the column in half



# Rinse Kit

> **Column must be bonded & cross linked!**

> Remove 15 – 30 cm from injector end

> Solvent flow should be from detector end to injector end

> Solvent order:

-from polar to non-polar

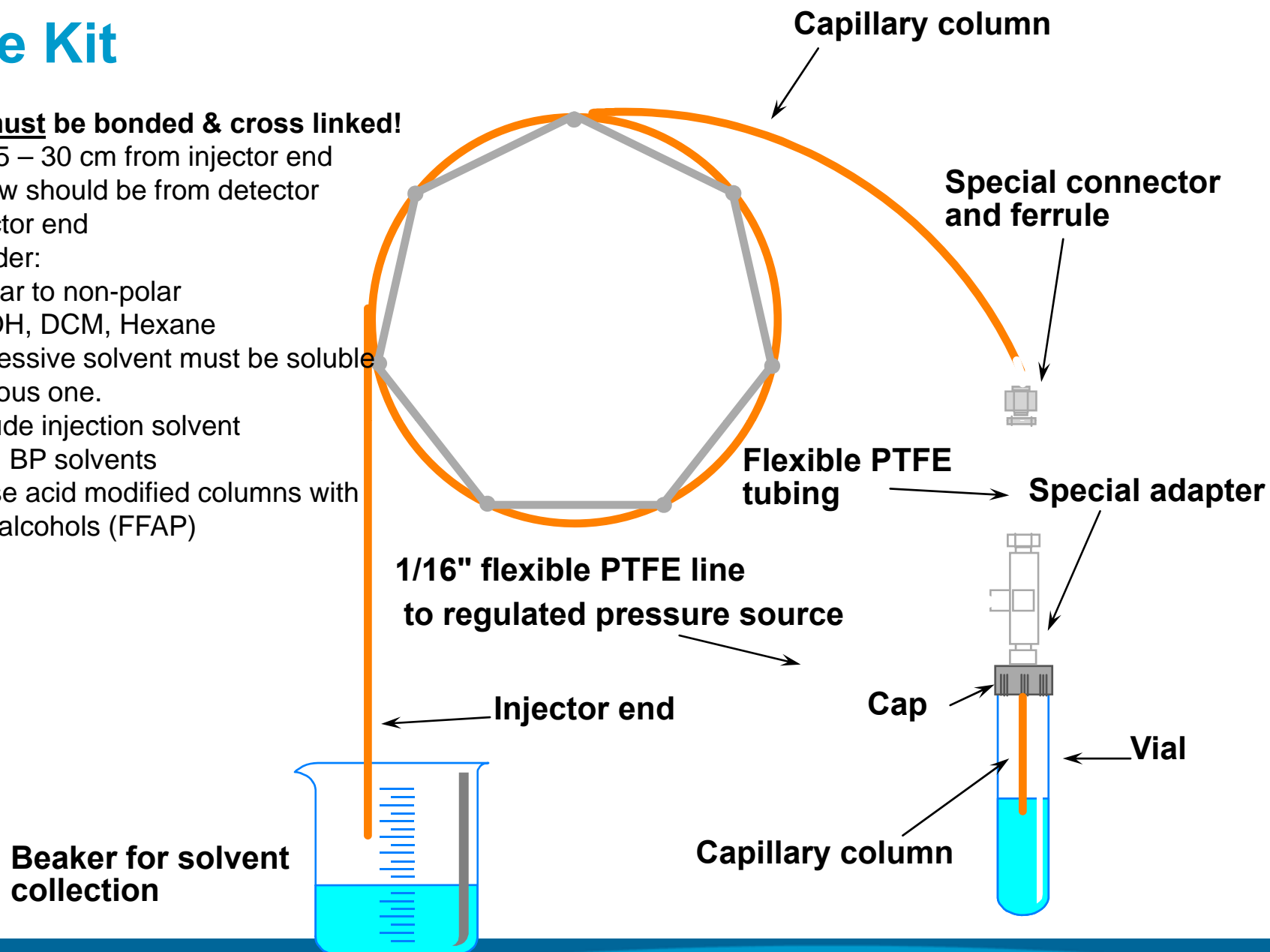
-i.e. MeOH, DCM, Hexane

> Each successive solvent must be soluble in the previous one.

> Try to include injection solvent

> Avoid High BP solvents

> Do not rinse acid modified columns with water or alcohols (FFAP)



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# Semi-Volatile Contamination

## What To Do If It Happens

- **“Bake out” the column**
  - **Limit to 1-2 hours**
  - **Longer times may polymerize some contamination and reduces column life**
- **Solvent rinse the column**



# Column Storage

- **Place septa over the ends**
- **Return to column box**





## Always Remember to:

- **Start with a good installation**
- **Maintain an oxygen free system**
- **Avoid physical, thermal, and chemical damage**
- **Take steps to prevent contamination**



# TECHNICAL SUPPORT

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

**866-422-5571 (FAX)**

**E-mail:**

**[gc-column-support@agilent.com](mailto:gc-column-support@agilent.com)**



# Wrap-up e-Seminar Questions

**Thank you for attending today's Agilent e-Seminar.**

**Our Seminar schedule is expanding regularly.**

**Please check our web site frequently at:**

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