## Gas Chromatography Cheat Sheet by shaylannxd via cheatography.com/149855/cs/32547/

Basic Theory		Column Type
Gas Chromatog- raphy	SP 🗲 Liquid	Wall Coated ( (WCOT)
	<ul><li>MP → Inert gas</li><li>No role in separation</li></ul>	
	<ul> <li>Only directs analyte down column (carrier gas)</li> </ul>	Fused Silica ( (FSOT)
	Dm >>> Ds	
	► CmU ~ 0	
	Flow rate <ul> <li>Dictate by choice of SP (thickness, proper-</li> </ul>	Column Diagr
	4: )	

- ties)
- Modest plate height ~1mm (↑ L = ↑N)





Column Type	
Packed	Packed full of particles
	Put SP on particles
	MP pushes through packed bed
	Tubing <ul> <li>Glass, stainless steel, etc.</li> </ul>

Inert = not part of separation

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Open Tubular Inside wall of quartz/glass tube Chemically roughen Surface area Coated with SP Open Tubular SP coating on wall of long thin tube Smooth wall

# Diameter ~ 75-200 um

#### ım

(cont)



GC Systems		
Sample	<ul> <li>Introduce into injector port</li> <li>Vaporize sample</li> <li>Vaporized analyte swept into column</li> </ul>	
Mobile phase	<ul><li>High pressure cylinders</li><li>Use a gas flow regulator</li><li>Regulate the pressure</li></ul>	
Detector	<ul><li>Detect components of the mixture being eluted off the chromatography column</li><li>Some may require a reference flow</li></ul>	
Oven	<ul> <li>Separation occurs</li> <li>Controlled temperature</li> <li>Fan → Circulates air and controls temperature</li> </ul>	

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GC System Diagram

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#### Split Flow Injector

Sample dissolved in volatile solvent

Collect sample into syringe and inject through rubber septum

- Seals injector for analyte to go into column
- Protects from outside atmosphere
- Bad peak shapes = hole in septum

Use a heat block to "flash" sample into vapour

- ~50-100C hotter than oven
- Need to vaporize sample

GC systems design to operate with 3 main columns

- Each column has a different flow rate
- Adjust based on column used

#### FSOT/WCOT

- Can't handle large sample mass
- Small diameter
- Limited SP
- Limited volume capacity
- Control by valve system

#### Split flow outlet

- Avoid overloading the column
- Packed > Set at 0 (closed)
- ▶ FSOT/WCOT → Split flow ratio → Depends on [analyte] in injection volume



# Requirements of SP (cont) Prefer a low volatile solvent → Don't want SP to vaporize in oven Thermal SP in column Stability Don't want thermal breakdown products Inert/Rea Don't want analyte to react with SP ctive Only want to interact

Stationany Dhase		
Stationary Phase		
Siloxane Polymer	Low volatility	
	Thermally stable bond	
	Contains a silicone backbone	
	Close to inert	
	<ul> <li>Can be derivatized</li> </ul>	
	Add pendant functional groups <ul> <li>Tune selectivity/solubility/retention</li> <li>Adjust polarity</li> </ul>	
Non-Polar	Poly(dimethyl)siloxane (PDMS)	
	<ul> <li>Good quality</li> </ul>	
	Flurocarbons	
Polar	Can replace dimethyl/methyl groups <ul> <li>CN,CO,OH</li> </ul>	
Phenyl Groups (Benzene Ring)	Non-polar	
	$\pi e^{-}$ delocalized	
	<ul> <li>When approach by polar molecules</li> <li>e<sup>-</sup> reorganized → Induced dipole interactions</li> <li>Can behave polar with polar molecules (vice versa)</li> </ul>	
Chiral Moiety	Chiral-chiral interactions on SP	
	Rise to selectivity of 1 enantiomer over another	

**Requirements of SP** 

	or solid	
	A substance with low volatili	ty is more likely to be a liquid
	vapour	
Volatility	A substance with high volati	lity is more likely to exist as a
	No separation	
	Affects R'~0	
	Unretained	
	Bad SP	
Solvent	Must dissolve analyte	

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Siloxane Polymer		Minimize Loss of SP	(cont)
			<ul> <li>Silanol</li> <li>Very polar</li> <li>Expose on surface of silica</li> <li>Disagree with SP polarity</li> <li>Competition for polar analyte</li> </ul>
	$\begin{array}{c} H_{5}C=\sum_{j=1}^{j-1} C=\sum_{j=1}^{j-1} C=\sum_{j=1}^{j-1} C=J_{3}\\ CH_{5}\\ poly(dmethylysiowane \end{array}$		<ul> <li>Deactivation chemistry</li> <li>Use dichloro dimethyl silane</li> <li>Use ethanol/MeOH</li> <li>Create less polar surface</li> </ul>
Minimize L	loss of SP	Inertness of Column	Residual silanols
Bonded Phase	<ul><li>Process of the SP polymer is attached to</li><li>Silica support particle</li><li>Wall of a capillary</li></ul>		<ul> <li>React strongly to polar compounds</li> <li>Produce tailing peaks</li> <li>Undesirable interactions in column</li> </ul>
	A liquid-liquid chromatography method in which a	Deactivation of Silan	
	stationary phase is covalently bound to a carrier particle		
Cross Linked Phase	Polymer attached to wall		
	<ul><li>Polymer cross-linked with each other</li><li>Critical for separation</li></ul>		Dichloro Dimethyl silane
	<ul> <li>Produce more rigidty, hardness and  Melting point</li> <li>Formation of covalent bonds</li> </ul>		
Issue	Most SP are non-polar and silica support surface are polar Not much intertaction		Ethanol MeOH
	Uses phases to prevent issue of contact		H <sub>3</sub> C O
	Use silane reaction to bond/cross-link		o~   `o~   `o~ `o
Silane Reaction	<ul><li>Use to anchor/bond silicones to silica surfaces</li><li>In packing materials (particles)</li><li>FS capillaries</li></ul>	Silane Reaction Mec	hanism
	Use to deactivate silanols		Terminal <u>silanols</u>
	Same chemistry for polymerization and cross-linking		

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Controlling Retention		Controlling Retention (cont)	
Retention	Controls resolving power (R') <ul> <li>R' depends on K'</li> </ul>		Adjust temperature during course of separation
	<ul> <li>K' depends on separation conditions</li> <li>Want all peaks to fall in "ideal range" of retention</li> </ul>		Resolution imporves under better retention conditions for the analyte
	<ul> <li>1-10</li> <li>MP is inert</li> <li>Only function to control retention</li> <li>Equilibrium constant = thermodynamic property</li> </ul>		<ul> <li>Change in gradient steep → Improves</li> <li>separation</li> <li>Shorten separation time</li> <li>Increase resolution</li> <li>As a function of temperature</li> </ul>
	Temperature <ul> <li>Alter overall retention</li> </ul>	Round-Up of T Programming	Powerful tool for controlling K'
	Type of SP		Directly affects distribution constant
	<ul> <li>Alter selectivity</li> </ul>		↑ Temperature = ↓ K'
Impact of Different	Isothermal separation <ul> <li>A thermodynamic process, in which the</li> </ul>		Ramped (gradient) temperature is used to adjust K'
Temperature temperature ↓ Tem Less Ana	<ul> <li>Temperature of the system remains constant</li> <li>Temperature =      Thermal energy available     Less thermal energy     Analyte spends more time in SP     </li> </ul>		<ul> <li>Make GC less intuitive</li> <li></li></ul>
	<ul> <li>More time in column</li> <li>Clearer separation</li> </ul>		Separation limited by $\Delta T/\Delta t$ (ramp rate)
	✓ Temperature =↑ Resolution = ↑ Overall		Column lifetime is shorter at higher temper- ature
	<ul> <li>Can become excessive</li> <li>Needs to adjust separation as it proceeds</li> </ul>	Other Factors	K'=K(Vs/Vm) ▶ SP thickness
	✤ Temperature ➔ Favors SP		
	↑ Temperature → Favors MP		<ul> <li>Calculate phase ratio (Vs/Vm)</li> </ul>
Different Ramp Rates	Altered tr and resolution independently		

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# Cheatography

#### Graphs with Different Temperature



#### **Ramp Rate Graphs**



#### **GC Detectors**

#### Requir- Sensitivity

ements  $\bullet$  10<sup>-8</sup>-10<sup>-15</sup> g analyte/s

Packed All sample used Decrease efficiency = broader peaks

► FSOT → Split flow injector (5-10% sample used) → Increase efficiency = narrow peaks

#### Stability

 Noise on baseline > Smooth {[fa-arrow-right}] Detect the smallest peaks > Minimal DL

Drift > No baseline (goes up and down)

#### LDR

▶ 5-8 orders of magnitude

#### GC Detectors (cont)

Can accept MP over a wide temperature range

- T Programming {[fa-arrow-right}] Improves separation
- Immune to T change
- ▶ Compensate T change → Require reference gas flow
- Fast response and independent of T

Simple to use, maintain, repair

#### Selective/Universal

- Detect analyte of interest (S)
- Detect all species (U)

Non-destructive

#### Flame Ionization Detector (FID)

Analyte elute from column

- Mix with H2 gas
- Combusted

#### Reduced carbons

Produce ions that alter conductivity of flame and alter current

Signal proportional to # of reduced carbons

#### Mass sensitive

Oxidized and e<sup>-</sup> capturing species

- No-little signal
- Cannot be oxidized further

#### Non-combustible gasses

- No signal
- Already oxidized

#### High sensitivity

- ▶ 10^-13& g/s
- use FSOT/WCOT

#### Large LDR

7 orders of magnitude

Destructive

No reference flow

#### FID Diagram





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Electron Capturing Detector (ECD)	Thermal Conductivity Detector(TCD) (cont)		
<ul> <li>response is based upon the ability of molecules with certain</li> <li>unctional groups to capture electrons generated by the radioactive ource</li> </ul>		Modest sensitivity ~ 10 <sup>-9 to -10</sup> g/ml <ul> <li>Less sensitive than FID</li> </ul> Modest LDR <ul> <li>Very short linearity</li> </ul>	
Radioactive source → <sup>63</sup> Ni ► Emits beta-particles		Non-destructive Require a reference flow	
<ul> <li>Vhen disintegration occurs</li> <li>Large energy release</li> <li>Beta particle emission</li> </ul>	Basic Theory	Based on ability of the gas exciting the column to absorb heat	
<ul> <li>Impacts any filler gas and/or MP present in detector and ionize it</li> <li>Jse a N2 make-up gas</li> <li>Get ionized by high energy</li> <li>Japired N2 gas an electric surrent through detector cell</li> </ul>		Contains thin filament <ul> <li>electrically heated</li> <li>As heat capacity of gas changes (MP vs</li> </ul> MP+analyte), so does the T of the filament	
<ul> <li>A constant current established through the detector</li> <li>When analyte with e<sup>-</sup> capturing groups</li> </ul>		<ul> <li>Resistance of thin filament</li> <li>T changes the resistance</li> <li>Resistance changes the current of the circuit</li> </ul>	
<ul> <li>Quench some ionization</li> <li>Reduce conductivity of gas = reduce current in cell</li> <li>Selective detector</li> <li>analytes with a high electron affinity</li> </ul>	Reference Flow (Type 1)	<ul> <li>Current is VERY sensitive to 1</li> <li>To compensate for the T of MP coming from the oven</li> <li>T is changing with T programmed elution</li> <li>Ieft section of diagram</li> </ul>	
<ul> <li>Sensitive for species that can disrupt ionization of N2 gas</li> <li>Pesticides → halides, peroxides, nitro groups</li> </ul>		Equation <ul> <li>Vout1 = Vapplied * (Rref/(Rcolumn+Rref))</li> </ul>	
CD Diagram		<pre>If Rcolumn = Rref   two resistors are "balanced"   The signal from the column is coming from the MP   Vout1=(1/2)Vapp</pre>	

#### Thermal Conductivity Detector(TCD)

Properties Signal proportional to change in heat capacity

 Difference between MP and MP+analyte are relatively small

Universal detector

- . . . . . .
- Detect solvent as well
- ► Undersirable → Solvent order of magnitude is more concentrated than analyte
- Result in large solvent peaks and small analyte

peaks

If analyte is not well retained > Interfered by

solvent

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Thermal Conductivity Detector(TCD) (cont)		Thermal Conductivity Detector(TCD) (cont)		
	If Rcolumn ≠ Rref Analyte's heat capacity changes T → heating or cooling of filament Vout1 increases as analyte elutes → As Rcolumn gets closer to 0, Vout1 gets closer to Vapp		<pre>If Rcolumn ≠ Rref    As analyte elutes    Vout1 increases    Vout2 decreases    Vout1 and Vout2 have same magnitude,    opposite signs</pre>	
Reference Flow (Type 2)	<ul> <li>W Opposite concept as reference flow type 1</li> <li>right section of diagram</li> </ul>		<ul> <li>Taking the difference of the two will double the</li> <li>V measured</li> <li>Double the signal for the same effort</li> </ul>	
	Equation <ul> <li>Vout2 = Vapplied * (Rcolumn/(Rcolum</li> <li>n+Rref))</li> </ul>	Reference Flow (Type 4)	<b>w</b> Same as Type 3, but with a single power supply	
	<pre>If Rcolumn = Rref   two resistors are "balanced"   The signal from the column is coming from   the MP</pre>		<ul> <li>Wheatstone bridge</li> <li>Name of this circuit</li> <li>Common approach for detecting VERY small change in resistance</li> <li>Advantage: Doubles the signal magnitude</li> </ul>	
	<pre>If Rcolumn ≠ Rref     Analyte's heat capacity changes T     Heating or cooling of filament     Vout2 decreases as analyte elutes     As Rcolumn gets closer to 0, Vout2 gets     closer to 0</pre>	TCD Diagram		
Reference Flow (Type 3)	Type 1 and Type 2 TCD operating together <ul> <li>With separate power supplies</li> </ul>	GC-MS Properties	Versatile	
	<pre>If Rcolumn = Rref   two resistors are "balanced"   The signal from the column is coming from   the MP</pre>		<ul> <li>Provide identification power</li> <li>Have to run known standards (spiked)</li> <li>Electron beam ionization</li> <li>M<sup>+</sup> and fragments</li> <li>Excellent DL</li> </ul>	
			<ul> <li>Depending on instrument and analyte</li> <li>~ 2-20 picog injected</li> </ul>	

Concentration DL in sample

Depends on sample work-up

#### Long LDR

- Dependent on instrument
- ▶ 4-6 orders of magnitude

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GC-MS (cont)		GC-MS (cont)	
	Selective <ul> <li>Less interferences</li> <li>Filters out MP signal</li> </ul> Destructive Expensive	<ul> <li>Isotopes</li> <li>▶ Parent ion</li> <li>→ Most prominent and heaviest</li> <li>▶ Isotopes</li> <li>→ Daughter peak from most prominent peaks</li> <li>&gt; can provide more inferdemending on its ratio with parent peak</li> </ul>	
Basic Theory	Basic       Quadruple MS         Fheory       • Contains 2 positive and 2 negative poles         Movement of M <sup>+</sup> • M <sup>+</sup> travels in a sinusoidal path	<ul> <li>Isotopically labelled analytes</li> <li>Replacing parts of molecule with deuterium</li> <li>Produces a known mass higher than the original mass</li> <li>Compare spectrum with orignal</li> </ul>	
<ul> <li>If M<sup>+</sup> is too light or too heavy, it is kicked out of quadrupole</li> <li>⇒ b/c they are not really able to respond to polarity change</li> <li>How to fix this</li> <li>⇒ Quickly change the frequency and voltage of the poles</li> <li>→ Can quickly scan through all m/z ratio to obtain mass spectrum</li> </ul>	<ul> <li>Positive identification</li> <li>Compare experimental spectrum with the "real" analyte spectrum</li> <li>3 steps</li> <li>Correct mass of molecule?</li> <li>Correct set of fragments?</li> <li>Correct fragment intensities?</li> </ul>		
	<ul> <li>Can quickly scan through all m/z ratio to obtain mass</li> <li>spectrum</li> </ul>	Quantitation  ► Usually multiple ions monitored/measured  ► Validate ratio of peaks at the correct m/z ratio	
<ul> <li>Spectrum generated</li> <li>Total ion current (TIC)</li> <li>Easiest way</li> <li>Sum of all ion signals that passes through</li> <li>Acts as a universal detector: does not filter out MP signal</li> <li>Tells you how many species are present</li> <li>Extracted mass spectra</li> <li>Take a slice of TIC peak and see its fragments</li> </ul>		<ul> <li>Column bleed</li> <li>SP is boiling and bleeding out</li> <li>Leads to a rise in baseline</li> <li>Not good</li> <li>How to fix it</li> <li>Running at low T</li> <li>→ Purchase column made specifically for MS (\$\$\$)</li> </ul>	
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#### **GC-MS Diagram**



#### **Key Factors and Applications** When will 1. Analyte GC be Needs to be volatile useful ▶ Not proteins Unstable at high temperature Silation reaction Produce volatile products (Risk of contamination, loss, produce new products) ▶ Needs to be stable → Stable enough to transit the column 2. High enough concentration to detect Packed columns: great sample capacity but low resolving power and resolution FSOT: lower capacity (split flow) but high resolving power and resolution Detectors: Has a good sensitivity 3. Does sample require high R' separation Depends on the type of detector Universal = high R Selective = low R 4. Generally faster than LC Applic-Anti-dopping and forensics ations BAC (Crime/forensics labs) Pharmaceuticals Process control Quality control Research and development

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#### Key Factors and Applications (cont)

Food and Beverages

- ▶ Wine/alcohol
- Pesticides

Environmental

- Pesticides
- PAH and industrial solvents
- Oil/hydrocarbon spills

#### R&D

- Organic synthesis
- Catalysis (monitor products)

Industrial

- Feedstock
- Off gassing