## Chromatography Theory Cheat Sheet by shaylannxd via cheatography.com/149855/cs/32519/

Separation The	eory	Fundamental Processes (cont)	
Analyze Comp Process of unn sample		als Dispersion Band Broadening	
Requirements		Depends on structure of column <ul> <li>▲ Analyte mix = ▲ Dispersion</li> </ul>	
Stationary Phase (SP)	Fixed in column Interacts with analyte	Depends on diffusion of analyte ↑ Diffusion Coefficient = ↑ Dispersion	
Mobile Phase (MP)	Moves through/over SP Carries analyte	Depends on total time in column <ul> <li>↑ Time ↑ Diffusion = ↑ Dispersion</li> </ul>	
Interactions	No interaction with SP <ul> <li>Travel same speed as MP</li> <li>No retention = No separation</li> </ul>	Separation Process	
	<ul> <li>Interaction with SP</li> <li>Analyte are retained → Dispersion</li> <li>Part time in SP (v=0) and MP (same speed</li> <li>All analyte spends same amount of time in MP but diff. time in SP</li> </ul>	Occurs in tube/plate (TLC) Drive MP through column	
Fundamental F	Processes	► Use gas pressure (GC)   Store MP in HP-cylinder + attach to gas regulator	
Retention	Peaks located in chromatogram Analyte interaction with column • stationary phase: strongint. = slow rate	Introduce sample at top of column Allow MP to drive sample through/over SP Detector at end (emerges vs. time)	
	Control by thermodynamic property • alter property = alter retention Example: • Temperature (GC) • MP (LC) • SP • Analyte	Process Mobile Phase Flow (J) analyte 2 analyte 1 Mobile Phase front (unretained) Detector	

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Time

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Retention		Efficiency	
Measure Retention       Variables         (K')       L=Length of column         U = MP velocity         V = Analyte velocity         tm = retention time of MP         tr=retention time of retained species         K=distribution constant         Cs = [Analyte] in SP         Cm= [Analyte] in MP		Quantify Efficiency	<ul> <li>Treat chromatographic peaks like "Gaussian" peaks</li> <li>Mean = Retention time</li> <li>Quantify width peak</li> <li>standard deviation</li> <li>peak width</li> <li>Smaller width = better efficiency</li> <li>Narrow peaks = Good efficiency</li> </ul>
	tr $\rightarrow$ use to identify analyte Simple matrix $\rightarrow$ 1< K' <10 Complex matrix $\rightarrow$ 0.5< K' <20		<ul> <li>Clear separation</li> <li>Broad peaks = Poor efficiency</li> <li>Overlapping</li> </ul>
	<ul> <li>K'</li> <li>Determined by chromatogram</li> <li>Controlled by equillibrium</li> <li>Judge separation by the last peak retention value</li> </ul>	Peak Shapes	<ul> <li>Sample volume ~ 1% column volume</li> <li>Various processes in column spread into larger volume</li> <li>Often significant &gt; starting volume</li> <li>Ex: Inj.volume = 25uL and detection volume = 200uL</li> </ul>
Contr Ac Ad Ad			Desirable  Narrow peaks and small volume  "Gaussian Peaks"  Peak could emerge with neighbor peak dilution can form broadening
Retention Equations		Measure Efficiency	<ul> <li>Variables</li> <li>N = # of theoretical plate</li> <li>H = Height of theoretical plate (HETP)</li> <li>L = Lenght of column</li> <li>W = peak width at baseline</li> <li>σ = Standard deviation (unit of lenght)</li> </ul>
	$\overline{U} = \frac{L}{t_m} \qquad \overline{v} = \overline{U} \frac{1}{(1+k')}$		Desirable

▶ ↑ N = ↓ H = ↓ σ

 $\bar{v} = \frac{L}{t_m} \qquad \bar{v} = \frac{U}{(1+k')}$   $\bar{v} = \frac{L}{t_r} \qquad \bar{v} = \frac{L}{t_r} = \frac{L}{t_m} \frac{1}{(1+k')}$   $K = \frac{C_s}{C_m} \qquad k' = \frac{t_r - t_m}{t_m} = K \frac{v_s}{v_m}$ 

C

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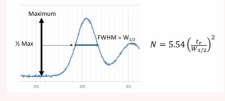
Efficiency (cont)	
	W range = -2t tp + 2t
	<ul><li>N should be consistent</li><li>tr and σ scale with each other</li></ul>
If Baseline not Accessible	Baseline peak width cannot be measured <ul> <li>nearby overlapping peaks</li> </ul>
	Use upper portion of peak that is undist- orted Use full-width at half maximum (FWHM) establish SD
	W1/2≠ 1/2 W

#### "Gaussian Peak" At W

$$\tau = \frac{\sigma}{\overline{v}} = \frac{\sigma}{L/t_r} = \frac{W}{4} \qquad \sigma = \frac{LW}{4t_r} = \frac{W}{4}$$

$$N = \frac{L}{H} = 16 \left(\frac{t_r}{W}\right)^2 = \left(\frac{t_r}{\sigma}\right)^2 \qquad H = \frac{\sigma^2}{L}$$
Baseline
Yest
Yest
W = 4T

#### Full-Width at Half Maximum



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Band Broaden	Band Broadening		
Occurrences	<ul> <li>Low efficiency</li> <li>Not fully separated peaks</li> <li>Interferences</li> <li>Dispersion is independent of retention</li> </ul>		
Van Deemeter Overview	<ul> <li>A-Term</li> <li>Associate with multiple flow paths through column</li> <li>Each unique distance</li> <li>Result in variety of times to transit column</li> </ul>		
	<ul> <li>B-Term</li> <li>Associate with longitudinal diffusion of analyte</li> <li>Some analyte will arrive sooner/later</li> <li>Depends on magnitude + direction of net diffusion during tr</li> </ul>		
	<ul> <li>C-Term</li> <li>Split into 2 sub-terms</li> <li>Relate to reality that chromatogram is carried out in non-equilibrium state</li> <li>Analyte in Mp will be out of equilibrium with those in SP (vice versa)</li> <li>Some analyte will arrive at detector earlier or later than true equilibrium would predicted</li> </ul>		
Van Deemeter Graph	<ul> <li>Produce the overall curve with distinct minimum</li> <li>Corresponding to Nmax and fixed L</li> </ul>		

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#### Band Broadening (cont)

#### **Overall Plate Height:**

- Equation:  $H = A + B/U + (C_S + C_m)U$
- Sum of 4 components (red line)

A-Term:

Constant (purple line)

#### B/U-Term:

Varies as 1/U (pink line)

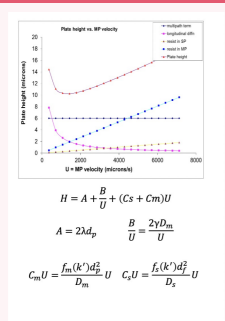
CsU-Term:

Linear increasing (blue line)

CmU-Term:

Linear increasing (yellow line)

#### Van Deemter Graph (copy)



#### A-Term: Multipath Band Broadening

All molecules start at top of column

As they move down

- Follow different paths through particles
- Irrespective of interaction with SP

Range of paths depends on size of particles

•  $\clubsuit$  Size =  $\clubsuit$  # of paths =  $\clubsuit$  Path length

Depends on how "packed: the bed is

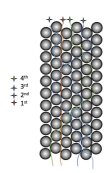
Crack, voids, etc

#### A-Term: Multipath Band Broadening (cont)

#### Equation:

- HA-term= 2λdp
- λ = quality/tortuosity factor
- ~0.5-0.6 (packed column)
- FSOT less

#### A-Term Diagram



#### B/U-Term: Longitudinal Diffusion

All molecules start at top of column

As they move down

- Molecules moves away from each other
- Process continues as long as they remain in column

Dispersion in all 3 directions

• Only longitudinal dispersion impacts peak width (  $\clubsuit$  and  $\clubsuit$ )

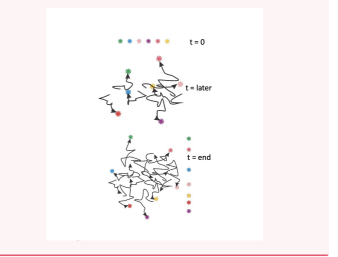
Packing column

- Reduce longitudinal diffusion = Plate height (Beneficial)
- Blocks molecules travel

#### Equation:

- $HB/U-Term = (2\gamma Dm)/U$
- Dm= Diffusion coefficient in MP
- γ= Obstruction factor
- ~ 0.6 (packed column)
- ~ 1.0 ( open tubular column)
- U= MP velocity

#### **B/U-Term Diagram**



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C-Term: Resistance to Mass Transfer		C-Term: Resistance to Mass Transfer (cont)	
C-Term	Ideal chromatography <ul> <li>Assumption that analyte can "instantly"</li> </ul>		Distance is proportional to size of particle
	equilibrate between 2 phases MP is always moving the analyte down		Equation: ▶ Hcm∪ = (fm(K')dp <sup>2</sup> *U)/Dm
	<ul> <li>Analyte in leading edge of peak are always moving over SP that is deficient in analyte</li> <li>Reverse for trailing edge</li> <li>Out of equilibrium</li> </ul>		<ul> <li>fm(K') = Quasi constant &gt; Depends on retention</li> <li>dp= Particle diameter (units)</li> <li>Dm= Diffusion coefficient of analyte in MP (cm<sup>2</sup>/s) &gt; 1 cm<sup>2</sup>= 10<sup>4</sup>mm</li> <li>U= MP velocity</li> </ul>
	Equilibrium established when there are analyte at:		
	► SP ► MP	CsU-Term: Resistance to Mass Transfer in SP	<ul><li>Space in SP depends on SP thickness</li><li>Distance required for diffusion</li></ul>
	<ul> <li>Interface</li> <li>Takes time for analyre to diffuse to/away from phases to match equilibrium constant</li> <li>In SP  Analyte gets further behind than expected</li> <li>In MP  Analyte gets further ahead than expected</li> </ul>		<ul><li>Analyte reach MP/SP interface</li><li>Equilibrium reach</li><li>Delays depends on distances</li></ul>
			Equation: <ul> <li>HCsU= (fs(K')df<sup>2</sup>*U)/Ds</li> <li>df= SP thickness</li> <li>Ds=Diffusion Coefficient of analyte</li> </ul>
	Rise to broadening		in SP
CmU-Term: Resistance to Mass Transfer in MP	<ul> <li>Space in-between particles depends on particles size/diameter</li> <li>Distance required for diffusion to move analyte</li> <li>Reach interface</li> </ul>		GC: ▶ ~0.1-0.5 µm film thickness ▶ Controls retention ▶ Impact resistance to mass transfer
	Delays in reaching equilibrium depends on distances	-	
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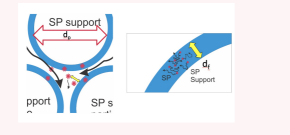
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#### C-Term: Resistance to Mass Transfer (cont)

#### LC:

- Never adjust to thickness Monolayer
- ▶ Resistance → Negligible
- Important in MP

#### CmU and CsU Term Diagram



Resolution	
Define Resolution	2 peaks of interest (critical pair) ▶ Peaks closest together
	Resolved <ul> <li>Clear separation</li> <li>No analyte mixing</li> <li>Pure peaks</li> </ul>
	W is not affected by plate height
Successful Separation	Isolated peaks
	See baseline between peaks
	Dependent on resolution
	Can use ruler to see if baseline from beginn- ing/end match to baseline between peaks
Quantify	Use W
Resolution (R/Rs)	<ul> <li>Captures +/- 2σ regions of "Gaussian" peaks</li> <li>Corresponds to ~ 95.5% of analyte</li> </ul>

#### Resolution (cont)

#### R improves:

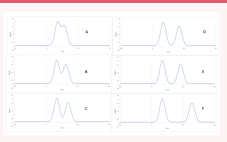
- ▶ Greater ∆tr
- Smaller Wa and/or Wb
- Narrow peaks = more baseline expose

Full W of peak does not matter

• Only back half (peak 1) and first half (peak 2)

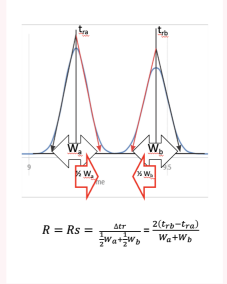
2 neighbouring peaks are resolved when  $\Rightarrow$  R ≥1.5

#### **Resolution Diagram**



Graph A-C has poor resolution → Overlapping peaks Graph D has the minimum resolution requirement Graph E-F has a good resolution → See baseline between peaks

#### **Resolution Equation**



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Controlling Resolving	Power	Controlling Resolving Powe	er (cont)
Control Resolution	<ul> <li>Proximity of 2 peaks is important to R</li> <li>Controlled by separation conditions</li> <li>Quantify proximity:         <ul> <li>Selectivity factor → Define as a ratio of distribution constant of 2 peaks</li> </ul> </li> <li>Peaks shares column → Same SP and</li> </ul>		N control by: Column (L) VD equations Type of column SP thickness Operating conditions
	<ul> <li>Peaks shares column - Same SP and MP</li> <li>α = ratio of retention factors</li> <li>Access from chromatogram</li> <li>Change in selectivity = change in resolution</li> </ul>	Effects of R' on Retention	<ul> <li>K' control by:</li> <li>SP type</li> <li>Phase Ratio (SP thickness)</li> <li>MP type (LC only)</li> <li>★ R'= ★ Total run time</li> </ul>
Effects of Retention and Selectivity on R'	Key variable that controls potential resolu- tions Resolving power (R')	Time	<ul> <li>Interplay</li> <li>Interplay between R' and tr as a function of K'</li> </ul>
	<ul> <li>R' dependent:</li> <li>Very sensitive to selectivity (α)</li> <li>Somewhat sensitive to retention (K')</li> <li>Moderately sensitive to efficiency of column (N)</li> <li>Choice of column</li> <li>Choice of MP (LC only)</li> </ul>		<ul> <li>R':</li> <li>N and α ~ constant when K' is alter</li> <li>Replace terms with Q</li> <li>trb:</li> <li>N,H,α, U ~ constant</li> <li>Assume R' is not changing</li> </ul>
	<ul> <li>α control by:</li> <li>Differential interactions between:</li> <li>Analyte ↔ MP↔SP</li> </ul>	Effects on Retention and S	dramatically <ul> <li>Replace constant terms with Q</li> </ul>
		$K = \frac{C_s}{C_m}$ $\alpha = \frac{\kappa_s}{\kappa_c}$	$k'_{a} = K_{a} \frac{v_{s}}{v_{m}} \qquad k'_{b} = K_{b} \frac{v_{s}}{v_{m}}$ $\frac{h}{a} \qquad \alpha = \frac{k'_{b}}{k'_{a}} = \frac{tr_{b} - t_{m}}{tr_{a} - t_{m}}$

$$\begin{split} R' &\approx \left(\frac{\alpha - 1}{\alpha}\right) \left(\frac{k'_b}{1 + k'_b}\right) \frac{\sqrt{N}}{4} & \longleftrightarrow \quad N \approx 16 R'^2 \left(\frac{\alpha}{\alpha - 1}\right)^2 \left(\frac{1 + k'_b}{k'_b}\right)^{\alpha} \\ & IF \; k'_\alpha \approx k'_b = k' \qquad THEN \; \alpha = \frac{k'_b}{k'_a} \approx 1 \\ R' &\approx (\alpha - 1) \left(\frac{k'}{1 + k'}\right) \frac{\sqrt{N}}{4} & \longleftrightarrow \quad N \approx 16 R'^2 \left(\frac{1}{\alpha - 1}\right)^2 \left(\frac{1 + k'}{k'}\right)^2 \end{split}$$

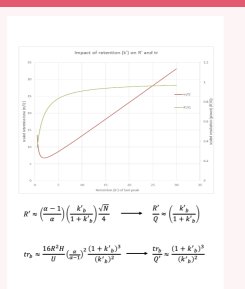
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#### Effects of R' on tr



Notice that R' increases significantly at low K' but plateaus at large K'

Don't use separations with small K' (low R')

Notice that tr increases linearly with increasing k' BUT R' plateaus at large k'

• Therefore there is no real benefit to sep'ns with large k's (b/c R'  $\approx$  constant)

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