

#### Liquid Chromatography

First chromatographic method described (as a non-instrumental method).

Since samples don't need to be initially vaporized, potentially any compound can be assayed by this method.

Instrumental development lagged behind that of GC because of difficulties in creating a stable solvent flow.



## Adsorption chromatography



The stationary phase is a solid. Separation is due to a series of adsorption/desorption steps.

# Adsorption chromatography

Silica and alumina are common stationary phases although there are a range of modified surfaces that have been evaluated.

Both solute and solvent are attracted to the polar sites on the stationary phase.

If solutes have differing degrees of attraction to the phase, a separation is possible.





## Partition chromatography

This approach comes closest to our countercurrent extraction model.

More highly retained species have a greater affinity (solubility) for the stationary phase - compared to the mobile phase (solvent)

Separation of solutes is based on differences in this relative solubility.









# Ion exchange chromatography For instrumental LC, weak exchange resins are typically used. These are exchange groups bound to a support. The traditional exchange resin beads would be crushed under normal HPLC conditions.





#### Size exclusion chromatography

Columns can be obtained that will separate specific size ranges.

Larger species will elute first - they can't pass through as many pores so their path is shorter.

Useful for determining size and size range for polymers, proteins, ...











#### Mode selection

#### In general

- If sample is water insoluble or non-polar use normal phase
- If sample is water soluble or not soluble but polar use reverse phase.
- It's not always this cut and dry but represents a good starting point.



lvent strer					
Solvent	63	P'	viscosity	RIU	V cutoff
n-pentane	0.00	~0.0	0.23	1.36	210
CCI4	0.18	1.6	0.97	1.47	265
toluene	0.29	2.4	0.59	1.50	285
ethyl ether	0.38	2.8	0.32	1.35	220
THF	0.45	4.0		1.41	220
мек	0.51	4.7		1.38	330
acetonitrile	0.65	5.8	0.37	1.34	210
methanol	0.95	5.1	0.60	1.33	210
E <sup>o</sup> is for alu	ımina.				













Sol	vent classes
Not	all solvents are truly usable.
•	Can't be mixed at any proportion
•	May interact chemically
•	UV absorption or viscosity is too high
•	Toxic, too flammable
•	High vapor pressure
•	Too expensive

Common reverse phase solvents	
methanol - acid acetonitrile - base tetrahydrofuran - large dipole water - polarity adjustment All are low viscosity available in high purity UV transparent miscible in each other	The four solvent methodA simple series of steps can be followed to obtain the optimum reverse phase solvent blend.1.Start with a single solvent and water. Adjust the % water from 0% on up until the best possible separation is obtained - optimum k' for peaks of interest.

### The four solvent method

- 2. Create blends using each of the other solvents and water that have the same solvent polarity.
- 3. Evaluate each solvent for improvements in peak shape or movement of selective peaks.
- 4. A mix of any of the blended solvents is then evaluated for optimum resolution.







#### Gradient elution

#### Can be produced

Stepwise - change from one solvent to another during a run.

Continuous - ramped - comparable to a temperature program.

Most common type is 'mixed stream' - solvents are pumped together with turbulent mixing. Each solvent is controlled by a programmer and the total flow rate is held constant.







