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Separation

Pure	Contain only ONE thing - have fixed melting and
substances	boiling points
Mixtures	Contain MORE THAN ONE thing, do not have exact properties, formed by a physical change so they can be easily separated

Purification as part of drug synthesis

Synthesis	Purification	Character-
		isation
Weigh up starting materials	Extraction	NMR
Set up a reaction	Recrystallisation	IR
Monitor reaction	Distillation	MS
Work up reaction to start	Chromatography	RF
purification	Solid-phase	Melting point
	extraction	

Separation techniques

1. Distil-	Used to separate the components of a LIQUID mixture by
lation	vapourising, condensing vapours and then collecting the
	liquid condensate.
	Separation is a result of different boiling points
2.	Physically separated solids from liquids
Filtration	Especially used for insoluble solids in liquids
3.	Uses centrifugal force that spins the samples fast
Centri-	Separated even fine solid matter form liquids
fugation	

Separation techniques (cont)

4. Recrystal-	1. Dissolve impure crude material in minimum
lisation	volume of solvent
	2. Filter any insoluble material
	3. Allow solution to cool slowly
	4. Crystals will form

Polarity

Definition of solubility

Ability of a solvent to dissolve in a solute, depending on the nature of the forces acting between the solute and solvent - LIKE DISSOLVES LIKE

e.g. **lonic** compounds are more soluble in **polar** solvents **Covalent** compounds are more soluble in **non-polar** solvents

Solvent mixtures

Immiscible: if the mixtures are too different, they will NOT mix, and will form separate layers or 'phases'

Partially miscible: mixtures are not too different but not too similar, they may mix at determinate proportions depending on the compositions

Miscible: if they are similar, they mix in EVERY proportion, forming one layer or 'phase'

Liquid-liquid Separation

What is liquid-liquid separation?

A method in which 2 molecules initially present in one sample can be separated by giving them the choice of different immiscible solvents

- Very common when isolating or purifying a product
- Can be used to extract natural products
- Normally done in workup stage after a reaction

Criteria of organic solvent for liquid-liquid extraction

SHOULD readily dissolve the substance to be extracted SHOULD NOT react with the substance to be extracted

SHOULD NOT react with or be miscible with water (usual second solvent)

SHOULD have a low boiling point so it can be easily removed from the products

Common extraction solvents are dichloromethane or ethyl acetate

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Chromatography

What is chromatography?

Physical method of separation where the components to be separated are distributed between 2 phases: the stationary and mobile phase

Common types:

- TLC (thin layer)
- Column
- HPLC (high performance liquid)
- GC (gas)
- SEC (size exclusion)
- Ion exchange
- Chiral

Chromatography - definitions

Analyte	Substance being separated (or analysed)
Mobile	Phase that moves in a definite direction - consists of
phase	sample being separated and the solvent that moves the
	sample through the column
Stationary	The substance fixed in place for the chromatography
phase	procedure
phase Eluent	procedure Solvent entering column
•	
Eluent	Solvent entering column

DIFFERENT METHODS & RETENTION OF ANALYTES

1.	Separation is based on differences between the	
Adsorption	adsorption affinities of the sample analytes for the	
chroma-	surface of a solid-stationary phase	
tography	Based on non-colavent interactions	
	Used for organic molecules	

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DIFFERENT METHODS & RETENTION OF ANALYTES (cont)

2.	Separation based on differences between the solubility
Partition	of the sample analytes in the mobile and stationary
chroma-	phases \rightarrow stationary phase = immobilised liquid
tography	Based on non-covalent interactions
	Used for organic polar molecules
3.	Separation based on exclusion effects e.g. differences in
Exclusion	size and shape
chroma-	used for proteins and nucleic acids
tography	
4. Gel	Specialised example of size-exclusion chromatography
electr-	which uses agarose/polyacrylamide and passes electr-
ophoresis	icity through
	Separates based on size and shape
	Used for proteins and nucleic acids
	Visualised by using UV and stains

DIFFERENT METHODS & RETENTION OF ANALYTES (cont)

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INSTRUMENTAL CHROMATOGRAPHY

	ETHODS & RETENTION OF ANALYTES (CONT)	INSTRUMENTAL	HRUMATUGRAPHY	
5. Ion exchange chromatog- raphy	Separation based on differences in ion exchange <pre>affinities → e.g. differences in charge size and type Based on ionic forces Used for cations, anions, proteins, peptides, amino acids and nucleic acids - Cation exchangers = negative stationary phase Anion exchangers = positive stationary phase</pre>	mance liquid chromatography 2. Gas chroma- tography	Analytical use: identiry complex mixtures Preparative use: purific Separated by their vola Forces the analyte thro stationary phase by the high pressure and tem	cation atility bugh a column of the e gas mobile phase at a
6. Thin layer chromatog- raphy	Separation based on Kx values Polar stationary phase = thin layer of silica spread over a glass plate Stationary phase is placed in the mobile phase Analyte moves up the plate and components are spread based on their Kx values Lower Rf = compound is MORE polar Higher Rf = compound is LESS polar Increased eluent polarity can increase the Rf More polar = stronger interactions in the normal phase Less polar = stronger interactions in the reverse phase	Quantitative analys Single point calibr-		Internal standard
		ation 1. A solution containing a known concentration of the compound to be measured in the sample is injected relation concen- tration obtained =AUC	e containing different concen- trations of	1. One solution containing a known concentration of the compound to be measured AND a known concentration of internal standards are injected to find a Response factor
7. Flash column chromatog- raphy	Used for large amounts of material (5-25mg) when TLC is not effective	2. The sample with unknown concen- tration is injected → new AUC is compared with previous to find	with unknown	2. The sample and the same known concen- tration of internal standard are injected to find the AUC and Response factor

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find concentration

in the sample

concentration in the

sample

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Separation in pharmacy

Drugs in the clinic must be pure:

- 1. Impurities can be harmful and cause side effects
- 2. Impurities can alter the ability to formulate a drug correctly
- 3. Impurities can also affect the stability and shelf-life of the drug

Drugs in development MUST be pure:

This is to ensure that it is the drug that brings about a therapeutic effect and not any impurities

- Before any new compound is tested, it is purified and characterised

Separation is also part of the quality control and monitoring process

Importance of ionisation

Some functional groups can be charged, and this depends on:

- whether the molecule is an acid or a base
- the pH of the molecule

Physiological pH = 7.4

we need to know what groups are charged at this pH as it will help us determine:

- types of drug-target binding interactions
- solubility (uptake and distribution)
- potential salt forms

e.g.1 - Carboxylic acids	e.g. 2 - Aliphatic amines
Acids react with water:	Bases react with water:
$HA + H2O \Rightarrow H3O+ + A-$	$B+H2O \rightleftharpoons BH+OH\text{-}$
pKa of carboxylic acids ~ <5	pKa of aliphatic amines ~ >8

These functional groups are almost ALWAYS ionised at physiological pH

pKa & pH = measures of dissociation

If the pKa = pH, then the functional group is 50% ionised

Partitioning

Partition Distribution of a solute between 2 solvents

Compound is present in BOTH phases according to its relative solubility in both

Solute distributes itself between the 2 liquids in accordance with its **partition coefficient**

Partitioning (cont)

Partition	Ratio of the concentration of the solute in one liquid	
coefficient	OVEr the concentration in the other	
	Dynamic equilibrium exists between the 2 liquids, temperature dependent	

Partitioning in pharmacy

Relative hydrophobic/hydrophilic properties of a drug are crucial as it influences:

- solubility
- adsorption
- distribution
- metabolism
- excretion

Drug too polar? It will not cross the cell membranes across the gut wall

Drug too lipophilic? Drug will be poorly absorbed, so will likely be taken up into the fat tissue and not circulated

LogP

P = Concentration of drug in octanol Concentration of drug in water

LogP: used as a measure of hydrophobicity of a drug

Hydrophobic/hydrophilic character is measured using partition

- Hydrophobic molecules prefer the $\textbf{octanol layer} \rightarrow \text{high P}$

- Hydrophilic molecules prefer the water layer \rightarrow low P

LogP measures only unionised forms of the drug between octanol and water

LogD is sued to represent relative distributions of all species, charged or uncharged

Distribution Coefficient (Kx)

$$K_x = \frac{C(stat)}{C(mob)}$$

C(stat) = conc of X in **stationary** phase C(mob) = conc of X in **mobile** phase

Each compound will have a different Kx Chromatographic separations can be altered by changing the nature of the stationary and/or mobile phase(s)

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Solid-phase extraction

Used in sample preparation to remove matrix interferences such as proteins

Concentrates sample of interest

Solid phase in SPE is similar to the stationary phase used in chromatography

Active substances can be retained or unretained

SPE vs. Chromatography

- Column is much smaller for SPE
- Specialised columns are more common
- Analytes are typically strongly retained on the SPE column, then impurities are washed away
- In chromatography, both the analyte and impurities are passed through at different rates
- SPE is used for sample preparation, then chromatographic methods follow

Stationary phases

Normal phase	Reverse phase	
Stationary phase = polar Mobile phase = non-polar	Stationary phase = non-polar Mobile phase = polar	
Components elute in order of increasing polarity	Components elute in order of decreasing polarity	
Used in TLC and flash column	Used in HPLC	
More polar compounds are retained stronger More polar solvents increase elution	Less polar compounds are retained stronger Less polar compounds increase elution (e.g. methanol, ethanol)	
Solvents = the complete range but rarely greater than 20% for very polar solvents	Solvents = very polar solvents (e.g. water, methanol)	
A strong solvent for the normal phase is a weak solvent for the		

stationary phase

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Component retention

For TLC, retention is expressed in terms of Rf

For column, retention is expressed in terms of column volumes (CV) The relationship between Rf and CV are reciprocal

Lower Rf = greater $CV \rightarrow Low RF$ if preferred as it increases analyte contact time and improves chances of component separation or resolution

To separate adjacent compounds, a large ${\bigtriangleup}\text{CV}$ is desired

Order of elution

TLC and HPLC depend on polarity

- Normal phase: polar retained more, non-polar elutes first
- Reverse phase: non-polar retained more, polar elites first

Gas chromatography depends on volatility or molecular weight

- Highly volatile or low MW elutes first

Size exclusion depends on molecular size

- Low size retained more, largest size elutes first
- Ion exchange depends on charge
- Complementary charges are retained more