





UNIT 4 CHEMISTRY – AREA OF STUDY 2 SUMMARY NOTES FOR THE VCAA EXAMS

WRITTEN BY A STUDENT WHO OBTAINED A NEAR PERFECT STUDY SCORE

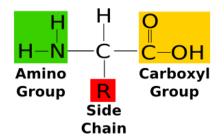
AREA FOOD CHEMISTRY

AMINO ACIDS

- the monomers that make up proteins (enzymes, hormones, keratin, antibodies)
- plants can manufacture all their amino acids, animals obtain these from food
- difunctional—contain both a weakly acidic carboxyl group COOH and a weakly basic amino group NH2
- both functional groups are attached to a central alpha carbon *distinguishing difference is the R group
- 20 amino acids
- some are essential (cannot be synthesised by the body) others are non essential (can be synthesised)
- both functional groups are polar, making amino acids soluble (extensive H bonding with water)
- all (except glycine) are chiral

STRUCTURE:

- carbons are numbered from right to left (carboxyl group is one)
- R-Groups can be:
 - ➤ non-polar (CH3)
 - > polar (CH2COOH)
 - >> proton donors (CH2COOH)
 - ➤ proton acceptors (CH2NH2)



AMPHIPROTIC ACTIONS:

- amino acids are amphiprotic (can act as acid or base) depending on pH of solution *as an equilibrium reaction, one form will be most dominant
- **zwitterion** (pH 5-7): electrically neutral form of an amino acid which contains both a positive and a negative charge (isoelectric point)
- Acid solutions: positive charge when COO- group gains a H+ *acts as a base*
- Basic solutions: negative charge when NH3 group loses a proton *acts as an acid*

Effect of R Group: some amino acids have basic or acidic R groups due to presence of NH2 or COOH which can influence charges e.g Asn @ pH 3 will have 2+ charge

PROTEIN FORMATION

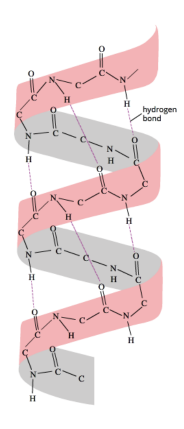
- formed via condensation polymerisation
- peptide links **AMIDE BONDS** -CO-NH- are formed to form a polypeptide (proteins have more than 50 amino acids in the polypeptide chain)
- can be dipeptides=two amino acids, trippedide= three amino acids
- during digestion, proteins are hydrolysed to produce the individual amino acids (requires water and enzymes)

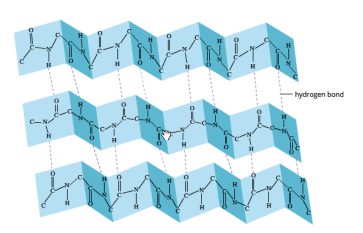
*always two isomers of the dipeptide depending on which way they condense (orientation)

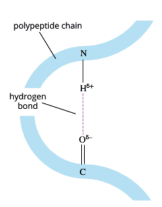
N terminal: NH3 endC terminal: COOH end

PROTEIN STRUTURES

- primary structure: the number, type and sequence of the amino acid residues in the protein—joined by strong covalent bonds *the length is determined by the gene that codes for the protein*
- **secondary structure:** three-dimensional structure, due to Hydrogen bonding between neighbouring amide groups
- hydrogen bonding between non-bonding electron pairs on an oxygen atom (C==O) and a hydrogen atom (NH2) from the same amino acid*results in an alpha helix
- interchain hydrogen bonding between polypeptides resulting in beta pleated sheets







tertiary structure: bending and folding of the protein

overall 3D structure

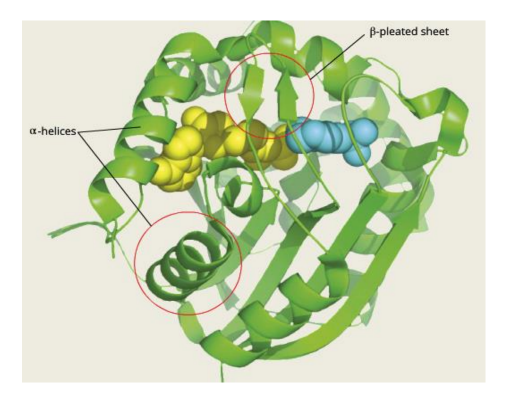
hydrophobic R groups fold inwards (away from water)

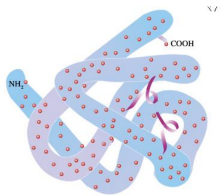
long and narrow—fibrous proteins responsible for structural functions

tightly folded alpha helix chains—globular proteins such as enzymes

arise due to bonding between R groups such as ion-ion interactions (NH3+/COO-), hydrogen bonding (C=O/OH), dipole-dipole interactions, dispersion forces

disulphide bridges: due to cysteine which bonds to another cysteine to form very strong S-S covalent bonds





denaturation: results in a loss of function due to...

pH change

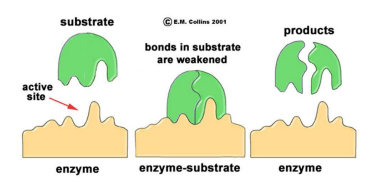
heat

presence of salts

involves uncoiling and unfolding of the helices (does not break peptide links) *primary structure left intact*

ENZYMES

Enzyme Action



- biological catalysts that speed up the rate of reaction bu providing an alternate reaction pathway with a lower Ea
- differ from inorganic catalysts as they are—much more efficient, much more selective, easily denatured
- lock and key model: substrate and active site have exact complimentary shapes
- enzyme-substrate complex achieved through either dispersion forces, dipoledipole interactions or hydrogen bonds *causes bonds within substrate to weaken and break

CONDITIONS AFFECTING ACTIVITY:

temperature: as temperature increases, rate of reaction increases due to higher KE of particles and higher proportion of collisions, enzymes will DENATURE past their optimal point as intermolecular bonds break

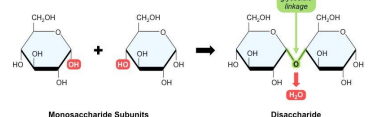
pH: each enzyme has an optimal pH based on acidic or basic nature of R group

CARBOHYDRATES

- most widely distributed and abundant organic compounds in the biosphere
- general formula is Cx(H2O)y
- monosaccharides are the most simple—C6H12O6
 - isomers are glucose, fructose, galactose
 - properties: crystalline solid, soluble (polar molecules), sweet tasting
 - role: cellular respiration to yield ATP
- * structure: energy and structural factors favour a more stable ring structure
 - rings produced through interactions within straight chain structure therefore have two geometric forms *differ in spatial arrangement of the hydroxyl and hydrogen groups on one carbon atom

Alpha-glucose and Beta-glucose, Plus Primary Structures of Amylose and Cellulose

- * disaccharides: formed by condensation reaction involving two monosaccharides e.g...
 - glucose + glucose -- maltose
 - glucose +fructose --> sucrose
 - glucose +galactose —> lactose



bond between monosaccharides is a glycosidic or ether bond (c-o-c)

 polysaccharides: complex, high Mr carbohydrates made from 10-10,000 monosaccharides e.g starch, cellulose, glycogen

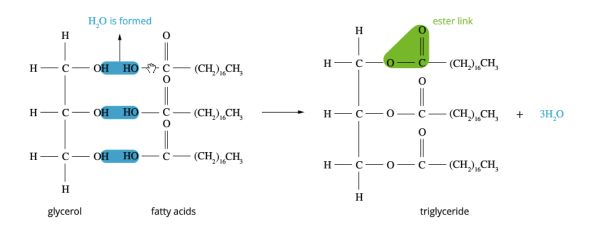
LIPIDS

- a class of compounds including fats and oils—primary function is a source of energy
- complex esters called glycerides formed via a condensation reactions between the alcohol glycerol C3H8O3 and fatty (carboxylic) acid molecules
- triglycerides are formed when three fatty acids are involved

Туре	R group	No of C=C	Example
saturated	CnH2n+1	0	C27H35COOH
monounsaturated	CnH2n-1	1	C17H33COOH
polyunsaturated	CnH2n-x	more than 1	C17H31COOH

- lipids are mixtures of all three types
 - animal fats are mainly saturated (solid at room temp) *due to stronger dispersion forces
 - vegetable oils are mainly unsaturated (liquid at room temp)

SYNTHESIS:



glycerol + 3 fatty acids —> triglyceride + 3H20

condensation reaction involving carboxyl group of a fatty acid and hydroxyl group of glycerol

DIGESTION:

- > lipids are large, non-polar and insoluble
- > bile (in intestine) converts lipids to a soluble form
- emulsified lipids then undergo hydrolysis to produce fatty acids and glycerol

SATURATED FATS:

made from only saturated fatty acids, generally unreactive and exist as waxy solids at room temp e.g cheese

UNSATURATED FATS:

- classified according to the position of the first double bond from the end of the hydrocarbon chain
- omega carbon is the carbon atom in the methyl group at the end of the hydrocarbon chain

e.g Linolenic Acid C17H31COOH is an **omega 3** fatty acid and has 3 C=C and the first is on the third carbon from the omega carbon

➤ Linoleic acid C17H31COOH is an omega 6 fatty acid, has 2 C=C and the first is on the sixth carbon atom from the omega carbon

MELTING POINTS:

fatty acids

- as chain length increases, strength of intermolecular (dispersion) forces increases, therefore melting point increases
- the presence of C=C in the R group means weaker dispersion forces (chains can't pack as closely together due to repulsions between electron dense double bond) + configuration of alkyl groups on the double bond in the cis arrangement
- melting point of saturated fatty acids is higher

lipids

- mixtures of saturated, mono and poly unsaturated triglycerides
- greater the percentage of saturated fatty acid residues, the stronger the dispersion forces therefore higher melting point *usually derived from animals fats and solid at room temp
- those predominately unsaturated have lower melting points, liquid at room temp and derived from plants

ESSENTIAL/NON-ESSENTIAL:

- most fats are non essential as they can be converted into other fatty acids
- all polyunsaturated fats are fatty acids such as linolenic and linleic acid

MACRONUTRIENTS + MICRONUTRIENTS

- Macronutrients: includes protein, carbohydrates and fats, minerals and water needed in high amounts for body functions
- Micronutrients: needed in the diet in very small amounts, are essential as cofactors for enzymes (such as digestive enzymes) e.g Fluorine, Zinc
- Vitamins: chemicals which cannot be synthesised by the body but are vital for metabolism, often cofactors for enzymes e.g Vitamin D needed for calcium absorption

SOLUBILITY OF VITAMINS:

> water soluble vitamins:

have more hydroxyl groups (form H bonds with water) allowing them to dissolve in blood

secreted by the body if not used

removed from foods if cooked in water e.g Vitamin B and C

fat soluble vitamins:

have few hydroxyl groups therefore fewer H bonds with water

build up on the fatty tissues of the body

can dissolve in non-polar environments due to dispersion forces between long non-polar chains e.g Vitamin D,E,A,K

METABOLISM OF FOOD

Nutrients:

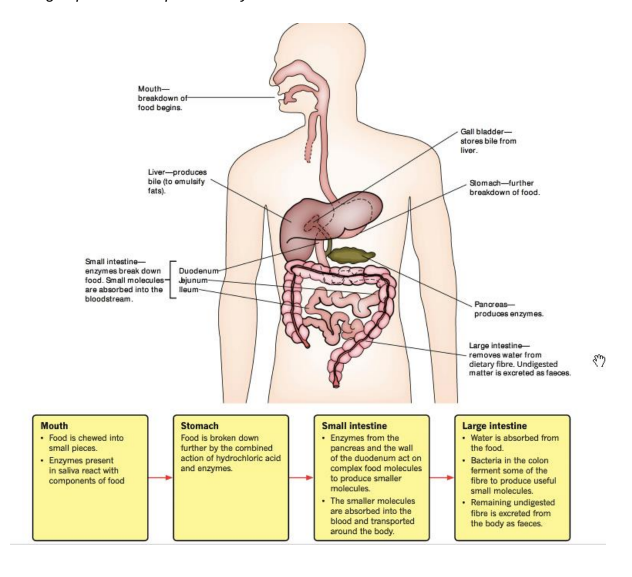
- → provide energy
- regulate growth and repair
- provide specialised roles eg disease prevention, cellular process

Metabolism:

- the chemical processes which occur within an organism that are necessary for the maintenance of life
- involves the breakdown of substances (nutrients from food) to yield energy for vital processes
- synthesis of larger molecules necessary for building structural tissue such as bone or muscle

DIGESTION:

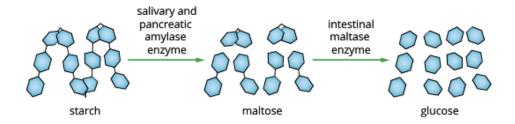
breakdown of large molecules in food into smaller molecules—a complex process involving separate and specific enzymes for each chemical reaction



CARBOHYDRATES: HYDROLYSIS

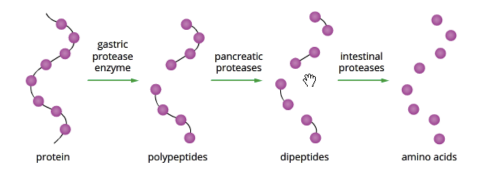
Polysaccharides e.g starch -> Disaccharides e.g maltose -> Glucose

• glucose produces glycogen (condensation polymerisation) to be an energy store for cell resp



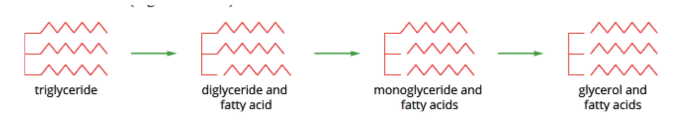
PROTEIN: HYDROLYSIS

- produces individual amino acids used to synthesise other proteins
- initially broken down by pepsin in stomach—produces smaller peptide chains which move to duodenum



TRIGLYCERIDES: FATS AND OIL

 catalysed by lipase—found in duodenum in small intestine and hydrolyses the ester links producing glycerol and fatty acids



^{*}digestion reactions are hydrolysis requiring water and enzymes and is exothermic

- lipids are insoluble in water—need to be converted into a soluble form before hydrolysis
- Bile emulsifies the fats—the hydrophobic tools of the bile is adsorbed at the surface of the fat, the hydrophilic head is then exposed to the aqueous solution
- lipase (water soluble) is involved in hydrolysis of fat to produce glycerol and fatty acids *only enzyme in this pathway
- glycerol and fatty acids then pass into bloodstream where they reform into triglycerides (stored in adipose tissue as an energy reserve)

RANCIDITY:

- fats deteriorate over time
- unsaturated fats are especially susceptible to rancidity—the C=C are reactive and undergo chemical reaction which may contain other functional groups (aldehydes and ketones)

hydrocarbon chain may break apart former smaller molecules *characterised by a change in smell or taste called **oxidative rancidity** (often forms aldehydes or ketones)

ANTIOXIDANTS:

- to decrease rate of deterioration food can be refrigerated, preservatives can be added or ANTIOXIDANTS can be used
- antioxidants—substances which slow the rate of oxidation of another substance by preventing the propagation
- act by intercepting the lipid peroxyl radical by donating a hydrogen
- able to slow down formation of free radicals (formed during autoxidation)
 e.g Vitamin C and E

AUTO-OXIDATION:

involves a free radical chain reaction—free radicals are highly reactive atoms or molecules with unpaired valence electrons

- 1) INITIATION:
 - cleavage of C-H bond in the fatty acid
 - requires energy due to strength of C-H bond
 - RH —> R + H
- 2) PROPAGATION:
 - free radical chain reaction in which free radicals react with O2 to form other free radicals
 - R + O2 -> RCOO -> RCOOH (hydroperoxide) + R
 - ROOH—> R'CHO (aldehyde) + R" +OH
- 3) TERMINATION:
 - process is completed when two radicals combine
 - R+ROO+ROOR+R2+O2

ENZYMES

- proteins and the biological catalysts for many chemical reactions in organisms
- not chemically altered in reaction
- more sensitive to changes in environment than inorganic catalysts
- highly specific—often can only catalyse one specific biochemical reaction

RELEVANCE OF OPTICAL ISOMERISM:

- amino acids are chiral molecules (except glycine where the R group is H)
- one of the two enantiomers of each amino acid is utilised in protein synthesis
- many substrate molecules have chiral centres, only one enantiomer will 'fit' into the active site, this is the biologically active isomer

MODELS OF ENZYME ACTION:

- Lock and key model (summarised early)
- Induced fit model: enzymes have a flexible 3-D shapes which can be slightly modified to fit the shape of the substrate, after reaction the enzymes shape is regained

COENZYMES:

- Cofactors (must be present for reaction to be catalysed) can be metal ions or non protein organic components called coenzymes which are required for enzyme activity e.g Mg2+ are cofactors in DNA replication to balance out negative charge of DNA
- interact with enzymes during catalysis and act as electron carriers or carriers of specific groups of atoms e.g acetyl CoA carries COCH3 or NADH
- this alters surface properties of the enzyme and enhance the binding properties of the enzyme, activating it
- may be chemically changes as a result of its involvement with an enzyme

GLYCAEMIC INDEX

- carbohydrate containing foods are rated on the glycemic index
- GI scale ranks foods according to their effect of blood sugar after two hours of consumption (100 is the standard, an equivalent amount of pure glucose)
- GI effect depends on:

type of starch whether starch polymers are entrapped within food fat and protein (and other substances) content of food

- useful to understand how the body breaks down carbohydrates, taking into account only available carbohydrates (total minus fibre)
- Low GI=values of 55 or less—release glucose more slowly and steadily due to slower digestion
- High GI =values greater than 70 release glucose very quickly
- linear polymer starch molecules (amylose) pack together more tightly and are less soluble than the branched polymer (amylopectin) —less access of enzymes to hydrolyse the ether links therefore amylopectin will digest MORE rapidly than amylose

ENERGY CONTENT

ENERGY FROM GLUCOSE:

- > a monosaccharides which is the bodies most preferred energy source
- starch (polysaccharide is hydrolysed to produce glucose)
- energy is obtained from glucose via CELLULAR RESPIRATION

Aerobic—C6H12O6 + 6O2 \rightarrow 6CO2 + 6H2O + 38 ATP \triangle =-2860kg/mol

Anaerobic—C6H12O6 —> 2CH3CH(OH)COO- + 2H+ (lactic acid) -120kg/mol or C6H12O6—> 2C2H5OH + 2CO2 (fermentation) -69kg/mol

ENERGY VALUES:

Nutrient	Energy content (kJ/g)	Energy value (available energy) kJ/g
Carbohydrates	17	17
Fats/Oil	39	37
Protein	24	17

- > high energy content of fats is due to a high degree of oxidation
- carbohydrates have more oxygen atoms then fats therefore have a higher degree of oxidation—fats have greater potential for oxidation therefore release more energy on combustion
- represented in kj/g as they are mixtures (no Mr) however glucose (pure substance) can have kj/mol

Energy available to the body:

- energy released when food is burned if often high then that after it has been digested due to...
 - incomplete absorption of nutrient by the body after digestion of food
 - incomplete oxidation of nutrients
 - heat loss—oxidation may result in energy being used up in heat loss

Working out energy content in foods:

e.g Find energy content of cashews that are 29% carbohydrates, 18% protein and 46% fats

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29x17 =493kj of carbs
18x17 =306kj of protein
46x37 =1702kj of fat
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$$(1702 + 306 + 493)/100 = 25 \text{kj/g}$$

CALORIMETRY

- energy released during a combustion reaction is transferred into the water in the calorimeter
- calibration factor: required so that energy change for the entire system for each 1degree celsius is known (units J/degrees celsius) once this is determined the calorimeter is CALIBRATED

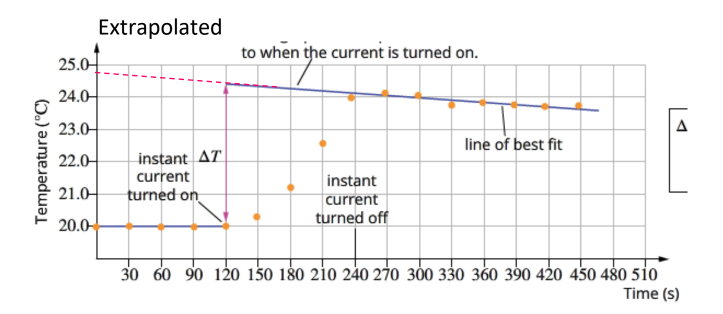
Electrical calibration:

- calibrates by using an electric heater to release a known quantity of thermal energy and measuring the resultant rise in temperature
- Energy = volts x current x time

$$CF = \frac{E}{\Lambda T} = \frac{VIt}{\Lambda T}$$

Temperature-time graphs:

- calorimeter is not perfectly insulated, it slowly loses heat during and after the heat is operating
- graphs at each time show accurately the change in temperature rather than finding delta T
- heat loss causes a decreasing gradient after heater is turned on—delta T is most accurately measure by extrapolating the line back to when heating commenced



Chemical calibration:

- performing a chemical reaction in the calorimeter that release known quantity of thermal energy
- often uses benzoic acid (C6H5COOH) enthalpy of combustion -3227kj/mol

$$E = n \times \Delta H_{c}$$
 $CF = \frac{E}{\Delta T}$