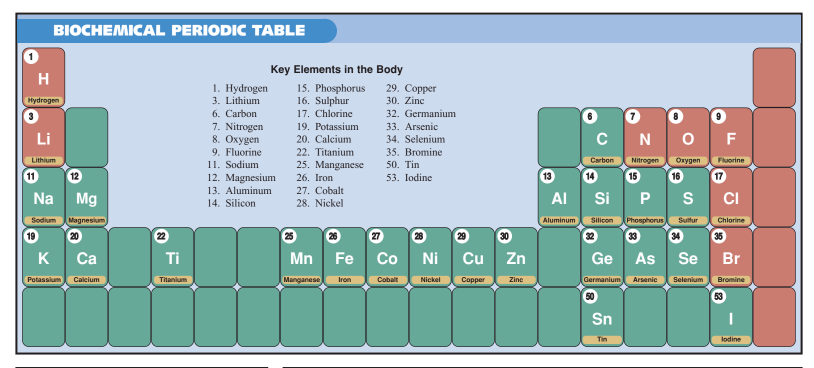
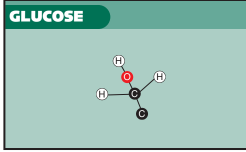
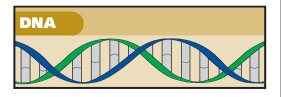
BarCharts,Inc.®

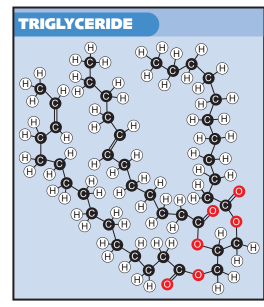
WORLD'S #1 ACADEMIC OUTLINE

Ouick Study Chemical principles BIOCCHEMICAL CONCEPTS organic compounds CHEMICAL CONCEPTS rates of reactions







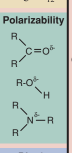


BROADER CHEMICAL PRINCIPLES

- A. Intermolecular Forces 1. Electrostatic: Strong interaction between ions; for charges q_1 and q_2 ; separated by r_{12} , and solvent dielectric constant, ε ; water has large ε ; stabilizes zwitterion formation Benergy = $\frac{1}{\varepsilon} \frac{q_1 \cdot q_2}{r_{12}}$ Polarizability R $\varepsilon = 0^{\delta}$
- 2. **Polarizability,** *α*: Measures distortion of electron cloud by other nuclei and electrons
- 3. **Dipole moment**, μ : Asymmetric electron distribution gives partial charge to atoms
- 4. London forces (dispersion): Attraction due to induced dipole moments; force increases with μ
- 5. **Dipole-dipole interaction:** The positive end of one dipole is attracted to the negative end of another dipole; strength increases with μ
- 6. Hydrogen bonding: Enhanced dipole interaction between bonded H and the lone-pair of neighboring O, N or S; gives "structure" to liquid water; solubilizes alcohols, fatty acids, amines, sugars, and amino acids
- B. Types of Chemical Groups
 - 1. Hydrophobic =
 - Lipophilic: Repelled

by polar group; insoluble in water; affinity for non-polar

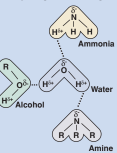
Examples: alkane, arene, alkene



Dipole Interaction $\delta^+ \delta^- \delta^+ \delta^$ stable $\delta^+ \delta^- \delta^- \delta^+ \delta^-$

 $\delta^+ \delta^- \delta^- \delta^+$ less stable

Hydrogen Bonding



 2. Hydrophilic = Lipophobic: Affinity for polar group; soluble in water, repelled by nonpolar Examples: alcohol, amine, carboxylic acid

3. **Amphipatic:** Polar and nonpolar functionality; common for most biochemical molecules: fatty acids, amino acids and nucleotides

C. Behavior of Solutions

- 1. **Miscible:** 2 or more substances form 1 phase; occurs for polar + polar or non-polar + non-polar
- 2. **Immiscible:** 2 liquids form aqueous and organic layers; compounds are partitioned between the layers based on chemical properties (acid/base, polar, nonpolar, ionic)

3. Physical principles:

- a. **Colligative** properties depend on solvent identity and concentration of solute; a solution has a higher boiling point, lower freezing point and lower vapor pressure than the pure solvent
- b.**Biochemical example: Osmotic pressure -** Water diffuses through a semi-permeable membrane from a hypotonic to a hypertonic region; the flow produces a force, the osmotic pressure, on the hypertonic side

Osmotic Pressure Π = IMRT

- $\Pi = IMRT$ Π : Osmotic pressure (in atm)
- **i:** Van't Hoff factor = # of ions per solute molecule
- M: Solution molarity (moles/L)
- **R:** Gas constant = 0.082 L atm mol⁻¹ K⁻¹
- **T:** Absolute temperature (in Kelvin)

4. Solutions of gases

- a. **Henry's Law:** The amount of gas dissolved in a liquid is proportional to the partial pressure of the gas b. Carbon dioxide dissolves in water to form carbonic acid c. Oxygen is carried by hemoglobin in the blood
- d.Pollutants and toxins dissolve in bodily fluids; react with tissue and interfere with reactions
 - **Examples:** Sulfur oxides and nitrogen oxides yield acids; ozone oxidizes lung tissue; hydrogen cyanide disables the oxidation of glucose

BONDS & STRUCTURE IN ORGANIC COMPOUNDS

A. Bonding Principles

1. Most bonds are polar covalent; the more electronegative atom is the "-" end of the bond **Example:** For >C=O, O is negative, C is positive 2. Simplest Model: Lewis Structure: Assign

Resonance =0 _0 N

valence electrons as bonding electrons and nonbonding lone-pairs; more accurate bonding models include valencebonds, molecular orbitals and molecular modeling

3. Resonance: The average of several Lewis structures describes the bonding

Example: The peptide bond has some >C=N< character **B. Molecular Structure**

Typical Behavior of C, N & O

· · · · · · · · · · · · · · · · · · ·							
Atom	sp ³	sp ²	sp				
C 4 e ⁻ 4 bonds	-C-C-	>C=C<	-C≡C-				
N 5 e ⁻ 3 bonds, 1 lone pair	>N-	R=N-	-C≡N				
O 6 e ⁻ 2 bonds, 2 lone pairs	-0-	>R=O					

1. Geometries of valence electron hybrids:

- sp² planar, sp³ tetrahedral, sp linear 2. Isomers and structure
- a. Isomers: same formula, different bonds
- b.Stereoisomers: same formula and bonds, different spatial arrangement
- c.Chiral = optically active: Produces + or rotation of plane-polarized light
- d.D: Denotes dextrorotary based on clockwise rotation for glyceraldehyde
- e.L: Denotes levorotary based on counter-clockwise rotation for glyceraldehyde; insert (-) or (+) to denote actual polarimeter results
- f. D/L denotes structural similarity with D or L glyceraldehyde
- g.Chiral: Not identical with mirror image
- h.Achiral: Has a plane of symmetry
- i. Racemic: 50/50 mixture of stereoisomers is L(-) - Glyceraldehyde optically inactive; + and - effects cancel
- j. R/S notation: The four groups attached to the chiral atom are ranked a,b,c,d by molar mass
- The lowest (d) is directed away from the viewer and the sequence of a-b-c produces clockwise (R) or counterclockwise (S) configurations •This notation is less ambiguous than
- D/L; works for molecules with >1 chiral centers
- k.Nomenclature: Use D/L (or R/S) and +/- in the compound name: Example: D (-) lactic acid

 CH_3

CH₂

Three-

dimensional

Cis

A H

Br

Alkene

Chain Positions

R

- 1. Fisher-projection: Diagram for chiral compound
- m. Molecular conformation: All molecules exhibit structural variation due to free rotation about C-C single bond; depict using a Newmandiagram
- n.Alkene: cis and trans isomers; >C=C< does not rotate; common in fatty acid side chains

C. Common Organic Terminology

- 1. Saturated: Maximum # of Hs (all C-C) 2. Unsaturated: At least one >C=C<
- 3. Nucleophile: Lewis base; attracted to the + charge of a nucleus or cation
- 4. Electrophile: Lewis acid; attracted to the electrons in a bond or lone pair

Carbon-chain Brofiver

			Carbon-cha	IIN I	refixes		
1	meth-	7	hept-	13	tridec-	19	nonadec-
2	eth-	8	oct-	14	tetradec-	20	eicos-
3	prop-	9	non-	15	pentadec-	22	docos-
4	but-	10	dec-	16	hexadec-	24	tetracos-
5	pent-	11	undec-	17	heptadec-	26	hexacos-
6	hex-	12	dodec-	18	octadec-	28	octacos-



 $\Delta G > 0$ endergonic

D. Standard-Free Energy

1. $\Delta \mathbf{G} = \Sigma \operatorname{prod} \Delta \mathbf{G}_{\mathrm{f}}^{0} - \Sigma \operatorname{react} \Delta \mathbf{G}_{\mathrm{f}}^{0}$

2. For coupled reactions: Hess's Law:

3. Combine reactions, add ΔG , ΔH , ΔS

4. An exergonic step can overcome an

Example: ATP/ADT/AMP reactions

are exothermic and exergonic; these

provide the energy and driving force

to complete less spontaneous

biochemical reactions; Example:

a.Equilibrium shifts to relieve the stress

due to changes in reaction conditions

b.Keq increases: Shift equilibrium to the

c.K_{eq} decreases: Shift equilibrium to the

2. Equilibrium and temperature

a.For an exothermic process, heat is a

b.For an endothermic process, heat is a

c.If $\Delta S > 0$, increasing T promotes

d.If ΔS < 0, decreasing T lessens

reactant; an increase in

temperature increases Keq

3. Entropy and Enthalpy factors

 $a.\Delta H < 0$ promotes spontaneity

 $b.\Delta S > 0$ promotes spontaneity

Note: T is always in Kelvin;

product; a decrease in temperature

 $ATP + H_2O => ADP + energy$

1. LeChatlier's Principle

product side

reactant side

increases K_{eq}

 $\Delta G = \Delta H - T \Delta S$

spontaneity

spontaneity

 $K = {}^{\circ}C + 273.15$

changes

equilibrium

Formation, ΔG_f^0 :

endergonic step

E. Equilibrium

not spontaneous small Kee

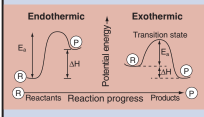
of

 $\Delta G = -RT \ln(K_{eq}) - \text{connection with}$

A. Mechanisms

QuickStudy

- 1. Biochemical reactions involve a number of simple steps that together form a mechanism
- 2. Some steps may establish equilibria, since reactions can go forward, as well as backward; the slowest step in the mechanism, the rate-determining step, limits the overall reaction rate and product formation
- 3. Each step passes through an energy barrier, the free energy of activation (E_a) , characterized by an unstable configuration termed the transition state (TS); E_a has an enthalpy and entropy component



B. Key Thermodynamic Variables

- 1. Standard conditions: 25°C, 1 atm, solutions = 1 M
- 2. Enthalpy (H): ΔH = heat-absorbed or produced
 - $\Delta H < 0$ exothermic $\Delta H > 0$ endothermic
- C. Standard Enthalpy of Formation, ΔH_{f}^{0}
 - 1. $\Delta \mathbf{H} = \Sigma \text{ prod } \Delta \mathbf{H}_{f}^{0} \Sigma \text{ react } \Delta \mathbf{H}_{f}^{0}$ 2. Entropy (S): $\Delta S =$ change in disorder
 - 3. Standard Entropy, S⁰:
 - $\Delta S = \Sigma \text{ prod } S^0 \Sigma \text{ react } S^0$
 - 4. Gibbs-Free Energy (G): $\Delta G = \Delta H - T\Delta S$; the capacity to complete a reaction $\Delta G = 0$ at equilibrium

steady state $\Delta G < 0$ exergonic

large Kee spontaneous

KINETICS: RATES OF REACTIONS

 $K_{ea} = 1$

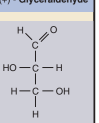
- A. Determination of Rate
 - For a generic reaction, $A + B \Rightarrow C$: 1. Reaction rate: The rate of producing
 - C (or consuming A or B) 2. Rate-law: The mathematical dependence
 - of the rate on [A], [B] and [C] 3. Multiple-step reaction: Focus on rate-determining step - the slowest
- step in the mechanism controls the overall rate **B. Simple Kinetics**

 - 1. First-order: Rate = $k_1[A]$ Examples: SN1, E1, aldose rearrangements
 - 2. Second order: Rate = $k_2[A]^2$ or $k_2[A][B]$ Examples: SN2, E2, acid-base,
 - hydrolysis, condensation

C. Enzyme Kinetics

2

- 1. An enzyme catalyzes the reaction of a substrate to a product by forming a
- stabilized complex; the enzyme reaction may be 103-1015 times faster than the uncatalyzed process 2. Mechanism: **Step 1.** $E + S = k_1 => ES$ **Step 2.** $ES = k_2 \implies E + S$ **Step 3.** $ES = k_3 \Rightarrow products + E$ [E] = total enzyme concentration, [S] = total substrate concentration, [ES] = enzyme-substrate complex concentration, k1 - rate ES formation, k2 - reverse of step 1, k₃ - rate of product formation
- 3. Data analysis: Michaelis-Menten Examine steady Equation: state of [ES]; rate state of [ES]; rate of ES formation $v = \frac{V_{max}[S]}{K_m + [S]}$ equal rate of disappearance $K_m = (k_2 + k_3)/k_1$ (Michaelis constant) v – reaction speed = $k_3[ES]$ $V_{max} = k_3[E]$



CH₃

CH₃

Fischer

projection

Trans

Br

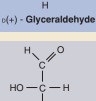
Br-

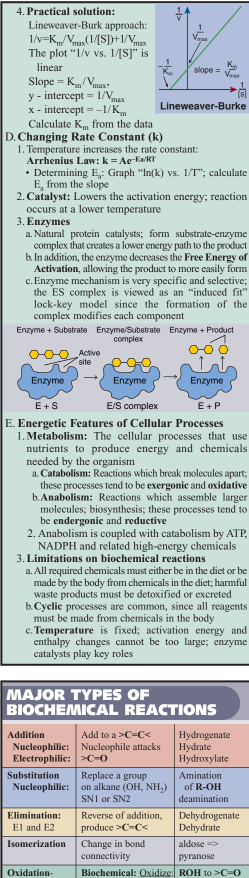
н-

110

— OH

-с — он





Add O or remove H

Reduce: Reverse of

Water breaks a bond,

add -H and -OH to

form new molecules

combine via bridging

R-NH or R-OH

Metals: Change

oxidize

valence

O or N

Hydrogenate

fatty acid

Hydrolyze

triglyceride

or amylose

Form peptide

peptide, sucrose

loss of e-

Reduction-

Hydrolysis

Condensation

gain of e-

Processes

Coupled

ORGANIC ACIDS & BASES

	Acid	Base
Arrhenius	aqueous H ₃ O ⁺	aqueous OH-
Brønsted-Lowry	proton donor	proton acceptor
Lewis	A A	electron-pr donor nucleophile

A. Amphoteric

- 1. A substance that can react as an acid or a base
- 2. The molecule has acid and base functional groups; Example: amino acids
- 3. This characteristic also allows amphoteric compounds to function as ОН single-component buffers for biological studies O = P - OH

B. Acids

- 1. $K_a = [A^-][H^+]/[HA]$ Phosphoric acid
- $p\bar{K}_a = -log_{10}(K_a)$ 2. Strong acid: Full dissociation: HCl, H₂SO₄ and HNO3: Phosphoric acid

ÓН

Purine

- 3. Weak acid: K_a << 1, large pK_a
- 4. Key organic acid: RCOOH
- Examples: Fatty acid: R group is a long hydrocarbon chain; Vitamin C is abscorbic acid; nucleic acids contain acid phosphate groups

Common Acids & pK_a Acid pK_a Acid pK. Acetic 4.75 Formic 3.75 6.35 Carbonic Bicarbonate 10.33 H₂PO₄ 7.21 HPO₄²⁻ 12.32 H₃PO₄ 2.16 NH_4^+ 9.25

- $pK_b = -\log_{10}(K_b)$
- 2. Strong base:

- 3. Weak base: K_b << 1,
- **Examples:** NH_3 (pK_b = 4.74), hydroxylamine $(pK_b = 7.97)$ and pyridine $(pK_b = 5.25)$
- 5. Purine: Nucleic acid component:
 - guanine (2-amino-6-hydroxypurine)

6. **Pyrimidine:** Nucleic acid component: cytosine (4-amino-2-hydroxypyrimidine), uracil (2,4-dihydroxypyrimidine) & thymine (5-methyluracil)

D. Buffers

- 1. A combination of a weak acid and salt of a weak acid; equilibrium between an acid and a base that can shift to consume excess acid or base
- 2. Buffer can also be made from a weak base and salt of weak base
- 3. The **pH of a buffer** is roughly equal to the pK_a of the acid, or pK_b of the base, for comparable
- amounts of acid/salt or base/salt 4. Buffer pH is approximated by the Henderson Hasselbalch equation

Note: This is for an acid/salt buffer

Henderson Hasselbalch Equation: $pH = pK_a + log (salt/acid)$

Common Buffers							
Buffer composition	approx. pH						
acetic acid + acetate salt	4.8						
ammonia + ammonium salt	9.3						
carbonate + bicarbonate	6.3						
diacid phosphate + monoacid phosphate	7.2						
E. Amino Acids	СООН						

1. Amino acids have amine (base) $H_0 N$ and carboxylic acid functionality; the varied chemistry arises from the chemical nature of the R- group L Amino acid

• Essential amino acids: Must be provided to mammals in the diet

- 2. Polymers of amino acids form proteins and peptides H_3N^{T}
- Natural amino acids adopt the L configuration
- 3. Zwitterion; self-ionization; the "acid" donates a proton to the "base"
 - · Isoelectric point, pI: pH that produces balanced charges in the Zwitterion

F ORGANIC COMPOUNDS

Type of Compound	Examples	
Alkane		ethane C_2H_{6} , methyl (Me) -CH ₃ , ethyl (Et) -C ₂ H ₅
Alkene	>C=C<	ethene C ₂ H ₄ , unsaturated fatty acids
Aromatic ring	-C ₆ H ₅	benzene - C6H6, phenylalanine
Alcohol	R-OH	methanol Me-OH, diol = glycol (2 -OH), glycerol (3 -OH)
Ether	R"-O-R'	ethoxyethane Et-O-Et, or diethyl ether
Aldehyde	O R-C-H	methanal H_2CO or formaldehyde, aldose sugars
Ketone	O R-C-R'	Me-CO-Me 2-propanone or acetone ketose sugars
Carboxylic acid	O RC-OH	Me-COOH ethanoic acid or acetic acid Me-COO ⁻ Acetate ion
Ester	O RC-OR'	Me-CO-OEth, ethyl acetate, Lactone: cyclic ester, Triglycerides
Amine	N-RR'R"	$\rm H_3C\text{-}NH_2,$ methyl amine, $\rm R\text{-}NH_2$ (1°) - primary, RR'NH (2°) - secondary, RR'R''N (3°) - tertiary
Amide	O R-C-NRR'	H ₃ C-CO-NH ₂ , acetamide Peptide bonds
Cyclic Ethers:		
Р	yran Furan	

- C. Organic Bases 1. K_b=[OH⁻][B⁺]/[BOH]
 - Full dissociation: NaOH, KOH
 - large pK_b
 - 4. Organic: Amines & derivatives

adenine (6-aminopurine) &

С-Н

COO

·с —н

R

Zwitterion

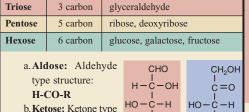
R

Pyrimidine

BIOCHEMICAL COMPOUNDS

- A. Carbohydrates: Polymers of Monosaccharides 1. Carbohydrates have the general formula
 - $(CH_2O)_n$ 2. Monosaccharides: Simple sugars; building
 - blocks for polysaccharides

Common Sugars



н-с-он structure: R-CO-R H-C-OH c. Ribose and deoxyribose: Key component in nucleic acids and ATP

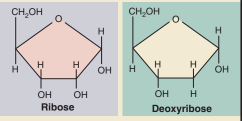
Aldose Ketose **D** Fructose D Glucose

ĊH₂OH

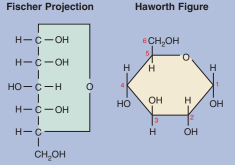
нċ−он

н-с-он

ĊH₂OH



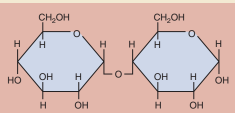
- d. Monosaccharides cyclize to ring structures in water •5-member ring: Furanose (ala furan)
 - •6-member ring: **Pyranose** (ala pyran)
 - •The ring closing creates two possible structures: α and β forms
 - •The carbonyl carbon becomes another chiral center (termed **anomeric**)
 - • α : -OH on #1 below the ring; β : OH on #1 above the ring
 - ·Haworth figures and Fischer projections are used to depict these structures (see figure for glucose below)



α-D-Glucopyronose

- 2.Polysaccharides
 - a. Glucose and fructose form polysaccharides b.Monosaccharides in the pyranose and furanose forms are linked to from polysaccharides;
 - dehydration reaction creates a bridging oxygen c.Free anomeric carbon reacts with -OH on opposite side of the ring
 - d.Notation specifies form of monosaccharide and the location of the linkage; termed a glycosidic bond

- e. Disaccharides •2 units
 - $\text{M-OH} + \text{M-OH} \rightarrow \text{M-O-M}$
 - •Lactose (β -galactose + β -glucose) β (1,4) link • Sucrose (α -glucose + β -fructose) α , β (1,2) link •Maltose (α -glucose + α -glucose) α (1,4) link



Maltose - Linked a D Glucopyronose

f. Oligosaccharides

- •2-10 units
- •May be linked to proteins (glycoproteins) or fats (glycolipids)
- •Examples of functions: cellular structure, enzymes, hormones
- g.Polysaccharides
 - •>10 units
 - **Examples:**
 - Starch: Produced by plans for storage
 - -Amylose: Unbranched polymer of α (1,4) linked glucose; forms compact helices
 - -Amylpectin: Branched amylose using α (1,6) linkage
 - -Glycogen: Used by animals for storage; highly branched polymer of α (1,4) linked glucose; branches use α (1,6) linkage
 - -Cellulose: Structural role in plant cell wall; polymer of β (1,4) linked glucose
 - Chitin: Structural role in animals; polymer of β (1,4) linked N-acetylglucoamine

3. Carbohydrate Reactions

- a. Form polysaccharide via condensation
- b.Form glycoside: Pyranose or furanose + alcohol
- c. Hydrolysis of polysaccharide
- d.Linear forms are reducing agents; the aldehyde can be oxidized
- e.Terminal -CH2-OH can be oxidized to carboxylic acid (uronic acid)
- f. Cyclize acidic sugar to a lactone (cyclic ester)
- g.Phosphorylation: Phosphate ester of ribose in nucleotides
- h.Amination: Amino replaces hydroxyl to form amino sugars
- i. Replace hydroxyl with hydrogen to form deoxy sugars (deoxyribose) Fatty Acid

 $\dot{c} = 0$

B. Fats and Lipids

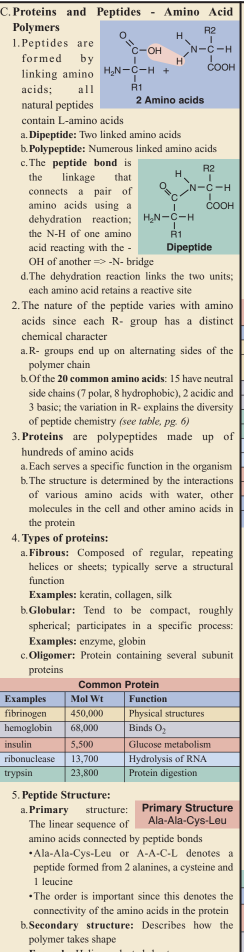
- 1. Lipid: Non-polar compound, insoluble in water Examples: steroids, fatty acids, нó triglycerides
- 2. Fatty acid: R-COOH
- Essential fatty acids cannot be synthesized by the body: linoleic, linolenic and arachidonic
- 3. Properties and structure of fatty acids: a. Saturated: Side chain is an alkane
 - b.Unsaturated: Side chain has at least one >C=C<; the name must include the position # and denote cis or trans isomer
 - c. Solubility in water: <6 C soluble, >7 insoluble; form micelles
 - d. Melting points: Saturated fats have higher melting points; cis- unsaturated have lower melting points

Common Fatty Acids							
Common Name	Systematic			Form	ıla		
Acetic acid	ethanoic			CH ₃ C	ООН		
Butyric	butanoic			C ₃ H ₇ C	СООН		
Valeric	pentanoic			C ₄ H ₉ C	СООН		
Myristic	tetradecanoic			C ₁₃ H ₂	7COOH		
Palmitic	hexadecanoic			C ₁₅ H ₃	1COOH		
Stearic	octadecanoic			C ₁₇ H ₃	5COOH		
Oleic	cis-9-octadecer	noic	;	C ₁₇ H ₃	3COOH		
Linoleic	cis, cis-9, 12 octadecadienoi	ic		C ₁₇ H ₃	1COOH		
Linolenic	9, 12, 15- octadecatrieno	ic (a	all cis)	C ₁₇ H ₂	9COOH		
Arachidonic	5, 8, 11, 14- eicosatetranoic	c (al	l trans)	C ₁₉ H ₃	1COOH		
	Saturated Stearic Acid						
4. Common fatty acid compounds							
a. Triglyceride or triacylglycerol: Three fatty acids bond via ester linkage to glycerol R1 - CO - O - CH2 R2 - CO - O - CH2 R3 - CO - O - CH2							
 b. Phospholipids: A Triglyceride phosphate group bonds to one of three positions of fatty acid/glycerol; R-PO₄ or HPO₄ group 5. Examples of other lipids 							
a.Stero	oids: Cholester nples: testoster	ol a	nd horr				
R = Nearly always methyl R' = Usually methyl R'' = Various groups R'' = Various grou							
Generic Steroid							
 b. Fat-soluble vitamins: Vitamin A: polyunsaturated hydrocarbon, all trans Vitamins D, E, K 							
-	6. Lipid reactions3 Fatty Acids + Glycerola. Triglyceride:R1-CO-OHHO-CH2						
		2-	co-o	нн	ю-сн		
-	ocess: vdration R	а-	co-o	нн	I IO-CH2		
dehydration no or							
b. The reverse of this reaction is hydrolysis of the							

- triglyceride
- c.Phosphorylation: Fatty acid + acid phosphate produces phospholipid
- d.Lipase (enzyme) breaks the ester linkage of triglyceride

Disaccharide

BIOCHEMICAL COMPOUNDS continued



- Example: Helix or pleated sheet
- Factors: H-bonding, hydrophobic interactions, disulfide bridges (cysteine), ionic interactions c. Tertiary structure: The overall 3-dimensional
- conformation

<u>QuickStudy</u>

- d.Quaternary structure: The conformation of protein subunits in an oligomer
- 6. Chemical reactions of proteins:
- a. Synthesis of proteins by DNA and RNA
- b. Peptides are dismantled by a hydrolysis reaction breaking the peptide bond
- c.Denaturation: The protein structure is disrupted, destroying the unique chemical features of the material
- d.Agents of denaturation: Temperature, acid, base, chemical reaction, physical disturbance
- 7. Enzymes
 - a.Enzymes are proteins that function as biological catalysts
 - b. **Nomenclature:** *Substrate* + *ase* Example: The enzyme that acts on phosphoryl groups (R-PO₄) is called phosphatase
- 8. Enzymes are highly selective for specific reactions and substrates

reactions and sub-	strates					
	Six Classes of Enzymes (Enzyme Commission)					
Туре	e Reaction					
1. Oxidoreductase	Oxidation-reduction					
Examples: oxidize CH- Oxygen acceptors: NAI						
2. Tranferase Functional group transfer						
Examples: transfer met	hyl, acyl- or amine group					
Hydrolysis reaction						
Examples: cleave carbo	xylic or phosphoric ester					
4. Lysase Addition reaction						
Examples: add to >C=C	C<, >C=O, aldehyde					
5. Isomerase Isomerization						
Example: modify carbohydrate, cis-trans fat						
6. Ligase Bond formation, via ATP						
Examples: form C-O, C-S or C-C						

Examples: form C-O, C-S or C-C

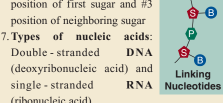
- 9. An enzyme may require a cofactor Examples: Metal cations (Mg²⁺, Zn²⁺ or Cu²⁺); vitamins (called **coenzymes**)
- 10. Inhibition: An interference with the enzyme structure or ES formation will *inhibit* or block the reaction
- 11. Holoenzyme: Fully functional enzyme plus the cofactors
- 12. Apoenzyme: The polypeptide component
- D. Nucleic Acids: Polymers of Nucleotides
 - 1. Nucleotide: A phosphate group and organic base (pyrimidine or purine) attached to a sugar (ribose or deoxyribose)
 - •Name derived from the base name
 - •Example: Adenylic acid = adenosine-5'monophosphate = 5' AMP or AMP
 - 2. Nucleoside: The base attached to the sugar •Nomenclature: Base name + idine (pyrimidine) or + osine (purine)
 - •Example: adenine riboside = adenosine; adenine deoxyriboside = deoxyadenosine

Nucleic Acid Components						
Base	Nucleoside	Nucleotide				
adenine	Adenosine	Adenylic acid, AMP				
	Deoxyadenosine	dAMP				
guanine	Guanasine	Guanylic acid, GMP				
	Deoxyguanisine	dGMP				
cytosine	Cytidine	Cytidylic acid, CMP				
	Deoxycytidine	dCMP				
uracil	Uridine	Uridylic acid, UMP				
thymine	Thymidine	Thymidylic acid, dTMP				
5						

- 3. Cyclic nucleotides: The Phosphate phosphate group attached to Sugar – Base the 3' position bonds to the Nucleotide 5' carbon 3', 5' cyclic AMP = cAMP and cGMP 4. Additional Phosphates
- a.A nucleotide can bond to 1 or 2 additional phosphate groups
- $b.AMP + P \implies ADP$ Adenosine diphosphate
- $ADP + P \Rightarrow ATP Adenosine triphosphate$
- c.ADP and ATP function as key biochemical energy-storage compounds
- 5. Glycosidic bond: Linkage between the sugar and base involve the anomeric carbon (carbon #1) >C-OH (sugar) + >NH (base) => linked sugar - base
- 6. Linking Nucleotides: The polymer forms as each phosphate links two sugars; #5 position of first sugar and #3 position of neighboring sugar

Double - stranded

single - stranded



B

Ρ

Ρ

Ρ

Ρ

S-T...A-S

S-C...G-S

S-G...C-S

Chargaff's

Rule

P

Р

Ρ

(ribonucleic acid) 8. Components of a nucleotide: sugar, base and phosphate

a.Sugar: ribose (RNA) or deoxyribose (DNA)

- b.Bases: purine (adenine and guanine) and pyrimidine (cytosine, uracil (RNA) and thymine (DNA))
- 9. In DNA, the polymer strands pair to form a double helix; this process is tied to base pairing

10. Chargaff's Rule for DNA:

- a.Adenine pairs with thymine (A:T) and guanine pairs with cytosine (C: G)
- b.Hydrogen bonds connect the base pairs and supports the helix
- c. The sequence of base pairs along the DNA strands serves as genetic for information reproduction and cellular control
- 11. DNA vs RNA: DNA uses deoxyribose, RNA uses ribose; DNA uses the pyrimidine thymine, RNA uses uracil
- 12. Role of DNA & RNA in protein synthesis a.DNA remains in the nucleus
 - b.Messenger-RNA (m-RNA): Enters the nucleus and copies a three-base sequence from DNA, termed a codon. m-RNA then passes from the nucleus into the cell and directs the synthesis of a required protein on a ribosome
 - c. Transfer-RNA (t-RNA): Carries a specific amino acid to the ribosomal-RNA (r-RNA) and aligns with the m-RNA codon
 - d.Each codon specifies an amino acid, STOP or START; a protein is synthesized as different amino-acids are delivered to the ribosome by t-RNA, oriented by m-RNA and r-RNA, then chemically connected by enzymes

COMMON AMINO ACIDS

COMMON AMINO ACIDS						
hydroph	obic = ye	ellow, b	asic = bl	ue, acidic = red, polar = green		
Amino acid MW essential - e		pK _a pK _b	pI R-pK _a	-R		
Alanine 89.09	Ala A	2.33 9.71	6.00	hydrophobic H ₃ C-		
Arginine e 174.20	Arg R	2.03 9.00	10.76 12.10	basic H_1 H_2 N-C-NH-CH ₂ -CH ₂ -CH ₂ -CH ₂ -		
Asparagine 132.12	Asn N	2.16 8.73	5.41	$\begin{array}{c} \mathrm{polar} & \mathrm{O} \\ \mathrm{H}_{2}\mathrm{N}-\mathrm{C}-\mathrm{CH}_{2}- \end{array}$		
Aspartate 133.10	Asp D	1.95 9.66	2.77 3.71	acidic HOOC-CH ₂ -		
Cysteine 121.16	Cys C	1.91 10.28	5.07 8.14	Polar HS-CH ₂ -		
Glutamate 147.13	Glu E	2.16 9.58	3.22 4.15	acidic HOOC-CH ₂ -CH ₂ -		
Glutamine 146.15	Gln Q	2.18 9.00	5.65	polar O H ₂ N-C-CH ₂ -CH ₂ -		
Glycine 75.07	Gly G	2.34 9.58	5.97	polar -H		
Histidine e 155.16	His H	1.70 9.09	7.59 6.04	basic		
Isoleucine e 131.18	Ile I	2.26 9.60	6.02	hydrophobic $CH_3 - CH_2$ HC - CH ₃		
Leucine e 131.18	Leu L	2.32 9.58	5.98	hydrophobic $CH_3 - CH_2$ $HC - CH_2$ $CH_3 - CH_2$		
Lysine e 146.19	Lys K	2.15 9.16	9.74 10.67	basic $H_2N-CH_2-CH_2-CH_2-CH_2-CH_2-CH_2-CH_2-CH_2$		
Methionine e 149.21	Met M	2.16 9.08	5.74	hydrophobic $CH_3 - S - CH_2 - CH_2 -$		
Phenylalanine e 165.19	Phe F	2.18 9.09	5.48	hydrophobic		
Proline 115.13	Pro P	1.95 10.47	6.30	hydrophobic $CH_2 - CH_2 + H$ $CH_2 - N - COOH$ H		
Serine 105.09	Ser S	2.13 9.05	5.68	polar HO – CH ₂ –		
Threonine e 119.12	Thr T	2.20 8.96	5.60	polar CH ₃ -CH- I OH		
Tryptophan e 204.23	Trp W	2.38 9.34	5.89	hydrophobic CH ₂ -		
Tyrosine 181.19	Tyr Y	2.24 9.04	5.66 10.10	HO-C ₆ H ₆ -CH ₂ -		
Valine - e 117.15	Val V	2.27 9.52	5.96	hydrophobic HC- CH ₃		

_	QuickStudy.
/ G	MINO ACID RNA CODONS
•	UUU UUC Thr
	ACU ACC ACA ACG Lys
	AAA AAG
	Leu UUA UUG CUU CUC CUA CUG
•	Ala GCU GCC GCA GCG
•	Asp GAU GAC
•	Glu GAA GAG
•	lle AUU AUC AUA
•	Tyr UAU UAC
•	Cys UGU UGC
•	Met START AUG
•	STOP UAA UAG UGA
•	Trp UGG
•	Val GUU GUC GUA GUG
•	His CAU CAC
•	Arg CGU CGC CGA CGG AGA AGG
•	Ser UCU UCC UCA UCG
•	Gln CAA CAG
•	Ser AGU AGC
•	Pro CCU CCC CCA CCG
•	Asn AAU AAC
•	Gly GGU GGC GGA GGG

ABI BIO	BREVIATIONS U LOGY & BIOCHE/	ISED MIST	TIN RY
aa	amino acid	Lys	aa lysine
А	aa alanine	М	aa methionine
	adenine - purine base		Molar (moles/L)
Ala	aa alanine	m	milli (10-3)
ADP	adenosine diphosphate	Man	mannose sugar
AMP	adenosine monophosphate	Met	aa methionine
Arg	aa arginine	mL	milliliter
Asn	aa asparagine	mm	millimeter
Asp	aa aspartate	N	aa asparagine
atm	atmosphere		Avogadro's number
	(pressure unit)		elemental nitrogen
ATP	adenosine triphosphate	n	nano (10-9)
С	aa cysteine	0	orotidine
	cytosine - pyrimidine		elemental oxygen
	elemental carbon	Р	aa proline
cal	calorie		phosphate group
Cys	aa cysteine		elemental phosphorous
D	aa aspartate	р	pico (10 ⁻¹²)
DNIA	Dalton	Phe	aa phenylalanine
DNA	deoxyribonucleic acid	Pro	aa proline
dRib	2-deoxyribose sugar	Q	aa glutamine
E	aa glutamate		coenzyme Q, ubiquinone
F	aa phenylalanine	R	aa arginine
Fru	fructose sugar		gas constant
G	aa glycine	Rib	ribose sugar
Gal	guanine - purine base	RNA	ribonucleic acid
Glc	galactose sugar	S	aa serine
Glu	aa glutamate		Svedberg unit
Н	aa histidine	s	second (unit)
н h	hour	Ser	aa serine
11	Planck's constant	Т	aa threonine
His	aa histidine		thymine - pyrimidine
I	aa isoleucine		absolute temperature
	inosine	Thr	aa threonine
	elemental iodine	Trp	aa tryptophan
Ile	aa isoleucine	Tyr	aa tyrosine
J	Joule (energy unit)	U	uracil - pyrimidine
К	aa lysine	V	aa valine
	Kelvin - absolute T		volt (electrical potential)
	elemental potassium	Val	aa valine
k	kilo (10 ³)	W	aa tryptophan
L	aa leucine		elemental tungsten
	liter (volume)	Х	xanthine
Lac	lactose sugar	Y	aa tyrosine
Leu	aa leucine	yr	year

Note: Source - CRC Handbook of Chemistry & Physics



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