Lecture 5 (9/16/24)

OUTLINE

Amino Acids

- Definition, Structure, and Properties
 - The 4 S's
 - · Common Properties
 - · Five Classes
 - Hydrophobic-aliphatic [6]
 - Hydrophobic-aromatic [3]
 - Special-sulfur [2]
 - Hydrophilic-polar [4]
 - Hydrophilic-charged [5]
 - · Other amino acids
 - · Linking amino acids
 - · Acid/base properties
- Titrations
- Isoelectric point
 - · Electrophoresis

Protein Purification

TODAY

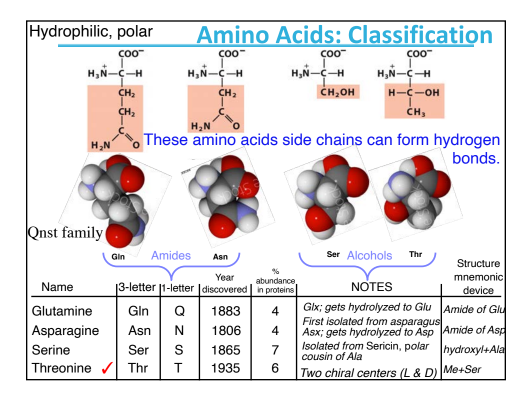
- Reading: Ch3; 76-82, 87-89
- Homework #5

NEXT

- Reading: Ch3; 83-87, 89-90
 - Ch1; Fig 7
 - Ch9; 313-314
- Homework #6

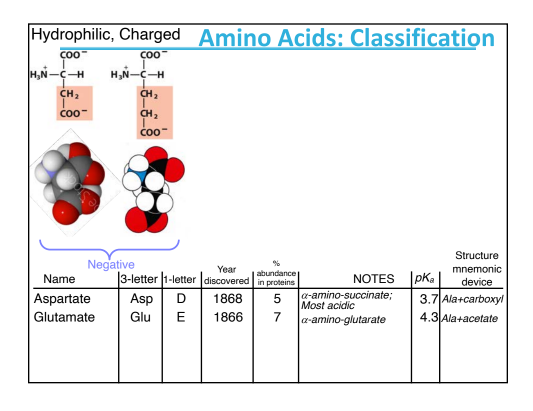
Amino Acids: Classification

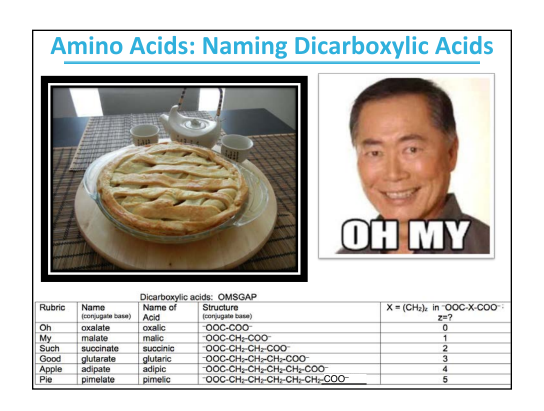
- The 20 amino acids found in proteins can be placed in five families based on the physical and chemical properties of their R groups:
 - Hydrophobic, aliphatic (6)
- Gavlip family
- Hydrophobic, aromatic (3)
- PTT family
- Special (hydrophobic/hydrophilic)(2) MC family
- Hydrophilic, polar (4)
- Hydrophilic, charged (5)

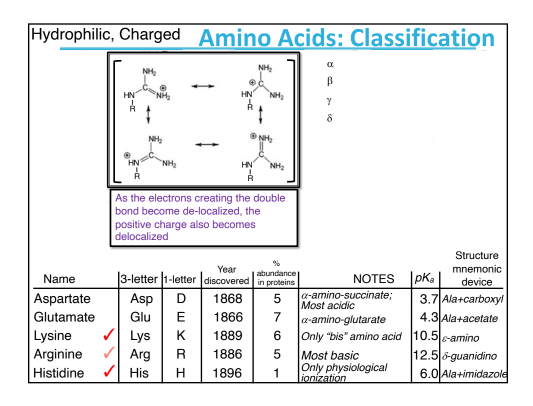


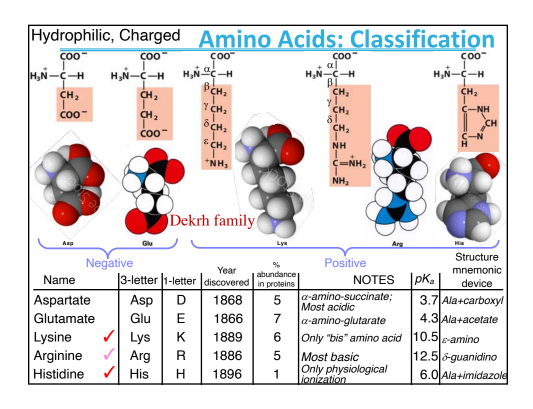
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Amino Acids: Classification

 The 20 amino acids found in proteins can be placed in five families based on the physical and chemical properties of their R groups:

• Hydrophobic, aliphatic (6) Gavlip family

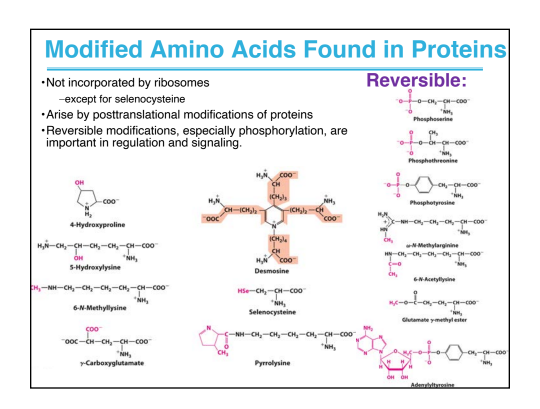
• Hydrophobic, aromatic (3) PTT family

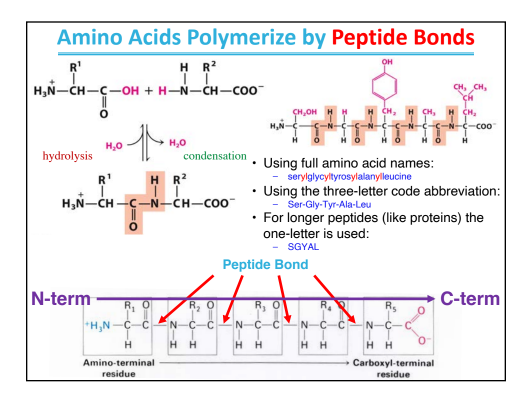
• Special (hydrophobic/hydrophilic)(2) MC family

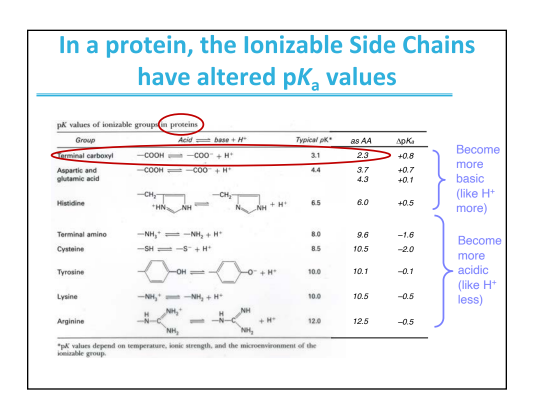
• Hydrophilic, polar (4) Qnst family

• Hydrophilic, charged (5) Dekrh family

		_	Year	% abundance i			Structure
Name	3-letter	1-letter	discovered	in proteins	NOTES	pK _a	mnemonic device
Glycine	Gly	G	1820	7	Smallest, not chiral		Н
Alanine	Ala	Α	1888	8	Foundational for ~10 other	er AA	Methyl
Valine 🗸	Val	V	1856	7	isopropyl		V-shaped
Leucine <	Leu	L	1819	10	Most abundant, domin	ant	Ala + Val
Isoleucine ✓	lle	I	1904	6	Two chiral centers (L &	& D)	Val + Me
Proline	Pro	Р	1901	5	Only imino acid (2° amine); sp bonds in proteins; is modified		5-membered ring; same # as Val; 3C
Phenylalanine	∕ Phe	F	1879	4	aromatic		Phenyl+Ala
Tyrosine	Tyr	Υ	1846	3	aromatic, can ionize; amphipathic	10.1	p-phenol+Ala
Tryptophan 🗸	Trp	W	1901	1	aromatic & fluorescent; least abundant		Indole+Ala
Methionine <	Met	M	1922	2	Most like straight-chain ali _l	ohatic	Ala+Me/ether
Cysteine	Cys	С	1899	2	can ionize; nucleophile	10.5	Ala+SH
Glutamine	Gln	Q	1883	4	Glx; gets hydrolyzed to G		Amide of Glu
Asparagine	Asn	N	1806	4	First isolated from aspara Asx; gets hydrolyzed to A	igus sp	Amide of Asp
Serine	Ser	S	1865	7	Isolated from Sericin, pola cousin of Ala	ar	hydroxyl+Ala
Threonine 🗸	Thr	Т	1935	6	Two chiral centers (L & D)		Me+Ser
Aspartate	Glu	D	1868	5	α-amino-succinate; Most acidic	3.7	Ala+carboxyl
Glutamate	Asp	E	1866	7	α-amino-glutarate	4.3	, , acciaic
Lysine 🗸	Lys	K	1889	6	Only "bis" amino acid	10.5	arepsilon-amino
Arginine /	Arg	R	1886	5	Most basic	12.5	δ -guanidino
Histidine <a>\checkmark	His	Н	1896	1	Only physiological ionization	6.0	Ala+imidazole

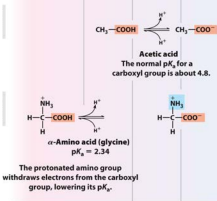






Chemical Environment Affects pK_a Values

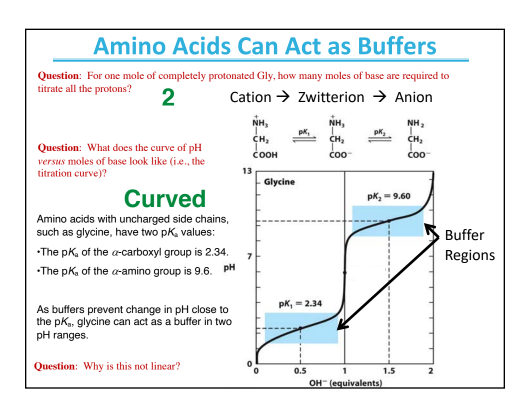
EXAMPLE: α -carboxy group is much more acidic than in carboxylic acids. PK_a 2 4 6

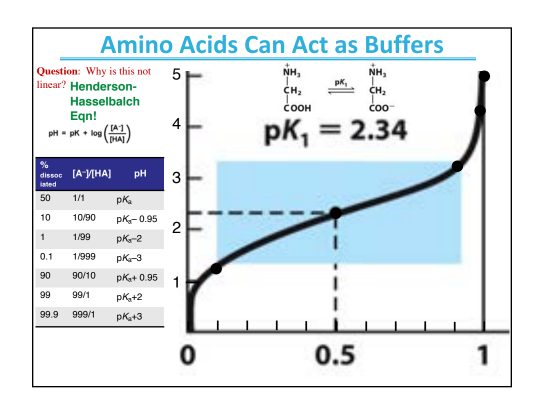


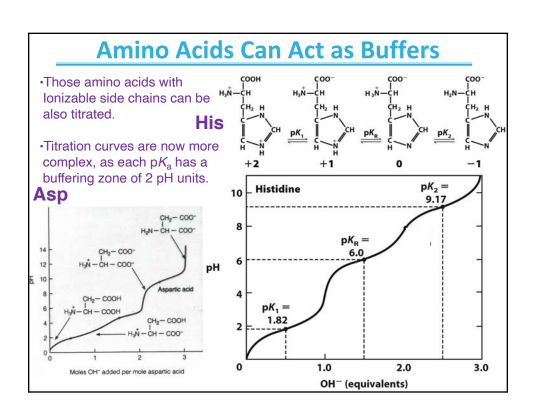
effect so that the carboxylate becomes more like acetic acid (p K_a = 3.1)

In proteins, the α -amino group is in the peptide bond and loses it's charge

IN PROTEINS, the environment can be much different than in bulk solution, and pK_a values can change by several orders of magnitude.







Amino Acids Carry a Net Charge of Zero at a Specific pH value (the pl)

- The Isoelectric Point (equivalence point, pl) is the pH value where the net charge is ZERO.
- Zwitterions predominate at pH values between the pK_a values of the amino and carboxyl groups.
- The exact value is the average of the two pK_a values forming or titrating the zwitterion.
- At the pH equal to the pI:

$$pI = \frac{pK_1 + pK_2}{2}$$

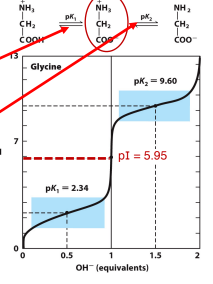
- AA is least soluble in water.
- AA does not migrate in electric field.
- AA does not bind well to other charged media/compounds

How to Calculate pl

What is the pl of glycine?

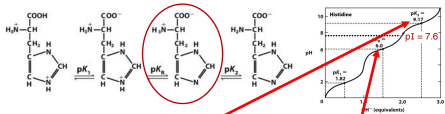
- Identify the zwitterion (species that carries a net charge of zero).
- Identify the pK_a value for the reaction that protonates the zwitterion.
- Identify the pK_a value for the reaction that titrates a proton from the zwitterion.
- Take the average of these two pK_a values.

$$pI = \frac{pK_1 + pK_2}{2}$$
$$pI = (2.3 + 9.6)/2$$
$$pI = 5.95$$



How to Calculate the pl When the Side Chain Is Ionizable His

· Identify species that carries a net zero charge.



- Identify the p K_a value for the reaction that protonates the zwitterion. For His this occurs on the R-group (p K_B).
- Identify the pK value for the reaction that titrates the next proton from the zwitterion. For His this is the α -amino group (pK_{NH}) .
- Take the average of these two pK_a values.

What is the pl of histidine?

pI = 2 pI = (6.0 + 9.2)/2pI = 7.6

How to Calculate the pl of a peptide

Estimate the pI value of the following hexapeptide:

Phe-Lys-Asp-Cys-Thr-Tyr

Step 1: Determine the total positive charge on the peptide when all acidic and basic groups are fully protonated (at low pH).

Step 2: Determine the total negative charge on the peptide when all the groups are titrated (at high pH).

Step 3: List the pK_a values of all acidic and basic groups in order from lowest (pK_{a1}) to highest.

Step 4: Calculate the pl as the average of the values for pK_a value of the proton dissociation forming a neutral species from a +1 species, and pK_a value of the proton dissociation forming a -1 species from the neutral species.

So for this peptide

Step 1: charge when fully protonated +2

Step 2: charge when fully de-protonated -4

Step 3: pK_a values are:

9.0(N-term), 10.5(Lys), 3.9(Asp), 8.4 (Cys), 10.5(Tyr), 3.5(C-term)

List from lowest to highest

pKa 3.5 3.9 8.4 9.0 10.5 10.5 Charges $+2 \leftrightarrows +1 \leftrightarrows 0 \leftrightarrows -1 \leftrightarrows -2 \leftrightarrows -3 \leftrightarrows -4$

Step 4: The pl is (3.9 + 8.4)/2 = 6.2

Proteins and their pl Values

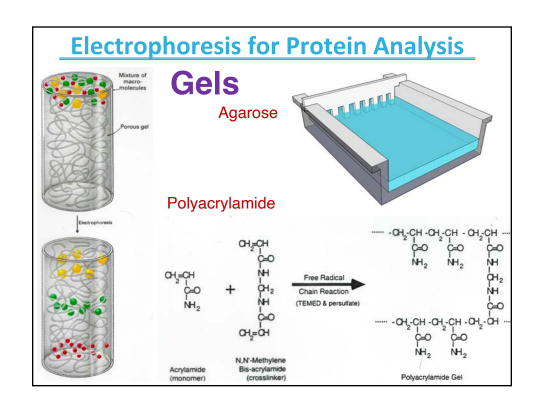
 $PH at: Cation \rightarrow Zwitterion \rightarrow Anion \\ pH at: <pI pI >pI$

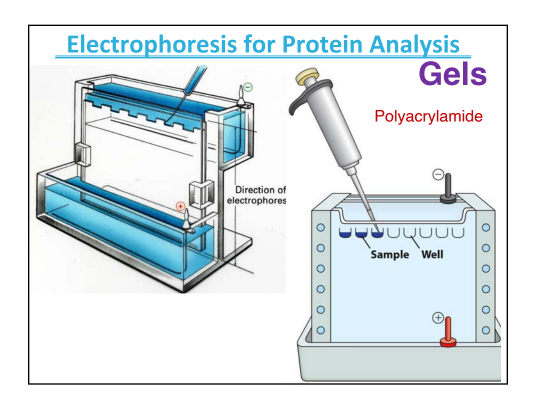
- IN GENERAL, if you take the % abundance of acidic and basic residues (Glu+Asp) and (Lys+Arg), you have ~12 and ~11%.
- So, there are slightly more acidic residues than there are basic residues.
- Half way between the most basic of these acidic residues lies the pl values for most proteins; most are below the average of 4.3 (Glu) + 10.5 (Lys) \div 2 = 7.4.
- Therefore, given that there is 1% more (Glu+Asp) than (Lys+Arg), most proteins are slightly more acidic than physiological pH.
 - ➤ That doesn't mean there are not many proteins that are very acidic (pl values <<7.4; negatively charged at pH 7.4):
 - Transcription factors
 - Pepsin
 - Ovalbumin
 - Serum albumin
 - ➤ Or, very basic (pl values >>7.4; positively charged at pH 7.4):
 - Cytochrome c
 - Lysozyme
 - Histones
 - Ribosomal proteins

- •Like AA, for proteins, at the pH equal to the pI:
 - Protein is least soluble in water, could precipitate.
 - Protein does not migrate in electric field.

Electrophoresis for Protein Analysis

- Electrophoresis is the migration of molecules in an electric field.
- Electrophoresis is one of the most commonly used analytical scale <u>separation</u> techniques
 - The electric field pulls proteins according to their charge.
- Gel electrophoresis adds a solid support in which the separation occurs. The gel matrix hinders mobility of proteins according to their size and shape.
 - The commonly used gels are either polyacrylamide (proteins) or agarose (nucleic acids).
 - separation of proteins via electrophoresis is often called polyacrylamide gel electrophoresis, or PAGE.
- For proteins to separate, they have to have a charge.

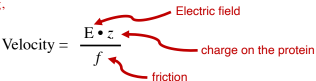




Electrophoresis for Protein Analysis

- For proteins to separate, they have to have a charge.
- The charge on a protein will depend on the pH.

But once they are moving, what does the velocity depend on?



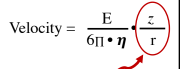
 $f \propto \text{mass}$, shape, viscosity of media

$$f = 6 \Pi \bullet \boldsymbol{\eta} \bullet \mathbf{r}$$

where:

 η = coefficient of viscosity

r = Stokes radius (mass and shape)
[this is from actual radius and specific volume (cm³/g)]



This is essentially the charge:mass ratio if all proteins are roughly the same shape (globular).

Electrophoresis for Protein Analysis

- For proteins to separate, they have to have a charge.
- pH and pl dependence:
 - o At pH near the pI, not much movement
 - At pH below the pl, proton concentration is higher, so charge becomes positive
 - At pH above the pI, protons will be titrated off, so the charge will become

Recall:

Isoelectric Focusing Takes advantage of the pl differences in Proteins for Separation

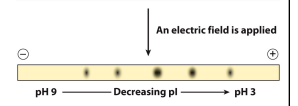
Can also be used to determine the pI values

Isoelectric Focusing

Velocity = $\frac{E}{6\pi \cdot \eta} \left(\frac{z}{r} \right)$

Ampholytes are highly charged small MW polymer with variable pI values. Due to their high z/r, they migrate rapidly setting up a buffered pH gradient. Once they reach the pH equal to their individual pl values, they STOP migrating, thus creating an immobilized pH gradient.

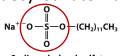
A protein sample may be applied to one end of a gel strip with an immobilized pH gradient. Or, a protein sample in a solution of ampholytes may be used to rehydrate a dehydrated gel strip.



After staining, proteins are shown to be distributed along pH gradient according to their pl values.

Sodium Dodecyl Sulfate-Polyacrylamide Gel Electrophoresis (SDS PAGE) Separates Proteins by Molecular Weight

• SDS – sodium dodecyl sulfate – a detergent



Velocity = $\frac{E}{6\Pi \cdot \eta} \left(\frac{z}{r}\right)$

Sodium dodecyl sulfate

- SDS micelles bind to proteins and facilitate unfolding.
 - SDS gives all proteins a uniformly negative charge and shape (micelle)
 - The native shape is perturbed; de-natured.
 - SDS binds proteins at a constant ratio of mass (1.4g SDS/g protein), coating them with a negative charge.
 - So much charge is added that all proteins have the same charge:mass ratio, and the rate of movement will only depend on the sieving properties of the gel: small proteins will move farther.

