

Lecture 5 (9/18/23)

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

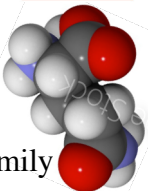

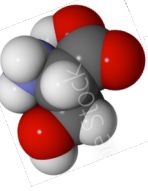
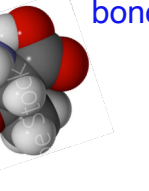
- Reading: Ch3; 76–82, 87–89
- Homework #5

NEXT

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

Amino Acids: Classification

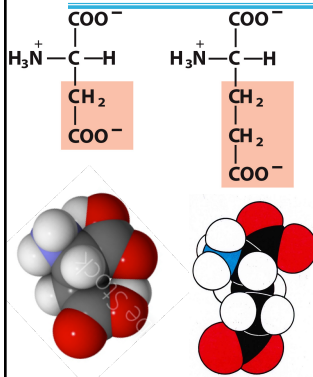
- The 20 amino acids found in proteins can be placed in five families based on the physical and chemical properties of their R groups:
 - Hydrophobic, aliphatic (6) Gavlip family
 - Hydrophobic, aromatic (3) PTT family
 - Special (hydrophobic/hydrophilic)(2) MC family
 - Hydrophilic, polar (4)
 - Hydrophilic,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

Hydrophilic, polar Amino Acids: Classification						
	$\begin{array}{c} \text{COO}^- \\ \\ \text{H}_3\text{N}^+-\text{C}-\text{H} \\ \\ \text{CH}_2 \\ \\ \text{CH}_2 \\ \\ \text{C}=\text{O} \\ \\ \text{H}_2\text{N} \end{array}$	$\begin{array}{c} \text{COO}^- \\ \\ \text{H}_3\text{N}^+-\text{C}-\text{H} \\ \\ \text{CH}_2 \\ \\ \text{C}=\text{O} \\ \\ \text{H}_2\text{N} \end{array}$	$\begin{array}{c} \text{COO}^- \\ \\ \text{H}_3\text{N}^+-\text{C}-\text{H} \\ \\ \text{CH}_2\text{OH} \end{array}$	$\begin{array}{c} \text{COO}^- \\ \\ \text{H}_3\text{N}^+-\text{C}-\text{H} \\ \\ \text{H}-\text{C}-\text{OH} \\ \\ \text{CH}_3 \end{array}$	<p style="color: #00AEEF;">These amino acids side chains can form hydrogen bonds.</p>	
						
	Gln	Amides	Asn	Ser	Alcohols	Thr
Name	3-letter	1-letter	Year discovered	% abundance in proteins	NOTES	Structure mnemonic device
Glutamine	Gln	Q	1883	4	<i>Glx; gets hydrolyzed to Glu</i>	<i>Amide of Glu</i>
Asparagine	Asn	N	1806	4	<i>First isolated from asparagus</i>	<i>Amide of Asp</i>
Serine	Ser	S	1865	7	<i>Isolated from Sericin, polar cousin of Ala</i>	<i>hydroxyl+Ala</i>
Threonine ✓	Thr	T	1935	6	<i>Two chiral centers (L & D)</i>	<i>Me+Ser</i>

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Hydrophilic, Charged Amino Acids: Classification



Name	3-letter	1-letter	Year discovered	% abundance in proteins	NOTES	pK _a	Structure mnemonic device
Aspartate	Asp	D	1868	5	<i>α</i> -amino-succinate; Most acidic	3.7	Ala+carboxyl
Glutamate	Glu	E	1866	7	<i>α</i> -amino-glutarate	4.3	Ala+acetate

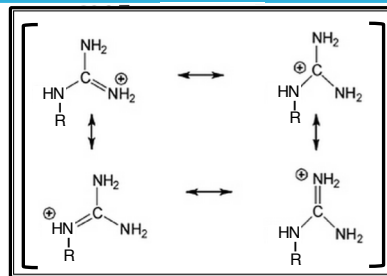
Amino Acids: Naming Dicarboxylic Acids



Dicarboxylic acids: OMSGAP

Rubric	Name (conjugate base)	Name of Acid	Structure (conjugate base)	X = (CH ₂) _z in ⁻ OOC-X-COO ⁻ ; z=?
Oh	oxalate	oxalic	⁻ OOC-COO ⁻	0
My	malate	malic	⁻ OOC-CH ₂ -COO ⁻	1
Such	succinate	succinic	⁻ OOC-CH ₂ -CH ₂ -COO ⁻	2
Good	glutarate	glutaric	⁻ OOC-CH ₂ -CH ₂ -CH ₂ -COO ⁻	3
Apple	adipate	adipic	⁻ OOC-CH ₂ -CH ₂ -CH ₂ -CH ₂ -COO ⁻	4
Pie	pimelate	pimelic	⁻ OOC-CH ₂ -CH ₂ -CH ₂ -CH ₂ -CH ₂ -CH ₂ -COO ⁻	5

Hydrophilic, Charged **Amino Acids: Classification**

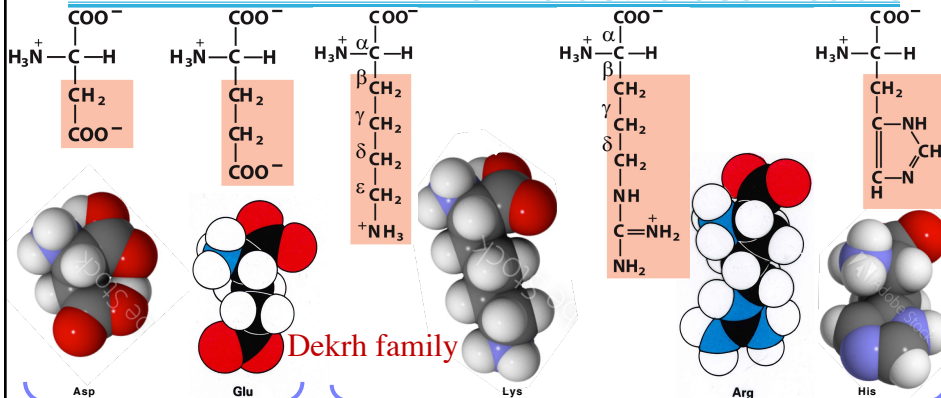


As the electrons creating the double bond become de-localized, the positive charge also becomes delocalized

α
 β
 γ
 δ

Name	3-letter	1-letter	Year discovered	% abundance in proteins	NOTES	pK_a	Structure mnemonic device
Aspartate	Asp	D	1868	5	α -amino-succinate; Most acidic	3.7	Ala+carboxyl
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Arginine ✓	Arg	R	1886	5	Most basic	12.5	δ -guanidino
Histidine ✓	His	H	1896	1	Only physiological ionization	6.0	Ala+imidazole

Hydrophilic, Charged **Amino Acids: Classification**



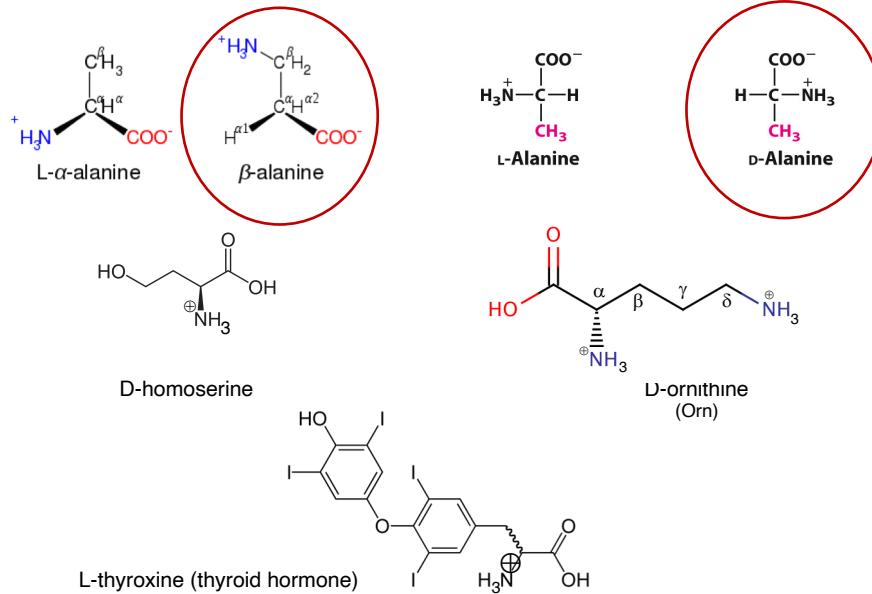
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Name	3-letter	1-letter	Year discovered	% abundance in proteins	NOTES	pK _a	Structure Mnemonic device
Glycine	Gly	G	1820	7	<i>Smallest, not chiral</i>		<i>H</i>
Alanine	Ala	A	1888	8	<i>Foundational for ~10 other AA</i>		<i>Methyl</i>
Valine ✓	Val	V	1856	7	<i>isopropyl</i>		<i>V-shaped</i>
Leucine ✓	Leu	L	1819	10	<i>Most abundant, dominant</i>		<i>Ala + Val</i>
Isoleucine ✓	Ile	I	1904	6	<i>Two chiral centers (L & D)</i>		<i>Val + Me</i>
Proline	Pro	P	1901	5	<i>Only imino acid (2° amine); special bonds in proteins; is modified by OH</i>		<i>5-membered ring; same # as Val; 3C</i>
Phenylalanine ✓	Phe	F	1879	4	<i>aromatic</i>		<i>Phenyl+Ala</i>
Tyrosine	Tyr	Y	1846	3	<i>aromatic, can ionize; amphipathic</i>	10.1	<i>p-phenol+Ala</i>
Tryptophan ✓	Trp	W	1901	1	<i>aromatic & fluorescent; least abundant</i>		<i>Indole+Ala</i>
Methionine ✓	Met	M	1922	2	<i>Most like straight-chain aliphatic</i>		<i>Ala+Me/ether</i>
Cysteine	Cys	C	1899	2	<i>can ionize; nucleophile</i>	10.5	<i>Ala+SH</i>
Glutamine	Gln	Q	1883	4	<i>Glx; gets hydrolyzed to Glu</i>		<i>Amide of Glu</i>
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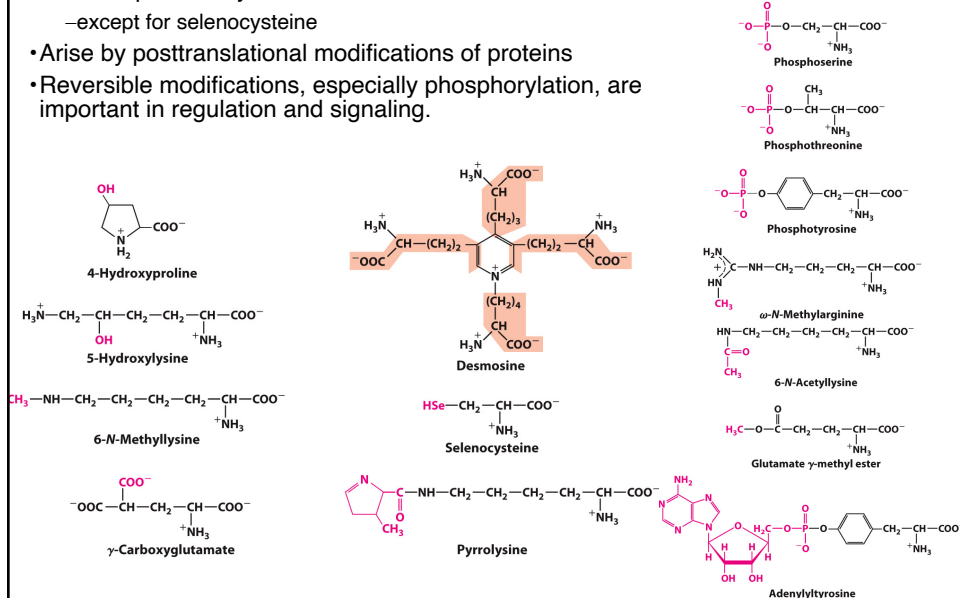
Other Cool Amino Acids



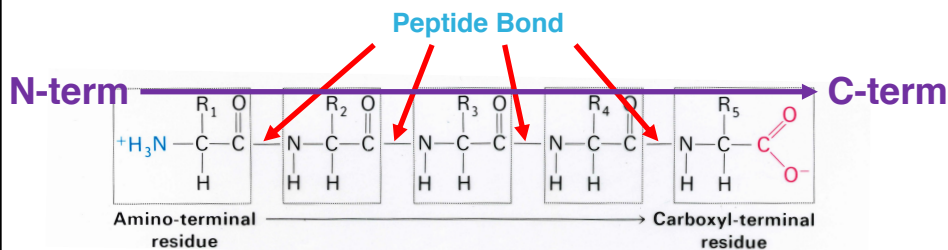
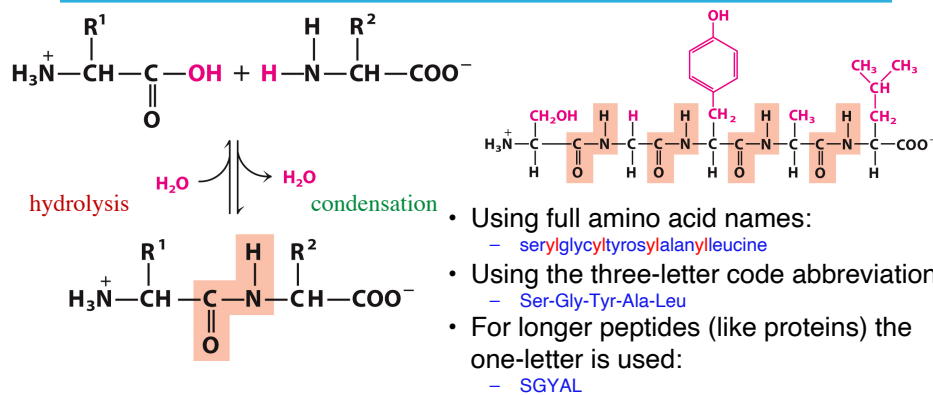
Modified Amino Acids Found in Proteins

- Not incorporated by ribosomes
 - except for selenocysteine
- Arise by posttranslational modifications of proteins
- Reversible modifications, especially phosphorylation, are important in regulation and signaling.

Reversible:



Amino Acids Polymerize by Peptide Bonds



In a protein, the Ionizable Side Chains have altered pK_a values

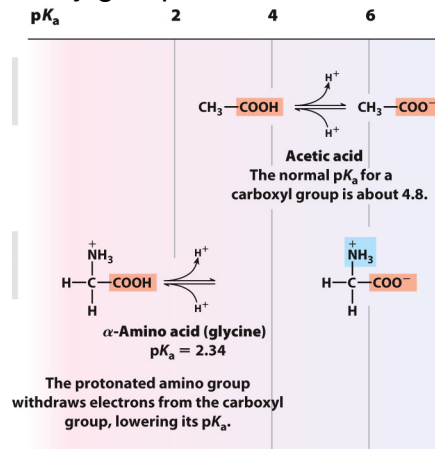
pK values of ionizable groups in proteins

Group	Acid \rightleftharpoons base + H ⁺	Typical pK*	as AA	Δ pK _a
Terminal carboxyl	$-\text{COOH} \rightleftharpoons -\text{COO}^- + \text{H}^+$	3.1	2.3	+0.8
Aspartic and glutamic acid	$-\text{COOH} \rightleftharpoons -\text{COO}^- + \text{H}^+$	4.4	3.7 4.3	+0.7 +0.1
Histidine	$-\text{CH}_2-\text{C}_5\text{H}_4\text{N}^+ \rightleftharpoons -\text{CH}_2-\text{C}_5\text{H}_4\text{N} + \text{H}^+$	6.5	6.0	+0.5
Terminal amino	$-\text{NH}_3^+ \rightleftharpoons -\text{NH}_2 + \text{H}^+$	8.0	9.6	-1.6
Cysteine	$-\text{SH} \rightleftharpoons -\text{S}^- + \text{H}^+$	8.5	10.5	-2.0
Tyrosine	$-\text{C}_6\text{H}_4\text{OH} \rightleftharpoons -\text{C}_6\text{H}_4\text{O}^- + \text{H}^+$	10.0	10.1	-0.1
Lysine	$-\text{NH}_3^+ \rightleftharpoons -\text{NH}_2 + \text{H}^+$	10.0	10.5	-0.5
Arginine	$-\text{N}(\text{H})-\text{C}(=\text{NH}_2^+)-\text{NH}_2 \rightleftharpoons -\text{N}(\text{H})-\text{C}(=\text{NH})-\text{NH}_2 + \text{H}^+$	12.0	12.5	-0.5

*pK values depend on temperature, ionic strength, and the microenvironment of the ionizable group.

Chemical Environment Affects pK_a Values

EXAMPLE: α -carboxy group is much more acidic than in carboxylic acids.



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 Can Act as Buffers

Question: For one mole of completely protonated Gly, how many moles of base are required to titrate all the protons?

2

Cation \rightarrow Zwitterion \rightarrow Anion

Question: What does the curve of pH versus moles of base look like (i.e., the titration curve)?

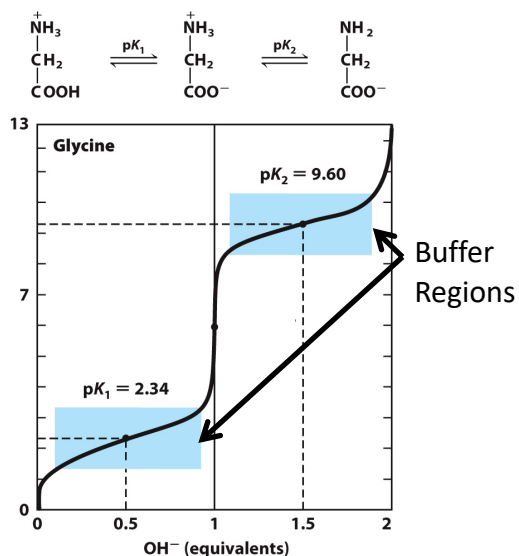
Curved

Amino acids with uncharged side chains, such as glycine, have two pK_a values:

- The pK_a of the α -carboxyl group is 2.34.
- The pK_a of the α -amino group is 9.6.

As buffers prevent change in pH close to the pK_a , glycine can act as a buffer in two pH ranges.

Question: Why is this not linear?



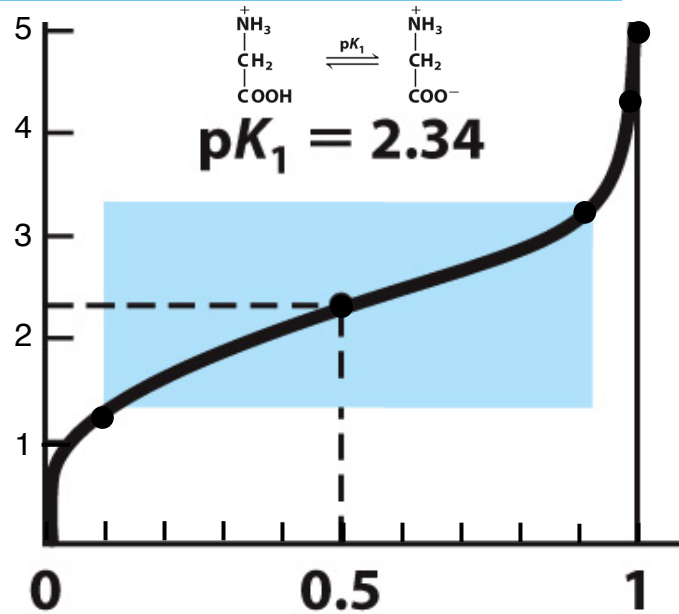
Amino Acids Can Act as Buffers

Question: Why is this not linear?

Henderson-Hasselbalch Eqn!

$$\text{pH} = \text{pK} + \log \left(\frac{[\text{A}^-]}{[\text{HA}]}\right)$$

% dissociated	[A ⁻]/[HA]	pH
50	1/1	pK _a
10	10/90	pK _a -0.95
1	1/99	pK _a -2
0.1	1/999	pK _a -3
90	90/10	pK _a +0.95
99	99/1	pK _a +2
99.9	999/1	pK _a +3



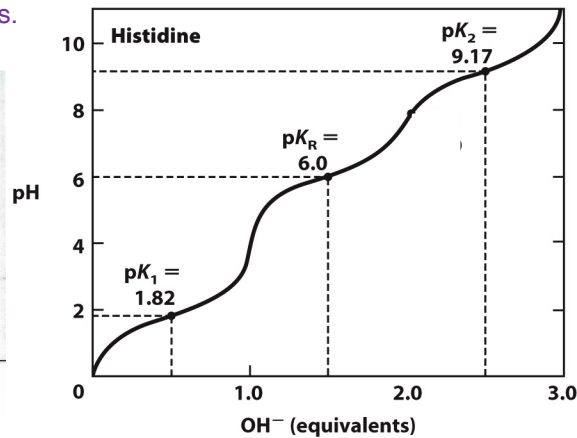
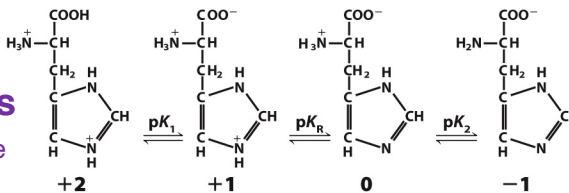
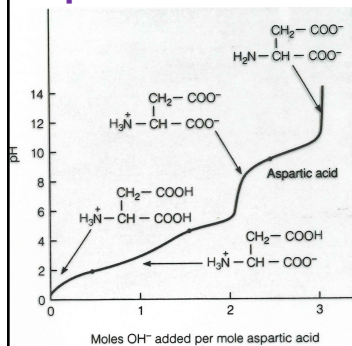
Amino Acids Can Act as Buffers

• Those amino acids with ionizable side chains can be also titrated.

His

• Titration curves are now more complex, as each pK_a has a buffering zone of 2 pH units.

Asp



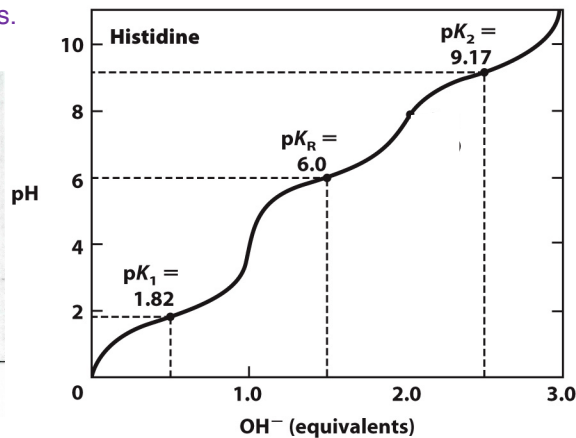
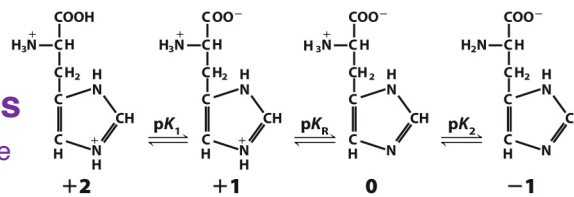
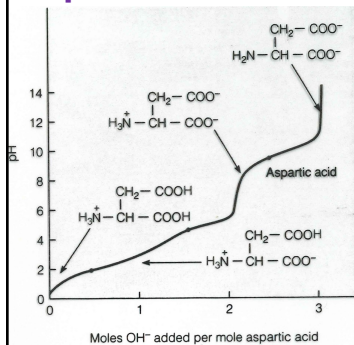
Amino Acids Can Act as Buffers

- Those amino acids with ionizable side chains can be also titrated.

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Asp



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

- The **Isoelectric Point** (equivalence point, **pI**) 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:
 - AA is least soluble in water.
 - AA does not migrate in electric field.
 - AA does not bind well to other charged media/compounds

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

How to Calculate pI

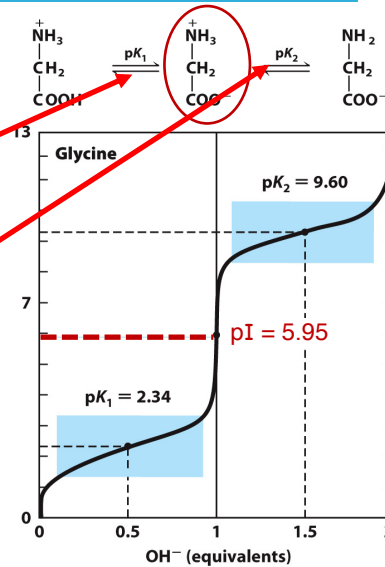
What is the pI of glycine?

- 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.

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

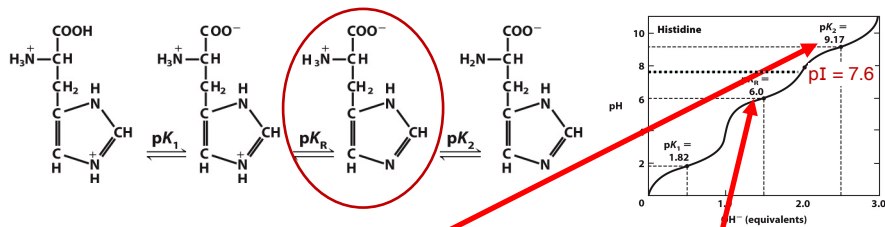
$$pI = (2.3 + 9.6)/2$$

$$pI = 5.95$$



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

- Identify species that carries a net zero charge.



- Identify the pK_a value for the reaction that protonates the zwitterion. For His this occurs on the R-group (pK_R).
- Identify the pK_a value for the reaction that titrates the next proton from the zwitterion. For His this is the α -amino group (pK_{NH_2}).
- Take the average of these two pK_a values.

What is the pI of histidine?

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

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

$$pI = 7.6$$

How to Calculate the pI 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 pI 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	↔ +1	↔ 0	↔ -1	↔ -2	↔ -3	↔ -4

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

Proteins and their pI Values

	Cation →	Zwitterion	→ 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 pI values for most proteins; most are below the average of $4.3 \text{ (Glu)} + 10.5 \text{ (Lys)} \div 2 = 7.4$.
- 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 (pI values $\ll 7.4$; negatively charged at pH 7.4):
 - Transcription factors
 - Pepsin
 - Ovalbumin
 - Serum albumin
 - Or, very basic (pI values $\gg 7.4$; 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.

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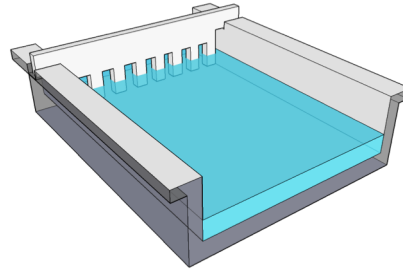
Electrophoresis for Protein Analysis

- **Electrophoresis** is the migration of molecules in an electric field.
- **Electrophoresis** is one of the most commonly used analytical scale **separation** 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.**

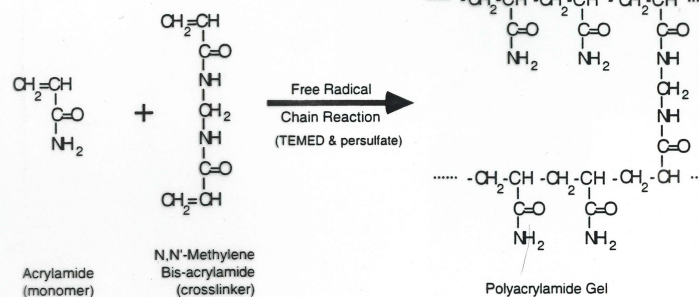
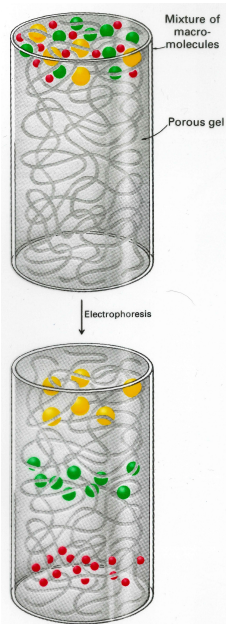
Electrophoresis for Protein Analysis

Gels

Agarose



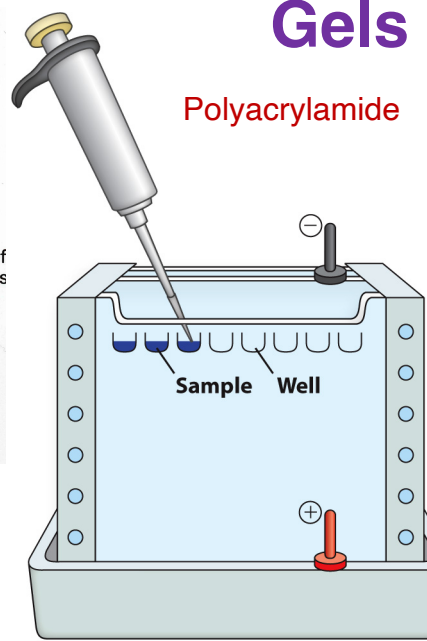
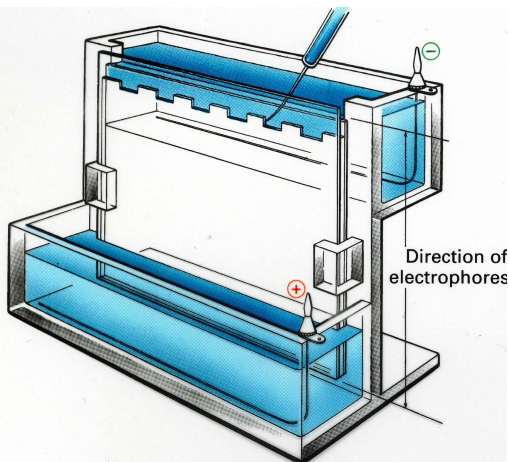
Polyacrylamide



Electrophoresis for Protein Analysis

Gels

Polyacrylamide



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?

$$\text{Velocity} = \frac{E \cdot z}{f}$$

Electric field
charge on the protein
friction

$f \propto$ mass, shape, viscosity of media

$$f = 6\pi \cdot \eta \cdot r$$

where:

η = coefficient of viscosity

r = Stokes radius (mass and shape)

[this is from actual radius and specific volume (cm^3/g)]

$$\text{Velocity} = \frac{E}{6\pi \cdot \eta} \cdot \frac{z}{r}$$

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 pI dependence:
 - At pH near the pI, not much movement
 - At pH below the pI, proton concentration is higher, so charge becomes positive \oplus
 - At pH above the pI, protons will be titrated off, so the charge will become \ominus

Recall:

	Cation	→	Zwitterion	→	Anion
pH at:	<pI		pI		>pI

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

Can also be used to determine the pI values

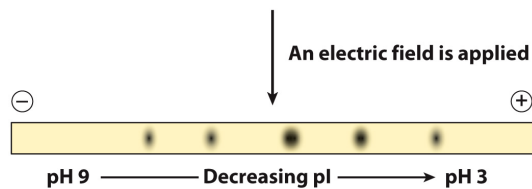
Isoelectric Focusing

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

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 pI values, they STOP migrating, thus creating an **immobilized pH gradient**.



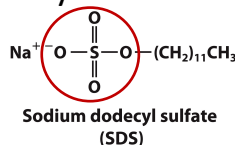
A protein sample may be applied to one end of a gel strip with an immobilized pH gradient. Or, a protein sample in a solution of ampholytes may be used to rehydrate a dehydrated gel strip.



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

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

- SDS – sodium dodecyl sulfate – a detergent

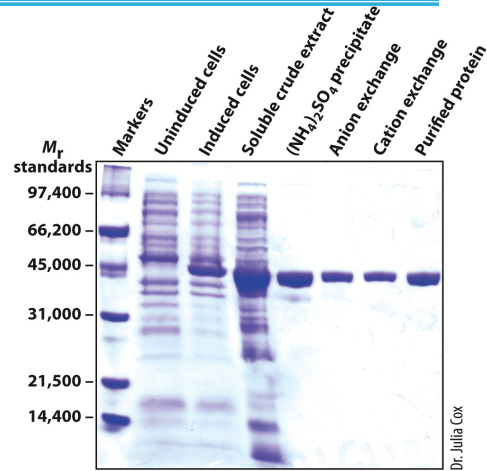
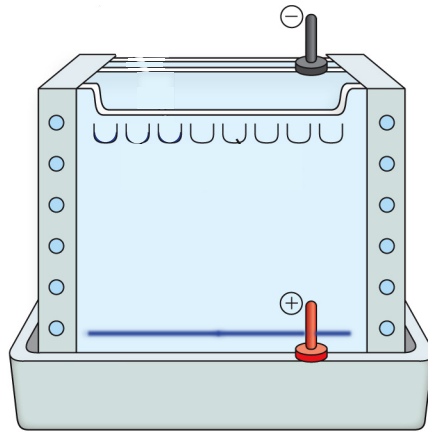


$$\text{Velocity} = \frac{E}{6\pi \cdot \eta} \cdot \left(\frac{z}{r} \right)$$

- 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.

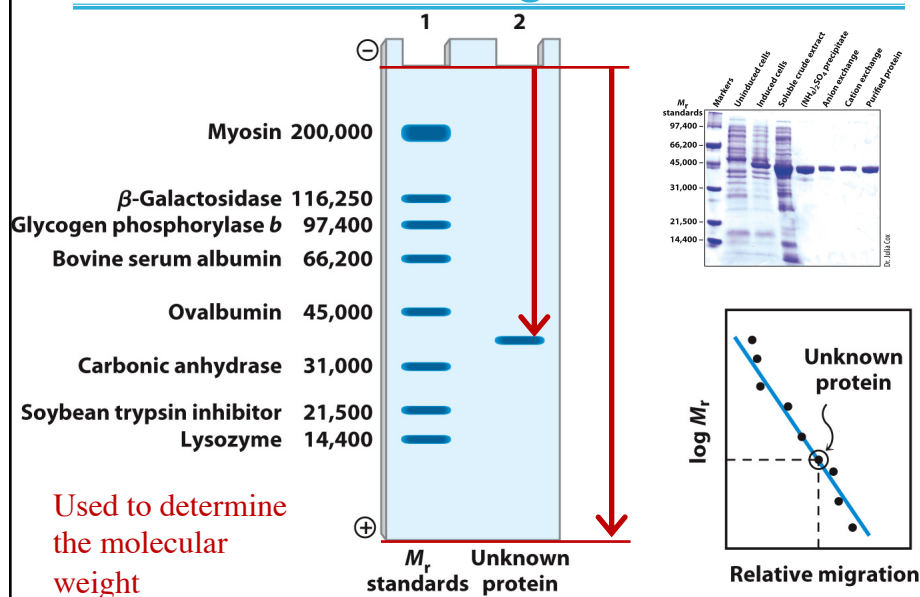
SDS PAGE Separates Proteins by Molecular Weight

SDS-PAGE

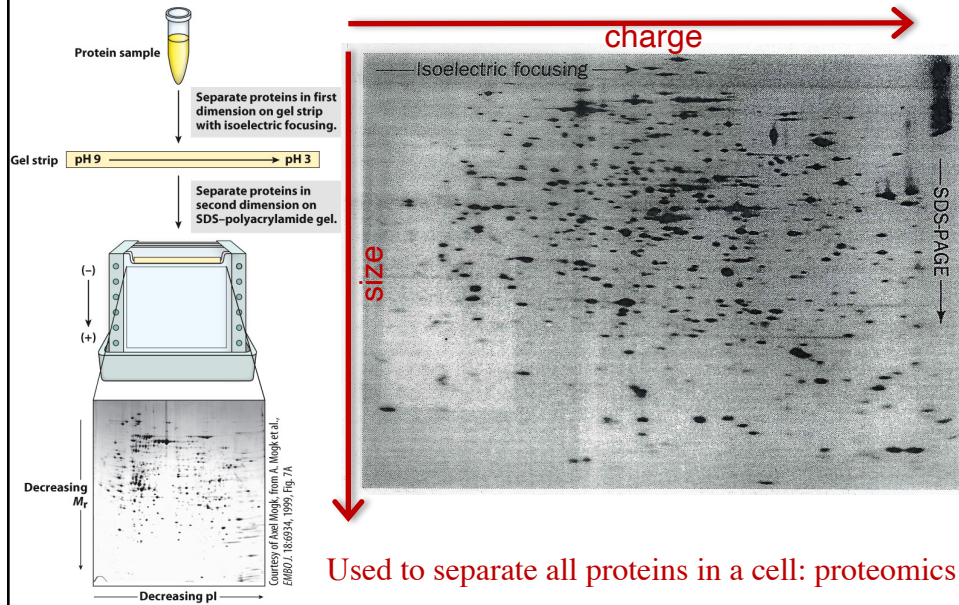


Used to detect purity or purification

SDS PAGE Can Be Used to Calculate the Molecular Weight of a Protein



SDS-PAGE + Isoelectric Focusing Can Separate nearly ALL the Proteins in *E. coli*









Protein Purification

Protein Purification

*Proteins are separated from each other (along with other macromolecules) due to the vast variability they have. The **basis** of the separation can be put into 4 categories:*

- Size, shape, density } Hydrodynamic properties
- Charge } Chemical properties
- Solubility } Biological properties
- Binding characteristics }

Protein Purification Procedures

Basis	Procedure	Covered
Hydrodynamics (size, shape, density)	Gel filtration <u>Chromatography</u>	Lab 
	SDS-PAGE	Lab 
	Centrifugation	Lab 
Charge	Ion exchange <u>Chromatography</u>	
	Isoelectric focusing	
	Native electrophoresis	Lab 
Solubility	Salting out	Lab 
	Organic extraction	
	Hydrophobic interaction	
	Chromatography	
Binding Specificity	Affinity <u>Chromatography</u>	Lab 

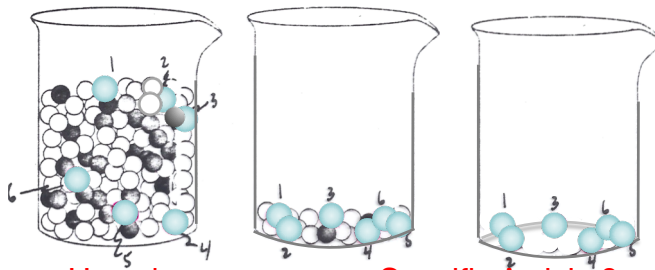
Before you can separate “your favorite protein (YFP)” from all the thousands of others, you need a way to “see” it..... An Assay!

- enzymes → activity assay
- binding proteins → binding assay
- other proteins → immunoblot or ELISA

How do you monitor the purification?

• Specific Activity =
$$\frac{\text{Activity of YFP}}{\text{Total protein}}$$

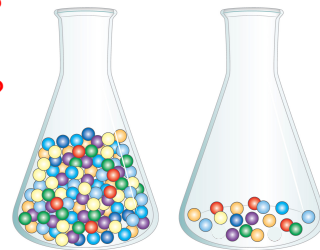
• Specific Activity =
$$\frac{\text{Activity of YFP}}{\text{Total protein}}$$



How do you measure Specific Activity?

Figure 6-3 Activity versus specific activity. The difference between these two terms can be illustrated by considering two jars of marbles. The jars contain the same number of blue marbles (representing an unknown protein), but different amounts of marbles of other colors. If the marbles are taken to represent proteins, both jars contain the same activity of the protein represented by the blue marbles. The second jar, however, has the higher specific activity because here the blue marbles represent a much higher fraction of the total.

What is the Yield?
(of red proteins?)



From [protein] (mg/mL) What is the specific activity? From [activity] (U/mL)

Purification of a hypothetical protein

Procedure or step	Fraction volume (mL)	Total protein (mg)	Activity (units)	Specific activity (units/mg)
1. Crude cellular extract	1,400	10,000	100,000	10
2. Precipitation	280	3,000	96,000	32
3. Ion-exchange chromatography	90	400	80,000	200
4. Size-exclusion chromatography	80	100	60,000	600
5. Affinity chromatography	6	3	45,000	15,000

* All data represent the status of the sample *after* the procedure indicated in the first column has been carried out.

- If protein was "pure" after step #5, what would the Specific Activity be after you performed a step #6?
- What is the Yield?