



Biochemistry

Title = Amino acid catabolism

Lec no =6

Done By Baraa Safi

Amino Acid Catabolism & Synthesis

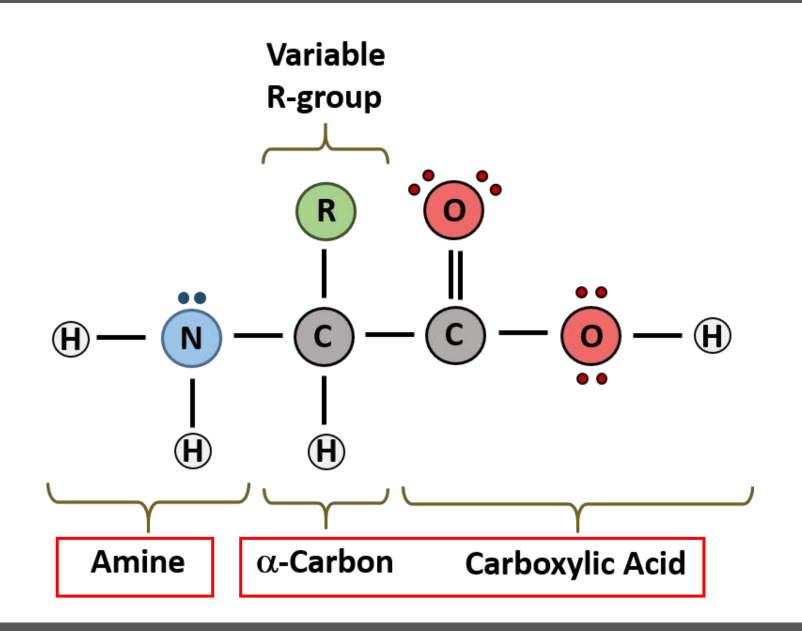
Nebras Melhem

برح نسمی الیوم عن کلیسرال (Amino acid) بیکروا ولیشی مینالد (Amino acid) بیکروا ولیشی (Mon-essential amino acid in our body) بیکروا ولیشی

Inroduction

• Unlike fats and carbohydrates, amino acids are **not stored by the body**.

- Amino acids must be obtained from the diet, synthesized de novo, or produced from the degradation of body protein.
- Any amino acids in excess of the biosynthetic needs of the cell are rapidly degraded.

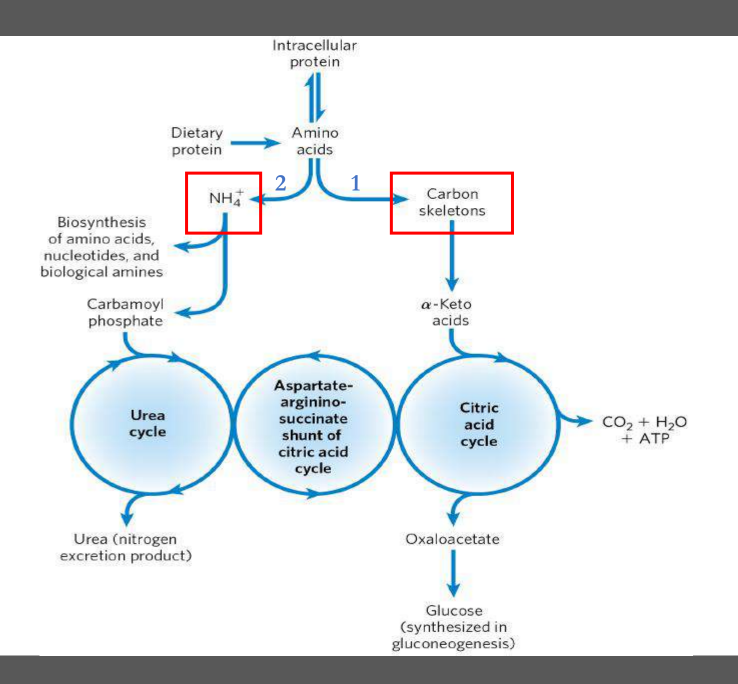


بعدا حضاً إنها بدي أكر الـ (Amino acid) فالله بحدهو الر مهم (Amine)وبغلالباقي يتعلى فالله بفله بعير

. ولا العاني (intermediate) الموند (d-Ketoncid). (ammonia) عداء في تناعد (amine group) والرا الموند المعانية في ا

(Am monia) سيئة الجسم لذا فأنارح اتخلف منها • فعماضة اليعم بدي اعمق كين بدي أكسر ال (a-Keto acid)

وال (Amonia)عنه بعم





Amino acids undergo oxidative degradation in three different metabolic circumstances:

- 1. Amino acids released during normal protein turnover are **not needed** for new protein synthesis.
- 2. Ingested amino acids **exceed the body's needs** for protein synthesis.
- 3. Cellular proteins are used as **fuel** because carbohydrates are either unavailable or not properly utilized due to **starvation** or **uncontrolled diabetes mellitus**.

First phase of AA catabolism

- Involves the **removal** of the α-amino groups, forming ammonia and the corresponding α-keto acids, the carbon skeletons of amino acids.
- A portion of the free ammonia is excreted in the urine, but most is used in the synthesis of urea which is quantitatively the most important route for disposing of nitrogen from the body.
- (next lecture)

Second phase of AA catabolism

- The carbon skeletons of the α-keto acids are converted to common intermediates of energy-producing metabolic pathways.
- These compounds can be metabolized to carbon dioxide (CO2) and water (H2O), glucose, fatty acids, or ketone bodies by the central pathways of metabolism.

3	Glucogenic	Glucogenic and Ketogenic	Ketogenic
Nonessential	Alanine/ Arginine Asparagine Aspartate Cysteine Glutamate Glutamine Glycine Proline Serine	Tyrosine	
Essential	Histidine Methionine Threonine Valine	Threenine Isoleucine Phenyl- alanine Tryptophan	Leucine Lysine

Principles of AA catabolism

• Four amino acids — alanine, glutamate, glutamine, and aspartate — play key roles in the transport and distribution of amino groups.

• All are present in relatively **high concentrations** in one or many tissues.

• All are readily converted to key citric acid cycle intermediates.



الى الله الما توكا ؟ (4) معمين كثير في عملية الـ (AA Catabo lism) طيبه ليشره وله معن والباقوكا ؟ سؤال من الكتورة ؟

Glutamate is the most abundant amino acid that can be synthesised in the body by awide variety of tissues and because it is non-essential and it synthesis various amino acids

ر المرا ((A)) من مؤلة عنه نفي التكسير (عربفي الما بفي التكسير (عربفي الما بنفي التكسير (عربفي الما بنفي التكسير (عربفي التكسير (عربفي التكسير)

because the process of catabolism requires specific enzymes to break down the complex structures of amino acids into simpler molecules. These enzymes are essential for the chemical reactions involved in the catabolic pathway. Without the appropriate enzymes, amino acids would remain relatively stable and cannot be efficiently broken down for energy production or other metabolic processes.

Principles of AA catabolism

• Metabolic pathways are not distinct. The various pathways for amino acid catabolism are elaborately intertwined with other catabolic and anabolic pathways.

• Free ammonia is toxic. Excess amino groups must be safely excreted. The urea cycle serves this purpose (next lecture).

Principles of AA catabolism

- Most amino acids are metabolized in the liver.
- Some of the ammonia generated in this process is recycled and used in **biosynthetic pathways** where glutamine, يعنى هلالوعنى عقده والمعلم والمعنى على المعنى الم

صن عنول الرا Am monia)

- Excess amino groups are either excreted directly or converted to urea or uric acid for excretion.
- Most excess ammonia generated in other (extrahepatic) tissues travels to the liver for conversion to urea.



• The first step in the catabolism of most L-amino acids, once they have reached the liver, is **removal of the α-amino groups**, promoted by enzymes called **aminotransferases** or **transaminases**.

• All amino acids can be transaminated, except lysine, threonine, proline
& hydroxyproline. (bansamination) ما بعد المحالية المعالية المع

Aminotransferase reactions

It is the transfer of an amino group from an α amino acid to an α keto acid, which is always α ketoglutarate (NH₂ acceptor).

- **Enzyme**: Aminotransferase (Transaminase)
- Coenzyme: Pyridoxal Phosphate (PLP) derived from vitamin B6, acts as intermediate-carrier of the amino group.
- Site: Present in the mitochondria and cytosol of all tissues, especially the liver.

Properties of aminotransferase reactions

- The reaction is reversible.
- Several aminotransferases exist, each specific for one amino acid and the corresponding α keto acid.
- An aminotransferase is called after the amino acid to which it is specific.
- Two important aminotransferases are those that catalyze aminotransferase reactions involving **aspartic acid** and **alanine**, they are respectively called, aspartate aminotransferase (AST) and alanine aminotransferase (ALT).
- α-Ketoglutarate acts as an amino group acceptor in most reactions, forming glutamate.
- It involves <u>neither the uptake nor the release of free ammonia</u>, the ammonia removed is being transferred to another amino acid.

(Amine groupe) المع الله بنقل الـ (PIP) X

Funneling Amino Groups to Glutamate

The first step in catabolism of most amino acids is the transfer of α -amino group to α -ketoglutarate, producing an α -keto acid (derived from the original amino acid) and glutamate.

Glutamate produced by transamination can be oxidatively deaminated or used as an armino group donor in the synthesis of nonessential amino acids.

C=0 CH_2 CH_2 CH_2 COO- COO^{-} α -Ketoglutarate L-Glutamate PLP aminotransferase α -Keto acid L-Amino acid

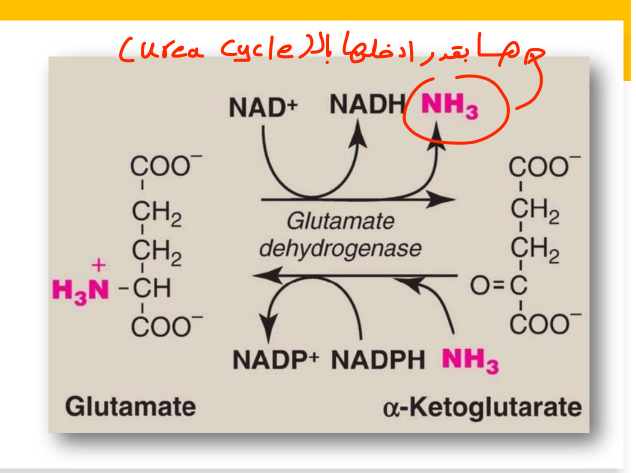
موجود كثير عندي منه

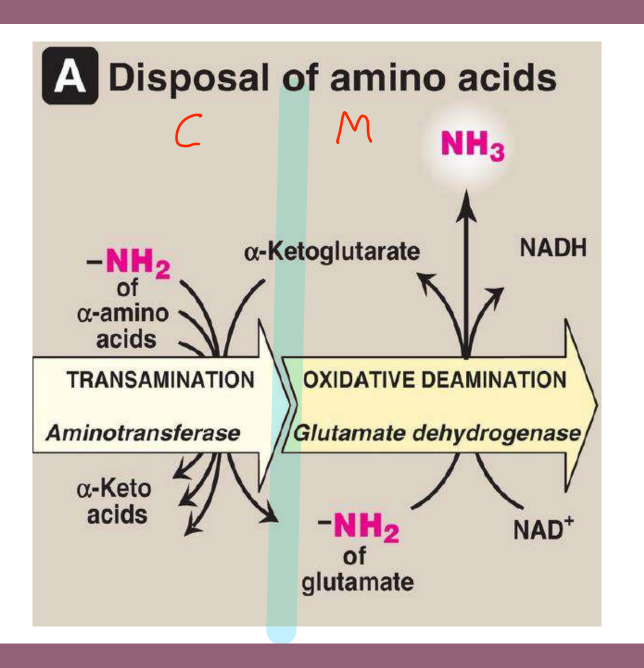
COO-

(mitochondria) عد ث في الر (cyto plasm) الر (transamination) المراد المعان الروبية (mitochondria) عد الروبية المرابية ال

Oxidative deamination: Amino group removal ((ابسیل (میساله (۲۰۰۰) (Amid group) وبیماله (۲۰۰۰)

- In hepatocytes, glutamate is transported from the cytosol into mitochondria.
- Here, it undergoes oxidative deamination catalysed by **L**-glutamate dehydrogenase to produce **NH**⁺₄ and α-ketoglutarate.
- These reversible reactions occur primarily in the liver and kidney.





Biological Significance of Transamination

1. First step of catabolism

• In this first step, **ammonia** is removed (or actually transferred as will be removed in deamination) and the carbon skeleton of the amino acid enters into catabolic pathway.

2. Synthesis of nonessential amino acids

- By means of transamination, all
- e.g. pyruvate → alanine; oxaloacetate → aspartic acid
- Those amino acids, which cannot be synthesized in this manner, are therefore essential; they should be made available in the food

Biological Significance of Transamination

3. Interconversion of amino acids

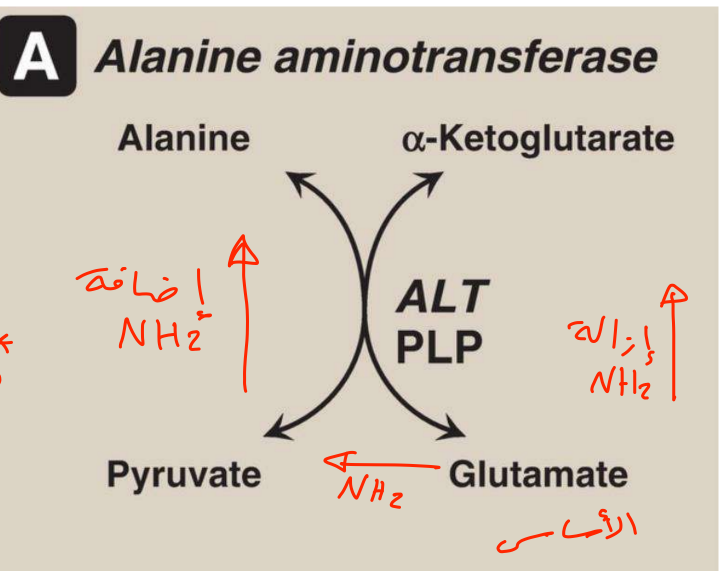
- If amino acid number 1 is high and number 2 is low, the amino group from number 1 may be transferred to alpha keto acid to give amino acid number 2; equalizing the quantity for both amino acids. Cnon-essential AA)
- This is called equalization of quantities of nonessential amino acids.

(Amine groupe) اهواللي بنقل الـ(PIP) *

The most active transaminases are:

1. Alanine transaminase (ALT)

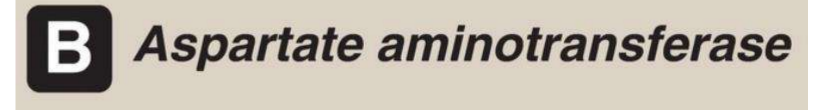
منه (Glutamate) منه (NH2) بتروح علی (Alanin) بتروح علی (pyruvate) فیتول دار (Alanin) بعید (Glutamate) بعید (Glutamate) عندطریقه (PIP) بوجود (PIP)

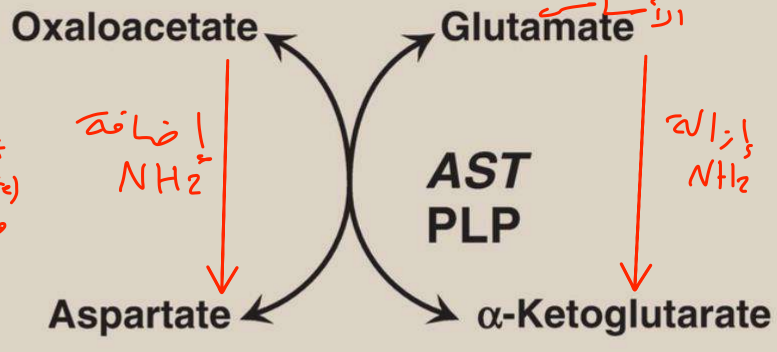


The most active transaminases are:

2. Aspartate transaminase (AST)

منه (Glutamate) منه (NH2) * (Aspartate) خبتصول له (Oxaloacetate) (اله اله Glutamate) بعب (Glutamate) واله (PIP) بعب (AST) بعبد (PIP)

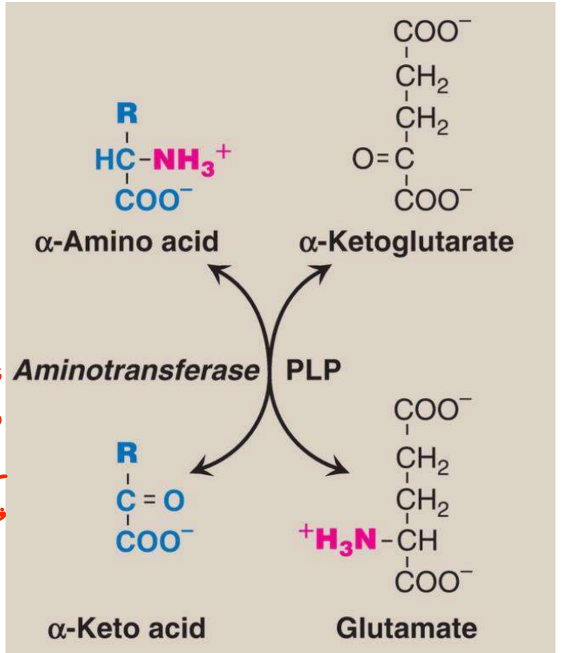




The most active transaminases are:

3. Glutamate transaminase

بكونه أنا بحكي (Glutamate trans aminuse) بككونه أنا بعك في المحكون الأسام Aminotransferase في المحلي المحلي المحلي (Glutamate) في المحلي المحلي (Amino Acid) في المحلي وبدورعل (Amino Acid) .



Clinical Significance of Transaminases

- Transaminases are intracellular enzymes; their levels in blood are **low** under **normal** conditions
- * ALT and AST are markers of liver injury
 - Elevated plasma levels of aminotransferases indicate damage to cells rich in these enzymes. For example, physical trauma or a disease process can cause cell lysis, resulting in release of intracellular enzymes into the blood.
- AST and ALT are induced by glucocorticoids which favors gluconeogenesis

بح صول بكونوا موجودي بكثرة بإلكبر وعنه عدوث الجرح فإن الكلالي تتعلل مطلقة مستوا حا منه عنه الا نديمات إلى جرى الم فرعه (Markes) لل (liver injury)

سوال موجدد شرح عنه النقطة رح يدغل بالإستعانه

(AST /ALT) أخرجه في عبد الوصول إلى (glucose) وكينه الله عند طريقه تكيير ال (AA) وال (AST /ALT)

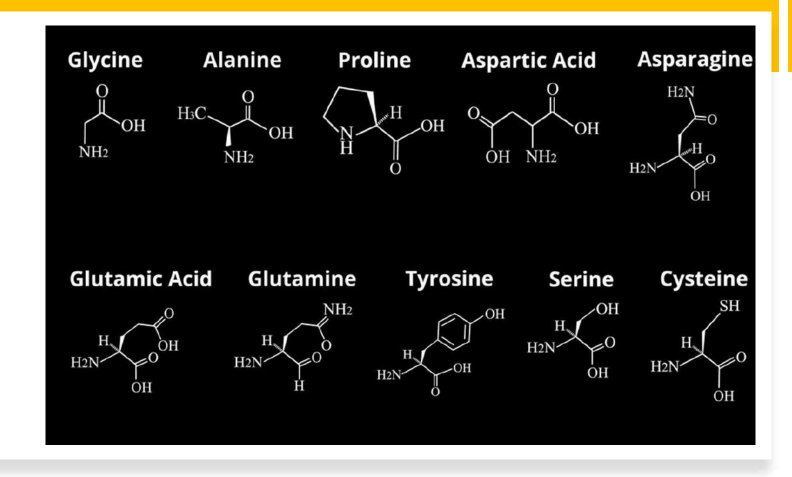
ب عدو الى ال (catabolism) فبالتالي على محونه عندي (gluconeogenesir) ويذيدوا بعفنوا (gluconeogenesir) وطاد التعفيذ

رواه cose) باستنام (intermediate) (ASTIALT) واستنام (catabolism of AA) بالمتناع (gla cose)

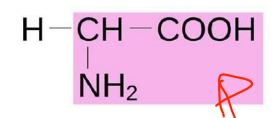
فأنامكا أحفنر ال(ata bolism)) بحنز صاي ال (Enzymes)

Synthesis of non-essential amino acids

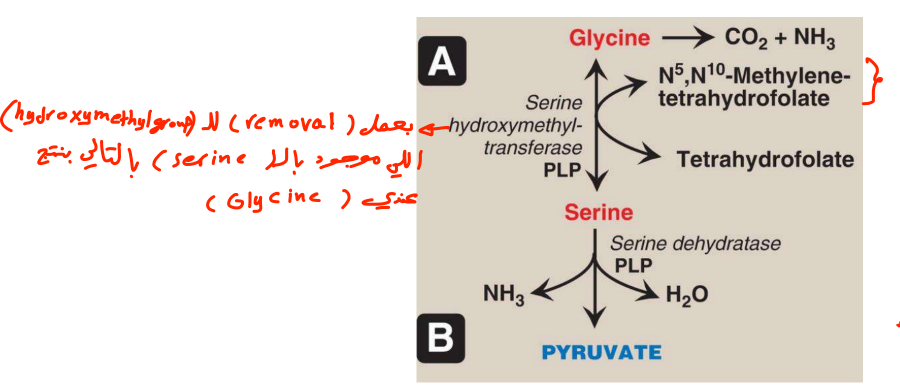
Non-essential amino acids are formed in the body in enough amounts for adults



Glycine

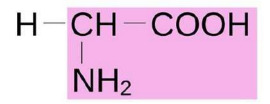


This amino acid is **synthesized** from **serine** by removal of a hydroxymethyl group, also by **serine** hydroxymethyltransferase.



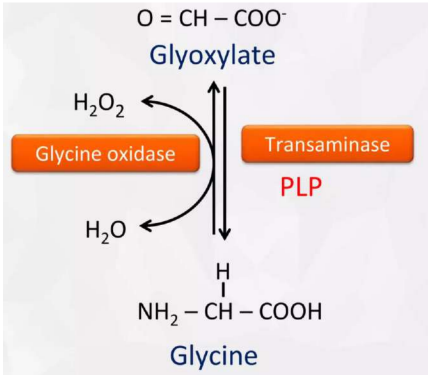
ساعد نصدا التفاعلات

Glycine



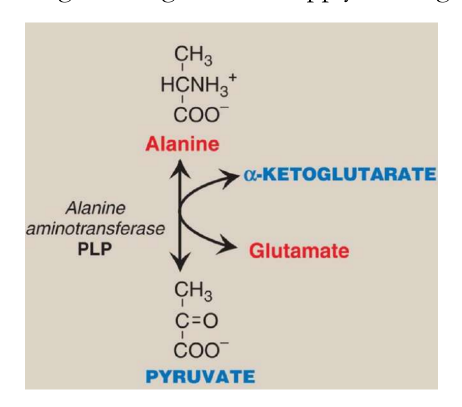
(Aminotransferase) Nie a (intermediate found in the body)

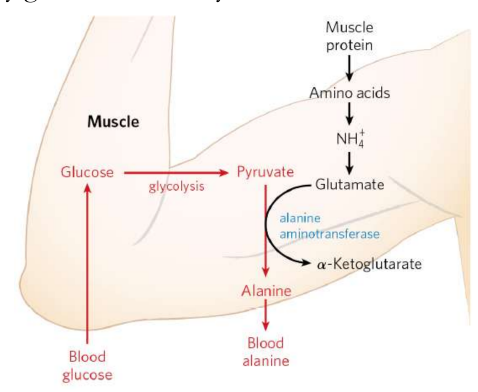
- Glycine can be deaminated to glyoxylate, which can be oxidized to oxalate or transaminated to glycine.
- Deficiency of the transaminase in liver peroxisomes causes **overproduction of oxalate**, the formation of oxalate stones, and kidney damage.

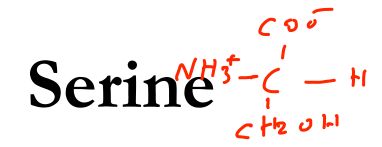


Alanine

- This amino acid loses its amino group by transamination to form pyruvate.
- It is released from the muscles during starvation and changed to glucose in the liver by gluconeogenesis to supply blood glucose by glucose alanine cycle.



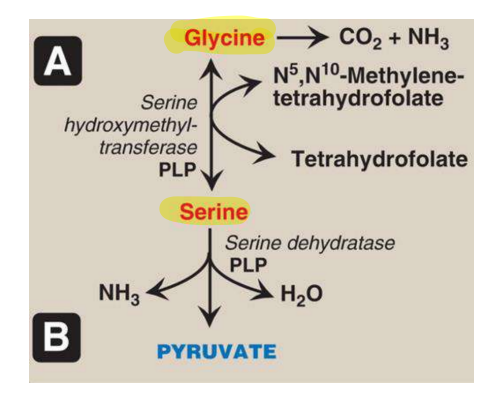




1. Formed from glycine through transfer of a hydroxymethyl group by serine hydroxymethyl transferase.

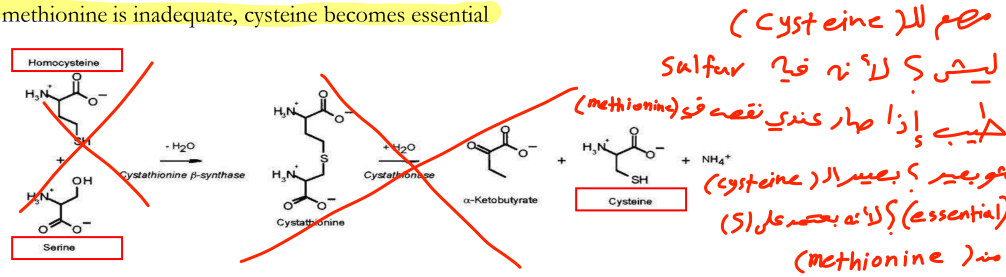
Formed from 3 phosphoglycerate that is oxidized to 3-phosphopyruvate, and then

transaminated to 3 phosphoserine.



Cysteine Cysteine معلوة منه العربة

- Y• It is formed from **serine** (provides the carbon skeleton) and **homocysteine** (provides the thiol group)
 - Homocysteine is formed from methionine
 - Dessential • Part of the dietary requirements for methionine is to provide sulfur for the biosynthesis of cysteine
 - If dietary methionine is inadequate, cysteine becomes essential



Importance of Cysteine

- 1. Synthesis of glutathione which is:
 - Powerful antioxidant
 - Used for removal of many toxic compounds
 - Acts as coenzyme for many enzymes
- 2. Synthesis of **thioethanolamine** which enters in the formation of:
 - phosphopantetheine of acyl carrier protein present in fatty acid synthase multienzyme complex coenzyme A
- 3. Synthesis of cystine; important in tertiary structure of proteins.
- 4. Cysteine desulfuration yields pyruvate (glucogenic).

Aspartic acid

NH3+- C-H
eHz

• It is formed from **oxaloacetic acid** by transamination, and also from asparagine by asparaginase enzyme.

• Importance:

- 1. urea formation
- 2. purine synthesis
- 3. pyrimidine synthesis

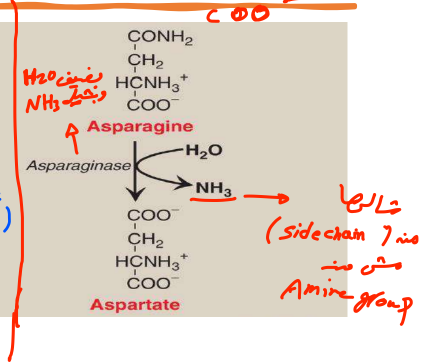


من (Glutamate) من (NH2)*

(Aspartate) المحتب (Oxaloacetate

(ه-الادtoglutarate) بعيد (Glutamate)

(واله (PIP) بعيد (AST) بعيد المحتبة (AST)



Amino acid α-Keto acid

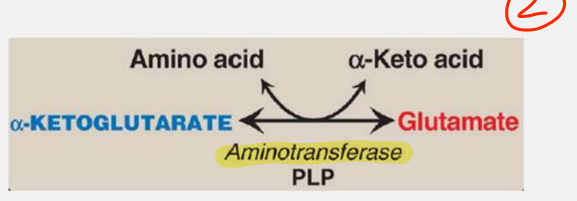
OXALOACETATE

Aminotransferase
PLP

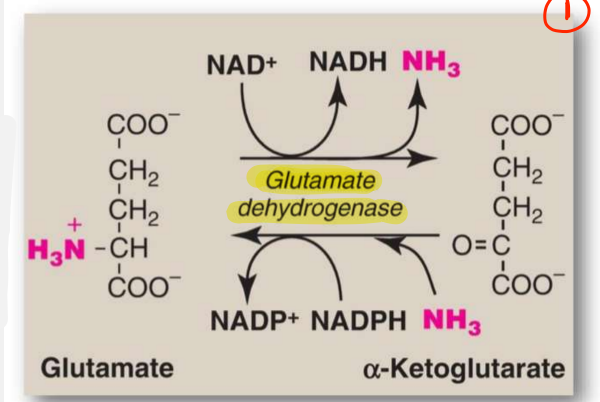
Glutamate

• Glutamate is unusual in that it can be synthesized by reversal of oxidative deamination, catalyzed by glutamate dehydrogenase, when ammonia levels

are high.

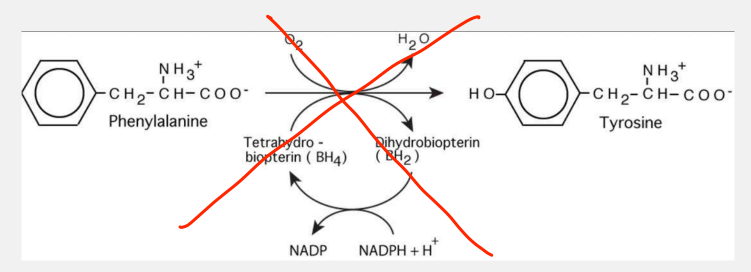


طريقتين نغس المبدأ

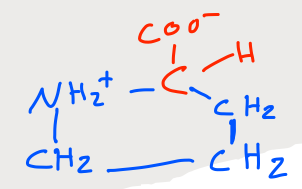


Tyrosine

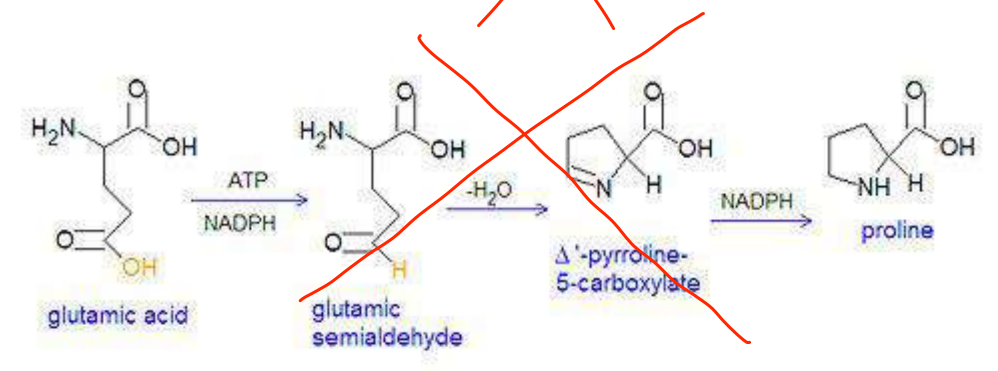
- N H3 C CH2
- Tyrosine is formed from phenylalanine, by phenylalanine hydroxylase. The reaction requires molecular oxygen and the coenzyme tetrahydrobiopterin (BH4), which is synthesized from guanosine triphosphate.
- Tyrosine, like cysteine, is formed from an essential amino acid and is, therefore, nonessential only in the presence of adequate dietary phenylalanine.



Proline

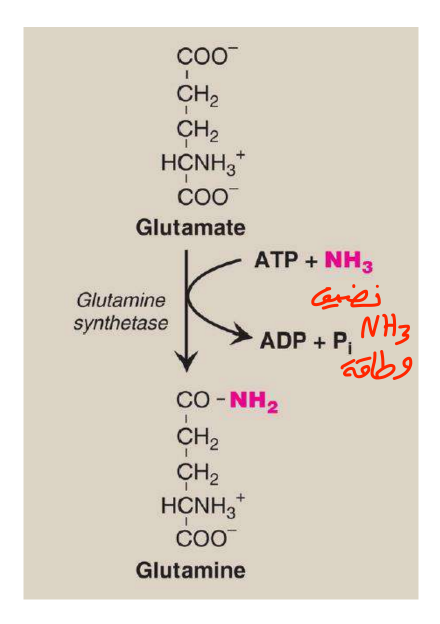


• Glutamate via glutamate semialdehyde is converted to proline by cyclization and reduction reactions



Glutamine

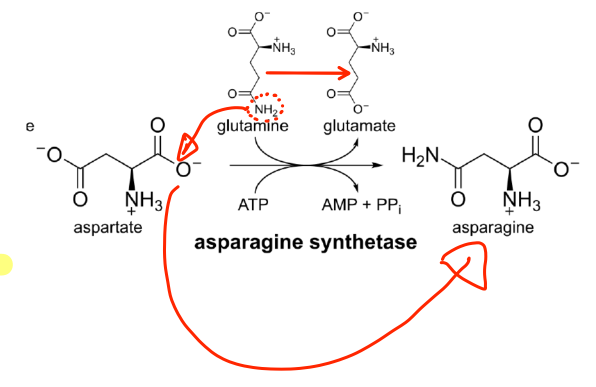
• Contains an amide linkage with ammonia at the γ -carboxyl, is NH_2 formed from glutamate by glutamine synthetase.



Asparagine

- NH3+- C-H1 CH2 CH2
- Asparagine is hydrolyzed by **asparaginase**, liberating ammonia and aspartate.
- This amino acid, which contains an amide linkage with ammonia at the β-carboxyl, is formed from aspartate by asparagine synthetase, using glutamine as the amide donor.

 (glutamate)



Importance of Asparaginase

- Some rapidly dividing leukemic cells are unable to synthesize sufficient asparagine to support their growth. This makes asparagine an essential amino acid for these cells, which, therefore, require asparagine from the blood.
- Asparaginase, which hydrolyzes asparagine to aspartate, can be administered systemically to treat leukemia.
- Asparaginase lowers the level of asparagine in the plasma, thereby depriving cancer cells of a required nutrient

مراجعي

