

GENITOURINARY SYSTEM



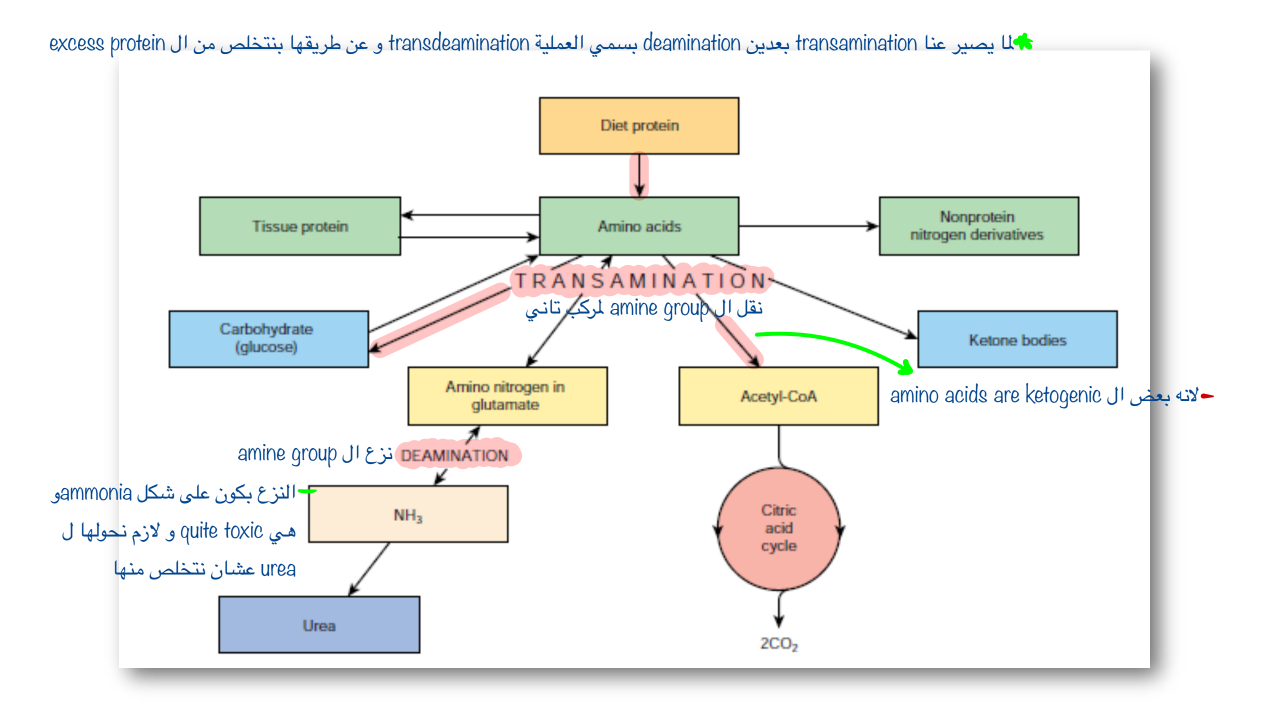




Urea and creatinine metabolism

Ahmed Salem, MD, MSc, PhD, FRCR

40	Special aspects of renal	1.	Discuss urea and creatinine metabolism/ cycle.
	metabolism. Role of kidney in	 Ney in 2. Understand the basic principles on the role of kidney in the regulation of hydrogen ions and bicarbonate buffer system to understand abnormalities in urine composition. 3. Discuss amino acids absorption by the kidney and 	
	acid base balance.		in the regulation of hydrogen ions and bicarbonate
	(Biochemistry 1+2)		buffer system to understand abnormalities in urine
			composition.
		3.	Discuss amino acids absorption by the kidney and
			their disorders
		4.	Discuss normal and abnormal composition of urine
		5.	Interpret the results of routine urine analysis



Nitrogen metabolism

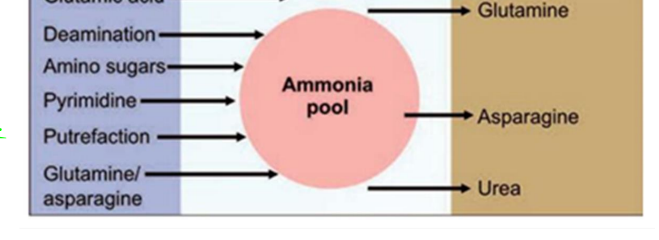
- An adult consuming 100g of protein/d excretes 16.5g nitrogen/d; 95% in urine and 5% in faeces (v small amounts in sweat & other routes)
- <u>Nitrogen balance: quantitative difference btwn nitrogen intake & output</u>
 - ال proteins مش زي ال carbohydrates و ال fats الي بناخدهم عشان نحرقهم proteins مش زي ال intake بروح للبناء
- المفروض يكون بالشخص الي ما بينموا كمية مدخوله نفس الكمية الي بتخرج (Positive nitrogen balance: intake > output
 - Growth, muscular training, pregnancy, recovery from –ve nitrogen balance
 - مثلا الناس الي عندهم uncontrolled cancer احد ال symptoms تعونهم هو weight
- Negative nitrogen balance: output > intake للبروتين catabolism نفسه بفرز شغلات بتعمل catabolism للبروتين

Inadequate protein diet, loss of protein, increased protein catabolism

- Nitrogen equilibrium \rightarrow output = intake
 - Normal healthy adult on an adequate diet

Ammonia

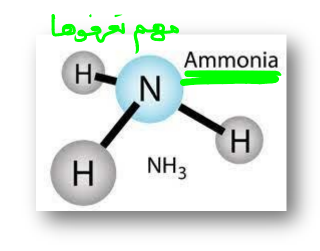
- Universal participant in amino acid synthesis and catabolism (deamination)
- Accumulation in abnormal concentrations → toxic effects
- Ammonia must be eliminated as soon as it is formed

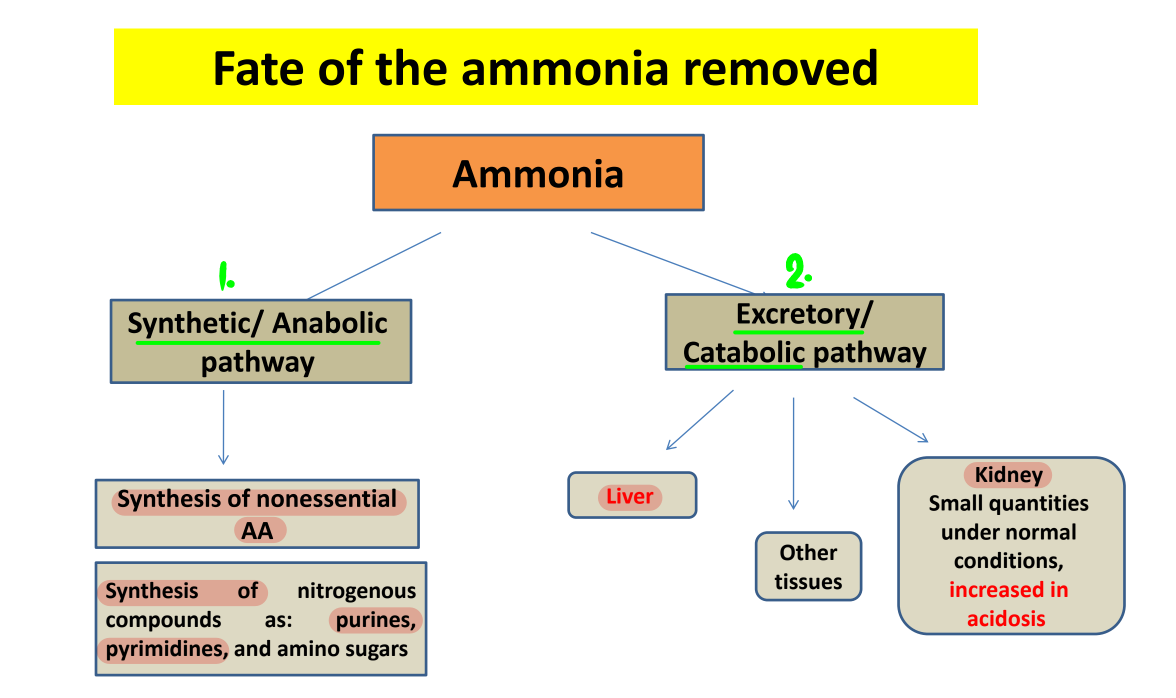


Sources and fate of ammonia

📌 و لازم بس تنتقل من مكان ل مكان تكون محمولة و بالعادة ما بنقدر نطلعها زي ما هي الا نسبة قليلة و المعظم بخرج من الجسم على هيئة urea

Glutamic acid





•Fate of products of deamination:

(A) Fate of the ammonia removed

لا يكون عندك amino acid بدك تتخلص منه بدك تفكر فيه انه مكون من شغلتين amino acid ولا ال amine group و او ال nitrogen و هاي راح تنزعها على هيئة amine group و بدك تفكر بال carbon skeleton الي ممكن تستخدمها بتصنيع اشياء تانية

(A) Fate of the carbon skeleton

Ammonia transport from tissues to liver/ kidney

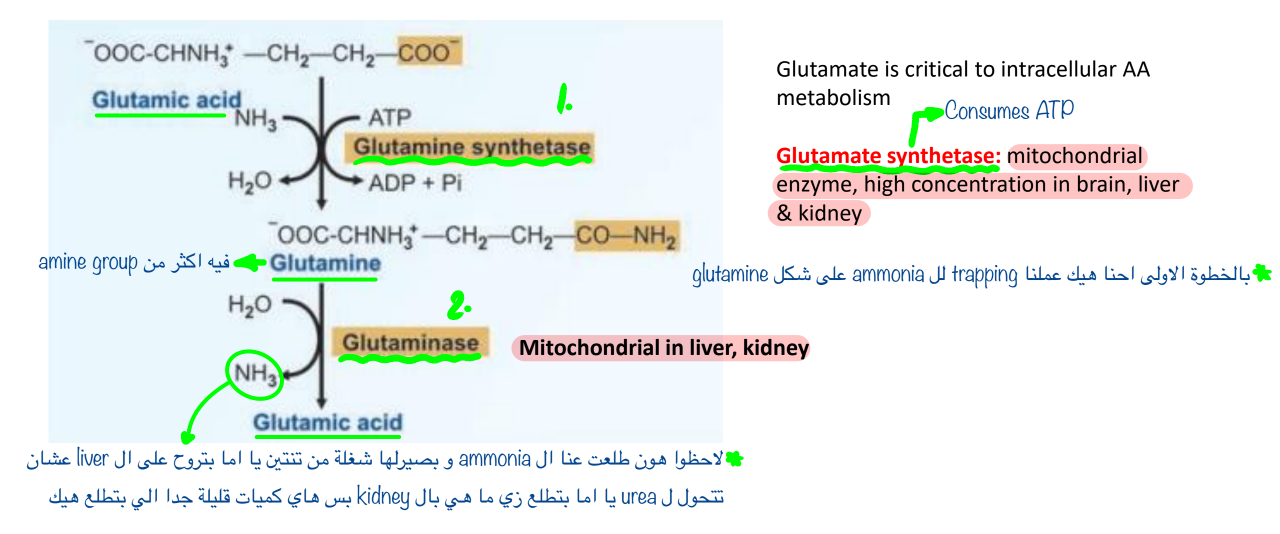
- Inside the cells of almost all tissues, the transamination of amino acids glutamic acid
 structure (ال structure بالسلايد الى تحت)

 ويعتبر acid
 ويعتبر bacid
 <l
- First line of Defense (Trapping of ammonia): Being highly toxic, ammonia should be eliminated or detoxified, as and when it is formed
 - Even very minute quantity of ammonia may produce toxicity in central nervous system
- Intracellular ammonia is immediately trapped by glutamic acid to form glutamine, especially in brain cells

• The glutamine is then transported to liver, where the reaction is reversed by the enzyme glutaminase

Aspartic acid may also undergo similar reaction to form asparagine

Ammonia trapping as glutamine



Catabolic and excretory pathways:

- Being highly toxic to tissues, the ammonia produced in excess of the requirements for anabolic purposes is rapidly disposed of
- The method of disposal depends upon the tissue in which deamination occurs

A- In the liver:

The liver is the main site of deamination of amino acids

- Most of the ammonia released (via glutaminase) is converted to urea
- The urea formed goes via the blood to the kidneys to be excreted in urine

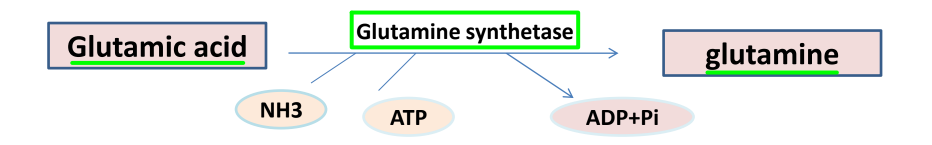
B- In the kidneys:

The ammonia resulting from the deamination of AA in the kidneys is directly excreted in urine

This accounts for about 40% of the urinary ammonia

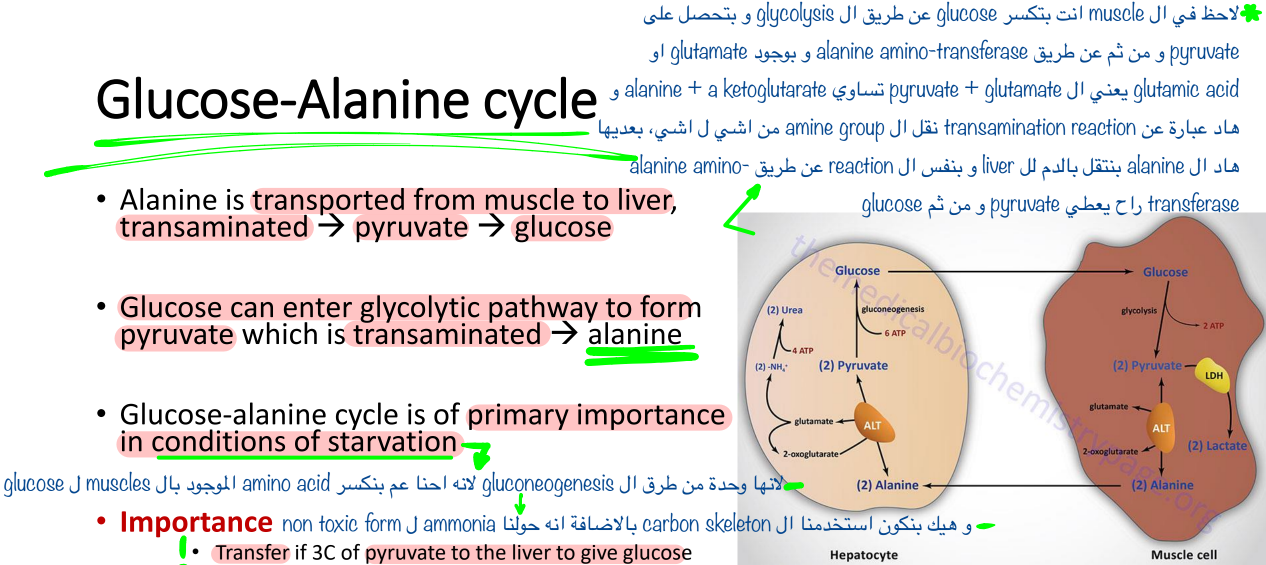
C- In extrarenal tissues:

The ammonia resulting from the deamination of AA in extrarenal tissues, <u>particularly the brain</u>, is converted to glutamine



- Glutamine goes, via the blood, to the kidneys where it becomes hydrolyzed by glutaminase into glutamic acid and ammonia
 - The ammonia is excreted in urine, accounting for about 60% of urinary ammonia
 - This amount increases in acidosis (forms salts with metabolic acids) → counteracting acidosis
- Glutamic acid acts as the link between amino groups of amino acids and ammonia
- The concentration of glutamic acid in blood is 10 times more than other amino acids

Glutamine is the transport forms of ammonia from brain and intestine to liver; while alanine is the transport form from muscle لینزل hydrogen بال ammonia is ا urine ف used to trappe it potentially کمیة ال ammonia الي بتنزل بال urine ممکن تزید کتیر بحالات ال acidosis



 Transfer of NH3 in non-toxic form from muscle to liver to be converted to urea

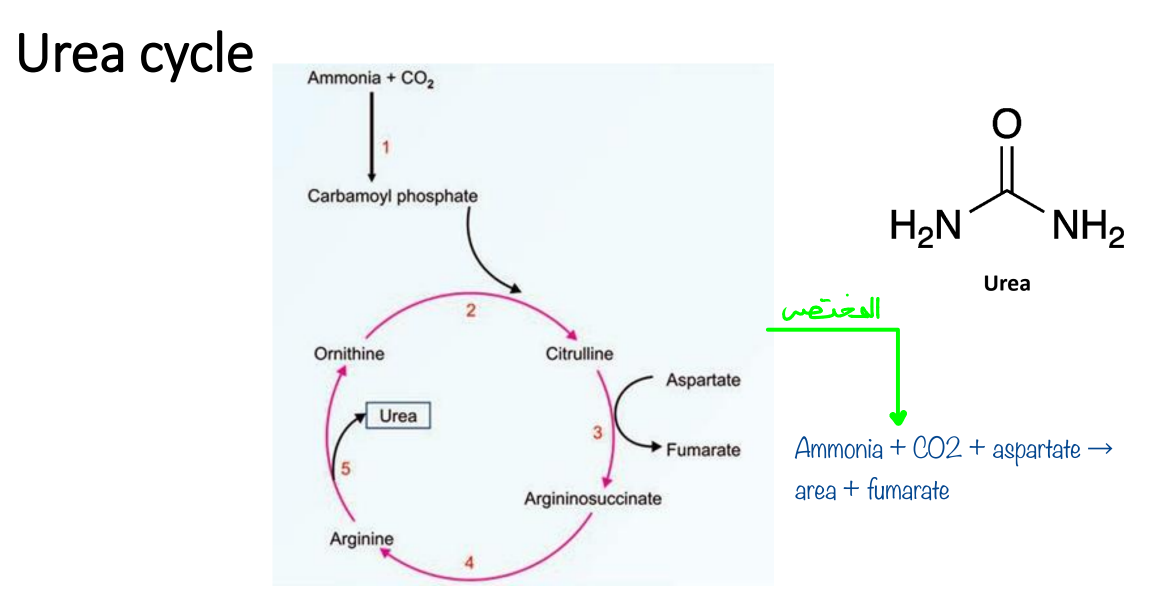
Urea cycle

- Urea is the main way of excretion of ammonia resulting from the deamination of AA
- on of AA which contains 2 nitrogen atoms

 H_2N

- Ammonia is highly toxic to the CNS; it is converted to non toxic urea in the liver only
 - Urea is water soluble easily excreted by the kidneys in urine. Urea is the main end product of protein (amino acids) metabolism
- Plasma urea is 15-45 (20-40) mg/dl, it is formed in the liver and transported in blood to the kidney to be excreted in urine (urinary urea is 15-45 (20-40) g/day)
 Keep in mind area is water soluble
- Urea cycle is known as Krebs–Henseleit cycle (5 reactions, 1-2 in mitochondria; 3-5 in cytosol)
- As ornithine is the first member of the reaction, it is also called as Ornithine cycle
- The two nitrogen atoms of urea are derived from two different sources:
 - one from ammonia; and
 - the other directly from the alpha amino group of aspartic acid

alpha carbon موجودة على ال alpha amino group موجودة على ال alpha carbon اسمها

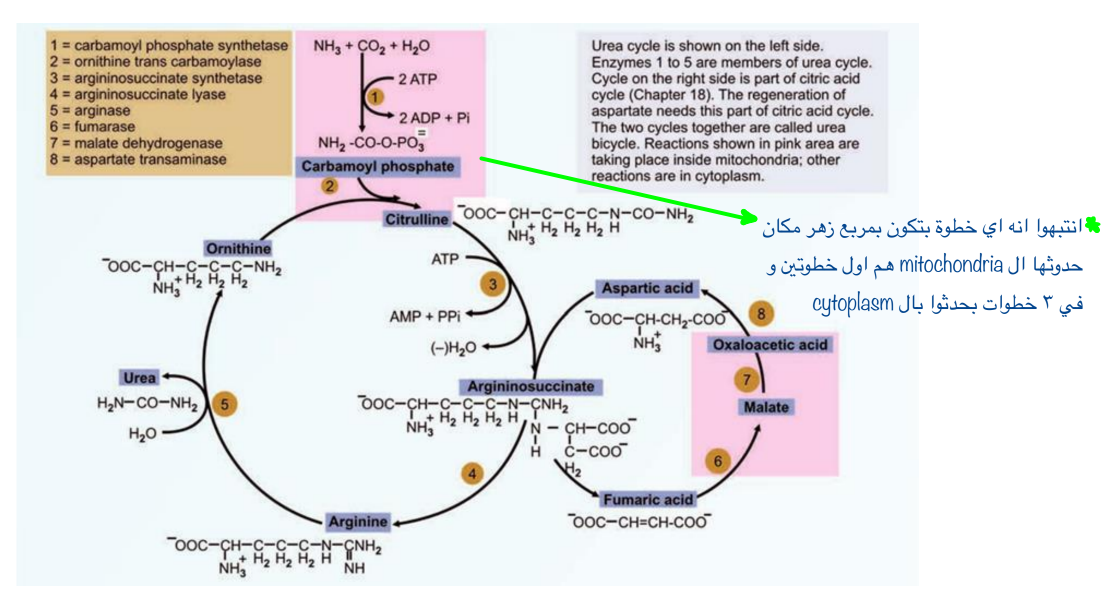


Urea cycle, summary. Note that aspartate enters and fumarate leaves at different steps

Step 1. Formation of Carbamoyl Phosphate

- One molecule of ammonia condenses with CO2 in the presence of two molecules of ATP → form carbamoyl phosphate
- The reaction is catalysed by the mitochondrial enzyme carbamoyl phosphate synthetase-I (CPS-I)
- An entirely different cytoplasmic enzyme, carbamoyl phosphate synthetase-II, (CPS-II) is involved in pyrimidine nucleotide synthesis
- CPS-I reaction is the rate-limiting step in urea formation (It is irreversible and allosterically regulated)

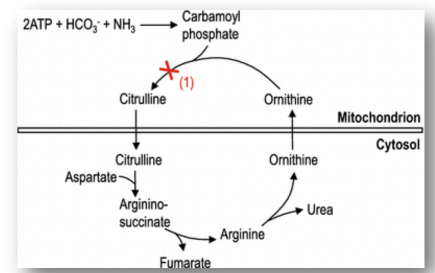
ال catalytic site يعني احنا عنا catalytic site يعني احنا عنا site تاني substrate و عنا site تاني rate of reaction الي بتحكم بال allosteric site تاني NH₃ Carbamoyl phosphate



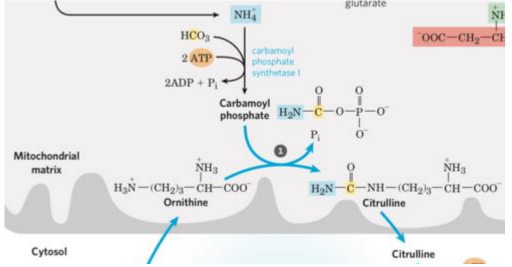
Urea cycle and its relation with citric acid cycle

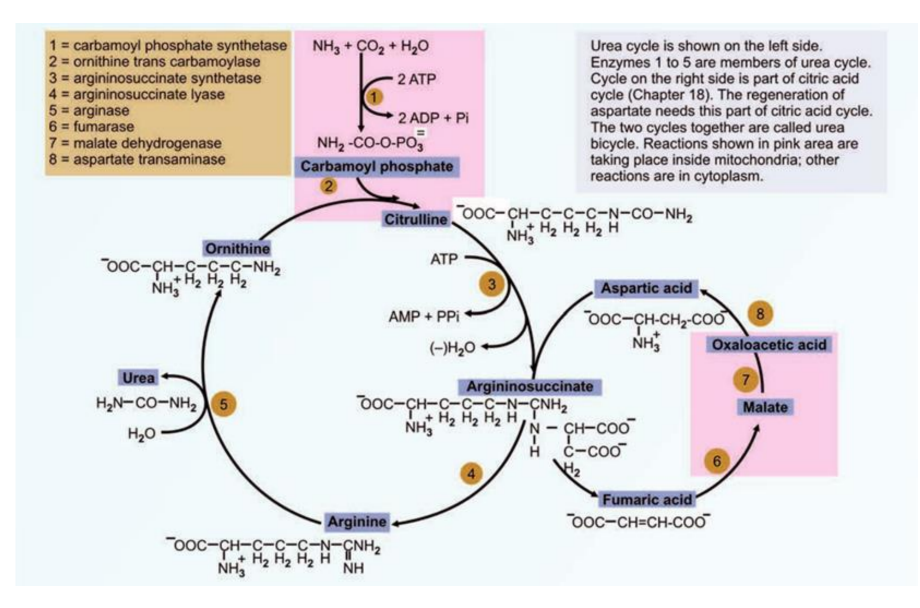
Step 2. Formation of Citrulline

• The second reaction is also mitochondrial



- The carbamoyl group is transferred to the NH₂ group of ornithine by ornithine transcarbamoylase (OTC)
- Citrulline leaves the mitochondria and further reactions are taking place in cytoplasm





Urea cycle and its relation with citric acid cycle

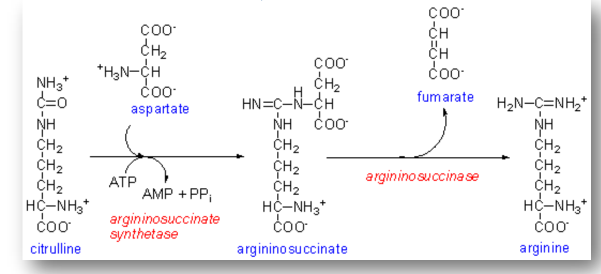
Step 3. Formation of Argininosuccinate

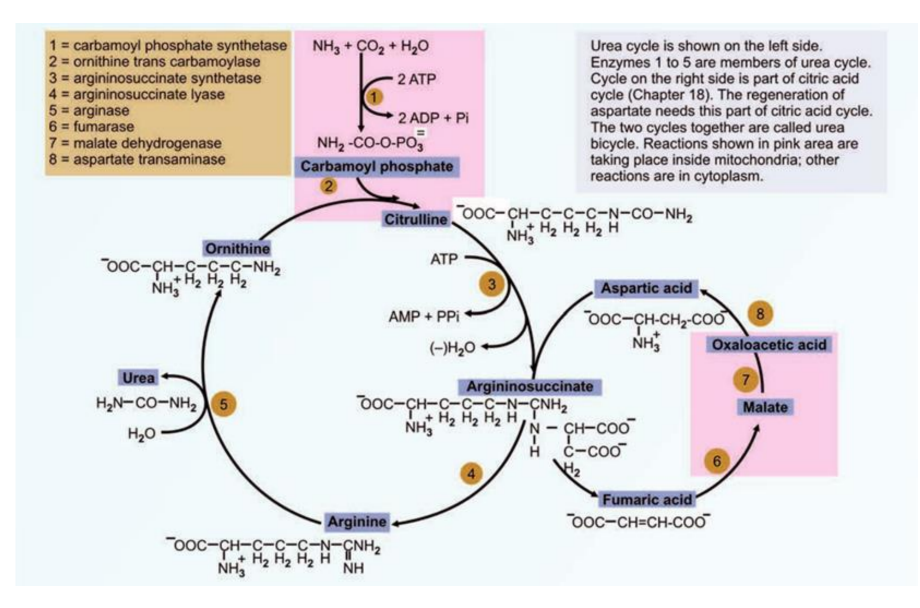
- One molecule of aspartic acid adds to citrulline forming a carbon to nitrogen bond which provides the 2nd nitrogen atom of urea
- Argininosuccinate synthetase catalyzes the reaction
- This needs hydrolysis of ATP to AMP level, so "two relatively high energy phosphate bonds" are utilized

🐥 يعني هون احنا حولنا من ATP ل ADP ثم ل AMP و هيك احنا بنكون استهلكنا one high

The PPi is an inhibitor of this step

energy bond و one relatively high energy bond بس احنا بنحسبهم تنتين relatively high

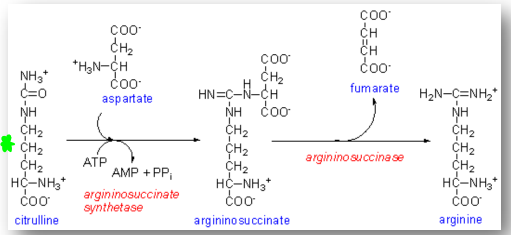


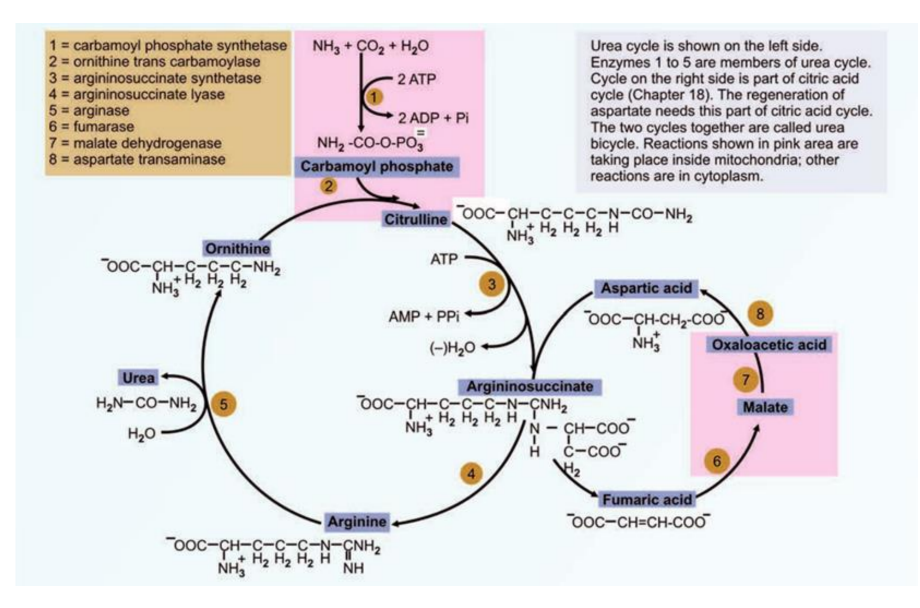


Urea cycle and its relation with citric acid cycle

Step 4. Formation of Arginine

- Argininosuccinate is cleaved by argininosuccinate lyase (argininosuccinase) to arginine and fumarate
- The enzyme is inhibited by fumarate
 - But this is avoided by the cytoplasmic localization of the enzyme
- The fumarate formed may be funneled into TCA cycle to be converted to malate and then to oxaloacetate to be transaminated to aspartate
- Thus the urea cycle is linked to TCA cycle through fumarate
- The 3rd and 4th steps taken together may be summarized as:
 Citrulline + aspartate → Arginine + fumarate





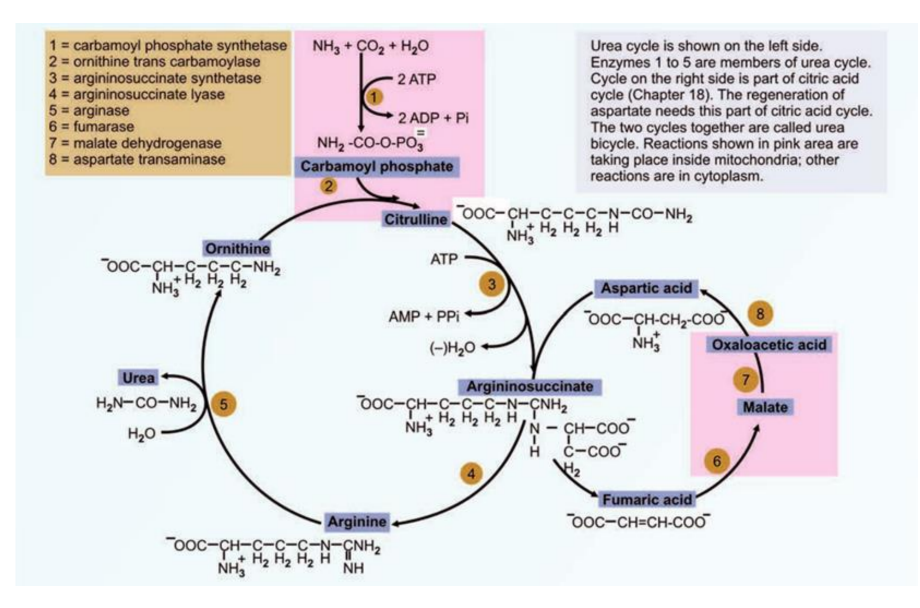
Urea cycle and its relation with citric acid cycle

Step 5. Formation of Urea

• The final reaction of the cycle is the hydrolysis of arginine to urea + ornithine by arginase

– ال ornithine ممكن نعتبره catalyst ب urea cycle

- The ornithine returns to the mitochondria to react with another molecule of carbamoyl phosphate so that the cycle will proceed
- Thus, ornithine may be considered as a catalyst which enters the reaction and is regenerated



Urea cycle and its relation with citric acid cycle

Energetics of Urea Cycle

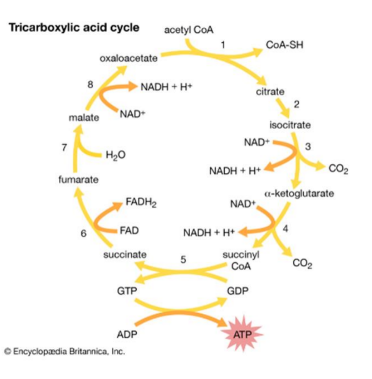
• The overall reaction may be summarized as:

 $NH_3 + CO_2 + Aspartate \rightarrow Urea + fumarate$

- During these reactions, 2 ATPs are used in the 1st reaction
- Another ATP is converted to AMP + PPi in the 3rd step, which is equivalent to 2 ATPs
- The urea cycle consumes 4 high energy phosphate bonds
 high energy منهم هم high energy وحدة relatively high energy بس بنعتبرهم كلهم high energy
- However, fumarate formed in the 4th step may be converted to malate
 - Malate when oxidised to oxaloacetate produces 1 NADH equivalent to 2.5 ATP (new system)
- So net energy expenditure is only 1.5 high energy phosphates
 - The urea cycle and TCA cycle are interlinked, and so, it is called as "urea bicycle"

Relationship between urea cycle and tricarboxylic acid cycle (Kerbs cycle):

- Fumarate produced in urea cycle can be oxidized in Kerbs cycle to oxaloacetate which by transamination give aspartate needed for urea synthesis
- Co2 needed in urea formation is derived mainly from TCA cycle
- ATP needed in urea formation is derived from TCA cycle



Regulation of urea cycle:

• Corse control → Effect of feeding and fasting: the enzymes of urea cycle are:

- (increased by high protein diet
- decreased by low protein diet

Fine control → N-acetylglutamate acts as activator for carbamoyl phosphate synthetase I (CPS I) which is inactive in its absence

Compartmentalization

- The urea cycle enzymes are located in such a way that the 1st two enzymes are in the mitochondrial matrix
 - The inhibitory effect of fumarate on its own formation is minimized because argininosuccinate lyase is in the cytoplasm, while fumarase is in mitochondria

 $\uparrow \uparrow AA \rightarrow \uparrow \uparrow transdeamination \rightarrow \uparrow \uparrow glutamate which$ combines with acetyl coA forming N-acety glutamate

NAG-		Dead	cylase		
Glutamate	CoA	N-Acetyl glutamate (NAG)	H20	Acetate Glutamate	

NAG synthesis and breakdown

Diagnostic importance of plasma urea determination:

- Plasma urea is one of the kidney function tests
- Plasma urea is increased in kidney diseases like renal failure (uremia)
- In liver failure: liver cells cannot convert ammonia to urea so there will be:
 - hyperammonemia (ammonia intoxication); and
 - urea is decreased

- كمية عالية من ال ammonia بالدم يعني مشكلة بال liver

- كمية عالية من ال urea بالدم يعني مشكلة بال kidney

- حكى انه كان يطلب من السنة القبل الماضية انه يحفظوهم بس احنا فش داعي بس مطلوب منا general principles

-راح اعمل هايلايت على الى بحكيه عشان التفريغ يكون شامل

Disorders of Urea Cycle

- Deficiency of any of the urea cycle enzymes would result in hyperammonemia
- When the block is in one of the earlier steps, the condition is more severe, since ammonia itself accumulates

مس مش كتير toxic عانت بالخطوات الى تحت ف الي راح يتراكم هو intermediates و هم مش كتير toxic زي ال ammonia

- As a general description, disorders of urea cycle are characterized by hyperammonemia, - بالمبدأ العام اهم واحد فيهم هو نقص orthinine trans carbomylase و هو x-linked condition و هو x-linked condition
- Clinical symptoms include vomiting, irritability, lethargy and severe mental retardation (if untreated) حكى انه المطلوب منا انه نكون فاهمين انه لما يكون المشكلة بالانزيمات الي فوق بتكون المشكلة اسوء لانه ال ammonia هي الي راح تتجمع و ال ammonia بشمل عام بكون الها

alcoholic, chronic hepatitis سببين يا اما بسبب نقص انزيمات الي هو inborn irror و هاي مش كتير common اما المشكلة لاكثر شيوعا انه يكون ال liver مثلا حدا endonic, chronic hepatitis • Infants appear normal at birth, but within days progressive lethargy sets in

🗕 في جزء تاني للعلاج غير ال diet هو انك تعطي اشي يمنع البكتريا بالامعاء انها تنتج ammonia 🗢

- **Treatment is** more or less similar in the different types of disorders
 - Low protein diet by frequent feeding can minimize brain damage since ammonia levels do not increase very high toxic effect of ammonia especially on the brain لانه اخطر عرض لهاد الاشی هو 😽

Disorders of Urea Cycle

• Brain is very sensitive to ammonia, \uparrow ammonia leads to:

- Ammonia will combine with α -ketoglutaric acid forming glutamate and glutamine \rightarrow
 - Image of the second seco
- \uparrow levels of glutamine \rightarrow \uparrow osmotic pressure in the astrocytes \rightarrow which become swollen

Cerebral edema

+ other mechanisms

neurotransmitter و التانية الها علاقة بال gaba aminobutyric acid و التانية الها علاقة ب gaba aminobutyric acid الي هو 🌪 وحدة منهم الها علاقة بال

Fate of carbon skeleton of amino acids

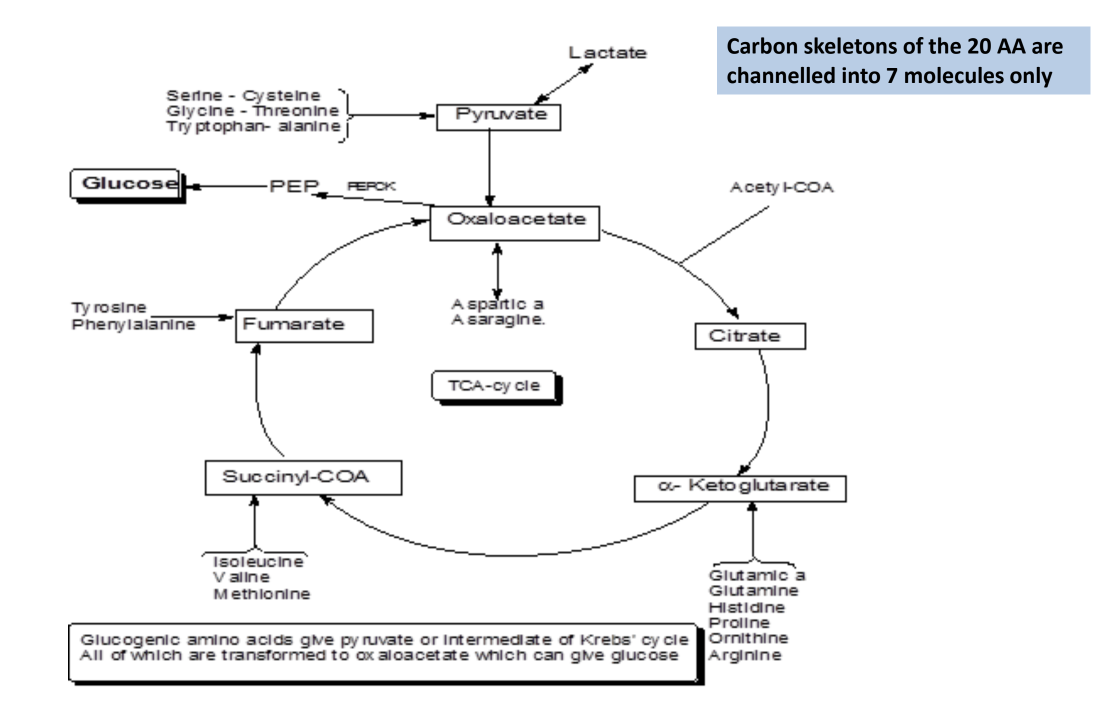
- Ketogenic AA: produce acetyl coA or aceto-acetyl coA used in ketogenesis
 Leucine
 - Lysine

Glucogenic and ketogenic AA: can give both glucose & ketone bodies

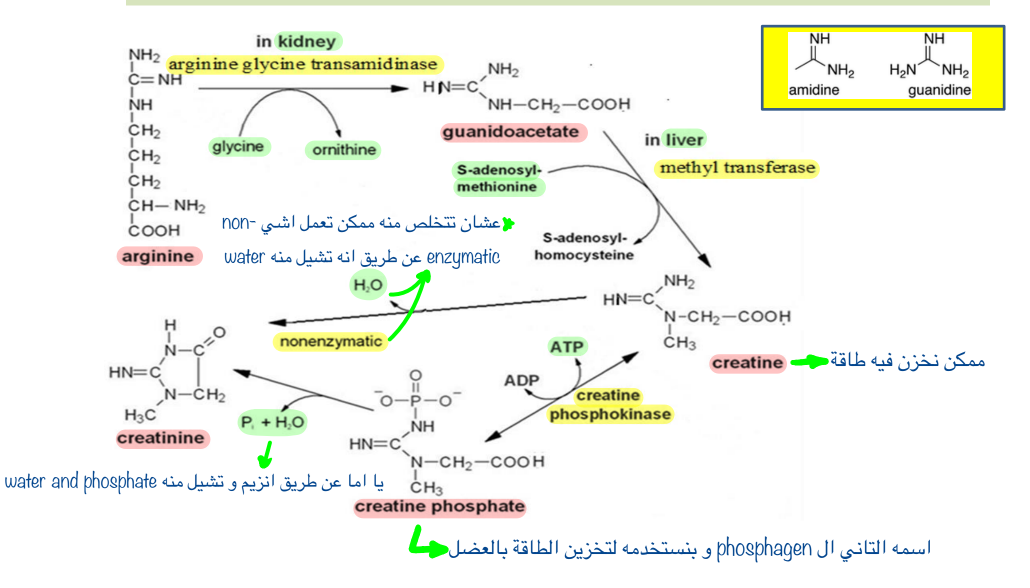
- Tyrosine
- Phenylalanine
- Tryptophan
- Isoleucine

7 العشرين amino acid الي بنستخدمهم لبناء البروتينات راح استخدمهم بتصنيع واحد من pyruvate, oxaloacetate, fumarate, succynil CoA, alpha : الي هم molecules الي مع ketogluterate,leucine, lysine





Creatine and Creatinine Metabolism



- <u>Creatine</u> is <u>methyl guanido acetic acid</u>. It is a NPN compound.
- It is widely distributed in our tissues: mainly (98%) in muscles as <u>phosphocreatine</u> (= phosphagen).
- <u>Creatinine</u> is creatine anhydride, it is the excretory product of creatine.
- The transamidinase reaction occurs in the kidney.
- The methyl transferase reaction occurs in the liver.

- The creatine goes via blood to different tissues mainly to the muscles (98% of the body creatine).
- Androgen (male sex hormones e.g. testosterone) increase the uptake and retention of creatine by muscles.

Function :

- Creatine forms creatine phosphate (phosphagen) which is the main storage form of energy in the cells.
- During muscular exercise, ATP is consumed rapidly to ADP. ATP is formed quickly at the expense of creatine phosphate by reversal of the CPK reaction. <u>This occurs</u> <u>before glycoysis starts to produce ATP.</u> i.e. maintain ATP during 1st few minutes of muscle contraction.

- Creatinine (the creatine anhydride) is formed from creatine <u>or</u> creatine phosphate.
- About 2% of body creatine is converted to creatinine. The amount of creatinine excreted for each individual is nearly constant (1-2 g/ day) and it is related to muscle mass.
- <u>The normal serum creatinine level</u> 0.6- 1.2 <u>mg/dl.</u>
- Plasma creatinine level increases in cases of kidney diseases and it is a good index for renal functions as its level is not affected by diet. hydration status اله علاقة بال diet و اشياء كتير زي مثلا ال

븆 هدول اسئلة حطهم الدكتور بمحاضرة اليوم نشكر محمد العمري انه صورهم و

بعتلي اياهم احطلكم اياهم بالتفريغ 🖊

In the urea cycle, which enzyme catalyzes the formation of fumarate • a) Arginase • b) Argininosuccinate synthetase • c) Argininosuccinate lyase • d) Ornithine transcarbamylase • e) Carbamoyl phosphate synthetase | • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • • <

A

- c) Ornithine
- d) Arginine
- e) Argininosuccinate
- Which urea cycle enzyme deficiency leads to the accumulation of argininosuccinate, resulting in citrullinemia and hyperammonemia?
- a) Ornithine transcarbamylase
- b) Arginase
- c) Carbamoyl phosphate synthetase I
- d) Argininosuccinate synthetase
- e) Argininosuccinate lyase

Answers : C, A, D