



# Brain Energy Metabolism II

major fuel source

- glucose → normal condition
- ketone bodies → starvation



ATP →  $\text{Na}^+$ -K channel

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# Glucose Metabolism Produce Energy



- **Glucose** metabolism in the brain is similar to that in other tissues. In an aerobic conditions, it includes three principle metabolic pathways:
  1. Glycolysis
  2. Tricarboxylic acid cycle (TCA) or Krebs cycle
  3. Oxidative phosphorylation and the electron transport chain (ETC)

# 1. Glycolysis → 2 pyruvate



- **Glycolysis** is the metabolic pathway which converts glucose (C6) into pyruvate (C3)
- It occurs in the **cell cytosol**. Glycolysis is a sequence of ten **oxygen-independent** and enzyme-catalyzed steps with the intermediates provide entry points to the cycle
- When the glycolysis end products (pyruvate and NADH) are disposed in presence of O<sub>2</sub>, the process is then called aerobic. Alternatively, in an anaerobic conditions, the products are removed via the lactic acid fermentation process

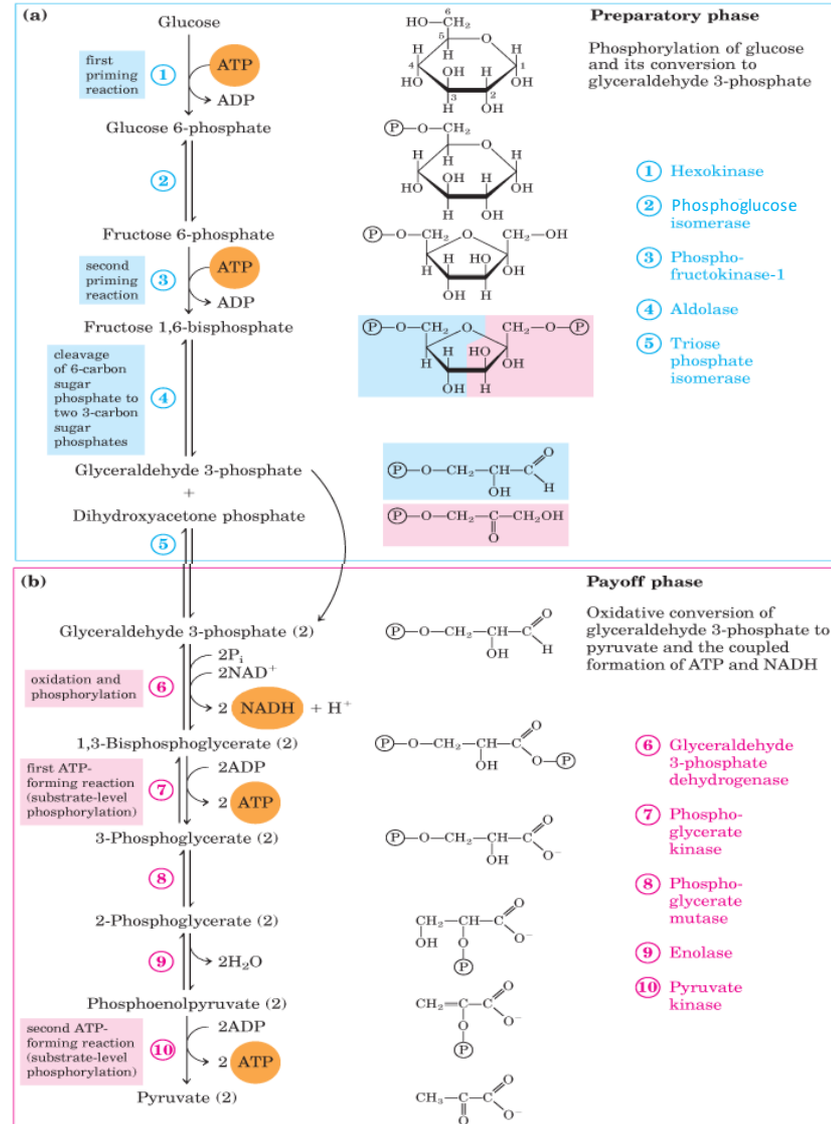
# 1. Glycolysis



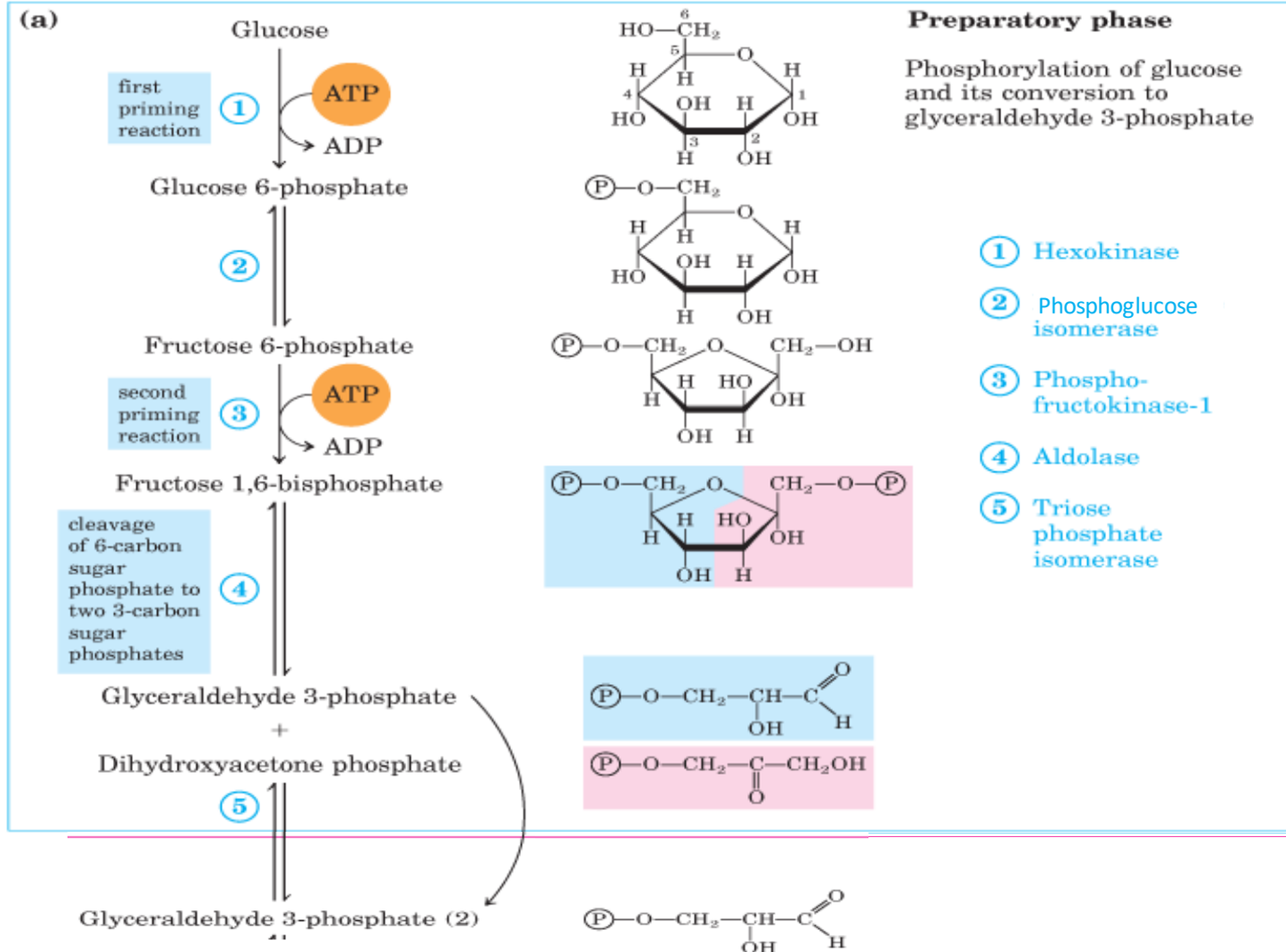
- The entire pathway is divided into two distinct phases:

- a) **Energy Investment Phase (Preparatory Phase)** - 2 ATP (استثمار) (استهلاك)
- b) **Energy Generation Phase (Pay Off Phase)** + 4 ATP (توليد) (النتج)

$$\sum \text{ATP} = +2 \text{ATP} \text{ (النتج)}$$



# a) Preparatory Phase



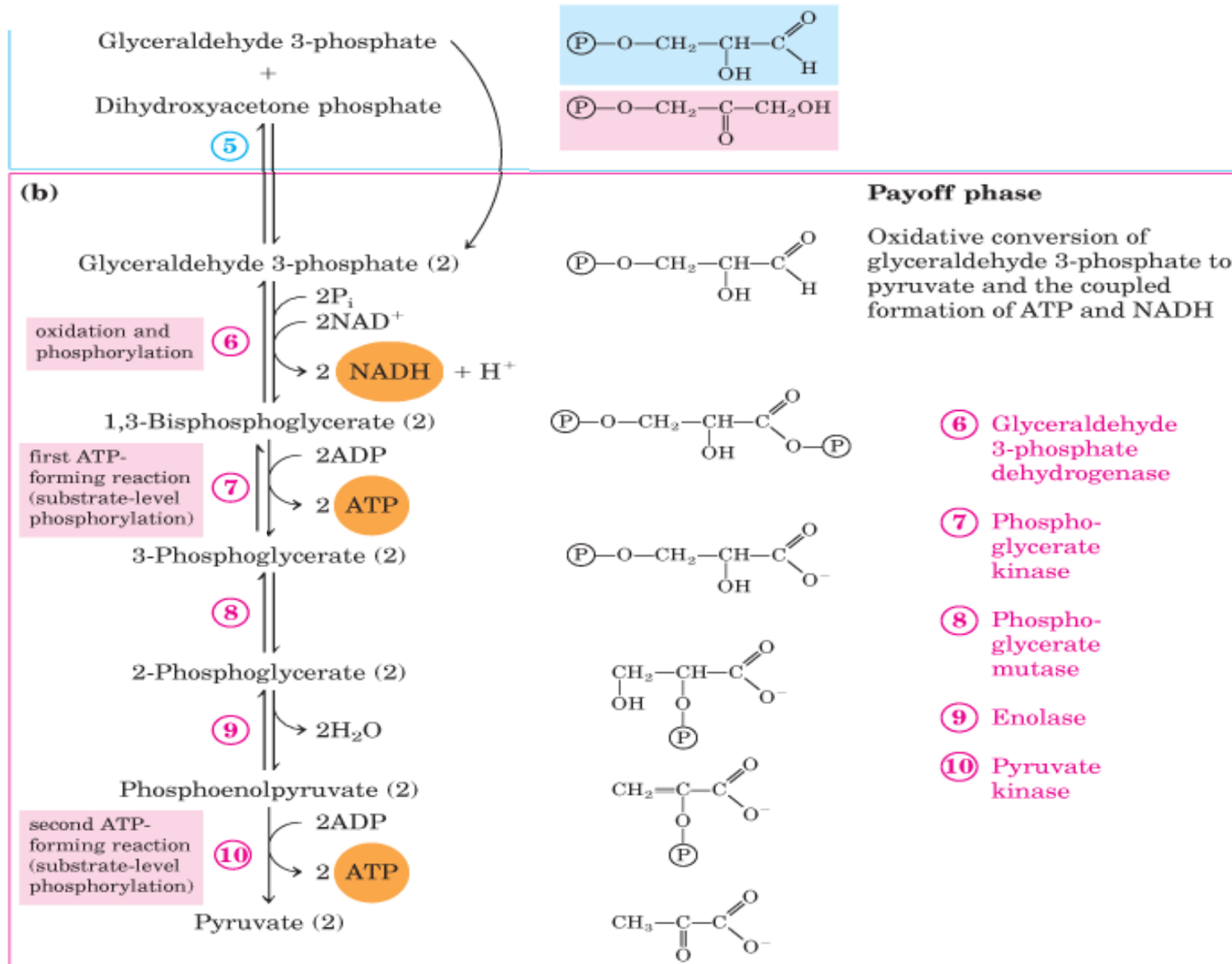
## a) Preparatory Phase

X



- **Step 1:** This first priming reaction is important to maintain the influx of glucose and at the same time to trap the transported glucose molecules inside the cell
- **Step 2:** Phosphoglucose isomerase (PGI) interconverts G6P and F6P. Indeed, Mannose and Fructose can enter the glycolytic pathway at this point
- **Step 3:** This is a **rate limiting** or key regulatory step because phosphofructokinase-1 (**PFK-1**) is an allosteric enzyme and its activity can be controlled
- **Step 4:** The cleavage to 2 triose phosphates: **DHAP** (dihydroxyacetone phosphate) and **GADP** (glyceraldehyde-3-phosphate)
- **Step 5:** Isomerization of DHAP by triose phosphate isomerase (TPI) to proceed further in glycolysis

# b) Pay Off Phase



## b) Pay Off Phase

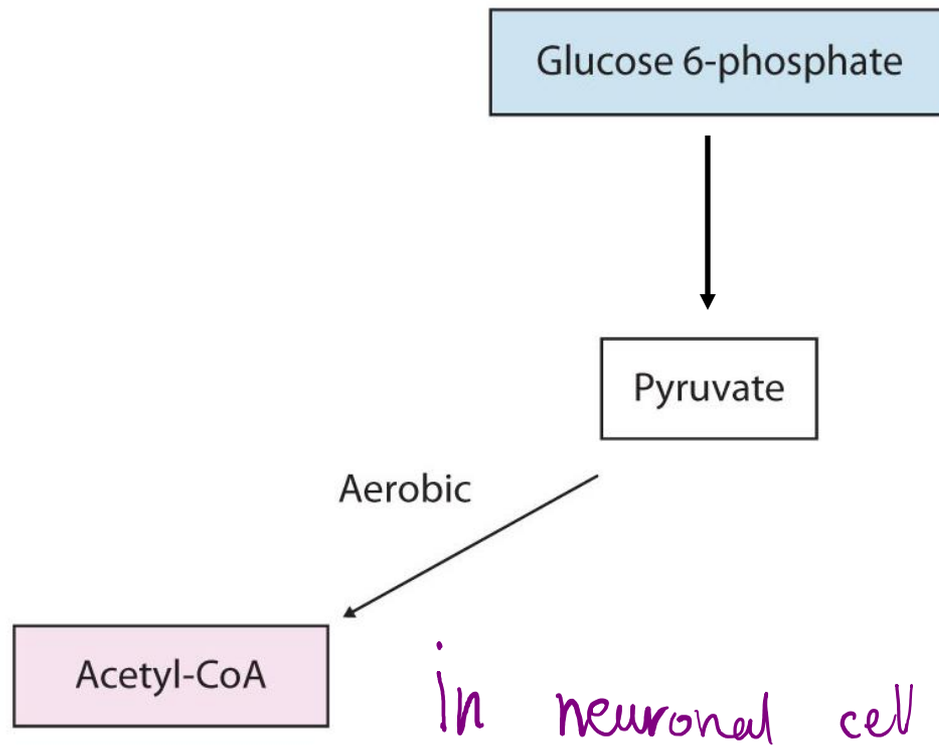


- **Step 6:** Dehydrogenase enzyme catalyzes the oxidative phosphorylation of GADP (electron donor) into super-high-energy compound (**1,3-BPG**) and the transfer of electrons into the coenzyme  $\text{NAD}^+$  (electron acceptor) forming **NADH**
- **Step 7:** The **first ATP** molecule is generated by the substrate-level phosphorylation process catalyzed by phosphoglycerate kinase (PGK)
- **Step 8:** Phosphoglycerate mutase (PGM) catalyzes the internal shifting of P group from C3 to C2
- **Step 9:** The synthesis of the second super-high-energy compound **phosphoenolpyruvate (PEP)** in a simple dehydration reaction catalyzed via enolase enzyme
- **Step 10:** The **second ATP** molecule is generated by the substrate-level phosphorylation process catalyzed by pyruvate kinase (PK). Pyruvate is the final product of glycolysis

# Metabolic Fates of Pyruvate in Brain



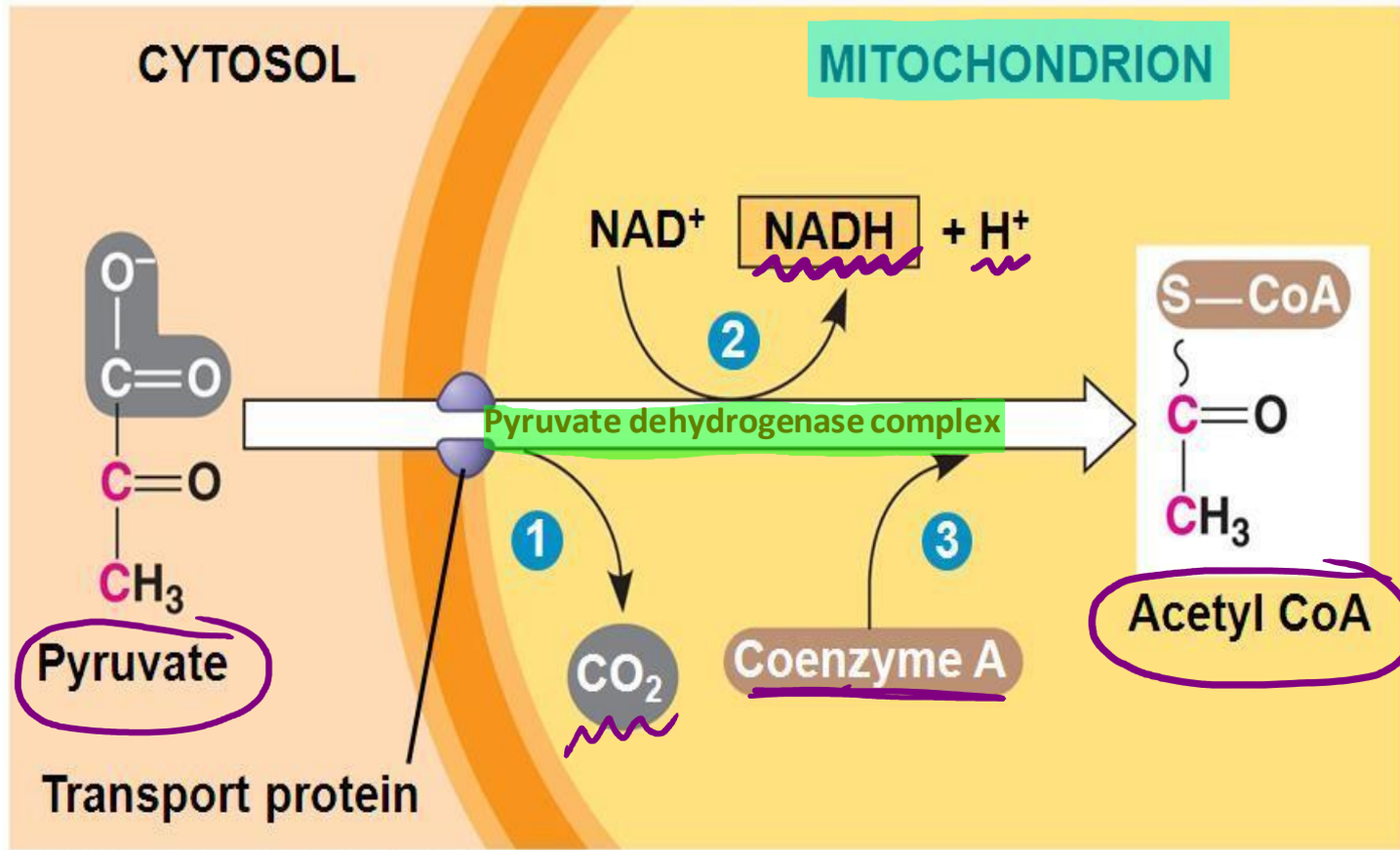
ATP in brain  $\rightarrow$  70%  $\rightarrow$   $\text{Na}^+/\text{K}^+$  ATPase Pump  $\rightarrow$  active transports



# Metabolic Fates of Pyruvate in Brain



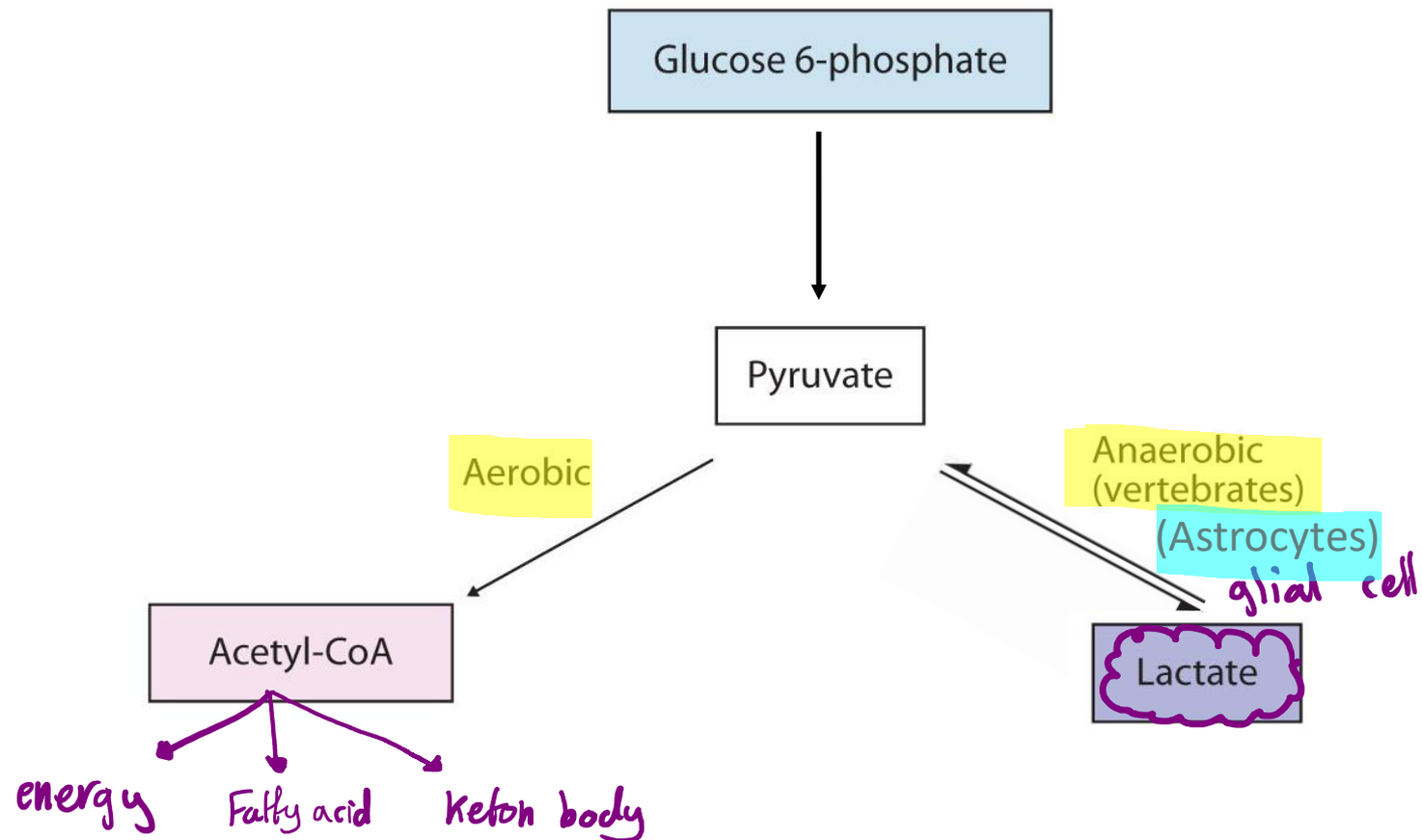
*irreversible*



# Metabolic Fates of Pyruvate in Brain



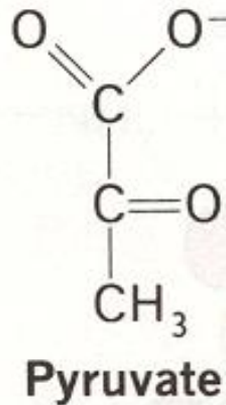
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# Metabolic Fates of Pyruvate in Brain



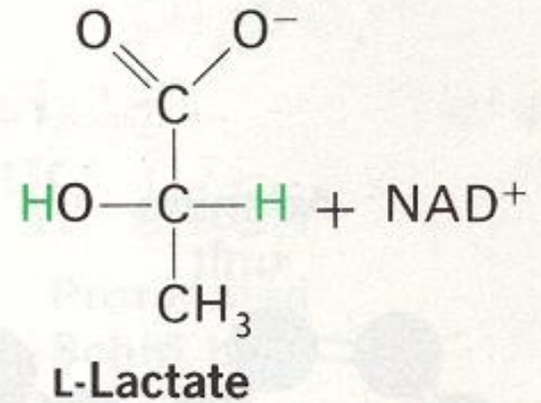
Cytosol



Lactate dehydrogenase

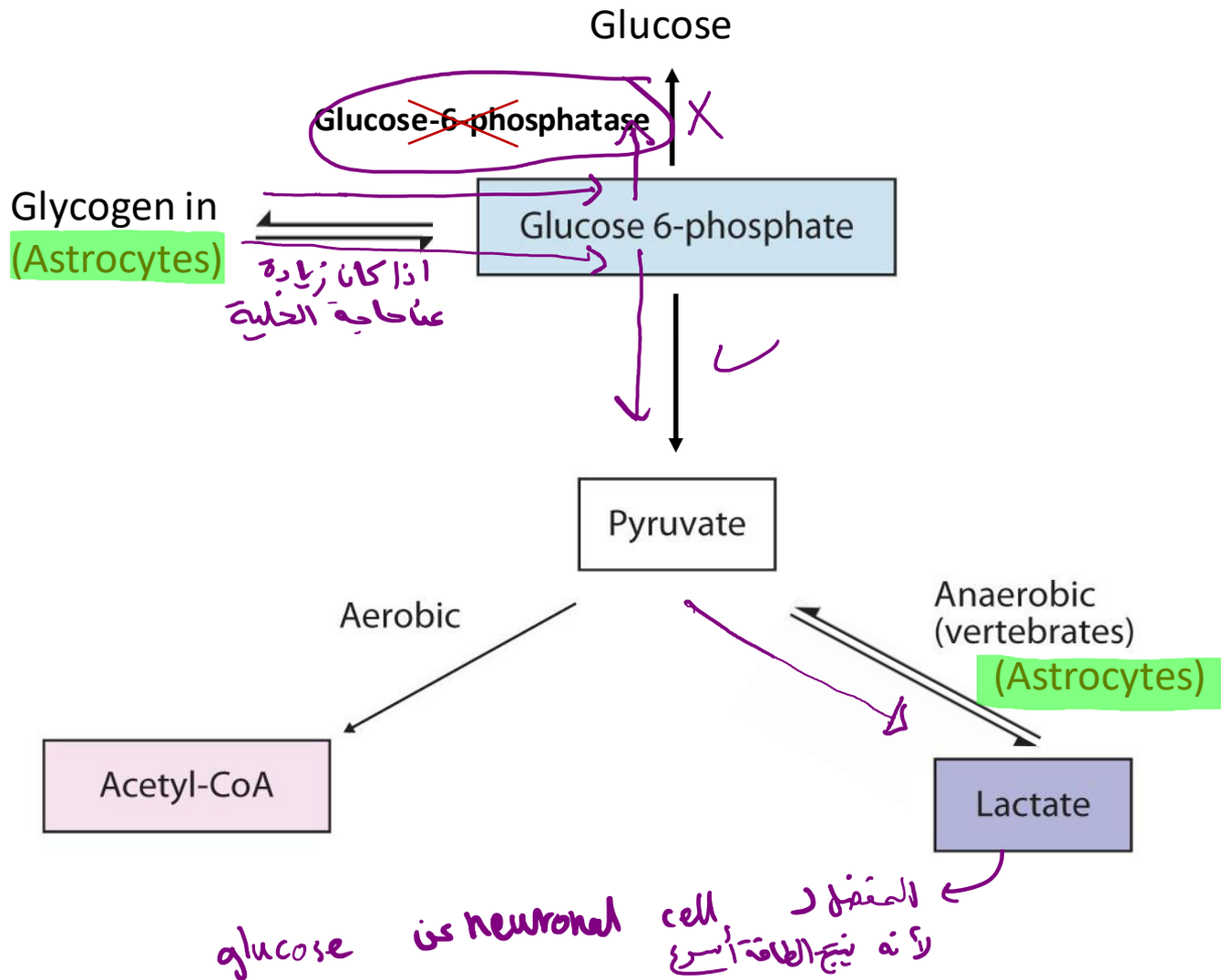
Fermentation

Astrocyte



↓  
neuronal cell    پینجی ر

# Metabolic Fates of Pyruvate in Brain



# Glycogen in Brain

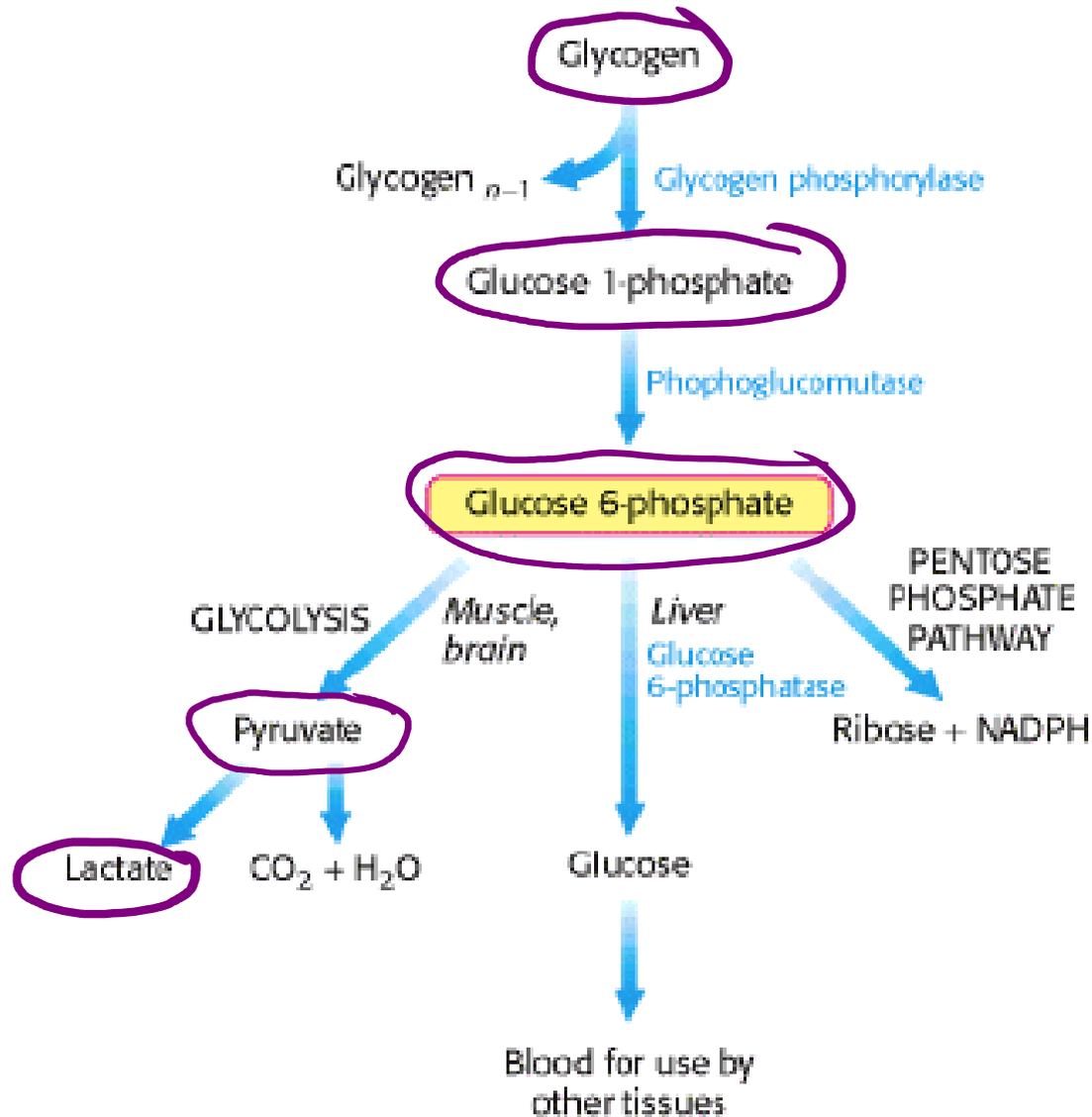


- Very limited amount of **glycogen** (3-12  $\mu\text{mol} / \text{g}$  tissue) is stored in brain predominantly in **astrocytes**
- Actually its role as energy source during hypoglycemia is restricted to few minutes
- **Glycogenolysis** or **glycogen mobilization** is the process of glycogen breakdown to glucose units. It occurs in the **cytosol** of glycogen-containing cells such as brain
- **Glycogenolysis** produces monomers of **glucose 1-phosphate** which is then converted to **G6p** by phosphoglucomutase enzyme

# Glycogenolysis



الانزيمات  
مس حفظ



# Fates of Glucose 6-phosphate



↓  
fuel

- The produced G6p has many possible routes. In brain, **G6p molecules** join the **glycolysis** and used as **fuel**.
- Indeed, due to the lack of **glucose 6-phosphatase** enzyme in brain and muscle tissues, **G6p** can't be converted back to glucose to be released in the blood as the case in liver cells

# Acetyl CoA Fate



- Sources of Acetyl CoA: fat metabolism (fatty acids  $\beta$ -oxidation, reversible) and CHO metabolism (pyruvate, irreversible)



- Fates of Acetyl CoA:

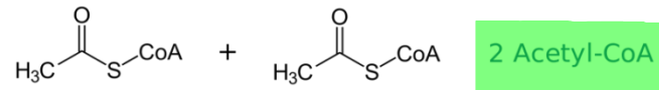
1. It can enter the Krebs cycle for energy production
2. Used for biosynthesis of fatty acids **but not CHO**
3. Formation of ketone bodies (Ketogenesis)

# Ketogenesis

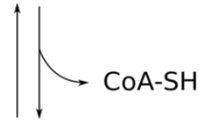


- Ketogenesis is the process of ketone bodies production from acetyl CoA mainly in **the mitochondrial matrix of hepatocytes**
- Ketogenesis occurs when acetyl CoA accumulates beyond its capacity to be oxidized (via Krebs cycle) or used for fatty acids synthesis (lipogenesis)
- When acetyl CoA level is high, 2 molecules of acetyl CoA undergo a reversal of thiolase reaction to acetoacetyl CoA which reacts with a third molecule of acetyl CoA to produce  $\beta$ -hydroxy- $\beta$ -methylglutaryl-CoA (HMG-CoA)
- HMG-CoA is converted to acetoacetate which undergoes either NADH-dependent reduction to  $\beta$ -hydroxybutyrate (reversible reaction) or spontaneous decarboxylation to acetone (in very small amounts)

# Ketogenesis



Thiolase



HMG-CoA synthase

Acetyl-CoA

CoA-SH

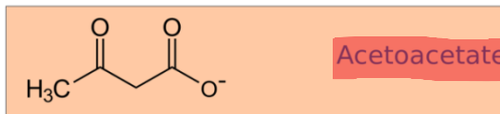
Rate limiting step



HMG-CoA lyase

Acetyl-CoA

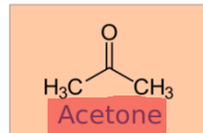
①



Non-enzymatic decarboxylation

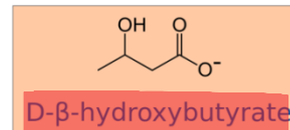
CO<sub>2</sub>

②



③

NADH + H<sup>+</sup>  
NAD<sup>+</sup>  
D-β-hydroxybutyrate dehydrogenase



3 keton bodies

موجود در هپاتوسیتس

# Ketogenesis



- HMG-CoA synthase is primarily expressed in hepatocytes and catalyzes the rate-limiting step in ketogenesis
- Ketone bodies are produced in the liver from accumulated Acetyl CoA during ketogenic conditions (uncontrolled diabetes and starvation) due to enhanced fat catabolism ( $\beta$ -oxidation of fatty acids)

# Ketone Bodies



- Ketone bodies (KB) are three water soluble molecules: acetoacetate,  $\beta$ -hydroxybutyrate ( $\beta$ -HB) and acetone
- Ketone bodies are important metabolic fuels for many peripheral tissues under normal conditions, particularly skeletal muscles, and during starvation they become the brain's major fuel source
- Ketone bodies transported from liver to other tissues where both acetoacetate and  $\beta$ -hydroxybutyrate can be reconverted to acetyl CoA for energy production, a process called ketolysis which occurs in mitochondria of extrahepatic tissues
- The reconversion first involves the transfer of all  $\beta$ -HB into acetoacetate followed by the enzymatic transfer of CoA moiety from succinyl-CoA to acetoacetate yielding acetoacetyl CoA and succinate (rate limiting step). Finally, thiolase converts acetoacetyl CoA to two molecules of acetyl CoA which enters Krebs cycle for energy production

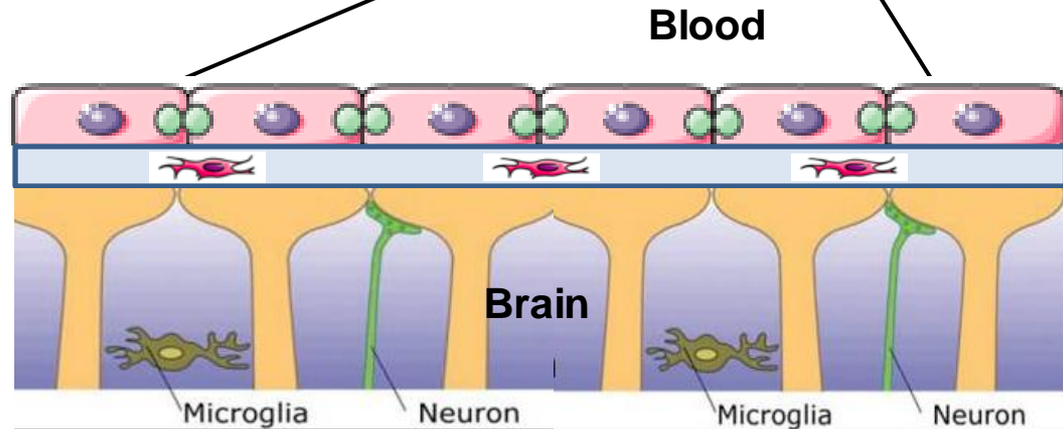
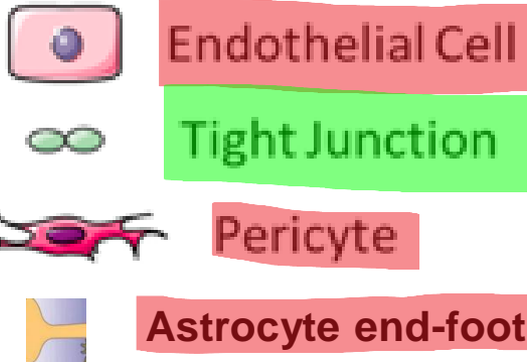
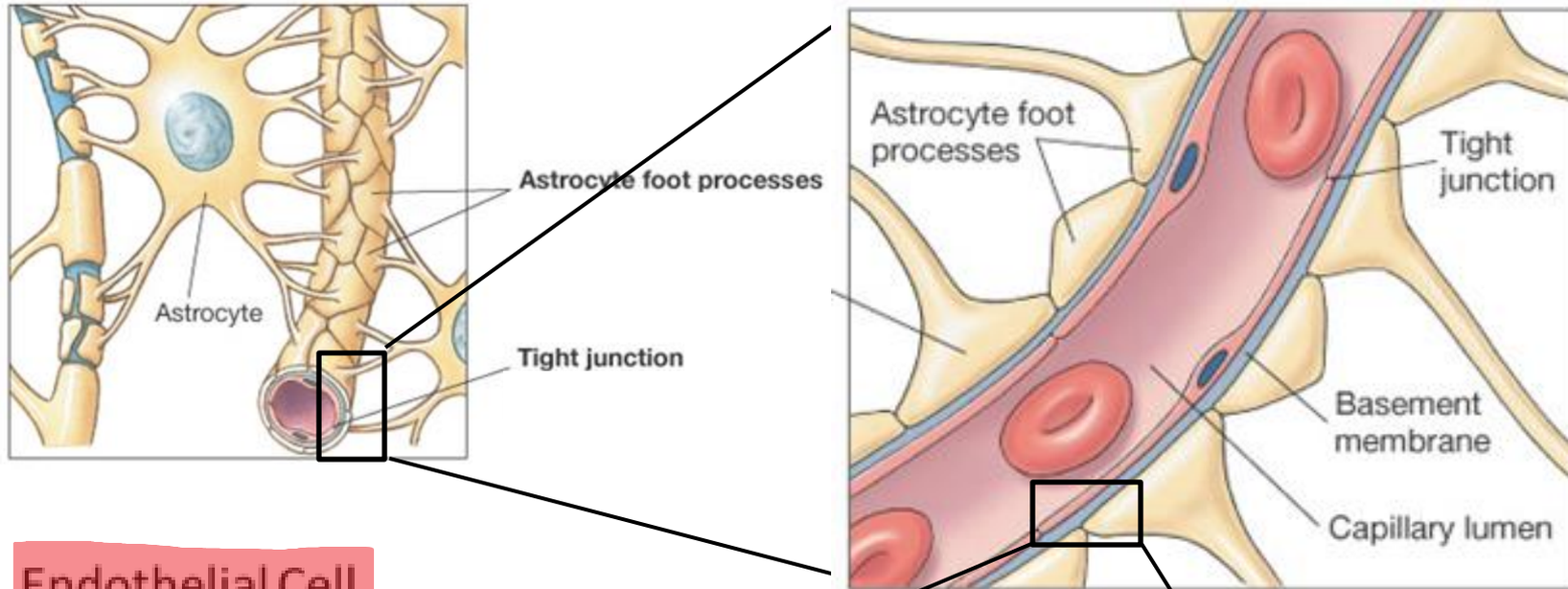


# Blood-Brain Barrier (BBB)



- BBB is a highly selective membrane which allows only very specific molecules to access the CNS so protecting the brain from circulating toxic substances and invading foreign bodies (e.g. bacterial infection)
- Therefore, BBB has a critical role in cerebral homeostasis
- The cellular and structural components of BBB:
  1. **Non-fenestrated endothelial cells (ECs)** which are connected via **tight junctions** thus prevent paracellular diffusion. Endothelial cells are supported by a **continuous basement membrane (BM)**
  2. **Astrocytes** send processes called **end-feet** which surround capillary walls to additionally support the ECs and maintain the BBB integrity
  3. **Pericytes** are embedded in the BM. They have a role in BBB development (e.g. formation of tight junction)

# Blood-Brain Barrier BBB



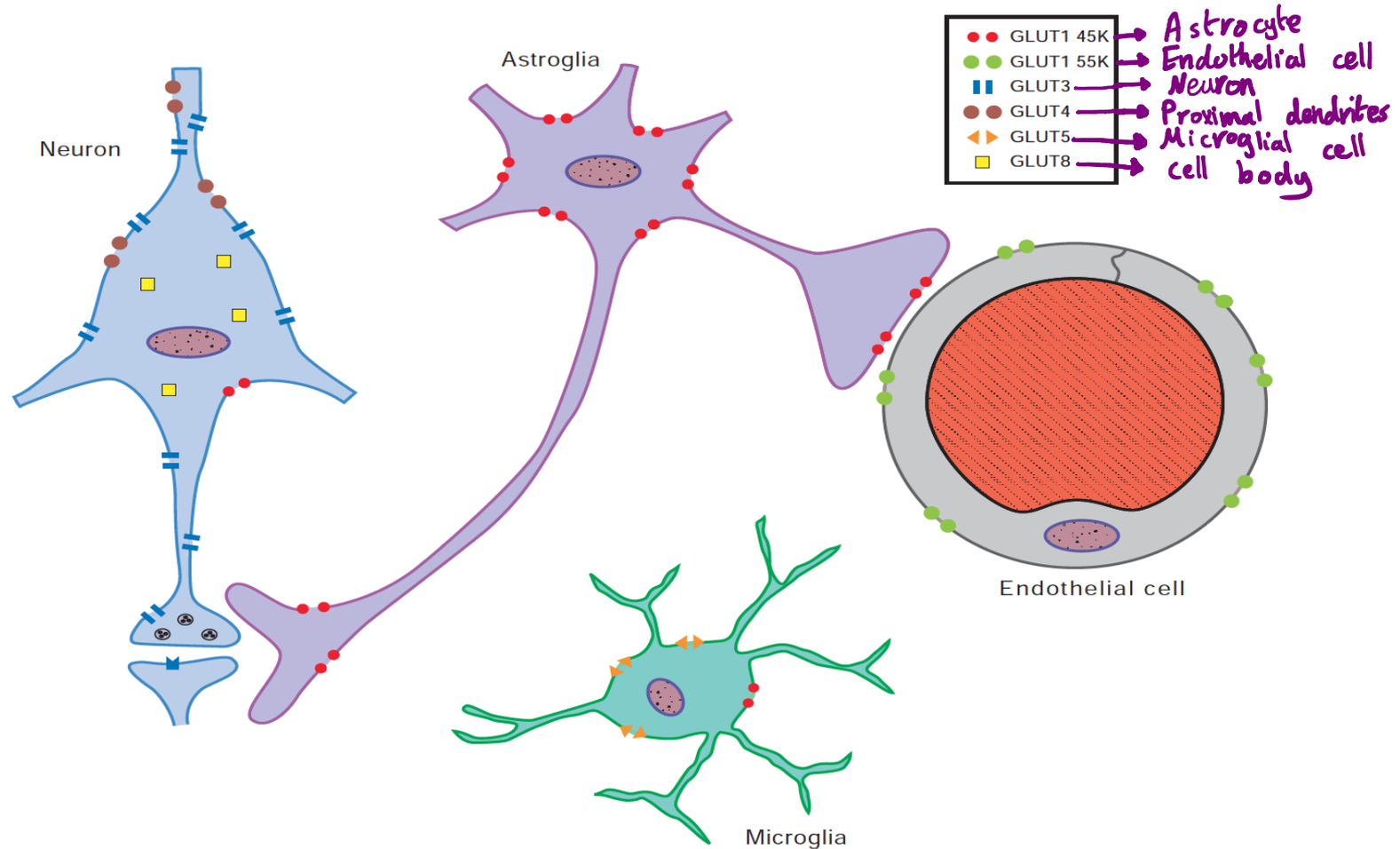
# Glucose Transporters



- Energy substrates cross BBB and reach brain cells via specific transport mechanisms. For example, glucose is hydrophilic molecule which enters the cells through glucose transporters (GLUT), a family of glycosylated transmembrane proteins
- In brain, seven transporters are expressed in a cell-specific manner:

- ❖ 55-KDa isoform of GLUT1 essentially localized on endothelial cells of BBB
- ❖ 45-KDa isoform of GLUT1 is localized predominantly in astrocytes (star-shaped non-neuronal cells)
- ❖ GLUT3 is specific for neurons with GLUT8 and 4 predominate on cell body and proximal dendrites respectively
- ❖ GLUT5 is localized in microglial cells (resident macrophages of the brain)

# Glucose Transporters



Cellular distribution of the principle glucose transporters in the nervous system.