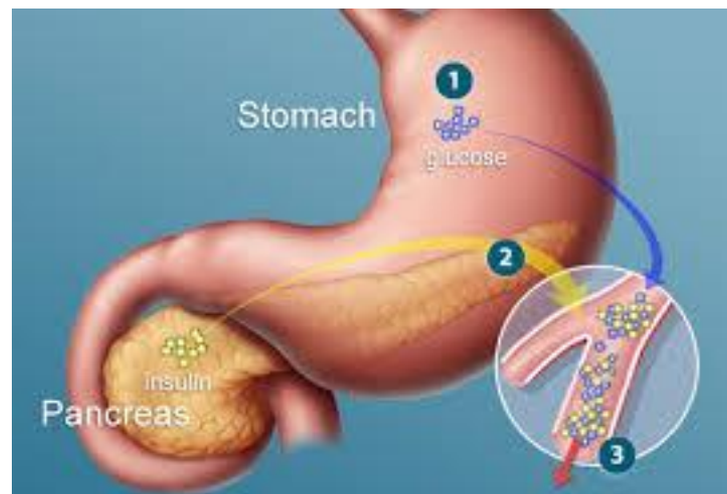




# Glycolysis I



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# Glucose as Energy Substrate



- To function properly, our cells are in need for energy which can be generated from the metabolism of various biomolecules such as carbohydrates, proteins and lipids
- Actually CHO particularly glucose is a major energy substrate in certain tissues like brain
- What are the metabolic pathways of glucose inside our cells?

# Glycolysis



- Glycolysis is the metabolic pathway which converts glucose (6C) into 2 pyruvate molecules (3C)
- It occurs in the cell cytosol
- Glycolysis takes place in nearly all organisms both aerobic and anaerobic (i.e. microorganisms live in O<sub>2</sub> free environments )
- Glycolysis is a sequence of ten oxygen-independent and enzyme-catalyzed steps
- The intermediates either provide entry points to the cycle or themselves directly useful (biosynthetic intermediates)

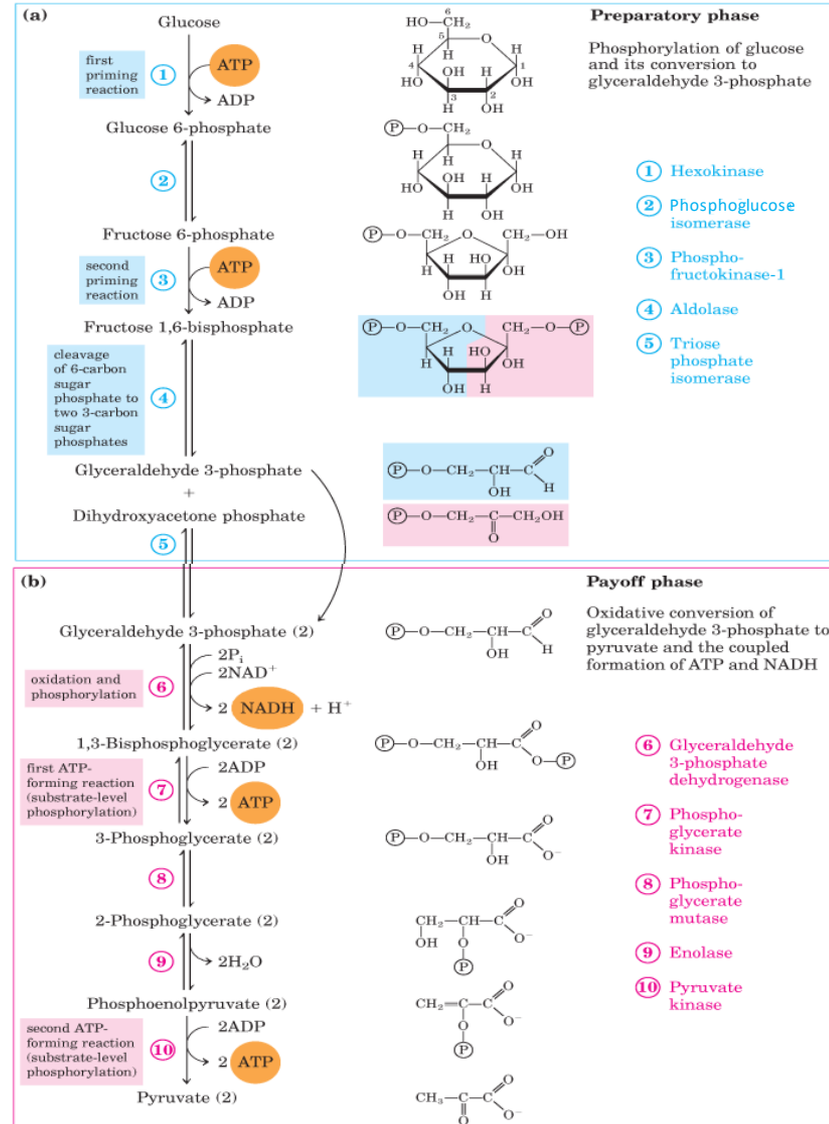
# Glycolysis



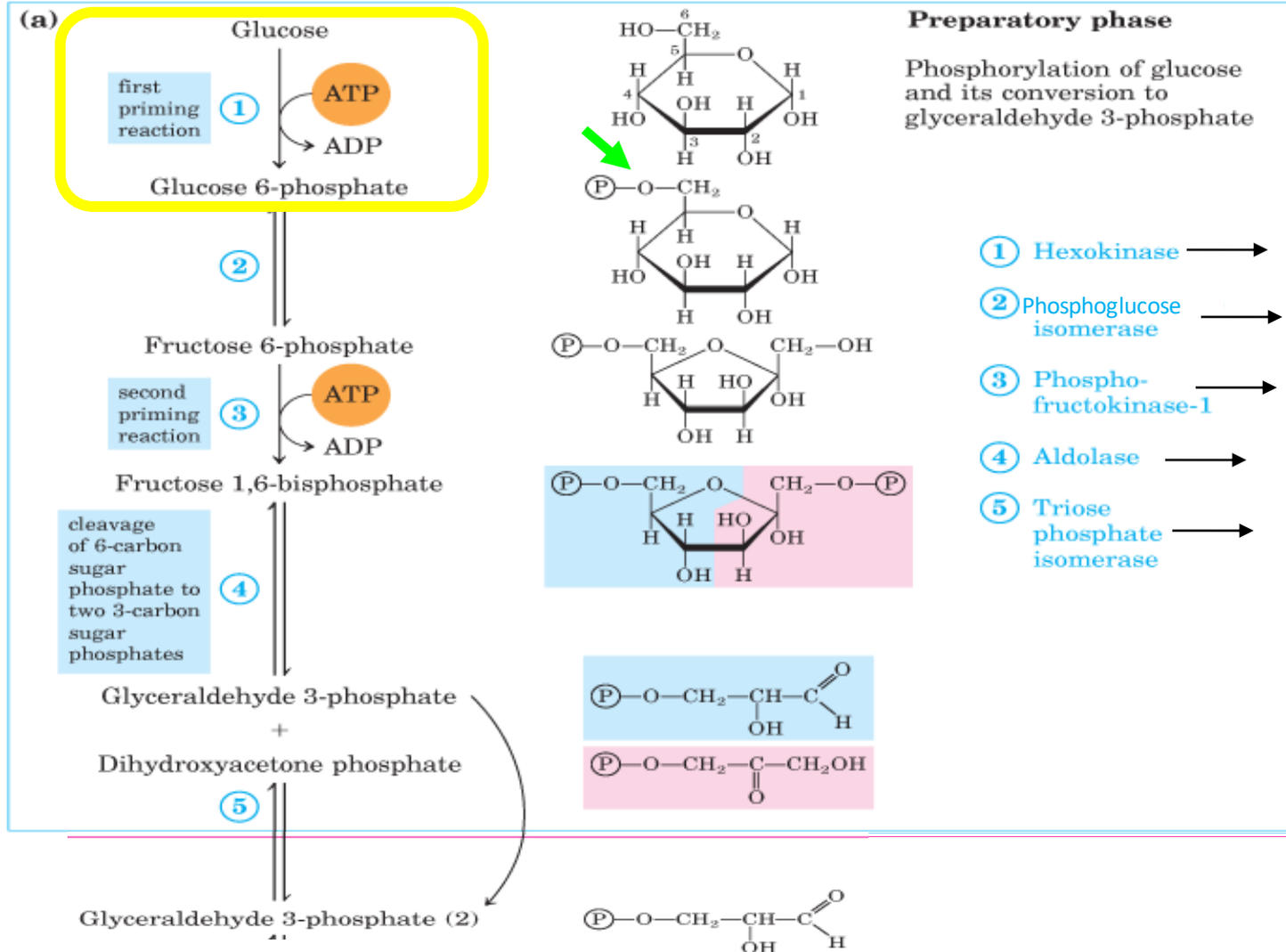
- The entire pathway is divided into two distinct phases:

## A. Energy Investment Phase (Preparatory Phase)

## B. Energy Generation Phase (Pay Off Phase)



# A. Preparatory Phase



# A. Preparatory Phase



- **Step 1:** Hexokinases catalyze the ATP- dependent phosphorylation of glucose to produce glucose-6-phosphate (G6P)
- Hexokinase is a transferase enzyme which phosphorylates hexoses by transferring an inorganic phosphate from ATP usually to hydroxyl O at C6
- Irreversible reaction (another enzyme catalyzes the dephosphorylation, only found in specific tissues). Therefore, it is a target site for cycle regulation
- This first priming reaction is important to maintain the influx of glucose through glucose transporters (**GLUTs**) and at the same time to trap the transported glucose molecules inside the cell

# Hexokinases



- 4 isoforms (isozymes) of hexokinase (I, II, III & IV) which differ in their **location**, **catalysis** and **regulation** thereby, contributing to different pattern of glucose metabolism in different tissues
- Hexokinase I, II & III are nonspecific and can phosphorylate a variety of hexoses (e.g. glucose, fructose, mannose) but type I is involved in **catabolic pathways** like glycolysis whereas type II & III are involved in **anabolic pathways** like glycogenesis
- Hexokinase IV is called glucokinase expressed in liver and pancreatic  $\beta$ -cells. It is specific for D-glucose

# Hexokinases

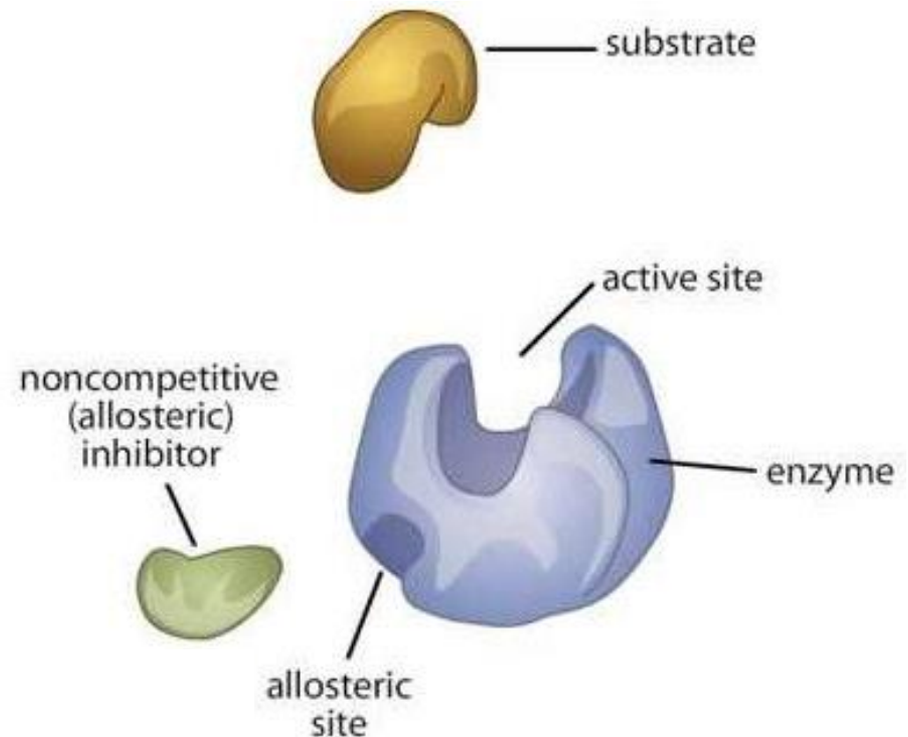


- Glucokinase has low affinity for glucose (high  $K_m$  value) compared to others (low  $K_m$  value)
- Therefore, glucokinase in **liver** is active only at high blood glucose level to accumulate G6P for glycogen synthesis but in the **pancreas** it acts as glucose sensor to control insulin release from beta cells
- Hexokinase isoforms (except isoform IV) are allosterically inhibited by G6P **only** at high level

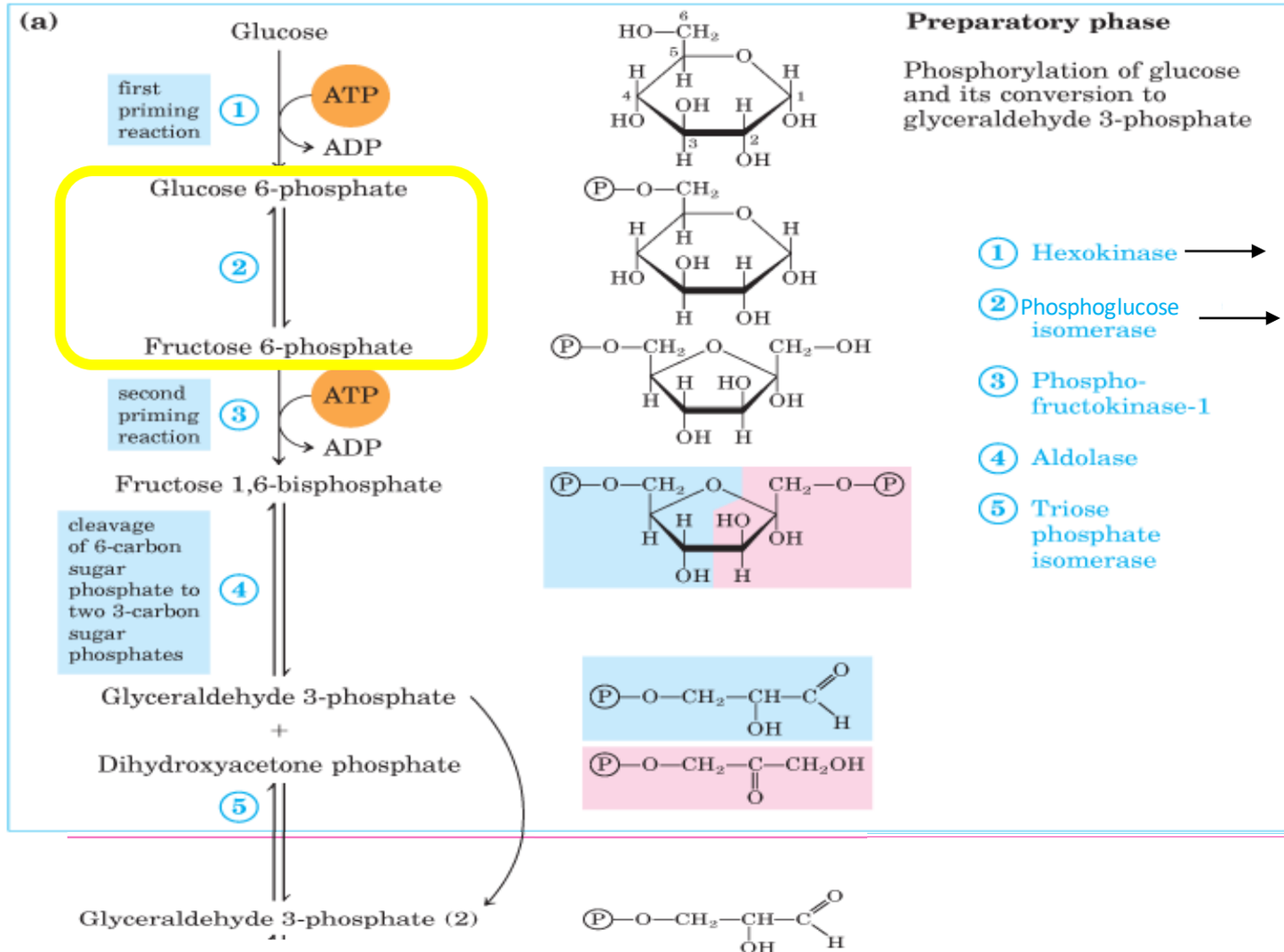
# Hexokinases



- Hexokinase is an allosteric enzyme with two binding sites: catalytic site (binds substrate) and regulatory site (allosteric site binds effectors)



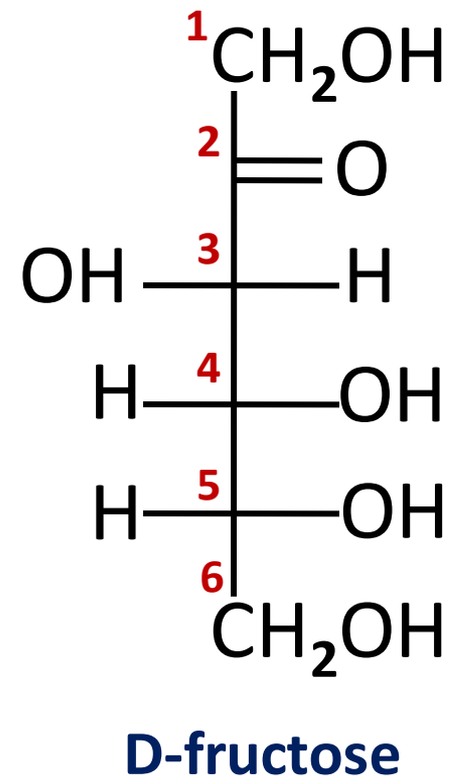
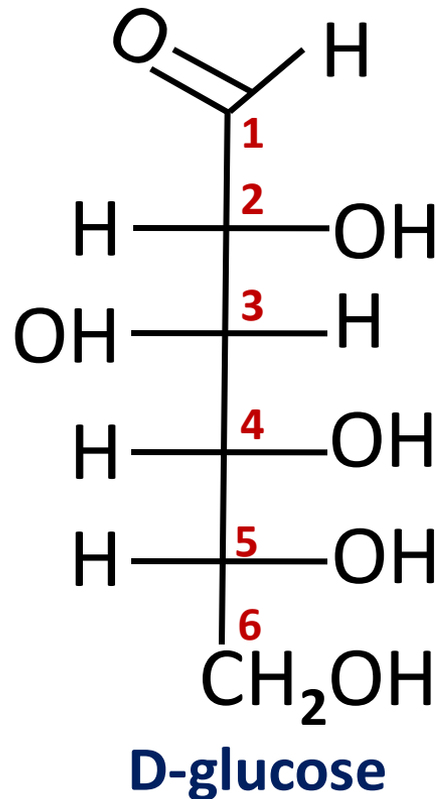
# A. Preparatory Phase



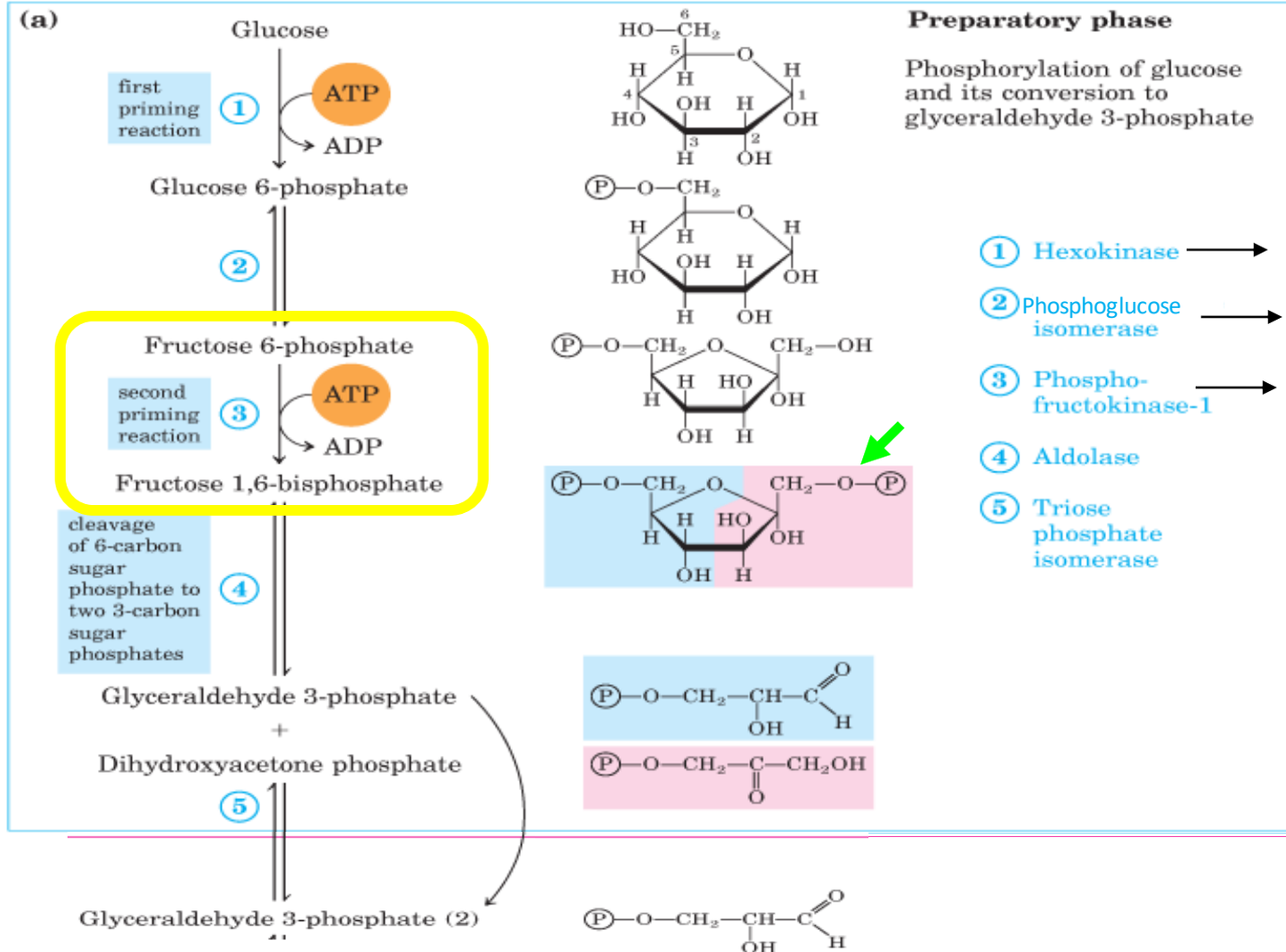
# A. Preparatory Phase



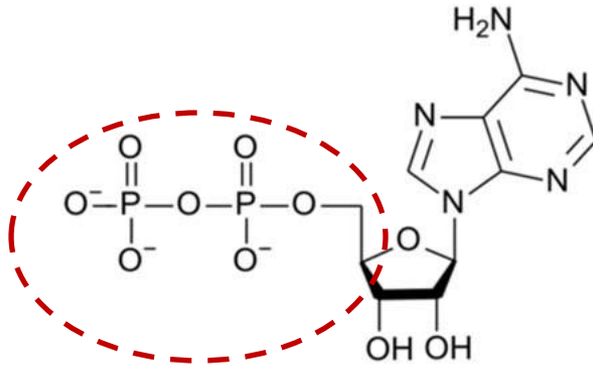
- **Step 2:** Phosphoglucose isomerase (PGI) interconverts G6P and F6P (reversible reaction).
- Indeed, **Mannose and Fructose** can enter the glycolysis pathway at this point



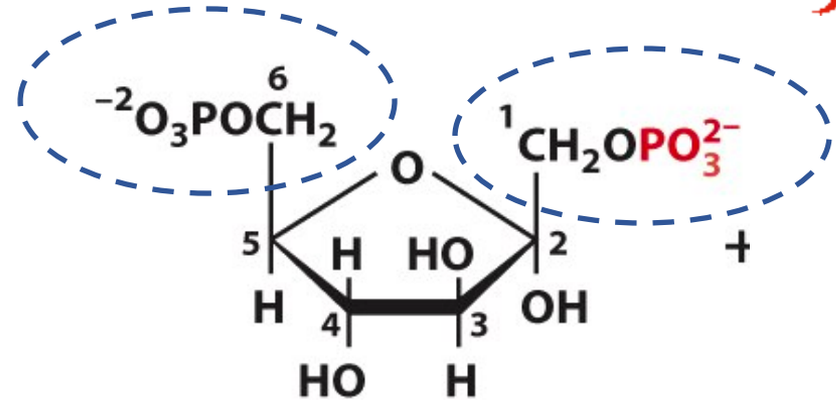
# A. Preparatory Phase



# A. Preparatory Phase

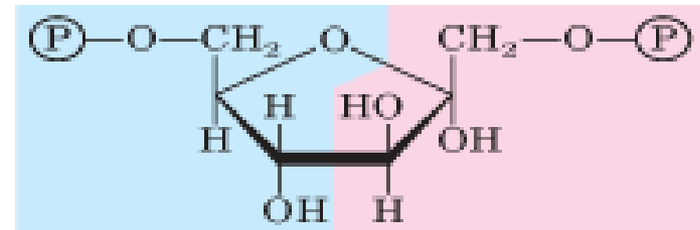
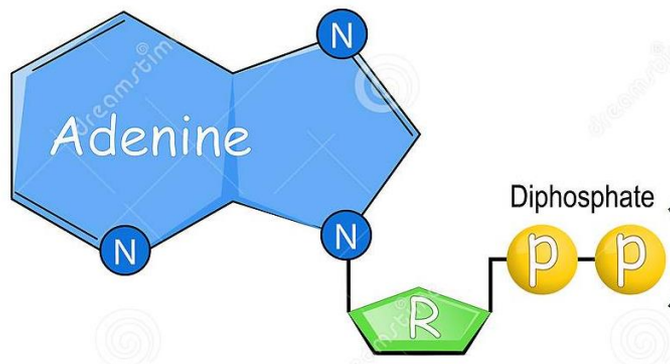


**ADP**



**Fructose 1,6 bisphosphate**

ADP (Adenosine diphosphate)

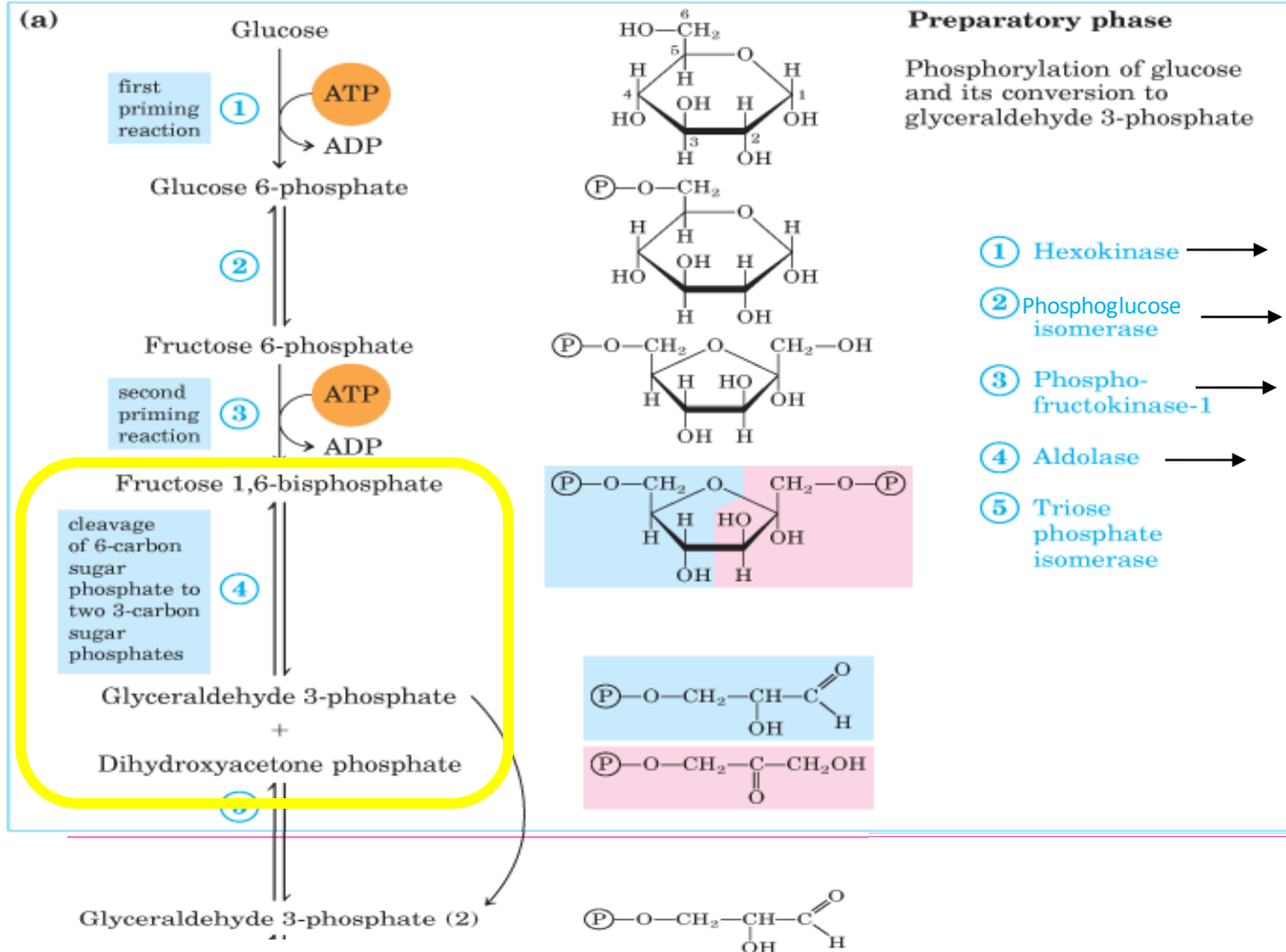


# A. Preparatory Phase



- **Step 3:** This is the **rate limiting** or key regulatory step. The activity of phosphofructokinase-1 (**PFK-1**) enzyme can be controlled. PFK-1 catalyzes the phosphorylation of hydroxyl oxygen at C1 to produce **fructose-1,6-bisphosphate**
- **Step 4:** Aldolase enzyme catalyzes the cleavage to two triose phosphates: **DHAP** (dihydroxyacetone phosphate) and **GAP** (glyceraldehyde-3-phosphate)
- The addition of the second phosphate group on C1 from the previous step **destabilizes** the hexose ring and facilitates the cleavage reaction

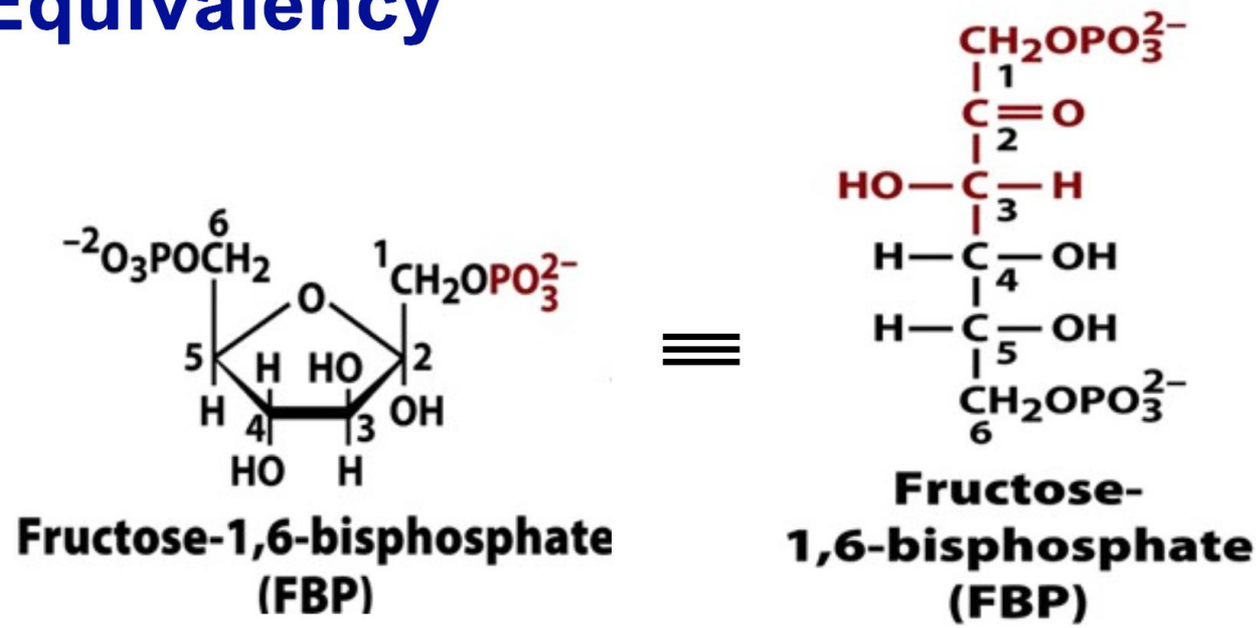
# A. Preparatory Phase



# Aldolase Mechanism of Action



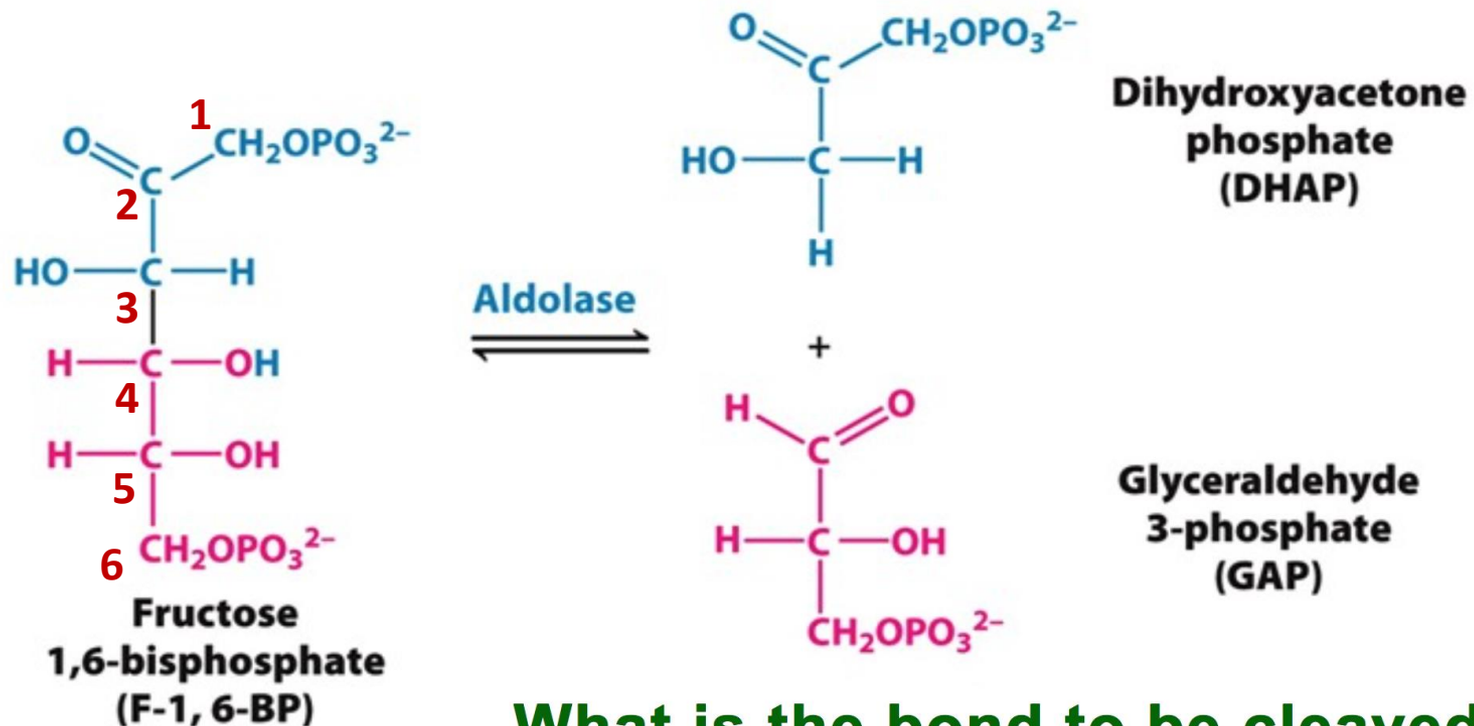
## Haworth and Fischer Projections Equivalency



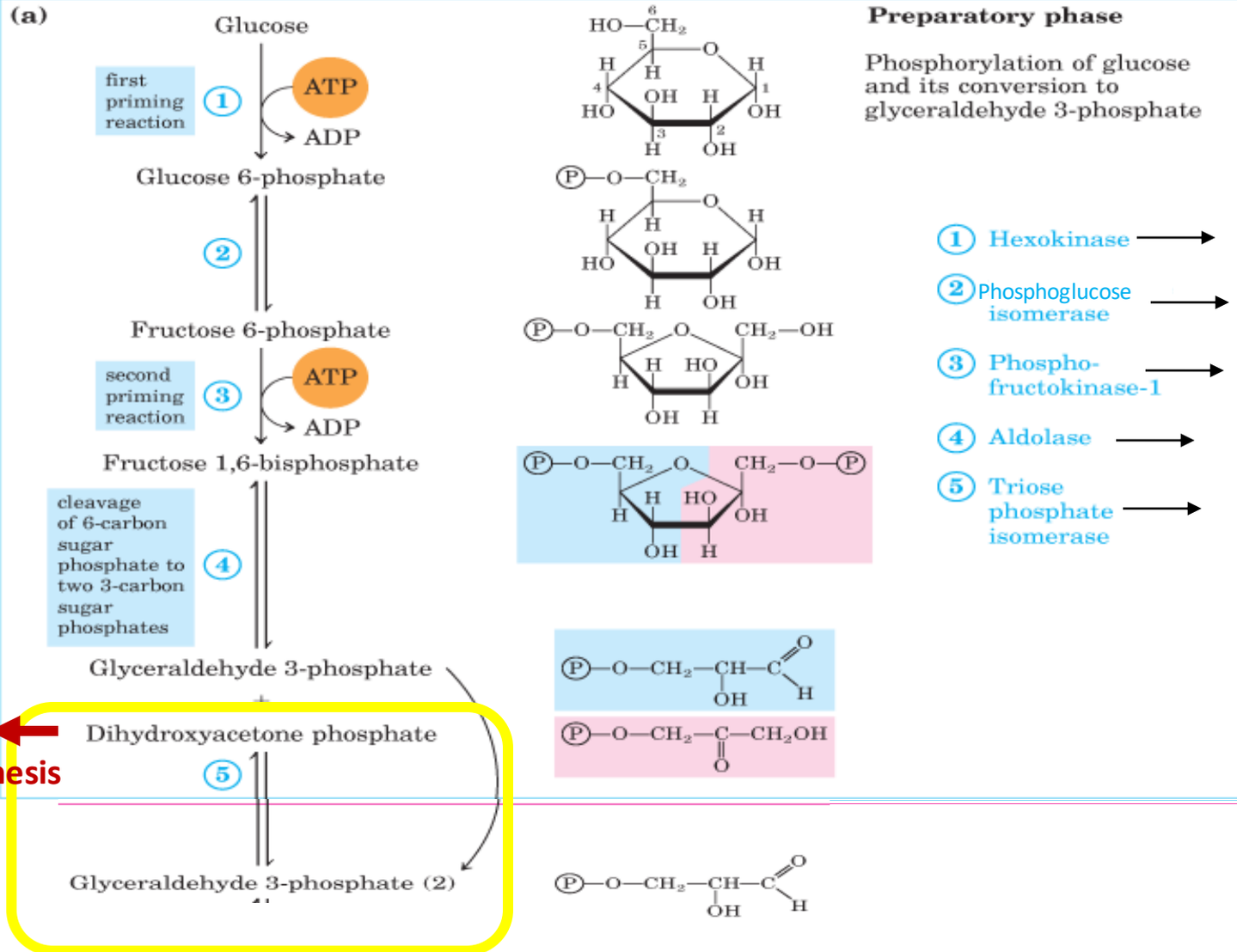
# Aldolase Mechanism of Action



## Six Carbon Sugar Cleaved to Two Three Carbon Units



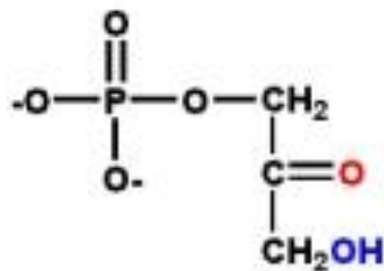
# A. Preparatory Phase



# A. Preparatory Phase



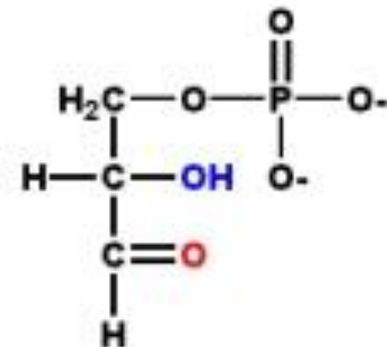
- **Step 5:** Isomerization of **DHAP** by triose phosphate isomerase (TPI) to **GAP** to proceed further in glycolysis as GAP is the substrate for the next reaction. This reaction is reversible



Dihydroxy acetone  
phosphate

**DHAP**

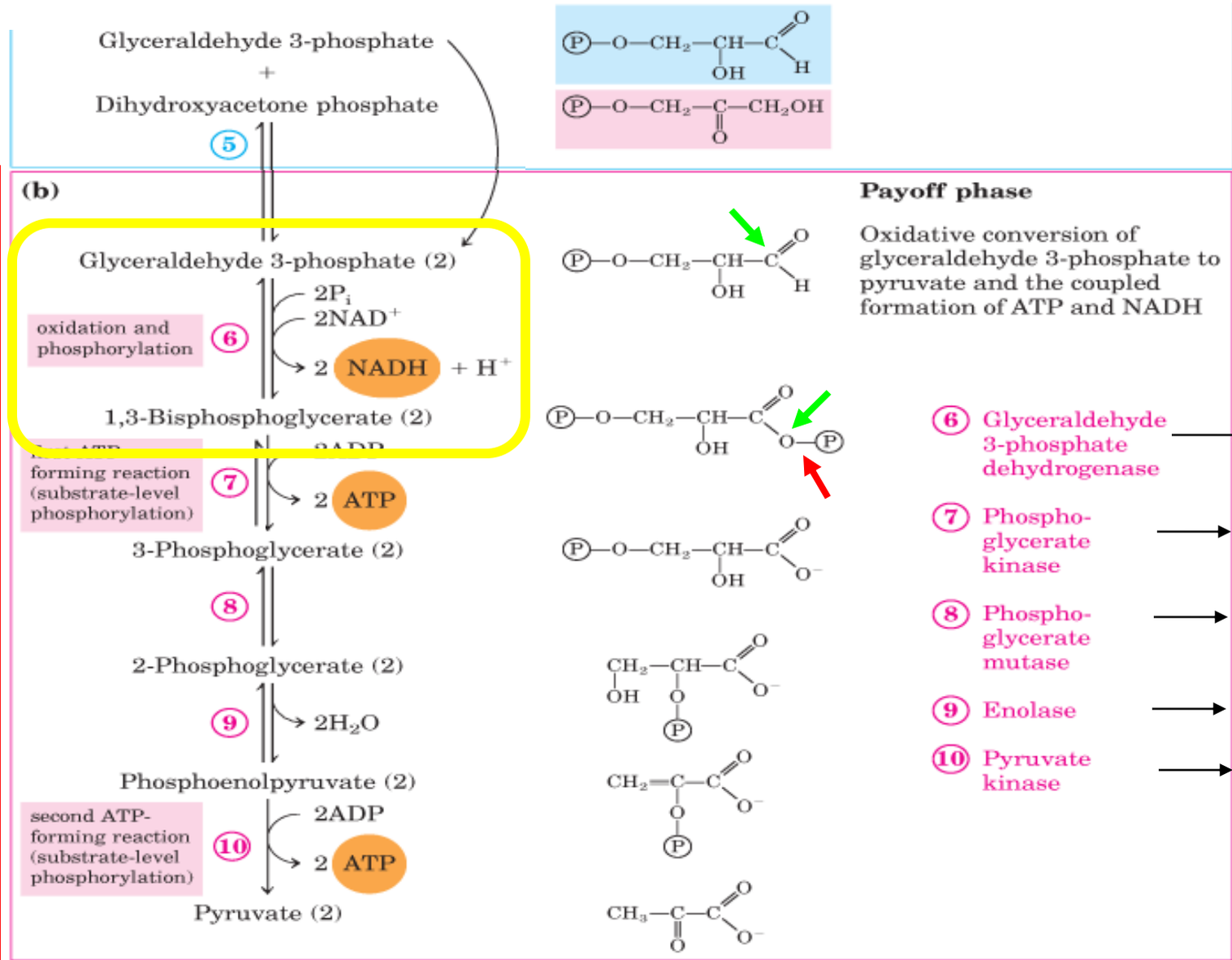
*Triose phosphate  
isomerase*



D-glyceraldehyde  
3-phosphate

**GAP**

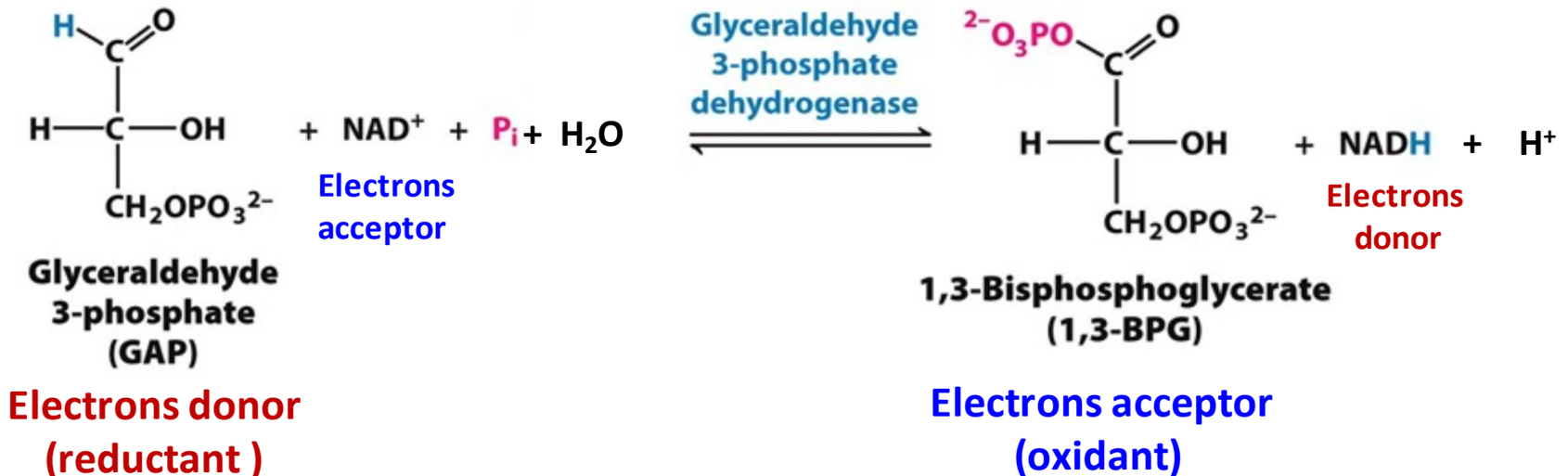
# B. Pay Off Phase



# B. Pay Off Phase



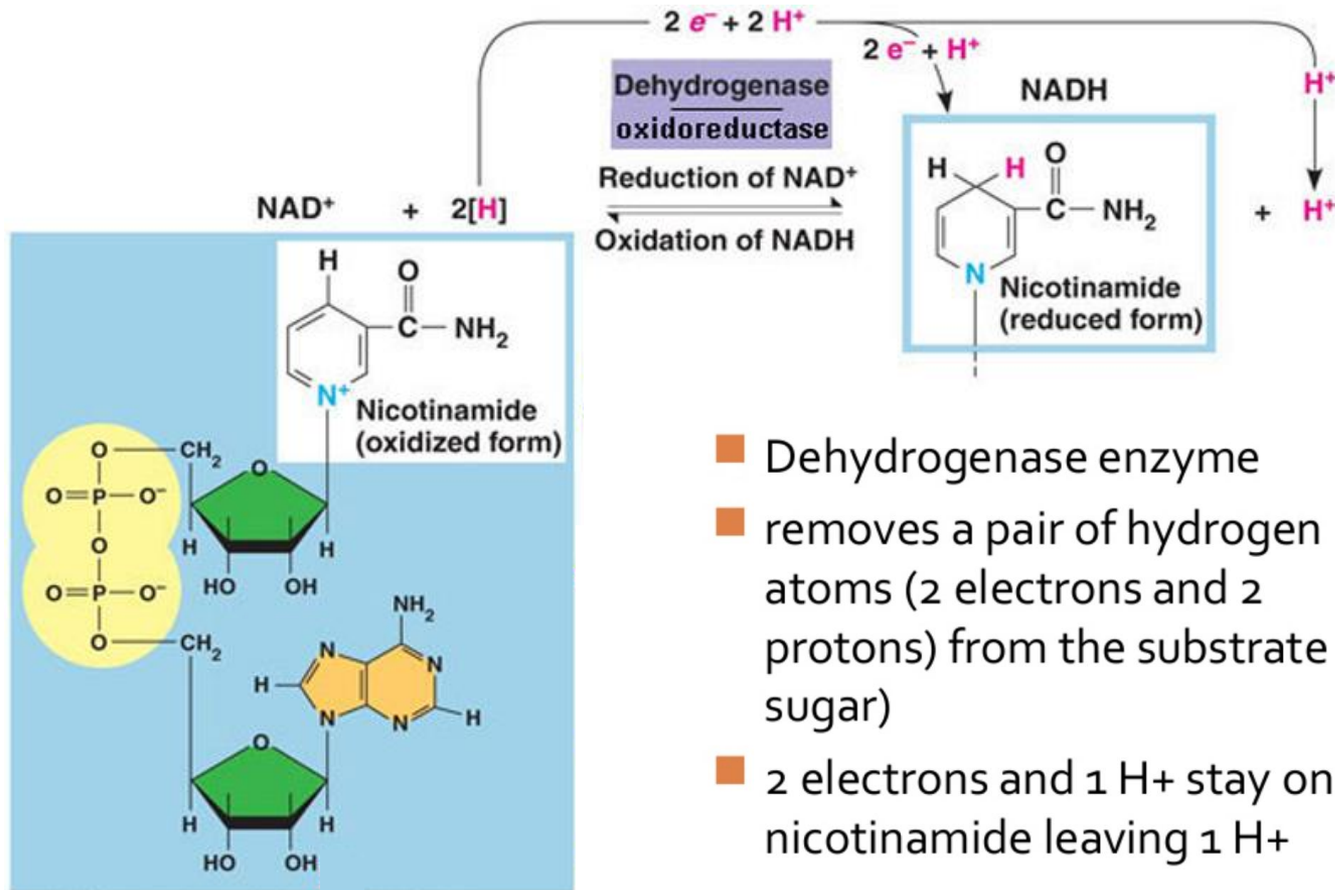
- **Step 6:** GAP dehydrogenase enzyme catalyzes the oxidative phosphorylation of GAP (electron donor) into super-high-energy compound (**1,3-BPG**) and the transfer of electrons into the coenzyme  $\text{NAD}^+$  (electron acceptor) forming **NADH**
- **Dehydrogenases** are named as electrons donor substrate -dehydrogenase





# Nicotinamide Adenine Dinucleotide

- **NAD (Nicotinamide adenine dinucleotide)** is a coenzyme which exists in two forms: NADH (the reduced form) and NAD<sup>+</sup> (the oxidized form)



# Nicotinamide Adenine Dinucleotide

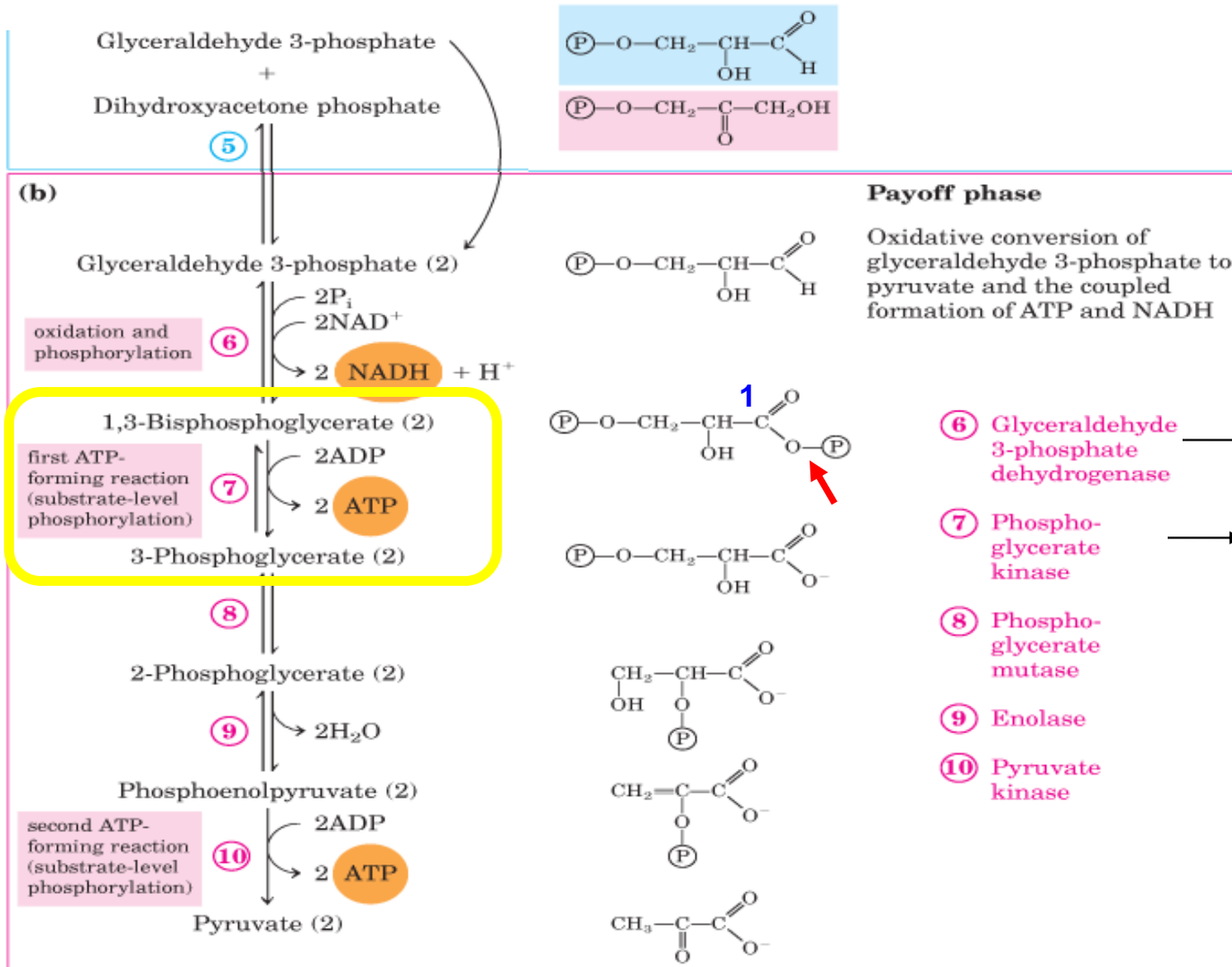


- **NAD (Nicotinamide adenine dinucleotide)** is a coenzyme of dehydrogenases
- The reduced form NADH is electrons carrier and it is called **energy rich molecule**. It is an indirect form of energy

1 NADH = 2.5 ATP



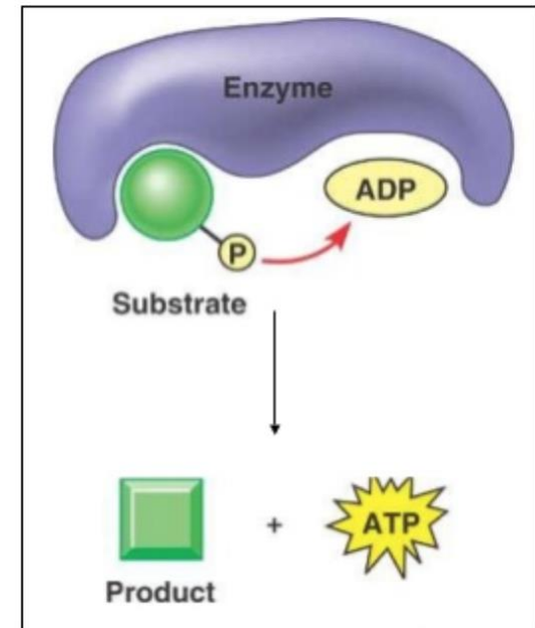
# B. Pay Off Phase



## B. Pay Off Phase

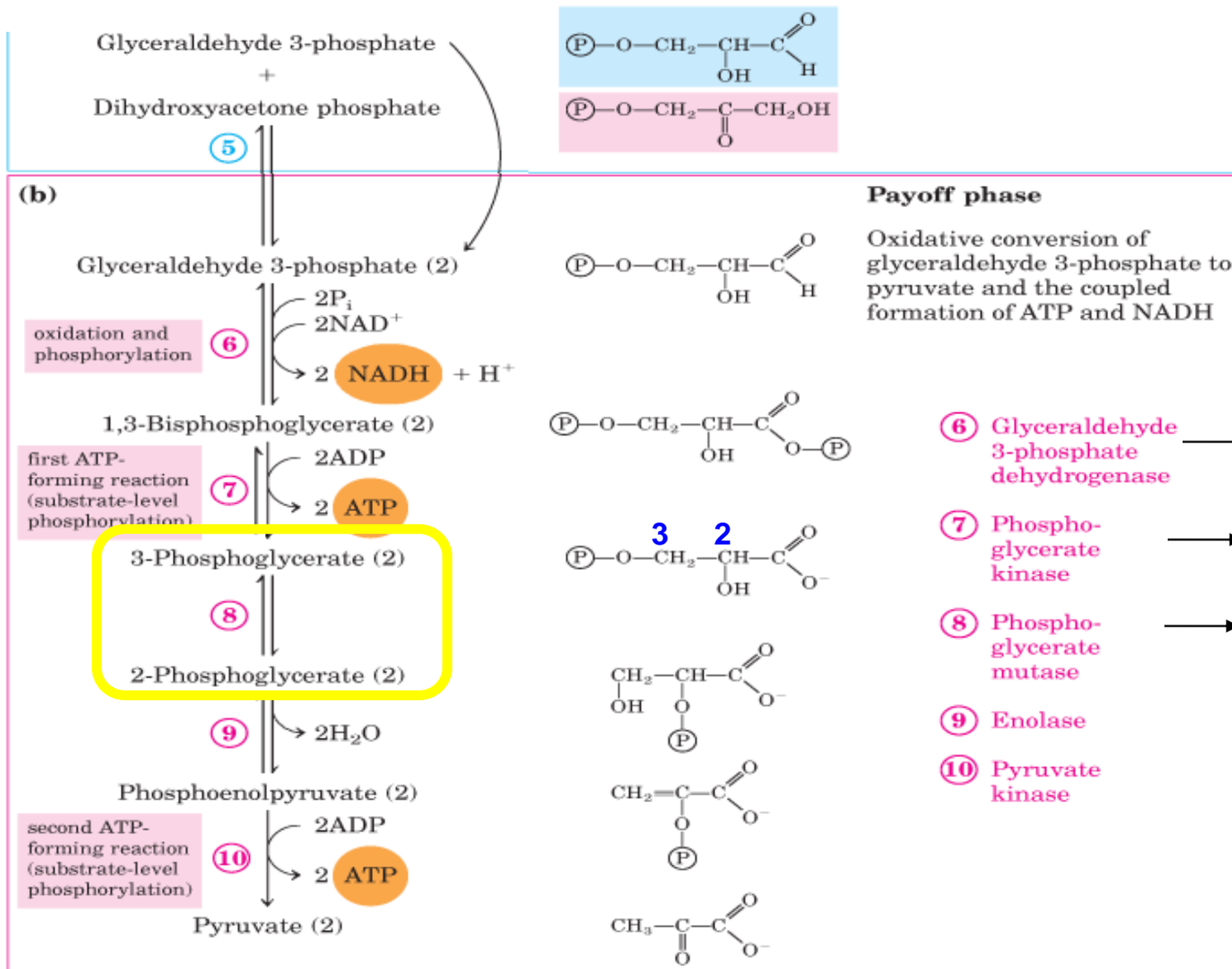


- **Step 7:** The **first ATP** molecule is generated by the substrate-level phosphorylation process catalyzed by phosphoglycerate kinase (PGK)
- **2 ATP molecules** will be generated in this step
- **Methods of ATP synthesis:**
  1. Substrate-level phosphorylation: it is a direct method of ATP synthesis by an enzyme which catalyzes the transfer of phosphate group from substrate to ADP
  2. Oxidative phosphorylation: indirect method of ATP synthesis in which the oxidation of NADH/FADH<sub>2</sub> and the subsequently transferred electrons indirectly drive ATP synthesis from ADP



An enzyme transfers phosphate from substrate to ADP

# B. Pay Off Phase

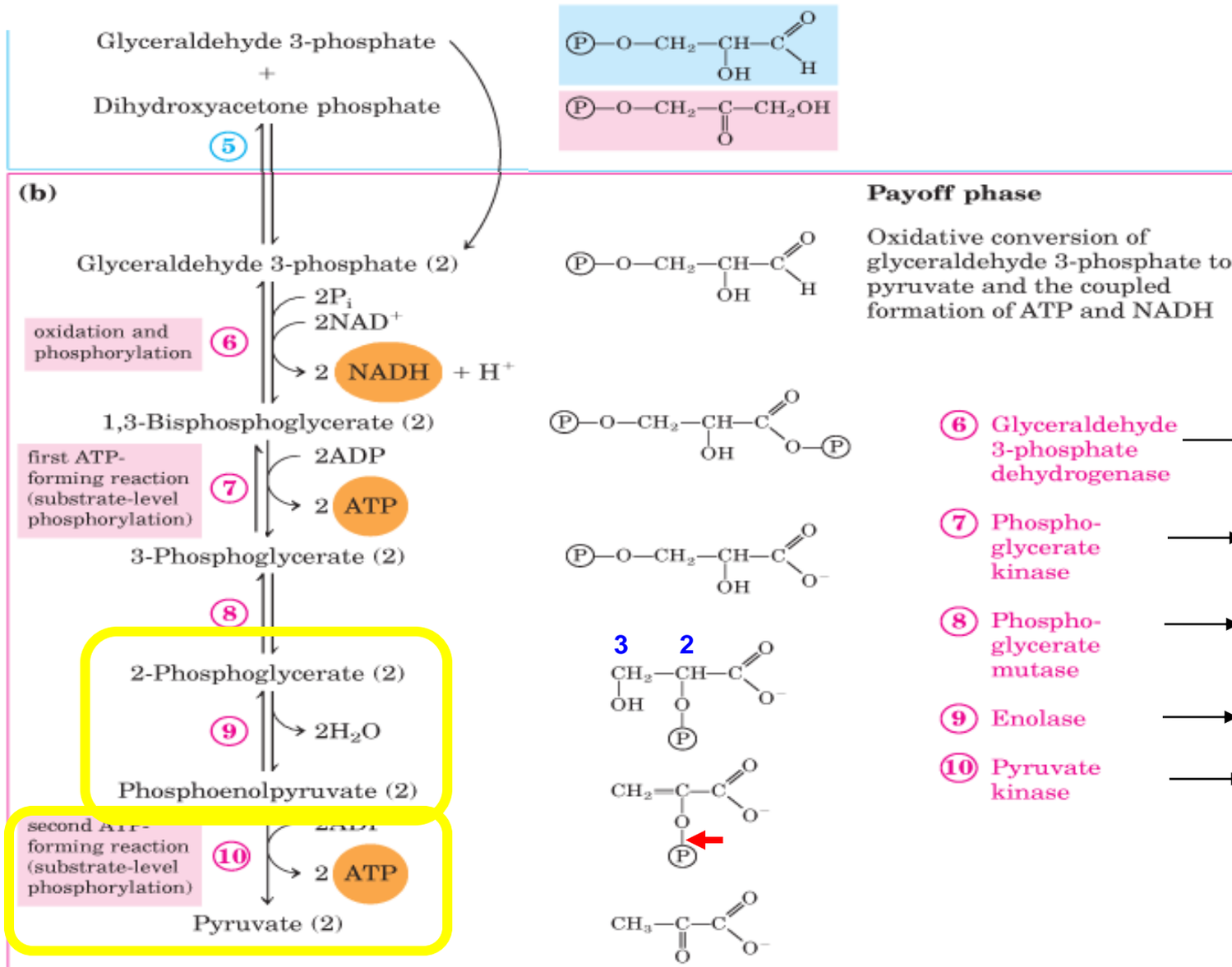


## B. Pay Off Phase



- **Step 8:** Phosphoglycerate mutase (PGM) is an **isomerase** which catalyzes the isomerization of 3-phosphoglycerate to 2-phosphoglycerate
- It is actually an **internal shifting of P** group from C3 to C2 within the same molecule
- The main purpose of this step is the **activation of the phosphate group** to prepare for the generation of the second ATP in the next reactions
- **Step 9:** The synthesis of the second super-high-energy compound **phosphoenolpyruvate (PEP)** in a simple dehydration reaction catalyzed by enolase enzyme
- Enolase acts by catalyzing the cleavage of bond between C3 and oxygen of OH group thus enhancing the formation of double bond between C3 & C2 and subsequently H atom elimination from C2

# B. Pay Off Phase



## B. Pay Off Phase



- The aim of this step is to increase the energy stored in the phosphate bond
- **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
- The activity of pyruvate kinase can be controlled (irreversible reaction) so this reaction is regulatory step
- The net result of glycolysis is the formation of:
  - 2 pyruvate
  - 2 ATP
  - 2 NADH