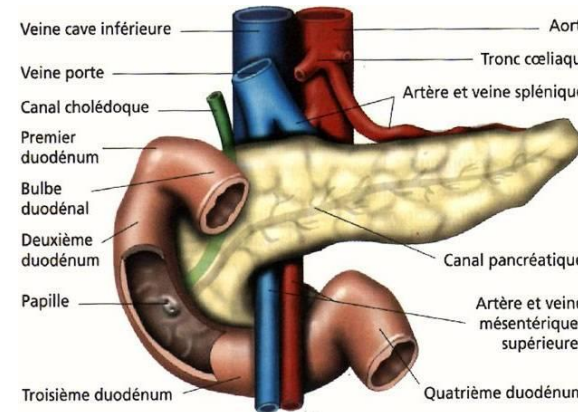




Endocrine Physiology

The Endocrine Pancreas

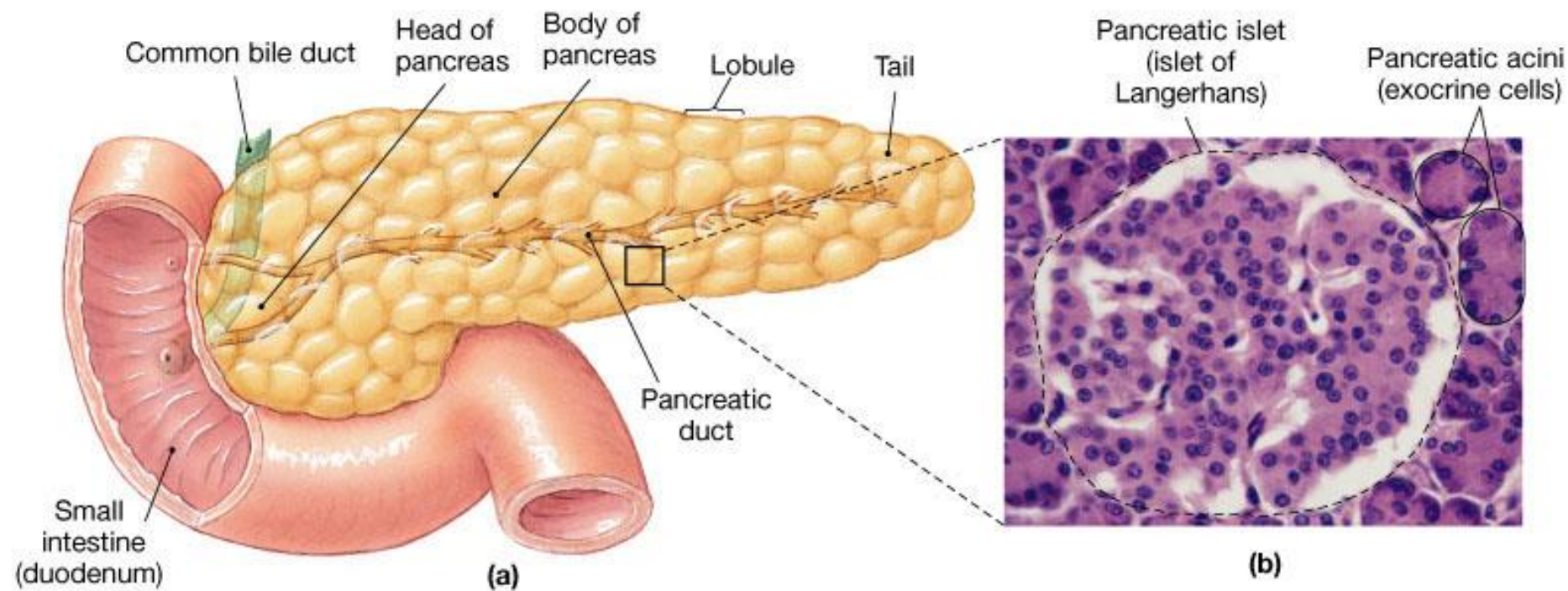
Dr. Khalid Al-Regaiey



Pancreas

- A triangular gland, which has both exocrine and endocrine cells, located behind the stomach
- Strategic location
- Acinar cells produce an enzyme-rich juice used for digestion (exocrine product)
- Pancreatic islets (**islets of Langerhans**) produce hormones involved in regulating fuel storage and use.

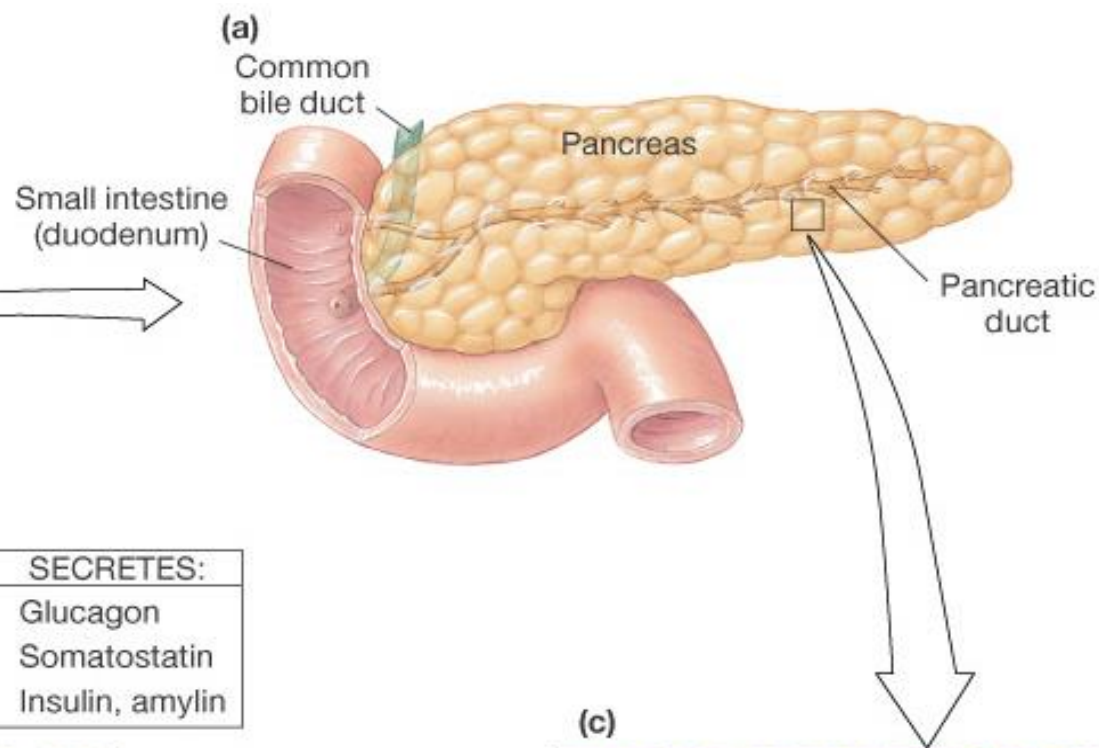
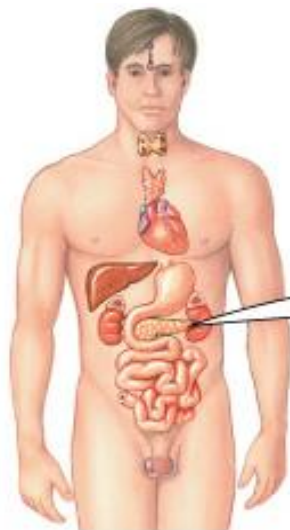
The Endocrine Pancreas



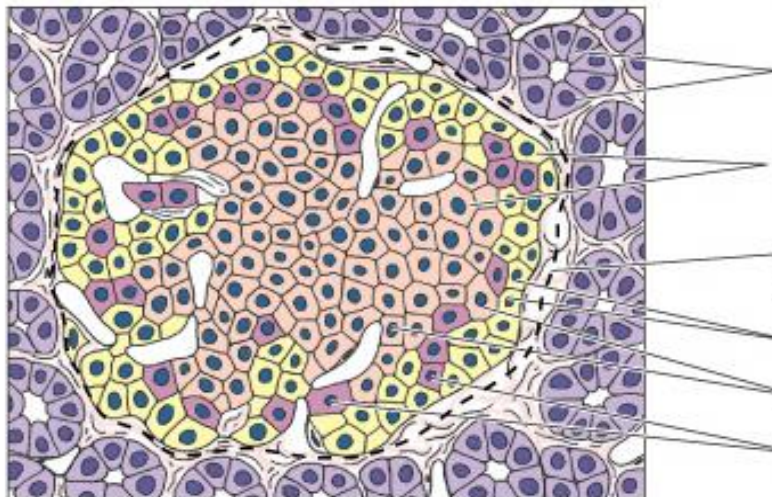
Islets of Langerhans

- 1-2 million islets
- Beta (β) cells produce insulin (60%)
- Alpha (α) cells produce glucagon (25%)
- Delta (δ) cells produce somatostatin (10%)
- F cells produce pancreatic polypeptide (5%)

Islets of Langerhans

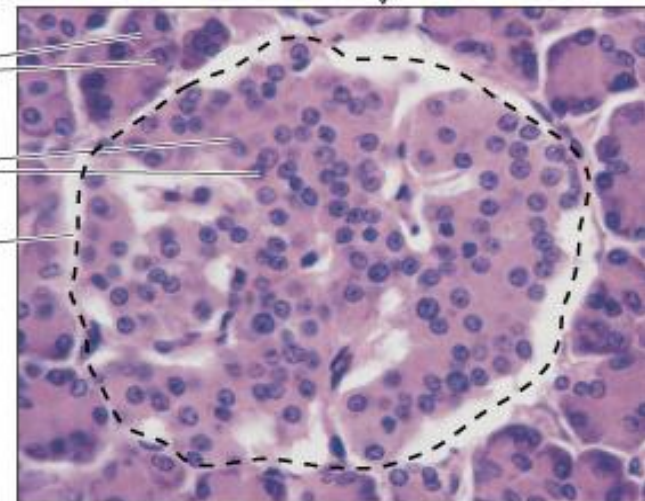


(b)



Exocrine cells
Endocrine cells
Islet of Langerhans
Alpha cells
Beta cells
D cells

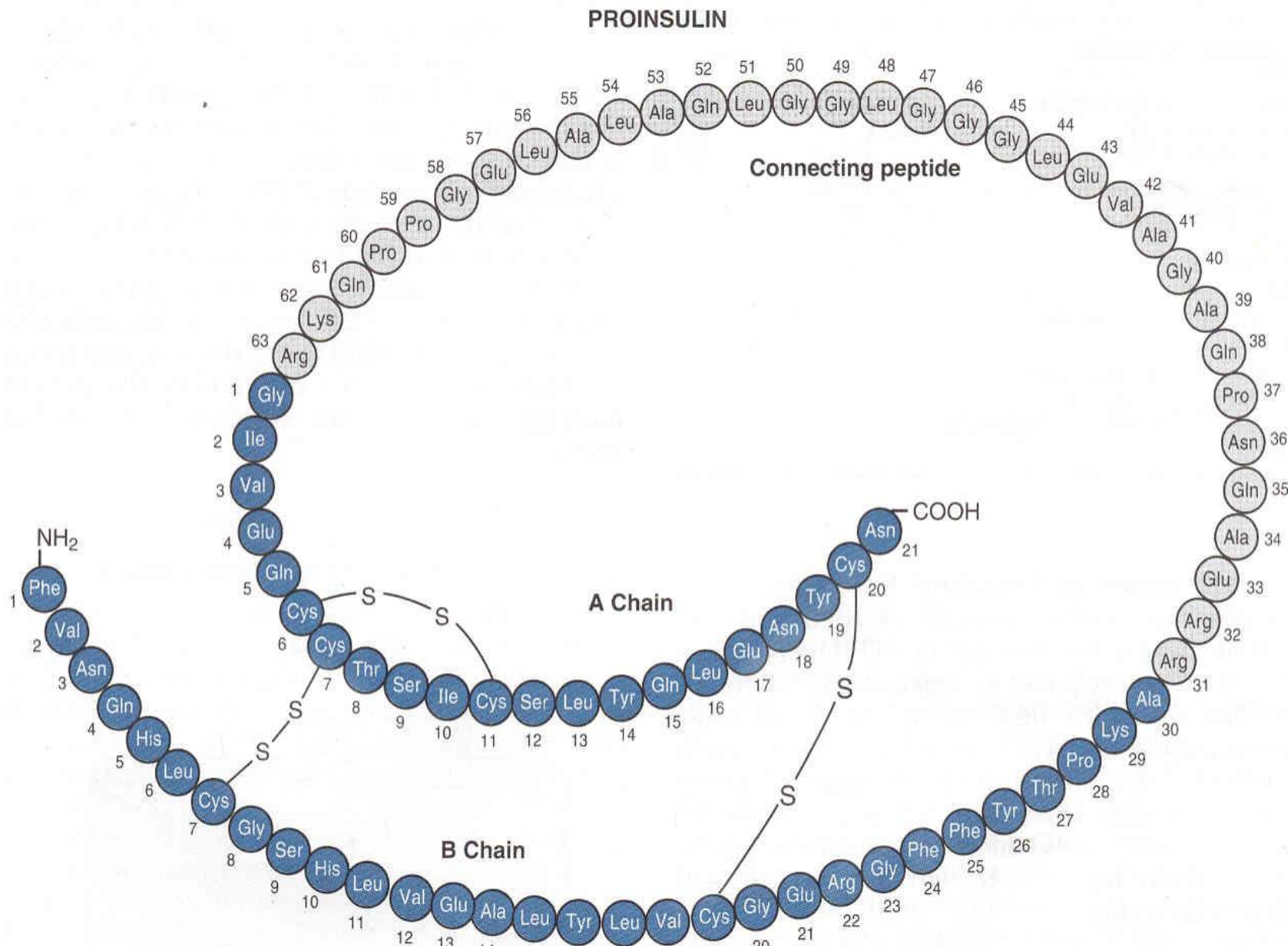
(c)



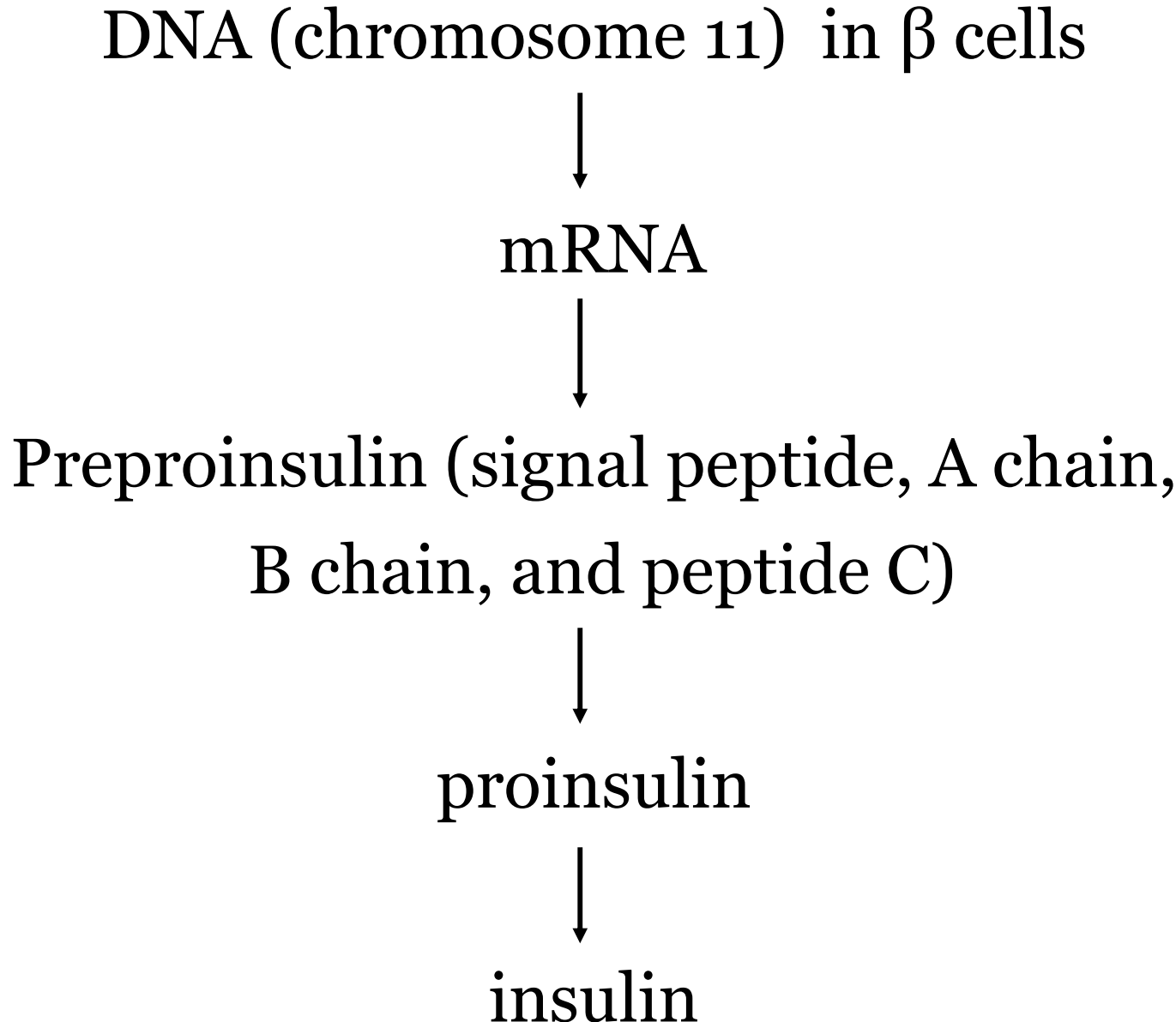
Insulin

- Hormone of nutrient abundance
- A protein hormone consisting of two amino acid chains linked by disulfide bonds
- Synthesized as part of proinsulin (86 AA) and then excised by enzymes, releasing functional insulin (51 AA) and C peptide (29 AA).
- Has a plasma half-life of 6 minutes.

Insulin Structure



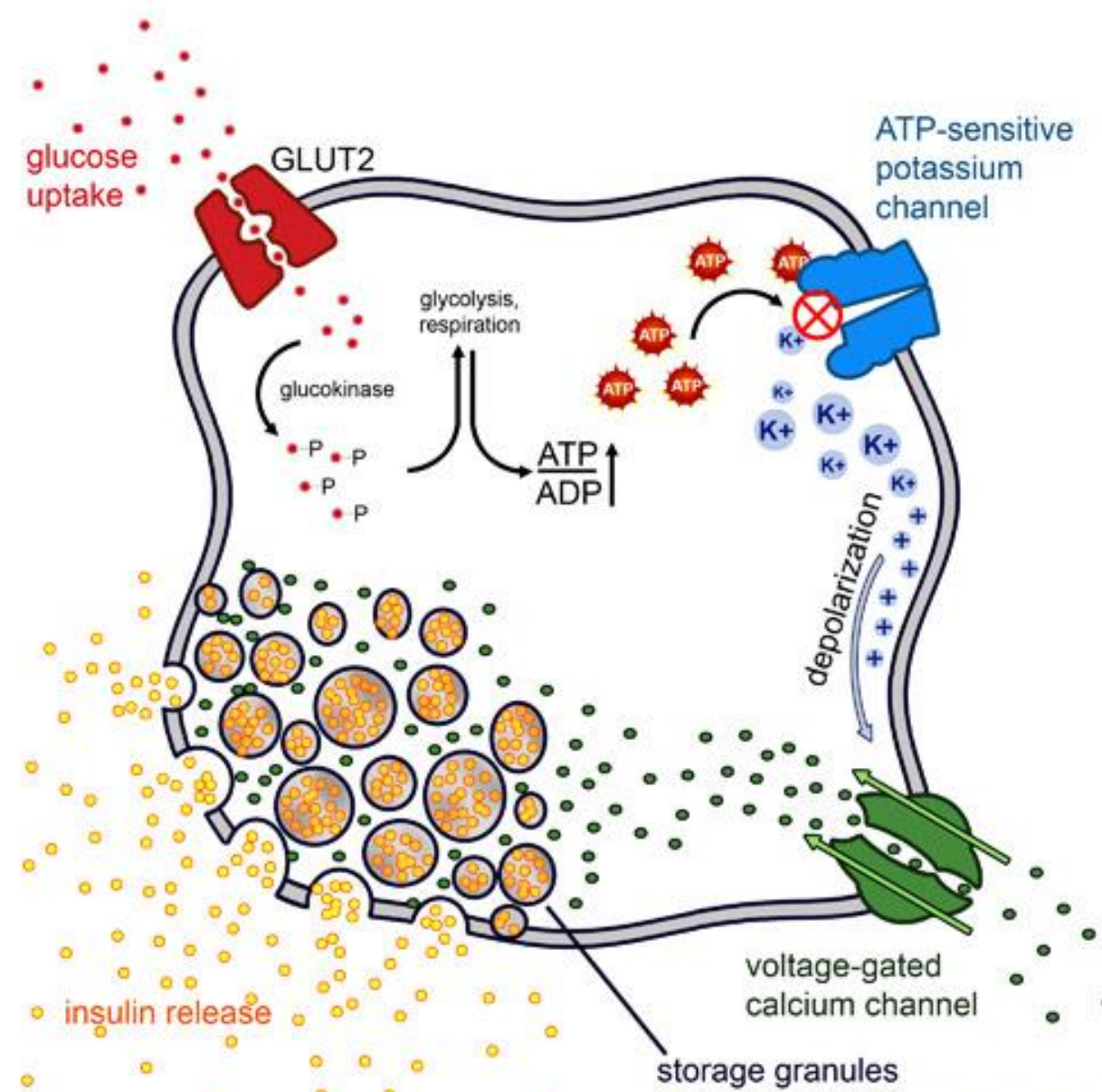
Insulin Synthesis



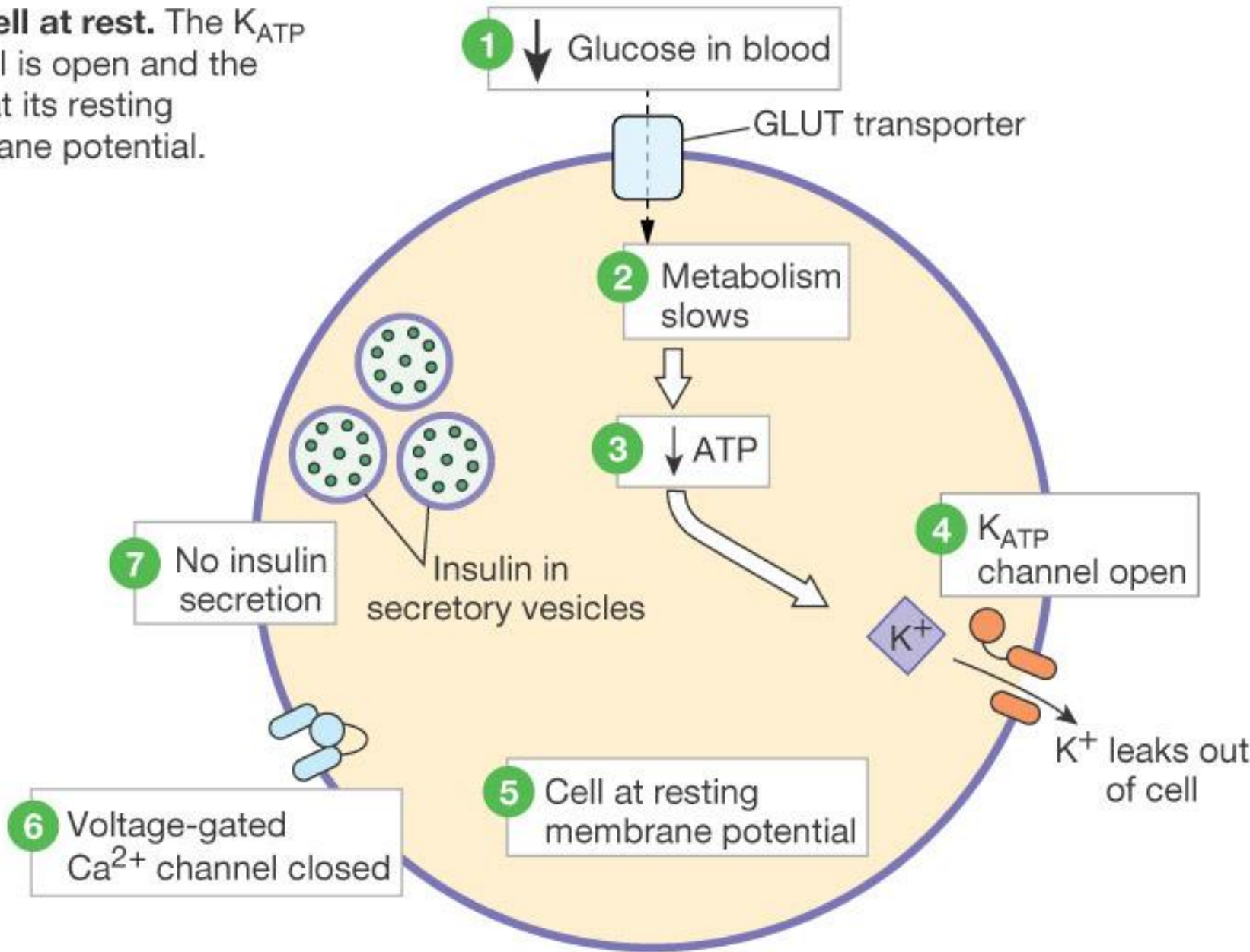
Insulin Synthesis

- Insulin synthesis is stimulated by glucose or feeding and decreased by fasting
- Threshold of glucose-stimulated insulin secretion is 100 mg/dl.
- Glucose rapidly increase the translation of the insulin mRNA and slowly increases transcription of the insulin gene

Glucose is the primary stimulator of insulin secretion

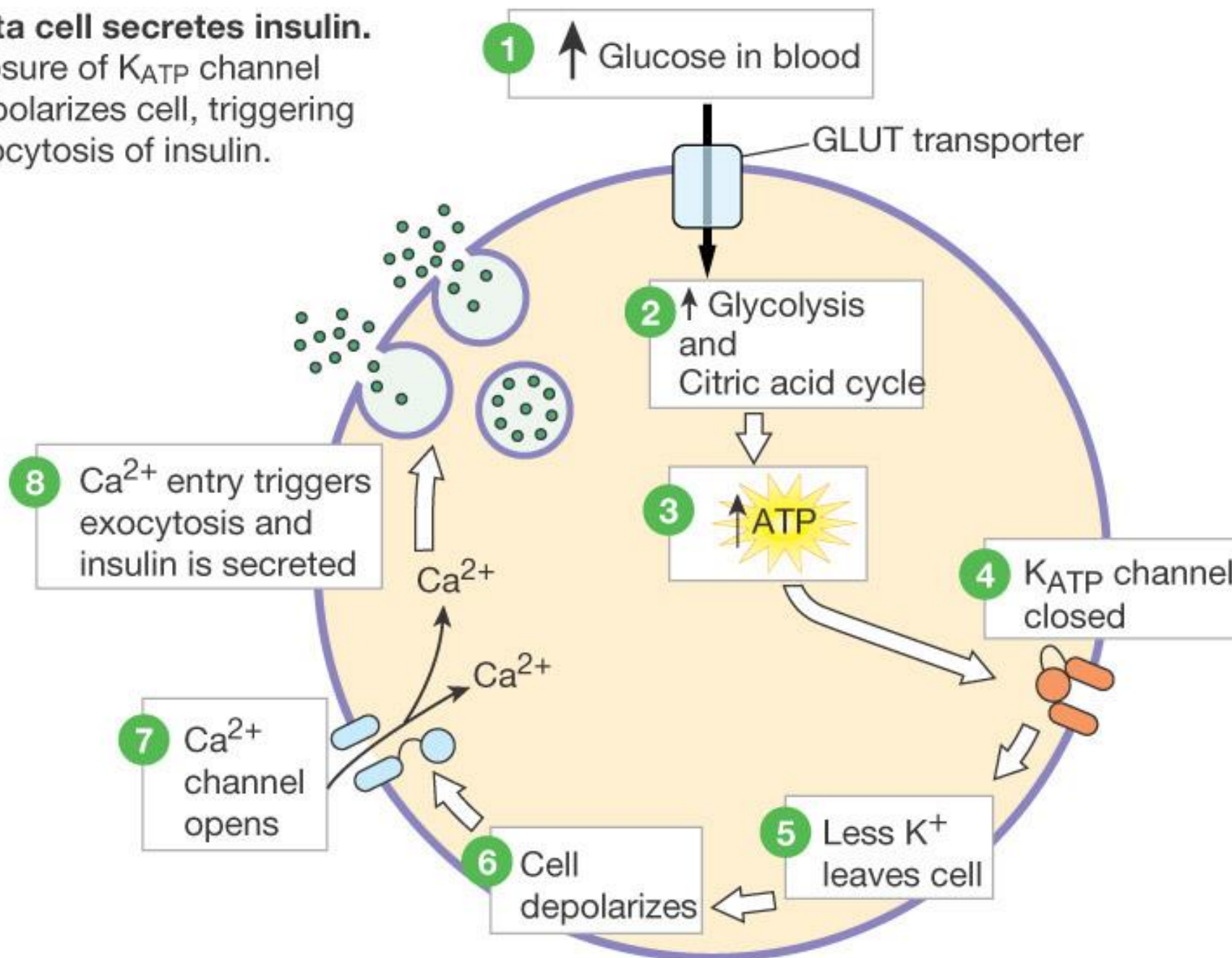


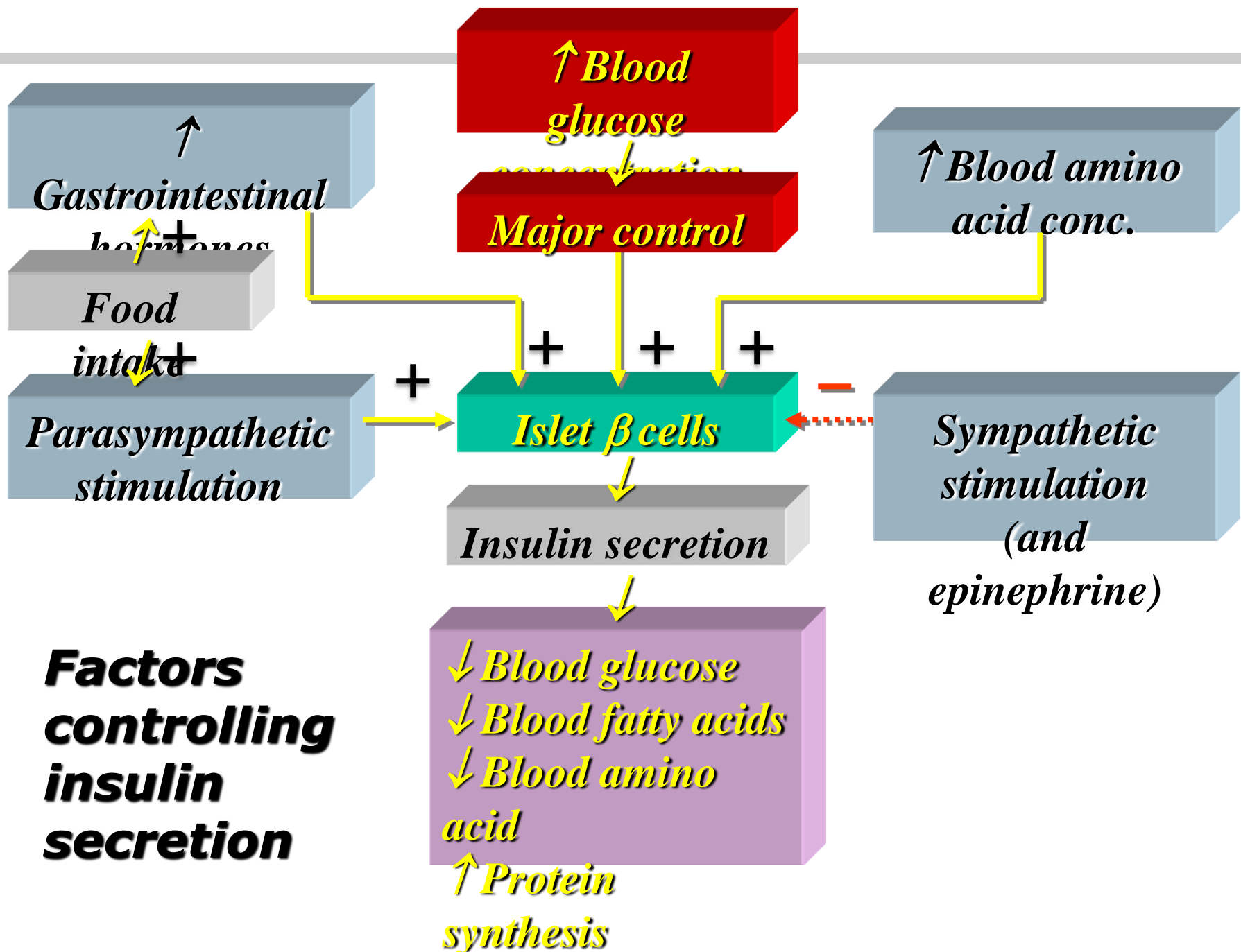
(a) Beta cell at rest. The K_{ATP} channel is open and the cell is at its resting membrane potential.



(b) Beta cell secretes insulin.

Closure of K_{ATP} channel depolarizes cell, triggering exocytosis of insulin.





Regulation of Insulin Secretion

Regulators of insulin secretion

Stimulators of insulin secretion

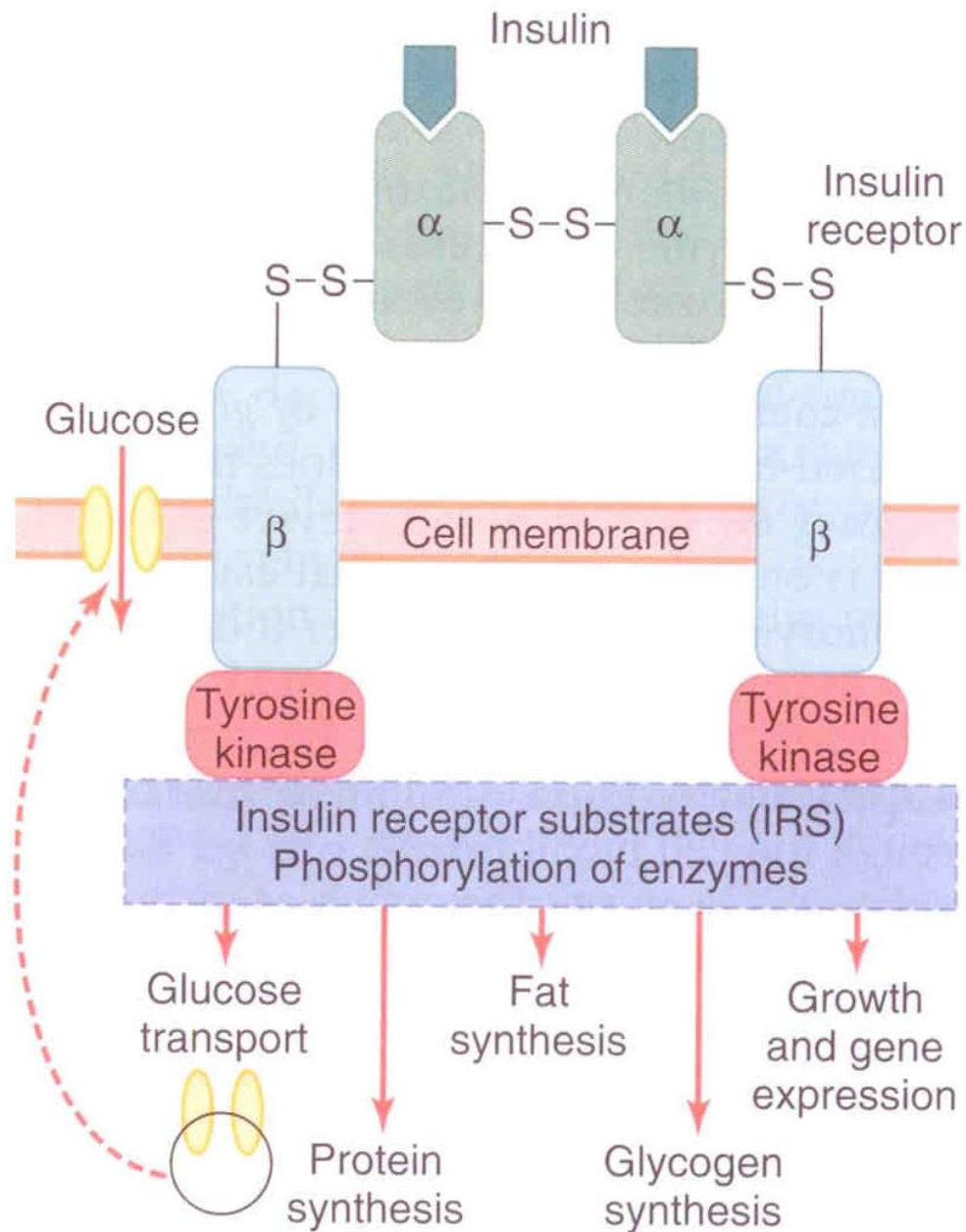
- ↑ Serum glucose
- ↑ Serum amino acids
- ↑ Serum free fatty acids
- ↑ Serum ketone bodies
- Hormones
 - Gastroinhibitory peptide (GIP)
 - Glucagon
 - Gastrin
 - Cholecystokinin (CCK)
 - Secretin
 - Vasoactive intestinal peptide (VIP)
 - Epinephrine (β -receptor)
- Parasympathetic nervous system

Inhibitors of insulin secretion

- ↓ Glucose
- ↓ Amino acids
- ↓ Free fatty acids
- Hormones
 - Somatostatin
 - Epinephrine (α -receptor)
- Sympathetic nervous system stimulation

Insulin Receptor

- the insulin receptor is a transmembrane receptor
- belongs to the large class of **tyrosine kinase receptors**
- Made of two alpha subunits and two beta subunits



Actions of insulin

Glucose regulation and metabolism terms

- **Gluconeogenesis** - Synthesis of glucose from noncarbohydrate precursors, Lactic acid, glycerol, amino acids, liver cells synthesis glucose when carbohydrates are depleted.
- **Glycogenesis** - Formation of glycogen, glucose stored in liver and skeletal muscle as glycogen, important energy reserve.
- **Glycogenolysis** – breakdown of glycogen (polysaccharide) into glucose molecules (monosaccharide)
- **Glycolysis** - the breakdown of glucose into pyruvate by cells for the production of ATP

- **Raapid (seconds)**
- (+) transport of glucose, amino acids, K⁺ into insulin-sensitive cells
- **Intermediate (minutes)**
- (+) protein synthesis
- (-) protein degradation
- (+) of glycolytic enzymes and glycogen synthase
- (-) phosphorylase and gluconeogenic enzymes
- **Delayed (hours)**
- (+) mRNAs for lipogenic and other enzymes

Action of insulin on Adipose tissue

- (+) glucose entry
- (+) fatty acid synthesis
- (+) glycerol phosphatase synthesis
- (+) triglyceride deposition
- (+) lipoprotein lipase
- (-) of hormone-sensitive lipase
- (+) K uptake

Action of insulin on Muscle:

- (+) glucose entry
- (+) glycogen synthesis
- (+) amino acid uptake
- (+) protein synthesis in ribosomes
- (-) protein catabolism
- (-) release of gluconeogenic amino acids
- (+) ketone uptake
- (+) K uptake

Action of insulin on Liver:

- (-) ketogenesis
- (+) protein synthesis
- (+) lipid synthesis
- (-) gluconogenesis, (+) glycogen synthesis, (+) glycolysis.

General

- (+) cell growth

Glucose Transport

- GLUT1 (erythrocytes, brain)
- GLUT2 (liver, pancreas, small intestines, kidney)
- GLUT3 (brain)
- **GLUT4**, insulin sensitive transporter (muscle, adipose tissue)

Insulin: Summary

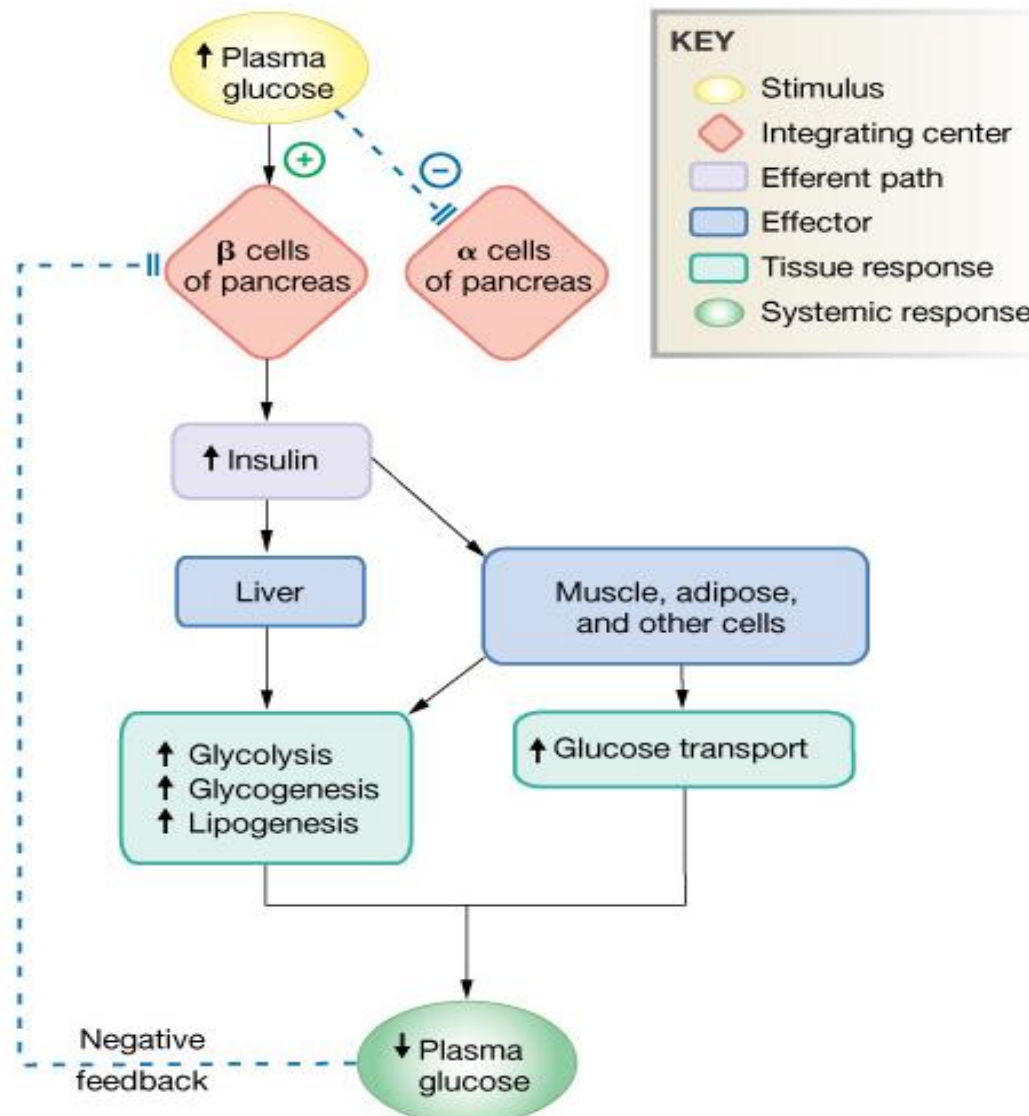


Table 22-3: Insulin

Cell of origin	Beta cells of pancreas
Chemical nature	51-amino acid peptide
Biosynthesis	Typical peptide
Transport in the circulation	Dissolved in plasma
Half-life	5 minutes
Factors affecting release	Plasma [glucose] > 100 mg/dL; ↑ blood amino acids; GI hormones (feedforward reflex) and parasympathetic amplify. Sympathetic inhibits.
Target cells or tissues	Liver, muscle, and adipose tissue primarily; brain, kidney, and intestine not insulin-dependent
Target receptor	Membrane receptor with tyrosine kinase activity; pathway with insulin-receptor substrates
Whole body or tissue action	↓ Plasma [glucose] by ↑ transport into cells or ↑ metabolic use of glucose
Action at cellular level	↑ Glycogen synthesis; ↑ aerobic metabolism of glucose; ↑ protein and triglyceride synthesis

Glucagon

- A 29-amino-acid polypeptide hormone that is a potent hyperglycemic agent
- Produced by α cells in the pancreas

SYNTHESIS

DNA in α cells (chromosome 2)



mRNA



Preproglucagon



proglucagon



glucagon

Factors Affecting Glucagon Secretion:

Effects on Glucagon Secretion

Stimuli for Glucagon Secretion

- ↓ Blood glucose
- ↑ Serum amino acids (arginine, alanine)
- Sympathetic nervous system stimulation
- Stress
- Exercise

Inhibitors of Glucagon Secretion

- Somatostatin
- Insulin
- ↑ Blood glucose

Glucagon Actions

- Its major target is liver:
 - Glycogenolysis
 - Gluconeogenesis
 - Lipid oxidation (fully to CO₂ or partially to produce keto acids “ketone bodies”).
 - Release of glucose to the blood from liver cells

Glucagon Action on Cells:

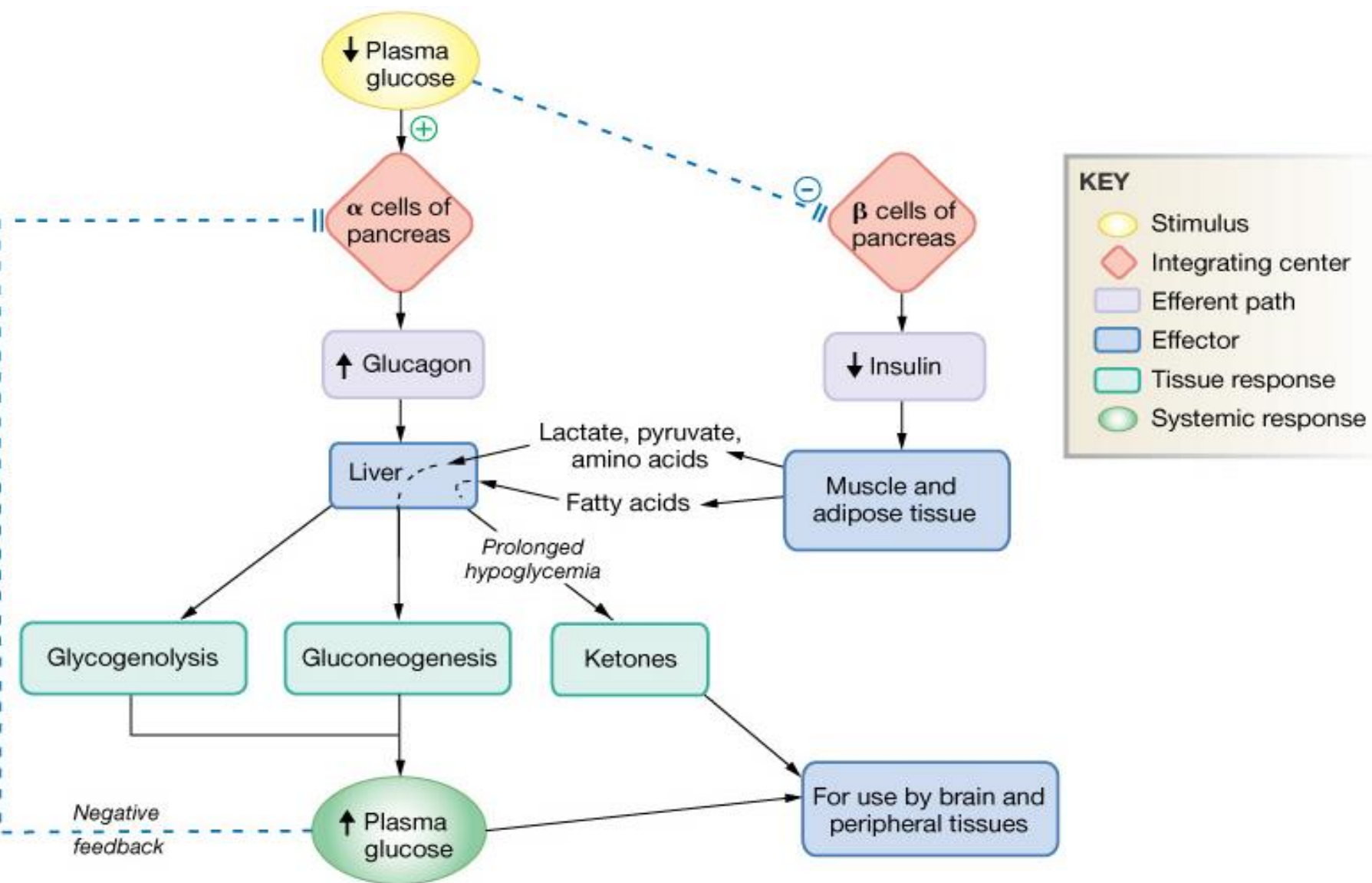
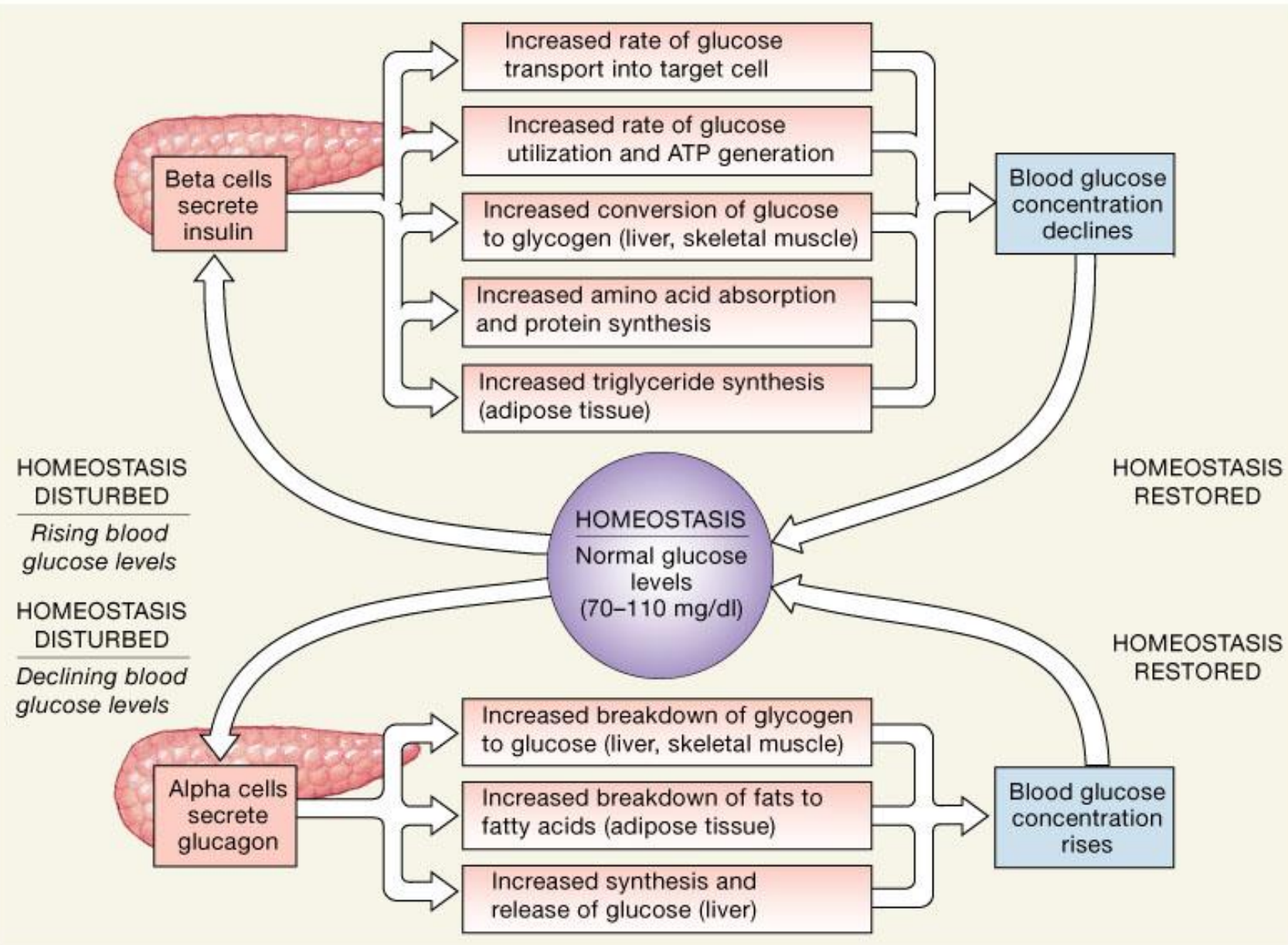


Table 22-5: Glucagon

Cell of origin	Alpha cells of pancreas
Chemical nature	29-amino acid peptide
Biosynthesis	Typical peptide
Transport in the circulation	Dissolved in plasma
Half-life	4–6 minutes
Factors affecting release	Stimulated by plasma [glucose] < 200 mg/dL, with maximum secretion below 50 mg/dL; ↑ blood amino acids.
Target cells or tissues	Liver primarily
Target receptor/second messenger	G protein-coupled receptor linked to cAMP
Whole body or tissue action	↑ Plasma [glucose] by glycogenolysis and gluconeogenesis; ↑ lipolysis leads to ketogenesis in liver
Action at molecular level	Alters existing enzymes and stimulates synthesis of new enzymes
Feedback regulation	↑ Plasma [glucose] shuts off glucagon secretion
Other information	Member of secretin family along with VIP, GIP, and GLP-1

The Regulation of Blood Glucose Concentrations





Diabetes

Olivia has a lot of baby fat on her leg so it's a good place to give her a shot.

~Olivia's Mom

Diabetes

- Diabetes is probably the most important metabolic disease.
- It affects every cell in the body and affects carbohydrate, lipid, and protein metabolism.
- characterized by the polytriad:
 - **Polyuria** (excessive urination)
 - **Polydypsia** (excessive thirst)
 - **Polyphagia** (excessive hunger).

Types of Diabetes

Type 1 Diabetes

Affects children

Cause:
inadequate
insulin secretion

Treatment :
insulin injection

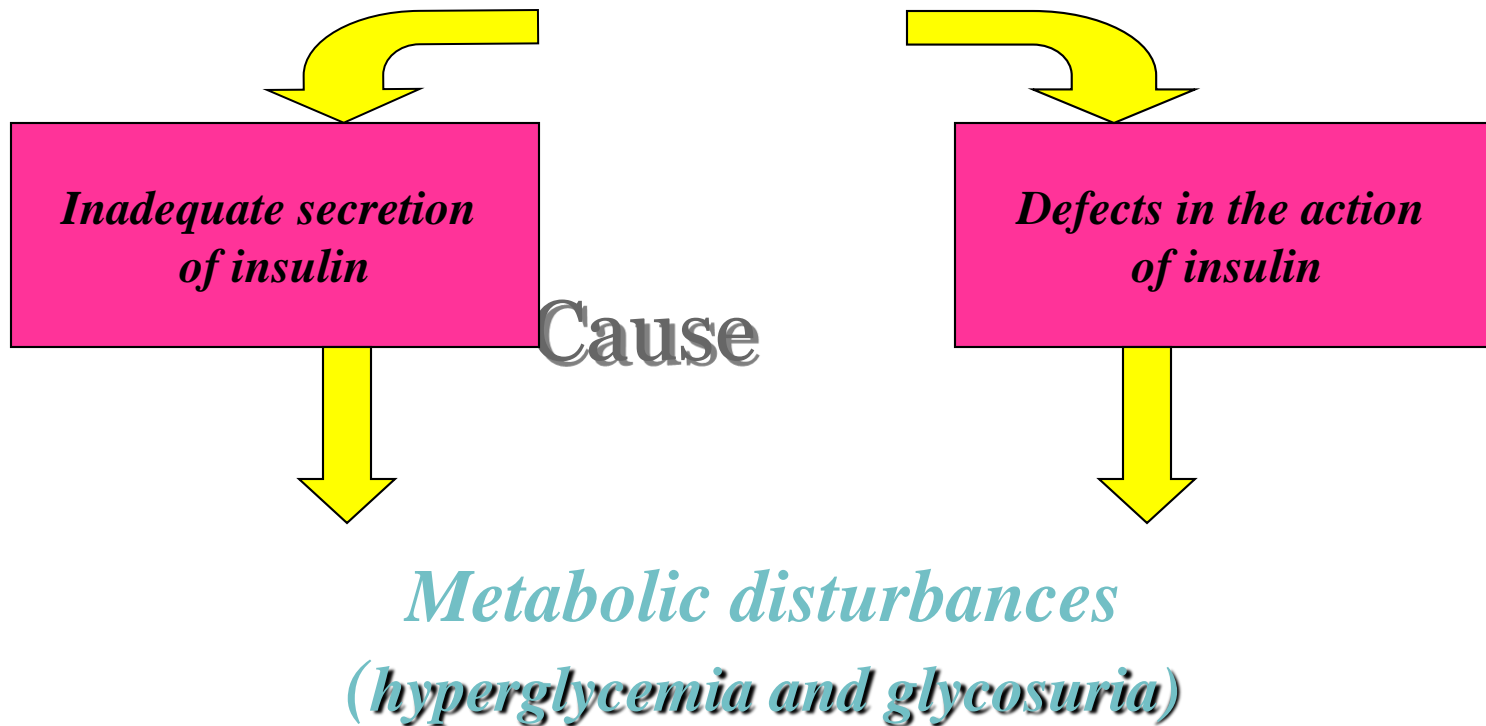
Type 2 diabetes

Affects adults

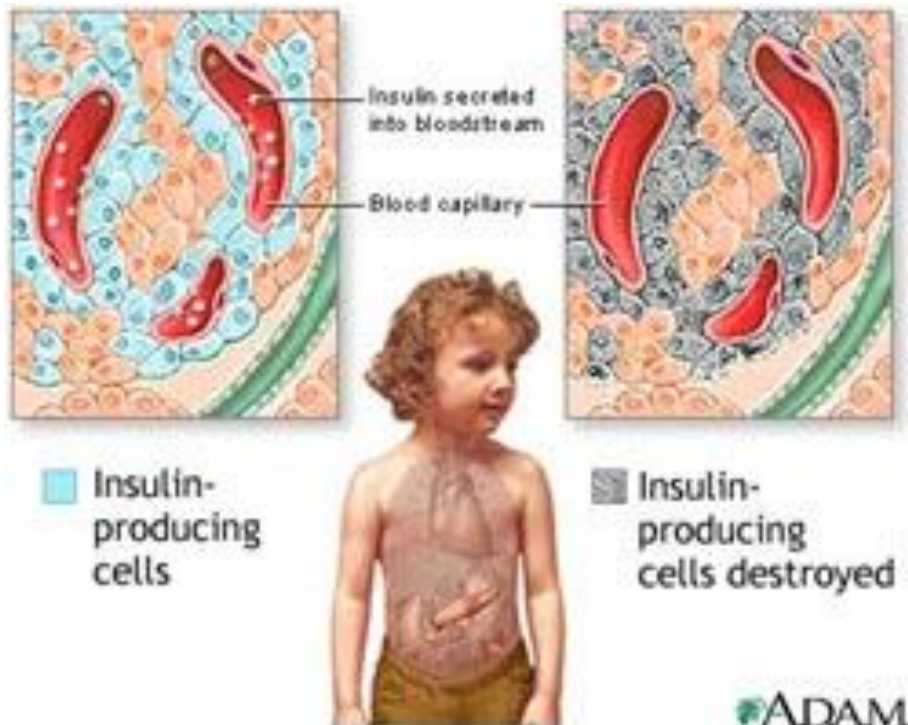
Cause defect in
insulin action

Treatment :
diet or OHA

Diabetes Mellitus



Type 1 diabetes



Diabetes Mellitus Type I

- Caused by an immune-mediated selective destruction of β cells
- β cells are destroyed while α cells are preserved:
No insulin :::: high glucagon \rightarrow high production of glucose and ketones by liver
glucose & ketones \uparrow \longrightarrow osmotic diuresis
keto acids \uparrow \longrightarrow diabetic ketoacidosis

Diabetes Mellitus: Type II

- More common in some ethnic groups
- Insulin resistance keeps blood glucose too high
- Chronic complications: atherosclerosis, renal failure & blindness

Long Term Complications of Uncontrolled Diabetes

- **MICROVASCULAR DISEASE**
- Hyperglycemia damages small blood vessels:
 - diabetic **retinopathy** → vision loss.
 - diabetic **neuropathy** → damage to nerves → most common cause of amputation in Western world.
 - diabetic **nephropathy** → kidney damage → chronic renal failure.



Oral glucose tolerance test

ADAM.

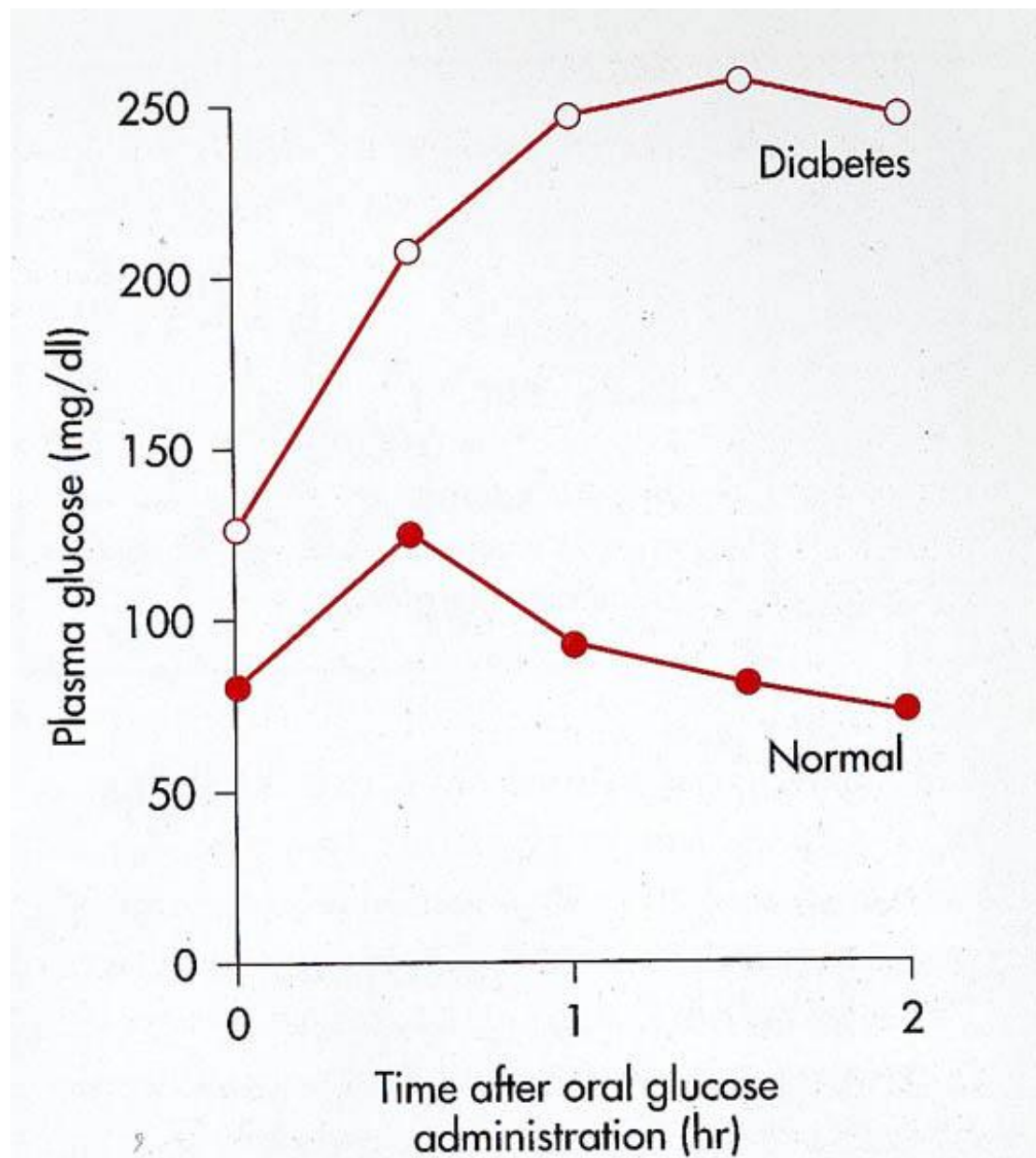
Glucose Tolerance Test

- Both the FPG and OGTT tests require that the patient fast for at least 8 hours (ideally 12 hr) prior to the test.
- The oral glucose tolerance test (OGTT):
 - FPG test
 - Blood is then taken 2 hours after drinking a special glucose solution

Glucose Tolerance Test (GTT)

- Following the oral administration of a standard dose of glucose, the plasma glucose concentration normally rises but returns to the fasting level within 2 hours.
- If insulin activity is reduced, the plasma glucose concentration takes longer than 2 hours to return to normal and often rises above 200 mg/dl.
- Measurement of urine glucose allows determination of the renal threshold for glucose.

GTT



Glucose Tolerance Test

The following results suggest different conditions:

- **Normal values:**
- FPG <100 mg/dl
- 2hr PPG < 140 mg/dL
- **Impaired glucose tolerance**
- 2hr PPG = 140 - 199 mg/dL
- **Impaired fasting glucose**
- FPG=100-125
- **Diabetes**
- FPG ≥ 126 mg/dl
- 2hr PPG levels ≥200 mg/dL

Symptoms of Diabetes Mellitus

Symptoms of Diabetes Mellitus

Hyperglycemia

Polyuria

Polydipsia

Polyphagia

Ketoacidosis (IDDM)

Hyperlipidemia

Muscle wasting

Electrolyte depletion

Diabetes Mellitus (DM)

Organs/tissue involved	Organ/tissue responses to insulin deficiency	Resulting condition of:		Signs and symptoms
		Blood	Urine	
  	Decreased glucose uptake and utilization	Hyperglycemia	Glycosuria	Polyuria - dehydration - soft eyeballs
	Glycogenolysis		Osmotic diuresis	Polydipsia Fatigue
 	Protein catabolism and gluconeogenesis			Weight loss Polyphagia
 	Lipolysis and ketogenesis	Lipidemia and ketoacidosis	Ketonuria Loss of Na ⁺ , K ⁺ ; electrolyte and acid-base imbalances	Acetone breath Hyperpnea Nausea/vomiting/abdominal pain Cardiac irregularities Central nervous system depression; coma

 = Muscle
  = Adipose tissue
  = Liver







The End

Thank You