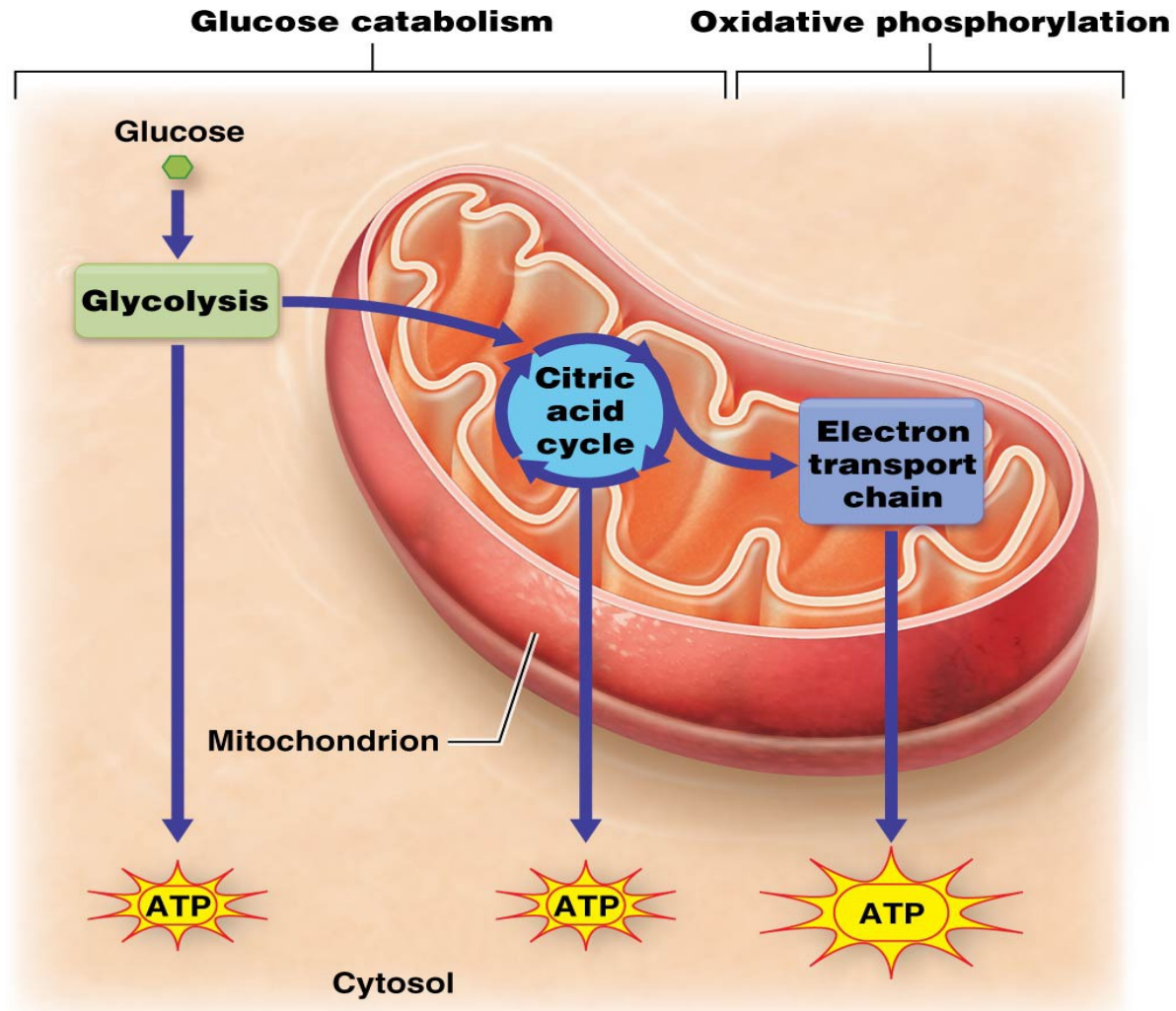
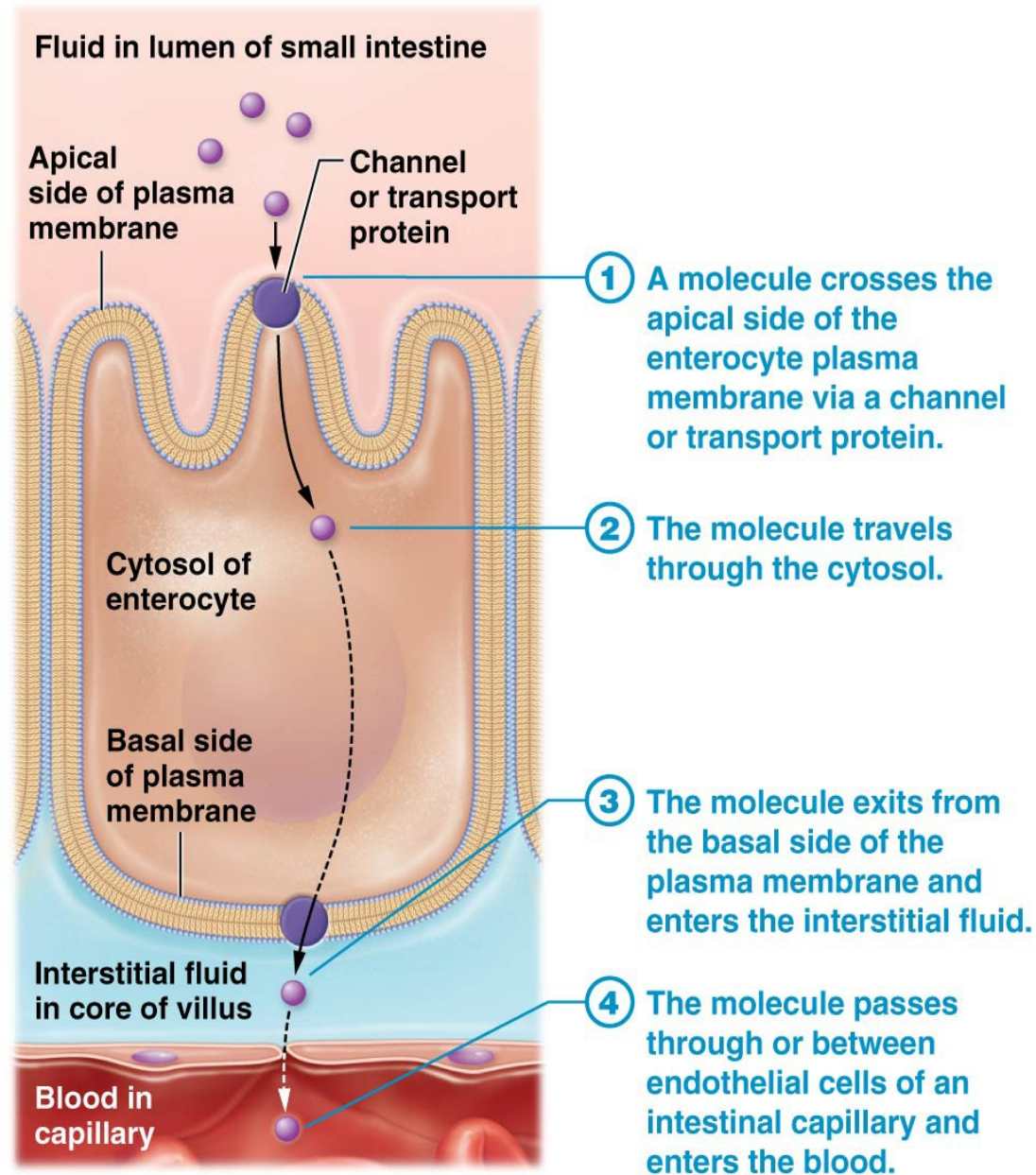


Metabolism (C26)



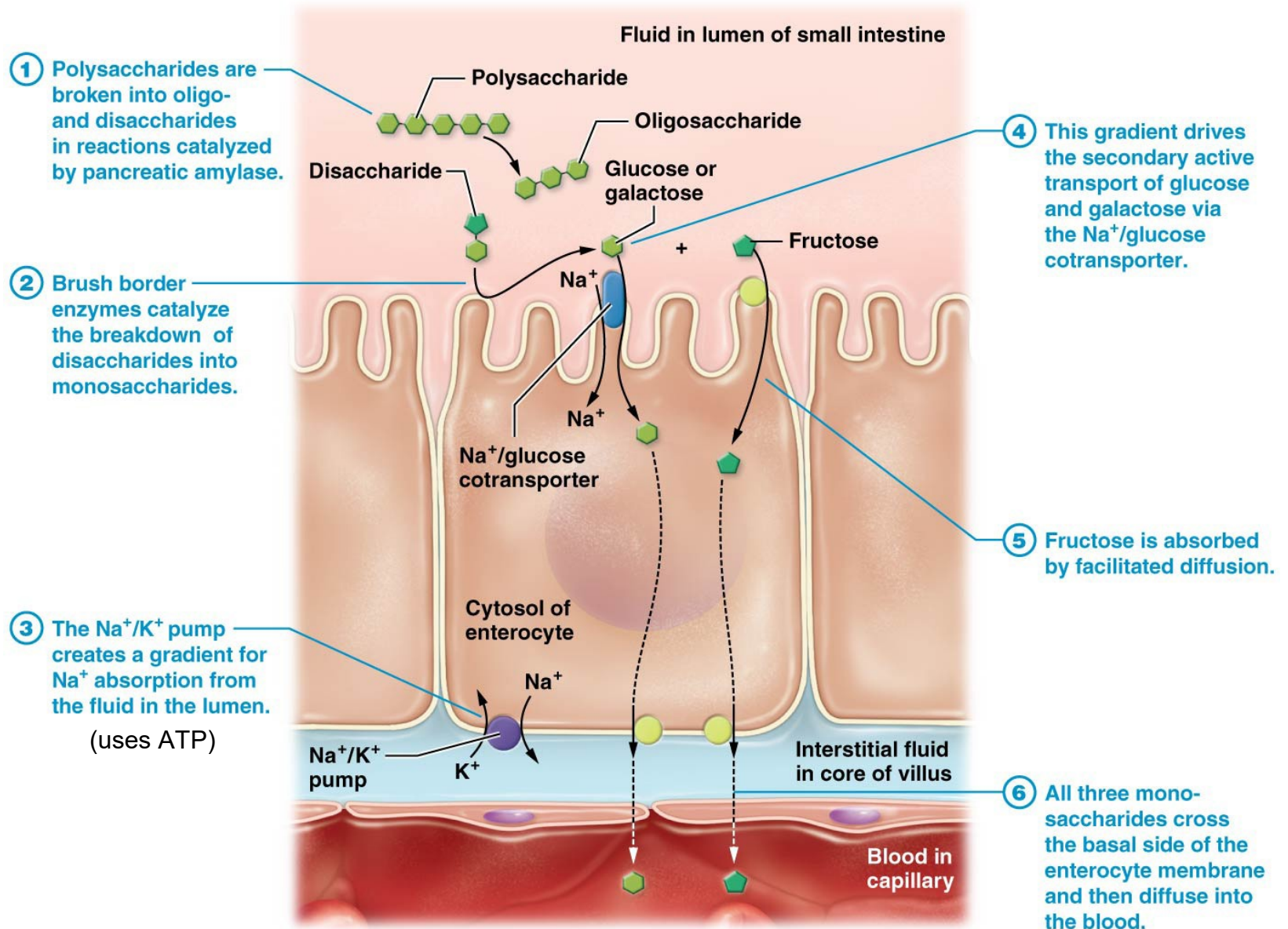
Path taken by hydrophilic **molecules** absorbed into blood through an enterocyte.



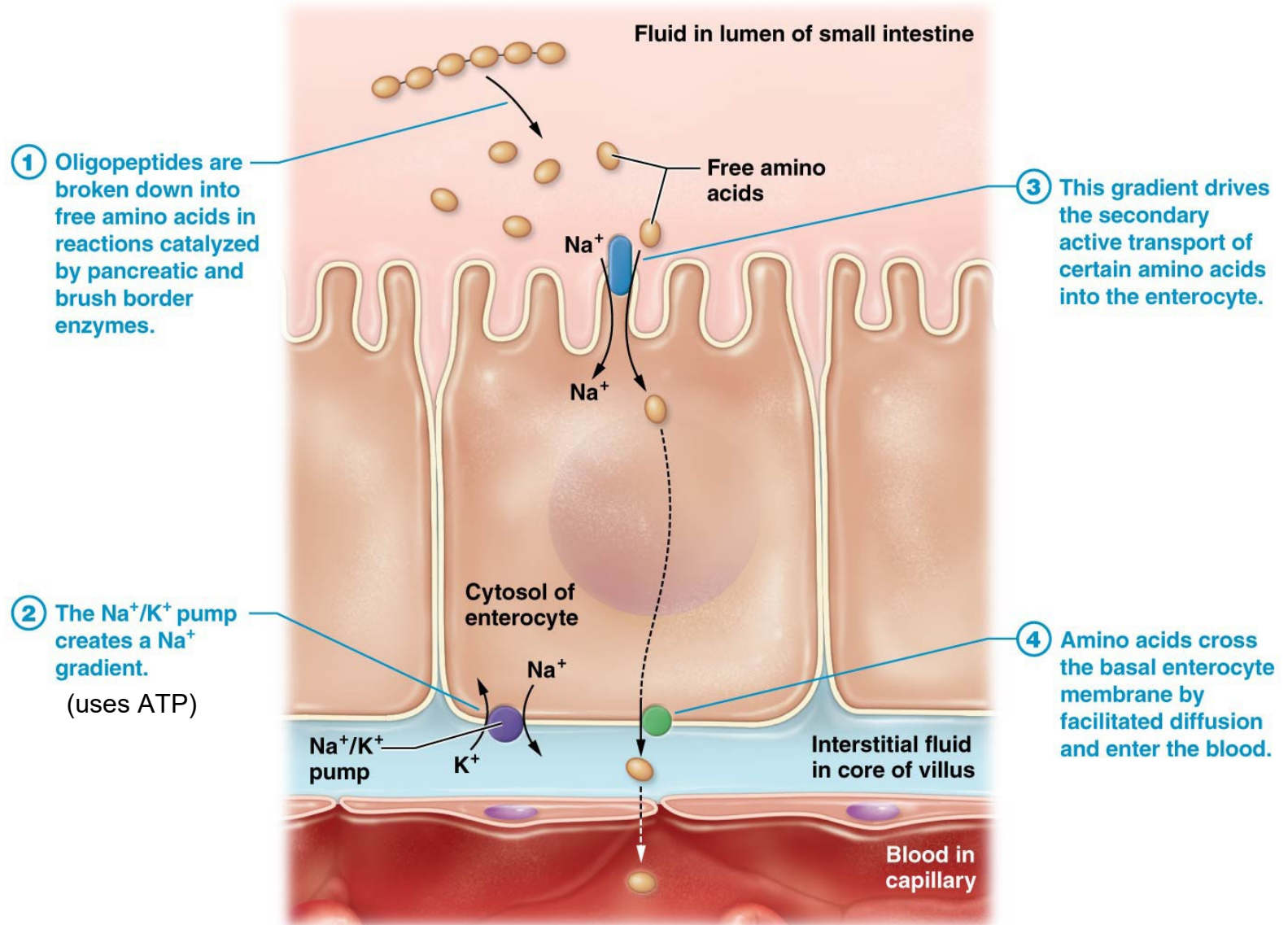
Carbohydrate (**hydrophilic molecule**) digestion and absorption in the small intestine



Can you find the symports, antiports, and uniports? Active vs passive transporters?

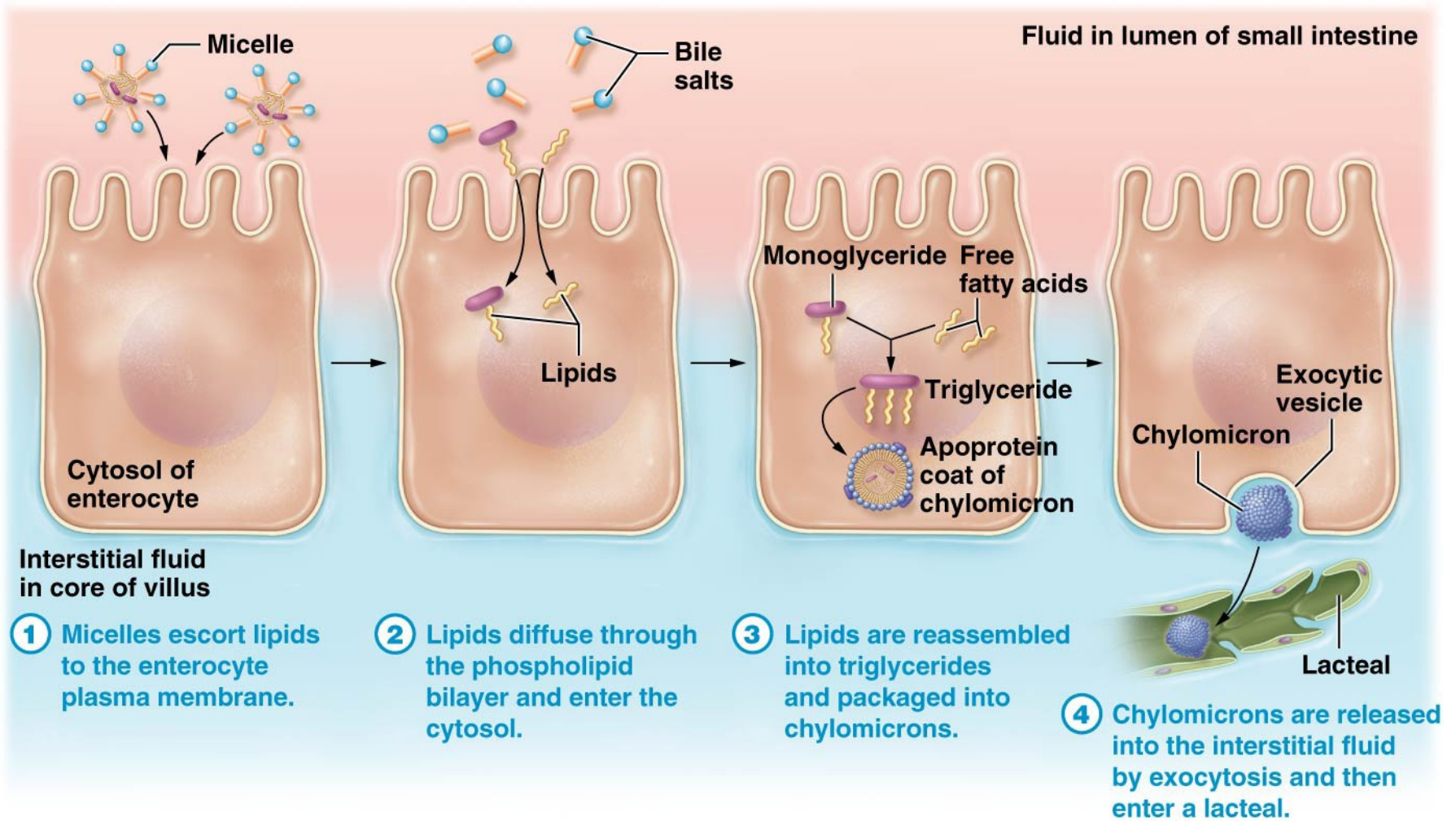


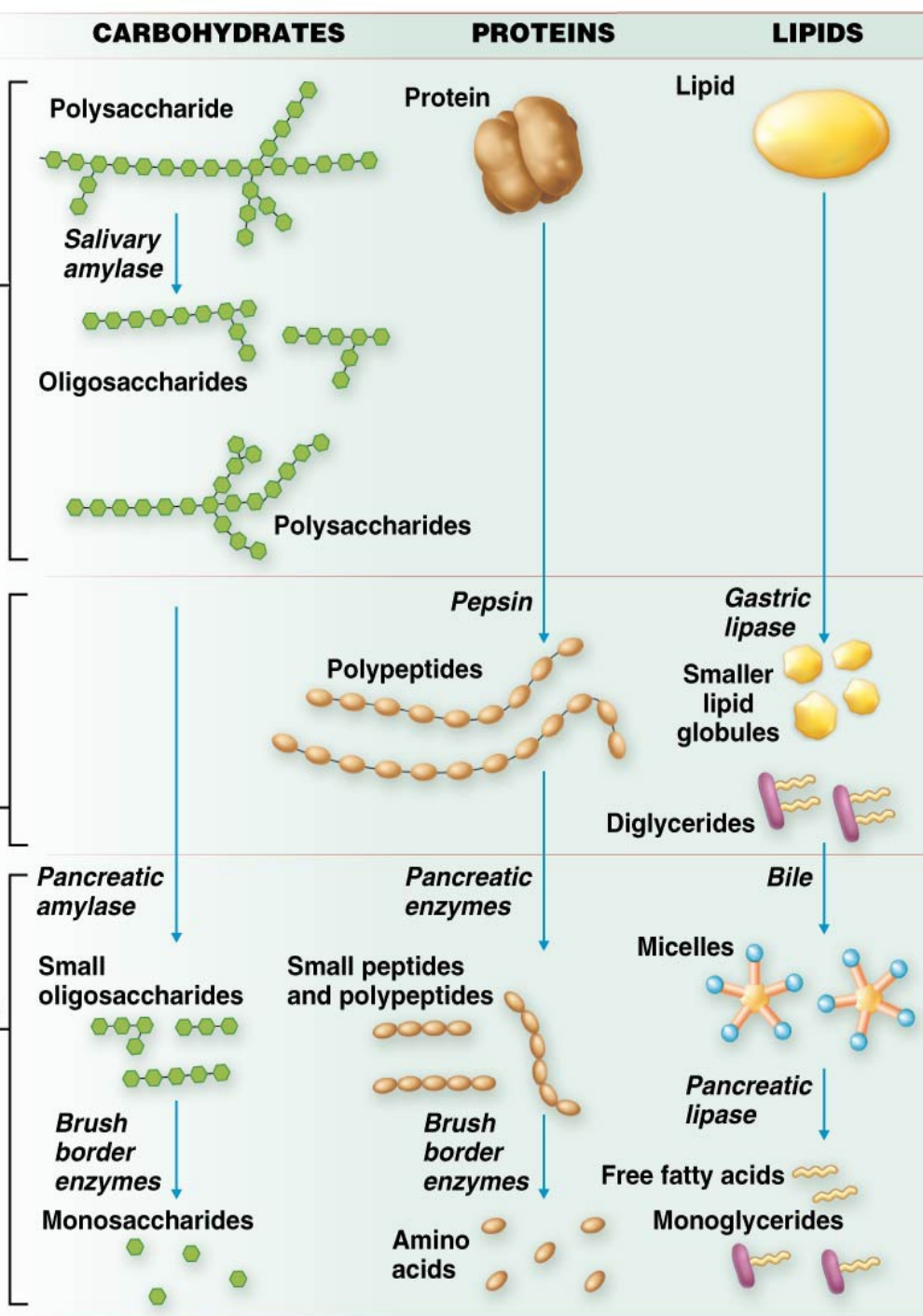
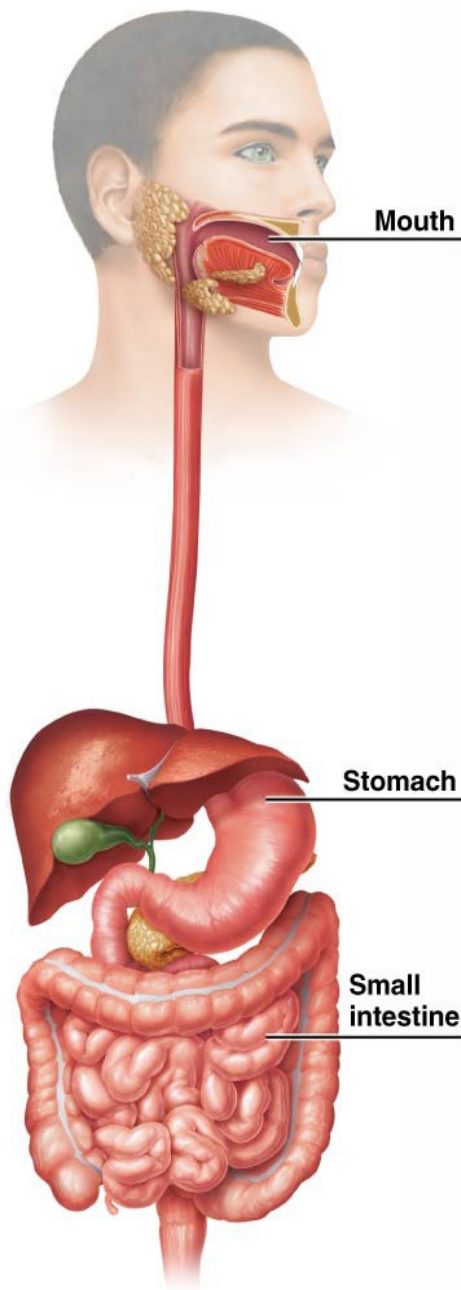
Amino acid digestion and absorption in the small intestine.



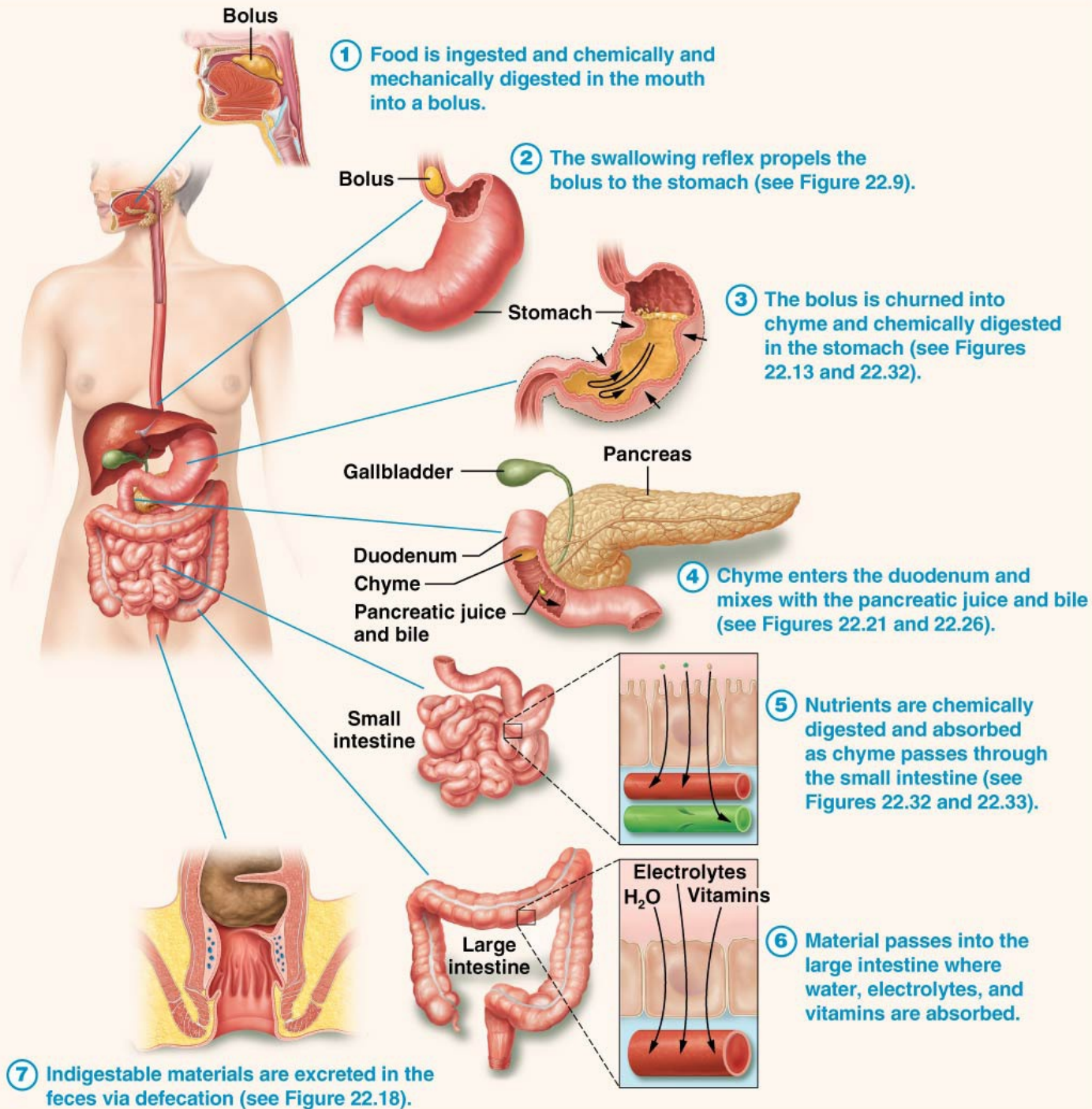
What is secondary active transport?

Lipid (hydrophobic molecule) digestion and absorption in the small intestine



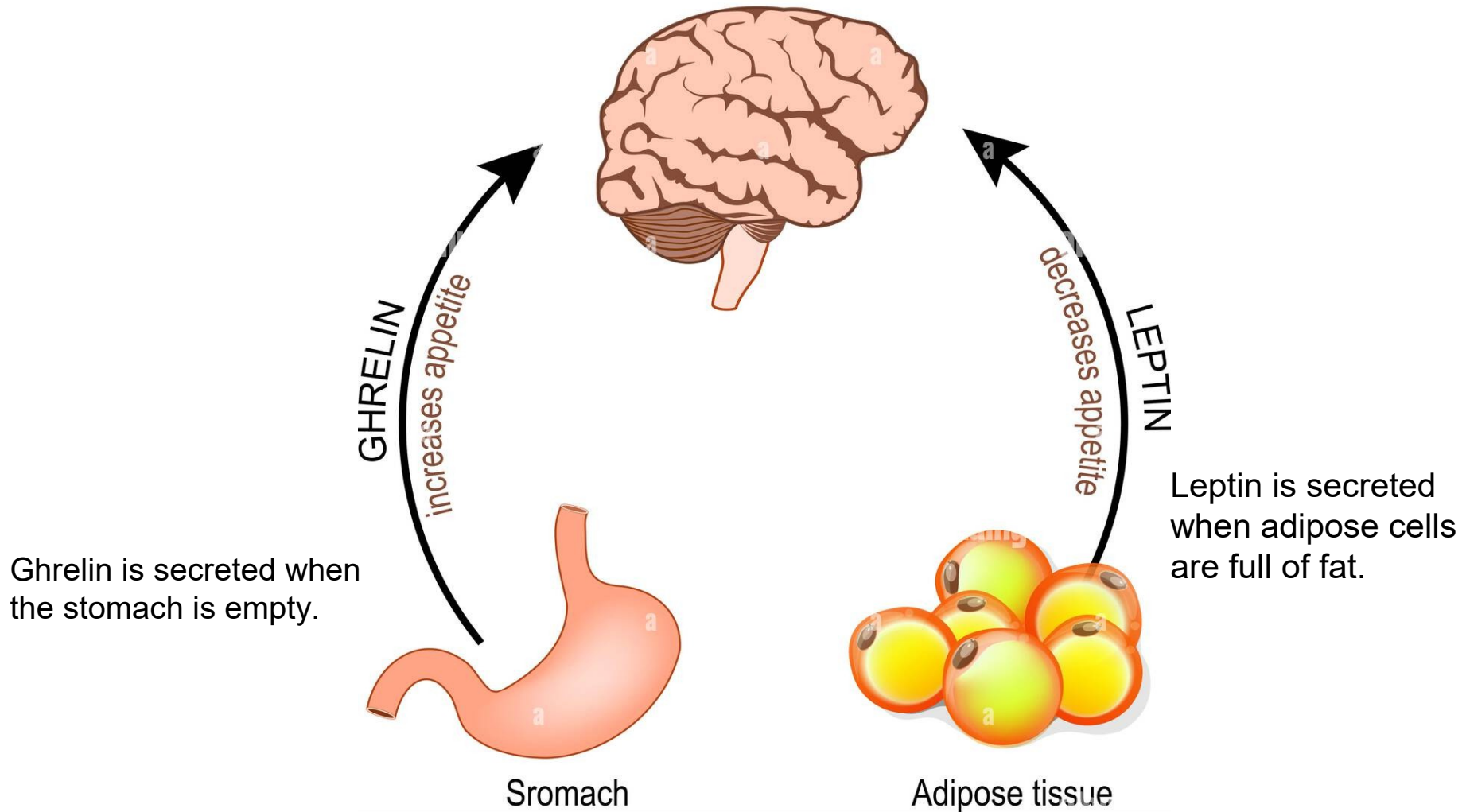


Summary of the digestion of carbohydrates, proteins, and lipids.

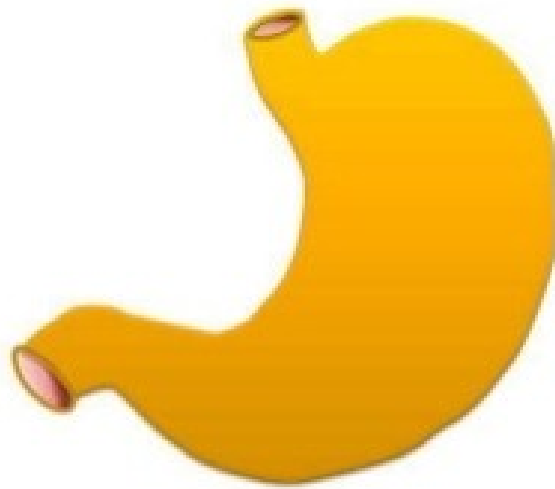


The Big Picture of Digestion.

Leptin & Ghrelin



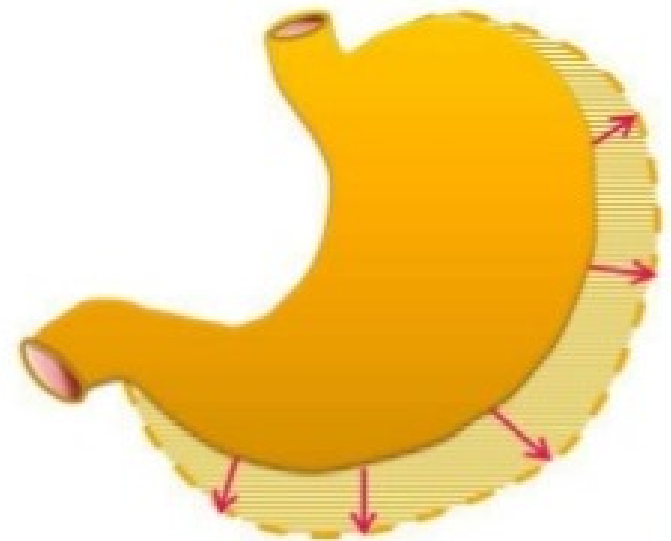
Stomach Empty



↑ Ghrelin = ↑ Appetite

CCK, GLP-1, PYY – all ↓

Stomach Full



↓ Ghrelin = ↓ Appetite

CCK, GLP-1, PYY – all ↑

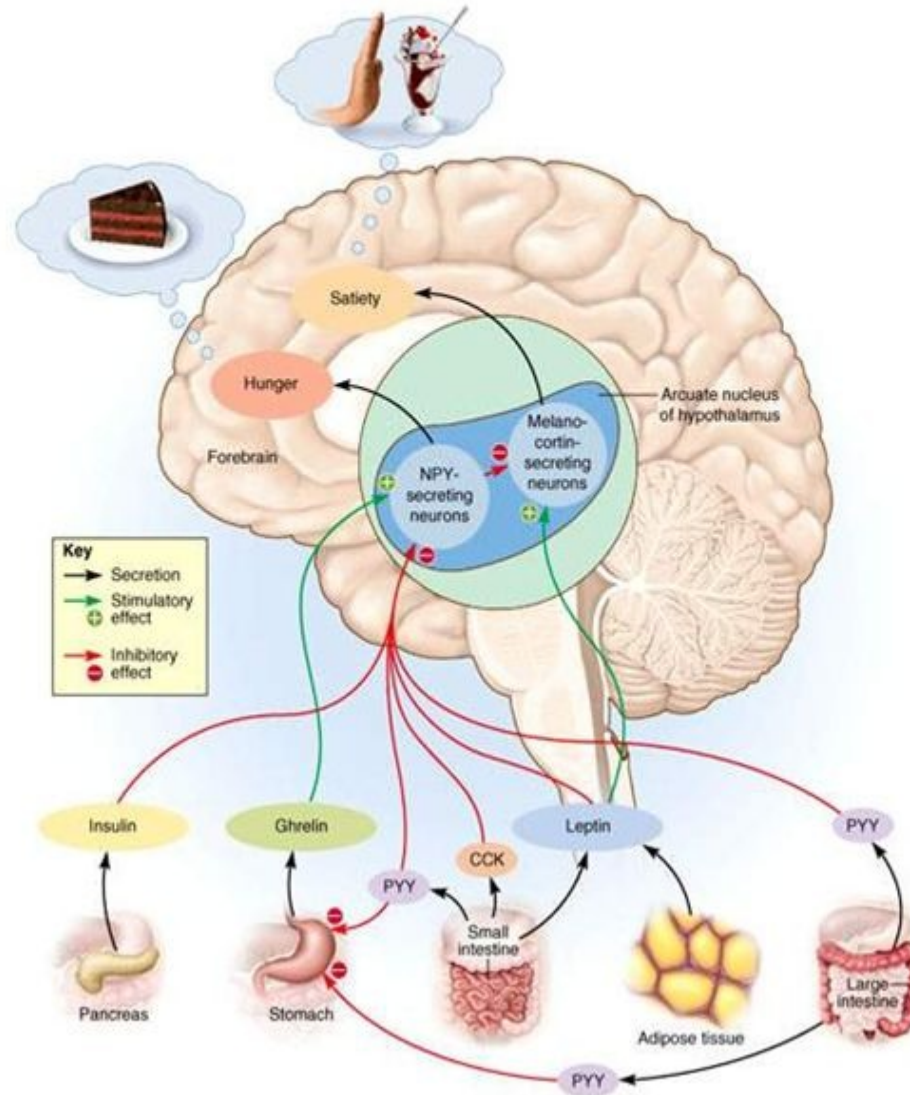
CCK – Cholecystokinin
GLP1 – Glucagon-like
peptide 1
PYY – Peptide YY

Primarily regulated
by feeding

(Crespo *et al.*,
2014)

Appetite Regulation

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Glucagon-Like Peptide (GLP-1)

New weight loss drugs (e.g. Semaglutide) work by mimicking the action of glucagon-like peptide 1 (GLP-1), a hormone that's naturally released in the body after eating:

Slow digestion: GLP-1-mimicking drugs slow down the rate at which food leaves the stomach, which helps people feel full longer.

Increase insulin production: These drugs stimulate the body to produce more insulin, which helps control blood sugar levels.

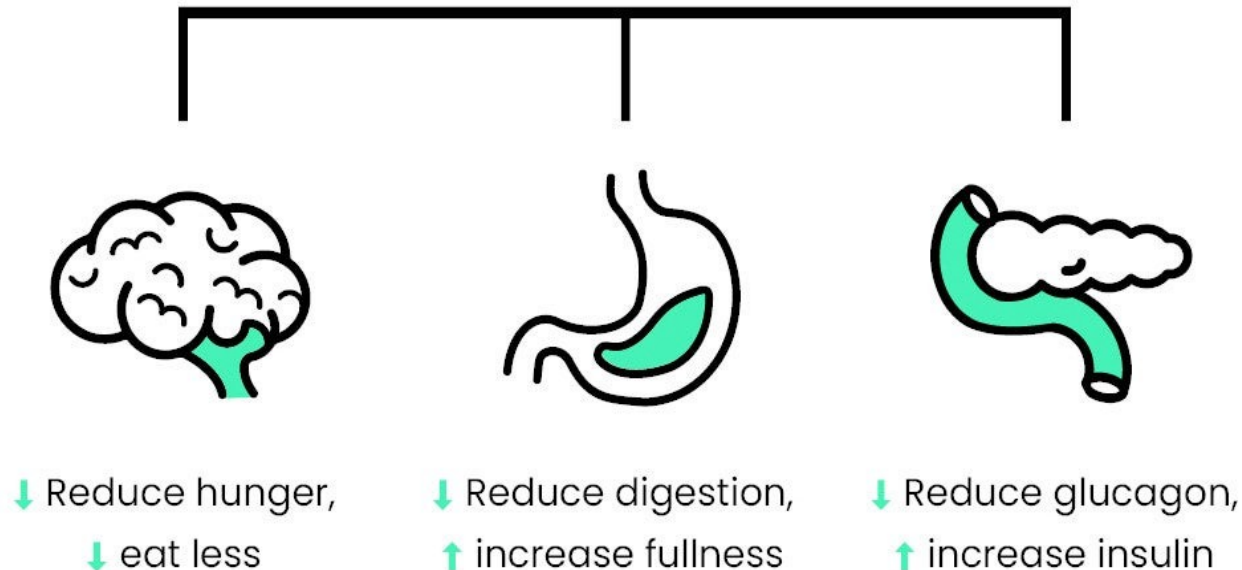
Reduce food cravings: GLP-1-mimicking drugs may also affect the brain's appetite pathway, which can help reduce food cravings.

Last longer than natural GLP-1: Unlike the natural hormone, which only lasts a few minutes, GLP-1-mimicking drugs can last for days.

Some examples of GLP-1-mimicking drugs include: Semaglutide: A synthetic drug that mimics GLP-1 // Ozempic: An injectable drug that mimics GLP-1 // Orforglipron: An oral drug that mimics GLP-1

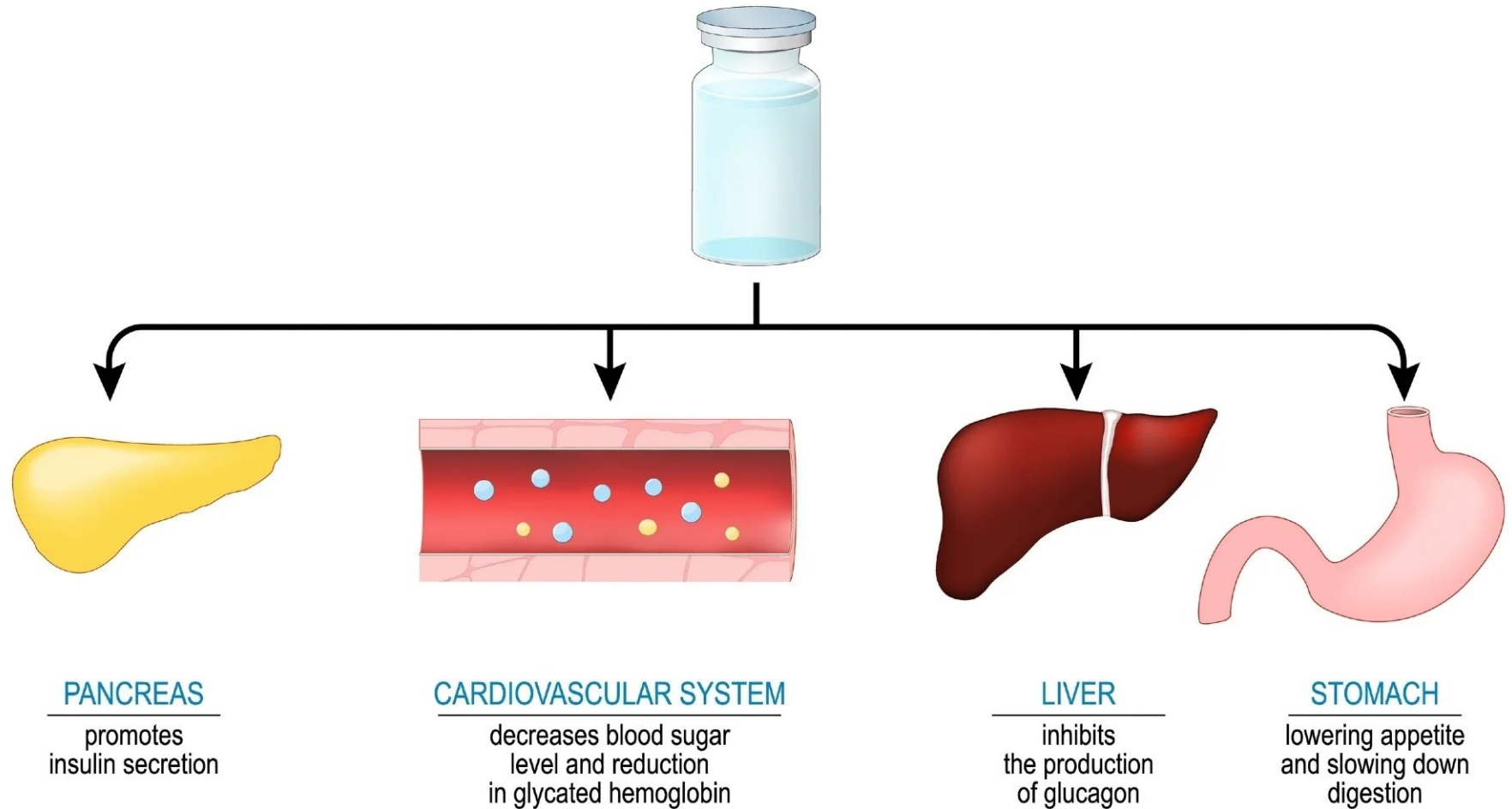
These drugs are only approved for certain people, such as those with obesity or type 2 diabetes. Some side effects of these drugs include nausea, diarrhea, constipation, and increased risk of gut-related side effects.

GLP-1



Semaglutide

(a peptide similar to the hormone GLP-1)



Glucagon-like peptide 1 (GLP-1) is released from the following cells in the body:

Intestinal L-cells: Located in the distal jejunum, ileum, and colon, these cells release GLP-1 in response to nutrients, hormones, and neurotransmitters. /// Pancreatic alpha cells: These cells release GLP-1. /// Neurons in the nucleus of the solitary tract:

GLP-1 is an incretin hormone that helps regulate blood glucose levels. It's released in response to eating, especially carbohydrates, and slows gastric emptying. GLP-1 also stimulates insulin secretion and can make people feel full, which can help reduce food intake.

Some things that can increase GLP-1 levels include:

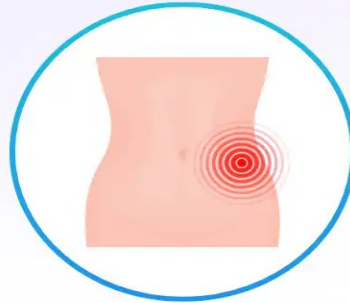
Eating fiber - Fiber from fruits, vegetables, and legumes can help the gut produce short-chain fatty acids (SCFAs) like acetate and butyrate, which can increase GLP-1 levels.

Eating arabinoxylan-oligosaccharide // This fiber is found in whole grains like wheat, rye, and barley, and can encourage the gut to produce more GLP-1.

Side Effects of GLP-1 Receptor Agonists



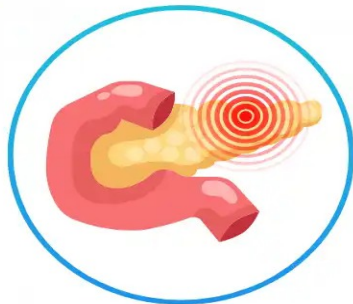
Gastrointestinal problems



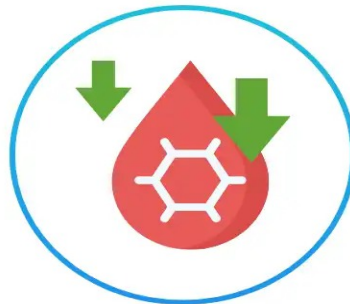
Injection site reactions



Decrease in appetite



Risk of pancreatitis



Hypoglycemia



Risk of thyroid cancer



Side effects are very serious, including gastric paralysis which may not be reversible. Some patients regret taking the medication.

Carbohydrate Digestion



- **Starch** – the most digestible carbohydrate
 - starch is first digested to:
 - **oligosaccharides** up to eight glucose residues long
 - then into the **disaccharide maltose**
 - finally to **glucose** which is absorbed by the small intestine
 - process **begins in the mouth**
 - **salivary amylase** hydrolyzes starch into oligosaccharides
 - amylase works best at pH of 6.8 – 7.0 of oral cavity
 - amylase quickly denatured on contact with stomach acid and digested by pepsin
- **Cellulose** – *indigestible glucose polymer!*
- **What is glycogen?**
- **Do we have enzymes to digest glycogen?**

Carbohydrate Digestion



- salivary amylase stops working in stomach at pH less than 4.5
 - 50% of dietary starch digested before it reaches small intestine
- when reaching the small intestine, **pancreatic amylase** converts **starch to oligosaccharides and maltose** within 10 minutes
- oligosaccharides and maltose contact **brush border enzymes** (dextrinase, glucoamylase, maltase, sucrase, and lactase) /// these act to turn oligosaccharides, maltose, sucrose, lactose, and fructose into glucose
- **lactose** becomes indigestible after age 4 in many humans due to decline in **lactase** production (lactose intolerance)

Carbohydrate Absorption



- plasma membrane of the **absorptive cells** has transport proteins that absorb monosaccharides as soon as the brush border enzymes creates monosaccharides
- 80% of absorbed sugar is the monosaccharide glucose
 - Moved from small intestine's lumen across apical enterocyte by **sodium-glucose transport proteins (SGLT)** and **secondary active transport of the sodium-potassium ATP pump in the basal membrane**
 - then glucose is transported out the base of absorptive cell into ECF by **facilitated diffusion**
 - sugar entering ECF increases its osmolarity /// draws water by osmotic drag from the lumen of the intestine, through **now leaky tight junctions** between epithelial cells
 - water carries additional glucose and other nutrients with it by **solvent drag through leaky junctions**
 - **See next slide**

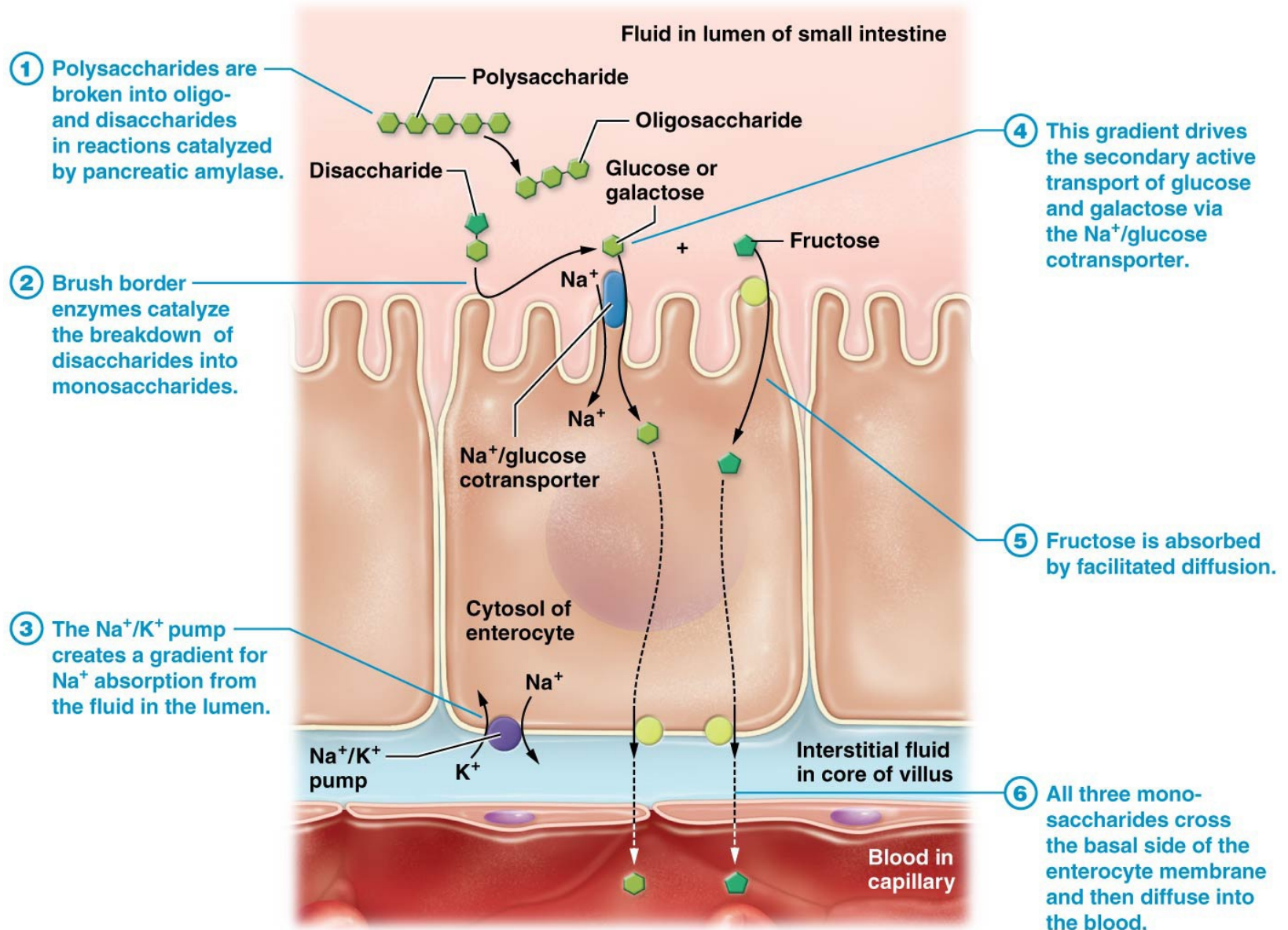
Carbohydrate Absorption

- **sodium-glucose transport proteins** (SGLT) (also transports galactose)
- The SGLT-1 was the first transporter studied. This protein is encoded by SLC5A1 gene (Turk et al., 1994). The SGLT1 protein controls the import of glucose in enterocytes and epithelial cells in the small intestine and proximal convoluted tubules in nephrons, respectively.
- fructose is absorbed by facilitated diffusion
- some glucose and galactose may also be transported out of the base of the cell by **facilitated diffusion**
- absorbed by blood capillaries in the **villus**
- hepatic portal system delivers these sugars to the **liver**

Carbohydrate digestion and absorption in the small intestine.



Can you find the symports, antiports, and uniports? Active VS passive transporters?



Lactose Intolerance

- At birth, all mammals (including humans) have the enzyme lactase to digest lactose (sugar in milk)
- Many will lose lactase as they age. At around age four the transcription factor to make lactase is inhibited. Now unable to digest lactose.
- If lactose passes undigested into large intestine
 - increases osmolarity of intestinal contents
 - causes water retention in the colon and diarrhea
 - gas production by bacterial fermentation of the lactose
- Lactose intolerance occurs in many populations //
 - 15% American whites
 - 90% of American blacks
 - 70% of Mediterranean's
 - nearly all of Asian descent
- Lactose intolerant people may consume dairy products like yogurt and cheese because bacteria in manufacturing process break down lactose

Proteins



- **amino acids** absorbed by the small intestine come from three sources:
 - **dietary proteins**
 - **digestive enzymes** digested by each other
 - **epithelial cells** sloughed off mucosa and digested by enzymes
- **endogenous amino acids** from last two sources total about 30g/day
- **exogenous amino acids** from our diet total about 44 to 60 g/day
- **proteases (peptidases)**
 - enzymes that digest proteins
 - begin their work in the stomach in optimum pH of 1.5 to 3.5
 - **pepsin** hydrolyzes any peptide bond between tyrosine and phenylalanine
 - **pepsin** digests 10-15% of dietary protein into shorter peptides and some free amino acids

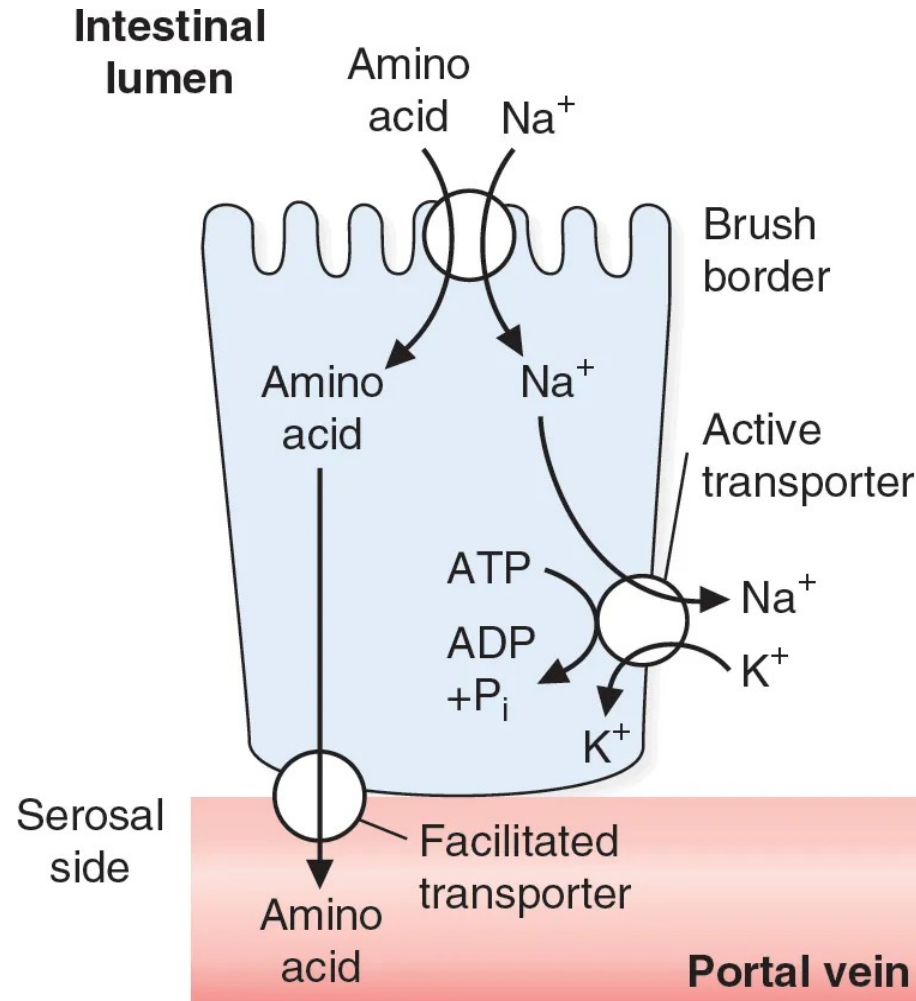
Proteins

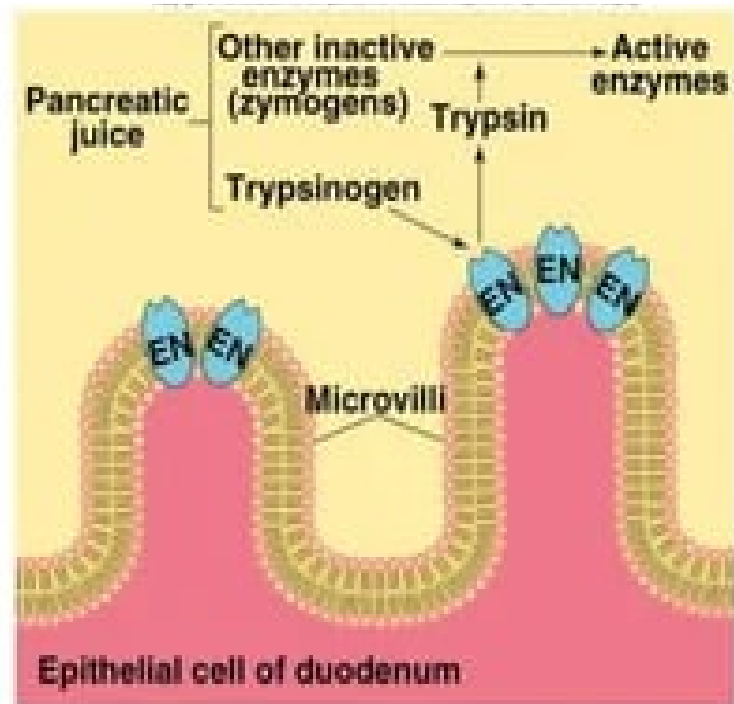


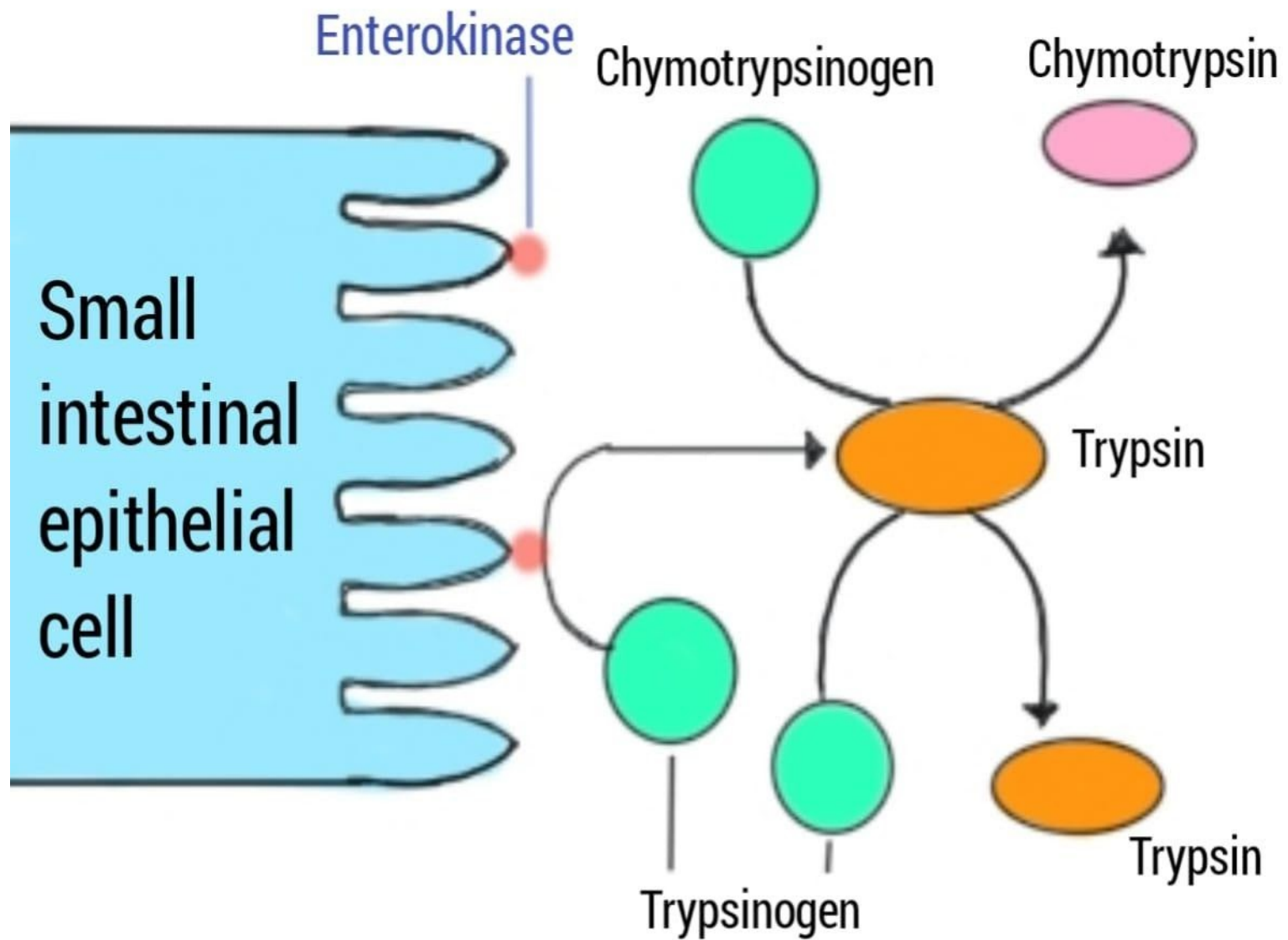
- Protein digestion started in stomach continue in the small intestine
 - Gastric pepsin inactivated when it passes into the duodenum and mixes with the alkaline pancreatic juice (pH 8)
 - pancreatic enzymes trypsin and chymotrypsin take over the process
 - hydrolyzing polypeptides into even shorter oligopeptides
 - oligopeptides taken apart one amino acid at a time by three more enzymes
 - carboxypeptidase – removes amino acids from -COOH end of the chain
 - aminopeptidase – removes them from the -NH_2 end
 - dipeptidase – split dipeptides in the middle and release two free amino acids
 - carboxypeptidase is a pancreatic secretion
 - aminopeptidase and dipeptidase are brush border enzymes

Protein Digestion and Absorption

Pancreatic enzymes take over protein digestion in small intestine by hydrolyzing polypeptides into shorter oligopeptides



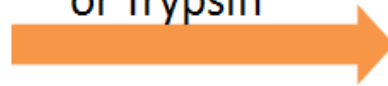




Trypsinogen

Enteropeptidase
or Trypsin

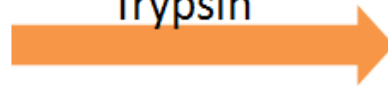
Trypsin



Chymotrypsinogen

Trypsin

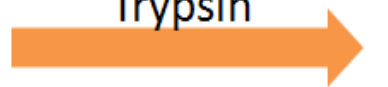
Chymotrypsin



Procarboxypeptidases

Trypsin

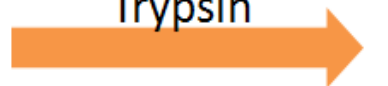
Carboxypeptidase A
Carboxypeptidase B
Aminopeptidases



Proelastase

Trypsin

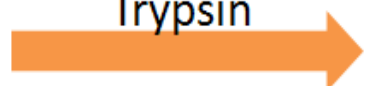
Elastase



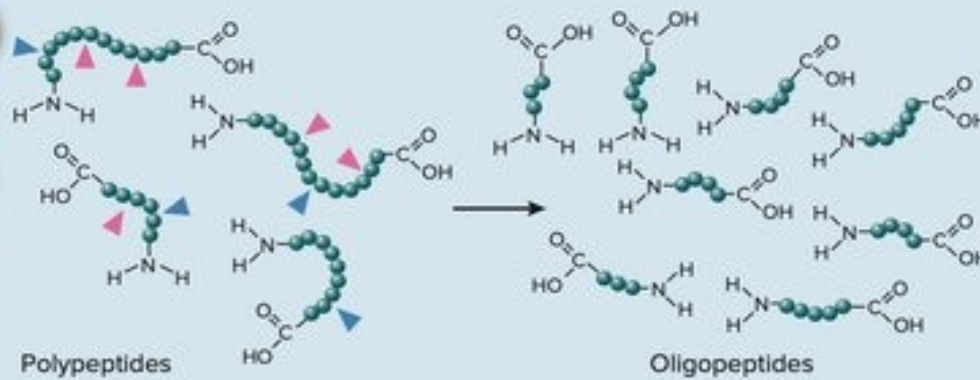
Procolipase

Trypsin

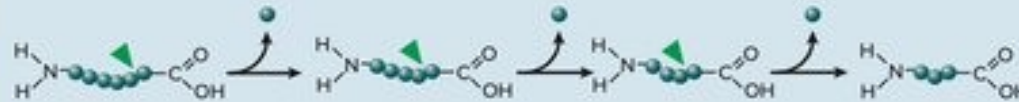
Colipase



Small intestine Actions of pancreatic enzymes

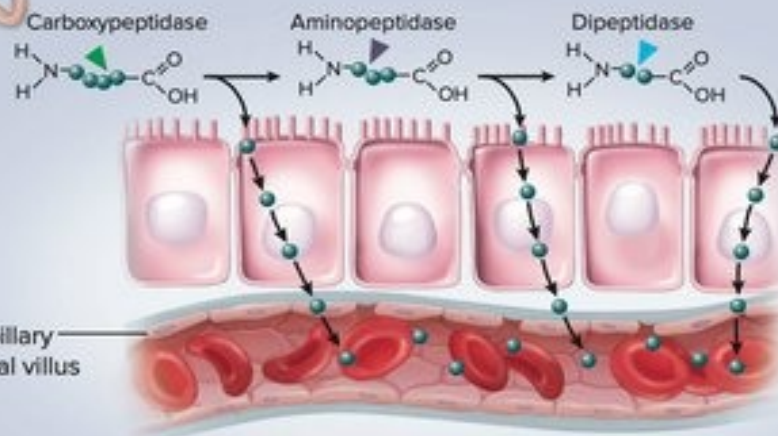


Trypsin (▲) and chymotrypsin (▲) hydrolyze other peptide bonds, breaking polypeptides down into smaller oligopeptides.



Carboxypeptidase (▲) removes one amino acid at a time from the carboxyl (—COOH) end of an oligopeptide.

Small intestine Actions of brush border enzymes (contact digestion)



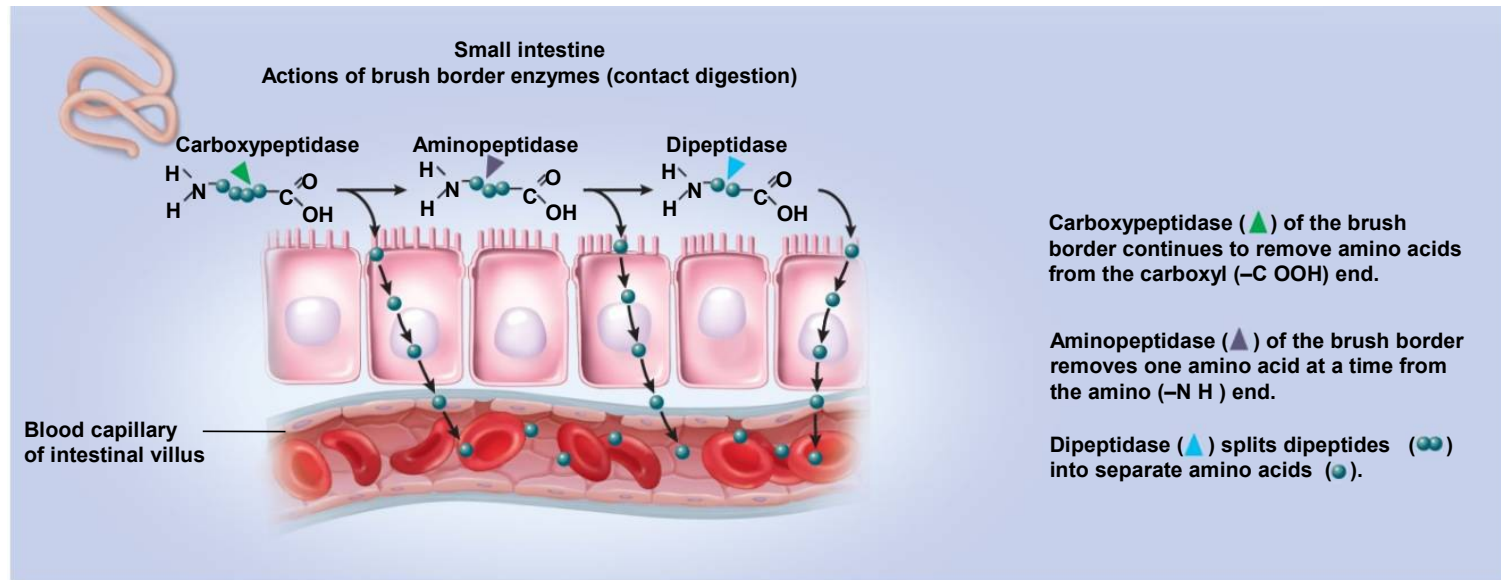
Carboxypeptidase (▲) of the brush border continues to remove amino acids from the carboxyl (—COOH) end.

Aminopeptidase (▲) of the brush border removes one amino acid at a time from the amino (—NH_2) end.

Dipeptidase (▲) splits dipeptides (●●) into separate amino acids (●).

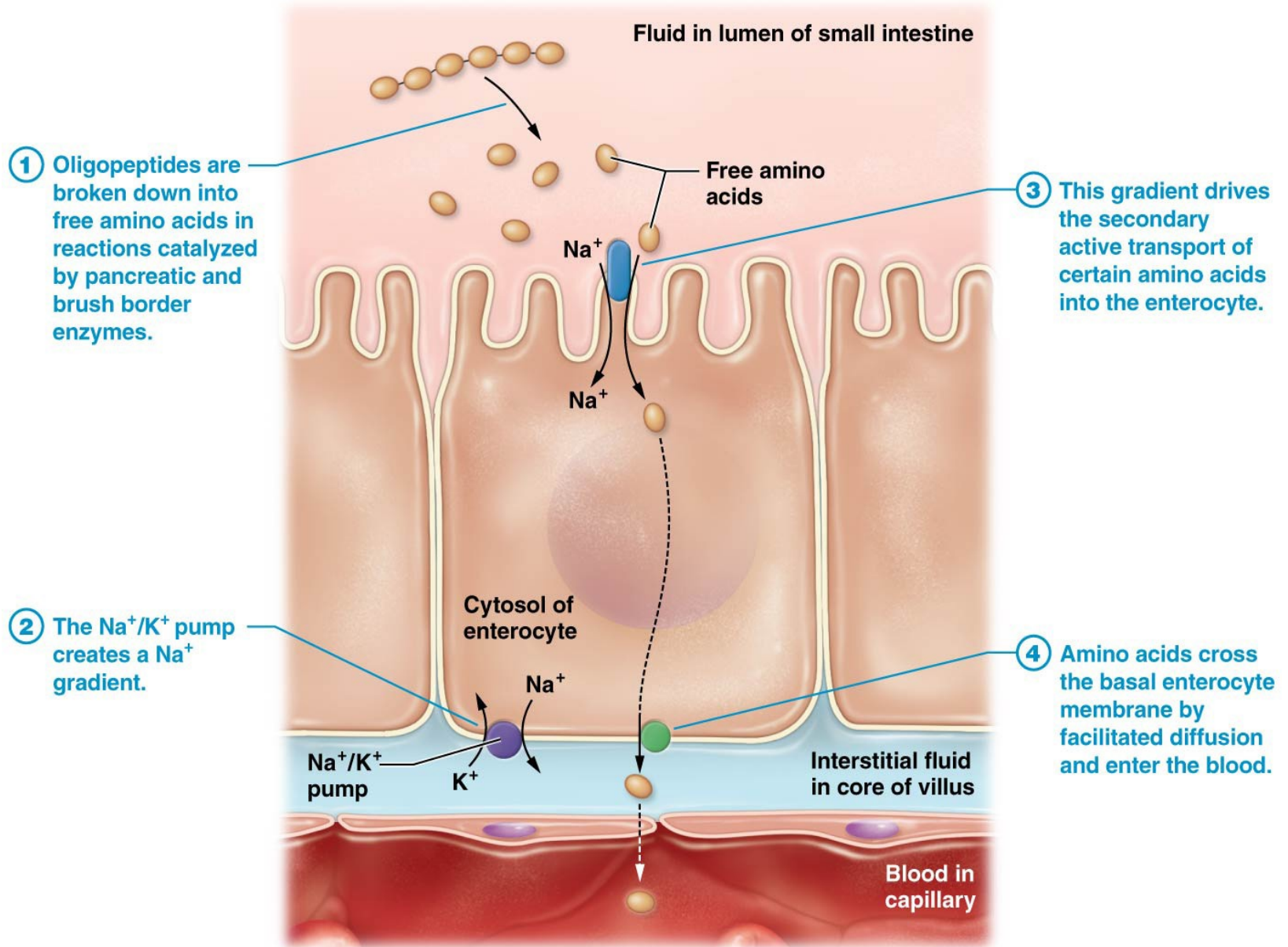


Protein Digestion and Absorption



- **brush border enzymes** finish task, producing free amino acids that are absorbed into intestinal epithelial cells
 - **sodium-dependent amino acid cotransporters** move amino acids into epithelial cells
 - **facilitated diffusion** moves amino acids into blood stream
- **infants absorb proteins by pinocytosis (maternal IgA)** and release into the blood by exocytosis

Amino acid digestion and absorption in the small intestine.



What is secondary active transport?

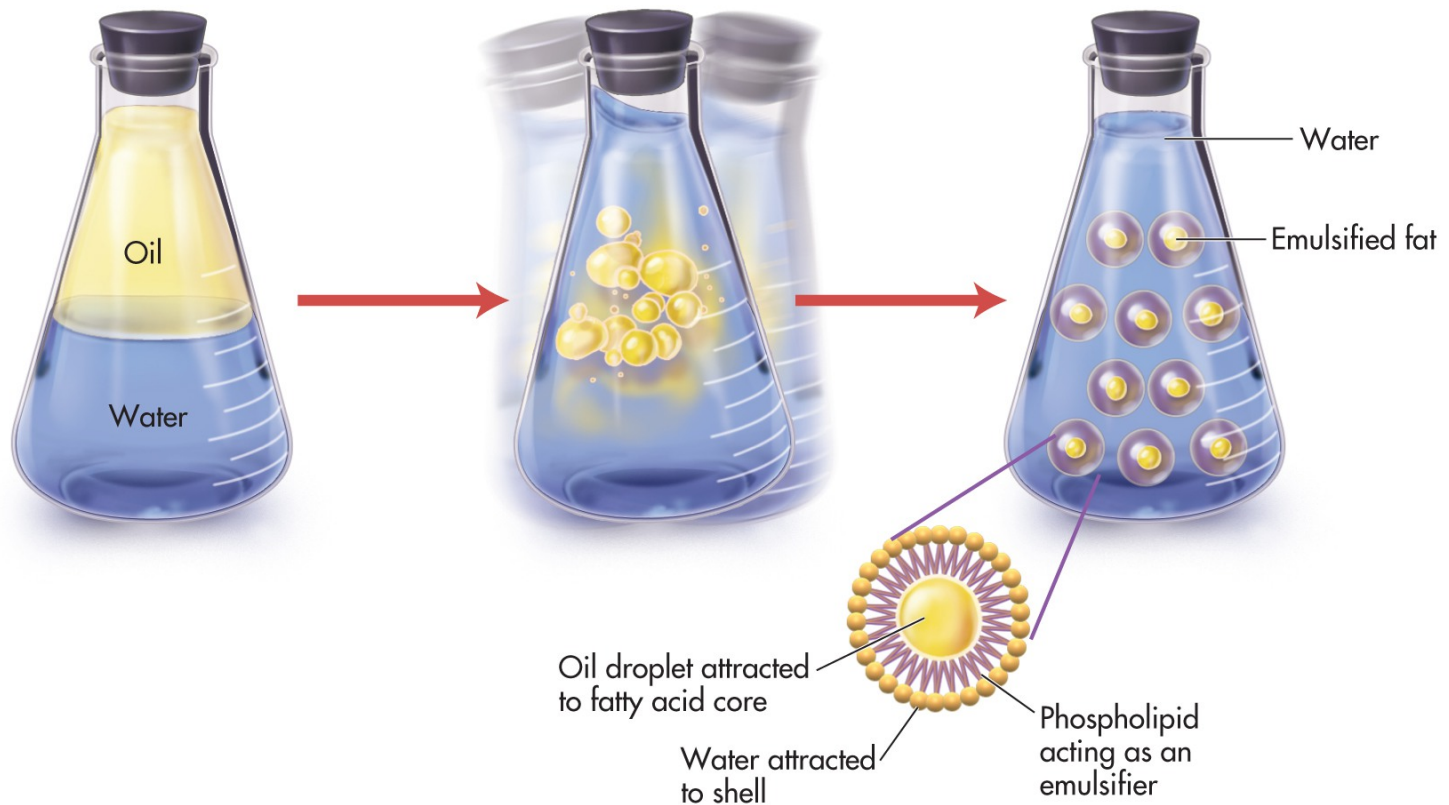
Lipids



- Hydrophobic quality of lipids makes their digestion and absorption more complicated than carbohydrates and proteins
- **Lipases** – fat digesting enzymes
 - **lingual lipase** secreted by the intrinsic salivary glands of the tongue /// generally thought to be inactive in mouth, but activated by stomach acids // acidic enzyme
 - stomach's chief cells also produce **gastric lipase** /// 10-15% of lipids digested before reaches duodenum // acidic enzyme // more important for neonates
 - **pancreatic lipase** – in the small intestine digest most of the fats in food
 - fat enters duodenum as large globules exposed to lipase only at their surface
 - globules broken up into smaller **emulsification droplets** by certain components of **bile** // **lecithin** and **bile acids**

Lipids

Emulsifiers and Agitation in Salad Dressing





Lipids

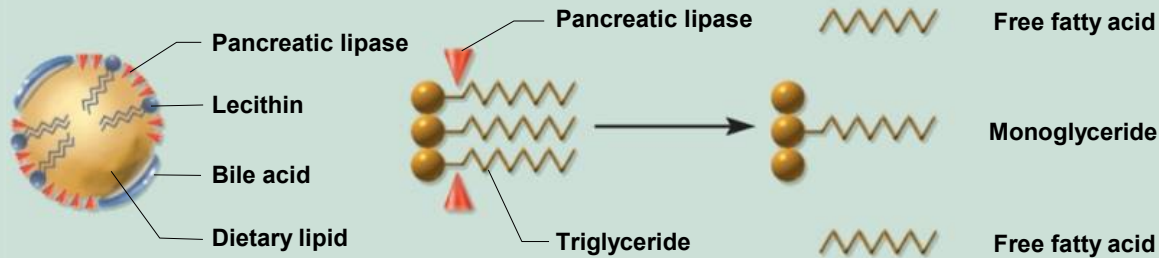
- agitation by **segmentation** breaks up the fats into droplets as small as 1 μm in diameter
- the coating of lecithin and bile acids keep it broken up
- exposing far more of its surface to enzymatic action
- there is enough pancreatic lipase in the small intestine after a meal to digest the average daily fat intake in as little as 1 to 2 minutes
- **lipase acts on triglycerides**
 - removes the first and third fatty acids from glycerol backbone
 - leaves the middle one
 - the product of lipase action are **two free fatty acids (FFAs)** and a **monoglyceride**



Fat Hydrolysis and Micelles

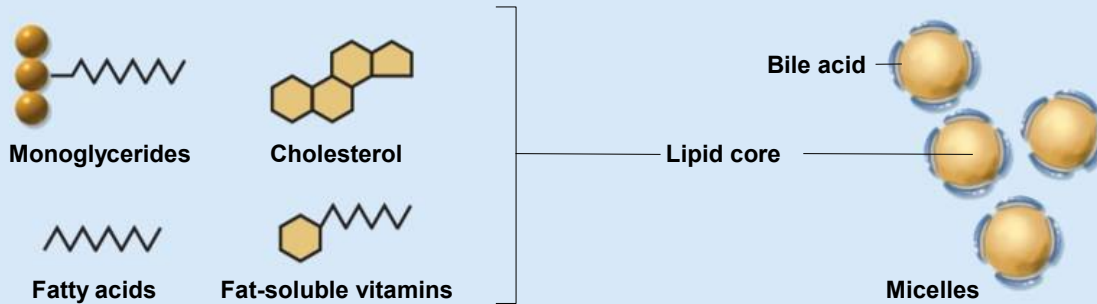
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Fat hydrolysis

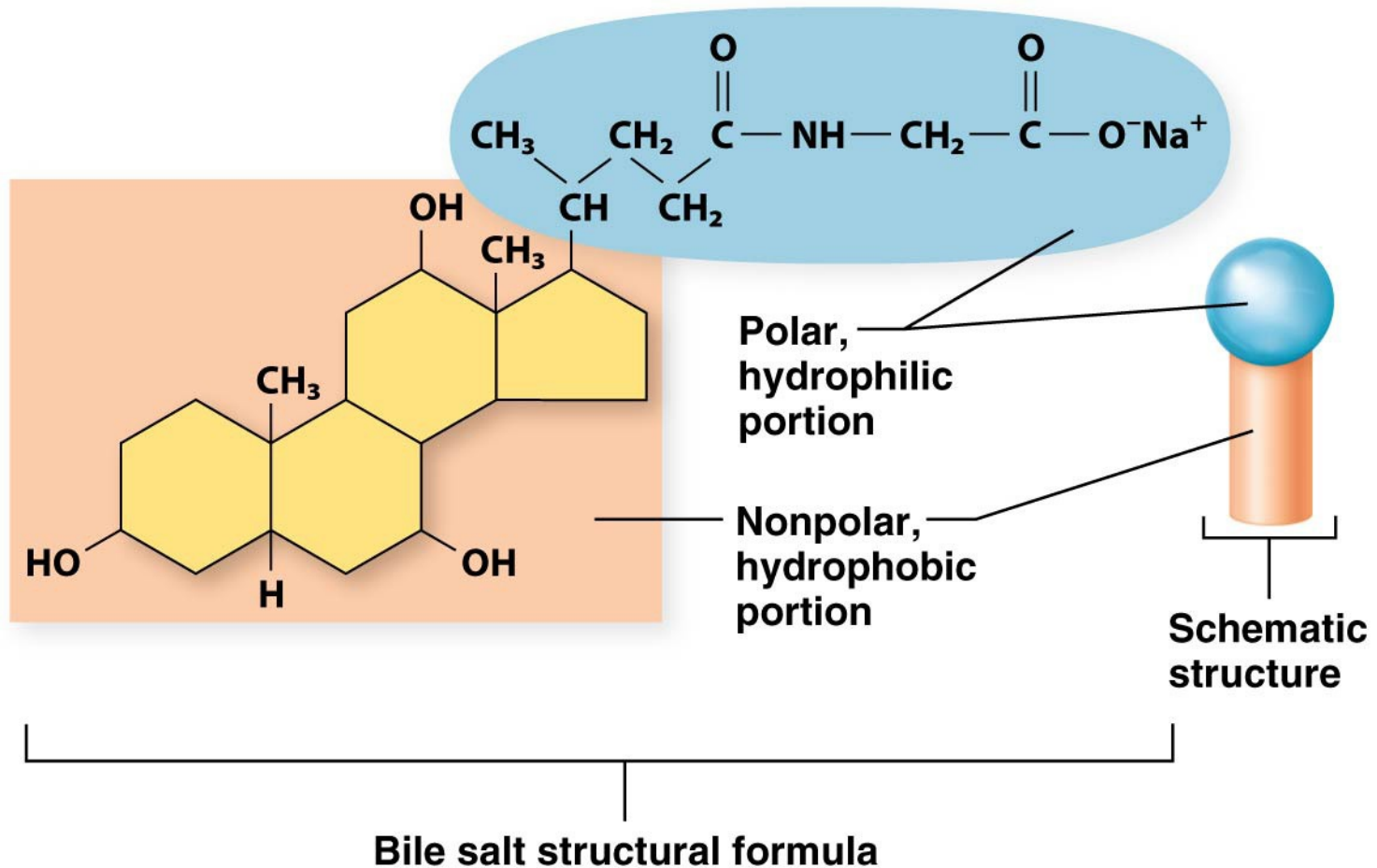


Emulsification droplets are acted upon by pancreatic lipase, which hydrolyzes the first and third fatty acids from triglycerides, usually leaving the middle fatty acid.

Lipid uptake by micelles



Micelles in the bile pass to the small intestine and pick up several types of dietary and semidigested lipids.



Micelles Role in Lipid Digestion and Absorption

- absorption of free fatty acids, monoglycerides, and other lipids depends on minute droplets in the bile called micelles
 - Micelles are formed in in the liver as well as in the duodenum
 - consist of **20 to 40 bile acid molecules** aggregated with their hydrophilic side groups facing outward and their hydrophobic steroid rings facing inward
 - **bile phospholipids and cholesterol** diffuse into the center of the micelle to form its core

Micelles Role in Lipid Digestion and Absorption

micelles pass down the bile duct into the duodenum /// where they absorb fat soluble vitamins, more cholesterol, and the FFAs and monoglycerides produced by fat digestion

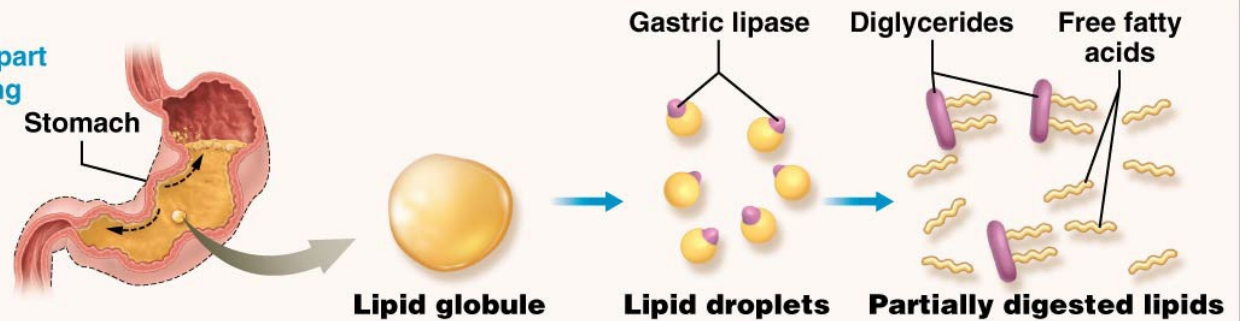
they transport lipids to the surface of the intestinal absorptive cells

lipids leave the micelles and diffuse through the plasma membrane into the cells

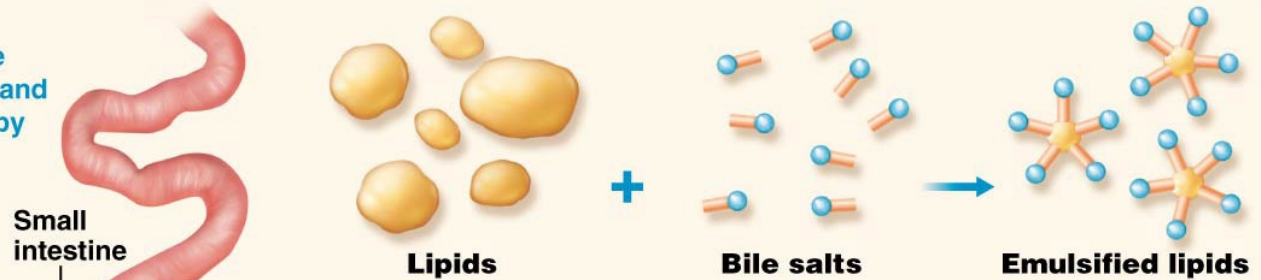
micelles are reused (**enterohepatic circulation**) /// Reabsorbed in ileum / go back to liver via hepatic portal circulation / returned to bile / secreted back into duodenum

Digestion of lipids.

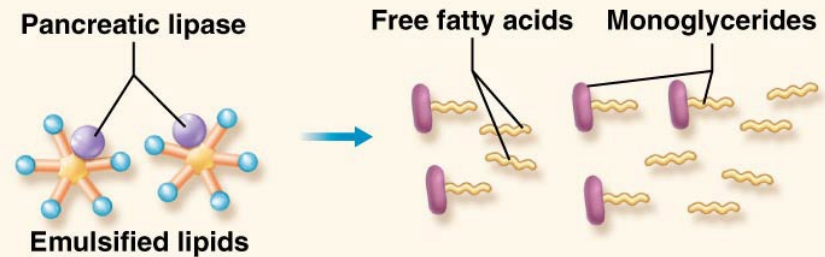
- ① Lipids are broken apart by stomach churning and broken down in reactions catalyzed by gastric lipase.



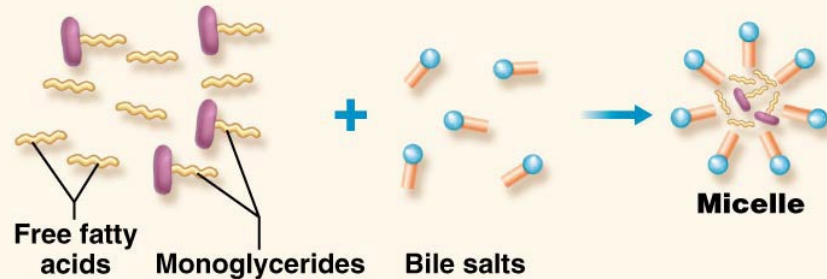
- ② Lipids enter the small intestine and are emulsified by bile salts.



- ③ Pancreatic lipase catalyzes reactions that digest the lipids into free fatty acids and monoglycerides.

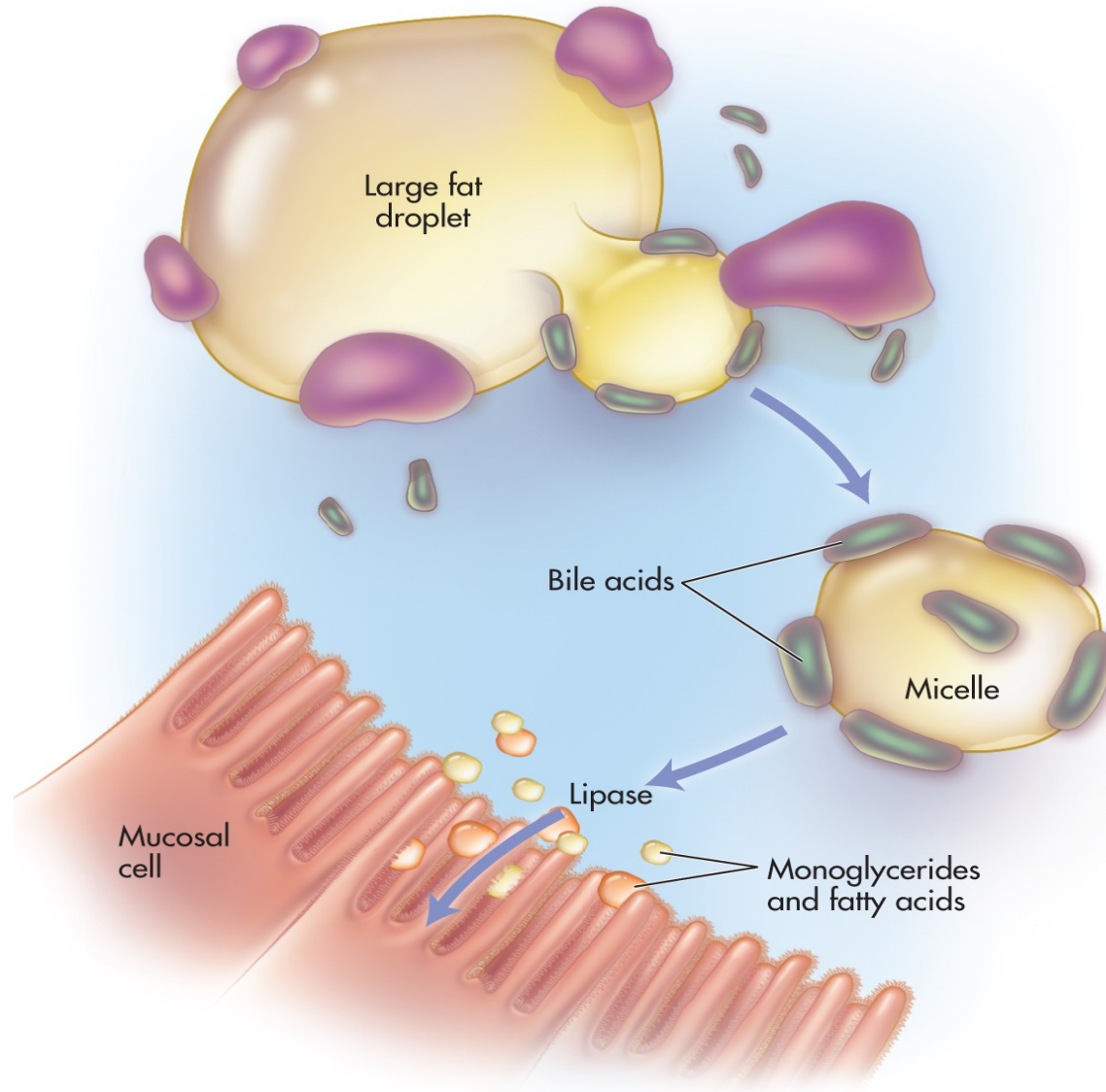


- ④ Bile salts remain associated with the digested lipids to form micelles.

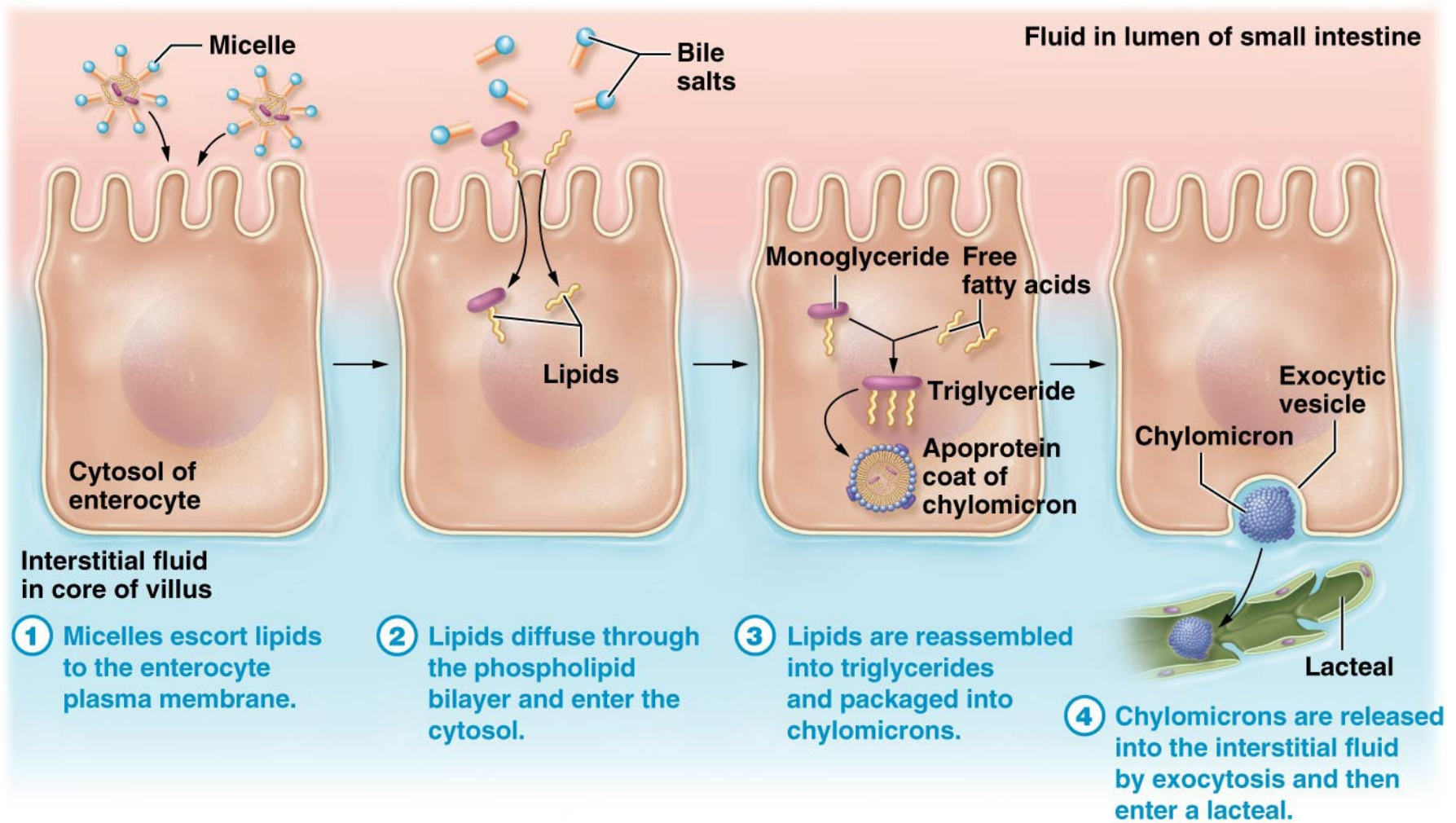


Fat Digestion and Absorption

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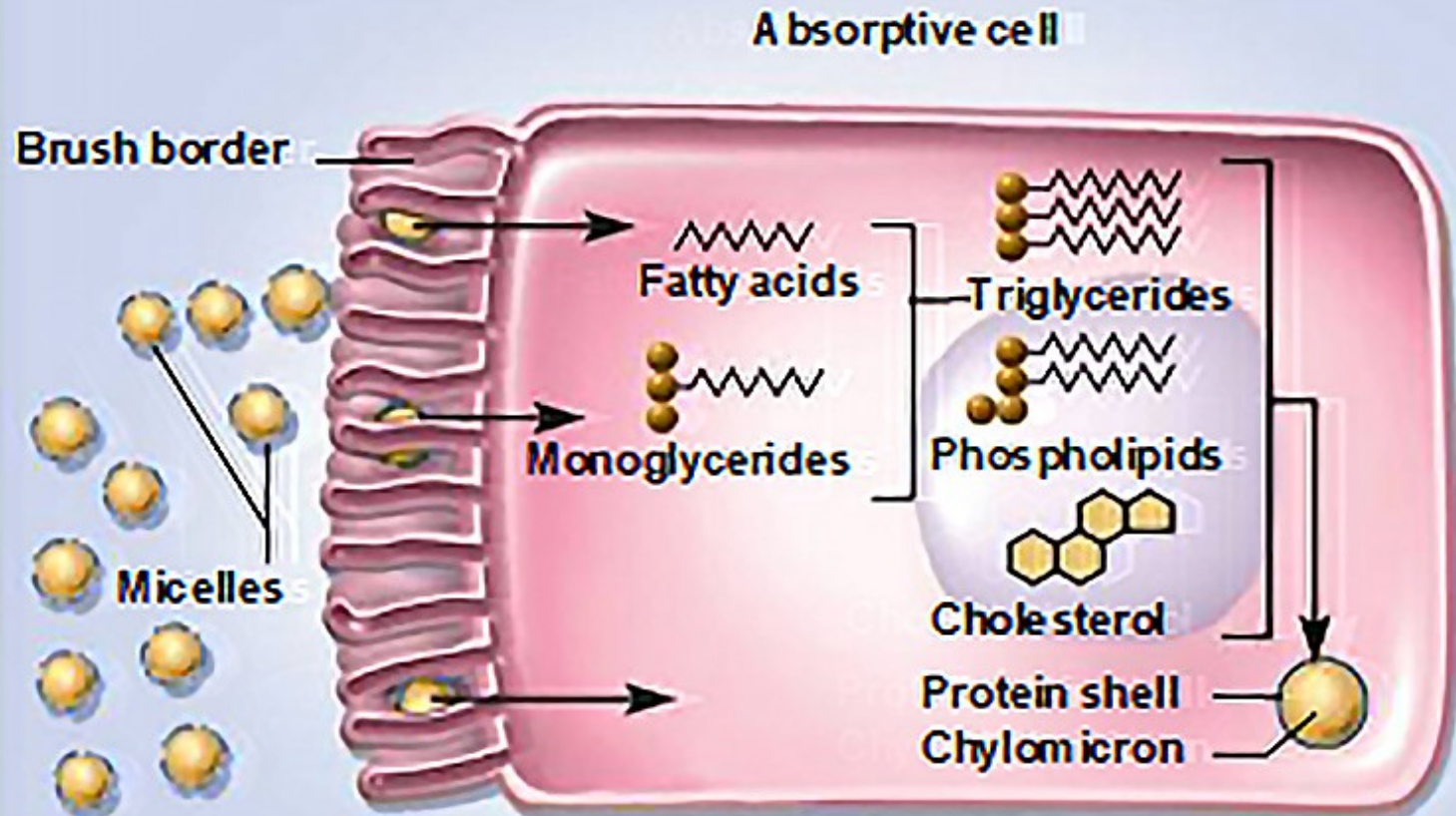


Lipid absorption in the small intestine.



Lipid Absorption

Chylomicron formation



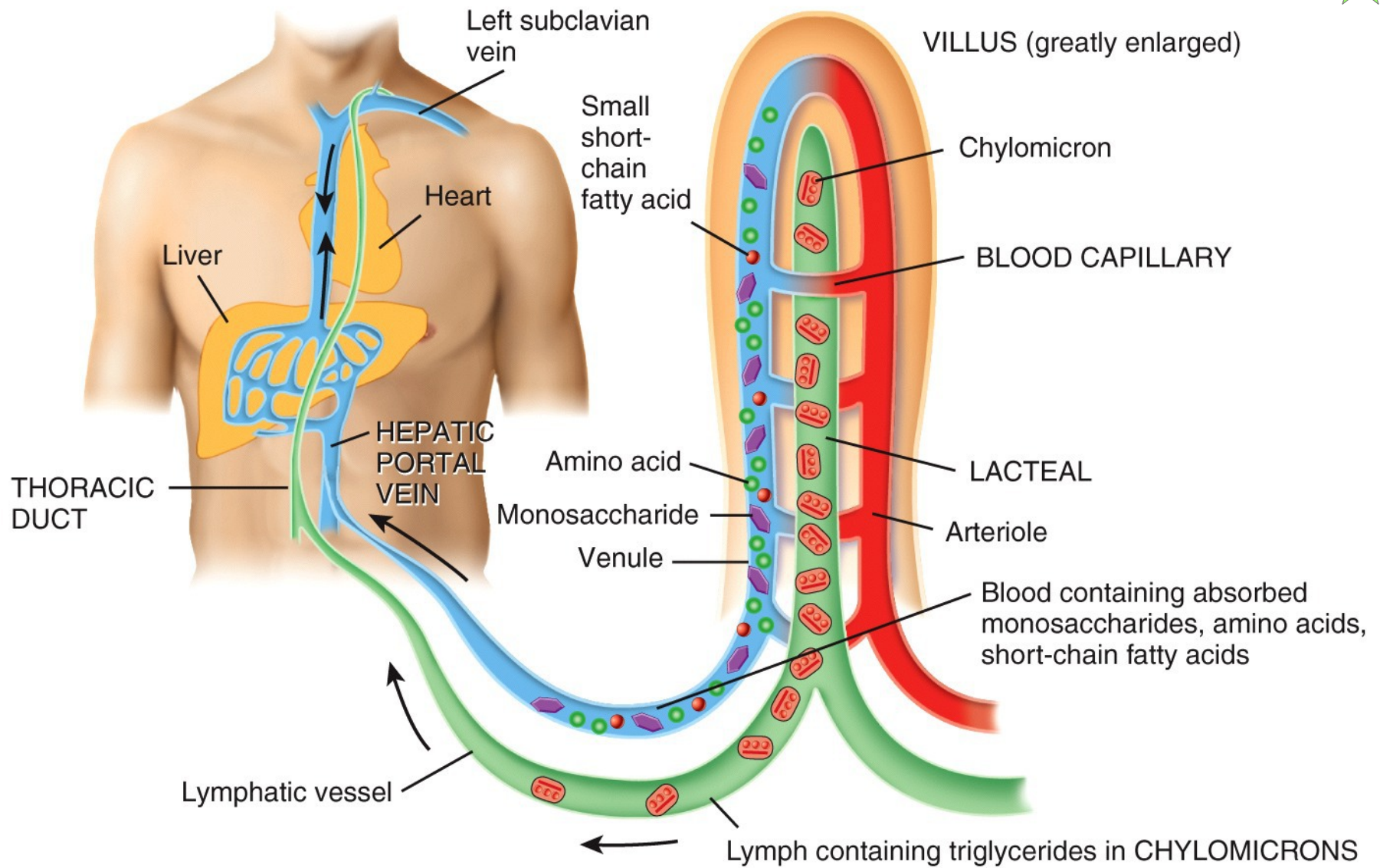
Intestinal cells absorb lipids from micelles, resynthesize triglycerides, and package triglycerides, cholesterol, and phospholipids into protein-coated chylomicrons.

Lipid Absorption



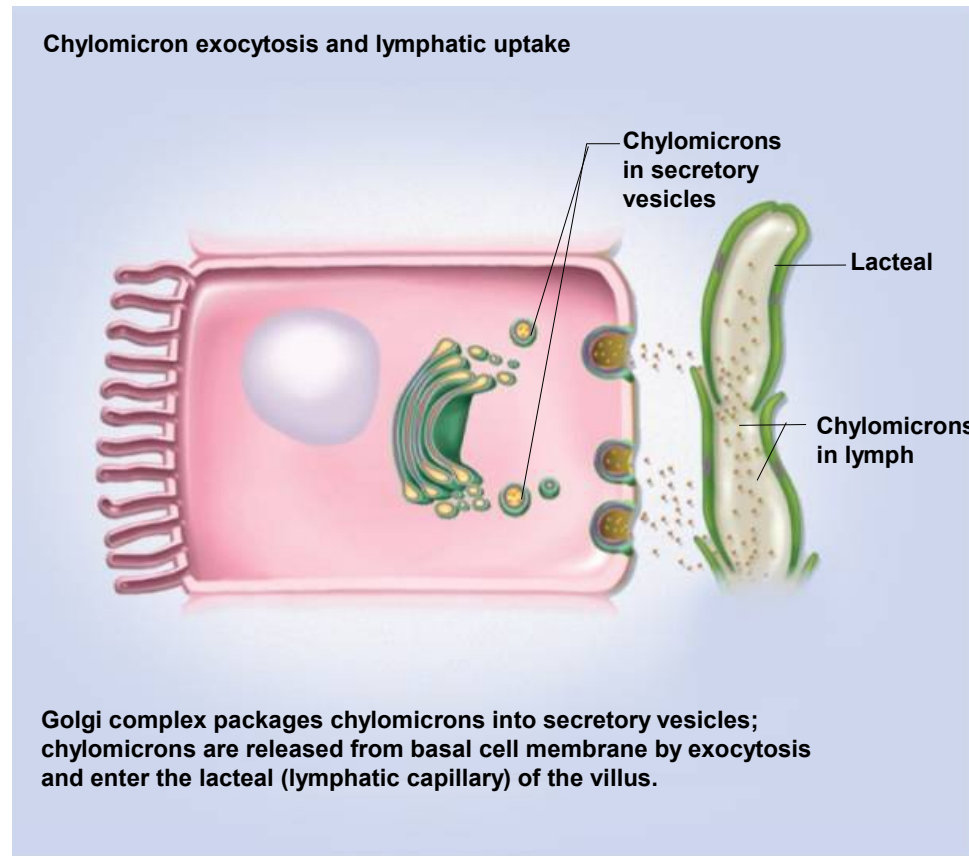
Within the jejunum absorptive cells

- free fatty acids and monoglycerides are transported to the smooth ER
- resynthesized into triglycerides
- golgi complex coats these with phospholipids and protein to form chylomicrons
- Chylomicrons loaded with fat soluble molecules including triglycerides, lecithin, fat soluble vitamins, etc.
 - packaged into secretory vesicles that migrate to basal surface of cell /// release their contents into the core of the villus (lamina propria) /// taken up by more porous lacteal into the lymph /// **white, fatty intestinal lymph (chyle)** flows into larger and larger lymphatic vessels until they reenter the bloodstream



(b) Movement of absorbed nutrients into blood and lymph

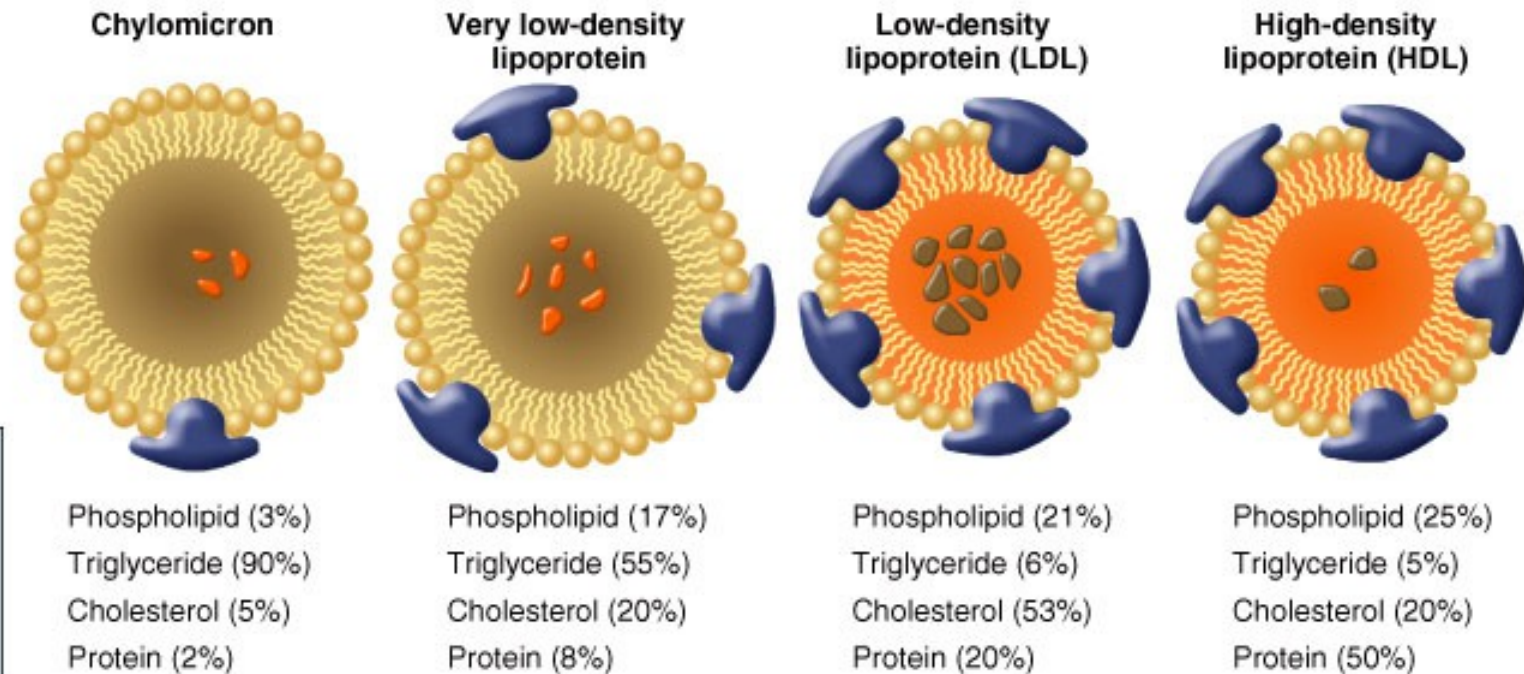
Chylomicrons and the Lymphatics



Chylomicrons are released into the lymphatic system in the lacteals of the villi. They enter the bloodstream when lymphatic fluid enters the subclavian vein via the thoracic duct.

Serum Lipoproteins

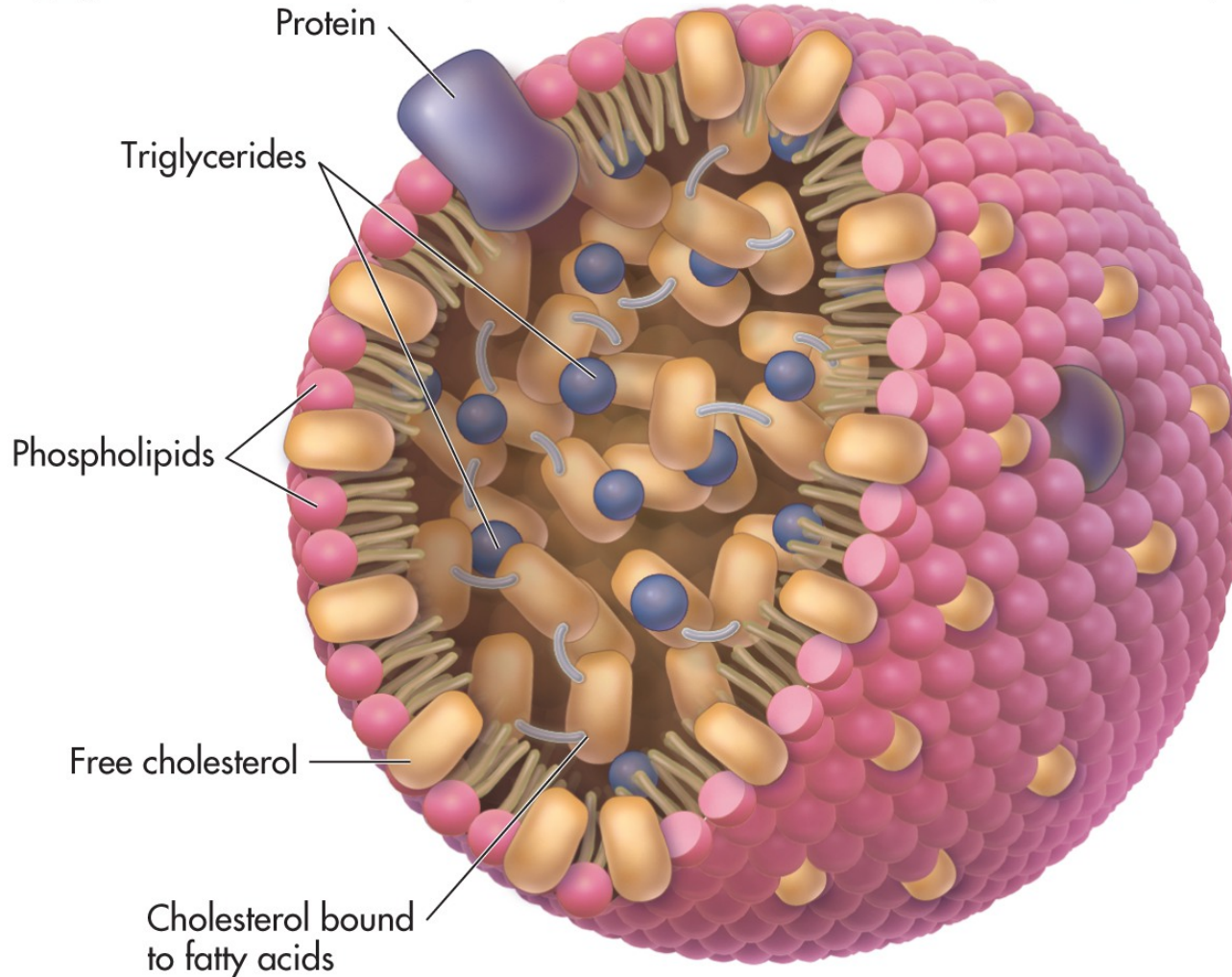
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(a) Lipoprotein types

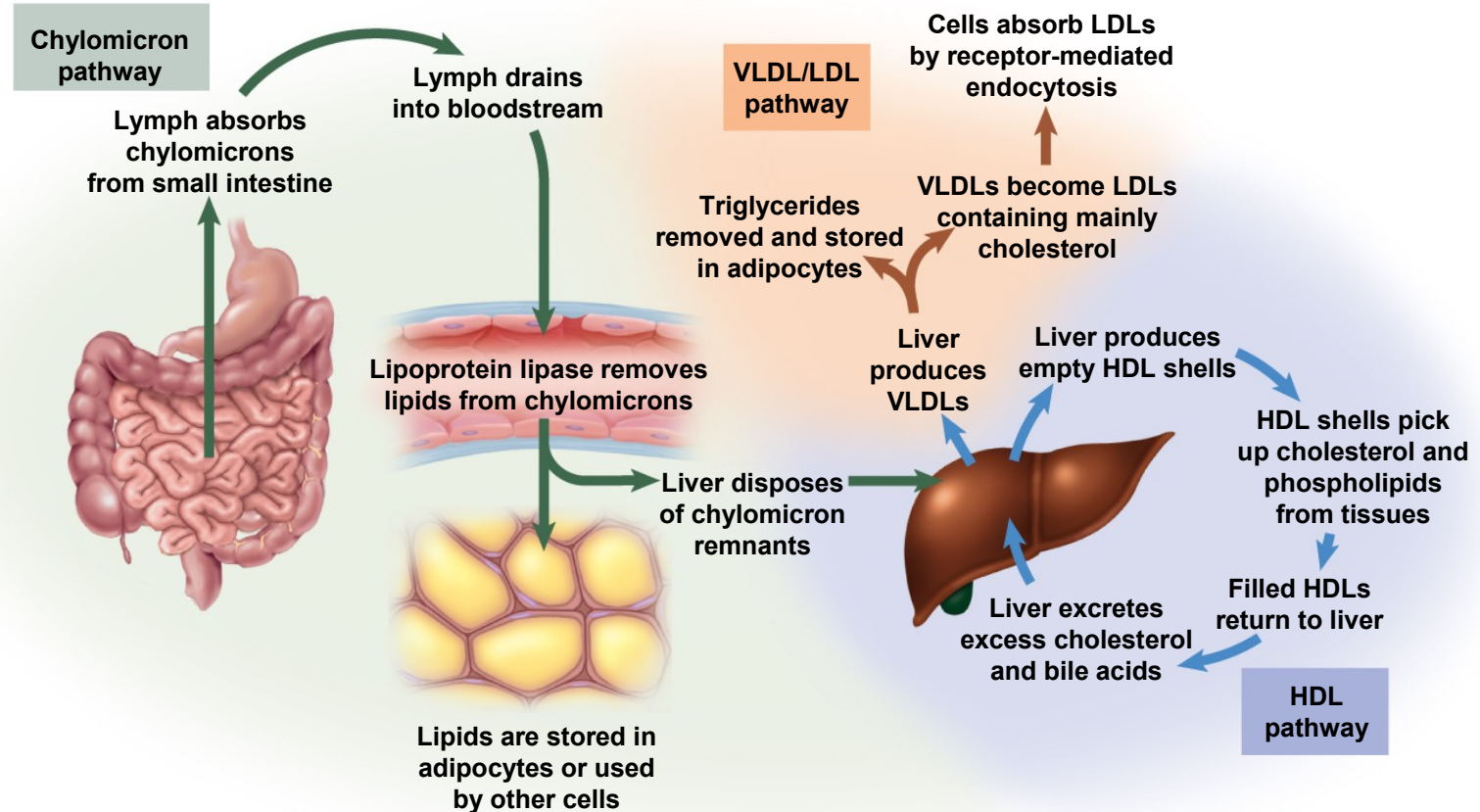
Composition and Roles of Lipoproteins

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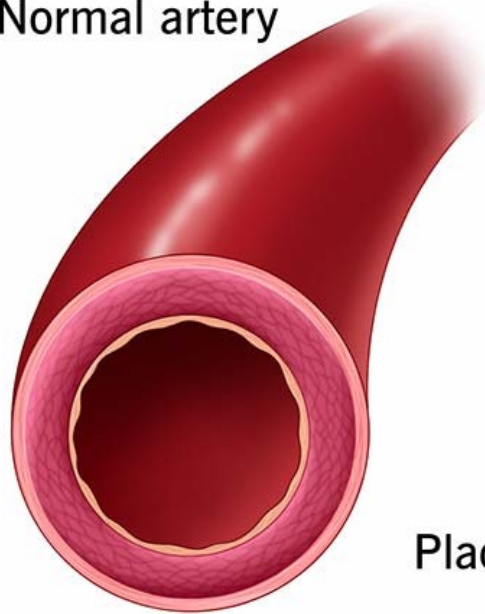


Lipoprotein Processing Pathways

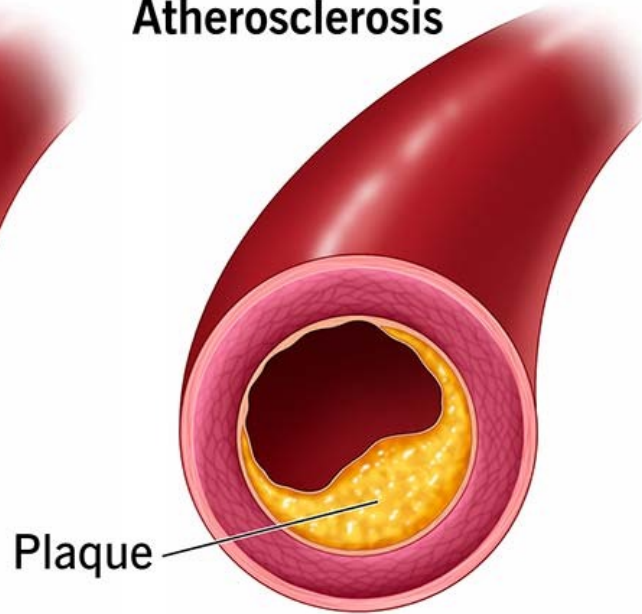


Atherosclerosis

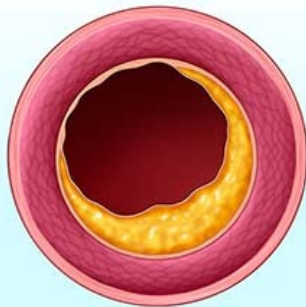
Normal artery



Atherosclerosis



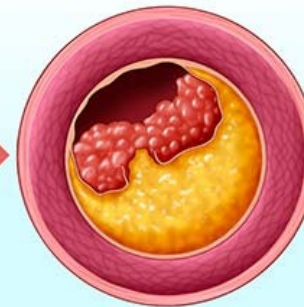
Progression



Formation

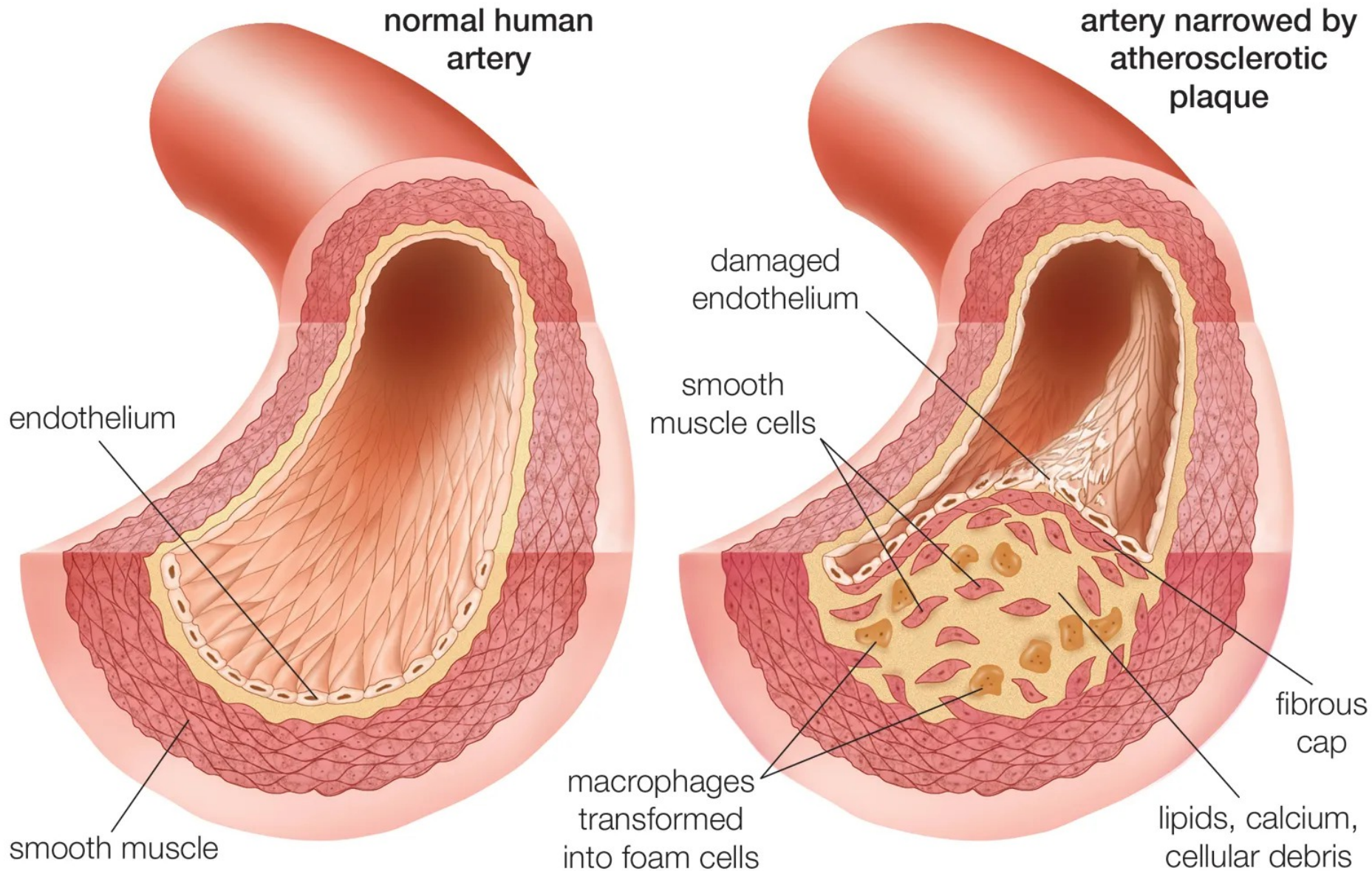


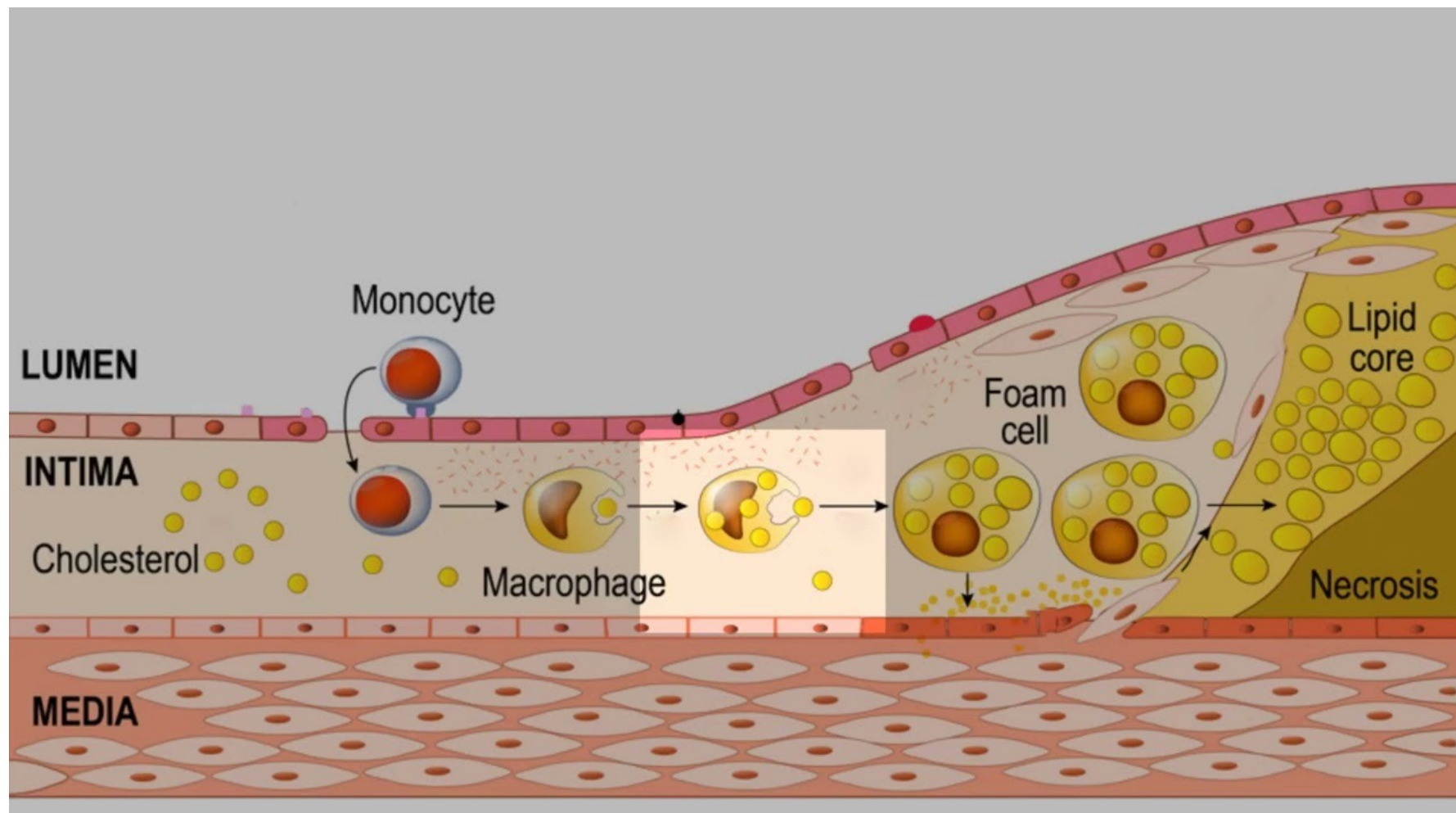
Growth and
rupture

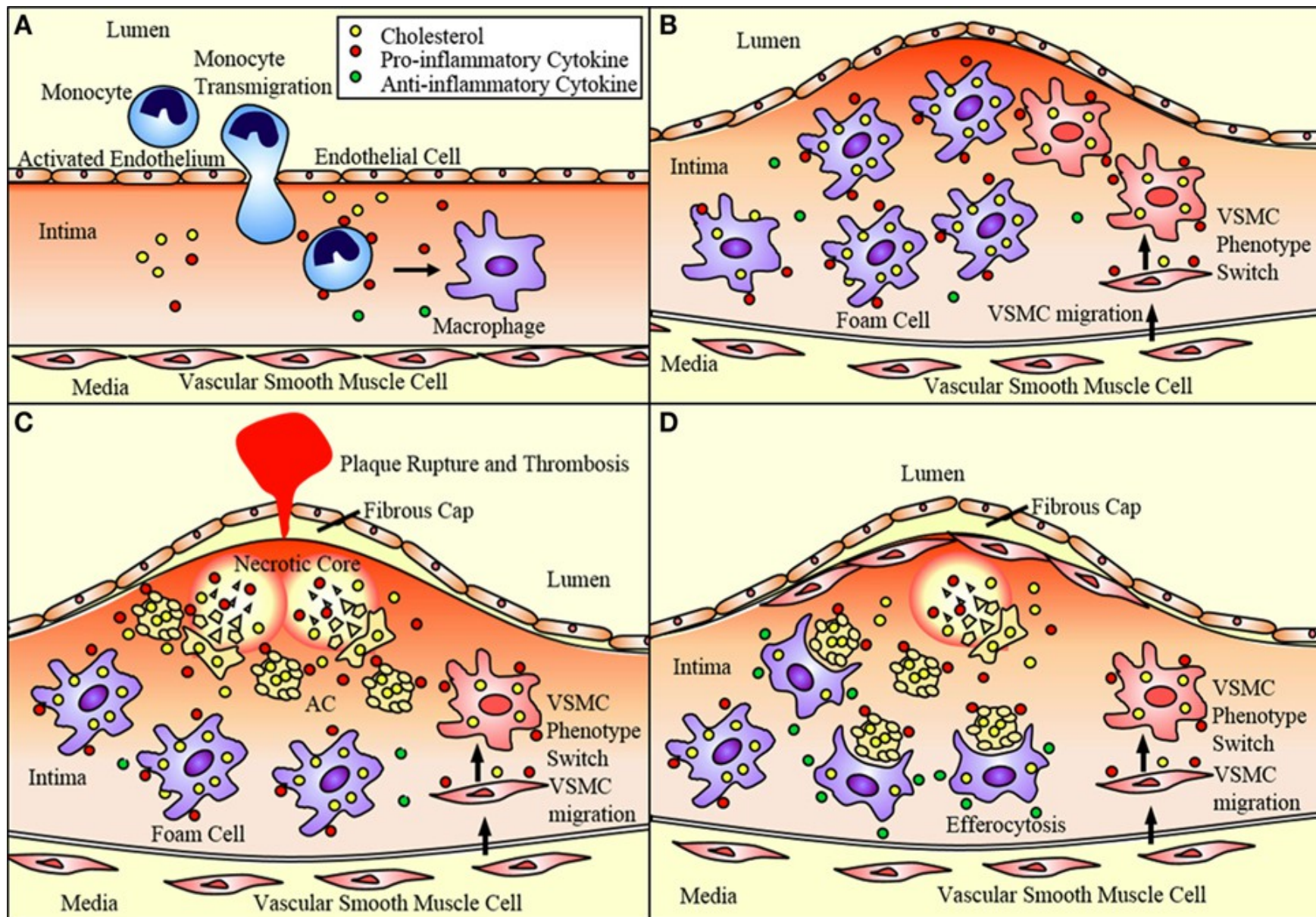


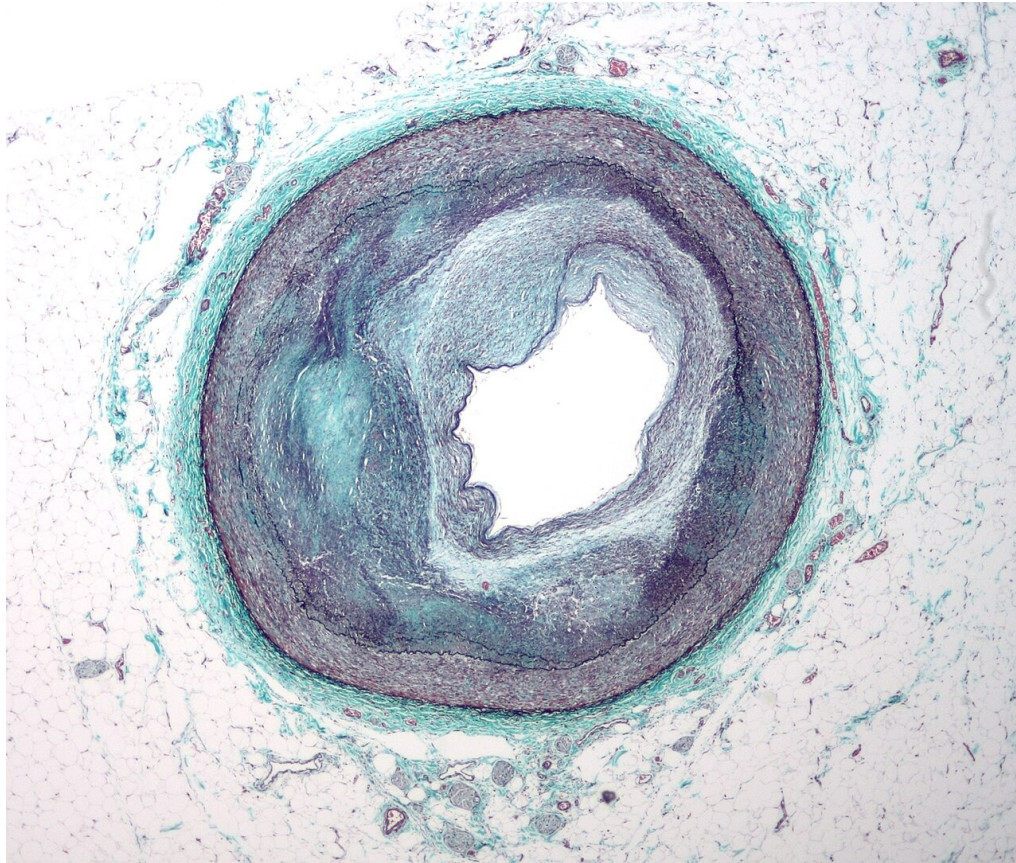
Blood clot

Atherosclerosis



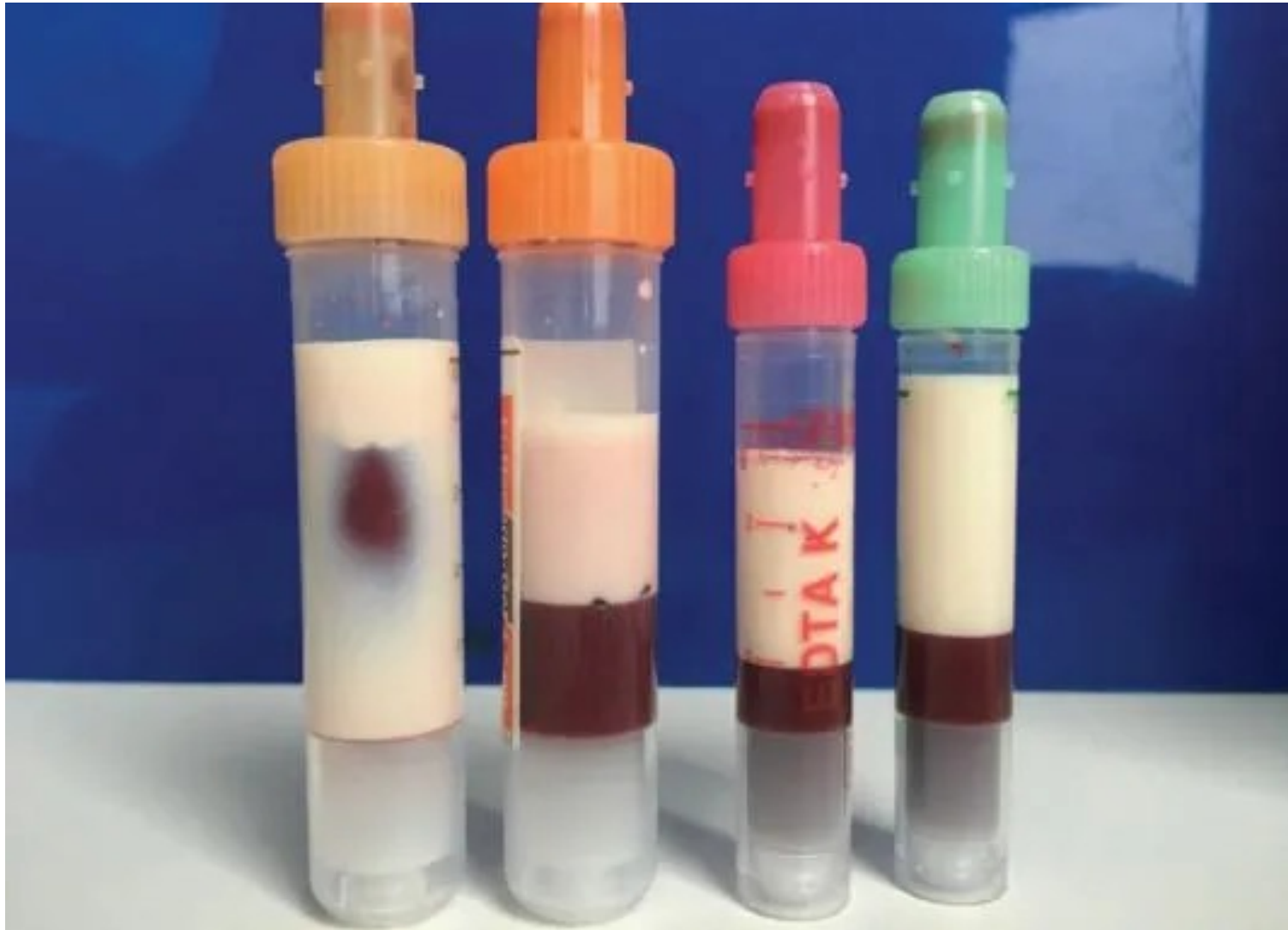






What's the difference between atherosclerosis and arteriosclerosis?

Atherosclerosis happens when your arteries become narrow due to a buildup of plaque. It's caused by inflammation in the arteries. Atherosclerosis progresses to arteriosclerosis, a condition in which your arteries become stiff. Arteriosclerosis is caused when arteries lose elasticity and calcium-phosphate salt move into the soft plaque to make it hard. (Oct 5, 2022)



A man in Germany had extraordinarily high levels of triglycerides, a type of fat, in his blood. Above, samples of the man's blood about two hours after they were drawn. The white is the fat.



A-72-37



Vitamins

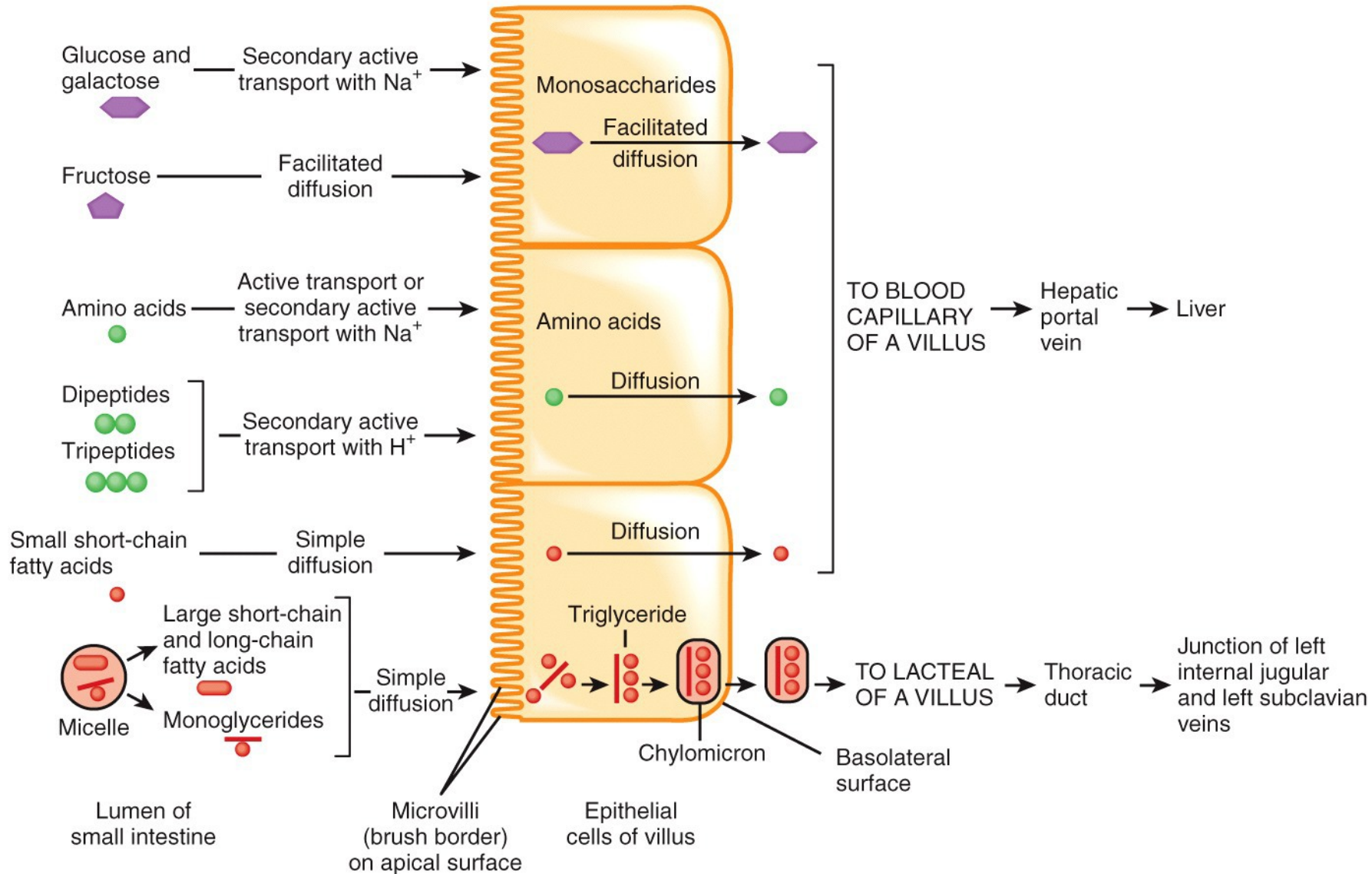


Vitamins are essential nutrients (must come from diet // we can not make them)

absorbed unchanged

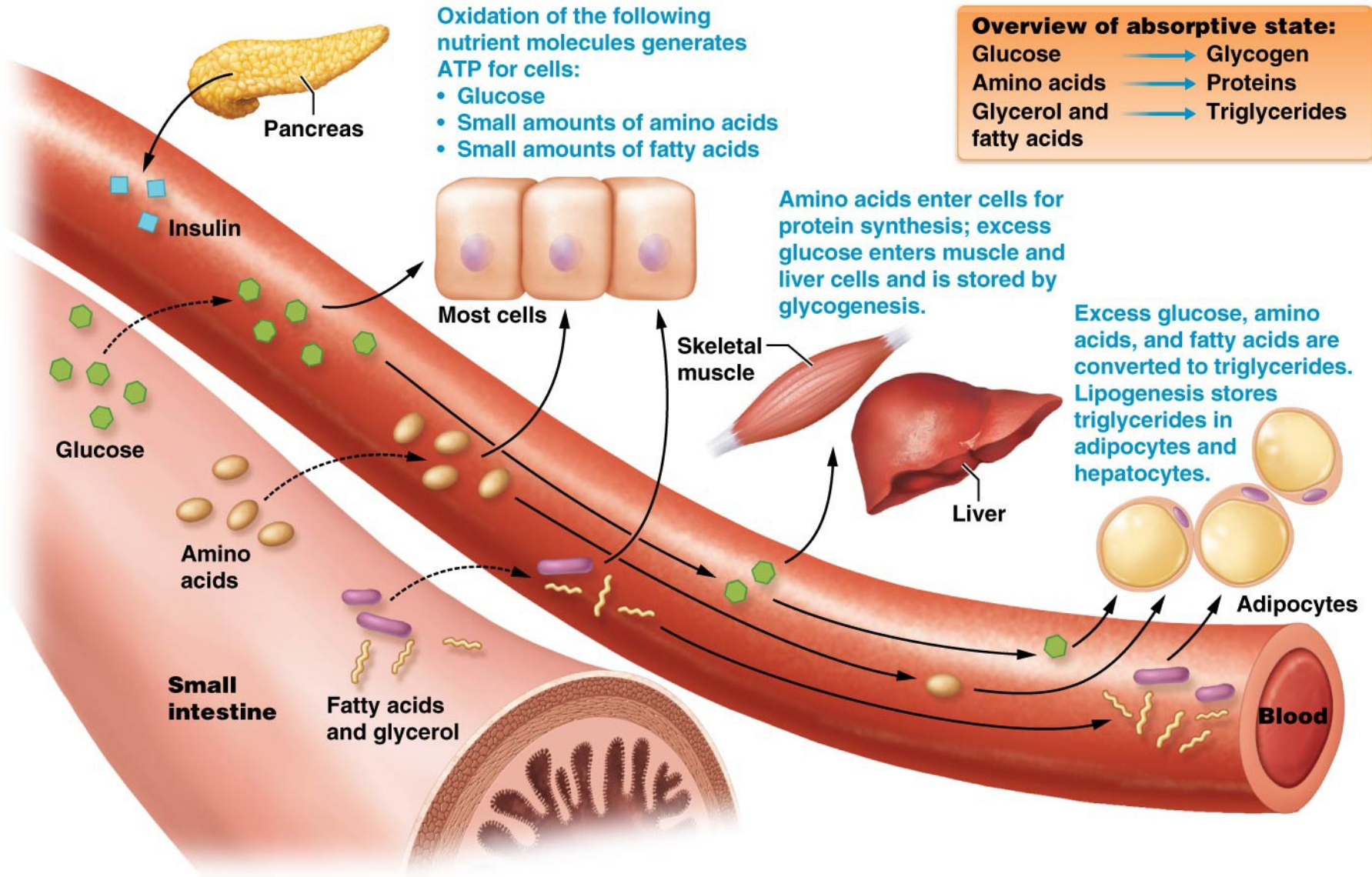
fat-soluble vitamins - A, D, E and K absorbed with other lipids // if they are ingested without fat-containing food, they are not absorbed at all, but are passed in the feces and wasted // Why do we put fatty dressing on our salads?

water-soluble vitamins, like B and C vitamins are absorbed by simple Diffusion /// B₁₂ requires gastric intrinsic factor to be absorbed into body



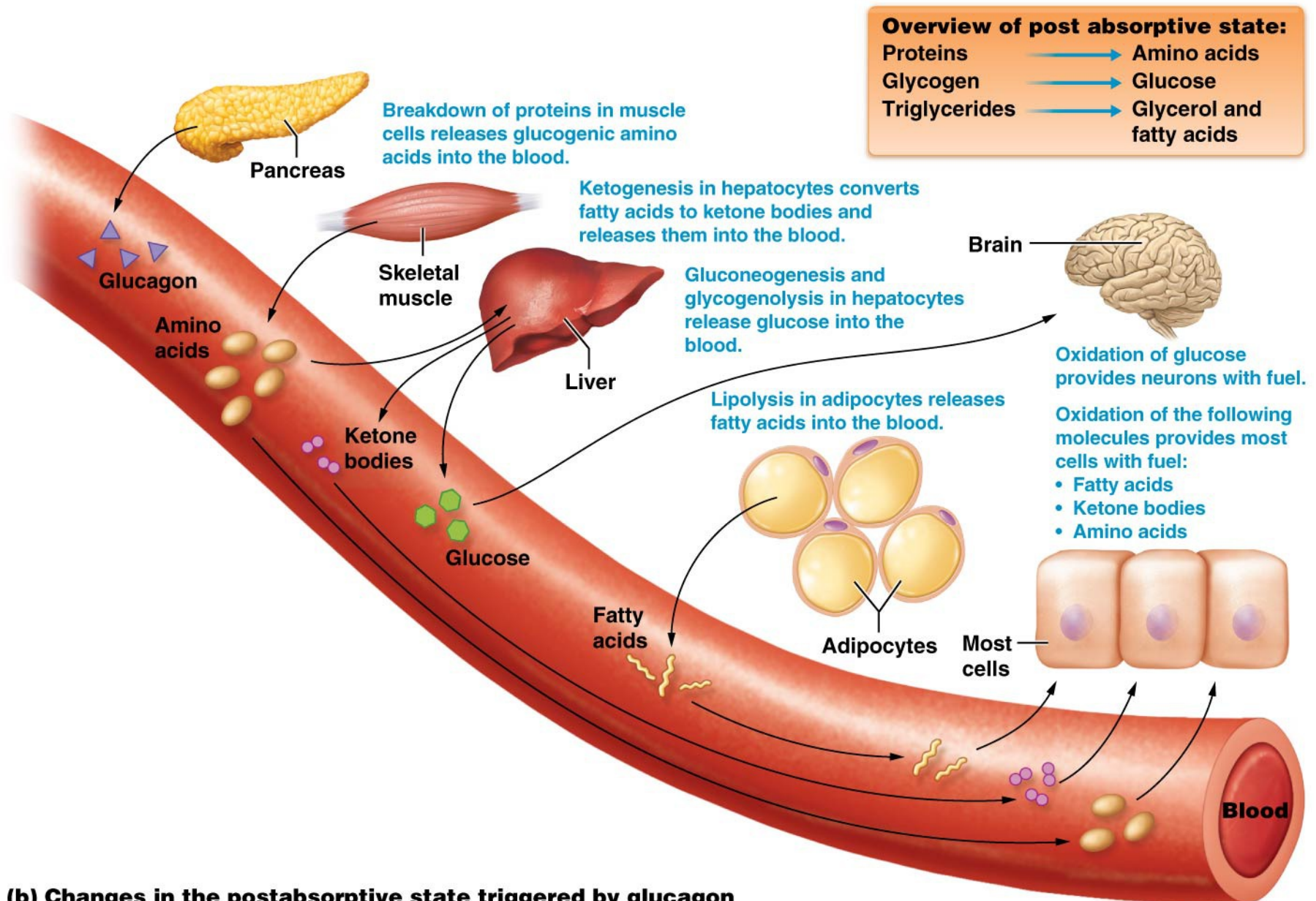
(a) Mechanisms for movement of nutrients through absorptive epithelial cells of villi

The Absorptive State



(a) Changes in the absorptive state triggered by insulin.

The Post Absorptive State



(b) Changes in the postabsorptive state triggered by glucagon



Hormonal Regulation of Digestion (1 of 3)

Gastrin // hormone // produced by G cells of gastric pit

Stimulate gastric secretions

Stimulate lower esophageal sphincter to contract //
prevents “heart burn”

Relax pyloric sphincter // allows small amount of chyme
to enter duodenum and

By relaxing pyloric sphincter and allowing only small
volumes of chyme to enters duodenum it then gives time
to process chyme



Hormonal Regulation of Digestion (2 of 3)

- **Chyme** leaving stomach stimulates duodenum's enteroendocrine cells to release **secretin** and **cholecystokinin**
 - These hormones stimulate different types of secretions from the pancreas and gall bladder
 - CCK suppress gastric secretion and interacts with hypothalamus to stimulate satiety // stimulate pancreas to produce and release more hormones // stimulate gall bladder to release bile
 - Another type of enteroendocrine cells secrete **glucose-dependent insulinotropic peptide (GIP)** /// this was called gastrin-inhibiting peptide
 - » stimulates insulin secretion in preparation for processing nutrients about to be absorbed by the small intestine // reduces gastric contractions



Hormonal Regulation of Digestion (2 of 2)

- **cholecystokinin (CCK)** - secreted by mucosa of duodenum in response to arrival of fats in small intestine
 - stimulate pancreatic acini to **secrete enzymes**
 - strongly stimulates gall bladder
 - induces **contractions of the gallbladder and relaxation of hepatopancreatic sphincter** causing discharge of bile into the duodenum
- **secretin** - released from duodenum in response to acidic chyme arriving from the stomach
 - stimulates ducts in both liver and pancreas **to secrete more sodium bicarbonate**
 - raising pH to level pancreatic and intestinal digestive enzymes require

Small Intestine VS Large Intestine

- Small intestine functions:
 - Digestion (mechanical and chemical)
 - Absorption of nutrients across mucosa /// active and passive
 - 99.9 percent of nutrients absorbed in small intestine
- Large intestine functions:
 - Eliminate food residue
 - Compaction
 - Recover water

INGESTED
AND SECRETED

ABSORBED

Saliva
(1 liter)

Ingestion
of liquids
(2.3 liters)

Gastric juice
(2 liters)

Bile
(1 liter)

Pancreatic
juice
(2 liters)

Intestinal
juice
(1 liter)

Total ingested
and secreted
= 9.3 liters

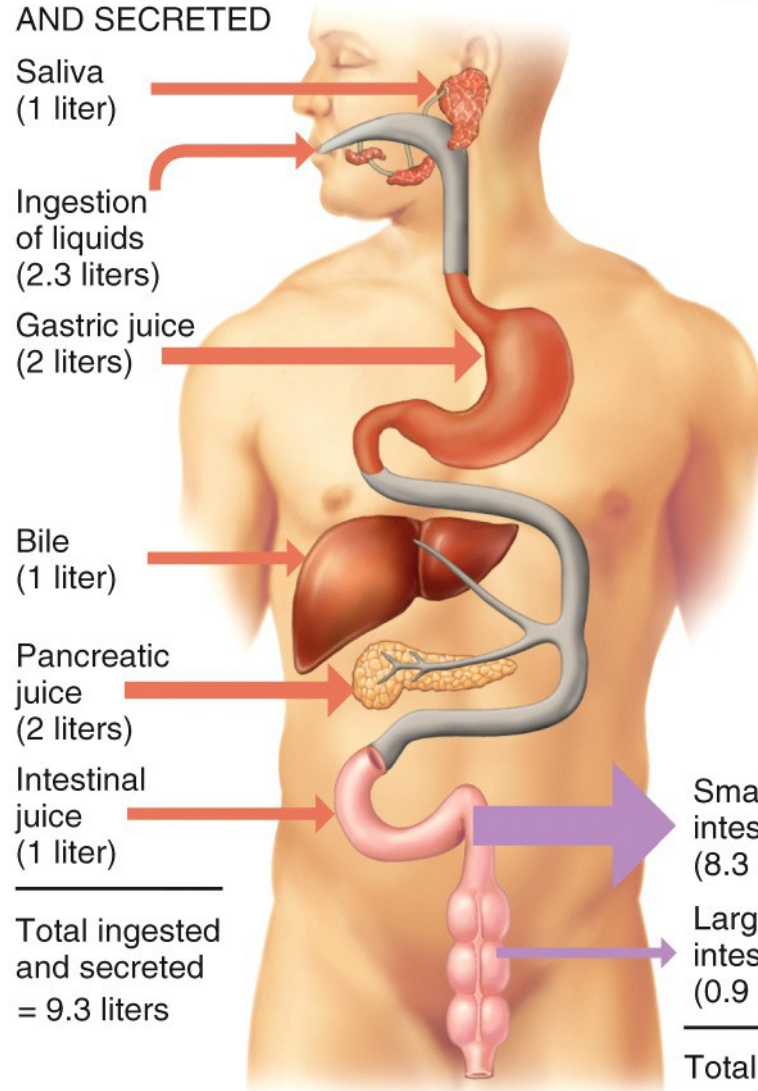
Small
intestine
(8.3 liters)

Large
intestine
(0.9 liters)

Total absorbed
= 9.2 liters

Excreted in feces
(0.1 liter)

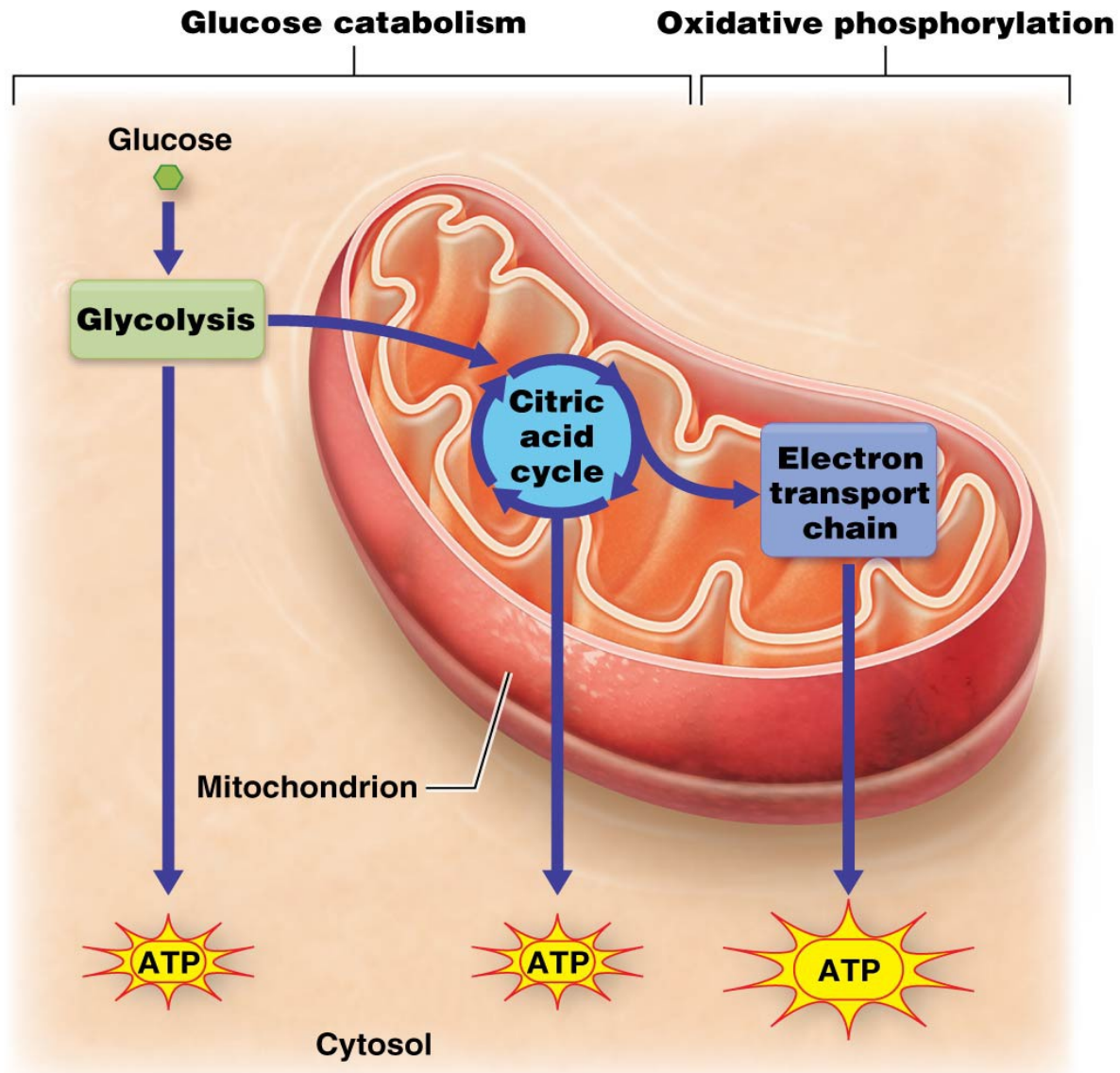
Fluid balance in GI tract



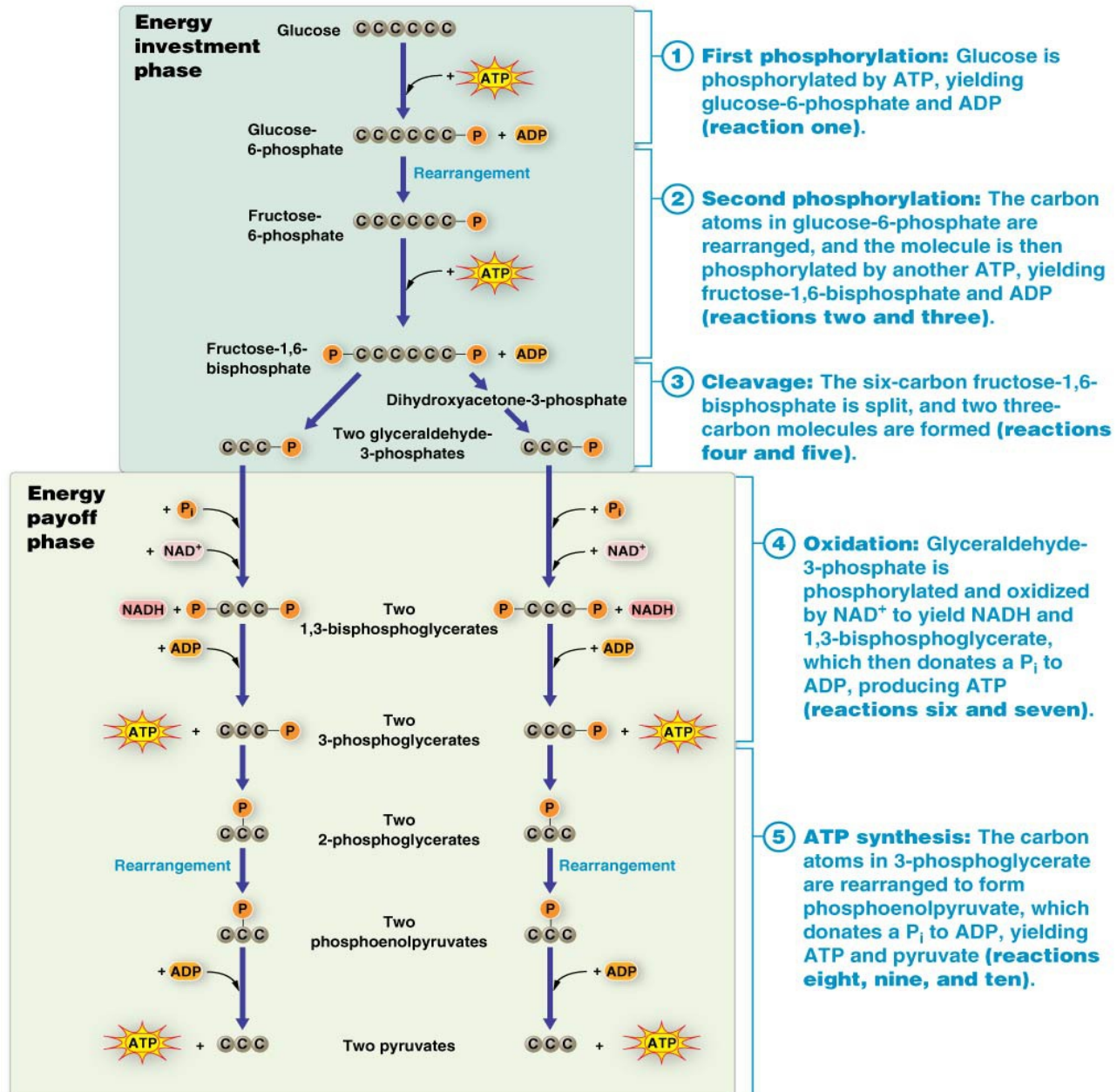
Review of Metabolism

- > Glycolysis
- > Krebs's Cycle
- > Electron Transport Chain
- > Anabolism VS Catabolism

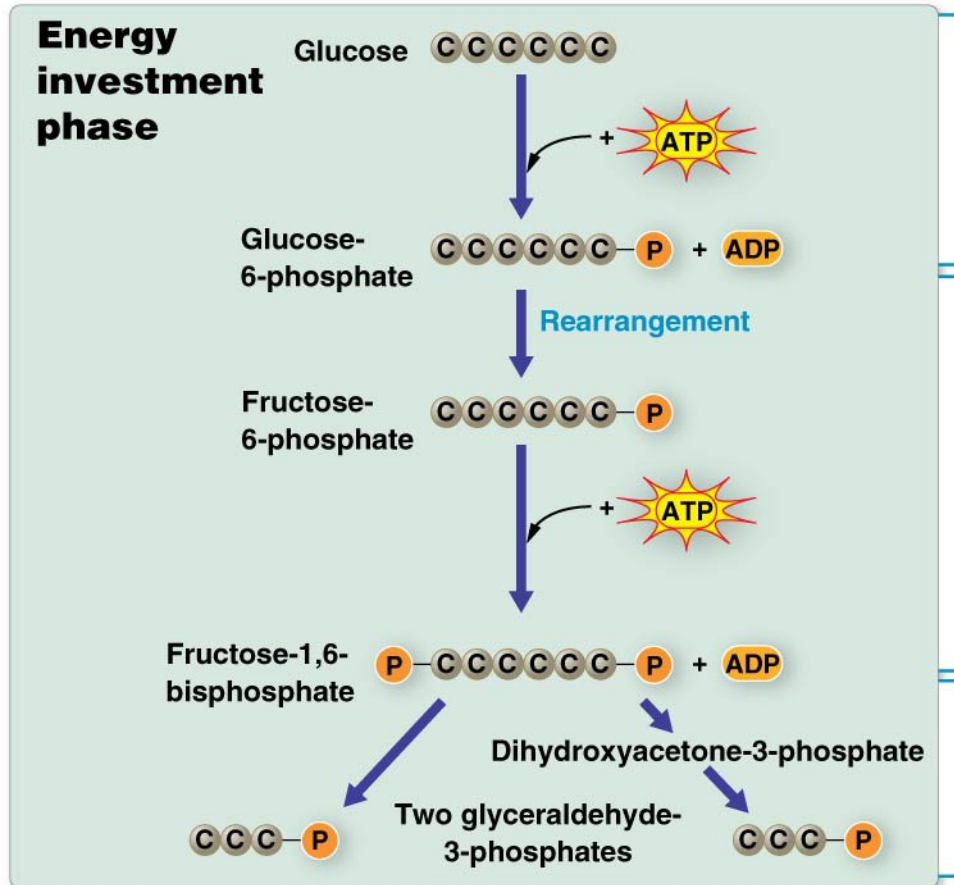
Overview of glucose catabolism and oxidative phosphorylation.



The reactions of glycolysis.

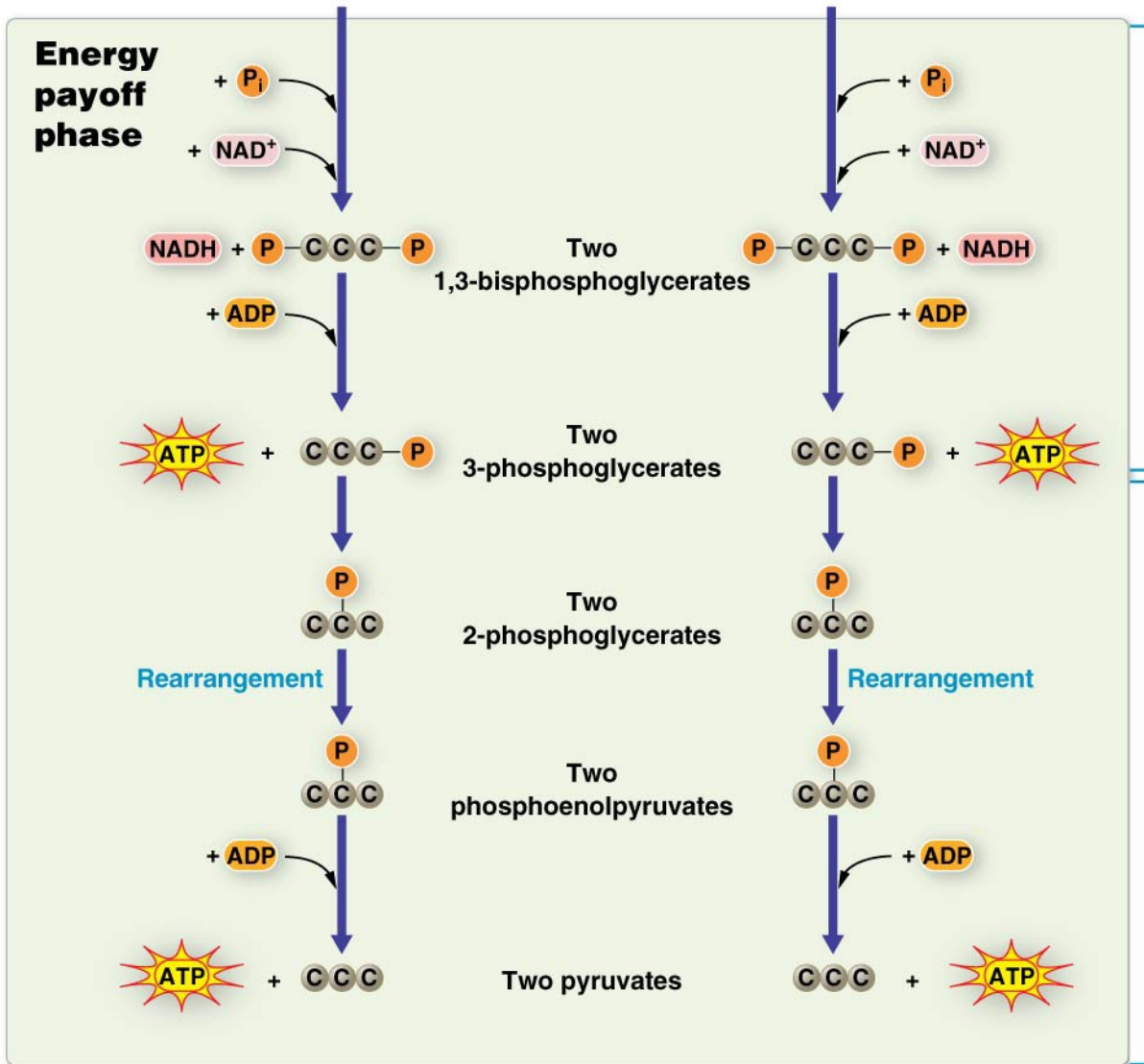


The reactions of glycolysis.



- 1 First phosphorylation:** Glucose is phosphorylated by ATP, yielding glucose-6-phosphate and ADP (reaction one).
- 2 Second phosphorylation:** The carbon atoms in glucose-6-phosphate are rearranged, and the molecule is then phosphorylated by another ATP, yielding fructose-1,6-bisphosphate and ADP (reactions two and three).
- 3 Cleavage:** The six-carbon fructose-1,6-bisphosphate is split, and two three-carbon molecules are formed (reactions four and five).

The reactions of glycolysis.



4 Oxidation: Glyceraldehyde-3-phosphate is phosphorylated and oxidized by NAD^+ to yield NADH and 1,3-bisphosphoglycerate, which then donates a P_i to ADP , producing ATP (reactions six and seven).

5 ATP synthesis: The carbon atoms in 3-phosphoglycerate are rearranged to form phosphoenolpyruvate, which donates a P_i to ADP, yielding ATP and pyruvate (**reactions eight, nine, and ten**).

The Big Picture of Glucose Catabolism and Oxidative Phosphorylation



Cytosol

- ① **Glycolysis:** Glucose is split and oxidized during glycolysis, yielding two pyruvate, two NADH, and two ATP (see Figure 23.4).

GLYCOLYSIS

Glucose

2 NADH

+

2 Pyruvate

- ② **Intermediate step:** The two pyruvates are oxidized to two acetyl-CoA, producing two NADH (see Figure 23.5).

- ③ **Citric acid cycle:** The two acetyl-CoA are further oxidized in the citric acid cycle, yielding six NADH, two FADH₂, and two ATP (see Figure 23.6).

Intermembrane space

- ④ **Electron transport chain:** Electrons from NADH and FADH₂ enter the electron transport chain (ETC), powering the pumping of H⁺ into the intermembrane space (see Figure 23.7a).

- ⑤ **Oxidative phosphorylation:** H⁺ re-enter the matrix through ATP synthase, which releases ATP from the synthase enzyme (see Figure 23.7b).

2 Acetyl-CoA
+
2 NADH

CITRIC
ACID
CYCLE

6 NADH
+
2 FADH₂

ETC

H⁺ H⁺ H⁺ H⁺
H⁺ H⁺

2 ATP

34 ATP

Mitochondrial matrix

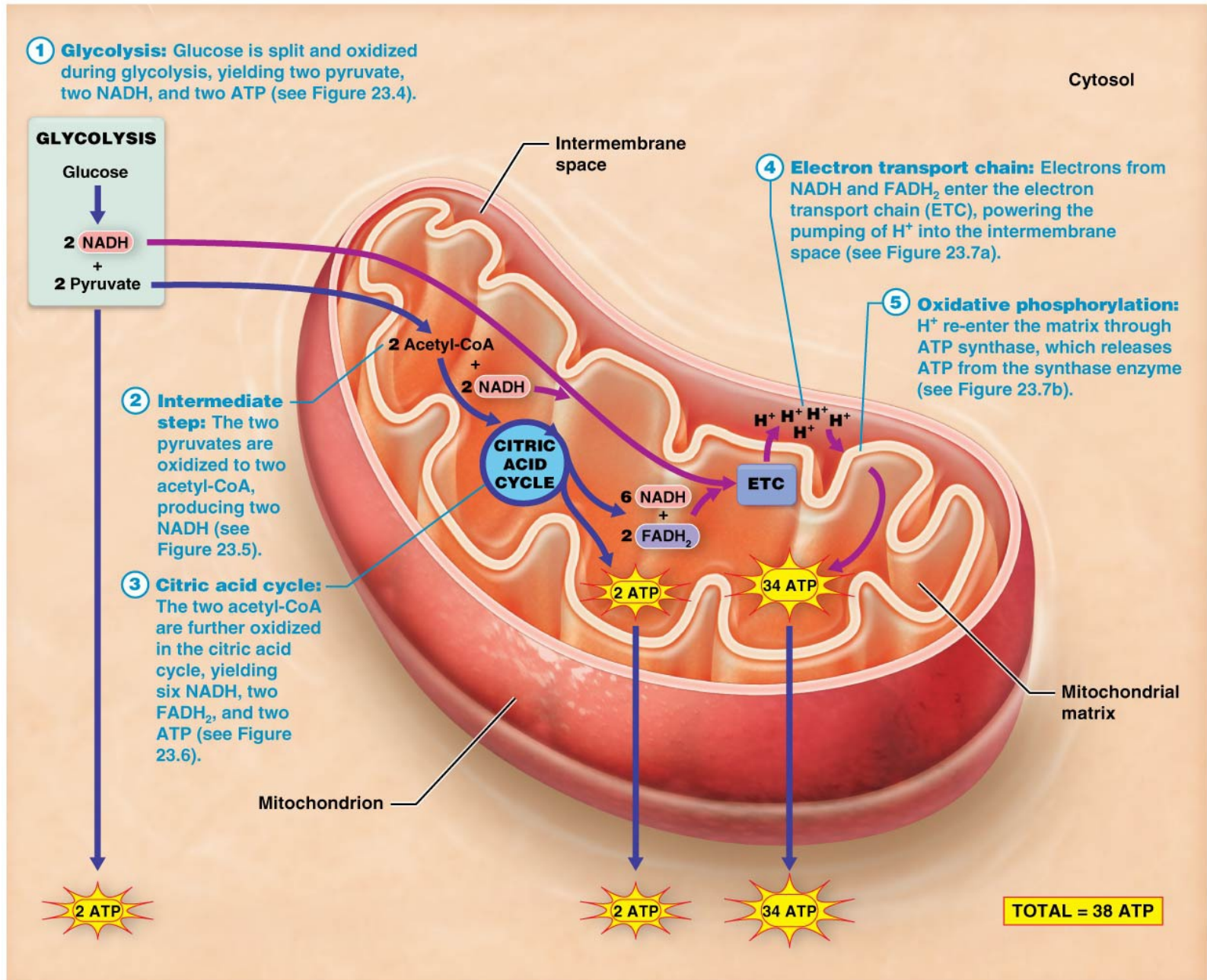
Mitochondrion

2 ATP

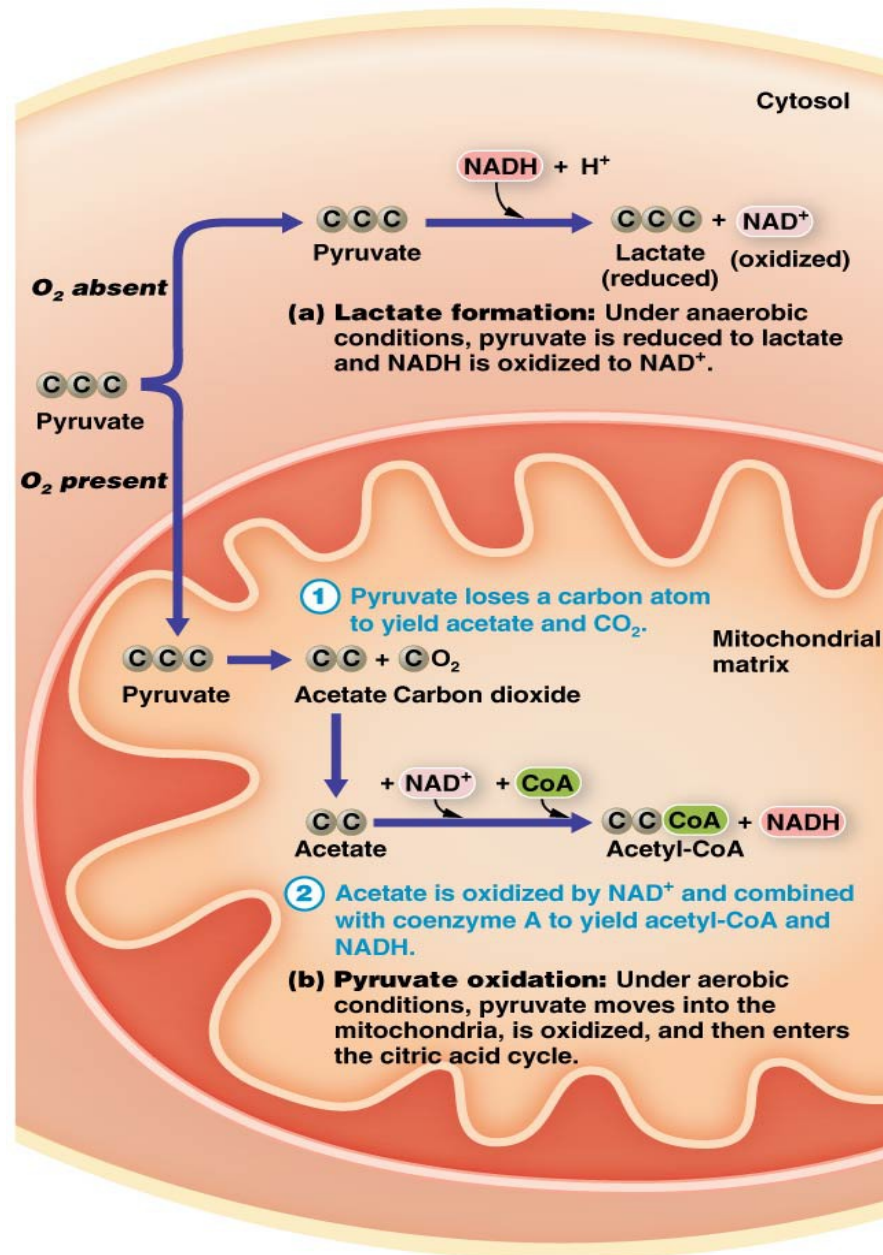
2 ATP

34 ATP

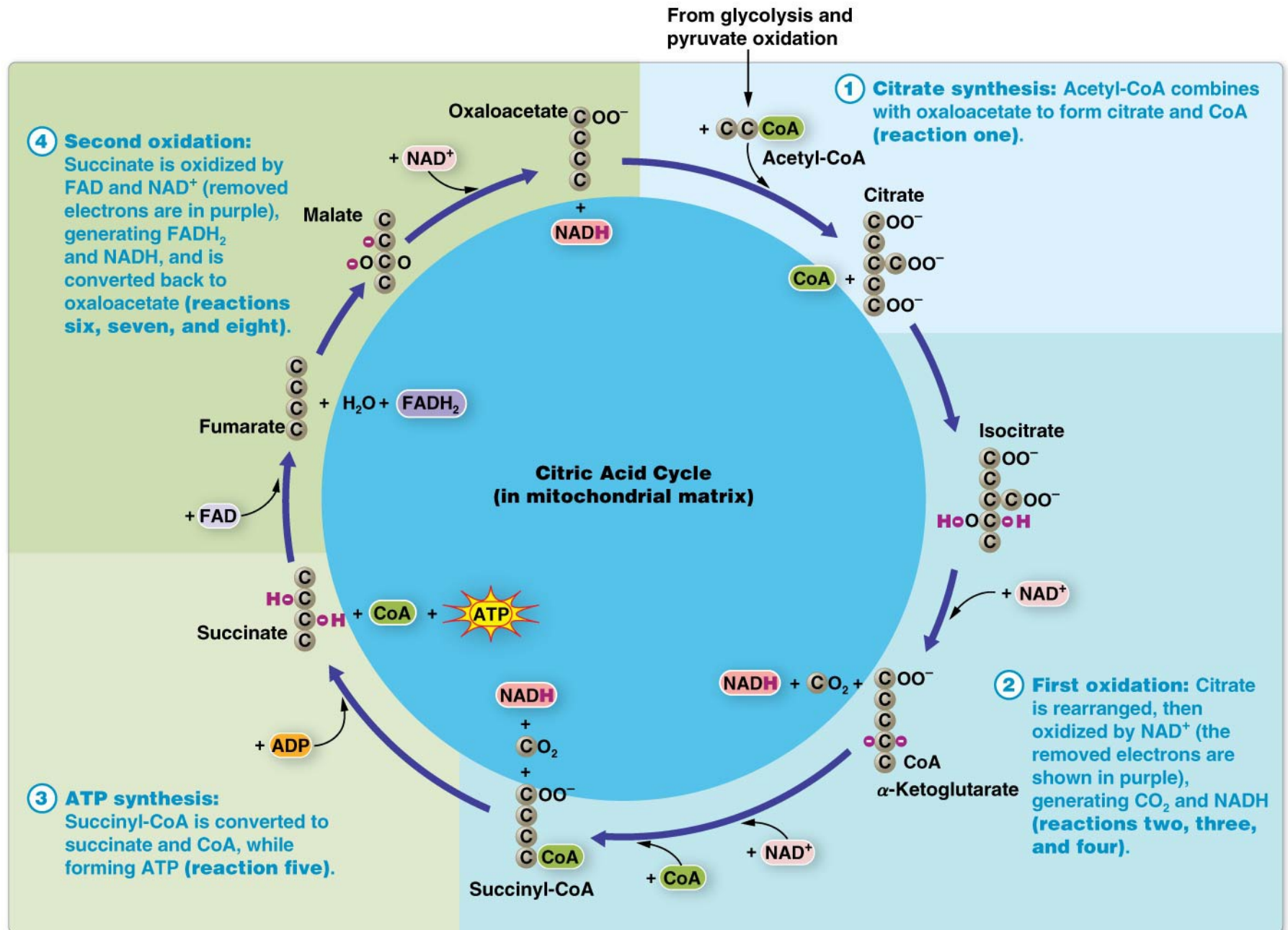
TOTAL = 38 ATP



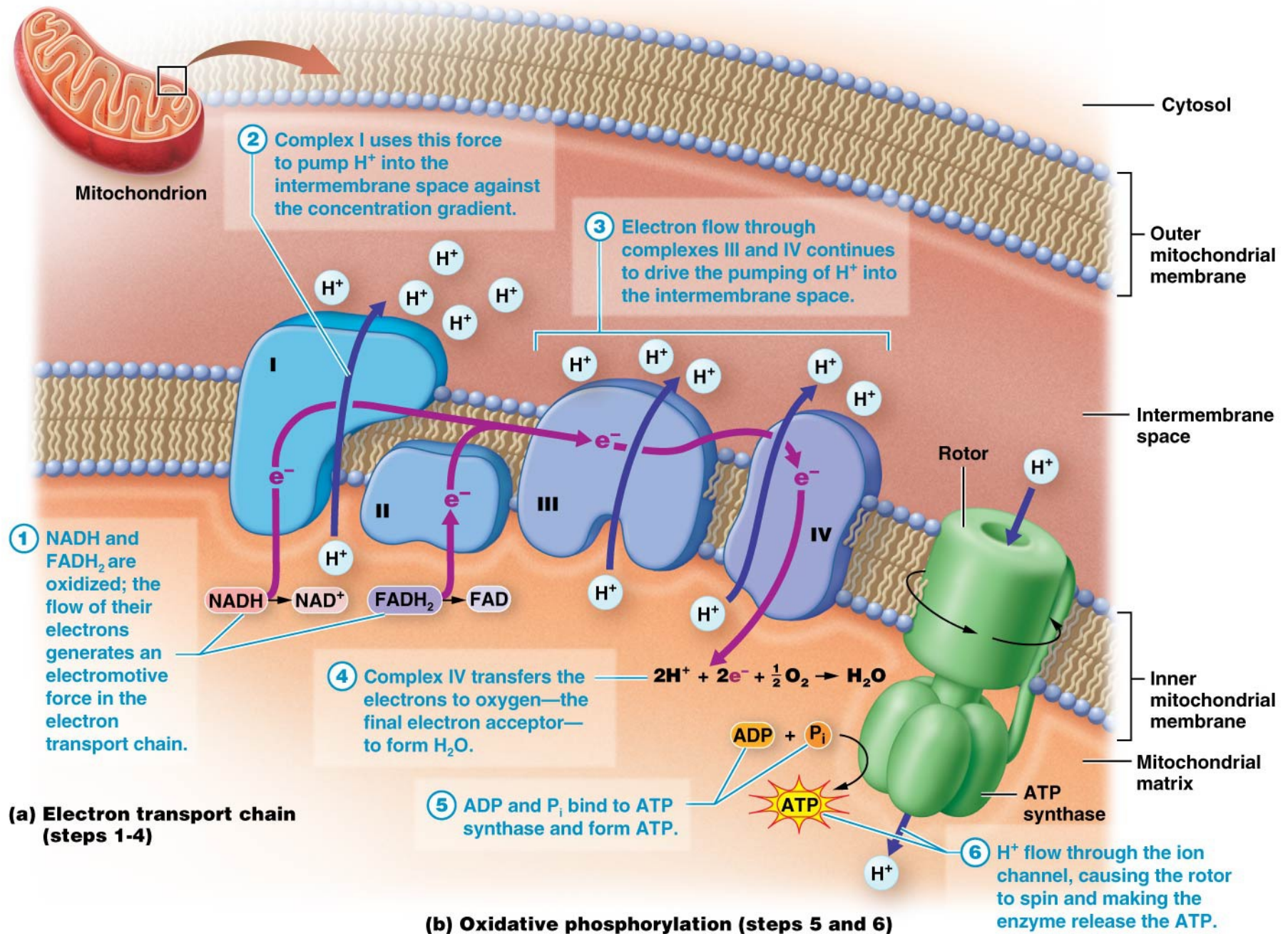
Intermediate step: the fate of pyruvate after glycolysis.



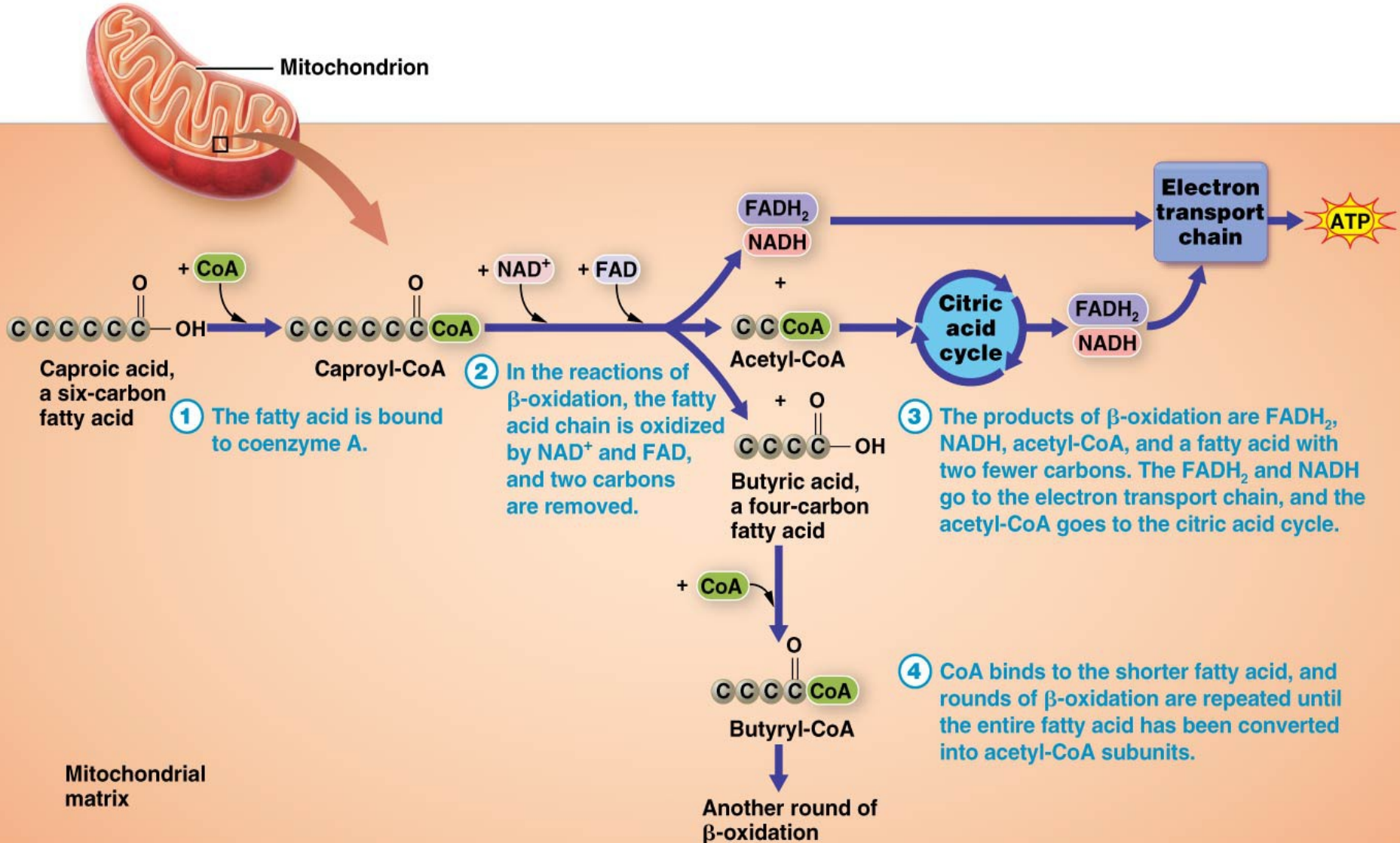
The citric acid cycle.



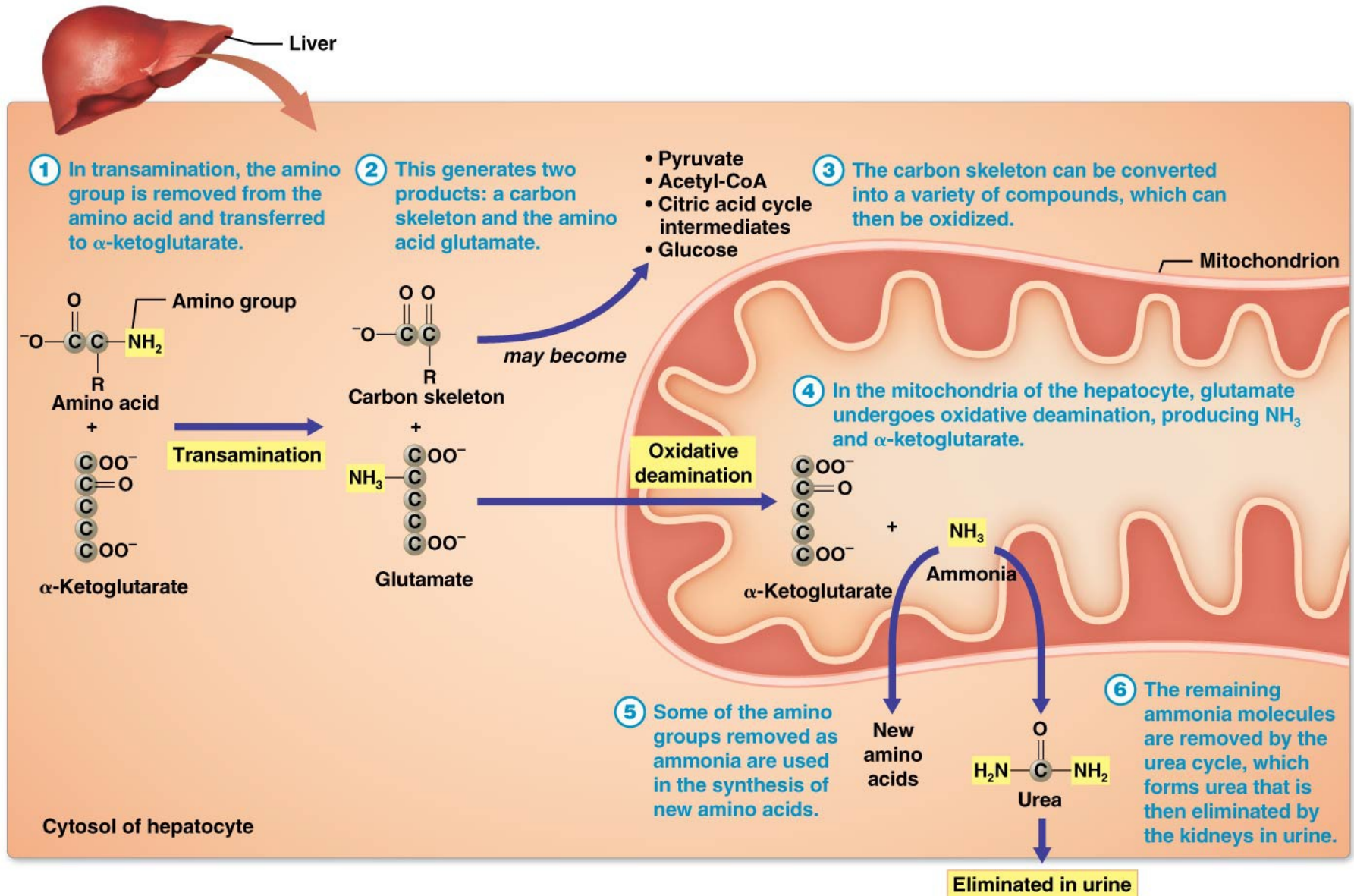
The electron transport chain and oxidative phosphorylation.



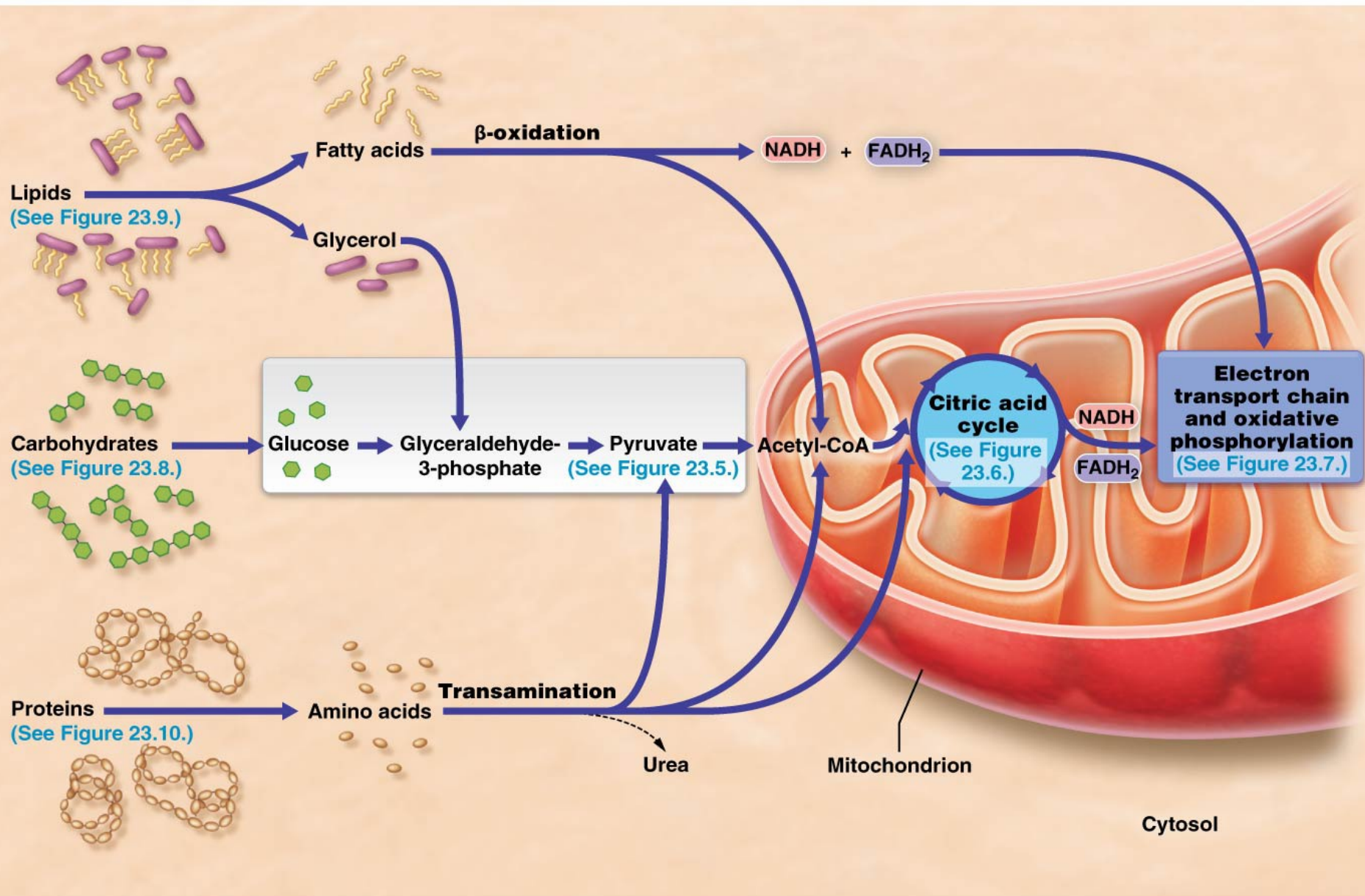
Fatty acid catabolism and β -oxidation.



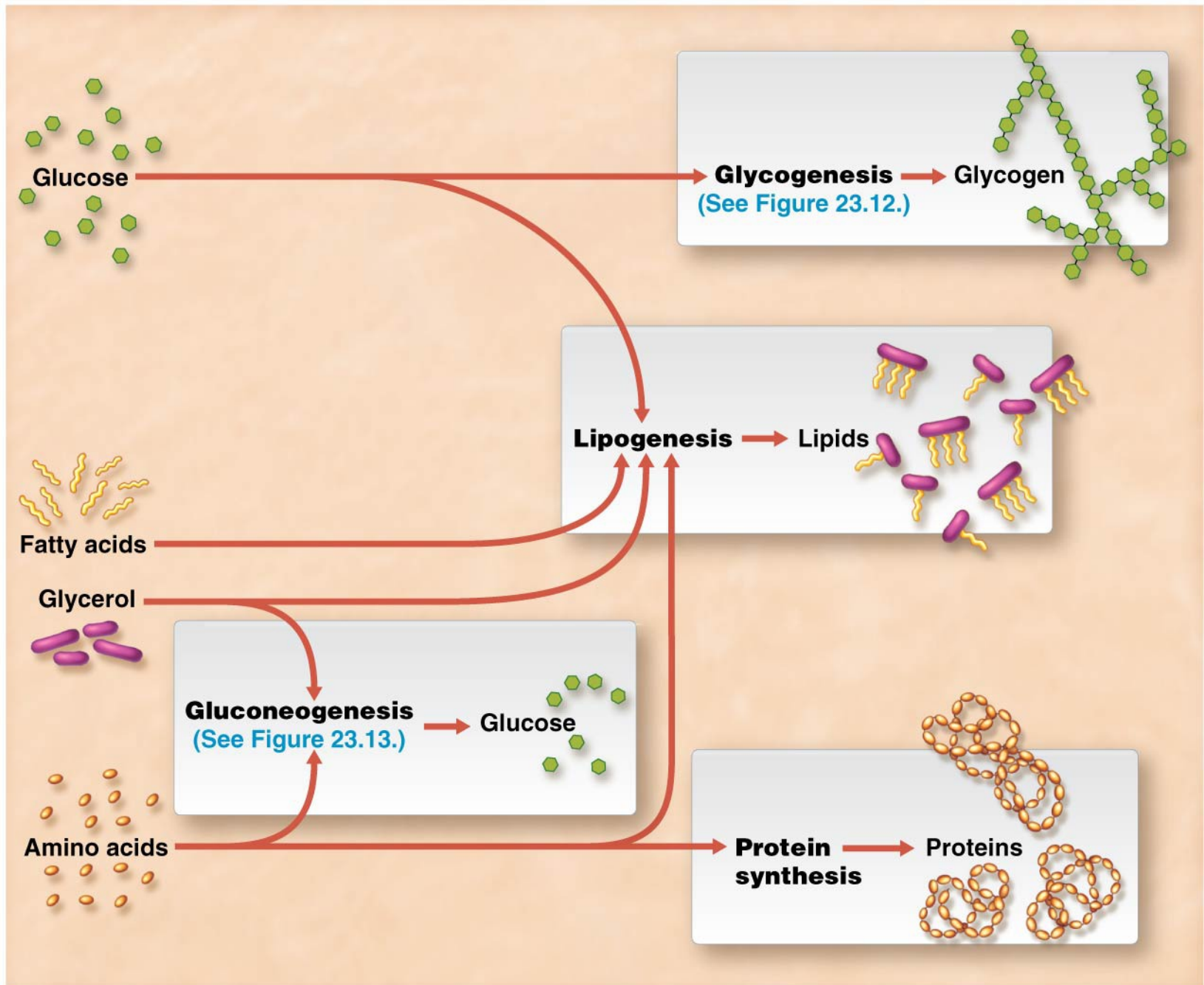
Amino acid catabolism.



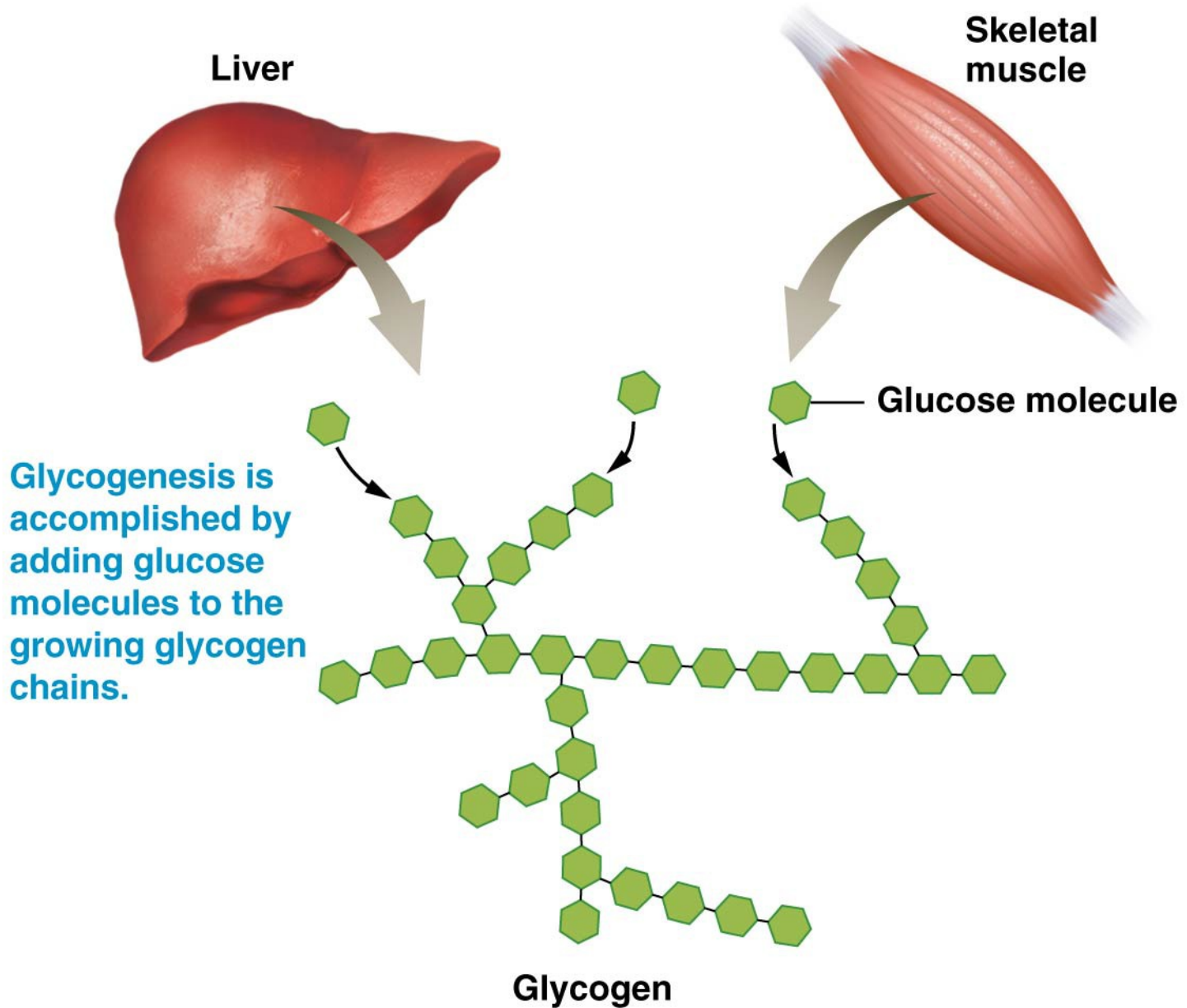
The Big Picture of Nutrient Catabolism



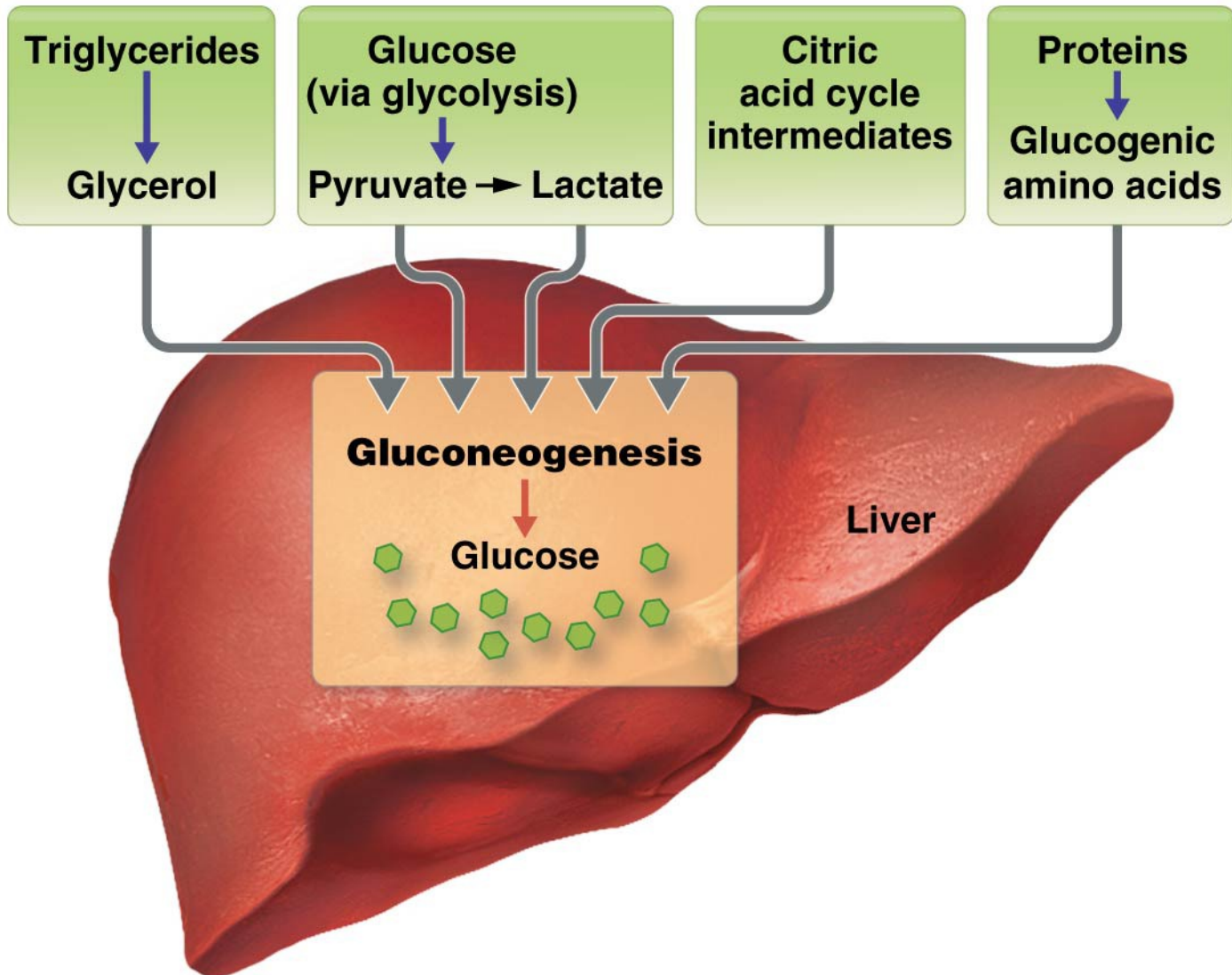
The Big Picture of Nutrient Anabolism



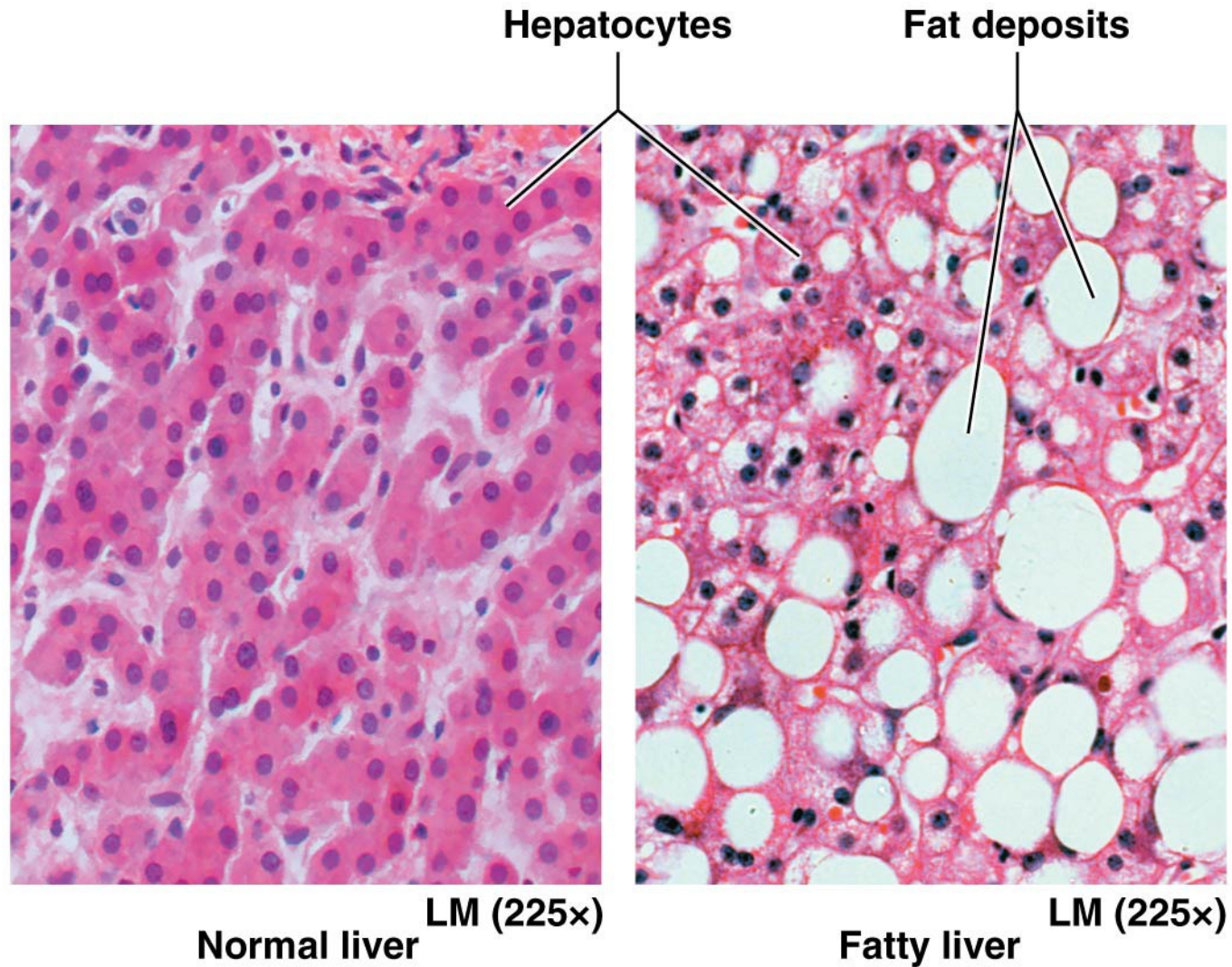
Glycogenesis.



Gluconeogenesis.



Too Much Lipogenesis in Hepatocytes

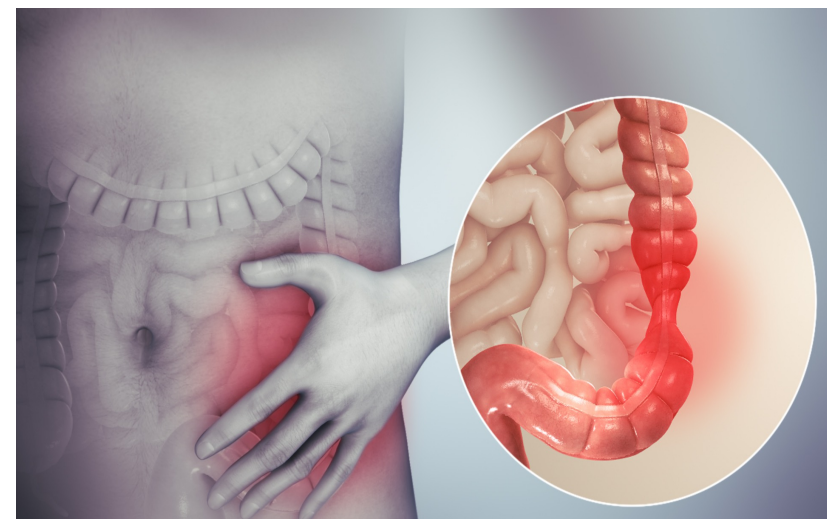


GI Tract Diseases

- > Irritable Bowel Syndrome
- > Crohn Disease
- > Ulceratis Colitis
- > Celiac Disease

Irritable Bowel Syndrome

Irritable bowel syndrome (IBS) is a common disorder that affects the **large intestine**. Signs and symptoms include cramping, abdominal pain, bloating, gas, and diarrhea or constipation, or both. IBS is a chronic condition that you'll need to manage long term.



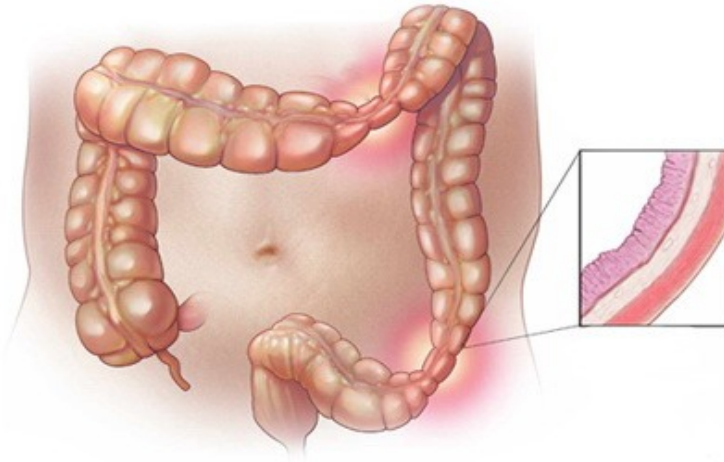
Only a small number of people with IBS have severe signs and symptoms. Some people can control their symptoms by managing diet, lifestyle and stress. More-severe symptoms can be treated with medication and counseling.

IBS doesn't cause changes in bowel tissue or increase your risk of colon-rectal cancer.

Symptoms - The signs and symptoms of IBS vary. The most common include: abdominal pain, cramping or bloating that is typically relieved or partially relieved by passing a bowel movement /// Excess gas /// Diarrhea or constipation — sometimes alternating bouts of diarrhea and constipation // Mucus in the stool

Most people with IBS experience times when the signs and symptoms are worse and times when they improve or even disappear completely.

WHAT ARE THE SYMPTOMS OF IRRITABLE BOWEL SYNDROME?



✓ BACK PAIN

✓ URINARY INCONTINENCE

✓ PAIN DURING SEXUAL INTERCOURSE

✓ STOMACH CRAMPS

✓ ABDOMINAL PAIN

✓ DIARRHEA

✓ CONSTIPATION

✓ BLOATING

✓ FLATULENCE

✓ INCONTINENCE

✓ STOOLS THAT CONTAIN MUCUS

✓ LETHARGY AND DROWSINESS

Crohn's Disease is a chronic, **inflammatory disease** of the **gastrointestinal tract**. It is an autoimmune disorder, meaning your body's immune system mistakenly attacks healthy tissue in your body.

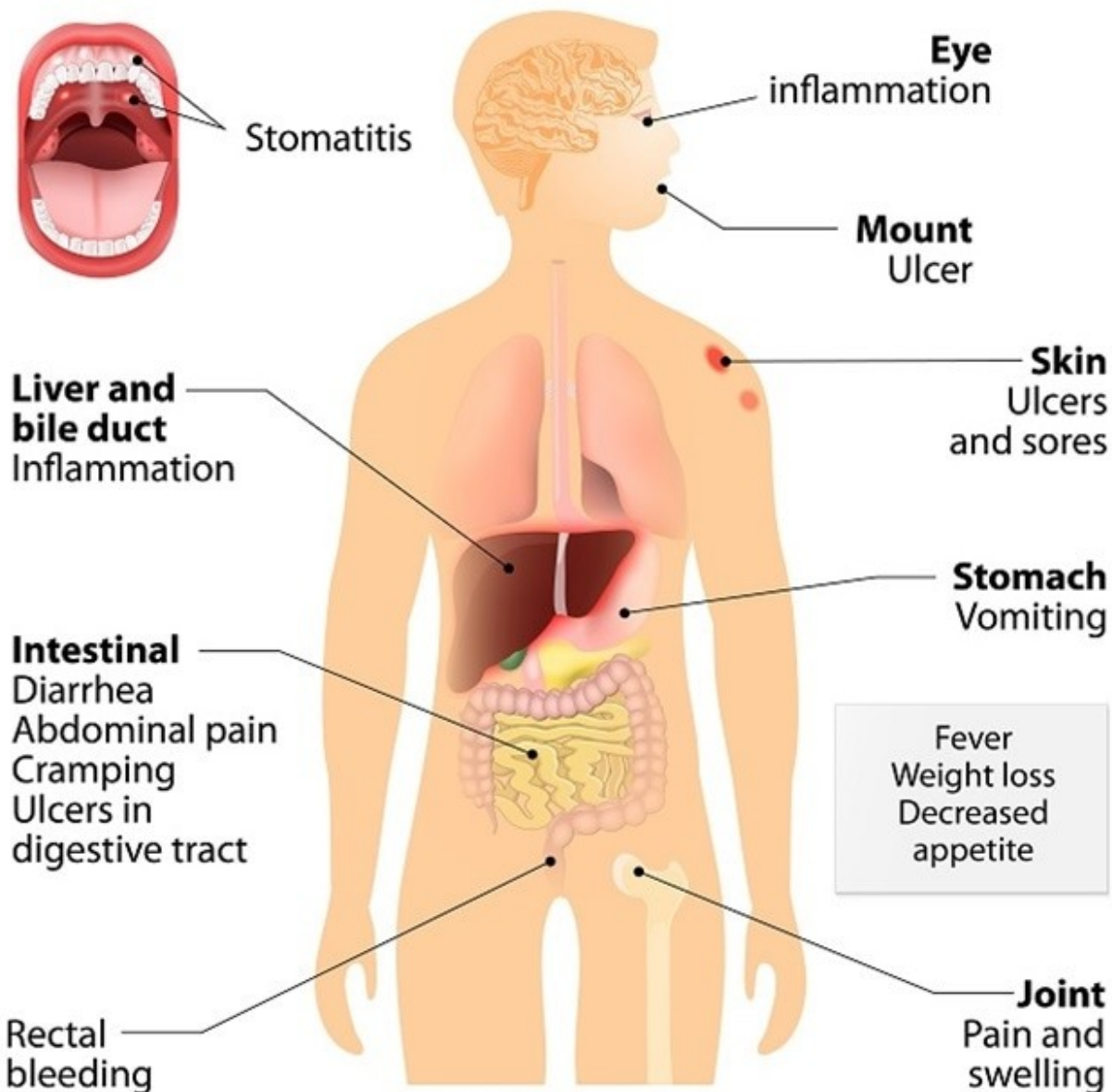
Crohn's disease is chronic (ongoing), and may appear and disappear at various times. Initially, it may affect only a small part of your gastrointestinal tract, but the disease has the potential to progress extensively.

Who Gets Crohn's Disease? When?

Crohn's disease **appears early in life**; approximately one-sixth of patients have symptoms before 15 years of age. Although the cause is unknown, doctors suspect a genetic influence, since many members of the same family may be affected. Crohn's disease affects the Jewish population more than the general population.

Crohn's disease often begins in a person's teens or twenties, though some patients experience symptoms even earlier. Most common symptoms include (see next slide)

Crohn's disease



Crohn disease

GI tract affected
in a patchy pattern

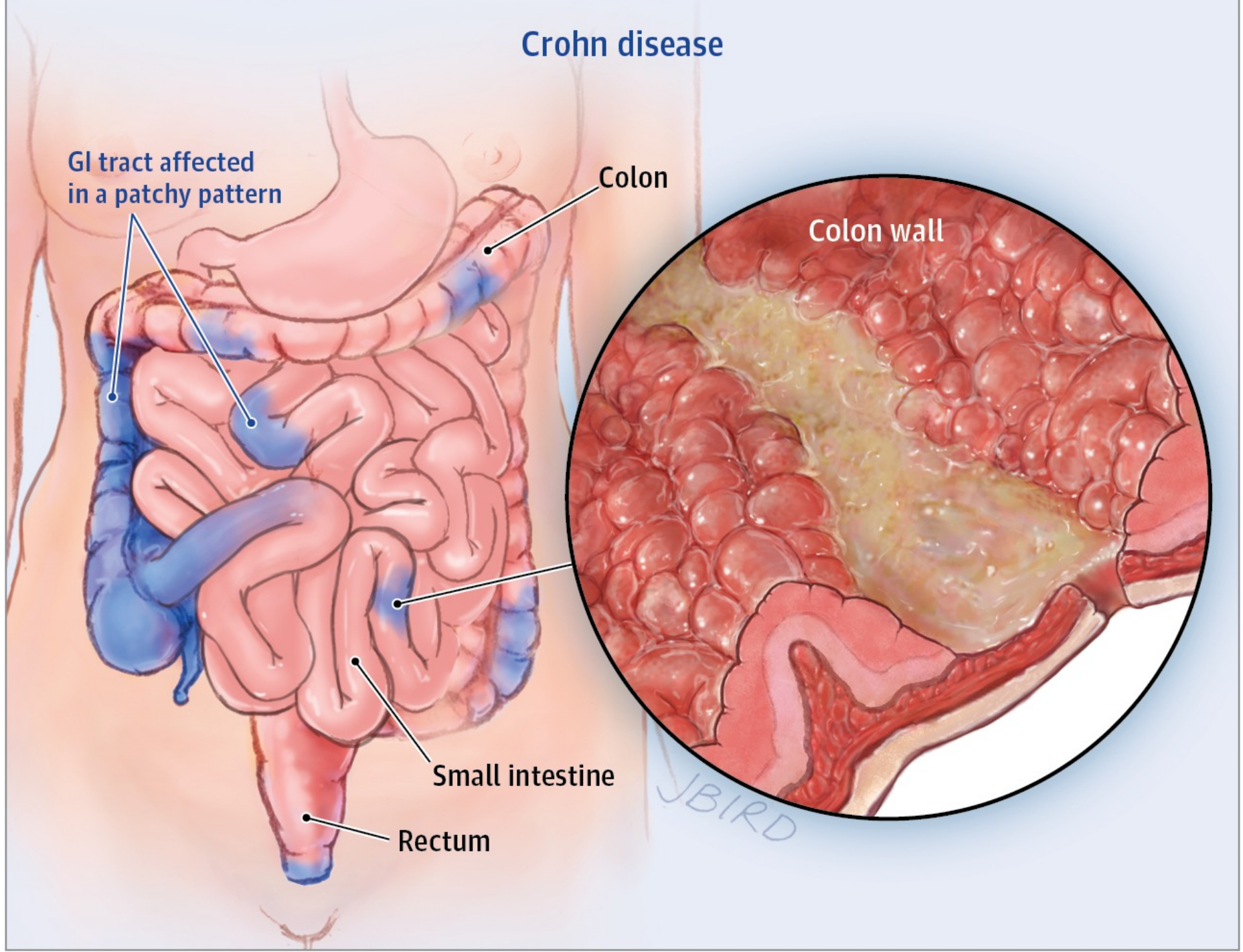
Colon

Colon wall

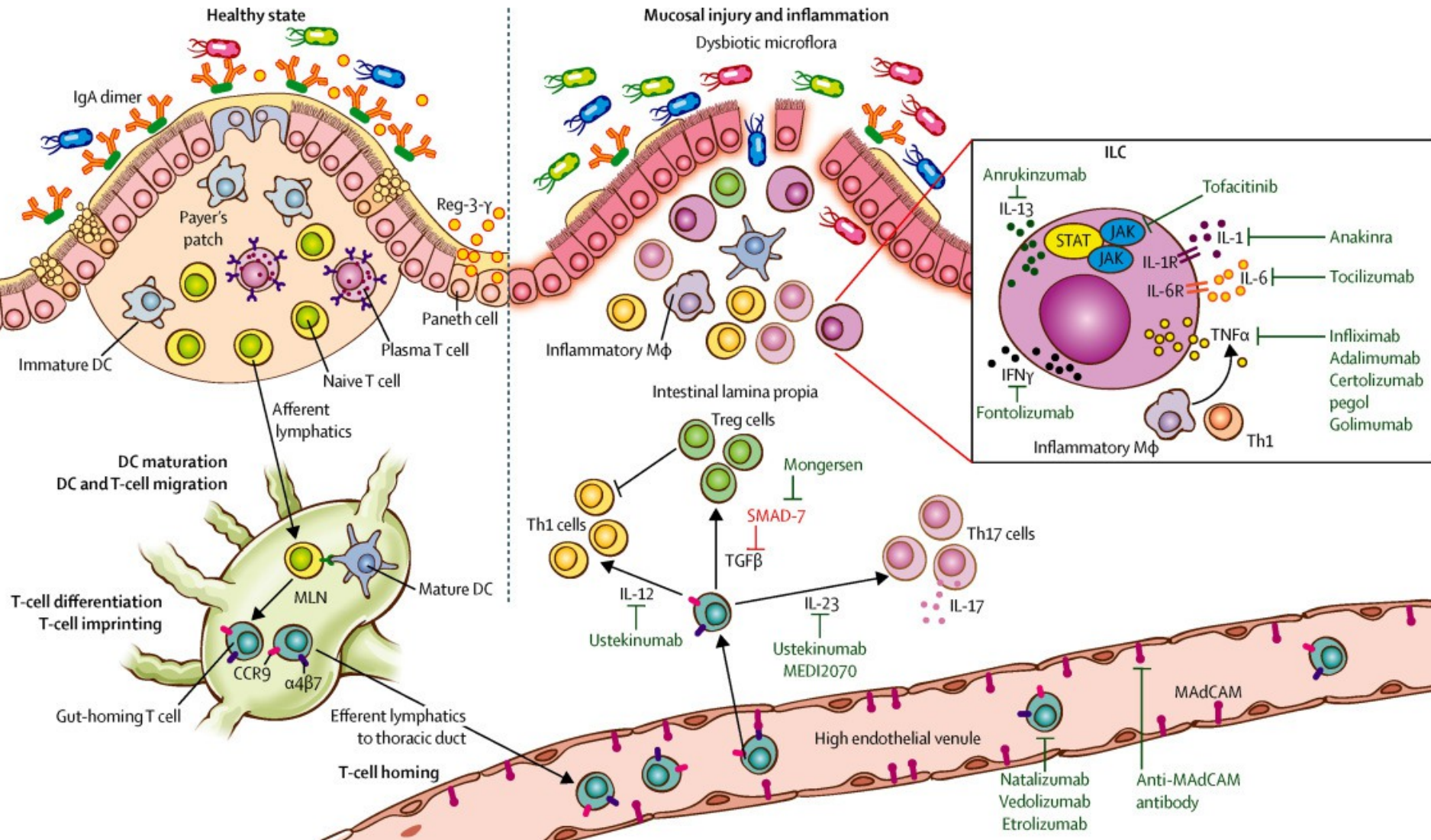
Small intestine

Rectum

JBIRD



Crohn Disease

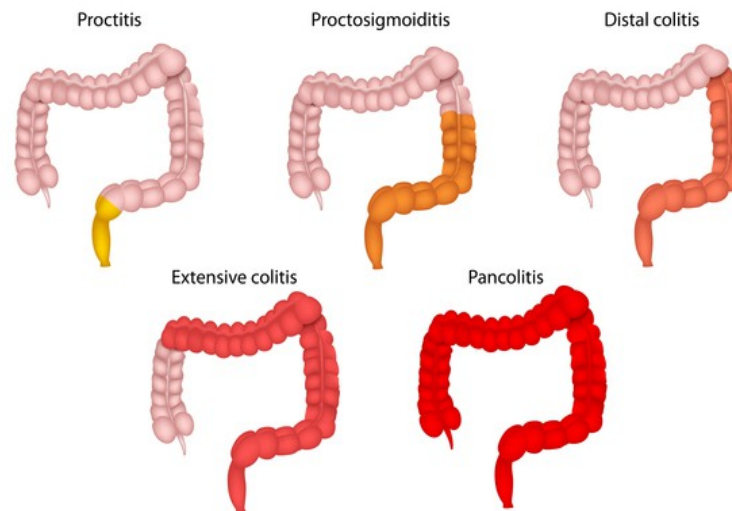


Ulcerative Colitis (UC) is a disease that affects your **large intestine**, or colon. It causes irritation and swelling called inflammation. ... UC is a type of **inflammatory bowel disease**, but it's different from other diseases with similar symptoms, like Crohn's disease or irritable bowel syndrome.

Ulcerative colitis is usually in the innermost lining of the large intestine (colon) and rectum. Forms range from mild to severe. Having ulcerative colitis puts a patient at **increased risk of developing colon cancer**.

Symptoms include rectal bleeding, bloody diarrhea, abdominal cramps, and pain.

TYPES OF ULCERATIVE COLITIS



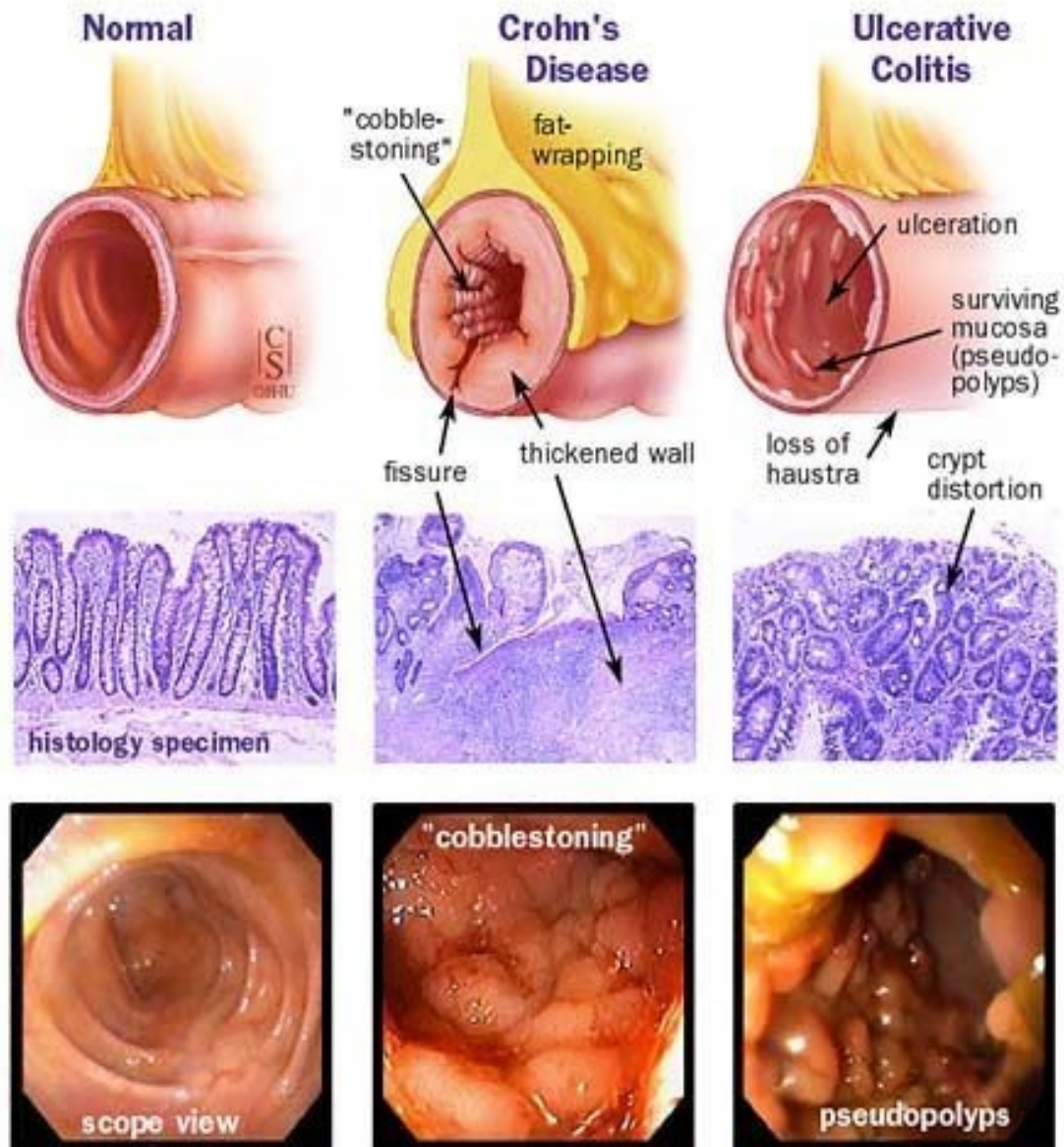
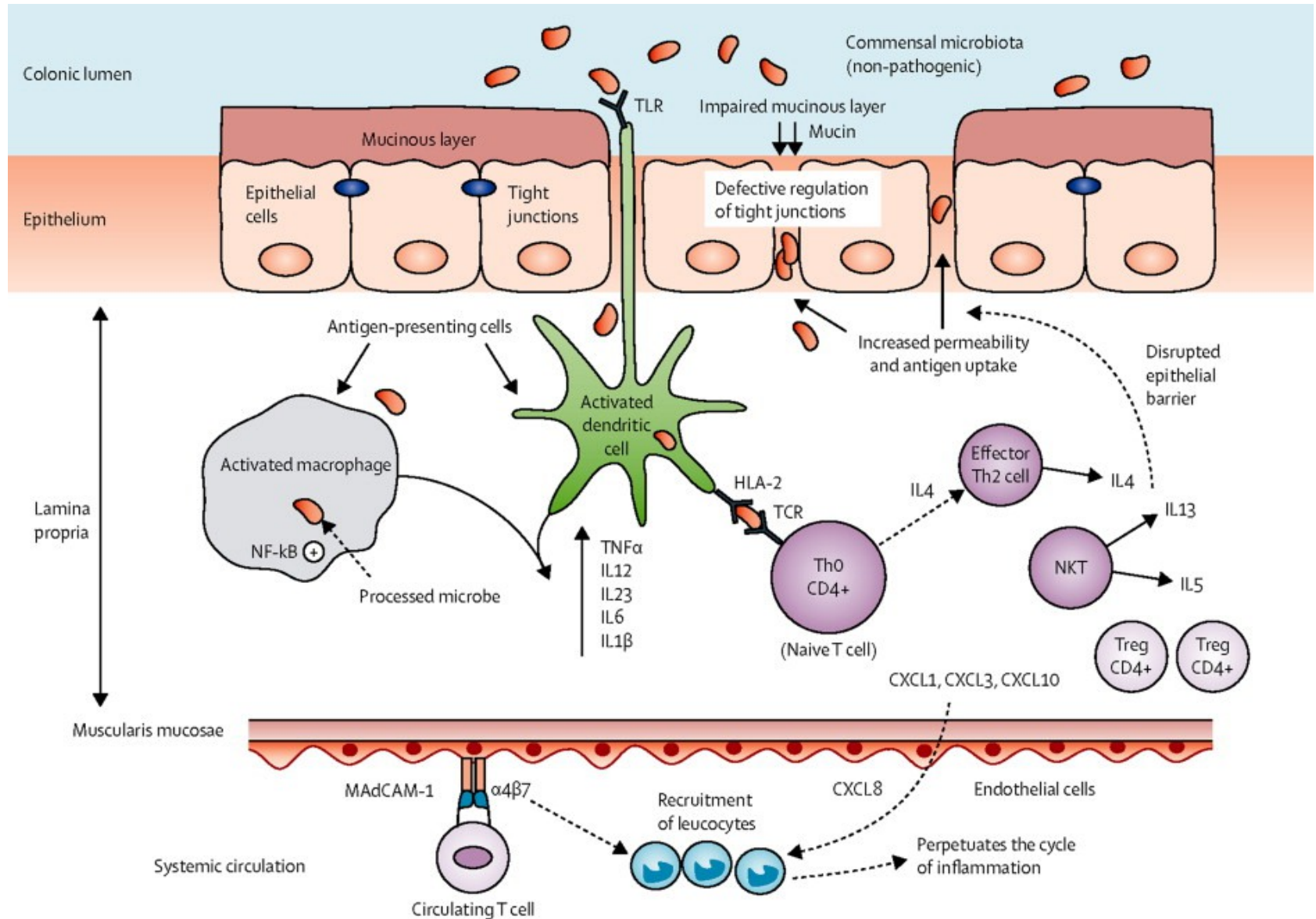


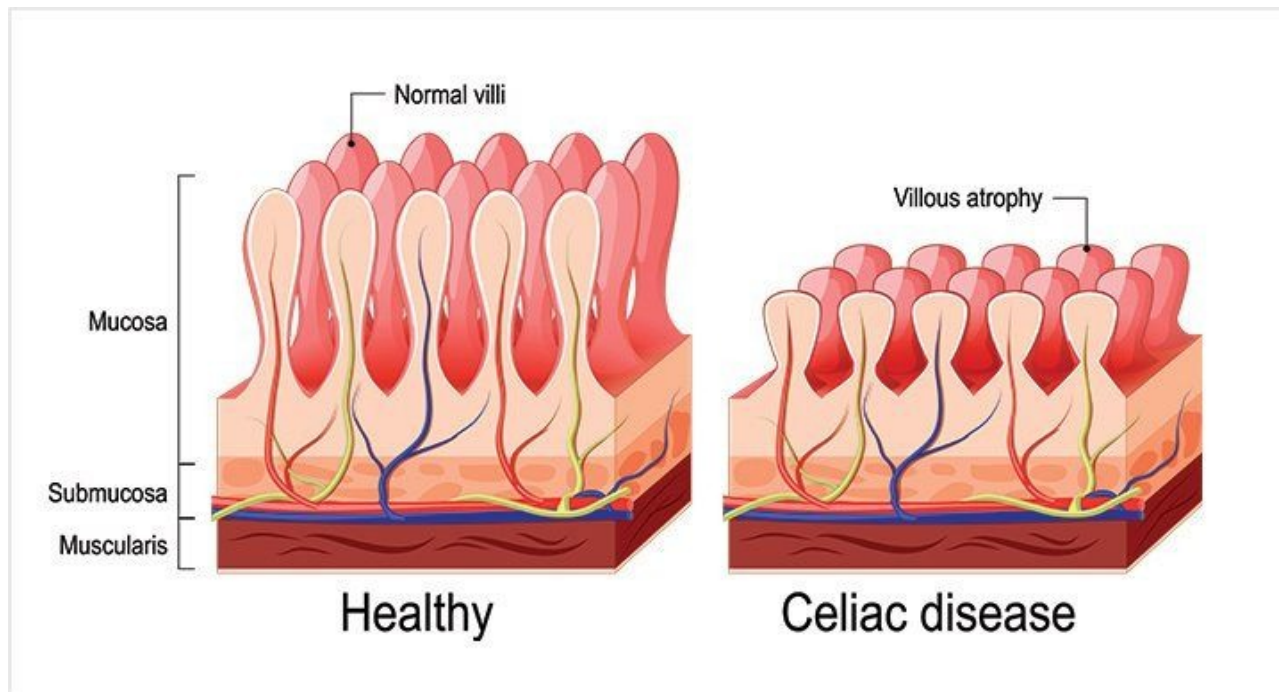
Figure 4. Gross (top), histological (center), and endoscopic (bottom) appearance of normal colon, Crohn's disease, and ulcerative colitis.

Ulcerative Colitis Pathophysiology



Celiac Disease

Celiac disease is caused by a **hypersensitivity** that causes an immune disease condition. The immune response causes damage to their **small intestine**. People with celiac disease are hypersensitive to **gluten**. To control the disease you need to eliminate gluten from their diet. Gluten is a protein found in wheat, rye, and barley. Loss of microvilli inhibit nutrient absorption.



SYMPTOMS OF CELIAC DISEASE



UNEXPLAINED WEIGHT LOSS



STOMACH & MUSCLE CRAMPS



NUMBNESS & TINGLING IN LEGS



TOOTH ENAMEL LOSS



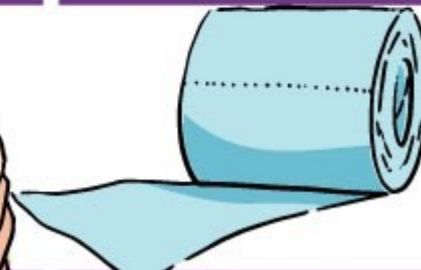
JOINT PAIN



WEAK BONES



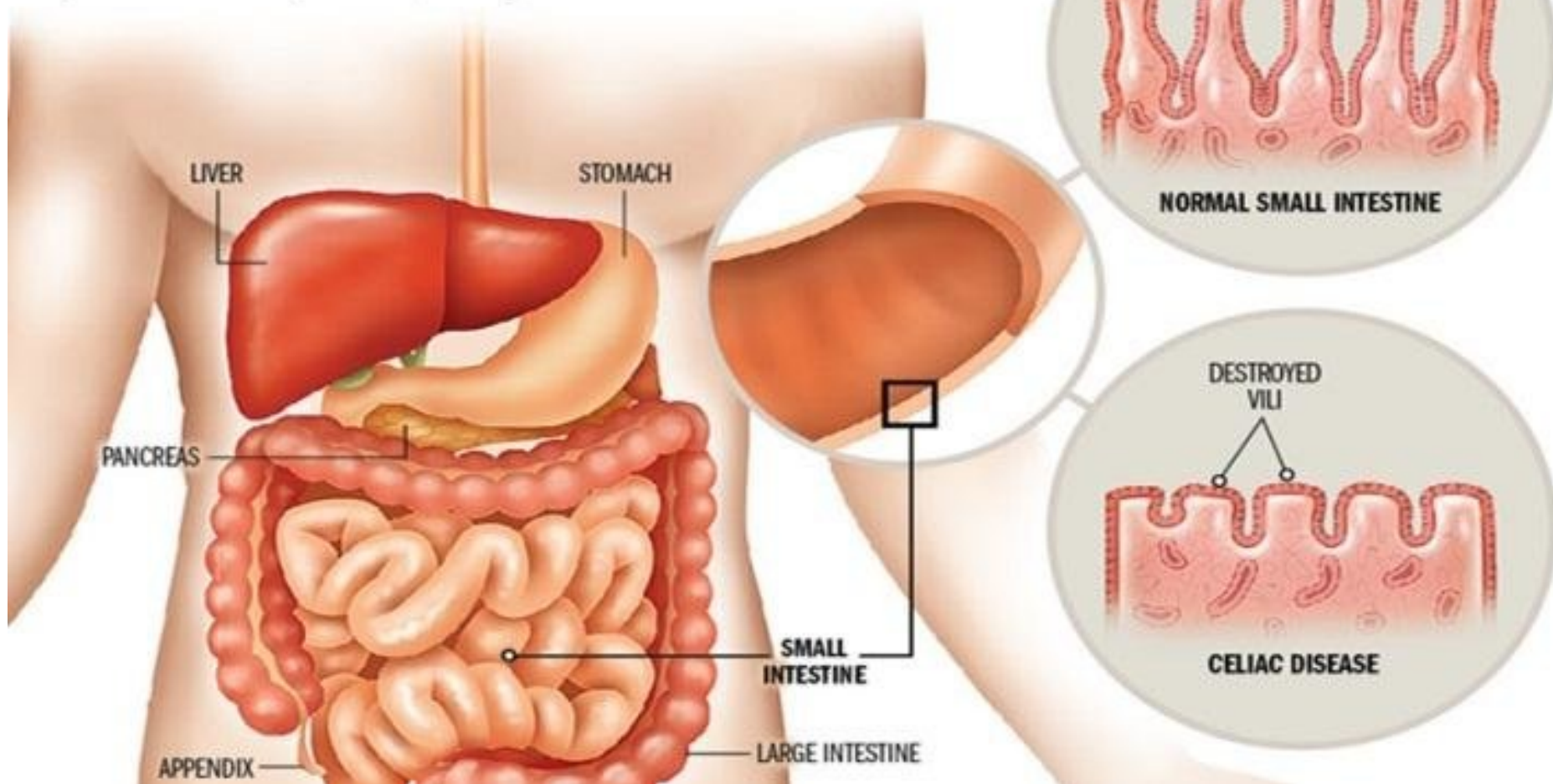
INFERTILITY

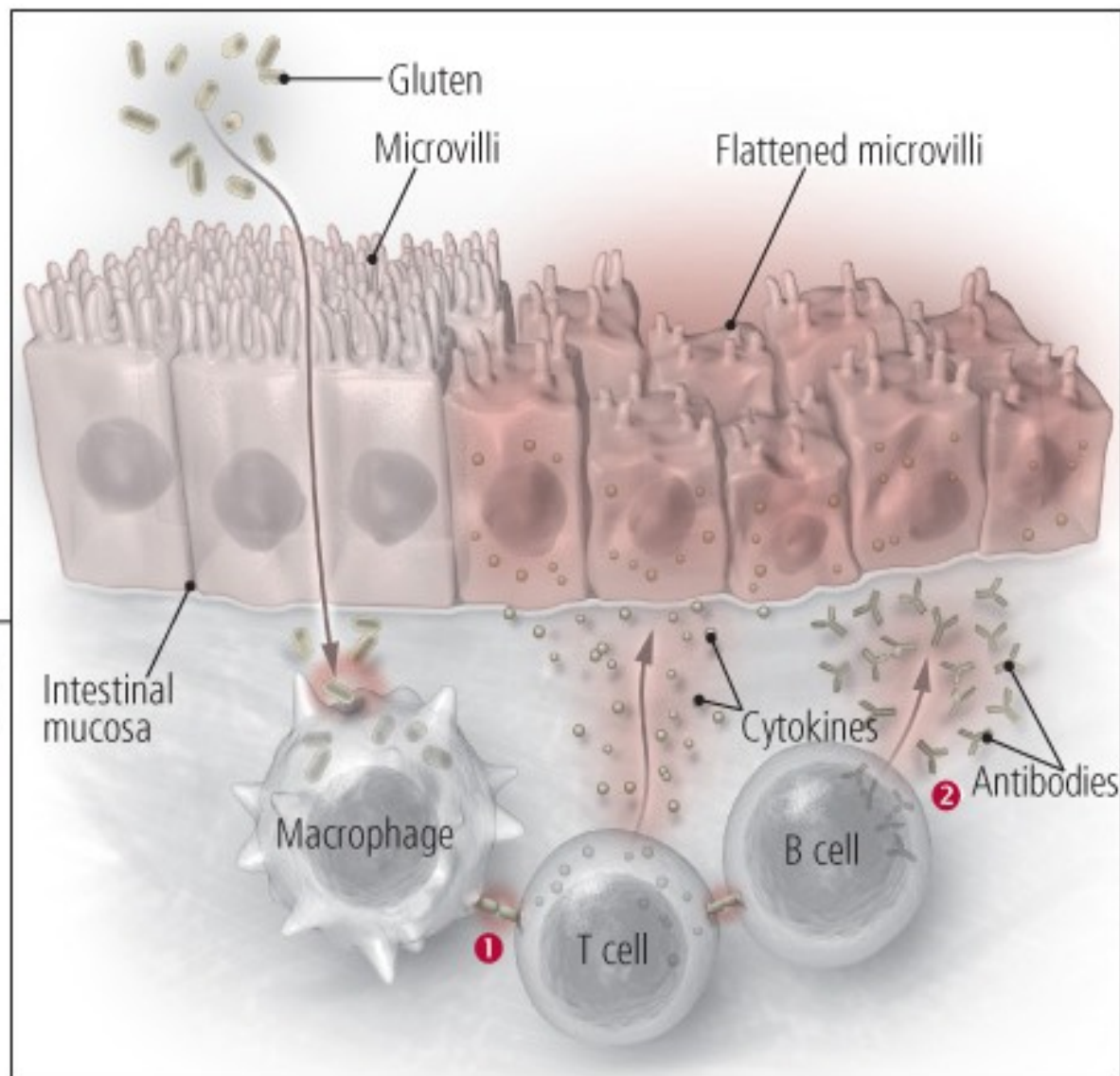
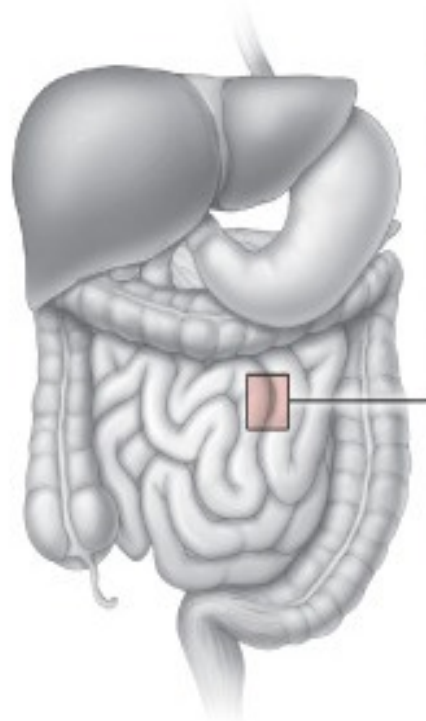


CHRONIC DIARRHEA
OR CONSTIPATION

Damage from celiac disease

In a healthy small intestine, tiny hairlike projections called villi absorb nutrients from food. When people with celiac disease eat foods containing wheat, barley, or rye, the body's immune system attacks the gluten proteins. This immune response also destroys the villi, leading to nutritional deficiencies.





Stage 1



Stage 2



Stage 3



Stage 4



A COMPLICATED CONVERSATION

The interplay between the digestive tract and the central nervous system is well-established. Signals produced by the gut microbiota are thought to influence this gut-brain axis. If they do, disruptions to this community of bacteria could have a role in exacerbating symptoms of irritable bowel syndrome.



The microbiota is thought to influence mood and cognition, possibly by signalling to the brain through the vagus nerve or stimulating production of neurotransmitters in the gut. But this has been shown only in animal studies.

The brain regulates digestive-tract activity.

The gut communicates with the brain through the vagus nerve.

Effects of environmental stress on the brain can have a lasting impact on gut health.

Disrupted gut function can change the microbiota's composition or metabolic activity.

Intestinal wall

Microbes

Signals from the microbiota maintain the intestinal wall.



Pathogenic bacteria or overexposure to antibiotics may kill off microbes that assist gut function.



Supplementation with 'good' bacteria, or probiotics, may restore the microbiota.

See home page link
Openbiome.

The Big Picture of Digestion.

