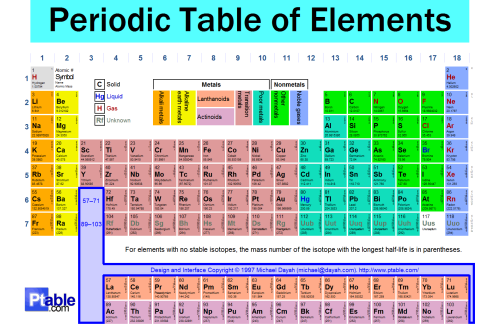
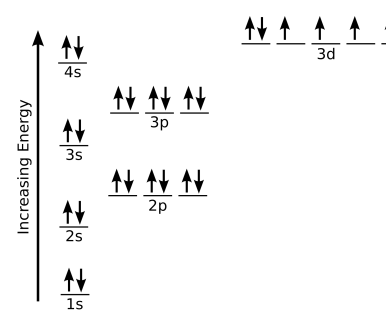
# 5. IRON METABOLISM

Reviews on Moodle

#### **IRON**

- 26th element in the periodic table
  - Chemical Symbol: Fe
  - MW = 55.85
  - Electron Configuration: 1s<sup>2</sup> 2s<sup>2</sup>2p<sup>6</sup> 3s<sup>2</sup>3p<sup>6</sup>4s<sup>2</sup>3d<sup>6</sup>
  - Fourth most abundant mineral O > Si > Al
  - Oxidation states = -2 to +6
- Readily interconverted, i.e. redox active
  - very useful for redox chemistry in the body
  - very dangerous chemistry





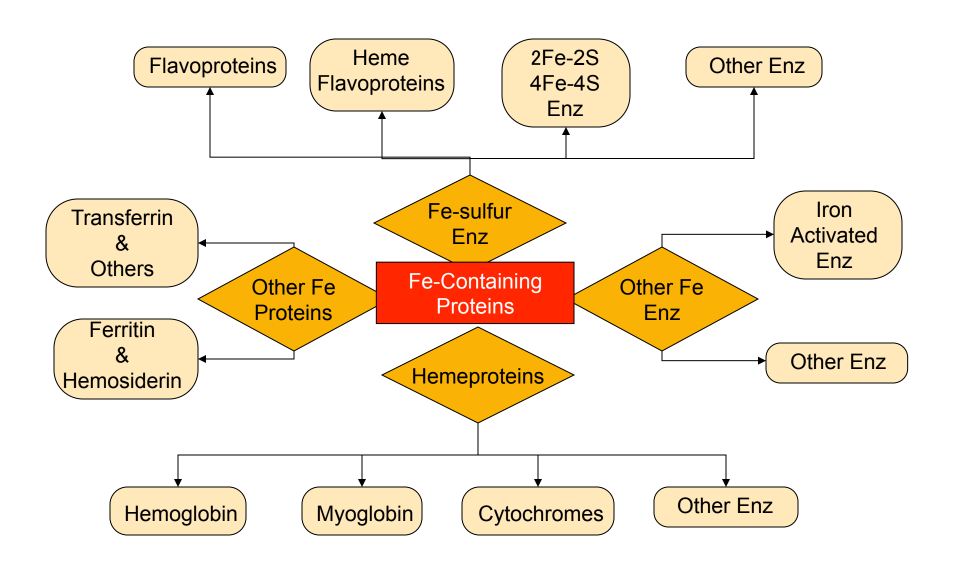
# IRON FUNCTIONS IN BIOLOGY

- Oxygen Transport and Storage
  - Hemoglobin
  - Myoglobin
- Electron Transport and Energy Metabolism
  - Cytochromes
  - Fe-S proteins
- Substrate Oxidation & Reduction

#### Iron dependent enzyme:

- Ribonucleotide reductase
- Amino acid oxidases
- Fatty acid desaturases
- Nitric oxide synthetase
- Peroxidases
- Regulation of intracellular iron

# **IRON IN BIOLOGY**



### IRON FORMS IN DIET

#### Heme

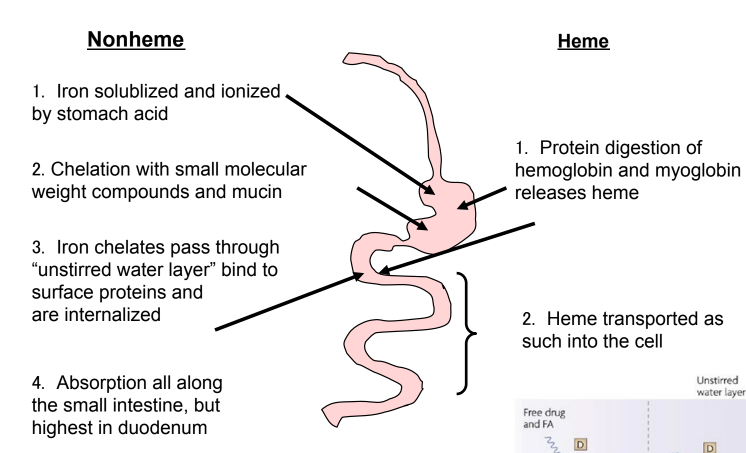
- Iron-porphyrin prosthetic group
- Hemoglobin, myoglobin, cytochromes
- Other iron-containing enzymes are
   ~3% body iron
  - Aconitase, peroxidases
- 5-35% is absorbed
- Exist as Fe<sup>2+</sup>

$$H_2C = CH$$
 $H_3C$ 
 $CH_2$ 
 $CH_2$ 
 $CH_3$ 
 $CH_3$ 
 $CH_4$ 
 $CH_5$ 
 $CH_5$ 
 $CH_7$ 
 $CH_8$ 
 $CH_8$ 
 $CH_9$ 
 $CH$ 

#### Non-heme

- >85% of iron in foods is non heme iron
- 2-20% is absorbed
- Exist as Fe<sup>3+</sup>

### **ABSORPTION**



The fluid in the unstirred water layer (UWL) is not stationary, but is rather a region of slow laminar flow parallel to the membrane in which the only mechanism of transport is by diffusion; the layers are often called "Nernst diffusion layers."

Enterocyte

Efflux

Passive diffusion Carriermediated

uptake

Absorption by collisional transfer

by vesicular

uptake

Binding followed

Apical

H\*

DOD

Micellar

drug and FA DOD

membrane

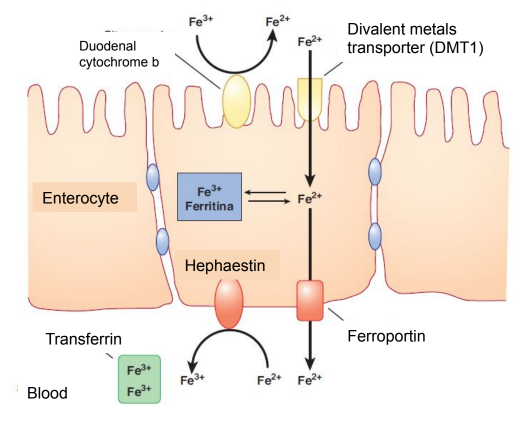
# **ABSORPTION**

- Ferrous, Fe<sup>2+</sup>, most soluble = most absorbable
- Each mechanism has 3 phases
  - Iron uptake
  - Intraenterocyte transport
  - Storage and extra-enterocyte transfer

#### Dietary Iron:

- Iron is essential element and must be precisely regulated.
- On the lumen side of small intestine iron is reduced from its ferric form (Fe<sup>3+</sup>) to ferrous form (Fe<sup>2+</sup>).
- Ferrous iron is then transported in enterocytes by DMT1 (divalent metal transporter).

#### Intestinal lumen



# **ABSORPTION**

- Heme iron is an important dietary sources of iron because it is more effectively absorbed than nonheme iron.
- From 5% to 35% of heme iron is absorbed from a single meal, whereas non-heme iron absorption from a single meal can range 2%-20%, depending on the iron status of the individual and the ratio of enhancers and promotors in the diet. Thus, although it constitutes about 10% of the iron found in the diet, heme iron may provide up to one-third of total absorbed dietary iron.
- The reason ascorbic acid promote iron absorption is ascorbic acid maintain iron in a reduced form and forms a soluble chelate with iron. These actions are shared by organic acids such as citric, lactic acids.
- A number of dietary factors influence iron absorption.

$$Fe^{3+} + e^{-}$$
 Fe<sup>2+</sup> Highly insoluble Soluble

#### Promotors

- Amino Acids
- Animal Proteins(for heme)
- Ascorbic Acid
- Hydrochloric Acid
- Organic Acids
- Sugars
- Mucin

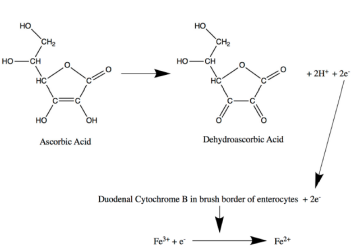


Figure 3. Low stomach pH and dietary ascorbic acid reduces non-haem iron from the highly insoluble Fe<sup>3+</sup> form to Fe<sup>2+</sup>, which is more readily absorbed. Duodenal cytochrome b accepts electrons intracellularly from oxidation of ascorbic acid into dehydroascorbic acid and uses these to catalyse the reduction of Fe<sup>3+</sup> to Fe<sup>2+</sup>.

#### Inhibitors

- Carbonates
- Calcium (for heme)
- Egg yolk phosvitin
- Fiber
- Oxalates
- Phosphates
- Phytates
- Plant polyphenols
- Soy proteins

#### **TRANSPORT**

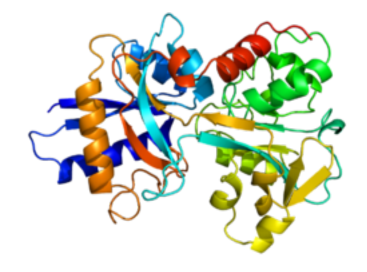
Free iron is toxic because it catalyses the Fenton reaction generating free oxygen radicals.

$$Fe^{2^{+}} + H_{2}O_{2}$$
  $\longrightarrow$   $Fe^{3^{+}} + OH^{*} + OH^{-}$ 

- In biological systems iron is always bound to proteins to limit its toxicity.
- In plasma, it is bound to transferrin that plays a key role in iron transport to sites where it is required.

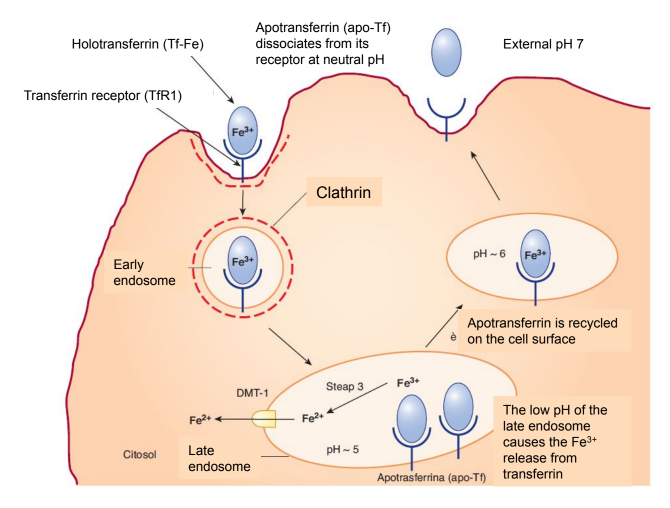
#### **Transferrin**: Transport iron

- Single polypeptides composed of two iron binding half-site motifs, ~679 aas, ~76 kDa MW;
- It is a glycoprotein synthesised in the liver;
- Bind <u>2 Fe<sup>3+</sup></u> and 2 HCO<sub>3</sub>
- Normally 25-50% saturated with iron
- Lactoferrin is an iron binding protein in milk, plasma and mucus secretion such as tears
- Ovotransferrin is an iron binding protein in bird's egg white



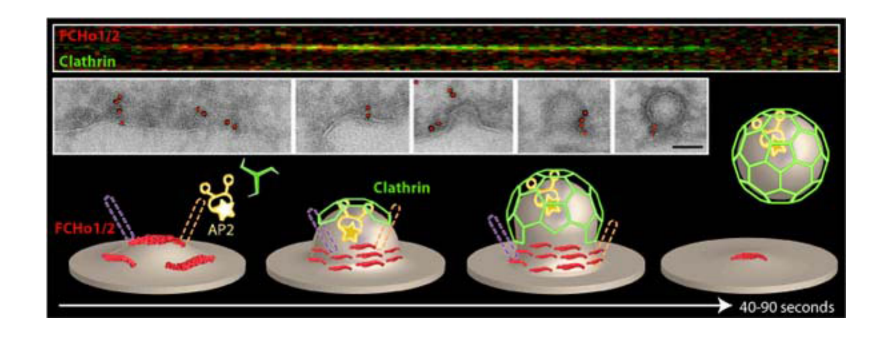
- Transferrin glycosilation is compromised in some disorders and in chronic alcholism, where carbohydrate-deficient transferrin, CDT, is decreased and can be monitored by isoelectric focusing (IEF).
- CDT is a marker for chronic alcholism

### IRON TRANSPORT: TRANSFERRIN CYCLE



Transferrin cycle: holotransferrin (Tf-Fe) binds to transferrin receptor 1 (TfR1) on cell surface. Clathrin-coated vescicles form and endocitosis occurs forming endosomes where pH is acidic. The acidic pH causes iron release from transferrin. Apotransferrin (Apo-Tf) still binds to TfR1. Ferric iron is converted to ferrous iron by a iron reductase (Step 3). Ferrous iron is transported in the cytosol trhough DMT-1. The TfR1-apo-Tf complex is recycled on cell surface where apo-Tf is released and TfR1 can bind another Tf-Fe.

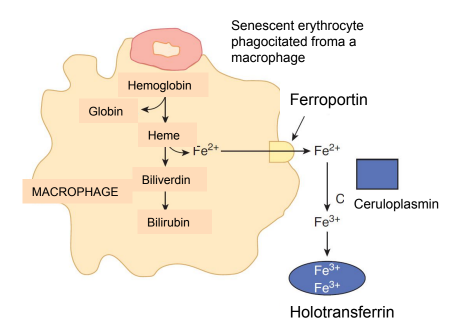
# IRON TRANSPORT: TRANSFERRIN CYCLE



- Clathrin constitutes the coat of vesicles involved in three receptor-mediated intracellular transport pathways: i) the export of aggregated material from the trans-Golgi network for regulated secretion, ii) the transfer of lysosomal hydrolases from the trans-Golgi network to lysosomes and iii) receptor-mediated endocytosis at the plasma membrane.
- The clathrin subunits and the other major coat constituents, the adaptor polypeptides, interact in specific ways to build the characteristic polygonal clathrin lattice and to attach the coat to integral membrane receptors.

#### IRON TRANSPORT: TRANSFERRIN CYCLE

- TfR1 is present in almost all cells, especially in erythrocytes precursors
- There is a TfR2 (TfR2) that is mainly expressed on hepatocyte cells surface but it has a low affinity for Tf-Fe and it does not seem to be involved in iron uptake.
- It is one of the sensors for body iron stored levels
- Erythrocites half-life is around 60 days and they are engulfed by macrophages, where heme is degraded and iron is recycled and it is the highest iron source for the body (25 mg per day) whereas diet iron contributes with only 1-2 mg



#### IRON TRANSPORT: SUMMARY

 Erythrocites half-life is around 60 days and they are engulfed by macrophages, where heme is degraded and iron is recycled and it is the highest iron source for the body (25 mg per day) whereas diet iron contributes with only 1-2 mg

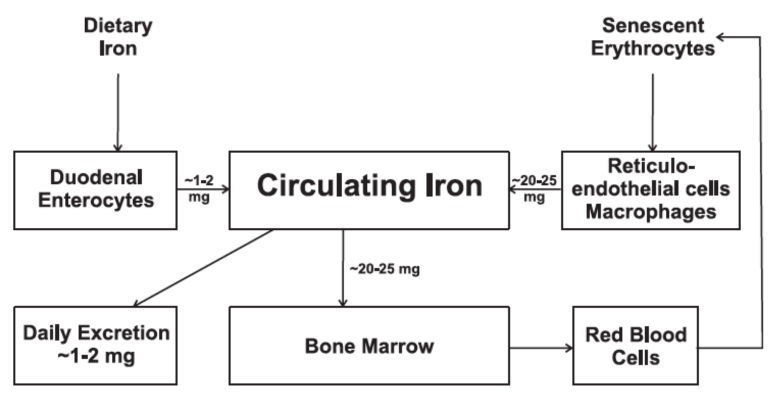


Fig. 7. Diagram showing the physiological turnover of iron in the body

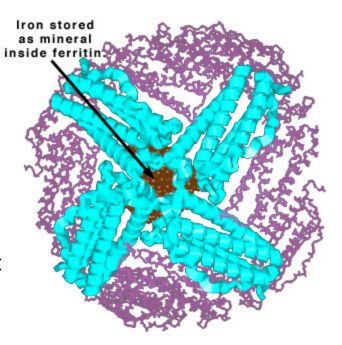
### STORAGE IRON

#### **Ferritin**

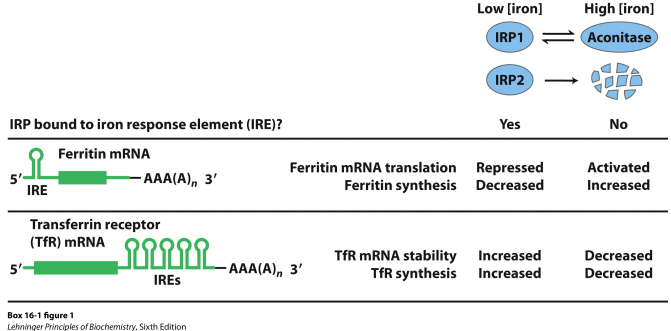
- Major iron storage protein
- Apoferritin 24 polypeptide units in raspberry-like cluster
- Surrounds spherical cluster of <u>hydrated</u> ferric phosphate within its hollow center
- Can contain up to 4500 Fe atoms
- Liver contains ~60% of ferritin in the body
- Two types of subunits:
  - H subunit: 22 kDa, 182 aa, predominant in heart
  - L subunit: 20 kDa, 174 aa, predominant in liver

#### Hemosiderin

- ~50% liver iron stores
- Reacts to ferritin antibodies likely a degradation product
- Less available for mobilization



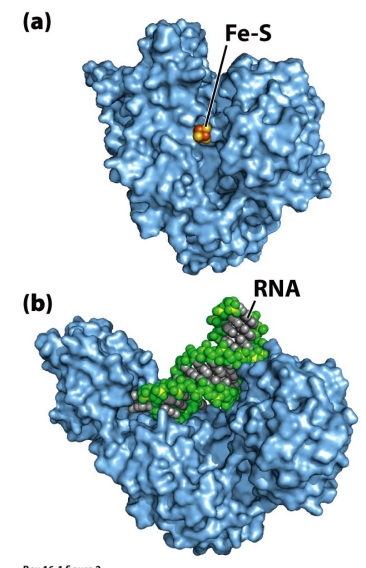
#### REGULATION OF INTRACELLULAR HOMEOSTASIS



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- Synthesis of TfR1 and ferritin are linked to the intracellular iron content
- When iron levels are high, ferritin is synthesised for iron storage and TfR1 synthesis is inhibited
- When iron levels are low, ferritin synthesis is blocked whereas TfR1 is active
- Regulation of mRNA stability is involved
- mRNA for ferritin and TfR1 contain iron response elements (IREs) forming hairpins in the untranslated regions at the 5' and 3', respectively
- IREs are linked to iron regulatory proteins (IRPs) that are sensitive to intracellular iron levels and induced by low levels of the metal
- IRPs bind to IREs when iron levels are low
- IRPs binding to 5' UTR mRNA blocks ferritin translation
- IRPs binding to 3' UTR mRNA stabilises mRNA and increases the synthesis of TfR1.

# REGULATION OF INTRACELLULAR HOMEOSTASIS

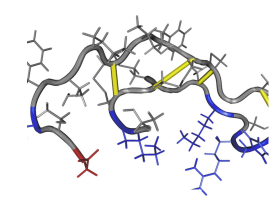
- Aconitase is a "moonlighting" protein since it has more than one role in the cell;
- Eukaryotes have 2 isoforms: the mithochondrial one is part of the TCA cycle converting citrate into isocitrate
- The cytosolic isoform has 2 roles:
  - 1- converts citrate into isocitrate providing the substrate for isocitrate dehydrogenase that generates NADPH
  - 2- it participated to iron homeostasis
- Aconitase has a Fe-S clusetr that detaches when iron levels are low
- The apoenzyme has a new activity, since it can bind to mRNA and regulates the expression of of ferritin and and Tfr
- Aconitase is structurally identical to IRP1 and similar to IRP2



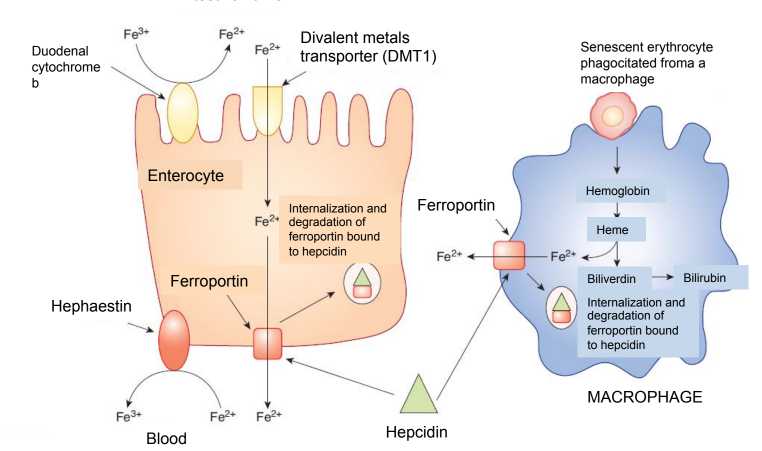
Box 16-1 figure 2
Lehninger Principles of Biochemistry, Sixth Edition
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### **HEPCIDIN**

- It is a protein with a key role in iron homeostasis
- It is synthesised by liver as a pro-peptide of 84 aa (pro-hepicidin)
- The bioactive peptide contains 25 aa
- This peptide binds to the cellular iron exporter, ferroportin, and initiates internalization and degradation

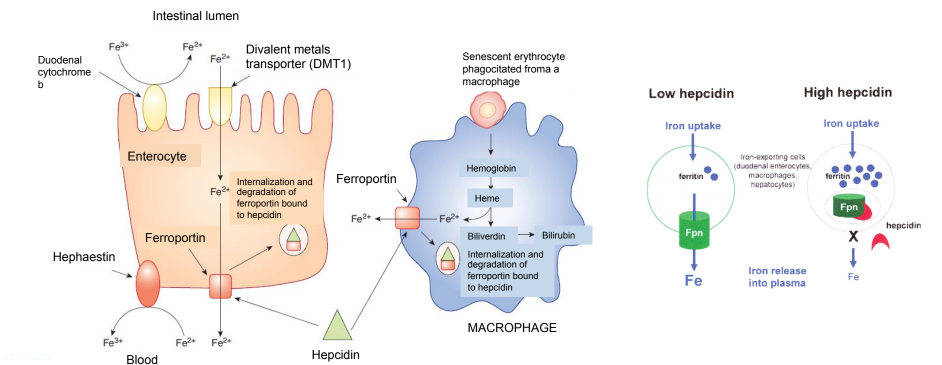


#### Intestinal lumen

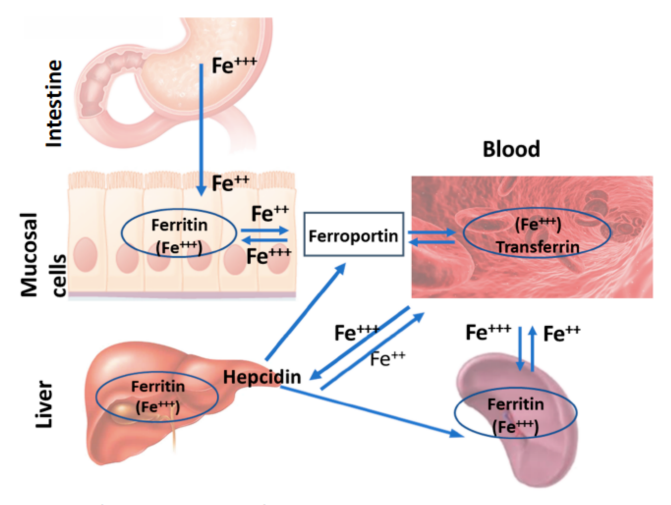


#### **HEPICIDIN: REGULATION**

- The major mechanism of hepicidin is THE REGULATION OF TRANSMEMBRANE IRON TRANSPORT
- It binds to FERROPORTIN, forms hepicidin-ferroportin complex ,which is degraded in the lysosomes and iron is locked inside the cells (mainly enterocytes, hepatocytes and macrophages).
- When iron levels are low, hepicidin levels are low as well and iron exporting cells have abundant ferroportin and thus releases iron into plasma.
- When iron levels are high, hepicidin concentration increases and it binds to ferroportin and thus iron is retained in the cells.



# REGULATION OF IRON HOMEOSTASIS



**Figure 1.** The main tissues involved in the regulation of systemic iron metabolism. Duodenal enterocytes are responsible for dietary iron absorption. Upon absorption, iron circulates around the body bound to the protein transferrin and is taken up by different tissues for utilisation. The reticuloendothelial system, which includes the splenic macrophages, recycles iron from senescent erythrocytes. Among many other functions, the liver produces the hormone hepcidin. Hepcidin controls the release of iron from enterocytes and macrophages into the circulation and is regarded as the master regulator of systemic iron metabolism.

# REGULATION OF IRON HOMEOSTASIS

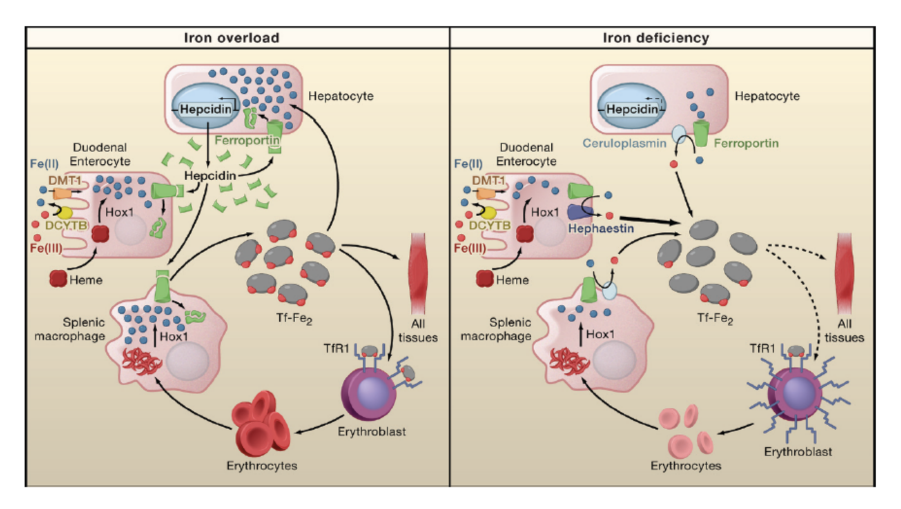


Figure 1. Regulation of Systemic Iron Homeostasis

Cells involved in systemic iron regulation are shown. Divalent metal transporter 1 (DMT1) at the apical membrane of enterocytes takes up iron from the lumen of the duodenum after DCYTB reduces Fe(II) to Fe(II). Ferroportin at the basolateral membrane cooperates with hephaestin that oxidizes Fe(II) to Fe(III). Iron-loaded (diferric) transferrin (Tf-Fe<sub>2</sub>), indicated by red dots, supplies iron to all cells by binding to the transferrin receptor 1 (TfR1) and subsequent endocytosis. TfR1 is highly expressed on hemoglobin-synthesizing erythroblasts. Hepatocytes sense transferrin saturation/iron stores and release hepcidin accordingly. Red cell iron is recycled by macrophages via ferroportin and the ferroxidase ceruloplasmin. In iron overload (left), high hepcidin levels inhibit ferroportin-mediated iron export by triggering internalization and degradation of the complex to reduce transferrin saturation. Hepcidin expression is high. In iron deficiency (right), iron is released by ferroportin into the circulation. Hemoglobin-derived heme is catabolized in macrophages by hemoxygenase-1 (HOX1). Hepcidin expression is low.