

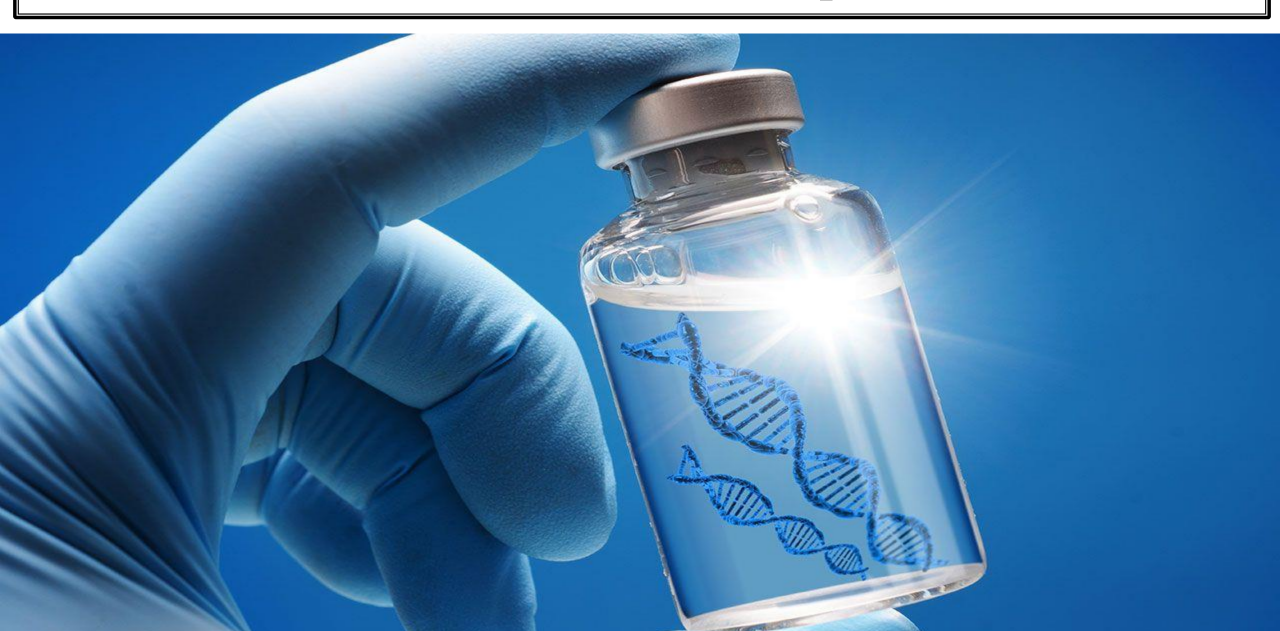
College of Pharmacy Fifth Stage

Pharmaceutical Biotechnology

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Lectutre 5
Routes of Administration and Absorption
Enhancement

Routes of Administration and Absorption Enhancement



Handling of Pharmaceutical Proteins Post-Production

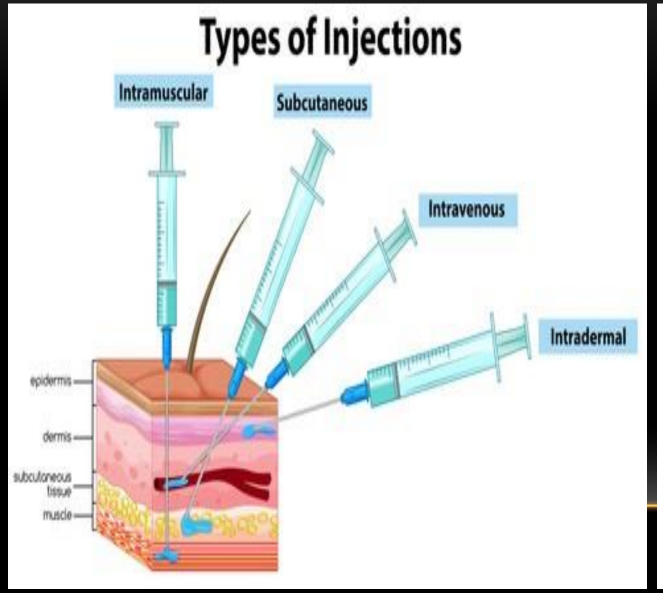
- Protein formulations undergo extensive testing during development and formulation to ensure their maximus stability and effectiveness.
- In spite of all these efforts, pharmaceutical proteins **remain sensitive** to 'real life' handling and may readily show degradation reactions that obviously affect both efficacy and safety.
- Set of instruction is generated for health professionals and patients about the conditions that should be maintained for the product, e.g., storage temperature window, avoidance of shaking/shear, exposure to light.
- As an example, the package insert of trastuzumab states: 'Swirl the vial gently to aid reconstitution. Trastuzumab may be sensitive to shear-induced stress, e.g., agitation or rapid <u>expulsion</u> from a syringe. **DO NOT SHAKE**.

Routes of Administration

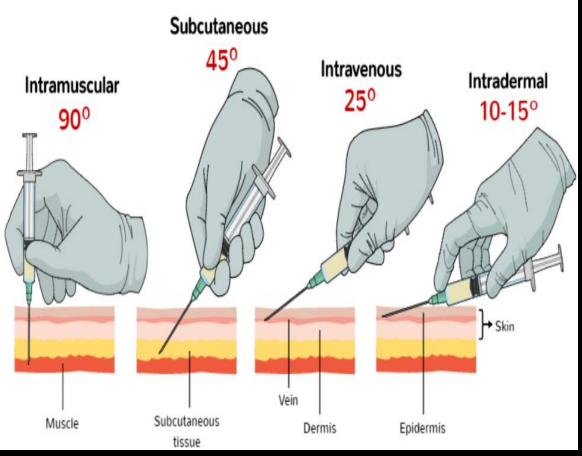
1. The Parenteral Route of Administration

- Parenteral administration is defined as administration via those routes where a needle is used, including intravenous (IV), intramuscular (IM), subcutaneous (SC), and intraperitoneal (IP) injections.
- It is Important to understand that the blood half-life of biotech products can vary over a wide range.
- For example, the circulation half-life of tissue plasminogen activator (t-PA) is a few minutes, while monoclonal antibodies reportedly have half-lives of a few days to weeks.

The Parenteral Route Of Administration



Injection technique



The Parenteral Route of Administration

- A simple way **to expand the mean residence** time for short half-life proteins is to switch from IV to IM or SC administration.
- ▶ But this may result in slow uptake to blood compartment and lower extent of absorption → this mean lower bioavailability.
- **This may be due to:**
- 1. Increase in the residence time at the IM or $SC \rightarrow$ increase in the exposure to inactivation reaction such as **peptidase**.
 - For instance, diabetics can become "insulin resistant" through high tissue peptidase activity.

The Parenteral Route of Administration

- Differences in disposition: Upon administration, the protein may reach the blood through the lymphatics or enter the blood circulation through the capillary wall at the site of injection.
 - **Lymphatic transport** takes time (**hours**), and uptake in the blood circulation is highly dependent on the injection site.
 - On its way to the blood, the lymph passes through draining lymph nodes, and contact is possible between lymph contents and cells of the immune system such as macrophages and B and T lymphocytes residing in the lymph nodes.

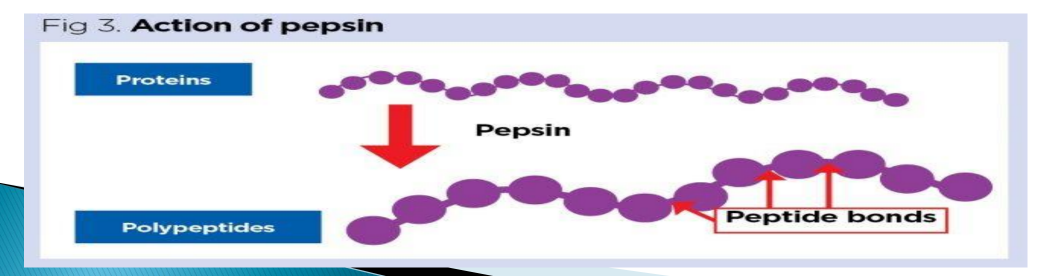
For several reasons, e.g., ease of administration, patient friendliness and cost, alternative administration routes to the parenteral route would be welcome for the successful systemic delivery of recombinant proteins.

Oral Route:

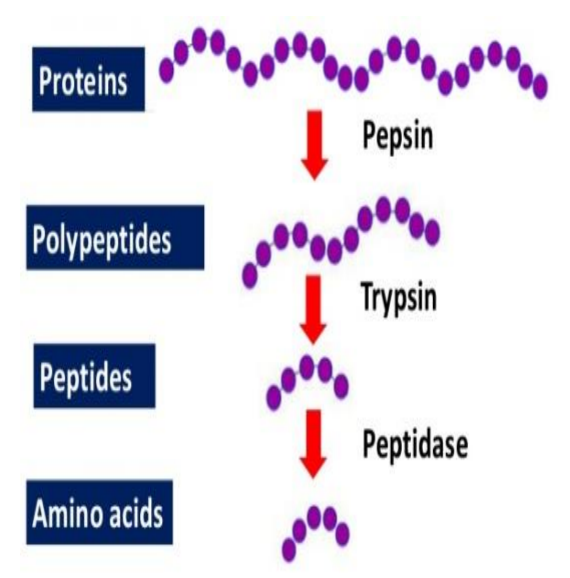
- Oral delivery of protein drugs would be preferable, because it is patient friendly and no intervention by a healthcare professional is necessary to administer the drug.
- Oral bioavailability, however, is usually very low for proteins.

- The two main reasons for this failure of uptake are (why oral bioavailability is low?):
- 1. The **efficient enzymatic system** that breakdown the protein:
 - The human body has developed a very efficient system to break down proteins in our food to amino acids or di- or tripeptides.
 - These building stones for body proteins are **actively** absorbed for use wherever necessary in the body.
 - Enzymes such as pepsin, trypsin and other are found in the GI tract.

- In the **stomach**, **pepsins**, a family of aspartic proteases, are secreted.
 - They are particularly active between **pH 3 and 5** and **lose activity** at higher pH values.
 - Pepsins are endo-peptidases capable of **cleaving peptide bonds distant from the ends** of the peptide chain.
 - They preferentially cleave peptide bonds between two hydrophobic amino acids [such as glycine (Gly), alanine (Ala), valine (Val)].



- II. Other endopeptidases are active in the gastrointestinal tract at neutral pH values, e.g., trypsin, chymotrypsin, and elastase.
- m. Exopeptidases, proteases degrading peptide chains from their ends, are present as well.
- Examples are carboxypeptidase A & B.
- In the GI lumen the proteins are cut into fragments that effectively further break down to **amino acids**, di and tripeptides by brush border, and cytoplasmic **proteases of the enterocytes**.



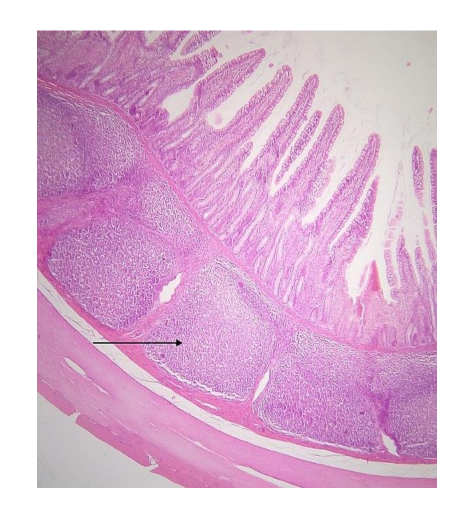
Second Reason for low oral bioavailability:

2. Permeability:

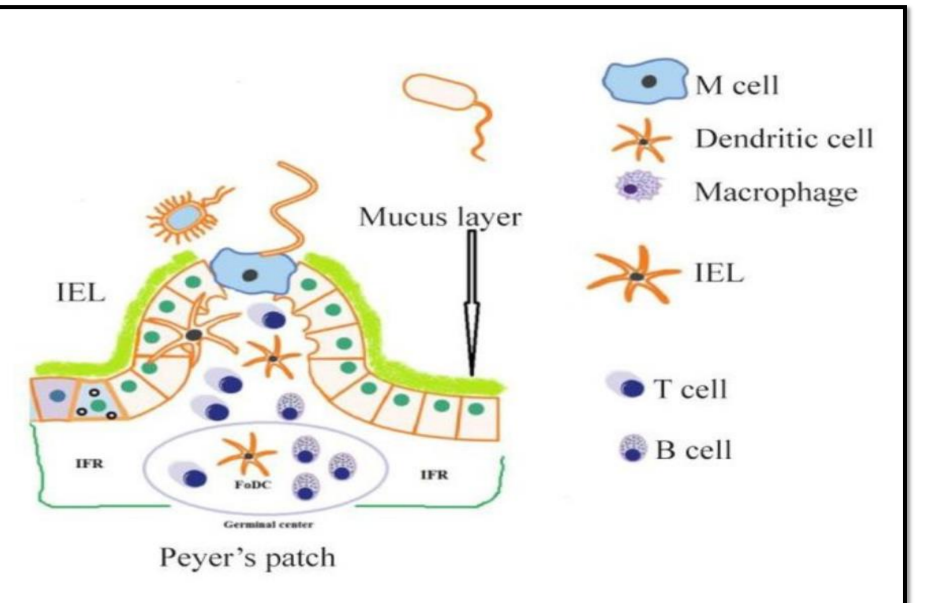
- ➤ High **molecular** weight molecules **do not readily penetrate** the intact and mature epithelial barrier.
- This leaves paracellular transfer and intracellular endocytosis into the enterocyte membrane as the sole pathway for mass transfer.

- The above analysis leads to the conclusion that nature, unfortunately, does not allow us to use the **oral route** of administration for therapeutic proteins if **high** and **constant** bioavailability is required.
- ► However, for the category of **oral vaccines**, the above mentioned hurdles of **degradation and permeation are not necessarily prohibitive**. Because: →
- 1. For oral immunization, **only a (small)** fraction of the antigen (protein) has to reach its target site to illicit an immune response.

- 2. The target cells are lymphocytes and antigen presenting accessory cells located in Peyer's patch.
 - These Peyer's patches are macroscopic identifiable follicular structures located in the wall of the GI tract.
 - Peyer's patches are overlaid with microfold (M) cells that separate the luminal contents from the lymphocytes.



- These M cells have little lysosomal degradation capacity and allow for antigen sampling by the underlying lymphocytes.
- Moreover, mucus-producing goblet cell density is reduced over Peyer's patches. This reduces mucus production and facilitates access to the M cell surface for luminal contents.



Schematic representation of Peyer's patches, M cells, and the different immune cell populations. M cells have no mucus. IFR: intra-follicular region, B: B cells, IEL: intraepithelial lymphocyte, T: T cells, FoDC: follicular dendritic cell, DC: dendritic cells.

Attempts to improve antigen delivery via the Peyer's patches and to enhance the immune response are made by using microspheres, liposomes or modified live vectors, such as attenuated bacteria and viruses.

