

*Use of Insulin in the treatment  
of diabetes mellitus*

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## Objectives:

*by the end of this lecture, students should be able to:*

- *Define diabetes and mention different types of diabetes*
- *Differentiate between difference in treating type I and type II diabetes.*
- *Understand mechanism of action, secretion, and actions of insulin.*
- *Describe different types of insulin analogues*
- *Be able to recognize the difference in pharmacokinetic profile between different types of insulin analogues.*
- *Know uses of different insulin analogues*

# Diabetes mellitus

- Is a **chronic metabolic disorder** characterized by high blood glucose level caused by caused by deficiency of insulin or by increased insulin resistance.

# Diabetes mellitus

**Fasting plasma glucose (no food for 8 hrs)**

**Normal** <100 mg/dl (5.6 mmol/l).

**Pre-diabetes** 100-125 mg/dl (5.6-6.9 mmol/L).

**Diabetes if**

**Fasting** >126 mg/dl (7 mmol/L)

**or 2h after a meal** > 200 mg/dl (11.1 mmol/L).

# Complications of diabetes

- **Cardiovascular problems**
  - **Micro- and macro-vascular disease**
- **Renal failure (nephropathy).**
- **Blindness (retinopathy).**
- **Neuropathy.**
- **Risk of foot amputation**

# Types of diabetes

- **Type I diabetes (IDDM)**  
**due to autoimmune or viral diseases**
- **Type II diabetes (NIDDM)**  
**due to genetic susceptibility and other factors (age, obesity).**

# Type I Diabetes

## IDDM

- **10-20% occurrence.**
- **During childhood or puberty**
- **$\beta$ -cells are completely destroyed.**
- **Absolute deficiency of insulin secretion**
- **Treated by insulin.**

# Type II Diabetes

## NIDDM

- **80-90% occurrence**
- **Over age 35**
- **Pancreatic  $\beta$ -cells are not producing enough insulin**
- **Obesity is an important factor.**
- **Insulin resistance in peripheral tissues.**
- **Treated by oral hypoglycemic drugs.**



<b>Characteristic</b>	<b>Type 1</b>	<b>Type 2</b>
<b>Onset (Age)</b>	Usually during childhood or puberty	Usually over age 40
<b>Type of onset</b>	Abrupt	Gradual
<b>Prevalence</b>	10-20%	80-90 %
<b>Genetic predisposition</b>	Moderate	Very strong
<b>Defects</b>	$\beta$ -cells are destroyed	$\beta$ -cells produce inadequate quantity of insulin
<b>Endogenous insulin</b>	Absent	Present (not enough)
<b>Insulin resistance</b>	absent	present
<b>Nutritional status</b>	Usually thin	Usually obese
<b>Ketosis</b>	Frequent	Usually absent
<b>Clinical symptoms</b>	Polydipsia, polyphagia, polyuria, weight loss	Often asymptomatic
<b>Related lipid abnormalities</b>	Hypercholesterolemia frequent	Cholesterol & triglycerides often elevated
<b>Treatment</b>	Insulin injection	Oral hypoglycemic drugs



# INSULIN

# Insulin receptors

- **Present on cell membranes of most tissues.**
- **Liver, muscle and adipose tissue**



# **Effects of insulin**

# I. Carbohydrate Metabolism:

- ↑ glucose uptake & utilization by peripheral tissues.
- ↑ Glycogen synthesis (**glycogen synthase**)
- ↑ Conversion of carbohydrate to fats.
- ↓ Gluconeogenesis.
- ↓ Glycogenolysis (liver).
- ↑ Glycolysis (muscle).

## II. Fat Metabolism:

- **Liver:**

- ↑ **Lipogenesis.**
- ↓ **Lipolysis.**
- **Inhibits conversion of fatty acids to keto acids.**

- **Adipose Tissue:**

- ↑ **Triglycerides storage.**
- ↑ **Fatty acids synthesis.**
- ↓ **Lipolysis**

# III. Protein Metabolism:

## Liver:

- ↓ protein catabolism.

## Muscle:

- ↑ amino acids uptake.
- ↑ protein synthesis.
- ↑ glycogen synthesis (**glycogenesis**).

## IV. potassium

- **↑ potassium uptake into cells.**



# Routes of administrations of exogenous insulin

- **Can not be given orally (why ?)**
- **Insulin syringes (s.c., arms, abdomen, thighs).**
- **Portable pin injector (pre-filled).**
- **Continuous S.C. infusion (insulin pump).**
  - **More convenient**
  - **Eliminate multiple daily injection**
  - **Programmed to deliver basal rate of insulin.**

# Routes of administrations of exogenous insulin

- **Intravenously (in a hyperglycemic emergency)**

## **Under Clinical Trials**

- **Inhaled aerosols, transdermal, intranasal.**

## Pin injector



## Insulin pump



# Insulin degradation

1. Basal level of endogenous insulin is 5-15  $\mu\text{U/ml}$ .
2. Half life of circulating insulin is 3-5 min.
3. 60% liver & 40% kidney (**endogenous insulin**)
4. 60% kidney & 40% liver (**exogenous insulin**)

# Sources of Exogenous Insulin

- **Beef Insulin**
  - Differs from human insulin by 3 amino acids (antigenic).
- **Porcine Insulin**
  - Differs by one amino acid (antigenic).

# Human Insulin analogues

- **Prepared by recombinant DNA techniques.**
- **Less immunogenic.**
- **Modifications of amino acid sequence of human insulin can change pharmacokinetics.**

# Types of insulin preparations

Differ in pharmacokinetic properties mainly

- Rate of absorption (Onset of action).
- Duration of action.

Variation is due to:

- Change of amino acid sequence.
- Size and composition of insulin crystals in preparations (monomers, dimers, hexamers).

# Types of insulin preparations

## Insulin Analogues

- **Ultra-short acting insulins**  
e.g. Lispro, aspart  
very fast onset of action and short duration
- **Short acting insulins**  
e.g. regular insulin  
fast onset of action and short duration.



# Types of insulin preparations

- **Intermediate acting insulins**
  - e.g. NPH, lente
  - **Slow onset, intermediate duration of action.**
- **Long acting insulins**
  - e.g. glargine, detemir
  - **Slow onset and long duration of action.**

# Ultra-short acting insulins

## Insulin lispro, insulin aspart

- **Clear** solutions at neutral pH.
- Do not aggregate or form dimers or hexamers (**monomeric analogue**).
- **Fast onset** of action (5-15 min)
- S.C. (5 -15 min before meal).
- **Short duration** of action (3-5 h)

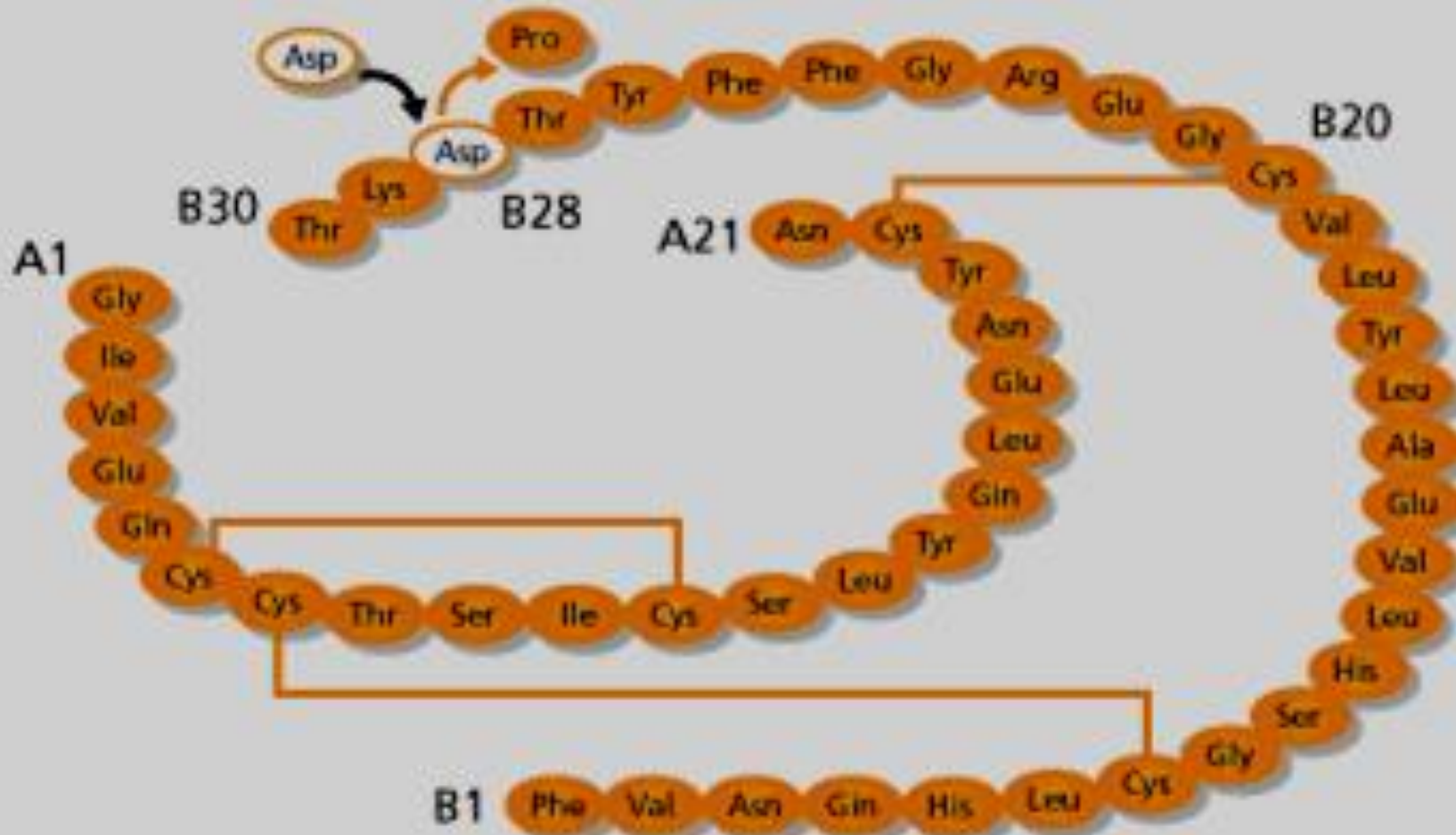
# Ultra-short acting insulins

## **Insulin Lispro, insulin aspart**

- **Reach peak level 30-90 min after injection.**
- **I.V. in emergency.**
- **When rapid-acting insulins (e.g., lispro, aspart) are mixed with another insulin, the preparation should be used immediately.**

# Insulin aspart

Structure



# Ultra-short acting insulins

## insulin lispro, insulin aspart

- 3 times/day.
- Mimic the prandial mealtime insulin release
- Preferred for external insulin pump (**Lispro does not form hexamers**)
- used to control post-prandial hyperglycemia (s.c.) and emergency diabetic ketoacidosis (i.v).

## Short acting insulins (regular insulin)

- **Soluble** crystalline zinc insulin
- **Clear** solutions at neutral pH.
- Forms hexamers.
- Onset of action 30-45 min (s.c.).
- **I.V.** in emergency situations.
- Peak 2-4 h.
- Duration 6-8 h.

# **Short acting insulins (regular insulin)**

- **2-3 times/day.**
- **Control postprandial hyperglycemia (s.c.) & emergency diabetic ketoacidosis (i.v.).**
- **Can be used in pregnancy**

	<b>Ultra-Short acting insulins e.g. Lispro, aspart, glulisine</b>
<b>Physical characteristics</b>	<b>Clear solution at neutral pH</b>
<b>chemistry</b>	<b>Monomeric</b> analogue
<b>Route &amp; time of administration</b>	<b>S.C. 5 min (no more than 15 min) before meal I.V. in emergency (e.g. diabetic ketoacidosis)</b>
<b>Onset of action</b>	<b>Fast 5 – 15 min ( S.C )</b>
<b>Peak level</b>	<b>30 – 90 min</b>
<b>Duration</b>	<b>3 – 5 hr Shorter</b>
<b>Usual administration</b>	<b>2 – 3 times/day</b>
	<b>postprandial hyperglycemia &amp; emergency diabetic ketoacidosis</b>

	<b>Short-acting (regular) insulins e.g. Humulin R, Novolin R</b>
	<b>Clear solution at neutral pH</b>
	<b>Hexameric</b> analogue
	<b>S.C. 30 – 45 min before meal I.V. in emergency (e.g. diabetic ketoacidosis)</b>
	<b>rapid 30 – 45 min ( S.C )</b>
	<b>2 – 4 hr</b>
	<b>6 – 8 hr longer</b>
	<b>2 – 3 times / day</b>
	<b>postprandial hyperglycemia &amp; emergency diabetic ketoacidosis</b>



# *Advantages of Insulin Lispro vs Regular Insulin*

- **Rapid onset of action (due to rapid absorption)**
- **Reduced risk of postprandial hypoglycemia and hyperinsulinemia (due to shorter duration of action, no more than 3-4 hrs regardless of dose).**

# Intermediate acting insulins

**Isophane (NPH) insulin**

**Lente insulin**

# Isophane (NPH) Insulin

- **NPH**, is a **Neutral Protamine Hagedorn** insulin in phosphate buffer.
- **NPH insulin** is combination of protamine & crystalline zinc insulin (1: 6 molecules).  
proteolysis release insulin.

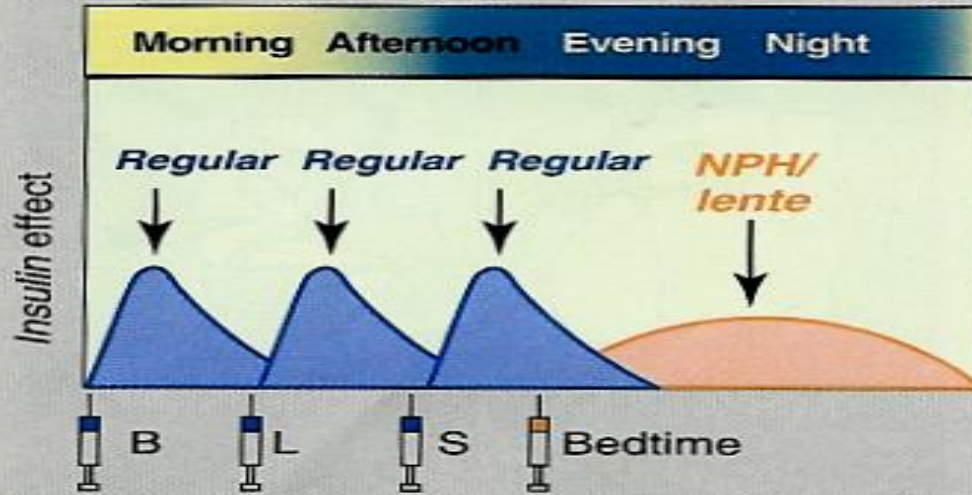
## Isophane (NPH) Insulin

- **Turbid suspension** at neutral pH.
- **Given S.C. only not i.v.**
- **Can not be used in ketoacidosis or emergency**
- **Onset of action 1-2 h.**
- **Peak serum level 5-7 h.**
- **Duration of action 13-18 h.**

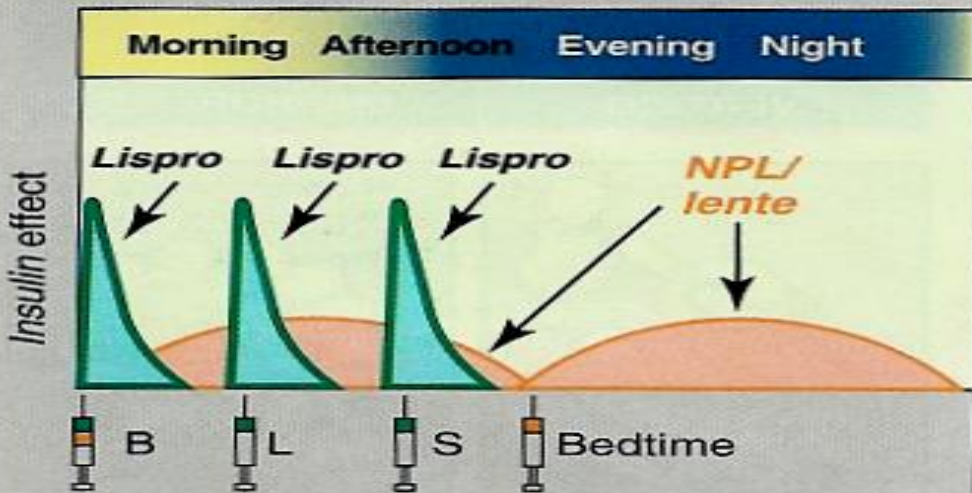
# Isophane (NPH) Insulin

## Insulin mixtures

- **NPH/regular insulin**
  - **75/25 , 70/30 , 50/50**
- **(NPL= NPH / lispro) (NPA= NPH / aspart)**
- **NPL & NPA have the same duration as NPH**
- **Have two peaks.**

**A**

**Prandial and basal insulin replacement**

**B**

# Lente insulin

- **Mixture of:**
  - **30% semilente insulin (amorphous precipitate of zinc insulin in acetate buffer)**
  - **70% ultralente insulin (poorly soluble crystal of zinc insulin)**
- **Turbid suspension at neutral pH**
- **Given S.C., not intravenously**

## Lente insulin (Humulin L, Novolin L)

- **Delayed onset of action (1-3 h)**
- **Peak serum level 4-8 h.**
- **Duration of action 13-20 h.**
- **Lente and NPH insulins are equivalent in activity.**
- **Lente is not used in diabetic ketoacidosis or emergency.**



# Long acting insulins

Insulin glargine (lantus), Insulin detemir (Levemir)

## Insulin glargine (Lantus)

- Clear solution **BUT** forms precipitate (hexamer) at injection site.
- Slow onset of action 2 h.
- absorbed less rapidly than NPH & Lente insulin.
- Given s.c., **not intravenously**
- Should not be mixed with other insulins in the same syringe.

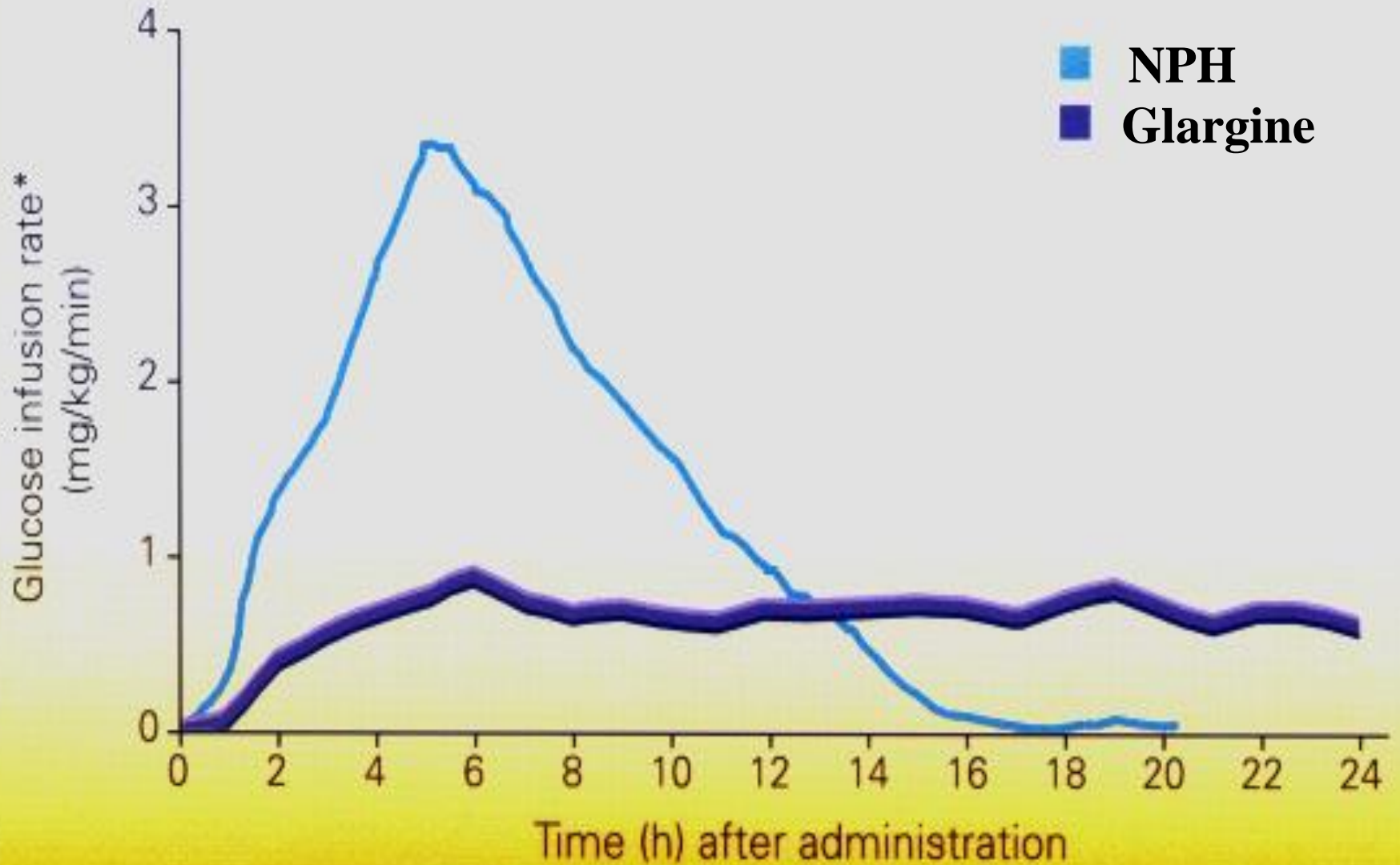
# Insulin glargine (Lantus)

- **Maximum effect after 4-5 h**
- **Prolonged duration of action (24 h).**
- **Once daily**
- **produce broad plasma concentration plateau (low continuous insulin level).**
- **Glargine must be used in regimens with rapid or short acting insulins.**

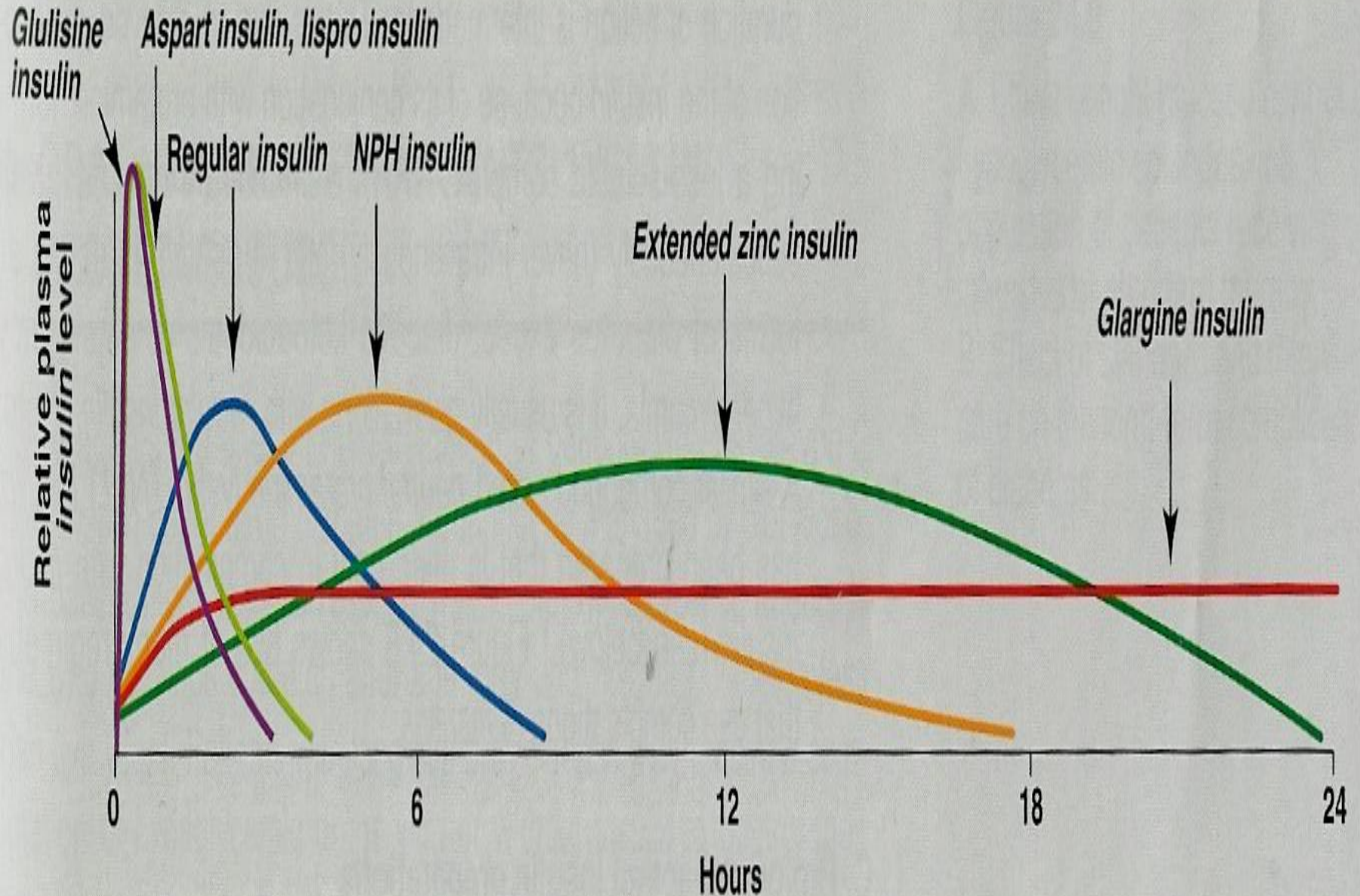
## Advantages over intermediate-acting insulins:

- **Constant circulating insulin over 24 hr with no peak (**peakless profile**).**
- **Produce flat prolonged hypoglycemic effect.**
- **Safer than NPH & Lente insulins  
( reduced risk of nocturnal hypoglycemia).**

# NPH vs Glargine



# Insulin preparations



# Insulin Dosing considerations

- **Blood glucose monitoring is required in all patients receiving insulin**
- **Rotate injection sites within the same region.**
- **Insulin should be stored in refrigerator and warm up to room temp before use.**

# Complications of Insulin Therapy:

- Hypoglycemia
- Hypersensitivity reactions.
- Lipodystrophy at injection site
- Weight gain (due to anabolic effects of insulin )
- Insulin resistance
- Hypokalemia

# Summary

- Insulin analogues are used to treat type I diabetes.
- *Fast acting insulins* (**lispro, aspart**), given s.c. or i.v., produce fast action, used to mimic postprandial insulin.
- *Short acting insulin* (**Regular insulin**), *given s.c. or i.v.* produce rapid action, used to mimic postprandial insulin.
- *Intermediate acting insulin* (**lente, Isophane**) produce slower action, than regular insulin, given s.c. not i.v.
- *Long acting insulins* (**glargine, detemir**) produce constant circulating insulin over 24 hr with no peak (peakless profile), s.c. not i.v.