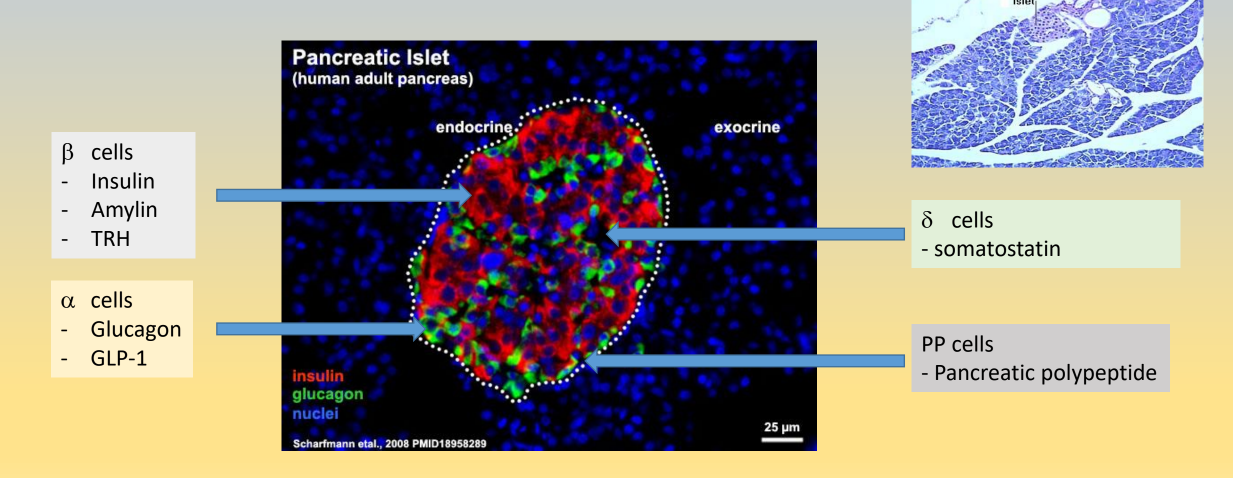
Endocrine versus exocrine pancreas

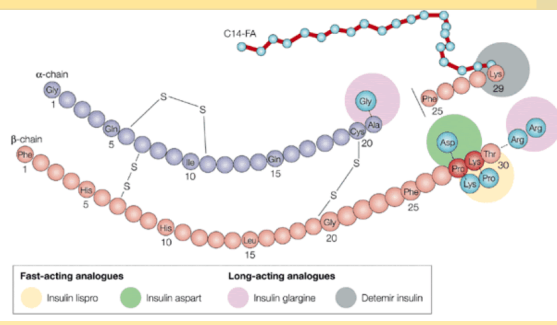


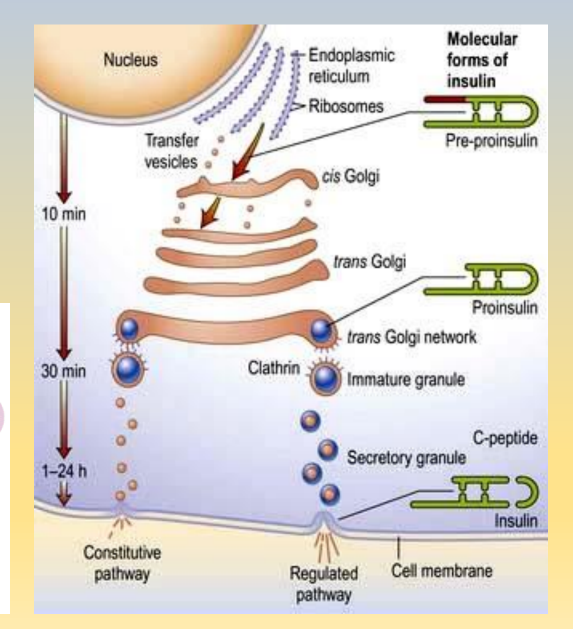
Pancreatic islets represent 1 - 2 % of pancreas, but blood flow through them represents 10 - 15 %.

Insulin

Characteristics

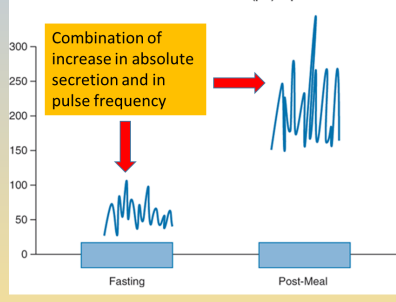
- Polypeptide
- Secretory granules free insulin and C-peptide
- Two types of secretory granules:
 - Quickly secretable (5 %)
 - Reserve pool (95 %)
- Half-time 3 8 min
- Degradation liver (up to 50 %), kidneys, target tissues (insulin proteases)



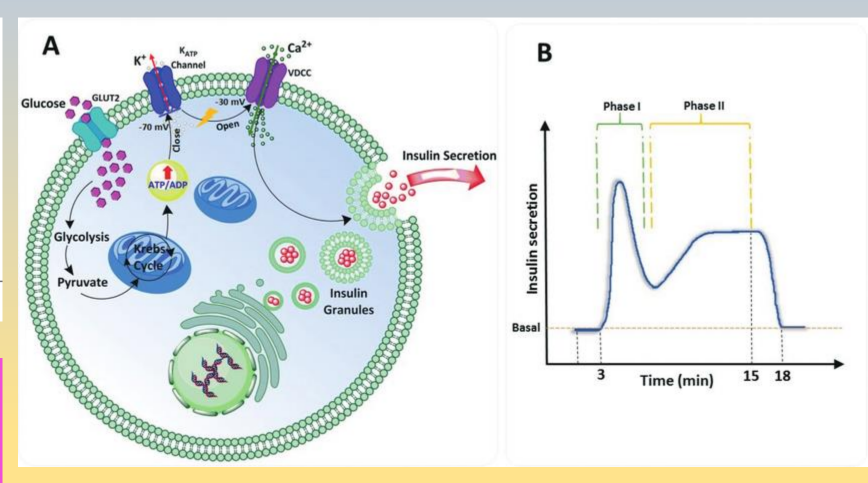


Insulin secretion

C Insulin concentrations (pM) in portal vein



Secretion of insulin is pulsatile and shows rhytmicity. Stimulation of insulin secretion by glucose is biphasic. Glucose exhibits incretin effect.



Regulation of insulin secretion

 β cells = neuroendocrine integrator, response to:

- Plasmatic concentrations of substrates (AA, Glu)
- PC of hormones (insulin, GLP-1, somatostatin, adrenaline)
- PC of neurotransmitters (noradrenaline, acetylcholine)

Glu

- Main mechanism!

AA – Leu, Arg, Lys

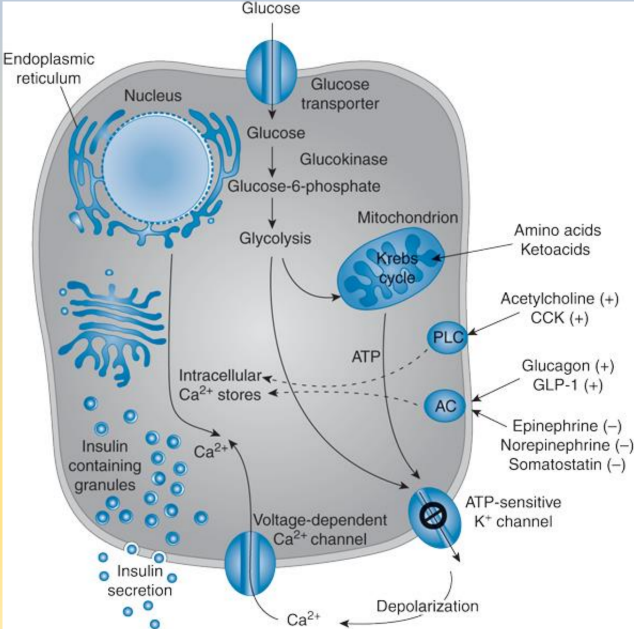
- Generation of ATP
- Direct depolarization of plasmatic membrane

Modification of mRNA translation

- Glu – (+) mRNA

Other: - GH, VIP, secretin, gastrin, glucocorticoids, prolactin, placental lactogene, sex hormones

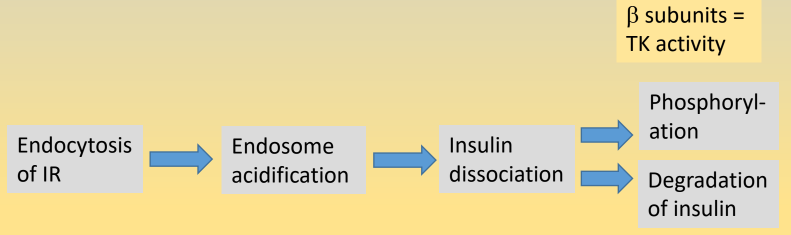
Glucose is the main stimulus for insulin secretion. Glucose has a permissive effect on secretion of other insulin secretion modulators.



Insulin receptor

Characteristics

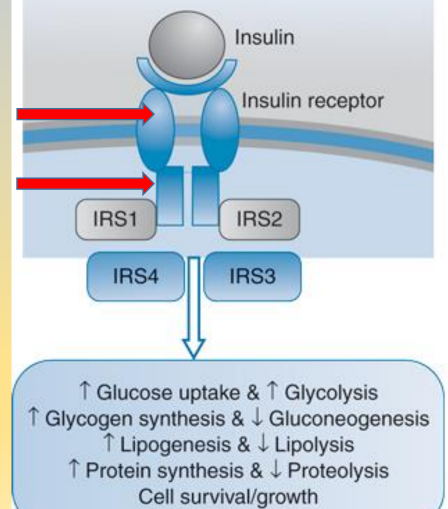
- 2 α and 2 β subunits
- TK activity
- Phosphorylation of IRS 1-4 (insulin receptor substrate)
- Interaction with other cell substrates
- PI3K (phosphatydylinositol-3-kinase)
- MAPK (mitogen-activated protein kinase)



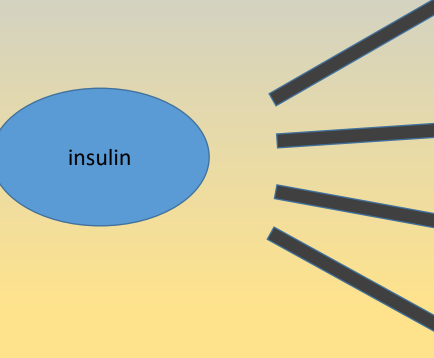
 α subunits =

Ligand binding

Number of available IR is influenced by exercise, diet, insulin itself and by other hormones. Obesity and chronic hyperinsulinemia causes significant decrease in number of IR, exercise and starvation significant increase in number of IR.



Physiologic effects of insulin



Immediate effects

- Seconds
- Modulation of K+ and Glu

transport

Early effects

- Several minutes
- Regulation of metabolic activity

Medium-term effects

- Minutes to hours
- Regulation of metabolic activity

Delayed effects

- Hours to days
- Cell growth
- Cell differentation

Effect of insulin on target tissue is anabolic and is mediated by insulin receptor.

Immediate effects of insulin on target tissues

 Utilization of glucose Approx. 40 % of glucose in body Approx. 80 – 90 % skeletal muscles Adipose tissue - adipocytes 	Transporter	Expression	Function
	GLUT1	 Ubiquitous Ery, endothelial cells (CNS), placenta, kidneys, colon Skeletal muscles and adipocytes 	- Basal uptake of Glu
- GLUT4 While GLUT1 is responsible for basal uptake of glucose by skeletal muscles and adipocytes, GLUT4 is stimulated by insulin and is responsible for insulin-stimulated uptake of glucose.	GLUT2	 β cells of pancreas Liver, small intestine, kidneys 	 Glu sensor Uptake of Glu during high concentrations of circulating Glu
	GLUT3	Primarily neuronsPlacenta, liver, epithelial cells of GIT	Basal uptake of GluEssential role in CNS
	GLUT4	Skeletal muscles and adipocytesVesicles!	 Insulin-stimulated uptake of Glu
	GLUT5	- Jejunum, sperms	- Transport of Fru

Utilization of glucose is the main immediate effect of insulin.

Early and medium-term effects of insulin

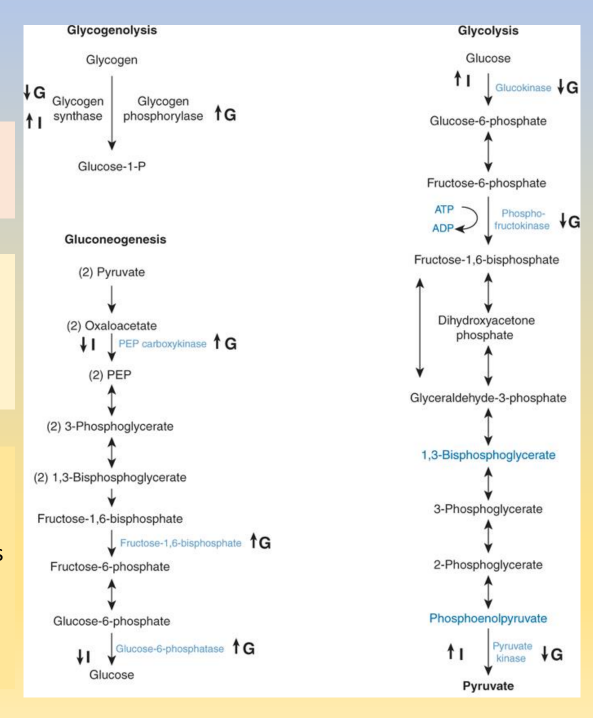
- Determined by phosphorylation of enzyme connected to metabolic pathways.
- Skeletal muscles, adipose tissue, liver

Production of ketone bodies (-)

- Dephosphorylation of hormone-sensitive lipase = inhibition of triglyceride utilization
- Activation of acetylcoenzyme A carboxylase (lipogenesis)
- Antagonization of catecholamines effect on lipolysis

Utilization of glucose

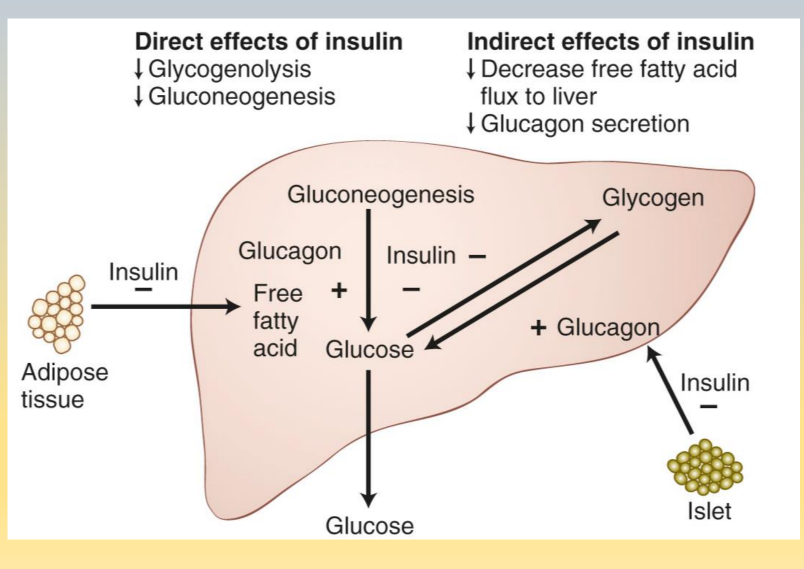
- liver
- Stimulation of expression of enzymes connected to Glu utilization (glucokinase, pyruvate kinase) and lipogenic enzymes
- Inhibition of enzymes connected to Glu production (phosphoenolpyruvate carboxykinase, glucose-6-phosphatase)
- Synthesis of glycogen
- Inhibition of production of keto bodies



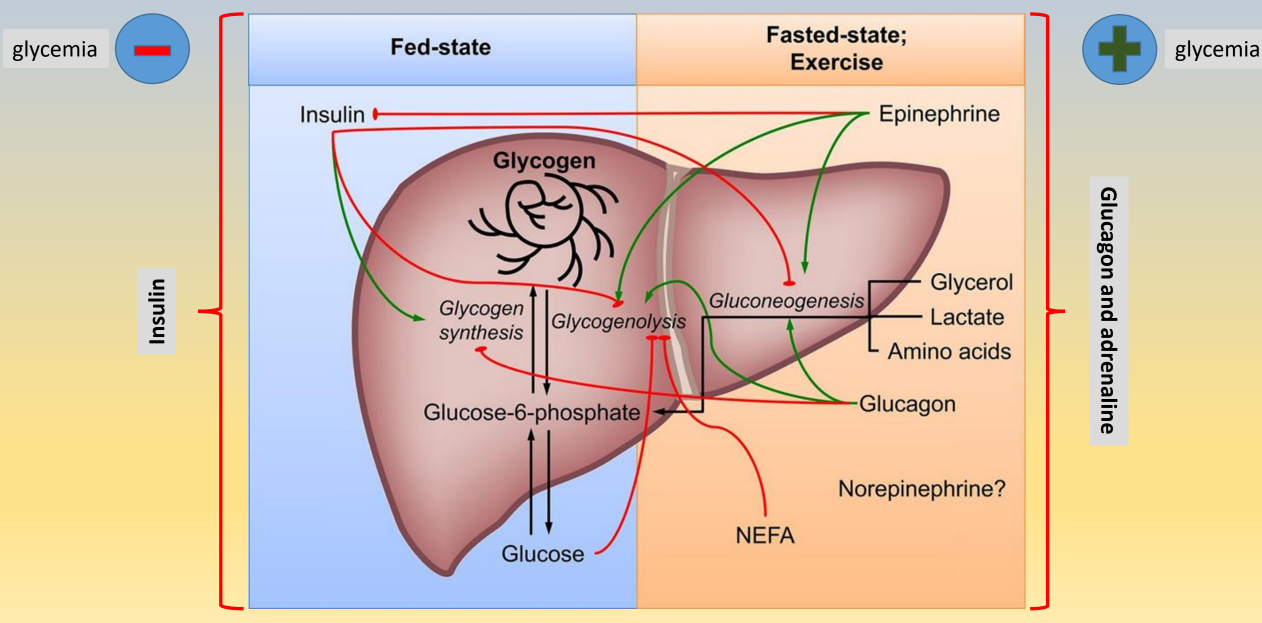
Insulin and skeletal muscles

- (+) uptake of glucose (GLUT4)
- (+) glycogen synthesis
- (+) transport of AA
- (+) translation of mRNA
- (-) degradation of proteins
- (+) preference of fat reserves
- mechanism mTOR phosphorylation

Insulin and liver

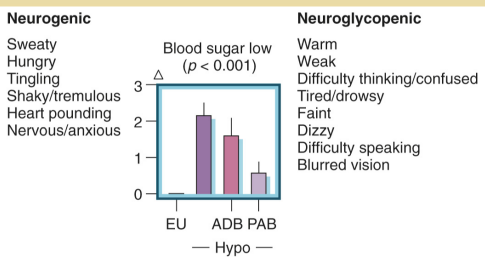


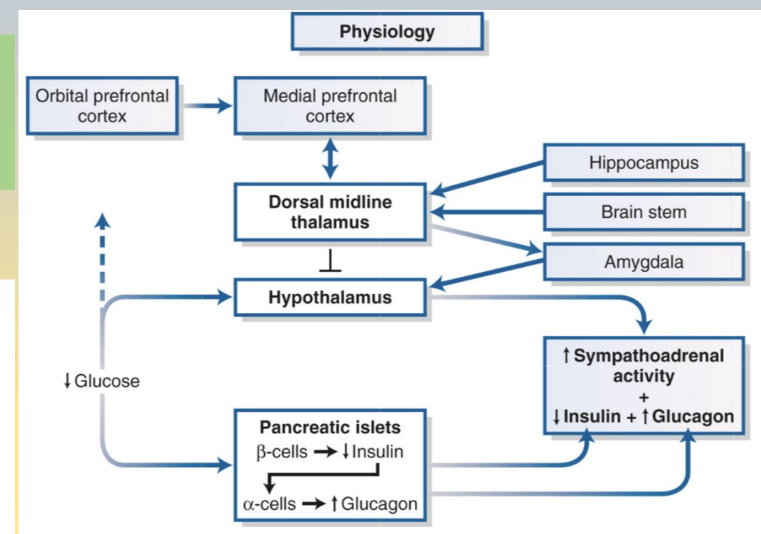
Fed-state versus fasted-state



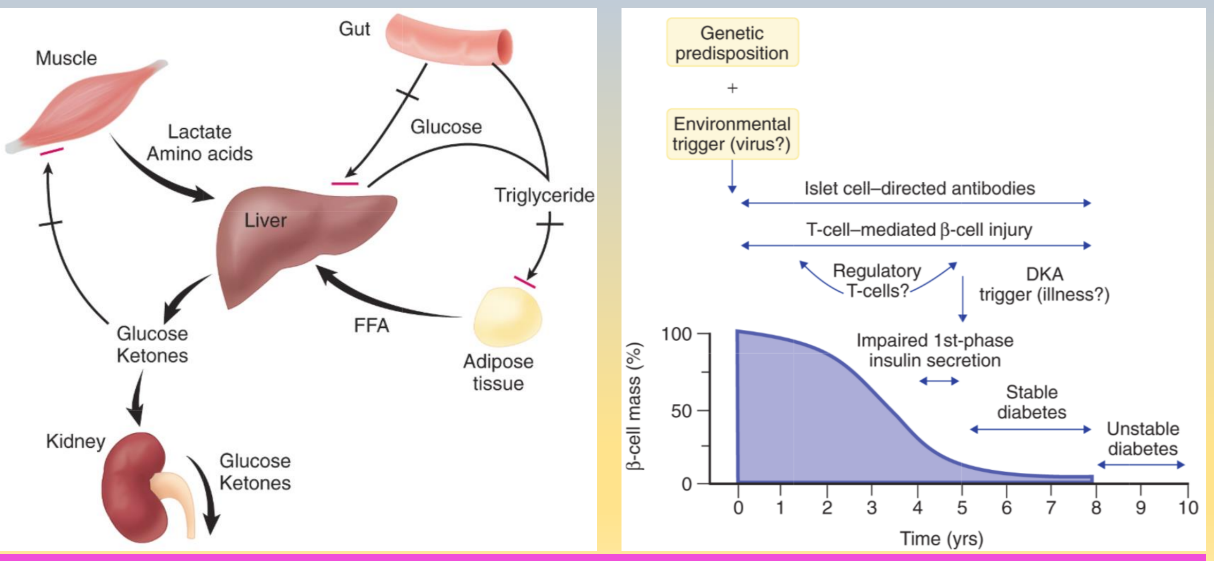
Hypoglycemia

- (-) insulin secretion
- (+) glucagon and adrenaline secretion (liver)
- (+) GH and cortisol (decreased utilization of Glu)





Diabetes mellitus type 1



DM1 is associated with mobilization of substrates for gluconeogenesis and ketogenesis from muscle and adipose tissue, increased gluconeogenesis and ketogenesis in the liver, as well as disturbed substrate intake by peripheral tissues.

Diabetes mellitus type 2

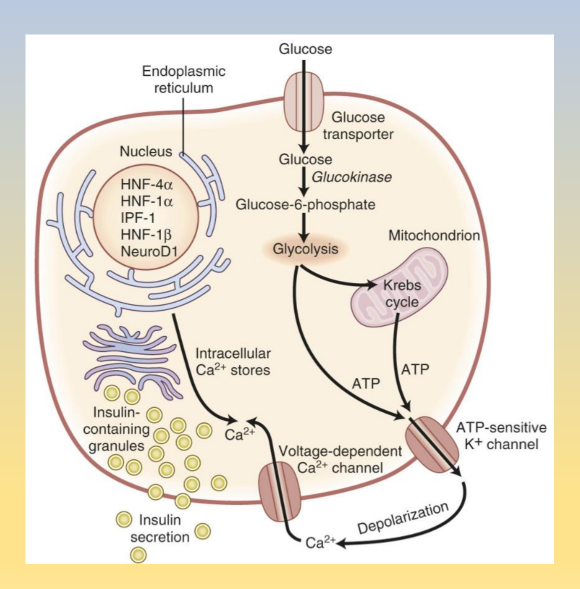
Clinical relevance

Insulin resistance

- Mutation in IR gene

Defects in insulin secretion

- Mutation in insulin gene (proinsulin)
- Mutation in mitochondrial genes
- MODY (Maturity-onset diabetes of the young)
 - HNF-4α (MODY 1)
 - Glucokinase (MODY 2)
 - HNF-1α (MODY 3)
 - IPF1 (MODY 4)
 - HNF-1β (MODY 5)
 - NeuroD1/BETA2 (MODY 6)



DM2 is multifactorial disease connected with resistence of peripheral tissues (muscles, adipose tissue) to insulin, disturbed insulin secretion (under glycemia influence) and increased glucose production in liver.

What are the consequences of DM?

Proteins

- Protein catabolism
- Negative nitrogen balance

Lipids

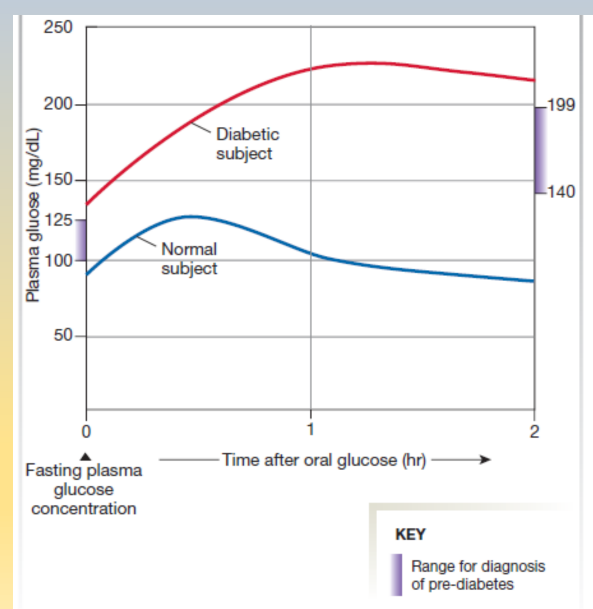
- Lipid catabolism with production of ketone bodies
- Decreased synthesis of FA and triglycerids
- Increased concentration of free FA
- FA catabolism, production of ketone bodies

Hyperglycemia

- Glycosuria, osmotic diuresis and polyuria
- Increased plasma osmolality, polydipsia, ADH
- Dehydratation
- Decreased blood pressure and volume of ECF
- Polyphagy

Ketoacidosis

- Metabolic acidosis
- Hyperventilation
- Acidification of urine
- Hyperkalemia



Glucagon

Characteristics

- Peptide hormone (29 AA)
- Syntesized as proglucagon
- Pancreas
- Enteroendocrine cels in GIT
- CNS
- Alternative splicing creates other peptides, most important GLP-1
- Short half-life (5 10 min)
- Degradation in liver

Secretion

- (+) AA

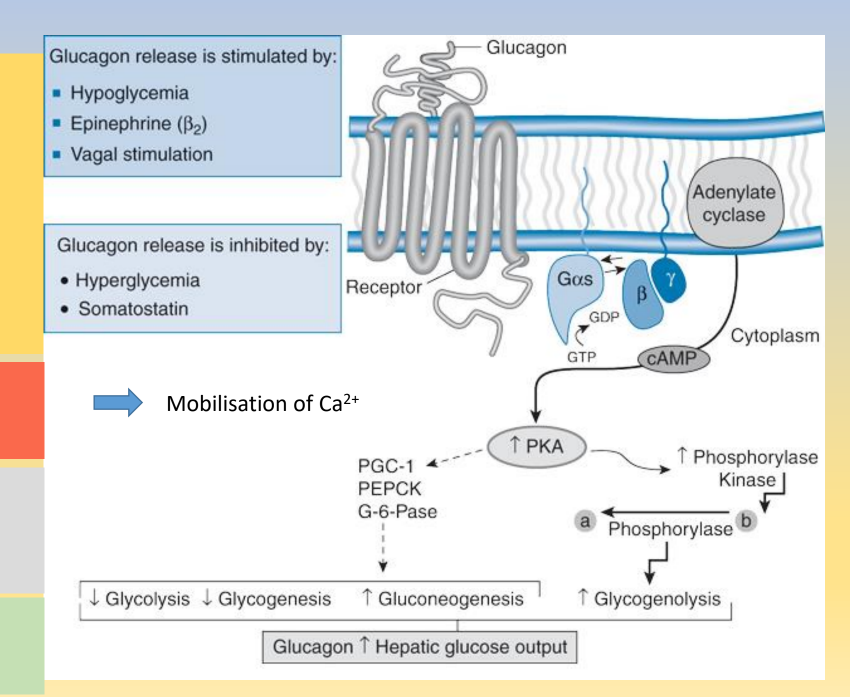
- (+) hypoglycemia

Receptors

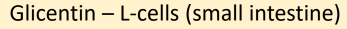
 Liver, β cells, kidneys, heart, adipose tissue, blood vessels, CNS, stomach, adrenal glands

Functions

 Glucose homeostasis – insulin antagonism



Proglucagon – alternative splicing



- Stimulation of insulin secretion
- Inhibition of stomach secretion
- Trophic effect in intestine

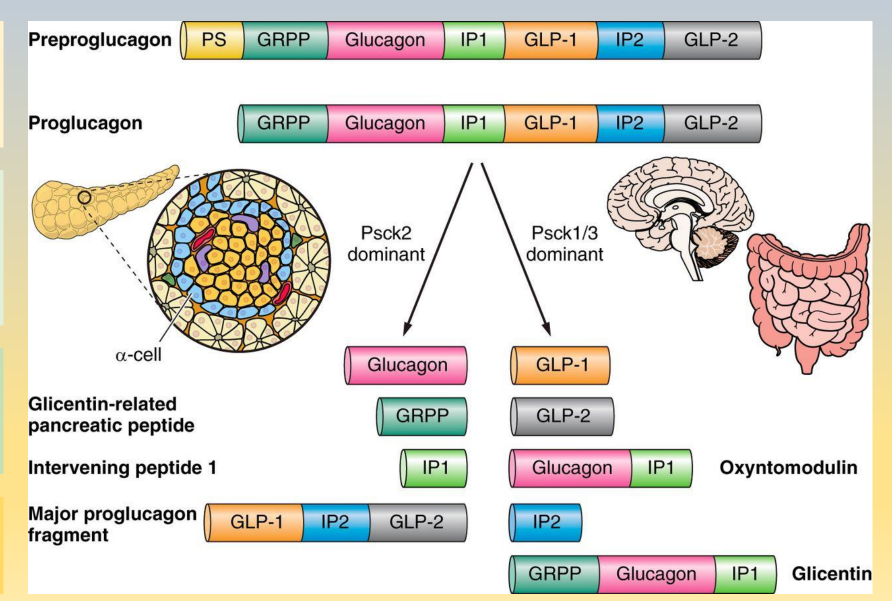
Oxyntomodulin – colon (anorexigenic factor)

- Postprandial secretion
- Increased energy expandituree
- (+) glucose tolerance

GRPP

(inhibition of Glu-stimulated insulin secretion, modulator of energy metabolism)

IP-1, IP-2 L-cells (modulation of insulin secretion?)



GLP-1 and GLP-2

Charakteristics

- Neuroendocrine L cells

Functions – GLP-1 (GLP1R)

- (+) insulin secretion
- (-) glucagon secretion
- (+) neogenesis and proliferation of pancreatic isles
- (-) β cell apoptosis

Functions – GLP-2 (GLP2R)

- (-) antrum motility
- (-) of gastric juice secretion stimulated by food
- Trophic effect (small intestine, colon)
- (-) enterocyte apoptosis
- (+) blood flow and nutrient absorption

CNS

- Caudal NTS viscerosensoric information
- Activation of POMC neurons
- Inhibition of food intake (anorexigenic factor)
- Induction of satiety

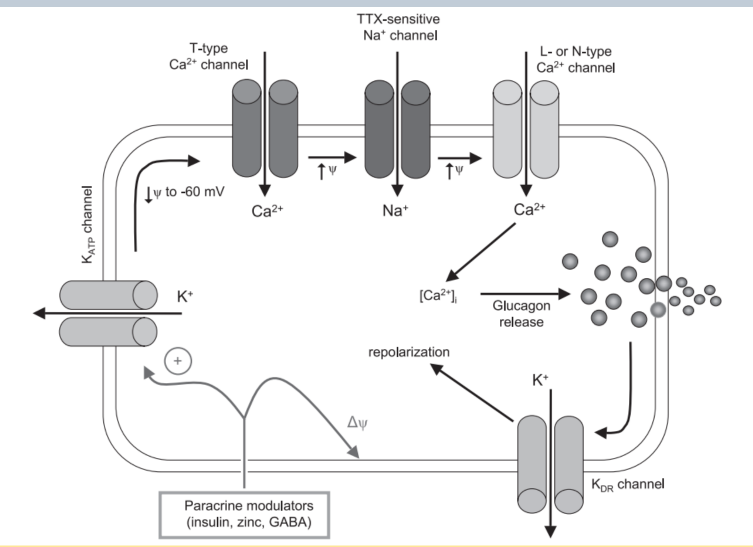
= quick modification of food intake based on metabolic substrates (glucose), hormones (leptin) and neuropeptides.

Clinical relevance

- Agonists of GLP1R treatment of DM2
 - Exenatid, lixisenatid
 - Liraglutid
 - Albiglutid, dulaglutid
- Inhibitors of dipeptidyl peptidase 4 (DPP4)
 - sitagliptin, vildagliptin, saxagliptin, alogliptin, linagliptin
 - DM2

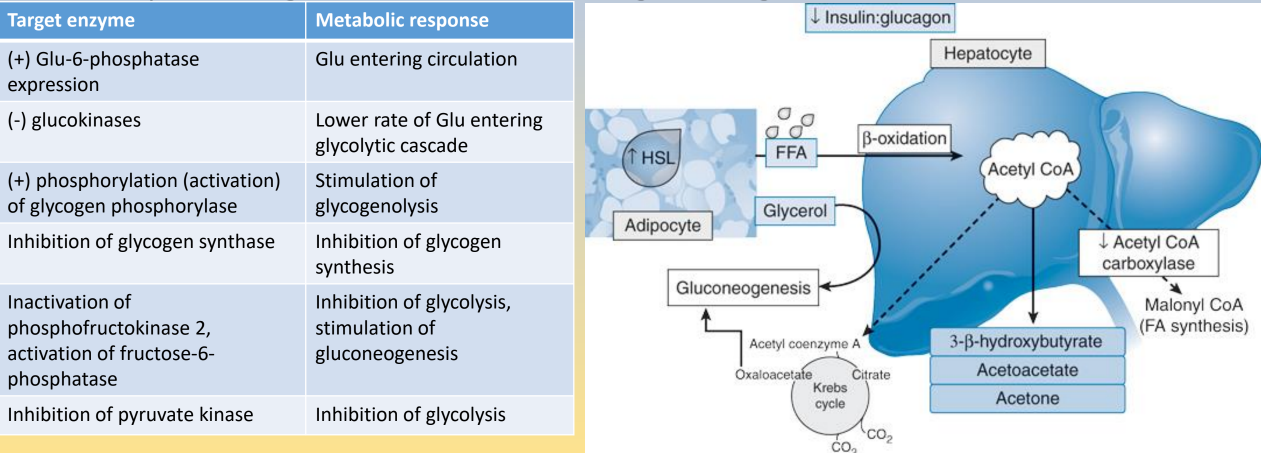
GLP-1 and GLP-2 show incretin effect preparing insulin secretion in dependence on glucose presence in GIT lumen.

Glucagon – secretion and its regulation



Glucagon secretion requires depolarizing cascade which ends with Ca²⁺ influx and glucagon secretion.

Physiological effects of glucagon



Other effects

- Stimulation of phosphorylation (activation) of hormone-sensitive lipase and lipolysis substrates for gluconeogenesis and antibody production
- FFA as a source of energy mainly for skeletal muscles

Target organ for glucagon effect is liver, where it stimulates gluconeogenesis and glycogenolysis, thus increasing glycemia.

Somatostatin

Characteristics

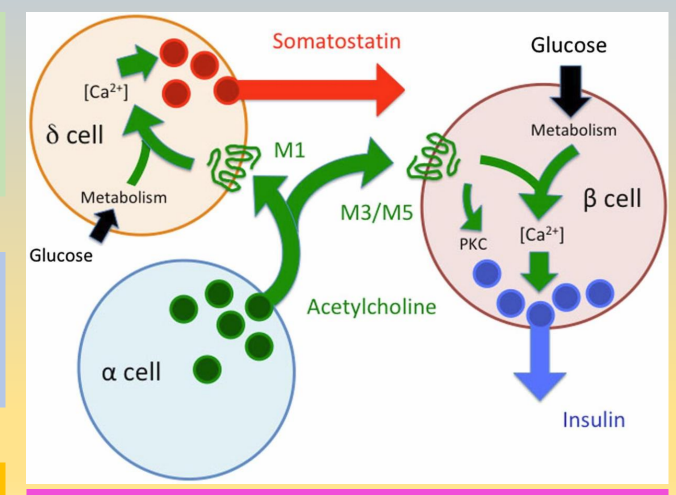
- Peptide hormone (14 AA)
- Secretion stimulated by:
 - food rich in lipids (FFA)
 - food rich in saccharides (Glu)
 - food rich in proteins (AA Leu, Arg)

Functions

- Paracrine effect (-) insulin, glucagon, PP
- Inhibition of practically all exocrine and endocrine GIT functions
- Inhibition of motility

Clinical relevance

- Somatostatin analogues and insulin/glucagonproducing tumors



Role of paracrine cholinergic signaling in somatostatin secretion – paracrine effect of acetylcholine stimulates insulin secretion, but also secretion of somatostatin.

Pancreatic polypeptide - PP

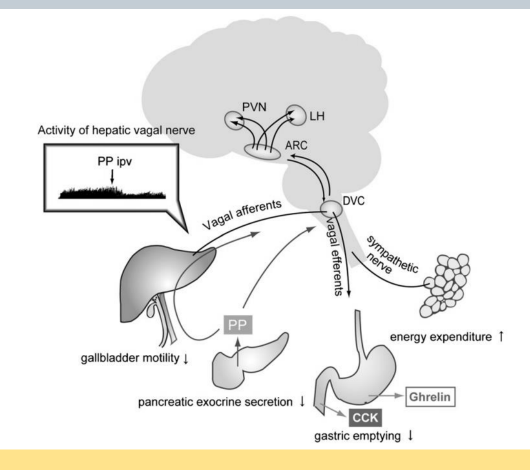
Characteristics

- Peptide hormone (36 AA)
- Secretion stimulated by:
 - Food (proteins), distention of stomach
 - Exercise
 - Direct vagal stimulation
 - Insulin-induced hypoglycemia
- Secretion inhibited by:
 - Hyperglycemia
 - Bombesin, somatostatin
- Receptors:
 - Stomach, small intestine, colon, pankreas, prostate, enteric NS, CNS

Functions

- Inhibition of pancreatic exocrine secretion
- Inhibition of gallbladder contraction
- Modulation of stomach secretion
- Modulation of stomach motility
- Regulation of food intake?

Pancreatic polypeptide stimulates energy consumption through sympathetic stimulation of brown adipose tissue. It also modulates secretion of CCK and inhibits ghrelin secretion.



Amylin

Characteristics

- Peptide hormone (37 AA)
- $-\beta$ cells, stomach, proximal small intestine
- Posttranslational modification (amidation)
- Secretion together with insulin and C-peptide
- Increase after application of:
 - p.o. and p.e. glucose

Function

- Slowing of emptying of stomach on vagal basis
- Inhibition of glucagon secretion (postprandial)
- Muscles
 - Inhibition of glycogen synthesis
 - Stimulation of glycogenolysis, glycolysis and lactate production

Clinical relevance

- Increased plasmatic concentration during obesity, gastric diabetes and DM2
- Analogue of amylin DM1 and DM2 therapy (pramlintid) amylin-deficient states

