Metabolic Changes in Diabetes Mellitus					
	DM type 1	DM type 2			
Age of onset	Childhood or puberty	Adult , frequently after age 35			
Symptoms develop	Rapidly	Gradually			
Nutritional status at the time of disease onset	Frequently undernourished	Obesity usually present			
Prevalence	10%	90%			
Defect & deficiency	Destroy β cells , eliminating production of insulin	No enough insulin or insulin resistance			
Ketosis	Common	Rare			
Plasma insulin	Low or absent	Reduce gradually (Early high —late low)			
Acute complication	Ketoacidosis	Hyperosmolar coma			
Genetic predisposition	Moderate	Very strong			
Using of oral hypoglycemic	No response	Response			
Treatment	Insulin	Diet, exercise, oral hypoglycemic, insulin			
	Actural course of T1DDA	Progression of t2DM			
Intertissue Relationship	Roce de la des l	ELCO			

Criteria for Diagnosis of DM	Increased risk of DM FPG (5.6-6.9) mmol/L OGTT (7.8-11) mmol/L A1c (5.7-6.4)%	A1c > 6.5% Random plasma glucos + Hyperglycemia sym In the absence of une	ptoms quivocal hyperglycemia ,		
		criteria 1-3 should be	confirmed by repeat testing.		
HEMOGLOBIN A1			audation of homeolohin		
 Hemoglobin A1C (A1C) is the result of non enzymatic covalent glycosylation of hemoglobin It is used to estimate glycemic control in the last 1-2 months Recently, A1C is recommended for the detection of T2DM A1C and fasting plasma glucose (FPG) were found to be similarly effective in diagnosing diabetes. A1C cut-off point of >6.5 % is used to diagnose diabetes. A1C values also correlate with the prevalence of retinopathy Assays for A1C has to be standardized according to the National Glycohemoglobin Standardization 					
Program (NGS	P).				
Metabolic Effects	Metabolic Effects of Diabetes Mellitus				
 Absolute or relative insulin deficiency → ↓ Glucose uptake (muscle & adipose tissue) ↑ Glucose production (liver) 					
		Metabolic changes in DM			
	ve insulin deficiency \rightarrow Mu				
CHO metabo	-	Lipid metabolism	Protein metabolism		
 ↓ Glucose up (adipose tissue) ↑ Glycogenol ↑ Gluconeog 	lysis	 ↑ Lipolysis ↑ Fatty acid oxidation ↑ Production of Ketone bodies 	 ↓ Protein synthesis ↑ Protein degradation 		
	Mechanisms of	f Increase Hepatic Glucose C	Dutput		
	bitory effect on glucagon s	. –			
\uparrow Glucagon → \uparrow Glu		nolysis (liver) →↑Plasma glu			
		ecrease of Peripheral Glucos			
	Muscle	Adipose Tissue			
↓ Insulin ↓ ↓Glucose & amino acid uptake ↑Protein breakdown ↓ ↑Plasma glucose ↑Plasma amino acids		↓ Insulin ↓ ↓ Glucose uptake ↓ ↑Plasma glucose			
Mechanisms of Diabetic Complications					
Typical Progression of T2DM					

General Mechanisms for Diabetic Microvascu	lar Complications				
Chronic hyperglycemia →					
1- \uparrow AGEs of essential cellular proteins \rightarrow cellular defects					
2- \uparrow Intracellular sorbitol \rightarrow \uparrow cell osmolality \rightarrow cellular swel					
3- \uparrow ROS \rightarrow oxidative stress \rightarrow cell damage	5				
Advanced Glycosylation End Produc	ts (AGEs)				
 ○ Chronic hyperglycemia →non-enzymatic combination bet 					
in proteins → formation of AGEs					
• AGEs may cross link with collagen \rightarrow microvascular comp					
• The interaction between AGEs and their receptor (RAGE)					
species (ROS) \rightarrow inflammation					
Polyol pathway					
• Glucose is metabolized to sorbitol within the cells by aldose	Sorbitol Metabolism				
reductase	(Polyol Pathway)				
The role of sorbitol in the pathogenesis of diabetic	A SEMINAL VESICLES Glycolysis Glycolysis				
complications is uncertain. Hypotheses are:					
 During sorbitol production, consumption of NADPH 	BLOOD GLOOP Glucose Glucose H+0-04 (elevated) NADPH + H*				
\rightarrow oxidative stress.	Glucose Addise Addise NADPH + H*				
• Sorbitol accumulation \rightarrow	CH2OH				
• Increase the intracellular osmotic pressure \rightarrow	HO-CH H-C-CH H-C-CH H-C-CH H-Q-CH CH/CH				
osmotic drag of fluid from extracellular space	Sorbitol Sorbitol				
 → cell swelling Alteration in the activity of PKC → altered 	Die generation app*				
VEGF activity \rightarrow altered vascular permeability	H0-5-01 H-5-01 H-5-04 CH,04				
	Fructose				
Diabetic Retinopathy					
A progressive microvascular complication of DM, affecting the retina of the eye					
A major cause of morbidity in DM (→ blindness)					
 Its prevalence 1 with increasing duration of disease in bot After 20 years of the disease: 	n type 1 & 2 Divi				
 Is present in almost all T1DM 					
 Is present in 50 – 80% of T2DM Diabetic Nephropathy 					
 Occurs in both type 1 & type 2 DM 					
 The earliest clinical finding of diabetic nephropathy is microa 	Ibuminuria:				
• (the persistent excretion of small amounts of albumin (30-300 mg per day) into the urine)					
 Microalbuminuria is an important predictor of progression to proteinuria: 					
 (the persistent excretion of >300 mg albumin per day into the urine) 					
• Once proteinuria appears, there is a steady \downarrow in the glomerular filtration rate (GFR)					
Finally, end-stage renal disease occurs					
Sequence of Events in Diabetic Nephropathy					
Glomerular hyperfiltration \rightarrow Microalbuminuria \rightarrow Proteinuria & \downarrow GFR \rightarrow End-stage renal disease					
Diabetic Neuropathy					
Loss of both myelinated and unmyelinated nerve fibers					
 Occurs in both type 1 & type 2 DM 					
It correlates with the duration of DM & with glycemic control					

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