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Diabetic Complications

Objectives:

- ★ Acute Diabetic Complications
 - Diabetic Ketoacidosis
 - Hyperglycemic Hyperosmolar State
 - Hypoglycemia
- ★ Chronic Diabetic Complications
 - Diabetic Retinopathy
 - Diabetic Nephropathy
 - Diabetic Neuropathy
 - Cardiovascular Disease
- ★ How to Screen and Prevent Diabetes Complications

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Glycated haemoglobin (HbA₁,)

Is a measure of an individual's average blood glucose concentration over the previous 6-8 weeks. The glycation occurs as a two-step reaction, the rate at which this reaction occurs is related to the prevailing glucose concentration. **HbA**_{1c}, is expressed as a percentage of the normal haemoglobin or as the mmol concentration of HbA_{1C} per mol of normal haemoglobin (**standardized range 4-6%**; **20-42 mmol/mol).** The result may be misleading if the lifespan of the red cell is altered.



Studies about diabetes complications

UKPDS: Type 2 diabetes complications

- A study done in multiple centers in UK from 1977 1997
- **Does intensive glucose control reduce risk of vascular complications?**
 - Is there going to be a difference in the incidence of diabetes complications if we lower A1C down to 7% versus if we keep it at 8%?

Intensive glucose therapy (lowering A1C to 7%) lowered risk of:

- Microvascular complications by 25% (after 15 years)
- Microalbuminuria by 33% after 12 years
- ♦ Any diabetes-related endpoint by 12%

There was a direct relationship between the glucose level and risk of vascular complications

- So we learned from this study that Intensive glucose control is essential in lowering the risk of diabetes complications
- Tight Blood Pressure control (144/82 mmHg) in patients with type 2 diabetes lowered the risk of:
 - Death by 32%
 - Stroke by 44%
 - ♦ Microvascular complications by 37%
 - ♦ Heart Failure by 56%
 - Retinopathy progression by 34%
 - ♦ Any diabetes-related endpoint by 24%
- ★ T2D: Start screening for complications at time of diagnosis: →They might have unrecognized diabetes for years thus when they are diagnosed they will have other complications(at least 10 years before diagnosis)
 - Yearly Dilated Eye Exam
 - Yearly Albumin: Cr ratio & Serum Creatinine¹
 - Yearly foot exam (ask the patient to examine feet, routinely)
 - o Other screening tests if clinically indicated
- ★ T1D: (because they present early) The same but start screening 5 years after the time of diagnosis

DCCT: Type 1 Diabetes & Complications

- Similar to UKPDS but in patients with T1D
- Would glucose control ameliorate the long-term complications of diabetes?





Chronic Complications of Diabetes (T1D)

1. The urine of all people with diabetes should be checked regularly for the presence of microalbuminuria. The albumin : creatinine ratio is less than 2.5mg/mmol in healthy men and less than 3.5 mg/mmol in healthy women. If microalbuminuria is detected, the test should be repeated twice because false-positive readings are common.

Diabetic Ketoacidosis



- ★ Status of <u>metabolic acidosis</u> due to absolute (or relative) insulin deficiency in association with increased levels of glucagon and other counter-regulatory hormones resulting in **increased ketone** production.
- ★ Mostly occurs in people with type 1 diabetes but may occasionally present in people with type 2 diabetes
- ★ It occurs more frequently in younger people but the mortality is higher in older people

Precipitating Causes of DKA

- \star It is usually seen in the following circumstances:
 - Previously undiagnosed diabetes
 - Non compliance with insulin therapy The most common
 - The stress of intercurrent illness and infection
- ★ Drugs: Corticosteroids, sympathomimetics, atypical anti-psychotics, SGLT-2 inhibitors

Table 1 Precipitating causes of diabetic ketoacidosis										
Precipitating cause	Australia ¹¹⁵	Brazil ¹¹⁶	China ¹¹⁷	Indonesia ¹¹⁸	Korea ¹¹⁹	Nigeria ¹²⁰	Spain ¹²¹	Syria ¹²²	Taiwan ¹²³	USA15,23
New diagnosis of diabetes mellitus, %	5.7	12.2	NR	3.3	NR	NR	12.8	NR	18.2	17.2-23.8
Infection, %	28.6	25.0	39.2	58.3	25.3	32.5	33.2	47.8	31.7	14.0-16.0
Poor adherence to treatment, %	40.0	39.0	24.0	13.3	32.7	27.5	30.7	23.5	27.7	41.0-59.6
Other, %	25.7	15.0	10.9	17.1	11.2	4.8	23.3	7.8	6.2	9.7-18.0
Unknown, %	NA	8.8	25.9	8.0	30.8	34.6	NA	20.9	16.2	3.0-4.2

Pathophysiology of DKA



1. Marked insulin deficiency is a necessary precondition for DKA since very little insulin is needed to inhibit hepatic ketogenesis and the breakdown of adipose triglycerides to non-esterified fatty acids (NEFAs).

2-The most important biochemical abnormality in is the **uncontrolled lipolysis** due to increased activity of hormone-sensitive lipase in adipose tissue and **uncontrolled ketogenesis** in the liver.

DKA



Laboratory Findings in DKA

 Hyperglycemia >250mg/dL + Hyperketonemia (or heavy ketonuria) +High anion gap (> 12 mmol\l) metabolic acidosis<18mEq/L

Other investigations:

- **ECG:** Cardiac rhythm should be monitored in severe DKA because of the risk of electrolyte-induced cardiac arrhythmia.
- Infection screen: full blood count, blood and urine culture, C-reactive protein, chest X-ray. Although leucocytosis invariably occurs in DKA, this represents a stress response and does not necessarily indicate infection.
- Blood electrolytes should be assessed as potassium abnormalities occur frequently (will be discussed in another slide)

Measure	DKA				
	Mild	Moderate	Severe		
Plasma glucose level, mmol/l	13.9	13.9	13.9		
Arterial or venous pH	7.25-7.30	7.00–7.24	<7.00		
Bicarbonate level, mmol/l	15-18	10-14	<10		
Urine or blood acetoacetate (nitroprusside reaction)	Positive	Positive	Positive		
Urine or blood β -hydroxybutyrate, mmol/l	>3	>3	>3		
Effective serum osmolality, mmol/kg*	Variable	Variable	Variable		
Anion gap, mmol/l	>10	>12	>12		
Alteration in sensorium	Alert	Alert or drowsy	Stupor or coma		

Based on the laboratory findings you can classify the cases into : mild, moderate or severe

DKA

Management of DKA

- Aggressive rehydration + Lowering glucose + Cessation of ketogenesis + Correcting electrolyte imbalances.
- Most patients with DKA are **treated in ICU**, it is associated with increased mortality.

The aim of treatment:

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- lower the blood ketone concentration by 0.5 mmol/L per hour
- increase the venous bicarbonate by 3.0 mmol/L per hour
- reduce capillary blood glucose by 3.0 mmol/L per hour (50mg/dL per hour)
- Maintaining potassium between 4.0 and 5.5mmol/L.

Rehydration

- IVF is the most critical step.
- Water deficit is ~ 100ml/kg of body weight equivalent to 7.5 litres in a 75kg adult. Considered a large deficit.
- Isotonic saline (0.9% sodium chloride) @ 500-1000 ml/hr during the 1st 2-4 h, Followed by isotonic saline 250—500 ml/h.
- Once the plasma glucose is **~250** mg/dl, **switch IVF to D5% IVF**. If the plasma sodium is greater than 155 mmol/L, 0.45% saline may be used initially.
- Hartmann's solution is an acceptable alternative, (potassium cannot added to Hartmann's solution).
- The aim of the first few litres of fluid is to correct any hypotension, replenish the intravascular deficit, and counteract the effects of the osmotic diuresis with correction of the electrolyte disturbance.
- **Over-rapid fluid** replace- may lead to **cerebral oedema**. The rate and volume of fluid replacement need to be modified in older people and in those with renal or heart failure.



- Start with normal saline and after 1-2 hours evaluate serum NA⁺
- **low**? continue with normal saline.
- **High or normal?** shift to half normal saline.
- In both cases continue IVF until the blood glucose reaches 250 mg/dL, now change to 5% dextrose with half normal saline as maintenance fluid.
 - We don't want to overcorrect hyperglycemia and cause hypoglycemia.

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Management of DKA (cont)

Insulin

- Insulin is the next step after IVF
- **Reduces** serum glucose, **suppresses** ketogenesis, and **correct** the electrolyte disturbance.
- Most of the time: we use IV insulin infusion
- Mild DKA can be treated with subcutaneous insulin (rarely)
- Most protocols: IV insulin bolus \rightarrow 0.1 unit/kg/hour
- Followed by: IV insulin infusion \rightarrow 0.1 unit/kg/h
- A rapid decrease in blood glucose should be avoided, as this might precipitate hypoglycaemia and the serious complication of cerebral oedema, particularly in children.
- Failure of blood glucose to fall within 1 hour of commencing insulin infusion should lead to a re-assessment of insulin dose.
- Should be continued until the ketosis has resolved.
- Once the person is able to eat and keep food down, subcutaneous insulin treatment should be resumed or initiated.

Electrolytes

- DKA is associated with total-body K+ deficit
- Serum **K+** is often **normal** or **high** (do not get fooled!).
- K+ Shift from intracellular to extracellular compartment with acidosis (serum K+ looks falsely normal). Metabolic acidosis causes hyperkalaemia as potassium is exchanged for hydrogen ions moving into the cell. As insulin promotes the co-transport of potassium along with glucose into cells. Although serum potassium may be elevated, there is a severe whole-body potassium deficiency as significant quantities of potassium are lost in vomit and urine.
- Insulin therapy moves K+ back into the cells (watch for a drop in K+).
 - After the initiation of treatment with insulin, potassium levels can fall rapidly. So commercially, 0.9% sodium chloride is available with premixed potassium chloride (40mmol/L (0.3%) allowing the potassium to be replaced safely.
 - We add k to the IV fluid even when serum k is normal (it should be high due to the shifting so don't miss this!) as it is not a reflection of total body k.
- K+ replacement starts early (when K+ is normal), and the rate of K infusion depends on K+ level and eGFR.
- Phosphate replacement is almost never required
- Hyperchloraemic acidosis may develop during treatment since a large variety of negatively charged electrolytes are lost in DKA, which are replaced with chloride.

Insulin		Potassium		
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IV route	SC route	If serum K ⁺ is <3.3 mmol/l hold insulin and give		
		until serum K ⁺ ≥ 3.3 mmol/l		
0.1U/kg IV bolus	0.2 U/kg SC bolus			
		+		
♦ 0.1U/kg/h IV insulin	♦ 0.2U/kg SC every	If serum K ⁺ is >5.0mmol/l do not give K ⁺ but check serum K ⁺ every 2h		
	2n	Ļ		
Check serum or capillary g every 1–2h When glucose reaches ~1 reduce insulin to 0.1U/kg to maintain glucose at 8.3 until resolution of ketoaci	If serum K ⁺ is <5.0mmol/l add 20–40mmol of KCL in each litre of IV fluid to keep serum K ⁺ at 4–5mmol/l			
Transition to SC insulin when the patient is alert and can eat Identify and treat precipitating cause				

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2 Hyperglycemic Hyperosmolar State (HHS)

- ★ Status of **severe hyperglycemia** due to insulin resistance (not absolute insulin deficiency) & relative insulin deficiency resulting in **increased serum osmolality.**
- ★ Severe hyperglycemia develops without significant ketosis, is the characteristic metabolic emergency of **uncontrolled type 2 diabetes.**
- ★ Although typically occurring in **older patients**, HHS is increasingly seen in younger adults.
- ★ People present in middle or later life, often with previously undiagnosed diabetes.
- ★ Precipitating factors include consumption of glucose-rich fluids, concurrent medication such as thiazide diuretics or steroids.
- ★ Evidence of underlying illness, such as pneumonia or pyelonephritis, may be present.

HHS

How HHS differs from DKA?

- ★ ~ 10 times higher mortality than DKA
- ★ Slower development (over several days)
- ★ No ketosis or minimal ketosis
 - The degree of insulin deficiency is less severe. Endogenous insulin levels are sufficient to inhibit hepatic ketogenesis but insufficient to inhibit hepatic glucose production.
- ★ Higher serum glucose level than DKA
- ★ More severe dehydration & higher plasma osmolality than DKA
 - Old people experience thirst less acutely and become dehydrated more readily.
 - In addition, the mild renal impairment associated with age results in increased urinary losses of fluid and electrolytes.
- ★ Gradual worsening of polydipsia, polyuria, & weight loss
- \star Impaired consciousness is more common than DKA

Pathophysiology of HHS



Laboratory Findings in HHS

Severe hyperglycemia (> 30 mmol/L (600 mg/dL)

Hyperosmolality (serum osmolality >320 mOsmol/kg)

Without significant ketonaemia (<3 mmol/L) or acidosis (pH >7.3 (H+ <50 nmol/L), bicarbonate >15 mmol/L)

Glycosuria, leading to an osmotic diuresis with loss of water, sodium, potassium and other electrolytes

Measure	DKA	HSS		
	Mild	Moderate	Severe	
Plasma glucose level, mmol/l	13.9	13.9	13.9	33.3
Arterial or venous pH	7.25–7.30	7.00–7.24	<7.00	>7.30 normal
Bicarbonate level, mmol/l	15–18	10-14	<10	>15 high
Urine or blood acetoacetate (nitroprusside reaction)	Positive	Positive	Positive	Negative or low positive
Urine or blood β -hydroxybutyrate, mmol/l	>3	>3	>3	<3
Effective serum osmolality, mmol/kg*	Variable	Variable	Variable	>320
Anion gap, mmol/l	>10	>12	>12	<12
Alteration in sensorium	Alert	Alert or drowsy	Stupor or coma	Stupor or coma

Hypovolemia

HHS

Management of HHS

- Management of HHS is similar to that of DKA
- The aims are to normalise osmolality, replace fluid and electrolyte losses, and normalise blood glucose, at the same time as preventing complications such as arterial or venous thrombosis, cerebral oedema and central pontine demyelinosis.
- Historically management of HHS has followed DKA guidelines, but increasing recognition of the differences between HHS and DKA has led to new approaches in HHS.
- The most important aspect of management is **fluid replacement**; **0.9% sodium chloride is the treatment of choice**, but <u>0.45% sodium chloride may be considered if the osmolality is</u> <u>not declining despite adequate fluid balance.</u>
- The rate of fall of plasma sodium should not exceed 10mmol/L in 24 hours because the resultant change in osmolality may cause cerebral damage.
- Fluid replacement is often enough to lower the glucose but insulin (0.05 units/kg per hour) should be used if the glucose is no longer falling with fluids alone or if the patient develops significant ketonaemia, when the diagnosis should be reconsidered.
- In order to prevent cerebral damage, the fall in blood glucose should be no more than 5 mmol/L per hour (90 mg/dL per hour)
- Prophylactic low-molecular-weight heparin should be given.

Hypoglycemia

3 Hypoglycemia

- ★ Plasma glucose <3.9 mmol/L (<70 mg/dl)</p>
- ★ Severe hypoglycemia: need for assistance from another person to correct glucose
- ★ Most frequent & serious adverse effect of glucose-lowering therapies
- ★ What is the most common adverse effect of insulin therapy ? Hypoglycemia.
- ★ Mainly due to Insulin & less frequently sulfonylureas.
- ★ Uncommon in people without diabetes but relatively frequent in people with diabetes
- ★ Hypoglycemia in a patient with diabetes is almost always due to glucose-lowering therapies
- ★ Major barrier to achieving desirable glucose control
- ★ Occurs in 30-40% of patients with T1DM
- ★ Occurs in 10-30% of patients with insulin-treated T2D

Hypoglycemia

Clinical features

- ★ Symptoms of hypoglycaemia are idiosyncratic, differing with age and duration of diabetes, and also depending on the circumstances in which hypoglycemia occurs.
- ★ They comprise two main group:
 - those related to acute activation of the autonomic nervous system
 - those secondary to glucose deprivation of the brain (neuroglycopenia).
- ★ Hypoglycemia also affects mood, inducing a state of increased tension and low energy.



Factors contributing to hypoglycaemia

- Insufficient patient education
- Medications (insulin, sulfonylureas, glinides, quinolones
- Aggressive treatment protocols targeting normoglycaemia
- Poor coordination of insulin administration and food delivery
- Abrupt changes in nutritional intake
- Abrupt discontinuation of parenteral or enteral nutrition among insulin - treated patients

Decline in renal or hepatic function

• Severe illness

- Tapering of steroid doses without appropriate reductions in insulin
- Inappropriate insulin dosing
- Counter regulatory hormone deficiencies
- Impaired awareness of hypoglycaemia
- Dementia
- Age > 65 years
- Sepsis

Treatment: (Rule of 15)

Give 15 grams of carbohydrates

- ➤ 4 glucose tablets
- > 1/2 cup of fruit juice or regular soda
- 1 tablespoon of sugar or honey

Wait 15 minutes and re-check glucose

Repeat the same if glucose is still less than 70 mg/dl

If glucose is above 70 mg/dl, have the patient eat a regular meal or a snack that **contains protein** (e.g. nuts, cheese, chicken, meat, etc) Oral carbohydrate usually suffices if hypoglycemia is recognised early. If parenteral therapy is required, then as soon as the patient is able to swallow, glucose should be given orally.



Remember, the patient **should not be driving** with hypoglycemia or (within 1 hour after treating hypoglycemia)

Chronic Diabetic Complications

- Micro-: Retinopathy, Neuropathy, and Nephropathy.
- Macro-: Ischemic Heart Disease (IHD), Cerebrovascular events, Peripheral Vascular Disease (PVD)
- Mortality
- Complications of Type 2 Diabetes:
- **Diabetes is the leading cause of:** Blindness, Renal failure, and Non-traumatic lower extremity amputation
- The presence of DM complication tremendously increases medical care cost
- Usually present after long period of hyperglycemia
- Fortunately, they can be delayed/prevented by early DM detection and better glucose control



Diabetic retinopathy

Retinopathy

Most commonly diagnosed diabetes-related complication, prevalence increases with the duration of diabetes
 Approximately 20% of people with type 1 diabetes will have retinal changes after 10 years, rising to 90% after 20 years; 20–30% of people with type 2 diabetes have retinopathy at diagnosis.

Non-proliferative (earliest change)	 Usually appears in the 1st decade of the disease or early 2nd decade. Characterized by retinal vascular microaneurysms, blot hemorrhage, and cotton-wool spots
Proliferative	 Hypoxemia & neovascularization leading to virtuous hemorrhage, fibrosis, and retinal detachment Some of these new vessels are inside the retina and give the appearance of intraretinal microvascular abnormalities (IRMAs), sometimes vessels induced to grow on the pupil margin (rubeosis) and give rise to a rapid increase in intraocular pressure (rubeotic glaucoma).
Macular edema	 can occur in non proliferative or proliferative stage



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Retinopathy

Other ways in which DM can affect the eye:

- **Cataract:** caused by the denaturation of the protein and other components of the lens of the eye, which renders it opaque.
- **Refractory defects:** result of osmotic changes within the lens (the absorption of water into the lens causes temporary hypermetropia). This presents as fluctuating difficulty in reading but people should be reassured that this resolves with better metabolic control of the diabetes.
- External ocular palsies
- Glaucoma: open angle glaucoma
- Blindness

Treatment of Diabetic Retinopathy



laser photocoagulation

- To treat new vessels of proliferative retinopathy
- Although laser treatment prevents blindness, the main adverse effects result from the destruction of retinal tissue. The visual field becomes permanently smaller and there is reduced dark adaptation. In essence, peripheral vision is sacrificed to maintain central vision.

Vitreoretinal surgery

- used if bleeding is recurrent and preventing laser therapy.
- It is also employed to try to salvage some vision if an intravitreal haemorrhage fails to clear and to treat fibrotic traction retinal detachment in advanced retinopathy.

ocular injection (intravitreal injection)

- Repeated injections of anti-VEGF drugs, such as bevacizumab, aflibercept and ranibizumab.
- therapy for macular edema, can control proliferative diabetic retinopathy and sight-threatening maculopathy.

Diabetic nephropathy

Nephropathy

- Characterized by gradually increasing urinary albumin excretion and blood pressure as the glomerular filtration rate falls insidiously towards end-stage renal disease. Glomerular hyperfiltration

 → microalbuminuria→ macroproteinuria→ progression to CKD.
- Always think about the other risk factors e.g HTN
- Patients with diabetic nephropathy, almost ALWAYS, have evidence of diabetic retinopathy
- If your patient with diabetes has nephropathy but no retinopathy; it is very likely that the nephropathy is NOT due to diabetes
- Manifests 15–25 years after the diagnosis of diabetes but affects 25–35% of people diagnosed under the age of 30 years. It is the leading cause of premature death in young people with diabetes.

Other ways in which DM can damage the kidney:

- ischaemic lesion: Arteriolar lesions, with hypertrophy and hyalinization of the vessels, can occur
- urinary tract infection: more common in women with diabetes, but not in men.

Investigations



The urine of all people with diabetes **should be checked regularly (at least annually)** for the presence of microalbuminuria.

- ★ Albuminuria (Albumin: Cr >30 mg/g)
- ★ Once proteinuria is present, other possible causes should be considered, but once these are excluded, a presumptive diagnosis of diabetic nephropathy can be made.
- ★ Clinical suspicion of a non- diabetic cause of nephropathy may be provoked by an atypical history, the absence of diabetic retinopathy and the presence of red-cell casts in the urine.
- ★ **Renal biopsy (Kimmelstiel-Wilson nodules)** should be considered in such cases but is rarely necessary or helpful.
 - ★ Plasma creatinine level and eGFR should be measured regularly.

Nephropathy

Treatment of Diabetic Nephropathy

Prevention (most effective treatment)

Aim is to slow the disease progression (or reverse it)

Glucose & BP (target BP <130/80 mmHg) control is key.

ACEI (or ARBs) are recommended to treat nephropathy.

SGLT-2 inhibitors can be used, decrease the risk and progression of diabetic nephropathy.

Remember to change doses (or stop) medications that are renally cleared if eGFR is low Oral antidiabetes agents, partially excreted via the kidney (e.g. glibenclamide and metformin), should be avoided. As insulin clearance is reduced in advanced renal disease, insulin dosage is usually reduced.

ESRD management: chronic ambulatory peritoneal dialysis may be preferable to haemodialysis, The failure rate of renal transplants is somewhat higher than in people without diabetes.

Diabetic neuropathy Neuropathy Treat with preventive foot care Most common form is distal symmetric polyneuropathy • Usually affects sensory nerves in a "stocking/glove pattern"—Typically begins in feet, later involves • hands (longest nerves affected first). Polyneuropathy Tingling, numbness . loss of sensation leads to the following: ulcer formation (patients do not shift their weight) with subsequent ischemia of pressure point areas; Charcot joints. Loss of fine touch, proprioception, and vibration. Loss of ankle deep reflex **Painful diabetic neuropathy**—hypersensitivity to light touch; severe "burning" pain, especially at night, that can be difficult to tolerate. Treatment is with gabapentin, tricyclic antidepressants, or pregabalin. Dysfunction of cranial or peripheral nerves . less common Mononeuropathies Most often involves **CN III**, but may also involve CN VI and IV. **Diabetic third nerve palsy:** eye pain, diplopia, ptosis, inability to adduct the eye; but the pupils are spared. Median nerve neuropathy, ulnar neuropathy, common peroneal neuropathy. Diabetic lumbosacral plexopathy—severe, deep pain in the thigh; atrophy and weakness in thigh and hip muscles; recovery takes weeks to months. Diabetic truncal neuropathy—pain in distribution of one of the intercostal nerves. • **Impotence in men** (most common presentation) neuropathy Neurogenic bladder-retention, incontinence Autonomic Gastroparesis—chronic nausea and vomiting, early satiety. Give metoclopramide or erythromycin Constipation and diarrhea (alternating) Postural hypotension

Summary

Hyperglycemic hyperosmolar

Hypoglyc

Retinonathv

Nephropathy

emia

state

Pathogenesis: Status of <u>metabolic acidosis</u> due to absolute (or relative) insulin deficiency in association with increased levels of glucagon and other counter-regulatory hormones resulting in **increased ketone** production.

Clinical features: polyuria , polydipsia , nausea , vomiting, weight loss, hypothermia, change in mental status, dehydration, Kussmaul respiration

Investigations: Hyperglycemia >250mg/dL + Hyperketonemia (or heavy ketonuria) +High anion gap (> 12 mmol\l) metabolic acidosis<18mEq/L

Management: Aggressive rehydration + Lowering glucose + Cessation of ketogenesis + Correcting electrolyte imbalances.

Pathogenesis: Status of **severe hyperglycemia** due to insulin resistance (not absolute insulin deficiency) & relative insulin deficiency resulting in **increased serum osmolality. without ketone body production**

Clinical features: Gradual worsening of polydipsia, polyuria, & weight loss , impaired consciousness

Investigation: Severe hyperglycemia (> 30 mmol/L (600 mg/dL), Hyperosmolality (serum osmolality >320 mOsmol/kg) Without significant ketonaemia (<3 mmol/L) or acidosis (pH >7.3 (H+ <50 nmol/L), bicarbonate >15 mmol/L), Glycosuria, osmotic diuresis with loss of water, sodium, potassium and other electrolytes, Hypovolemia Management: Management of HHS is similar to that of DKA

Pathogenesis: Plasma glucose <3.9 mmol/L (<70 mg/dl) , Mainly due to Insulin & less frequently sulfonylureas.</th>Clinical features: sweating , hunger, anxiety, headache, nausea, tiredness, speech difficulty, deliriumManagement: (Rule of 15) Give 15 grams of carbohydrates and Wait 15 minutes and re-check glucose . Repeat the same if glucose is still less than 70mg/dL

	Non-proliferative	 Characterized by retinal vascular microaneurysms, blot hemorrhage, and cotton-wool spots
	Proliferative	• Hypoxemia & neovascularization leading to virtuous hemorrhage, fibrosis, and retinal detachment
	Macular edema	• can occur in non proliferative or proliferative stage

Pathogenesis: Characterized by gradually increasing urinary albumin excretion and blood pressure as the glomerular filtration rate falls insidiously towards end-stage renal disease

Investigation:

- Screen with Urinary Albumin: Creatinine & eGFR Albuminuria (Albumin: Cr >30 mg/g)
- Renal biopsy (Kimmelstiel-Wilson nodules) should be considered in such cases but is rarely necessary or helpful.
- Plasma creatinine level and eGFR should be measured regularly.

Management: ACEI (or ARBs) are recommended to treat nephropathy ,SGLT-2 inhibitors can be used

Polyneuropa thy	 Most common form is distal symmetric polyneuropathy Tingling, numbness, loss of sensation leads to the following: ulcer formation (patients do not shift their weight) with subsequent ischemia of pressure point areas; Charcot joints. Loss of fine touch, proprioception, and vibration. Loss of ankle deep reflex
Mononeuropathies	 Dysfunction of cranial or peripheral nerves Most often involves CN III, but may also involve CN VI and IV. Diabetic third nerve palsy: eye pain, diplopia, ptosis, inability to adduct the eye; but the pupils are spared. Median nerve neuropathy, ulnar neuropathy, common peroneal neuropathy. Diabetic lumbosacral plexopathy—severe, deep pain in the thigh; atrophy and weakness in thigh and hip muscles; recovery takes weeks to months. Diabetic truncal neuropathy—pain in distribution of one of the intercostal nerves.
Autoimmune neuropathy	 Impotence in men (most common presentation) Neurogenic bladder—retention, incontinence Gastroparesis—chronic nausea and vomiting, early satiety. Give metoclopramide Constipation and diarrhea (alternating) Postural hypotension

Lecture Quiz

Q1: A 29-year-old woman is found unconscious by her partner and rushed to accident and emergency. She is a type 1 diabetic and has maintained excellent glucose control using insulin injections. Blood biochemistry results demonstrate a moderately raised level of insulin, no detectable C-peptide and very low blood glucose. Her partner mentions she is a lawyer and has been working particularly hard in the last week, eating quick meals and occasionally missing meals. The most likely diagnosis is:

- A. Hyperosmolar coma
- B. Diabetic ketoacidosis
- C. Insulin overdose
- D. Hypoglycaemic coma

Q2: A 55-year-old diabetic woman presents with altered sensations in her hands and feet. She finds it difficult to turn pages of books and discriminating between different coins. When walking, the floor feels different and she likens the sensation to walking on cotton wool. The most likely diagnosis is:

- A. Autonomic neuropathy
- B. Diabetic amyotrophy
- C. Acute painful neuropathy
- D. Symmetrical sensory neuropathy
- E. Diabetic mononeuropathy

Q3:A 49-year-old woman presents to her physician's office with a long-standing history of polydipsia, polyuria, central obesity, and hyperlipidemia. She is currently taking metformin, a sulfonylurea, and an angiotensin-converting enzyme (ACE) inhibitor. ACE inhibitors are most beneficial in preventing or slowing the progression of which of the following diabetic complications?

- A. Diabetic ketoacidosis
- B. Diabetic nephropathy
- C. Diabetic neuropathy
- D. Diabetic retinopathy
- E. Peripheral vascular disease

Q4: What is the most common adverse event of insulin in type 1 diabetes?

A. HypoglycemiaB. LipohypertrophyC.Skin allergyD.Anxiety or depression

Answers: Q1:D | Q2:D | Q3:B | Q4:A |





Females co-leaders:

Raghad AlKhashan Amirah Aldakhilallah Males co-leaders: Mashal AbaAlkhail Nawaf Albhijan

Send us your feedback: We are all ears!

