

## Foundations Frameworks Approach to Hyperglycemia and Insulin

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**Diabetes** is the most common cause of hyperglycemia. There are four major types:

- 1. **Type 1 Diabetes Mellitus:** This results from absent or diminished insulin production, usually due to autoimmune destruction of pancreatic beta cells. These patients require insulin supplementation
- 2. **Type 2 Diabetes Mellitus:** These patients have insulin resistance resulting from decreased receptor production or receptor defects. There are a wide variety of therapies for this, including insulin, oral hypoglycemics, and metformin
- 3. **Gestational Diabetes:** This is pathophysiologically like DM2 but resolves in the post-partum period
- 4. **Secondary Diabetes:** This may be like Type 1 or Type 2 DM but is due to other causes (e.g. pancreatitis-induced beta cell destruction, medication-induced insulin resistance, etc.). Exogenous steroids may also cause hyperglycemia and mimic diabetes

**Hyperglycemia** by itself is not usually dangerous and can be treated with IV fluids or supplemental insulin. Acute complications of hyperglycemia, however, need to be ruled out

- **New-onset DM2:** In the asymptomatic but hyperglycemic patient without a formal diagnosis of DM, workup and initiation of oral medications is appropriate
  - o Labs: CBC, chem, UA, HgbA1C, LFTs, lipase
  - <u>Diagnosis</u>: HgbA1c > 6.5%
  - o <u>Meds</u>: Start metformin 500 mg q12 hr or 850 mg qD
  - *Disposition*: Usually home, with PCP follow-up within 1 week
- Hyperglycemia in DM: In the hyperglycemic patient with a known diagnosis of DM, adverse consequences such as DKA and HHS (below) must be ruled out. This will often have a precipitant as well, most frequently med non-compliance or infection, but consider cardiac causes or other systemic stressors
  - *Labs*: VBG, CBC, chem, LA, workup for infectious precipitant (e.g. UA, CXR), consider troponin, BNP, etc.
  - o *Diagnosis*: Hyperglycemia without other metabolic abnormalities
  - <u>Meds</u>: IV normal saline, insulin per pt's own sliding scale regimen
  - *Disposition*: Usually home, with PCP follow-up
- **Diabetic Ketoacidosis:** Mainly a complication of DM1, this does exist to a lesser degree in DM2 as well
  - Labs: VBG, CBC, chem, Mg, UA, LA, acetone (if available), workup for infectious precipitant (e.g. CXR)
  - <u>Diagnosis</u>: Hyperglycemia with metabolic anion gap acidosis and ketosis make sure to apply Winter's formula to detect concomitant respiratory/metabolic alkaloses. Rarely, DKA may occur with euglycemia, but acidemia and ketosis must be present to make this diagnosis
  - <u>Meds</u>: IV normal saline/LR bolus (corrects dehydration from hyperglycemia-driven osmotic diuresis), replete K as needed, then start insulin

gtt (DKA results from insulin deficiency, with the ketosis being driven by unopposed glucagon)

- Once the patient's glucose hits 250, change IVF to D5NS and titrate insulin for goal glucose 150-200 until acidosis/anion gap resolves
- Check the K every hour and replete when it gets to 4.0
- *Disposition*: Generally the ICU
- Hyperglycemic Hyperosmolar State: Mainly a complication of DM2 (sometimes concurrently with DKA) and *usually involves mental status changes* due to fluid shifts
  - <u>Labs</u>: VBG, serum/urine osms, CBC, chem, LA, UA, workup for precipitant
    - Note that electrolytes will need to be monitored here too, but less aggressively than for DKA, as you don't need to give insulin (see below)
  - *Diagnosis*: Marked hyperglycemia **without ketosis** (but acidemia may be present if an acidemic state, e.g. sepsis, is the precipitant), longer prodrome than DKA
  - <u>Meds</u>: Copious IV fluids (fixes profound dehydration from hyperglycemia-driven osmotic diuresis), can consider insulin but this is not essential as insulin is still secreted in HHS
  - <u>Disposition</u>: ICU

## **References:**

- Maloney G and Glauser J. *Rosen's Emergency Medicine: Concepts and Clinical Practice*, 9e. Chapter 118: "Diabetes Mellitus and Disorders of Glucose Homeostasis." Elsevier 2018, pp. 1533-1547.
- Evans M, Schumm-Draeger PM, Vora J, King AB. A review of modern insulin analogue pharmacokinetic and pharmacodynamic profiles in type 2 diabetes: improvements and limitations. *Diabetes Obes Metab* 2011; 13:677.≥