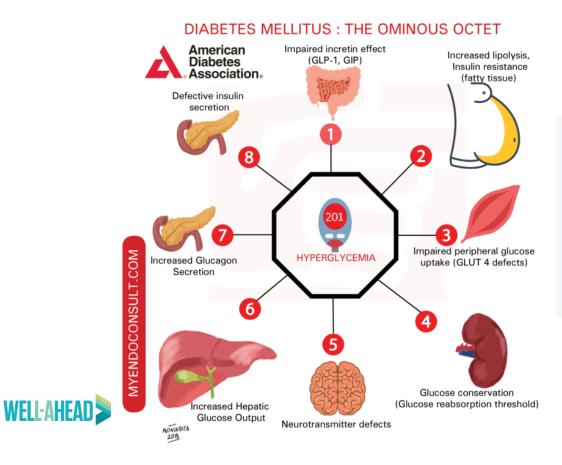
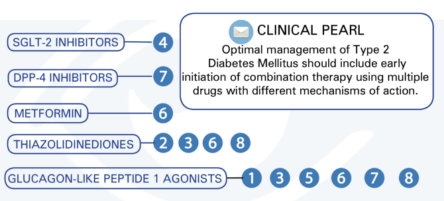
Pharmacologic Therapy for Type 2 Diabetes





Ominous Octet

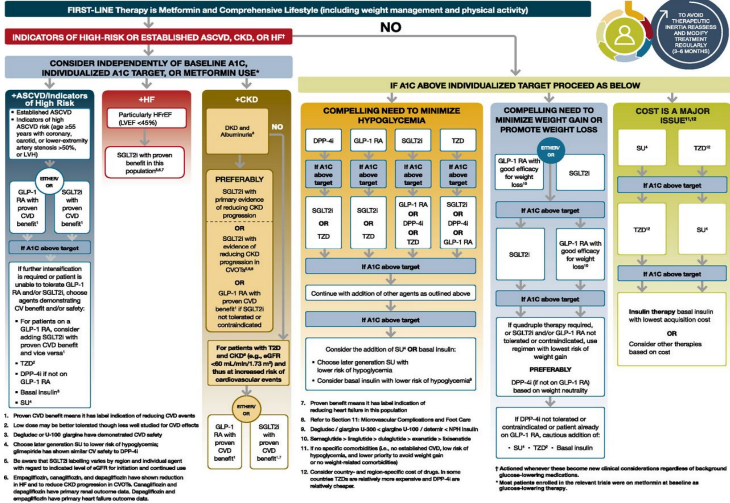




Novel Agents for the Treatment of Type 2 Diabetes. Diabetes Spectr. 2014 May; 27(2): 100-112.

Project
(ECHO [®])

Treatment Algorithm – 2021 ADA Standards







Pharmacologic Considerations

WELL-AHEAD

		Efficacy	Hypoglycemia	Weight change	CV effects		Cost	Oral/SQ	Renal effects		Additional considerations
					ASCVD	HF	con		Progression of DKD	Dosing/use considerations*	
Metformir	1	High	No	Neutral (potential for modest loss)	Potential benefit	Neutral	Low	Oral	Neutral	Contraindicated with eGFR <30 mL/min/1.73 m ²	 Gastrointestinal side effects common (diarrhea, nausea) Potential for B12 deficiency
SGLT-2 inh	ibitors	Intermediate	No	Loss	Benefit: empagliflozin†, canagliflozin	Benefit: empagliflozin†, canagliflozin, dapagliflozin‡	High	Oral	Benefit: canagliflozin§, empagliflozin, dapagliflozin	 Renal dose adjustment required (canagilflozin, dapagliflozin, empagliflozin, ertugliflozin) 	Should be discontinued before any scheduled surgery to avoid potential risk for DKA DKA risk (all agents, rare in T2D) Risk of bone fractures (canagliflozin) Genitourinary infections Risk of volume depletion, hypotension TLDL cholesterol Risk of Fournier's gangrene
GLP-1 RAs		High	No	Loss	Neutral: exenatide once weekly, lixisenatide Benefit: dulaglutidet, liraglutidet, semaglutidet	Neutral	High	SQ; oral (semaglutide)	Benefit on renal end points in CVOTs, driven by albuminuria outcomes: liraglutide, semaglutide, dulaglutide	Exenatide, lixisenatide: avoid for eGFR <30 mL/min1.73 m ² No dose adjustment for dulaglutide, liraglutide, semaglutide Caution when initiating or increasing dose due to potential risk of nausea, vomiting, diarrhea, or dehydration. Monitor renal function in patients reporting severe adverse GI reactions when initiating or increasing dose of therapy.	FDA Black Box: Risk of thyroid C-cell tumors in rodents; human relevance not determined (liraguluida, albiglutida, dulaglutida, exenatida extended release, semaglutida) Gi sida effects common (nausea, vomiting, diarrhea) Injection site reactions Pancreatitis has been reported in clinical trials but causality has not been established. Discontinue if pancreatitis is suspected.
DPP-4 inhi	ibitors	Intermediate	No	Neutral	Neutral	Potential risk: saxagliptin	High	Oral	Neutral	Renal dose adjustment required (sitagliptin, saxagliptin, alogliptin); can be used in renal impairment No dose adjustment required for linagliptin	Pancreatitis has been reported in clinica trials but causality has not been established. Discontinue if pancreatitis is suspected. Joint pain
Thiazolidinediones		High	No	Gain	Potential benefit: pioglitazone	Increased risk	Low	Oral	Neutral	No dose adjustment required Generally not recommended in renal impairment due to potential for fluid retention	FDA Black Box: Congestive heart failure (pioglitazone, rosiglitazone) Fluid retention (edema; heart failure) Benefit in NASH Risk of bone fractures Bladder cance (pioglitazone) TLDL cholesterol (rosiglitazone)
Sulfonylureas (2nd generation)		High	Yes	Gain	Neutral	Neutral	Low	Oral	Neutral	 Glyburide: not recommended Glipizide and glimepiride: initiate conservatively to avoid hypoglycemia 	 FDA Special Warning on increased risk of cardiovascular mortality based on studies of an older sulfonylurea (tolbutamide)
ins	Human insulin	Highest	fighest Yes	Gain	Neutral	Neutral	Low (SQ)	SQ; inhaled	Neutral	 Lower insulin doses required with a decrease in eGFR; titrate 	Injection site reactions Higher risk of hypoglycemia with human insulin (NPH or premixed formulations) vs. analogs
	Analogs						High	SQ	pe	per clinical response	



Other Considerations

- Available Combinations

 Oral
 Insulin and GLP-1 agonists
- Cost
- Optimizing therapy
- Side effects and counseling

 Metformin Diarrhea
 GLP1 Agonist Nausea
- Adherence
- Adding and Removing Therapy



