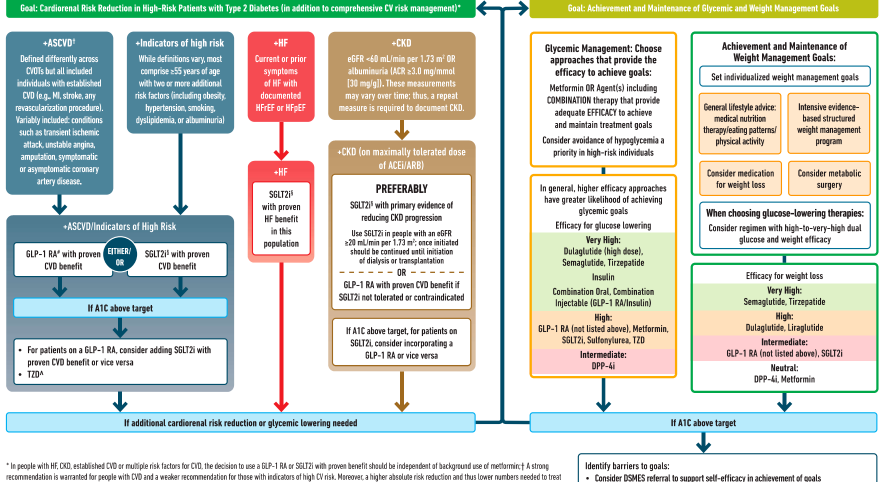
## USE OF GLUCOSE-LOWERING MEDICATIONS IN THE MANAGEMENT OF TYPE 2 DIABETES

## HEALTHY LIFESTYLE BEHAVIORS; DIABETES SELF-MANAGEMENT EDUCATION AND SUPPORT (DSMES); SOCIAL DETERMINANTS OF HEALTH (SDOH)





are seen at higher levels of baseline risk and should be factored into the shared decision-making process. See text for details; ^ Low-dose TZD may be better tolerated and similarly effective; & For SGLT2i, CV/ renal outcomes trials demonstrate their efficacy in reducing the risk of composite MACE, CV death, all-cause mortality, MI, HHF, and renal outcomes in individuals with T2D with established/high risk of CVD; # For GLP-1 RA, CVOTs demonstrate their efficacy in reducing composite MACE, CV death, all-cause mortality, MI, stroke, and renal endpoints in individuals with T2D with established/high risk of CVD.

- Consider technology (e.g., diagnostic CGM) to identify therapeutic gaps and tailor therapy
- Identify and address SDOH that impact achievement of goals

Figure 9.3—Use of glucose-lowering medications in the management of type 2 diabetes. ACEi, angiotensin-converting enzyme inhibitor; ACR, albumin-to-creatinine ratio; ARB, angiotensin receptor blocker; ASCVD, atherosclerotic cardiovascular disease; CGM, continuous glucose monitoring; CKD, chronic kidney disease; CV, cardiovascular; CVD, cardiovascular disease; CVOT, cardiovascular outcomes trial; DPP-4i, dipeptidyl peptidase 4 inhibitor; eGFR, estimated glomerular filtration rate; GLP-1 RA, glucagon-like peptide 1 receptor agonist; HF, heart failure; HFpEF, heart failure with preserved ejection fraction; HFrEF, heart failure with reduced ejection fraction; HHF, hospitalization for heart failure; MACE, major adverse cardiovascular events; MI, myocardial infarction; SDOH, social determinants of health; SGLT2i, sodium-glucose cotransporter 2 inhibitor; T2D, type 2 diabetes; TZD, thiazolidinedione, Adapted from Davies et al. (45).

Table 9.2—Medications for lowering glucose, summary of characteristics

		Efficacy <sup>1</sup> Hypogly- cemia	Hypogly-	. Weight change <sup>2</sup>	CV effects		Renal effects				
					Effect on MACE	HF	Progression of DKD	Dosing/use considerations*	Oral/SQ	Cost	Clinical considerations
Metformin		High	No	Neutral (potential for modest loss)	Potential benefit	Neutral	Neutral	Contraindicated with eGFR <30 mL/min per 1.73 m <sup>2</sup>	Oral	Low	Gl side effects common; to mitigate Gl side effects, consider slow dose titration, extended release formulations, and administration with food Potential for vitamin B12 deficiency; monitor at regular intervals
SGLT2 inhibitors		Intermediate to high	No	Loss (intermediate)	Benefit: canagliflozin, empagliflozin	Benefit: canagliflozin, dapagliflozin, empagliflozin, ertugliflozin	Benefit: canagliflozin, dapagliflozin, empagliflozin	See labels for renal dose considerations of individual agents     Glucose-lowering effect is lower for SGLT2 inhibitors at lower eGFR	Oral	High	DKA risk, rare in T2DM: discontinue, evaluate, and treat promptly if suspected; be aware of predisposing risk factors and clinical presentation (including euglycemic DKA); discontinue before scheduled surgery (e.g., 3–4 days), during critical illness, or during prolonged fasting to mitigate potential risk Increased risk of genital mycotic infections Necrotizing fascitits of the perineum (Fournier gangrene), rare reports: institute prompt treatment if suspected Attention to volume status, blood pressure; adjust other volume-contracting agents as applicable
GLP-1 RAs		High to very high	No	Loss (intermediate to very high)	Benefit: dulaglutide, liraglutide, semaglutide (SQ)  Neutral: exenatide once weekly, lixisenatide	Neutral	Benefit for renal endpoints in CVOTs, driven by albuminuria outcomes: dulaglutide, lirglutide, semaglutide (SQ)	See labels for renal dose considerations of individual agents     No dose adjustment for dulaglutide, liraglutide, semaglutide     Monitor renal function when initiating or escalating doses in patients with renal impairment reporting severe adverse GI reactions	SQ; oral (semaglutide)	High	Risk of thyroid C-cell tumors in rodents; human relevance not determined (liraglutide, dulaglutide, exenatide extended release, semaglutide) Counsel patients on potential for GI side effects and their typically temporary nature; provide guidance on dietary modifications to mitigate GI side effects (reduction in meal size, mindful eating practices [e.g., stop eating once full], decreasing intake of high-fat or spicy food); consider slower dose titration for patients experiencing GI challenges Pancreatitis has been reported in clinical trials but causality has not been established. Discontinue if pancreatitis is suspected Evaluate for gallbladder disease if cholelithiasis or cholecystitis is suspected
GIP and GLP-1 RA		Very high	No	Loss (very high)	Under investigation	Under investigation	Under investigation	See label for renal dose considerations     No dose adjustment     Monitor renal function when initiating or escalating doses in patients with renal impairment reporting severe adverse GI reactions	SQ	High	Risk of thyroid C-cell tumors in rodents; human relevance not determined  Counsel patients on potential for GI side effects and their typically temporary nature; provide guidance on dietary modifications to mitigate GI side effects (reduction in meal size, mindful eating practices [e.g., stop eating once full), decreasing intake of high-fat or spicy food); consider slower dose titration for patients experiencing GI challenges  Pancreatitis has been reported in clinical trials but causality has not been established. Discontinue if pancreatitis is suspected  Evaluate for gallbladder disease if cholelithiasis or cholecystitis is suspected
DPP-4 inhibitors		Intermediate	No	Neutral	Neutral	Neutral (potential risk, saxagliptin)	Neutral	Renal dose adjustment required (sitagliptin, saxagliptin, alogliptin); can be used in renal impairment     No dose adjustment required for linagliptin	Oral	High	Pancreatitis has been reported in clinical trials but causality has not been established. Discontinue if pancreatitis is suspected     Joint pain     Bullous pemphigoid (postmarketing): discontinue if suspected
Thiazolidinediones		High	No	Gain	Potential benefit: pioglitazone	Increased risk	Neutral	No dose adjustment required     Generally not recommended in renal impairment due to potential for fluid retention	Oral	Low	Congestive HF (pioglitazone, rosiglitazone) Fluid retention (edema; heart failure) Benefit in NASH Risk of bone fractures Weight gain: consider lower doses to mitigate weight gain and edema
Sulfonylureas (2nd generation)		High	Yes	Gain	Neutral	Neutral	Neutral	Glyburide: generally not recommended in chronic kidney disease     Glipizide and glimepiride: initiate conservatively to avoid hypoglycemia	Oral	Low	FDA Special Warning on increased risk of CV mortality based on studies of an older sulfonylurea (tolbutamide); glimepiride shown to be CV safe (see text)     Use with caution in persons at risk for hypoglycemia
Insulin	Human Analogs	High to very high	Yes	Gain	Neutral	Neutral	Neutral	Lower insulin doses required with a decrease in eGFR; titrate per clinical response	SQ; inhaled	Low (SQ) High	Injection site reactions     Higher risk of hypoglycemia with human insulin (NPH or premixed formulations) vs. analogs

CV, cardiovascular; CVOT, cardiovascular outcomes trial; DKA, diabetic ketoacidosis; DKD, diabetic kidney disease; DPP-4, dipeptidyl peptidase 4; eGFR, estimated glomerular filtration rate; FDA, U.S. Food and Drug Administration; GI, gastrointestinal; GIP, gastric inhibitory polypeptide; GLP-1 RA, glucagon-like peptide 1 receptor agonist; HF, heart failure; NASH, nonalcoholic steatohepatitis; MACE, major adverse cardiovascular events; SGLT2, sodium–glucose cotransporter 2; SQ, subcutaneous; T2DM, type 2 diabetes mellitus. \*For agent-specific dosing recommendations, please refer to manufacturers' prescribing information. <sup>1</sup>Tsapas et al. (62). <sup>2</sup>Tsapas et al. (114). Reprinted from Davies et al. (45).