

Sodium-glucose Co-transporter (SGLT) Inhibitors and Combinations Step Therapy with Quantity Limit Program Summary

Florida Blue/Truli does not accept the use of samples to satisfy prerequisite requirements in any of the step therapy or prior authorization programs. Samples would also include any type of prescription or copay assistance rendered to the patient by the drug manufacturer or any other organization that would allow the patient to get the drug without a paid claim through billing Florida Blue/Truli.

For a complete list of Florida Blue step therapy programs, please refer to the following link: http://www.bcbsfl.com/DocumentLibrary/Providers/Content/Rx_ResponsibleSteps.pdf

For a complete list of Truli step therapy programs, please refer to the following link: https://www.myprime.com/en/forms/coverage-determination/step-therapy.html

POLICY REVIEW CYCLE

Effective Date 04-01-2025

Date of Origin

FDA LABELED INDICATIONS AND DOSAGE

Agent(s)	FDA Indication(s)	Notes	Ref#
Brenzavvy®, Bexagliflozin	An adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus		19
Tablet	Limitation of Use: Not recommended in patients with type 1 diabetes mellitus. May increase the risk of diabetic ketoacidosis in these patients.		
Farxiga®, Dapagliflozin	As an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus		2
Tablet	To reduce the risk of hospitalization for heart failure in adults with type 2 diabetes mellitus and established cardiovascular disease or multiple cardiovascular risk factor		
	To reduce the risk of hospitalization for heart failure in adults with type 2 diabetes mellitus and either established cardiovascular disease or multiple cardiovascular risk factors.		
	To reduce the risk of sustained eGFR decline, end stage kidney disease, cardiovascular death, and hospitalization for heart failure in adults with chronic kidney disease at risk of progression		
	Limitations of Use:		
	 Not recommended for use to improve glycemic control in patients with type 1 diabetes mellitus. Not recommended for use to improve glycemic control in adults with type 2 diabetes mellitus with an eGFR less than 45 mL/min/1.73 m^2. Farxiga is likely to be ineffective in this setting based upon its mechanism of action Not recommended for the treatment of chronic kidney disease in patients with polycystic kidney disease or patients requiring or with a recent history of immunosuppressive therapy for the 		

Agent(s)	FDA Indication(s)	Notes	Ref#
	treatment of kidney disease. Farxiga is not expected to be effective in these populations.		
Glyxambi®	To improve glycemic control in adults with type 2 diabetes mellitus.	DPP-4 Inhibitor Combinations	14
(empagliflozin /linagliptin)	To reduce the risk of cardiovascular death in adults with type 2 diabetes mellitus and established cardiovascular disease.		
Tablet	Limitations of Use:		
	 Not recommended in patients with type 1 diabetes mellitus. It may increase the risk of diabetic ketoacidosis in these patients. Has not been studied in patients with a history of pancreatitis. Not recommended for use to improve glycemic control in adults with type 2 diabetes mellitus with an eGFR less than 30 mL/min/1.73 m^2. 		
Inpefa®	To reduce the risk of cardiovascular death, hospitalization for heart failure, and urgent heart failure visit in adults with:		18
(sotagliflozin) Tablet	 heart failure or type 2 diabetes mellitus, chronic kidney disease, and other 		
. 43.00	cardiovascular risk factors		
Invokamet®	As an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus		3
(canagliflozin/ metformin) Tablet	Canagliflozin is indicated to reduce the risk of major adverse cardiovascular events in adults with type 2 diabetes mellitus and established cardiovascular disease		
	Canagliflozin is indicated to reduce the risk of end-stage kidney disease, doubling of serum creatinine, cardiovascular death, and hospitalization for heart failure in adults with type 2 diabetes mellitus and diabetic nephropathy with albuminuria		
	Limitations of Use: Not recommended for use to improve glycemic control in patients with type 1 diabetes mellitus		
Invokamet® XR	As an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus		3
(canagliflozin/ metformin ER)	Canagliflozin is indicated to reduce the risk of major adverse cardiovascular events in adults with type 2 diabetes and established cardiovascular disease		
Tablet	Canagliflozin is indicated to reduce the risk of end-stage kidney disease, doubling of serum creatinine, cardiovascular death, and hospitalization for heart failure in adults with type 2 diabetes mellitus and diabetic nephropathy with albuminuria		
	Limitation of Use: Not recommended in patients with type 1 diabetes mellitus or diabetic ketoacidosis		
Invokana®	As an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus		1

Agent(s)	FDA Indication(s)	Notes	Ref#
(canagliflozin) Tablet	To reduce the risk of major cardiovascular events in adults with type 2 diabetes mellitus and established cardiovascular disease To reduce the risk of end-stage kidney disease, doubling of serum creatinine, cardiovascular death, and hospitalization for heart failure in adults with type 2 diabetes mellitus and diabetic nephropathy with albuminuria		
	Limitations of Use:		
	Not recommended in patients with type 1 diabetes mellitus		
	 Not recommended for use to improve glycemic control in adults with type 2 diabetes mellitus with an eGFR less than 30 mL/min/1.73 m^2 		
Jardiance®	To reduce the risk of cardiovascular death and hospitalization for heart failure in adults with heart failure.		4
(empagliflozin) Tablet	To reduce the risk of sustained decline in eGFR, end-stage kidney disease, cardiovascular death, and hospitalization in adults with chronic kidney disease at risk of progression.		
	To reduce the risk of cardiovascular death in adults with type 2 diabetes mellitus and established cardiovascular disease.		
	As an adjunct to diet and exercise to improve glycemic control in adults and pediatric patients aged 10 years and older with type 2 diabetes mellitus.		
	Limitations of Use:		
	 Not recommended for use to improve glycemic control in patients with type 1 diabetes mellitus. It may increase the risk of diabetic ketoacidosis in these patients. Not recommended for use to improve glycemic control in patients with type 2 diabetes mellitus with an eGFR less than 30 mL/min/1.73 m^2. Not recommended for the treatment of chronic kidney disease in patients with polycystic kidney disease or patients requiring or with a recent history of intravenous immunosuppressive therapy or greater than 45 mg of prednisone or equivalent for kidney disease. 		
Qtern®	Indicated as an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus	DPP-4 Inhibitor Combinations	15
(dapagliflozin/ saxagliptin)	Limitation of use: Not recommended for patients with type 1 diabetes mellitus.		
Tablet			
Segluromet®	As an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus		9
(ertugliflozin/ metformin)	Limitation of use: Not for the treatment of type 1 diabetes mellitus or diabetic ketoacidosis		
Tablet			

Agent(s)	FDA Indication(s)	Notes	Ref#
Steglatro®	As an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus		8
(ertugliflozin) Tablet	Limitation of Use: Not recommended in patients with type 1 diabetes mellitus		
Steglujan®	Indicated as an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus	DPP-4 Inhibitor Combinations	16
(ertugliflozin/s itagliptin)	1		
Tablet	Not recommended for patients with type 1 diabetes mellitus		
	Has not been studied in patients with a history of pancreatitis.		
Synjardy® (empagliflozin	As an adjunct to diet and exercise to improve glycemic control in adults and pediatric patients aged 10 years and older with type 2 diabetes mellitus		6
/metformin)	Empagliflozin, when used as a component of Synjardy or Synjardy XR,		
Tablet	is indicated in adults with type 2 diabetes mellitus to reduce the risk of:		
	Cardiovascular death in adults with established cardiovascular disease.		
	 Cardiovascular death and hospitalization for heart failure in adults with heart failure. 		
	Limitations of Use:		
	Not recommended for use in patients with type 1 diabetes mellitus. It may increase the risk of diabetic ketoacidosis in these patients.		
	Because of the metformin component, Synjardy and Synjardy XR are not recommended for use in patients with heart failure without type 2 diabetes mellitus.		
Synjardy® XR	As an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus		7
(empagliflozin /metformin)	Empagliflozin, when used as a component of Synjardy or Synjardy XR, is indicated in adults with type 2 diabetes mellitus to reduce the risk		
Tablet	of:		
	 Cardiovascular death in adults with established cardiovascular disease. Cardiovascular death and hospitalization for heart failure in 		
	adults with heart failure.		
	Limitations of Use:		
	 Not recommended for use in patients with type 1 diabetes mellitus. It may increase the risk of diabetic ketoacidosis in these patients. 		

Agent(s)	FDA Indication(s)	Notes	Ref#
	 Because of the metformin component, Synjardy and Synjardy XR are not recommended for use in patients with heart failure without type 2 diabetes mellitus. 		
Trijardy® XR (empagliflozin /linaglipin/me tformin) Tablet	Indicated as an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus Empagliflozin is indicated to reduce the risk of cardiovascular death in adults with type 2 diabetes mellitus and established cardiovascular disease Limitations of Use: Not recommended in patients with type 1 diabetes. Has not been studied in patients with a history of pancreatitis	DPP-4 Inhibitor Combinations	17
Xigduo® XR, Dapagliflozin/ metformin Tablet	As an adjunct to diet and exercise to improve glycemic control in adults and pediatric patients aged 10 years and older with type 2 diabetes mellitus. Dapafliflozin is indicated to reduce: • The risk of hospitalization for heart failure in adults with type 2 diabetes mellitus and established cardiovascular disease (CVD) or multiple cardiovascular (CV) risk factors. • The risk of cardiovascular death and hospitalization for heart failure in adults with heart failure with reduced ejection fraction (NYHA class II through IV). • The risk of sustained eGFR decline, end stage kidney disease, cardiovascular death, and hospitalization for heart failure in adults with chronic kidney disease at risk of progression. Limitations of Use: • Not recommended for use to improve glycemic control in patients with type 1 diabetes mellitus. • Because of the metformin HCl component, the use of Xigduo XR is limited to patients with type 2 diabetes mellitus for all indications. • Not recommended for the treatment of chronic kidney disease in patients with polycystic kidney disease or patients requiring or with a recent history of immunosuppressive therapy for the treatment of kidney disease. Xigduo XR is not expected to be effective in these populations.		5

See package insert for FDA prescribing information: https://dailymed.nlm.nih.gov/dailymed/index.cfm

CLINICAL RATIONALE

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Overview	The American Diabetes Association (ADA) recommends the following guidelines:(10,11)
	 Healthy lifestyle behaviors, diabetes self-management education and support, avoidance of clinical inertia, and social determinants of health should be considered in the glucose-lowering management of type 2 diabetes.

- Pharmacologic therapy should be guided by person-centered treatment factors, including comorbidities and treatment goals.
- In adults with type 2 diabetes and established/high risk of atherosclerotic cardiovascular disease, heart failure, and/or chronic kidney disease, the treatment regimen should include agents that reduce cardiorenal risk.
- Pharmacologic approaches that provide adequate efficacy to achieve and maintain treatment goals should be considered, such as metformin or other agents, including combination therapy.
- Weight management is an impactful component of glucose-lowering management in type 2 diabetes. The glucose-lowering treatment regimen should consider approaches that support weight management goals.
- Metformin should be continued upon initiation of insulin therapy (unless contraindicated or not tolerated) for ongoing glycemic and metabolic benefits.
 A Early combination therapy can be considered in some individuals at treatment initiation to extend the time to treatment failure.
- The early introduction of insulin should be considered if there is evidence of ongoing catabolism (weight loss), if symptoms of hyperglycemia are present, or when A1C levels (>10% [86 mmol/mol]) or blood glucose levels (greater or equal to 300 mg/dL) are very high.
- A person-centered approach should guide the choice of pharmacologic agents.
 Consider the effects on cardiovascular and renal comorbidities, efficacy, hypoglycemia risk, impact on weight, cost and access, risk for side effects, and individual preferences.
- Among individuals with type 2 diabetes who have established atherosclerotic
 cardiovascular disease or indicators of high cardiovascular risk, established
 kidney disease, or heart failure, a sodium-glucose cotransporter 2 inhibitor
 and/or glucagon-like peptide 1 receptor agonist with demonstrated
 cardiovascular disease benefit is recommended as part of the glucose-lowering
 regimen and comprehensive cardiovascular risk reduction, independent of A1C
 and in consideration of person-specific factors.
- In adults with type 2 diabetes, a glucagon-like peptide 1 receptor agonist is preferred to insulin when possible.

Healthy lifestyle behaviors, diabetes self-management, education, and support, avoidance of clinical inertia, and social determinants of health should be considered in the glucose-lowering management of type 2 diabetes. Pharmacologic therapy should be guided by person-centered treatment factors, including comorbidities and treatment goals. Pharmacotherapy should be started at the time type 2 diabetes is diagnosed unless there are contraindications. Pharmacologic approaches that provide the efficacy to achieve treatment goals should be considered, such as metformin or other agents, including combination therapy, that provide adequate efficacy to achieve and maintain treatment goals. In adults with type 2 diabetes and established/high risk of atherosclerotic cardiovascular disease (ASCVD), heart failure (HF), and/or chronic kidney disease (CKD), the treatment regimen should include agents that reduce cardiorenal risk.(11)

Pharmacologic approaches that provide the efficacy to achieve treatment goals should be considered, specified as metformin or agent(s), including combination therapy, that provide adequate efficacy to achieve and maintain treatment goals. In general, higher-efficacy approaches have greater likelihood of achieving glycemic goals, with the following considered to have very high efficacy for glucose lowering: the GLP-1 RAs dulaglutide (high dose) and semaglutide, the gastric inhibitory peptide (GIP) and GLP-1 RA tirzepatide, insulin, combination oral therapy, and combination injectable therapy. Weight management is an impactful component of glucose-lowering management in type 2 diabetes. The glucose-lowering treatment regimen should consider approaches that support weight management goals, with very high efficacy for weight loss seen with semaglutide and tirzepatide.(11)

Metformin is effective and safe, is inexpensive, and may reduce risk of cardiovascular events and death. Metformin is available in an immediate-release form for twice-daily dosing or as an extended-release form that can be given once daily. Compared with sulfonylureas, metformin as first-line therapy has beneficial effects on A1C, weight,

and cardiovascular mortality. For people with type 2 diabetes and established ASCVD or indicators of high ASCVD risk, HF, or CKD, an SGLT2 inhibitor and/or GLP-1 RA with demonstrated CVD benefit is recommended as part of the glucose-lowering regimen independent of A1C, independent of metformin use and in consideration of person-specific factors. For people without established ASCVD, indicators of high ASCVD risk, HF, or CKD, medication choice is guided by efficacy in support of individualized glycemic and weight management goals, avoidance of side effects (particularly hypoglycemia and weight gain), cost/access, and individual preferences.(11)

Dapagliflozin and empagliflozin have been shown to significantly reduce the risk of worsening heart failure or cardiovascular death independently of diabetes status.(2,4) Angiotensin-converting enzyme inhibitors (ACEIs), angiotensin receptor blockers (ARBs), angiotensin receptor-neprilysin inhibitors (ARNIs), funny current channel inhibitors (e.g., Corlanor), aldosterone antagonists, beta blockers, isosorbide dinitrate and hydralazine are all medications commonly used for heart failure with reduced ejection fraction (HFrEF).(12,13)

Safety

Invokamet, Invokamet XR, Segluromet, Synjardy, Synjardy XR, Trijardy XR, and Xigduo XR all have a black box warning for lactic acidosis due to their metformin component:(3,5-7,9,17)

- Postmarketing cases of metformin-associated lactic acidosis have resulted in death, hypothermia, hypotension, and resistant bradyarrhythmias. Symptoms included malaise, myalgias, respiratory distress, somnolence, and abdominal pain. Laboratory abnormalities included elevated blood lactate levels, anion gap acidosis, increased lactate/pyruvate ratio; and metformin plasma levels generally >5 mcg/mL.
- Risk factors include renal impairment, concomitant use of certain drugs, age
 more than 65 years old, radiological studies with contrast, surgery and other
 procedures, hypoxic states, excessive alcohol intake, and hepatic impairment.
 Steps to reduce the risk of and manage metformin-associated lactic acidosis in
 these high-risk groups are provided in the Full Prescribing Information.
- If lactic acidosis is suspected, discontinue the medication and institute general supportive measures in a hospital setting. Prompt hemodialysis is recommended.

Brenzavvy, Farxiga, Invokana, Jardiance, Steglatro, and Glyxambi are contraindicated in patients on dialysis.(1,2,4,8,14,19)

Inpefa is contraindicated in history of serious hypersensitivity reaction to Inpefa. (18)

Invokamet and Invokamet XR are contraindicated in patients with severe renal impairment, acute or chronic metabolic acidosis, including diabetic ketoacidosis.(3)

Segluromet, Synjardy, Synjardy XR, Xigduo XR, and Trijardy XR are contraindicated in patients with severe renal impairment, end stage renal disease (ESRD), patients on dialysis, and patients with acute or chronic metabolic acidosis, including diabetic ketoacidosis. (5-7,9,17)

Steglujan and Qtern are contraindicated in patients with severe renal impairment, end stage renal disease (ESRD), or on dialysis.(5-7,9,17)

REFERENCES

	<u> </u>
Number	Reference
1	Invokana prescribing information. Janssen Pharmaceuticals, Inc. August 2024.
2	Farxiga prescribing information. Astra Zeneca. June 2024.
3	Invokamet and Invokamet XR prescribing information. Janssen Pharmaceuticals, Inc. August 2024.
4	Jardiance prescribing information. Boehringer Ingelheim Pharmaceuticals, Inc. September 2023.

Number	Reference
5	Xigduo XR prescribing information. AstraZeneca Pharmaceuticals LP. June 2024.
6	Synjardy prescribing information. Boehringer Ingelheim. October 2023.
7	Synjardy XR prescribing information. Boehringer Ingelheim. March 2022.
8	Steglatro prescribing information. Merck & Co, Inc. June 2024.
9	Segluromet prescribing information. Merck Sharp & Dohme Corp. June 2024.
10	American Diabetes Association. Standards of Medical Care in Diabetes-2022. Available at: https://care.diabetesjournals.org/content/45/Supplement_1.
11	ElSayed NA, Aleppo G, Bannuru RR, et al. 9. Pharmacologic Approaches to Glycemic Treatment: Standards of Care in Diabetes-2024. Diabetes Care. 2023;47(Supplement_1):S158-S178. doi:10.2337/dc24-s009.
12	ElSayed NA, Aleppo G, Bannuru RR, et al. 10. Cardiovascular Disease and Risk Management: Standards of Care in Diabetes-2024. Diabetes Care. 2023;47(Supplement_1):S179-S218. doi:10.2337/dc24-s010
13	Heidenreich PA, Bozkurt B, Aguilar D, et al. 2022 AHA/ACC/HFSA Guideline for the Management of Heart Failure: A report of the American College of Cardiology/American Heart Association Joint Committee on Clinical Practice Guidelines. Circulation. 2022;145(18). doi:10.1161/cir.00000000000001063
14	Glyxambi prescribing information. Boehringer Ingelheim Pharmaceuticals, Inc and Eli Lilly and Company. October 2023.
15	Qtern prescribing information. Astra Zeneca. September 2023.
16	Steglujan prescribing information. Merck & Co., Inc. June 2024.
17	Trijardy XR prescribing information. Boehringer Ingelheim International GmbH. October 2023.
18	Inpefa Prescribing Information. Lexicon Pharmaceuticals, Inc. January 2024.
19	Brenzavvy prescribing information. TheracosBio, LLC. September 2023.

POLICY AGENT SUMMARY STEP THERAPY

Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	Targeted MSC	Availabl e MSC	Final Age Limit	Preferred Status
Bexagliflozin ; Brenzavvy	bexagliflozin tab	20 MG	M;N;O	М		
Dapagliflozin propanediol ; Farxiga	dapagliflozin propanediol tab	10 MG ; 5 MG	M;N;O	М		
Dapagliflozin propanediol ; Xigduo xr	dapagliflozin prop-metformin hcl tab er	10-1000 MG; 10-500 MG; 2.5-1000 MG; 5-1000 MG; 5-000 MG	M;N;O	M ; N		
Glyxambi	empagliflozin-linagliptin tab	10-5 MG ; 25-5 MG	M;N;O	N		
Inpefa	sotagliflozin tab	200 MG ; 400 MG	M;N;O	N		
Invokamet	canagliflozin-metformin hcl tab	150-1000 MG; 150-500 MG; 50-1000 MG; 50-500 MG	M;N;O	N		
Invokamet xr	canagliflozin-metformin hcl tab er	150-1000 MG; 150-500 MG; 50-1000 MG; 50-500 MG	M;N;O	N		
Invokana	canagliflozin tab	100 MG ; 300 MG	M;N;O	N		
Jardiance	empagliflozin tab	10 MG ; 25 MG	M;N;O	N		
Qtern	dapagliflozin-saxagliptin tab	10-5 MG ; 5-5 MG	M;N;O	N		

Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	Targeted MSC	Availabl e MSC	Final Age Limit	Preferred Status
Segluromet	ertugliflozin-metformin hcl tab	2.5-1000 MG; 2.5-500 MG; 7.5-1000 MG; 7.5-500 MG	M;N;O	N		
Steglatro	ertugliflozin l-pyroglutamic acid tab	15 MG ; 5 MG	M;N;O	N		
Steglujan	ertugliflozin-sitagliptin tab	15-100 MG ; 5- 100 MG	M;N;O	N		
Synjardy	empagliflozin-metformin hcl tab	12.5-1000 MG ; 12.5-500 MG ; 5-1000 MG ; 5-500 MG	M;N;O	N		
Synjardy xr	empagliflozin-metformin hcl tab er	10-1000 MG; 12.5-1000 MG; 25-1000 MG; 5-1000 MG	M;N;O	N		
Trijardy xr	empagliflozin-linaglip-metformin tab er ; empagliflozin-linagliptin- metformin tab er	10-5-1000 MG ; 12.5-2.5- 1000 MG ; 25- 5-1000 MG ; 5- 2.5-1000 MG	M;N;O	N		

POLICY AGENT SUMMARY QUANTITY LIMIT

Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strengt h	QL Amount	Dose Form	Day Supply	Duratio n	Addtl QL Info	Allowed Exceptions	Targete d NDCs When Exclusi ons Exist
Bexagliflozin ; Brenzavvy	bexagliflozin tab	20 MG	30	Tablets	30	DAYS			
Dapagliflozin propanediol ; Farxiga	dapagliflozin propanediol tab	10 MG ; 5 MG	30	Tablets	30	DAYS			
Dapagliflozin propanediol ; Xigduo xr	Dapagliflozin- Metformin HCl Tab ER 24HR 10-1000 MG	10-1000 MG	30	Tablets	30	DAYS			
Dapagliflozin propanediol ; Xigduo xr	Dapagliflozin- Metformin HCl Tab ER 24HR 5-1000 MG	5-1000 MG	60	Tablets	30	DAYS			
Glyxambi	empagliflozin- linagliptin tab	10-5 MG ; 25-5 MG	30	Tablets	30	DAYS			
Inpefa	sotagliflozin tab	200 MG ; 400 MG	30	Tablets	30	DAYS			
Invokamet	canagliflozin- metformin hcl tab	150- 1000 MG; 150-500 MG; 50-1000 MG; 50-500 MG	60	Tablets	30	DAYS			
Invokamet xr	canagliflozin- metformin hcl tab er	150- 1000 MG; 150-500 MG; 50-1000 MG; 50-500 MG	60	Tablets	30	DAYS			

Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strengt h	QL Amount	Dose Form	Day Supply	Duratio n	Addtl QL Info	Allowed Exceptions	Targete d NDCs When Exclusi ons Exist
Invokana	canagliflozin tab	100 MG ; 300 MG	30	Tablets	30	DAYS			
Jardiance	empagliflozin tab	10 MG ; 25 MG	30	Tablets	30	DAYS			
Qtern	dapagliflozin- saxagliptin tab	10-5 MG ; 5-5 MG	30	Tablets	30	DAYS			
Segluromet	Ertugliflozin- Metformin HCl Tab 2.5-1000 MG	2.5- 1000 MG	60	Tablets	30	DAYS			
Segluromet	Ertugliflozin- Metformin HCl Tab 2.5-500 MG	2.5-500 MG	120	Tablets	30	DAYS			
Segluromet	Ertugliflozin- Metformin HCl Tab 7.5-1000 MG	7.5- 1000 MG	60	Tablets	30	DAYS			
Segluromet	Ertugliflozin- Metformin HCl Tab 7.5-500 MG	7.5-500 MG	60	Tablets	30	DAYS			
Steglatro	Ertugliflozin L- Pyroglutamic Acid Tab 15 MG (Base Equiv)	15 MG	30	Tablets	30	DAYS			
Steglatro	Ertugliflozin L- Pyroglutamic Acid Tab 5 MG (Base Equiv)	5 MG	60	Tablets	30	DAYS			
Steglujan	ertugliflozin- sitagliptin tab	15-100 MG;5- 100 MG	30	Tablets	30	DAYS			
Synjardy	empagliflozin- metformin hcl tab	12.5- 1000 MG; 12.5- 500 MG; 5- 1000 MG; 5- 500 MG	60	Tablets	30	DAYS			
Synjardy xr	Empagliflozin- Metformin HCl Tab ER 24HR 10-1000 MG	10-1000 MG	60	Tablets	30	DAYS			
Synjardy xr	Empagliflozin- Metformin HCl Tab ER 24HR 12.5-1000 MG	12.5- 1000 MG	60	Tablets	30	DAYS			
Synjardy xr	Empagliflozin- Metformin HCl Tab ER 24HR 25-1000 MG	25-1000 MG	30	Tablets	30	DAYS			
Synjardy xr	Empagliflozin- Metformin HCl Tab ER 24HR 5-1000 MG	5-1000 MG	60	Tablets	30	DAYS			
Trijardy xr	Empagliflozin- Linaglip-Metformin Tab ER 24HR 12.5- 2.5-1000MG	12.5- 2.5- 1000 MG	60	Tablets	30	DAYS			
Trijardy xr	Empagliflozin- Linagliptin-Metformin Tab ER 24HR 10-5- 1000 MG	10-5- 1000 MG	30	Tablets	30	DAYS			
Trijardy xr	Empagliflozin- Linagliptin-Metformin	25-5- 1000 MG	30	Tablets	30	DAYS			

Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strengt h	QL Amount	Dose Form	Day Supply	Duratio n	Addtl QL Info	Allowed Exceptions	Targete d NDCs When Exclusi ons Exist
	Tab ER 24HR 25-5- 1000 MG								
Trijardy xr	Empagliflozin- Linagliptin-Metformin Tab ER 24HR 5-2.5- 1000MG	5-2.5- 1000 MG	60	Tablets	30	DAYS			
Xigduo xr	Dapagliflozin- Metformin HCl Tab ER 24HR 10-500 MG	10-500 MG	30	Tablets	30	DAYS			
Xigduo xr	Dapagliflozin- Metformin HCl Tab ER 24HR 2.5-1000 MG	2.5- 1000 MG	60	Tablets	30	DAYS			
Xigduo xr	Dapagliflozin- Metformin HCl Tab ER 24HR 5-500 MG	5-500 MG	30	Tablets	30	DAYS			

CLIENT SUMMARY - STEP THERAPY

Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	Client Formulary
Bexagliflozin ; Brenzavvy	bexagliflozin tab	20 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Dapagliflozin propanediol ; Farxiga	dapagliflozin propanediol tab	10 MG ; 5 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Dapagliflozin propanediol ; Xigduo xr	dapagliflozin prop-metformin hcl tab er	10-1000 MG; 10-500 MG; 2.5-1000 MG; 5-500 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Glyxambi	empagliflozin-linagliptin tab	10-5 MG ; 25-5 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Inpefa	sotagliflozin tab	200 MG ; 400 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Invokamet	canagliflozin-metformin hcl tab	150-1000 MG ; 150-500 MG ; 50-1000 MG ; 50- 500 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Invokamet xr	canagliflozin-metformin hcl tab er	150-1000 MG ; 150-500 MG ; 50-1000 MG ; 50- 500 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Invokana	canagliflozin tab	100 MG ; 300 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Jardiance	empagliflozin tab	10 MG ; 25 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Qtern	dapagliflozin-saxagliptin tab	10-5 MG ; 5-5 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Segluromet	ertugliflozin-metformin hcl tab	2.5-1000 MG; 2.5-500 MG; 7.5-1000 MG; 7.5- 500 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Steglatro	ertugliflozin l-pyroglutamic acid tab	15 MG ; 5 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Steglujan	ertugliflozin-sitagliptin tab	15-100 MG ; 5-100 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Synjardy	empagliflozin-metformin hcl tab	12.5-1000 MG ; 12.5-500 MG ; 5-1000 MG ; 5-500 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Synjardy xr	empagliflozin-metformin hcl tab er	10-1000 MG ; 12.5-1000 MG ; 25-1000 MG ; 5- 1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Trijardy xr	empagliflozin-linaglip-metformin tab er ; empagliflozin-linagliptin-metformin tab er	10-5-1000 MG ; 12.5-2.5- 1000 MG ; 25-5-1000 MG ; 5-2.5-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers

CLIENT SUMMARY - QUANTITY LIMITS

Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	Client Formulary
Bexagliflozin ; Brenzavvy	bexagliflozin tab	20 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Dapagliflozin propanediol ; Farxiga	dapagliflozin propanediol tab	10 MG ; 5 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Dapagliflozin propanediol ; Xigduo xr	Dapagliflozin-Metformin HCl Tab ER 24HR 10-1000 MG	10-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Dapagliflozin propanediol ; Xigduo xr	Dapagliflozin-Metformin HCl Tab ER 24HR 5-1000 MG	5-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Glyxambi	empagliflozin-linagliptin tab	10-5 MG ; 25-5 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Inpefa	sotagliflozin tab	200 MG ; 400 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Invokamet	canagliflozin-metformin hcl tab	150-1000 MG ; 150-500 MG ; 50-1000 MG ; 50- 500 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Invokamet xr	canagliflozin-metformin hcl tab er	150-1000 MG ; 150-500 MG ; 50-1000 MG ; 50- 500 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Invokana	canagliflozin tab	100 MG ; 300 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Jardiance	empagliflozin tab	10 MG ; 25 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Qtern	dapagliflozin-saxagliptin tab	10-5 MG ; 5-5 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Segluromet	Ertugliflozin-Metformin HCl Tab 2.5-1000 MG	2.5-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Segluromet	Ertugliflozin-Metformin HCl Tab 2.5-500 MG	2.5-500 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Segluromet	Ertugliflozin-Metformin HCl Tab 7.5-1000 MG	7.5-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Segluromet	Ertugliflozin-Metformin HCl Tab 7.5-500 MG	7.5-500 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Steglatro	Ertugliflozin L-Pyroglutamic Acid Tab 15 MG (Base Equiv)	15 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Steglatro	Ertugliflozin L-Pyroglutamic Acid Tab 5 MG (Base Equiv)	5 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Steglujan	ertugliflozin-sitagliptin tab	15-100 MG ; 5-100 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Synjardy	empagliflozin-metformin hcl tab	12.5-1000 MG; 12.5-500 MG; 5-1000 MG; 5-500 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Synjardy xr	Empagliflozin-Metformin HCl Tab ER 24HR 10-1000 MG	10-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Synjardy xr	Empagliflozin-Metformin HCl Tab ER 24HR 12.5-1000 MG	12.5-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Synjardy xr	Empagliflozin-Metformin HCl Tab ER 24HR 25-1000 MG	25-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Synjardy xr	Empagliflozin-Metformin HCl Tab ER 24HR 5-1000 MG	5-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Trijardy xr	Empagliflozin-Linaglip-Metformin Tab ER 24HR 12.5-2.5-1000MG	12.5-2.5-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Trijardy xr	Empagliflozin-Linagliptin-Metformin Tab ER 24HR 10-5-1000 MG	10-5-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Trijardy xr	Empagliflozin-Linagliptin-Metformin Tab ER 24HR 25-5-1000 MG	25-5-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Trijardy xr	Empagliflozin-Linagliptin-Metformin Tab ER 24HR 5-2.5-1000MG	5-2.5-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Xigduo xr	Dapagliflozin-Metformin HCl Tab ER 24HR 10-500 MG	10-500 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers
Xigduo xr	Dapagliflozin-Metformin HCl Tab ER 24HR 2.5-1000 MG	2.5-1000 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers

Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	Client Formulary
Xigduo xr	Dapagliflozin-Metformin HCl Tab ER 24HR 5-500 MG	5-500 MG	Commercial ; HIM 5 and 6 Tiers ; HIM 7 Tiers

STEP THERAPY CLINICAL CRITERIA FOR APPROVAL

Module	Clinical Criteria for Approval
All other	TARGET AGENT(S)
target	TARGET AGENT(S)
agent(s)	Brenzavvy, Bexagliflozin Glyxambi (empagliflozin/linagliptin) Invokana (canagliflozin) Invokamet (canagliflozin/metformin) Invokamet XR (canagliflozin/metformin ER) Qtern (dapagliflozin/saxagliptin) Segluromet (ertugliflozin/metformin) Steglatro (ertugliflozin) Steglujan (ertugliflozin/sitagliptin) Synjardy (empagliflozin/metformin) Synjardy XR (empagliflozin/metformin ER) Trijardy XR (empagliflozin/linagliptin/metformin ER) Xigduo XR, Dapagliflozin/metformin ER
	Target Agent(s) will be approved when ONE of the following is met:
Farxiga	 The patient is currently being treated with the requested SGLT inhibitor within the past 90 days OR The prescriber states the patient is currently being treated with the requested SGLT inhibitor within the past 90 days AND is at risk if therapy is changed OR The patient's medication history includes use of an agent containing metformin or insulin in the past 90 days OR The patient has an intolerance or hypersensitivity to ONE of the following agents: metformin or insulin OR The patient has an FDA labeled contraindication to ALL of the following agents: metformin and insulin OR The patient has a diagnosis of type 2 diabetes with or at high risk for atherosclerotic cardiovascular disease, heart failure, and/or chronic kidney disease Length of Approval: 12 months NOTE: If Quantity Limit applies, please refer to Quantity Limit Criteria. TARGET AGENT(S)
	Farxiga, Dapagliflozin
	Target Agent(s) will be approved when ONE of the following is met:
	 The patient is currently being treated with the requested SGLT inhibitor within the past 90 days OR The prescriber states the patient is currently being treated with the requested SGLT inhibitor within the past 90 days AND is at risk if therapy is changed OR The patient has a diagnosis of heart failure OR The patient has a diagnosis of type 2 diabetes with or at high risk for atherosclerotic cardiovascular disease, heart failure, and/or chronic kidney disease OR The patient has a diagnosis of chronic kidney disease (CKD) OR The patient's medication history includes use of an agent containing metformin, insulin, ACE inhibitors, angiotensin receptor blockers (ARBs), angiotensin receptor-neprilysin inhibitors (ARNIs), If channel inhibitors (e.g., Corlanor), aldosterone antagonists, beta blockers, isosorbide dinitrate or hydralazine in the past 90 days OR

Module	Clinical Criteria for Approval
	7. The patient has an intolerance or hypersensitivity to ONE of the following agents: metformin or
	insulin OR 8. The patient has an FDA labeled contraindication to ALL of the following agents: metformin and insulin OR
	9. The patient has an intolerance or hypersensitivity to ONE of the following agents: ACE inhibitors, angiotensin receptor blockers (ARBs), angiotensin receptor-neprilysin inhibitors (ARNIs), If channel inhibitors (e.g., Corlanor), aldosterone antagonists, beta blockers, isosorbide dinitrate, or hydralazine OR
	10. The patient has an FDA labeled contraindication to ALL of the following agents: ACE inhibitors, angiotensin receptor blockers (ARBs), angiotensin receptor-neprilysin inhibitors (ARNIs), If channel inhibitors (e.g., Corlanor), aldosterone antagonists, beta blockers, isosorbide dinitrate, and hydralazine
	Length of Approval: 12 months
	NOTE: If Quantity Limit applies, please refer to Quantity Limit Criteria.
and	TARGET AGENT(S)
Inpefa	Jardiance (empagliflozin) Inpefa (sotagliflozin)
	Target Agent(s) will be approved when ONE of the following is met:
	1. Information has been provided that indicates the patient is currently being treated with the requested SGLT inhibitor within the past 90 days OR
	2. The prescriber states the patient is currently being treated with the requested SGLT inhibitor within the past 90 days AND is at risk if therapy is changed OR
	 If the requested agent is Jardiance, then BOTH of the following: A. The patient has a diagnosis of chronic kidney disease (CKD) AND B. The patient is at high risk for progression of CKD, including, risk of sustained decline in eGFR, end-stage kidney disease, cardiovascular death, and hospitalization OR
	4. The patient has a diagnosis of heart failure OR
	5. The patient has a diagnosis of type 2 diabetes with or at high risk for atherosclerotic cardiovascular disease, heart failure, and/or chronic kidney disease OR
	6. The patient's medication history includes use of an agent containing metformin, insulin, ACE inhibitors, angiotensin receptor blockers (ARBs), angiotensin receptor-neprilysin inhibitors (ARNIs), If channel inhibitors (e.g., Corlanor), aldosterone antagonists, beta blockers, isosorbide dinitrate or hydralazine in the past 90 days OR
	7. The patient has an intolerance or hypersensitivity to ONE of the following agents: metformin or insulin OR
	8. The patient has an FDA labeled contraindication to ALL of the following agents: metformin and insulin OR
	 The patient has an intolerance or hypersensitivity to ONE of the following agents: ACE inhibitors, angiotensin receptor blockers (ARBs), angiotensin receptor-neprilysin inhibitors (ARNIs), I_f channel inhibitors (e.g., Corlanor), aldosterone antagonists, beta blockers, isosorbide dinitrate, or hydralazine OR
	10. The patient has an FDA labeled contraindication to ALL of the following agents: ACE inhibitors, angiotensin receptor blockers (ARBs), angiotensin receptor-neprilysin inhibitors (ARNIs), If channel inhibitors (e.g., Corlanor), aldosterone antagonists, beta blockers, isosorbide dinitrate, and hydralazine
	Length of Approval: 12 months
	NOTE: If Quantity Limit applies, please refer to Quantity Limit Criteria.

OUANTITY LIMIT CLINICAL CRITERIA FOR APPROVAL

Module	Clinical Criteria for Approval				
Universa	Quantity limit for the Target Agent(s) will be approved when ONE of the following is met:				
l QL					

Module	Clinical Criteria for Approval
Module	1. The requested quantity (dose) does NOT exceed the program quantity limit OR 2. The requested quantity (dose) exceeds the program quantity limit AND ONE of the following: A. BOTH of the following: 1. The requested agent does NOT have a maximum FDA labeled dose for the requested indication AND 2. There is support for therapy with a higher dose for the requested indication OR B. BOTH of the following: 1. The requested quantity (dose) does NOT exceed the maximum FDA labeled dose for the requested indication AND 2. There is support for why the requested quantity (dose) cannot be achieved with a lower quantity of a higher strength that does NOT
	exceed the program quantity limit OR C. BOTH of the following: 1. The requested quantity (dose) exceeds the maximum FDA labeled dose for the requested indication AND 2. There is support for therapy with a higher dose for the requested indication Length of Approval: up to 12 months