

Request for Prior Authorization



VERICIGUAT (VERQUVO)

Provider Help Desk I (844) 236-1464

(PLEASE PRINT – ACCURACY IS IMPORTANT)

Fax Completed Form To 1 (877) 733-3195

IA Medicaid Member ID #	Patient name		DOB		
Patient address					
Provider NPI	Prescriber name		Phone		
Prescriber address	,		Fax		
. Pharmacy name	Address	1	Phone		
Prescriber must complete all inform	ation above. It must be legible, correct, an	d complete or fo	orm will be returned.		
Pharmacy NPI	Pharmacy fax	NDC			
Prior authorization is required for vericiguat (Verquvo). Payment will be considered when patient has an FDA approved or compendia indicated indication for the requested drug under the following conditions: 1) Request adheres to all FDA approved labeling for requested drug and indication, including age, dosing, contraindications, warnings and precautions, drug interactions, and use in specific populations; and					
 Patient has a diagnosis of symptomatic chronic heart failure (NYHF Class II-IV) with a left ventricular ejection fraction (LVEF) ≤ 45%; and 					
3) Patient meets one of the following:					
a. Recent hospitalization for heart failure (within the last 6 months); or					
b. Recent need for outpatient intravenous diuretics (within the last 3 months); and					
4) Female patients of reproductive potential have been advised to use effective contraception during treatment and for at least one month after the last dose; and					
5) Will not be used concomitantly with other soluble guanylate cyclase (sGC) stimulators (e.g. riociguat) or phosphodiesterase type 5 (PDE-5) inhibitors (e.g. sildenafil, tadalafil, vardenafil); and					
6) Documentation of prior or current therapy, at a maximally tolerated dose, with one drug from each category below:					
 a. Renin-angiotensin system inhibitor (angiotensin converting enzyme [ACEI], angiotensin receptor blocker [ARB], or angiotensin receptor-neprilysin inhibitor [ARNI]; and 					
b. Evidence-based beta-blocker (carvedilol, metoprolol succinate, or bisoprolol); and					
c. Mineralcorticoid receptor antagonist (MRA); and					
 d. Sodium-glucose cotransporter 2 inhibitor (SGLT2i) indicated for the treatment of heart failure (empagliflozin or dapagliflozin); and 					
 Initial requests for vericiguat (Verquvo) 2.5mg and 5mg tablets will be limited to one 14-day supply for each strength. 					
The required trials may be overr medically contraindicated.	idden when documented evidence is p	rovided that us	e of these agents would be		
Non-Preferred					
☐ Verquvo					
Strength Dosag	ge Instructions	Quantity	Days Supply		
Diagnosis:					

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Document LVEF:		
Patient meets one of the following:		
Recent hospitalization for heart failure: Provide date:		
Recent need for outpatient intravenous diuretics: Provide date & drug name	me:	_
Female patient of reproductive potential has been advised to use eff treatment and for at least one month after last dose?	ective contraception during	☐ No
Will Verquvo be used in combination with sGC stimulators or PDE-	5 inhibitors?	☐ No
Document prior or current therapy, at maximally tolerated dose, w below:	ith one drug from each categ	ory
Renin-angiotensin system inhibitor (ACEI, ARB, ARNI):		
Name/Dose:Tria	ıl Dates:	
Failure reason:		
Evidence-based beta-blocker (carvedilol, metoprolol succinate, or bi	soprolol):	
Name/Dose:Tria	ıl Dates:	
Failure reason:		
Mineralocorticoid receptor antagonist (MRA):		
Name/Dose:Tria	ıl Dates:	
Failure reason:		
Sodium-glucose cotransporter 2 inhibitor (SGLT2i) indicated for the (empagliflozin or dapagliflozin):	e treatment of heart failure	
Name/Dose:Tria	ıl Dates:	
Failure reason:		
Medical or contraindication reason to override trial requirements:		
Attach lab results and other documentation as necessary.		
Prescriber signature (Must match prescriber listed above.)	Date of submission	
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IMPORTANT NOTE: In evaluating requests for prior authorization the consultant will consider the treatment from the standpoint of medical necessity only. If approval of this request is granted, this does not indicate that the member continues to be eligible for Medicaid. It is the responsibility of the provider who initiates the request for prior authorization to establish by inspection of the member's Medicaid eligibility card and, if necessary by contact with the county Department of Human Services, that the member continues to be eligible for Medicaid.

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