SENTARA COMMUNITY PLAN (MEDICAID)

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions</u>: <u>The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request</u>. All other information may be filled in by office staff; <u>fax to 1-800-750-9692</u>. No additional phone calls will be necessary if all information (<u>including phone and fax #s</u>) on this form is correct. <u>If the information provided is not complete, correct, or legible, the authorization process can be delayed.</u>

Drug Requested: Kerendia® (finerenone)

MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.			
Member Name:			
Member Sentara #:			
Prescriber Name:			
Prescriber Signature:	Date:		
Office Contact Name:			
Phone Number:			
DEA OR NPI #:			
DRUG INFORMATION: Authoriza	tion may be delayed if incomplete.		
Drug Form/Strength:			
Dosing Schedule:	Length of Therapy:		
Diagnosis:	ICD Code, if applicable:		
Weight:	Date:		
Quantity Limit: 30 tablets per 30 days			

Kerendia[®] **Initial Dosing Recommendations**:

eGFR (mL/min/1.73m ²)	Starting Dose
≥ 60	20 mg once daily
\geq 25 to < 60	10 mg once daily
< 25	Not Recommended

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Dose Adjustment Based on Current Serum Potassium Concentration and Current Kerendia® Dose				
		10 mg once daily	20 mg once daily	
Current Serum Potassium (mEq/L)	≤ 4.8	Increase the dose to 20 mg once daily*	Maintain 20 mg once daily	
	> 4.8 – 5.5	Maintain 10 mg once daily	Maintain 20 mg once daily	
	> 5.5	Withhold Kerendia [®] . Consider restarting at 10 mg once daily when serum potassium ≤ 5.0 mEq/L	Withhold Kerendia®. Restart at 10 mg once daily when serum potassium ≤ 5.0 mEq/L	

^{*}If eGFR has decreased by more than 30% compared to previous measurement, maintain 10 mg dose.

CLINICAL CRITERIA: Check below all that apply. <u>All criteria must be met for approval</u>. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.

Initial Authorization: 6 months

- ☐ Member is 18 years of age or older and has a diagnosis of chronic kidney disease associated with type 2 diabetes
- ☐ Must submit lab test results documenting **BOTH** of the following obtained within the past 60 days
 - ☐ Members' current eGFR is > 25mL/minute/1.73 m²
 - \square Member's current Urinary Albumin-to -Creatinine Ratio (UACR) is $\ge 30 \text{ mg/g}$
- □ Member's current serum potassium is \leq 5 mEq/L along with <u>BOTH</u> of the following (submit current lab documentation obtained within the past 60 days):
 - ☐ Therapy will **NOT** be initiated if serum potassium >5 mEq/L
 - □ Initiation with increased serum potassium monitoring during the first 4 weeks will be preformed if serum potassium is >4.8 to 5 mEq/L
- ☐ Member is established on treatment with maximally tolerated dose of angiotensin-converting enzyme inhibitor (ACE) or angiotensin receptor blocker (ARB) medication and will continue to take along with Kerendia® (finerenone)
- □ Member tried and failed, has a contradiction, or intolerance to at least <u>ONE</u> sodium glucose transport protein 2 (SGLT2) inhibitor that is indicated for use in patients with chronic kidney disease (e.g., Farxiga[®], Jardiance[®])
- ☐ Member doses <u>NOT</u> have a diagnosis of adrenal insufficiency or a diagnosis of know significant non-diabetic renal disease, including clinically relevant renal artery stenosis
- ☐ Member is **NOT** receiving simultaneous treatment with strong CYP3A4 inhibitors
- ☐ For initial therapy, member will be dosed as follows:
 - \Box eGFR \geq 60 mL/minute/1.73 m²: starting dose will be 20 mg once daily
 - \Box eGFR \geq 25 to <60 mL/minute/1.73 m²: starting dose will be 10 mg once daily

Reauthorization: 12 months. Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.

- ☐ Member's current eGFR is >25 mL/minute/1.73 m² (submit current lab documentation)
- ☐ Member's Urinary Albumin-to-creatinine Ratio (UACR) has decreased by $\geq 30\%$ from baseline level and/or been sustained at $\geq 30\%$ below baseline level since last approval (submit current lab documentation)
- ☐ Member's current serum potassium level does <u>NOT</u> exceed 5.5 mEq/L (submit current lab documentation)
- □ Provider attests Kerendia[®] will be withheld if serum potassium is >5.5 mEq/L and will consider restarting therapy when serum potassium normalizes ($\leq 5.0 \text{ mEq/L}$)

**Use of samples to initiate therapy does not meet step edit/ preauthorization criteria. **

Previous therapies will be verified through pharmacy paid claims or submitted chart notes.