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)</u> on this request. 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 (including phone and fax #s) on this form is correct. <u>If the information provided is not complete, correct, or legible, the authorization process can be delayed.</u>

Continuous Glucose Monitors (CGM)

<u>Device Requested</u>: (Check below the CGM that applies, only **<u>ONE</u>** prior authorization form is required)

Formulary 1	Preferred CGMs		
☐ FreeStyle Libre 14 Day System (Reader/Sensors)	□ FreeStyle Libre 2 System (Reader/Sensors/Plus Sensors)		
 □ Dexcom G6[™] System (Receiver/Transmitter/Sensors) 	□ FreeStyle Libre 3 System (Reader/Sensors/Plus Sensors)		
 □ Dexcom G7[™] System (Receiver/Transmitter/Sensors) 			
Non-Fo	rmulary CGMs		
Provider please note: Medical Exception is required			
□ Eversense® (Sensor/Transmitter)	□ Guardian [™] 3 (Transmitter/Sensors)		
☐ Guardian [™] 4 (Transmitter/Sensors)			
PROVIDER PLEASE NOTE:			
• Only one prior authorization form is required to be submitted for the request of any formulary or non-formulary CGM. If approved, an authorization will be entered for corresponding CGM supplies (e.g., readers, sensors, transmitters).			
MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.			
Member Name:			
Member Sentara #:	Date of Birth:		
Prescriber Name:			
Prescriber Signature:	Date:		
Office Contact Name:			

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Phone Number: _____ Fax Number: _____

DRUG INFORMATION: Authorization may be delayed if incomplete. Drug Name/Form/Strength:			
ICD Code, if applicable:			
Weigh	nt (if applicable):	Date weight obtained:	
Ouantity Limits:			
	<u>Dexcom</u>	<u>Freestyle</u>	
• 3	receiver per year sensors per 30 days transmitter per 90 days	1 reader kit per year2 sensors per 28 days2 plus sensors per 30 days	
CLINICAL CRITERIA: 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. ☐ Has the member been approved for a CGM device previously through the Sentara Health Plans medical department? ☐ Yes ☐ No			
□ C	Continuous Glucose Monitors		
Length of Authorization: 12 months (1 year)			
	The member is 2 years of age or older.		
	The member has been diagnosed with diabetes by their primary care physician, or another licensed health care practitioner authorized to make such a diagnosis.		
	The member is being treated with insulin and/or	the member has a history of problematic hypoglycemia.	
	The member's treating practitioner concluded that the member (or member's caregiver) has had sufficient training using the continuous glucose monitor prescribed as evidenced by the prescription provided.		
	The continuous glucose monitor has been present Administration indications for use.	ribed in accordance with the Food and drug	

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ı I	Replacement Device
	Member has been previously approved for a CGM
	Member has been benefiting from the use of the CGM
	At least ONE of the following problems have occurred which limits the use of the member's current CGM:
	☐ Reagent or instrument failure/defective devices
	□ Defects in product design
	□ Product instability
	☐ Failure to perform according to performance characterized in package insert
	☐ Incorrect test results (falsely elevated or decreased glucose results) that cause or contributed to an incorrect patient diagnosis and/or treatment
	☐ Unexplained quality control (QC) failures
	☐ Any other device problems that may compromise patient health or safety
	Provider or member must submit documentation that the member's current CGM is not under warranty, including the date of warranty expiration
upp	uthorization: 12 months. Check below all that apply. All criteria must be met for approval. To out each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be ided or request may be denied.
	The member continues to meet the relevant criteria identified in the initial criteria.
	Member has participated in follow-up care with their treating health care practitioner, in person or through telehealth, at least once every 6 months during the first 18 months after the first prescription of the CGM to assess the efficacy of using the monitor for the treatment of diabetes OR
	The member has, after the first 18 months of CGM use, followed up with their treating health care practitioner at least once every 12 months.

**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.