# AvMed

# PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST\*

**Directions:** The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; <u>fax to 1-305-671-0200</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</u> complete, correct, or legible, the authorization process can be delayed.

# **Continuous Glucose Monitors (CGM)**

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

Formulary Preferred CGM's			
	FreeStyle Libre 14 Day System	FreeStyle Libre 2 System	
	(Reader/Sensors)	(Reader/Sensors)	
	Dexcom G6 <sup>™</sup> System	□ FreeStyle Libre 3 (Reader/Sensors)	
	(Receiver/Transmitter/Sensors)		
	Dexcom G7 <sup>™</sup> System		
	(Receiver/Transmitter/Sensors)		
Non-Formulary – Provider please note: A pharmacy medical necessity request form must be submitted for all Non-Formulary CGM requests			
	Eversense <sup>®</sup> (Sensor/Transmitter)	□ Guardian <sup>™</sup> 3 (Transmitter/Sensors)	
	Guardian <sup>™</sup> 4 (Transmitter/Sensors)		

<u>PROVIDER PLEASE NOTE</u>: 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).

(Continued on next page)

MEMBER & PRESCRIBER INFO	<b>RMATION:</b> Authorization may be delayed if incomplete.		
Member Name:			
Member AvMed #:	Date of Birth:		
Prescriber Name:			
Prescriber Signature:			
Office Contact Name:			
Phone Number:	ne Number: Fax Number:		
DEA OR NPI #:			
DRUG INFORMATION: Authorizat	ion may be delayed if incomplete.		
Drug Form/Strength:			
Dosing Schedule:	Length of Therapy:		
Diagnosis:	ICD Code, if applicable:		
Weight:	Date:		

## **<u>Quantity Limits</u>:**

1	it per lifetime per 28 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.

### **Continuous Glucose Monitors – Long Term Use**

### Length of Authorization: Indefinite

- □ Member requires Long-term CGM device indicated by <u>ALL</u> the following:
  - Diagnosis of <u>ONE</u> of the following:
    - **D** Type 1 Diabetes Mellitus
    - □ Type 2 Diabetes Mellitus
    - Gestational Diabetes

- □ Member requires a demanding insulin regimen of at least three or more insulin injections per day or current use of an insulin pump
- □ Member or guardian consistently monitors blood glucose three or more times per day
- Provider attests that the member is motivated and knowledgeable about use of CGMs, is adherent to diabetic treatment plan, and participates in ongoing education and support

#### **Continuous Glucose Monitors – Short Term Use**

#### Length of Authorization: 1 month (30 days)

- □ Member requires Short-term CGM device indicated by <u>ALL</u> the following:
  - □ Diagnosis of <u>ONE</u> of the following:
    - □ Type 1 Diabetes Mellitus
    - □ Type 2 Diabetes Mellitus
    - **Gestational Diabetes**
  - □ Member must demonstrate at least <u>ONE</u> of the following:
    - Observed increase in blood glucose levels that takes place in the early morning (also known as The Dawn Phenomenon), known or suspected
    - □ Hypoglycemia unawareness (i.e., member does not have symptoms with hypoglycemia)
    - □ Nocturnal hypoglycemia, known or suspected
    - Destprandial hyperglycemia, known or suspected
    - □ Significant change to diabetes treatment regimen (e.g., initiation of insulin, change from multiple- dose insulin to insulin pump therapy)
    - □ Unexplained hyperglycemia
  - □ Member requires short term blood glucose monitoring (i.e., 7-14 days)

#### **D** Replacement Device

- □ Member has been previously approved for a CGM or insulin pump device
- □ At least <u>ONE</u> of the following problems have occurred which limits the use of the member's current CGM or insulin pump device
  - □ Abuse of equipment
  - □ Misuse of equipment
  - □ 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 or insulin pump device is not under warranty, including the date of warranty expiration

### Not all drugs may be covered under every Plan

If a drug is non-formulary on a Plan, documentation of medical necessity will be required. \*\*Use of samples to initiate therapy does not meet step edit/ preauthorization criteria.\*\* \*<u>Previous therapies will be verified through pharmacy paid claims or submitted chart notes.</u>\*