

**Continuous Glucose Monitor (CGM) Prior Authorization Form**

Member Name: \_\_\_\_\_ Date of Birth: \_\_\_\_\_ Member ID#: \_\_\_\_\_

**System Information**

<input type="checkbox"/> <b>Dexcom G5®</b> Receiver NDC: _____ Transmitter NDC: _____ Sensor NDC: _____	<input type="checkbox"/> <b>Dexcom G6®</b> Receiver NDC: _____ Transmitter NDC: _____ Sensor NDC: _____	<input type="checkbox"/> <b>Freestyle Libre®</b> (age 18 years & older) Reader NDC: _____ Sensor NDC: _____
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<b>Please indicate quantity:</b> Sensor qty: _____ per _____ days Transmitter qty: _____ per _____ days	<b>Please indicate quantity:</b> Sensor qty: _____ per _____ days Transmitter qty: _____ per _____ days	<b>Please indicate quantity:</b> Sensor qty: _____ per _____ days
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**Billing Provider Information**

Pharmacy NPI: \_\_\_\_\_ Pharmacy Name: \_\_\_\_\_  
Fill Date: \_\_\_\_\_ Pharmacy Phone: \_\_\_\_\_ Pharmacy Fax: \_\_\_\_\_

**Prescriber Information**

Prescriber NPI: \_\_\_\_\_ Prescriber Name: \_\_\_\_\_  
Prescriber Phone: \_\_\_\_\_ Prescriber Fax: \_\_\_\_\_

**Clinical Information**

Page 1 of 2 - Please complete and return all pages. Failure to complete all pages will result in processing delays.  
**For Initial Authorization:**

1. Please indicate diagnosis:
  - Type I diabetes mellitus (T1DM) meeting the criteria of American Diabetes Association (ADA) Standards of Medical Care in Diabetes, 2019
  - Other: \_\_\_\_\_
2. Has member been using self-monitoring blood glucose (SMBG; finger sticks)? Yes \_\_\_\_\_ No \_\_\_\_\_
3. Has member been performing frequent blood glucose testing (≥4/day)? Yes \_\_\_\_\_ No \_\_\_\_\_
4. Please indicate how member is receiving insulin therapy:
  - a. Is member insulin-treated with multiple daily injections (≥3/day)? Yes \_\_\_\_\_ No \_\_\_\_\_
  - b. Is member using insulin pump therapy? Yes \_\_\_\_\_ No \_\_\_\_\_
5. Does member's insulin treatment regimen require frequent adjustment by the member or provider on the basis of SMBG or continuous glucose monitoring (CGM) testing results? Yes \_\_\_\_\_ No \_\_\_\_\_
6. In the past 6 months, has member experienced 2 or more Level 2 hypoglycemic episodes [glucose <54mg/dL (3.0mmol/L)] in spite of appropriate therapy? Yes \_\_\_\_\_ No \_\_\_\_\_
  - a. If "Yes" to Question 6 above, please provide the following:
    - i. Glucose: \_\_\_\_\_ mg/dL Date Taken: \_\_\_\_\_
    - ii. Glucose: \_\_\_\_\_ mg/dL Date Taken: \_\_\_\_\_
7. In the past 6 months, has member experienced 1 Level 3 glucose episode (severe event characterized by altered mental and/or physical status requiring assistance as a result of hypoglycemia or ketoacidosis, hyperglycemia) in spite of appropriate therapy? Yes \_\_\_\_\_ No \_\_\_\_\_
  - a. If "Yes" to Question 7 above, please describe: \_\_\_\_\_
8. Has the treating practitioner had an in-person or telehealth visit with the member and/or family within in the 6 months prior to ordering the CGM to evaluate their diabetes control and determined that the above criteria are met? Yes \_\_\_\_\_ No \_\_\_\_\_
9. Has the member and/or family member participated in age-appropriate diabetes education, training, and support prior to beginning CGM? Yes \_\_\_\_\_ No \_\_\_\_\_

PLEASE PROVIDE THE INFORMATION REQUESTED AND RETURN TO:

University of Oklahoma College of Pharmacy  
Pharmacy Management Consultants  
Product Based Prior Authorization Unit  
Fax: 1-800-224-4014  
Phone: 1-800-522-0114 Option 4

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State of Oklahoma  
Oklahoma Health Care Authority  
Continuous Glucose Monitor (CGM) Prior Authorization Form

Member Name: \_\_\_\_\_ Date of Birth: \_\_\_\_\_ Member ID#: \_\_\_\_\_

**Clinical Information**

Page 2 of 2 - Please complete and return all pages. *Failure to complete all pages will result in processing delays.*

**For Continued Authorization:**

1. Has member been seen at least every 6 months following the initial prescription of the continuous glucose monitoring (CGM), by the CGM prescriber, to assess adherence to their CGM regimen and diabetes treatment plan? Yes \_\_\_ No \_\_\_
2. Has member received ongoing instruction and regular evaluation of technique, results, and their ability to use data from self-monitoring of blood glucose to adjust therapy? Yes \_\_\_ No \_\_\_
3. Do the member's prescriber records include documentation (i.e. trend graphs or CGM reports) demonstrating member's daily use of the CGM? Yes \_\_\_ No \_\_\_
4. Does member continue to meet **Initial Authorization** criteria #1-5 (including criteria #3 when CGM is not being utilized)? Yes \_\_\_ No \_\_\_

**Prescriber Signature:** \_\_\_\_\_ **Date:** \_\_\_\_\_  
(By signature, the physician confirms the criteria information above is accurate and verifiable in patient records.) *Please do not send in chart notes. Specific information/documentation will be requested if necessary.*

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