

PATIENT INFORMATION

Name: _____ Date of Birth: _____
Primary Phone: _____ Secondary Phone: _____
Address: _____ City/State/Zip: _____
Primary Insurance: _____ ID# _____
Secondary Insurance: _____ ID# _____
Contact person (if other than patient): _____

REQUIRED INFORMATION FOR ALL PATIENTS

PLEASE COMPLETE THIS SECTION IN FULL

Date of Last Office Visit: _____ Duration of Need (12 months unless other wise noted): _____
Type 1— IDDM Type 2—Pills, Diet, and/or Insulin Treated HbA1c: _____
E10.9 ___ E11.9 **Currently using a pump?** YES NO
E10.65 ___ E11.65 **Currently on CGM Therapy?** YES NO
Other _____ **Other** _____ **Fasting Hyperglycemia:** _____
Testing Frequency: _____ **Fluctuation of blood glucose values:**
Using insulin shots to control? Yes No **Low:** _____ **High:** _____
Number of injections a day: _____

- | | |
|--|---|
| <input type="checkbox"/> Test Strips | <input type="checkbox"/> Transparent Tape |
| Product Name: _____ | <input type="checkbox"/> Prep Wipes |
| <input type="checkbox"/> Lancets | <input type="checkbox"/> Adhesive Remover |
| Product Name: _____ | <input type="checkbox"/> Infusion Sets |
| <input type="checkbox"/> Lancing Device PRN | <input type="checkbox"/> Reservoirs |
| <input type="checkbox"/> Ketostix | <input type="checkbox"/> Omnipod Insulin Pump PDM |
| <input type="checkbox"/> Alcohol Wipes | <input type="checkbox"/> Omnipod Pods |
| <input type="checkbox"/> Meter Battery PRN | <input type="checkbox"/> Other: _____ |
| <input type="checkbox"/> Control Solution PRN | Other: _____ |
| <input type="checkbox"/> Meter PRN | |

CGM Products

DEXCOM G6

Receiver: Dispense 1: DME Only : 1/5 years
Sensors: Quantity 13 boxes (units): DME Only: 9 units/90 days
Transmitter Dexcom G6 (3 month use): 4/365 Days

FREESTYLE LIBRE 2

Receiver: Dispense 1: DME Only: 1/5 years
Sensors: Quantity 26.00 units: 6 units/90 days

USE PER MANUFACTURER RECOMMENDATIONS

My signature below denotes to the best of my knowledge the patient/caregiver is able to follow instructions for controlling diabetes and is able to use the ordered items which are designed for home use. The patient/caregiver has successfully completed training or is scheduled to begin training in the use of supplies or equipment ordered. I am a provider who manages patients with diabetes, insulin pump, or CGM therapy and work closely with a team including nurses, diabetic instructors, and dietitians who are knowledgeable in the use of subcutaneous insulin infusion therapy.

Physician Staff Contact:

Provider Name (print): _____ **Provider NPI #** _____
Provider Phone Number: _____ **Provider Fax:** _____
Provider Signature: _____ **Date:** _____

CLINICAL DOCUMENTATION CHECKLIST FOR FLORIDA MEDICAID WELLCARE/STAYWELL

The following must be completed and signed by the physician with supporting documentation.

Therapeutic Continuous Glucose Monitors are considered medically necessary when all of the following criteria are met. Please check all that apply and include supporting documentation.

- 1. The member has diabetes mellitus (type 1DM and type 2DM); **AND,**
- 2. The member requires twice day BGM (for device calibration for DexCom G5 mobile); **AND,**
- 3. The member is insulin-treated with multiple daily injections (MDI) of insulin or a continuous subcutaneous insulin infusion (CSII) pump; **AND,**
- 4. The member's insulin treatment regimen requires frequent adjustment by the member on the basis of therapeutic continuous glucose monitoring (CGM) testing results, **AND,**
- 5. The member requires insulin injections 3 or more times per day or a medically necessary insulin pump is used for maintenance of blood sugar control.
- 6. A comprehensive glucose level log is maintained, documenting significant changes in diabetic management as a result of the continuous monitoring

Pregnant Members- Please check all that apply and send accompanying documentation

Continuous (long term) monitoring of glucose levels in interstitial fluid in during pregnancy is considered medically necessary if ALL of the below criteria are met.

- 1. The member has diabetes mellitus; **AND,**
- 2. The member has been using a home blood glucose monitor (BGM) and performing frequent (four or more times a day) BGM testing; **AND,**
- 3. The member is insulin-treated with multiple daily injections (MDI) of insulin or a continuous subcutaneous insulin infusion (CSII) pump; **AND,**
- 4. The member's insulin treatment regimen requires frequent adjustment by the beneficiary on the basis of therapeutic CGM testing results; **AND,**
- 5. The member has used best practices and was compliant with 4 or more finger sticks per day; **AND,**
- 6. Continuous glucose monitoring has led to a beneficial series of behavioral modifications resulting in a reduction of hypoglycemic events; **AND,**
- 7. Insulin injections are required 3 or more times per day or a medically necessary insulin pump is used for maintenance of blood sugar control; **AND,**
- 8. A comprehensive glucose level log is maintained, documenting significant changes in diabetic management as a result of the continuous monitoring.

External insulin pumps are considered medically necessary if the Member:

- 1. Has a diagnosis of insulin dependent type I diabetes mellitus; **AND**
- 2. Completed a comprehensive diabetes education program. (This may include, but is not limited to, leading the Member to demonstrate the ability and commitment to comply with the regimen of pump care, frequent self-monitoring of blood glucose, and careful attention to diet and exercise, and has received appropriate training on pump usage); **AND**
- 3. Has been on a program of multiple daily injections of insulin (e.g., at least 3 injections per day) with frequent self-adjustments of insulin dose for at least 6 months prior to initiation of the insulin pump; **AND**
- 4. Member has been on an external insulin infusion pump prior and has documented frequency of glucose self-testing an average of at least 4 times per day during the month prior to enrollment; **AND**
- 5. Has an endocrinologist or physician experienced in providing insulin infusion therapy orders the insulin pump and states that he/she will monitor the members status while he/she uses the pump; **AND**
- 6. Has Provider documentation of a history of poor glycemic control on multiple daily injections of insulin, including a persistently elevated glycosylated hemoglobin level (HBA1C>7.0%). Additional history of poor control may be documented in the medical record, including but not limited to:
 - Widely fluctuating blood glucose levels before bedtime **OR**
 - History of severe hypoglycemia (<60mg/dL) or hyperglycemia (>300 mg/dL) **AND/OR**
 - Treatment of secondary diabetic complications requiring more extensive blood glucose control

I certify that this treatment is indicated and necessary and meets the guidelines for use as outlined by Staywell/Wellcare Florida Medicaid. I will be supervising the patient's treatment. Required supporting documentation from the patient's medical record is attached.

Prescribing Practitioner Signature (Required)

Date

Stamps/copies of the physician's signature will not be accepted