#### PATIENT INFORMATION

Name:	_ Date of Birth
Primary Phone:	Secondary Phone:
Address:	City/State/Zip:
Primary Insurance:	ID#
Contact Person:	Email:
<b>REQUIRED INFORMATION FOR ALL PATIEN</b>	TS - PLEASE FULLY COMPLETE THIS SECTION
Date of Last Office Visit: Duration of Need: mo. (12 unless noted)	
Type 1 -IDDM E 10.9 E 10.65 E 10.649 Other:	
Type 2- Pills, Diet, and/or Insulin Treated E 11.9 E 11.65 Other:	
Testing Frequency: X per Day Using Insulin treatment to control? Yes No	
Number of insulin treatments: X Per Day A1c: Currently Using a Pump? Yes No	
Currently on CGM Therapy? YesNo Fasting Hyp	perglycemia:
Fluctuation of Blood Glucose Values: Low High	
PRODUCTS PRN- USE PER MANL	JFACTURERS RECOMMENDATION
Testing Supplies: Glucometer, test strips, lancing device	e, lancets, ketone strips, control solution, alcohol wipes
CGM, Dexcom G6: Sensors, transmitter, receiver, prep wipes, adhesive remover, dressing	
CGM, Dexcom G7: Sensors (transmitter included), rece	iver, prep wipes, adhesive remover, dressing
CGM,FreeStyle Libre 2 orFreeStyle Libre 3: S	ensors, reader, prep wipes, adhesive remover, dressing
Insulin Pump: Tandem T-slim Control IQ Ta	ndem T-slim Basal IQiLet by Beta Bionics
Medtronic MiniMed 630	Medtronic MiniMed 770G
Insulin Pump Supplies: Reservoirs, infusion sets, prep wipes, adhesive remover, dressing	
Omnipod 5 Starter Kit Omnipod 5 Pods: Pod (5 p	er box), prep wipes, adhesive remover, dressing
Omnipod Dash Supplies: Pods, prep wipes, adhesive re wipes, adhesive remover, dressing- PRN	mover, dressing Omnipod Eros Supplies: Pods, prep
Other:	
and is able to use the ordered items, which are designed for hom respond as needed. The parent/caregiver has successfully comple equipment ordered. I am a provider who manages patients with team including nurses, diabetic instructors, and dietitians knowled	eted training or is scheduled to begin training in using supplies or diabetes, insulin pump, or CGM therapy and works closely with a edgeable in the use of subcutaneous insulin infusion therapy. For nd is followed by our clinic. I am writing to support the continued
Provider Name (Print):	Provider NPI:
Provider Phone Number:	Provider Fax:
Provider Signature: 500 Patriot Pkwy, Ste B, Tuscaloosa, AL 35405 P	Date: hone: 866-919-1246
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#### Criteria Checklist MUST ACCOMPANY THE PRIOR AUTHORIZATION FORM



# Medicaid Agency/Commerical Insurance External Ambulatory Insulin Infusion Pump (E0784)

AIM Plus Medical Supplies 866-919-1246

Children under 21 years of age and EPSDT eligible

### **PREREQUISITE CRITERIA** The patient **must** meet all of the following:

- □ Patient must be Medicaid eligible, less than 21 years of age, and EPSDT eligible.
- □ Patient must have a documented\* diagnosis of insulin dependent diabetes mellitus (IDDM, also known as type I).
- A board certified endocrinologist must have evaluated the patient and ordered the insulin pump.
- Patient must have been on a program of multiple daily injections (MDI) of insulin (i.e., at least three injections per day) for at least six months prior to initiation of the insulin infusion pump. Supporting documentation\* must be submitted.
- Patient has documented frequency of glucose self-testing (i.e. patient "logs") an average of at least four times per day during the three months prior to initiation of the insulin pump. Patients in Alabama must include six consecutive weeks' worth of logs within the three months prior to the prior authorization request.
- Patient and/or caregiver must be capable, physically and intellectually, of operating the pump. Patient/caregiver must demonstrate ability and commitment to comply with regimen of pump care, diet, exercise, medications, and

glucose testing at least four times a day. Supporting documentation\* must be submitted.

- Education on insulin pump MUST have been conducted prior to prior authorization request, and each the patient, caregiver if child, and educator signed to document\* their understanding.
- Documentation\* of active and past recipient compliance with medications and diet, appointments, and other treatment recommendations must be provided.

### ADDITIONAL CRITERIA The patient must also meet one or more of the following, supported by documentation\*:

- □ Two elevated glycosylated hemoglobin levels (HbA1c> 7.0%) within a 120-day time span, while on multiple daily injections of insulin.
- □ History of severe glycemic excursions (commonly associated with brittle diabetes, hypoglycemic unawareness, nocturnal hypoglycemia, extreme insulin sensitivity and/or very low insulin requirements).
- □ Widely fluctuating blood glucose levels before mealtime (i.e., pre-prandial blood glucose level consistently exceeds 140 mg/dL).
- Dawn phenomenon with fasting blood sugars frequently exceeding 200 mg/dL.

### **DIAGNOSIS CODES**

Approval will be given for only type I diabetes mellitus diagnosis codes. Please refer to Chapter 14 of the Provider Manual for the ICD-10 crosswalk codes.

### **PROCEDURE CODES**

E0784, A4221, A4232, A4230, A9274

Maximum yearly limits apply to each of the procedure codes indicated above. Requests for replacement of E0784 will be limited to once every five years based on a review of submitted documentation requested.

\*Documentation may include notes from the patient chart and/or pharmacy printouts (to support medication compliance history).

Please ask the endocrinologist prescribing the insulin pump to sign below. This form should be returned to AIM Plus Medical Supplies by email at documents@aimplusonline.com or fax to 866-496-7054 to the attention of the medical device dept.

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

Prescribing Practitioner Signature (Required) (Stamps/copies of physician's signature will not be accepted)

Patient's Name

Date of Birth

# Medicaid Agency/Commerical Insurance Continuous Glucose Monitoring

### **PREREQUISITE CRITERIA** All of the following **must** be met with supporting documentation\*:

- □ Patient is a child diagnosed with \*Type 1 diabetes mellitus or pregnant female (Type 1 or 2); and \*For patients in MS, poorly controlled Type 1 diabetes is defined as:
  - a) Unexplained hypoglycemic episodes,
  - b) Nocturnal hypoglycemic episode(s),

c) Hypoglycemic unawareness and/or frequent hypoglycemic episodes leading to impairments in activities of daily living,

- d) Suspected postprandial hyperglycemia,
- e) Recurrent diabetic ketoacidosis, or

f) Unable to achieve optimum glycemic control as defined by the most current version of the American Diabetes Association (ADA).

- □ Patient is insulin-treated with multiple (three or more) daily injections of insulin or a Medicaid-covered continuous subcutaneous insulin infusion (CSII) pump.
- Patient's insulin treatment regimen requires frequent adjustment by the patient and/or caregiver on the basis of BGM or CGM testing results (AL Medicaid patients only).
- □ Within six (6) months prior to ordering the CGM, the treating practitioner has an in-person visit with the patient to evaluate their diabetes control (to include HbA1c) and determined that criteria (1-4) above are met.
- □ Every six (6) months following the initial prescription of the CGM, the treating practitioner has an inperson visit with the patient to assess adherence to their CGM regimen and diabetes treatment plan.
- □ For patients on Mississippi Medicaid, there is an additional requirement of documentation proving selfmonitoring of blood glucose at least four times a day.

# **RECERTIFICATION/RENEWAL:**

For patients who have received CGM equipment and supplies through AL Medicaid and are in need of a Prior Authorization Renewal, an updated prescription and an attestation from the patient's prescribing provider, stating their recommendation for continued CGM therapy, is required. A request for replacement of the Receiver (A9278) will be considered for approval every five years upon review of submitted medical documentation. If a replacement request is submitted within less than five years and the replacement is due to a natural disaster and not the result of misuse, neglect or malicious acts by the user, the request may be considered for approval and payment.

### Limitations

Approval will be given for only type I diabetes mellitus diagnosis codes. Please refer to Chapter 14 of Provider Manual for the ICD-10 crosswalk codes.

# **PROCEDURE CODES**

A9276, A9277, A9278

Maximum limits apply to each of the procedure codes indicated above. Requests for replacement of A9278 will be limited to once every five years based on a review of submitted documentation requested.

This checklist is provided for informational purposes in order to guide you in submitting supporting documentation. The checklist does not have to be signed by a provider or returned to AIM Plus Medical Supplies. Please email supporting documents to documents@aimplusonline.com or fax them to 866-496-7054 to the attention of the medical device team. Thank you for your referral. We will take excellent care of your patient!