Field Name	Field Description
Prior Authorization	Inculin Dumps
Group Description	Insulin Pumps
Drugs	Omnipod Dash, insulin delivery pods only
	(Notes: The Omnipod Dash PDM (Personal Diabetes Manager) is
	provided direct by Insulet and should not be requested by the
	prescriber/billed to the plan.)
	This policy does not apply to pumps reviewed and/or covered by
	the Medical Benefit including, but not limited to V-Go 24-hour
	disposable system and t:slim X2, and continuous glucose
	monitor/insulin pumps such as MiniMed. Requests for these
	products are referred to the plan's Utilization Management team
	for review.
Covered Uses	Medically accepted indications are defined using the following sources:
	the Food and Drug Administration (FDA), Micromedex, American
	Hospital Formulary Service (AHFS), United States Pharmacopeia Drug
	Information for the Healthcare Professional (USP DI), the Drug
	Package Insert (PPI), or disease state specific standard of care
	guidelines.
Exclusion Criteria	None
Required Medical	See "Other Criteria"
Information	
Age Restrictions	None
Prescriber	Prescribed by or in consultation with an endocrinologist, <u>a certified</u>
Restrictions	diabetic educator, or an obstetrician/gynecologist
Coverage Duration	If all of the criteria are met, the request will be approved for 12 months.
	If the criteria are not met, the request is referred to a Medical
	Director/Clinical Reviewer for medical necessity review.
Other Criteria	Initial Authorization
	• Diagnosis – diabetes
	<ul> <li>One of the following</li> <li>○ ≤ 18 years with type 1 diabetes or other insulin-deficient forms</li> </ul>
	$\circ \leq 18$ years with type 1 diabetes or other insulin-deficient forms of diabetes (i.e. cystic-fibrosis related diabetes)
	<ul> <li>Continuation of therapy for patient new to plan</li> </ul>
	• Treatment with multiple daily doses ( $\geq 3$ ) of insulin and one of
	the following
	• Persistently inadequate glycemic control (i.e. $HbA1C \ge$
	7% on multiple consecutive readings with one being within the last 3 months, frequent houts of hypoglycemia
	within the last 3 months, frequent bouts of hypoglycemia,

	<ul> <li>overt microvascular complications)</li> <li>History of acutely dangerous symptoms (i.e. severe glycemic excursions; brittle diabetes; nocturnal hypoglycemia; hypoglycemia unawareness, ketosis)</li> <li>Other difficult to manage symptoms/scenarios (i.e. "dawn" phenomenon; extreme insulin sensitivity; very low insulin requirements)</li> <li>Pregnancy</li> </ul>
	<ul> <li>One of the following:         <ul> <li>Child or adolescent with type 1 diabetes or other insulin- deficient form of diabetes</li> <li>Documentation of positive clinical response (i.e. improved HbA1C; reduced frequency of severe hypoglycemia episodes; target time in range [TIR] &gt; 70% or time below range &lt; 4%) with 1<sup>st</sup> reauthorization</li> <li>Initial approval was based on continuation of therapy for patient new to plan.</li> </ul> </li> </ul>
	• Continuation of therapy based on a diagnosis of pregnancy alone is not eligible for reauthorization
Revision/Review Date <u>5/20/22</u> <del>10/20/21</del>	Medical Director/clinical reviewer must override criteria when, in his/her professional judgement, the requested item is medically necessary.