





FAX Completed Form To 1.833.404.2392

## Prescriber Help Desk 1.833.587.2012

Online

covermymeds.com/main/ prior-authorization-forms/

## Request for Prior Authorization GLP-1 Agonist/Basal Insulin Combinations

(PLEASE PRINT – ACCURACY IS IMPORTANT)

IA Medicaid Member ID #	A Medicaid Member ID # Patient name			DOB	
Patient address					
Provider NPI	Prescriber name		Phone		
Prescriber address			Fax		
Pharmacy name	Address		Phone		
Prescriber must complete all information above. It must be legible, correct, and complete or form will be returned.					
Pharmacy NPI	Pharmacy fax		NDC		
Prior authorization is required for GLP-1 agonist receptor/basal insulin combination products. Payment will be considered for patients when the following criteria are met:  1) A diagnosis of Type 2 Diabetes Mellitus, and					
2) Patient is 18 years of age or older; and					
<ol> <li>The patient has not achieved HgbA1C goals after a minimum three month trial with metformin at a maximally tolerated dose, unless evidence is provided that use of this agent would be medically contraindicated; and</li> </ol>					
<ul> <li>4) Documentation of an adequate trial and inadequate response with at least one preferred GLP-1 receptor agonist and one preferred long-acting insulin agent concurrently; and</li> <li>5) Will not be used concurrently with prandial insulin; and</li> </ul>					
<ol> <li>Clinical rationale is provided as to why the patient cannot use a preferred GLP-1 receptor agonist and a preferred long-acting insulin agent concurrently; and</li> </ol>					
<ul> <li>7) Medication will be discontinued and alternative antidiabetic products will be used if patients require a daily dosage of:</li> <li>a) Soliqua below 15 units or over 60 units, or</li> <li>b) Xultophy persistently below 16 units or over 50 units.</li> </ul>					
Non-Preferred					
☐ Soliqua ☐ Xultophy					
Strength	Dosage Instruction		uantity	Day's Supply	
Diagnosis:					
Most Recent HgbA1C Level: Date this level was obtained:					
Metformin Trial: Trial start date:		Trial end date:		Trial dose:	
Reason for failure:					
Medical or contraindication reason to override trial requirements:					

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Preferred GLP-1 Receptor Agonist Trial: Drug	name/dose:			
Trial start date:	Trial end date:			
Reason for failure:				
Preferred Long-Acting Insulin Trial: Drug nam	e/dose:			
Trial start date:	_ Trial end date:	Trial end date:		
Reason for failure:				
Clinical rationale as to why patient cannot use acting insulin agent concurrently:				
Is prandial insulin being used concurrently?	☐ Yes	□ No		
Medication will be discontinued and alternative daily dosage of:	e antidiabetic prod	ducts will be used if patients require a		
☐ Soliqua – below 15 units or over 60 units	☐ Yes	□ No		
☐ Xultophy – persistently below 16 units or over	50 units 🗌 Yes	□ No		
Attach lab results and other documentation as nec	<u> </u>			
Prescriber signature (Must match prescriber listed above.	.)	Date of submission		

**IMPORTANT NOTE:** In evaluating requests for prior authorization the consultant will consider the treatment from the standpoint of medical necessity only. If approval of this request is granted, this does not indicate that the member continues to be eligible for Medicaid. It is the responsibility of the provider who initiates the request for prior authorization to establish by inspection of the member's Medicaid eligibility card and, if necessary by contact with the county Department of Human Services, that the member continues to be eligible for Medicaid.

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