





Fax Completed Form To 1.833.404.2392

**Prescriber Help Desk** 1.833.587.2012

covermymeds.com/main/

Online

## **Request for Prior Authorization Incretin Mimetics for Non-Diabetes Indications**

	(PLEASE PRINT – ACCURACY IS IMPORTA	ANT)	prior-authorization-forms/		
IA 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 (PA) is required for incretin mimetics not otherwise covered by the Anti-Diabetics Non-Insulin Agents PA criteria for covered FDA approved for compendia indications. Payment for excluded medical use(s) (e.g. weight loss), as defined in the Iowa State Plan and Iowa Administrative Code 441 – 78.2(4) will be denied. Payment will be considered under the following conditions:

- Reguest adheres to all FDA approved labeling for requested drug and indication, including dosing, contraindications, warning and precautions, drug interactions, and use in specific populations; and
- 2. Patient has been screened for and does not have type 1 or type 2 diabetes mellitus (attach current lab results, obtained within 6 months of request, documenting an A1C < 6.5% or a fasting plasma glucose < 126 mg/dL); and
- The requested drug will be used to reduce the risk of major adverse cardiovascular events (MACE) (cardiovascular death, non-fatal myocardial infarction, or non-fatal stroke) in an adult with established cardiovascular disease (CVD) and either obesity or overweight; and
  - a. Patient has established CVD with history of one of the following (attach chart notes documenting diagnosis):
    - i. Prior myocardial infarction (MI);
    - ii. Prior stroke (ischemic or hemorrhagic);
    - iii. Symptomatic peripheral arterial disease (PAD), as evidenced by intermittent claudication with ankle-brachial index (ABI) less than 0.85 (at rest), peripheral arterial revascularization procedure, or amputation due to atherosclerotic disease; and
  - b. Patient has a baseline body mass index (BMI) ≥ 27kg/m², obtained within 6 months of request; and
  - Patient has been evaluated for cardiovascular standard of care treatment; and
  - d. For Wegovy:
    - i. Patient is ≥ 45 years of age; and
    - ii. Initiation and escalation dosages will be permitted for a maximum of 8 weeks for each dosage; and
    - iii. Maintenance dosages other than 1.7 mg or 2.4 mg once weekly will not be approved for maintenance treatment; or
- Patient has a diagnosis of moderate to severe obstructive sleep apnea (OSA); and
  - Patient has a baseline BMI ≥ 30kg/m<sup>2</sup>; and a.
  - Prescriber attests patient has a recent (within prior three years) apnea/hypopnea index (AHI) ≥ 15 events b. per hour, as documented by a polysomnography (PSG) or at-home sleep study (document AHI); and
  - For Zepbound: C.
    - i. Patient meets the FDA approved age for OSA; and
    - ii. Initiation and escalation dosages will be permitted up to a maximum of 20 weeks prior to reaching the recommended maintenance dosage of 10 mg to 15 mg once weekly; and

(7/25)Page 1 of 3







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 Incretin Mimetics for Non-Diabetes

(PLEASE PRINT - ACCURACY IS IMPORTANT)

- iii. Maintenance dosages other than 10 mg to 15 mg once weekly will not be approved for maintenance treatment; and
- 5. Patient will use medication in combination with a reduced calorie diet and increased physical activity; and
- 6. The requested agent will not be used in combination with other incretin mimetics.

The required trials may be overridden when documented evidence is provided that use of these agents would be medically contraindicated.

Requests will be considered for initiation and appropriate dosage escalation. Requests for continuation of therapy, once at an established maintenance dose, will be considered when:

- 1. The requested drug will be used to reduce the risk of MACE; and
  - a. Patient has been evaluated for cardiovascular standard of care treatment; and
  - b. For Wegovy, a maintenance dose of 1.7 mg or 2.4 mg weekly is requested; or
- 2. The requested drug will be used to treat moderate to severe OSA; and
  - a. Documentation of a positive response to therapy is provided; and
  - b. The maintenance dose is requested and maintained (Zepbound 10 mg to 15 mg once weekly); and
- 3. Patient does not have type 1 or type 2 diabetes; and
- 4. Patient continues to use medication in combination with a reduced calorie diet and increased physical activity; and
- 5. The requested agent will not be used in combination with other incretin mimetics.

<u>Preferred</u>	Non-Preferred						
Zepbound	☐ Wegovy						
Strength	<b>Usage Instructions</b>	Quantity	Day's Supply				
Diagnosis:							
Initial Requests:  Does patient have Type 1 or Type 2 Diabetes (attach lab results documenting current A1C or fasting plasma glucose)?							
Requests for Wegovy:							
Patient has established (diagnosis):  Prior myocardia	CVD documented by one of the fo	llowing (attach cha	rt notes documenting				

(7/25) Page 2 of 3







#### Fax Completed Form To 1.833.404.2392

# **Prescriber Help Desk**

Online covermymeds.com/main/

1.833.587.2012

### **Request for Prior Authorization Incretin Mimetics for Non-Diabetes Indications**

(PLEASE PRINT - ACCURACY IS IMPORTANT)

	(PLEASE PRINT – ACCURAC	Y IS IMPOR'	TANT)	prior-authorization-forms/			
	Prior stroke (ischemic or hemorrhagic)		·				
	Symptomatic PAD, as evidenced by:						
ш	Intermittent claudication with ABI less than 0.85	(at rest) or					
		` ,					
	Peripheral arterial revascularization procedure, o	ı					
	Amputation due to atherosclerotic disease						
Provide p	patient's baseline BMI:	Da	ate Obtained:				
Has patie	ent been evaluated for cardiovascular standard of	care treat	ment?	☐ No			
Requests	s for Zepbound:						
Provide <sub>I</sub>	patient's baseline BMI:	Da	ate Obtained:				
D		- //	- !  (ALII) > 4	F			
	ient have a recent (within prior three years) apne nented by a PSG or at-home sleep study?	a/nypopne	a index (AHI) ≥ 1	<u>5 events per nour,</u>			
∐ Yes	Document AHI:	. No	)				
Renewal	Requests:						
-	ient have Type 1 or Type 2 Diabetes (attach lab re lucose)?	esults docu	menting current	A1C or fasting			
	ontinues to use medication in combination with a	a reduced o	calorie diet and i	ncreased physical			
Will the requested agent be used in combination with other incretin mimetics?   Yes   No							
Wegovy:							
Has patient been evaluated for cardiovascular standard of care treatment?   Yes   No							
Zepboun	d:						
Document positive response to therapy:							
————Attach lab	results and other documentation as necessary.						
Prescriber	signature (Must match prescriber listed above.)		Date of submission	٦			

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 Health and Human Services, that the member continues to be eligible for Medicaid.

(7/25)Page 3 of 3