





Fax Completed Form To 1.833.404.2392

1.833.587.2012

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Prescriber Help Desk

Request for Prior Authorization ORAL CONSTIPATION **AGENTS**

	(PLEASE PRINT - ACCURACY IS IMPOR	(TANT) <u>prior-authorization-forms</u>				
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 oral constipation agents subject to clinical criteria. Payment for nonpreferred oral constipation agents will be authorized only for cases in which there is documentation of a previous trial and therapy failure with a preferred oral constipation agent. Payment will be considered when patient has an FDA approved or compendia indication for the requested drug when the following criteria are

contraindications, warnings and precautions, drug interactions, and use in specific populations; and						
2) Patien		ntation of adequate trials and there	herapy failures with	both of the following:		
 i. Stimulant laxative (senna) plus saline laxative (milk of magnesia); and ii. Stimulant laxative (senna) plus osmotic laxative (polyethylene glycol or lactulose); or b. Members 17 years of age or younger: i. Polyethylene glycol; and ii. One other preferred generic laxative, such as lactulose or senna; and 						
3) Patient does not have a known or suspected mechanical gastrointestinal obstruction.						
treatment.	Requests for contin	net, initial authorization will be uation therapy may be provided s to meet the age for indication	d if the prescriber d			
Preferred Linzess	s ☐ Lubiprostor	ne				
Non-Prefe	rred	rucalopride	Symproic 🗌 Trula	nce		
	Strength	Dosage Instructions	Quantity	Days Supply		
Treatmen	nt failures:					
Members	18 years of age an	d older:				
Trial 1: St	timulant Laxative (s	senna) plus Osmotic Laxativ	e (polyethylene g	lycol / lactulose)		
Stimulant Laxative Trial: Name/Dose:				_Trial Dates:		
Failure rea	ason:					

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Trial 2: Stimulant Laxative (senna) plus Saline Laxative (milk of magnesia) Stimulant Laxative Trial: Name/Dose: Trial Dates: Failure reason: Members 17 years of age and younger: Polyethylene Glycol Trial: Name/Dose:______ Trial Dates:_ Failure reason: Additional Preferred Generic Laxative Trial: Name/Dose:______Trial Dates: Failure reason: Does patient have a known or suspected mechanical gastrointestinal obstruction: \(\square \) Yes □ No Chronic Idiopathic Constipation: (Linzess, Lubiprostone, Motegrity or Trulance) • Patient has less than 3 spontaneous bowel movements (SBMs) per week: ☐ Yes ☐ No Patient has two or more of the following symptoms within the last 3 months: Straining during at least 25% of the bowel movements Lumpy or hard stools for at least 25% of bowel movements Sensation of incomplete evacuation for at least 25% of bowel movements • Documentation the patient is not currently taking constipation causing therapies: Medication review completed: ☐ Yes ☐ No Current constipation causing therapies: ☐ Yes (please list) □ No Irritable Bowel Syndrome with Constipation: (Ibsrela, Linzess, Lubiprostone, or Trulance) Patient is female (Lubiprostone requests only): \(\begin{align*} \text{Yes} & \quad \text{No} \end{align*} \) Patient has recurrent abdominal pain on average at least 1 day per week in the last 3 months associated with two (2) or more of the following: Related to defecation Associated with a change in stool frequency Associated with a change in stool form Opioid-Induced Constipation with Chronic, Non-Cancer Pain: (Lubiprostone, Movantik, Relistor, or Symproic) • Patient has been receiving stable opioid therapy for at least 30 days as seen in the patient's pharmacy claims: Yes No Patient has less than 3 spontaneous bowel movements (SBMs) per week, with at least 25% associated with one or more of the following: Hard to very hard stool consistency Moderate to very severe straining Sensation of incomplete evacuation **Functional Constipation:** (Linzess) Patient has less than 3 spontaneous bowel movements (SBMs) per week: • Patient has one or more of the following criteria at least once per week for at least 2 months: History of stool withholding or excessive voluntary stool retention History of painful or hard bowel movements History of large diameter stools that may obstruct the toilet Presence of a large fecal mass in the rectum At least one episode of fecal incontinence per week

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Other Diagnosis:					
Renewal Requests: Provide documentation of adequate response to treatment:					
Requests for Non-Preferred Oral Constipation Agent: Document trial of preferred agent					
Drug Name/Dose:	Trial Dates:				
Failure reason:					
Possible drug interactions/conflicting drug therapies:					
Attach lab results and other documentation as necessary.					
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 Health and Human Services, that the member continues to be eligible for Medicaid.

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