SENTARA HEALTH PLANS

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

Directions: The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; fax to <u>1-800-750-9692</u>. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. If information provided is not complete, correct, or legible, authorization may be delayed.

Gastrointestinal (GI) Motility Drugs

Drug Requested: (select one drug below)				
Non-Preferred				
□ Ibsrela [®] (tenapanor)		Motegrity [®] (prucalopride)	□ Relistor [®] (methylnaltrexone	
			bromide)	
Trulance [®] (plecanatide)		Zelnorm [™] (tegaserod)		

MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.

Member Name:	
Member Sentara #:	
Prescriber Name:	
	Date:
Office Contact Name:	
Phone Number:	
DEA OR NPI #:	
DRUG INFORMATION: Auth	norization may be delayed if incomplete.
Drug Form/Strength:	
Dosing Schedule:	Length of Therapy:
Diagnosis:	ICD Code:

CLINICAL CRITERIA: Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.

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□ Approval of Motegrity[®] for diagnosis of Chronic Idiopathic Constipation (CIC)

□ Member has had trial and failure, contraindication, or intolerance to <u>ONE</u> of the following generic prerequisite therapies:

AND

□ Member has had trial and failure, contraindication, or intolerance to lubiprostone (Amitiza[®])

AND

□ Member has had trial and failure, contraindication, or intolerance to Linzess[®]

□ Approval of Zelnorm[™]

D Diagnosis of Irritable Bowel Syndrome with Constipation (IBS-C)

AND

Member is < 65 years of age with no history of ischemic cardiovascular disease and has no more than one CVD risk factor. CVD risk factors are defined as active smoking, current hypertension/history of antihypertensive treatment, current hyperlipidemia/history of lipid lowering medication, history of diabetes mellitus, age >55 years, or obesity (BMI >30 kg/m²)

AND

- □ Provider attests that member does <u>NOT</u> have any of the following contraindications to therapy:
 - History of myocardial infarction (MI), stroke, transient ischemic attack (TIA), or angina
 - History of ischemic colitis or other forms of intestinal ischemia
 - Severe renal impairment (eGFR $< 15 \text{ mL/min}/1.73 \text{ m}^2$) or end-stage renal disease
 - Moderate and severe hepatic impairment (Child-Pugh B or C)
 - History of bowel obstruction, symptomatic gallbladder disease, suspected sphincter of Oddi dysfunction, or abdominal adhesions

AND

□ Member has had trial and failure, contraindication, or intolerance to <u>ONE</u> of the following generic prerequisite therapies:

$\square \text{lactulose} \square \text{polyethylene glycol (generic MiraLAX}^{\textcircled{R}})$
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AND

□ Member has had trial and failure, contraindication, or intolerance to lubiprostone (Amitiza[®])

AND

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□ Member has had trial and failure, contraindication, or intolerance to Linzess[®]

Approval of Trulance[®] for diagnosis of Irritable Bowel Syndrome with Constipation (IBS-C) or Chronic Idiopathic Constipation (CIC)

□ Member has had trial and failure, contraindication, or intolerance to <u>ONE</u> of the following generic prerequisite therapies:

$\Box polyethylene glycol (generic MiraLAX®)$

AND

□ Member has had trial and failure, contraindication, or intolerance to lubiprostone (Amitiza[®])

AND

□ Member has had trial and failure, contraindication, or intolerance to Linzess[®]

Approval of Ibsrela[®] for diagnosis of Irritable Bowel Syndrome with Constipation (IBS-C)

□ Member has had trial and failure, contraindication, or intolerance to <u>ONE</u> of the following generic prerequisite therapies:

□ lactulose	$\Box polyethylene glycol (generic MiraLAX®)$
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AND

□ Member has had trial and failure, contraindication, or intolerance to lubiprostone (Amitiza[®])

AND

□ Member has had trial and failure, contraindication, or intolerance to Linzess[®]

AND

□ Member has had trial and failure, contraindication, or intolerance to Trulance[®] (requires prior authorization)

□ Approval of Relistor[®]

Recommended Dosing:

Weight of Adult Patient	Subcutaneous Dose	Injection Volume
Less than 38kg	0.15 mg/kg	See below
38kg to less than 62 kg	8mg	0.4 mL
62kg to 114kg	12mg	0.6 mL
More than 114kg	0.15 mg/kg	See below

Calculate injection volume by multiplying member weight in kilograms by 0.0075, then round up the volume to the nearest 0.1 mL

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- □ Select <u>ONE</u> of the following:
 - □ Member has a diagnosis of opioid-induced constipation (OIC) with advanced illness or pain caused by active cancer who require opioid dosage escalation for palliative care
 - Member has a diagnosis of opioid-induced constipation (OIC) with chronic non-cancer pain, including members with chronic pain related to prior cancer or its treatment who do not require frequent (e.g., weekly) opioid dosage escalation

AND

Member has been on an opioid within the last 60 days of prior authorization request but not less than 4 weeks. Provider please note: Members receiving opioids for less than 4 weeks may be less responsive to Relistor[®]

AND

□ Member has had trial and failure, contraindication, or intolerance to <u>ONE</u> of the following generic prerequisite therapies:

	\Box polyethylene glycol (generic MiraLAX [®])
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AND

□ Member has had trial and failure, contraindication, or intolerance to lubiprostone (Amitiza[®])

AND

Member has had trial and failure, contraindication, or intolerance to both Movantik[®] AND Symproic[®]

Not all drugs may be covered under every Plan

If a drug is non-formulary on a Plan, documentation of medical necessity will be required.

Use of samples to initiate therapy does not meet step edit/ preauthorization criteria. *<u>Previous therapies will be verified through pharmacy paid claims or submitted chart notes.</u>*