SENTARA COMMUNITY PLAN (MEDICAID)

MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions:</u> 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; <u>fax to 1-844-305-2331</u>. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. <u>If information provided is not complete, correct, or legible, authorization can be delayed</u>.

<u>Drug Requested</u>: Onapgo[™] (apomorphine hydrochloride) (J3490) (Medical)

MEMBER & PRESCRIBER INF	ORMATION: Authorization may be delayed if incomplete.			
Member Name:				
Member Sentara #:				
Prescriber Name:				
Prescriber Signature:	Date:			
Office Contact Name:				
Phone Number:				
NPI #:				
DRUG INFORMATION: Authoriz	cation may be delayed if incomplete.			
Drug Name/Form/Strength:				
Dosing Schedule:	Length of Therapy:			
Diagnosis:	ICD Code, if applicable:			
Weight (if applicable):	Date weight obtained:			
Standard Review. In checking this box, the timeframe does not jeopardize the life or health of the member or the member's ability to regain maximum function and would not subject the member to severe pain.				

Recommended Dosage:

- Initial continuous dosage is 1 mg/hr with a maximum of 6 mg/hr for up to 16 hours per day. An extra dose may be titrated in increments of 0.5 mg or 1 mg based on clinical response and tolerability. Subsequent extra doses are between 0.5 mg and 2 mg, with at least 3 hours between extra doses and a maximum of 3 extra doses per day.
- Maximum recommended total daily dosage of Onapgo, including the continuous dosage and any extra dose(s), is 98 mg (1 cartridge per day) generally administered over the waking day (e.g., 16 hours).

Quantity Limit: 6 cartons (30 cartridges; 600 mL) per 30 days

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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.

Initial Authorization: 12 months

Member must be 18 years of age or older				
Medication must be prescribed by, or in consultation with a neurologist				
Member has a diagnosis of advanced Parkinson's disease (PD) with complicated motor fluctuations				
Member does NOT have a diagnosis of atypical PD or secondary PD				
Member is experiencing "OFF" episodes such as muscle stiffness, slow movements, or difficulty starting movements				
Provider has submitted documentation which confirms member's symptoms have <u>NOT</u> been adequately controlled with optimal medical therapy using <u>ALL</u> the following agents:				
Dopamine agonist (e.g., Apokyn [®] or apomorphine hydrochloride) (*both prerequisite drugs options require prior authorization – see www.sentarahealthplans.com for prior authorization form)				
ONE agent from any of the following classes:				
☐ Catechol-0-methyl transferase (COMT) inhibitor (e.g., entacapone, Ongentys®, tolcapone)				
☐ Monoamine oxidase B (MAO-B) inhibitor (e.g., rasagiline, selegiline, Xadago®)				
☐ Adenosine receptor antagonist (e.g., Nourianz®)				
Member is receiving concurrent therapy with carbidopa/levodopa within the past 30 days AND will b used in combination with continuous carbidopa/levodopa treatment (verified by chart notes and/or pharmacy paid claims)				
Provider has made adjustments to members carbidopa/levodopa dose in order to manage symptoms without success				
Member must be started on anti-emetic therapy with trimethobenzamide 3 days prior to beginning treatment (<u>NOTE</u> : trimethobenzamide is the only antiemetic that has been studied and can be use with apomorphine)				
Member is <u>NOT</u> currently taking a 5-HT3 antagonist medication such as Zofran [®] (ondansetron), Kytril (granisetron), Aloxi [®] (palonostron), Lotronex [®] (alosetron), or Anzemet [®] (dolasetron) which can result profound hypotension and loss of consciousness (verified by chart notes and/or pharmacy paid claims)				
Member has received a starting dose of requested medication and did <u>NOT</u> develop clinically significated orthostatic hypotension				
Member does NOT have hypersensitivity to apomorphine, its excipients or sodium metabisulfite				

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supp		documentation, incl		oria must be met for approval. To ostics, and/or chart notes, must be		
	Provider must submit chart notes confirming a positive clinical response to therapy, such as continued success in reversing "OFF" episodes, improved motor function, or clinically significant improvement or stabilization in the signs and symptoms of the disease					
	Member continues to me	eet all initial criteria	a and has an absence of o	drug toxicity		
Me	dication being provid	led by (check app	plicable box(es) below):			
	Physician's office	OR	□ Specialty Phar	rmacy		
standa urgent	ard review would subject the	he member to adve t could seriously je	rse health consequences	orization Department if they believe a . Sentara Health Plan's definition of h of the member or the member's		
	•		•	/preauthorization criteria.** laims or submitted chart notes.		