SENTARA HEALTH PLANS

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

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

Drug Requested: Antiparkinson Agents

indrija (levodopa innalation powder)	u Nourianz (istraderylline)
□ Ongentys [®] (opicapone)	□ tolcapone (Tasmar)
MEMBER & PRESCRIBER INFORMATI	ON: Authorization may be delayed if incomplete.
Member Name:	
Member Sentara #:	Date of Birth:
Prescriber Name:	
Prescriber Signature:	Date:
Office Contact Name:	
Phone Number:	Fax Number:
DEA OR NPI #:	
DRUG INFORMATION: Authorization may be	e delayed if incomplete.
Drug Form/Strength:	
Dosing Schedule:	
Diagnosis:	
Weight:	Date:

RECOMMENDED DOSAGE:

- **Inbrija**[™]: Oral inhalation: 84 mg up to 5 times daily as needed when symptoms of an OFF period return. Maximum quantity limit: 84 mg/dose and 420 mg/day.
- **Nourianz**[™]: Oral: 20 mg once daily; may further increase dose based on response and tolerability to a maximum dose of 40 mg once daily. Maximum quantity limit: 30 tablets /30 days
- Ongentys[®]: Oral: 50 mg once daily at bedtime. Maximum quantity limit: 30 tablets/30 days.
- **tolcapone (Tasmar):** Oral: Initial: 100 mg 3 times daily always as an adjunct to levodopa/carbidopa; may increase as tolerated to 200 mg 3 times daily. Maximum quantity limit: 180 tablets/30days

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 Approval: 6 months

	Me	ember must be 18 years of age or older	
		AND	
	Me	edication must be prescribed by, or in consultation with a neurologist	
		<u>AND</u>	
	inte	ember must have a confirmed diagnosis of Parkinson's disease in an individual who is having ermittent OFF episodes while on continuous carbidopa/levodopa therapy and all of the following teria has been met: (must submit chart notes)	
		Provider has made adjustments to adjust the carbidopa/levodopa's dose in order to manage symptoms without success	
		AND	
		Member is receiving concurrent therapy with carbidopa/levodopa within the past 30 days AND the requested medication will be used in combination with continuous carbidopa/levodopa treatment	
		AND	
		For ALL Antiparkinson Agents , member is currently not taking or has not recently (within 2 weeks) taken a nonselective MAO inhibitor such as Nardil [®] (phenelzine), Parnate [®] (tranylcypromine), or Marplan [®] (isocarboxazid)	
F	or t	tolcapone (Tasmar). Check below all that apply. All criteria must be met for approval. To	
		rt each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be ded or request may be denied.	
	Me	ember must have documentation of trial and failure of TWO (2) of the following:	
		COMT inhibitor: generic entacapone	
		AND	
		Dopamine agonist: ropinirole/ropinirole ER, pramipexole/pramipexole ER; OR	
		Monoamine oxidase type B inhibitors: rasagiline; OR	
		Ongentys® (opicapone) requires prior authorization; OR	
AND			
		ovider attestation to monitor for liver failure/hepatic dysfunction and should discontinue tolcapone if LT/AST levels exceed 2 times the upper limit of normal	
ch	eck	Ongentys [®] . Check below all that apply. All criteria must be met for approval. To support each line ed, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or st may be denied.	
	Me	ember must have documentation of trial and failure of TWO (2) of the following:	
		COMT inhibitor: generic entacapone	
		AND	
		Dopamine agonist: ropinirole/ropinirole ER, pramipexole/pramipexole ER; OR	
		Monoamine oxidase type B inhibitors: rasagiline; OR	
		(Continued on next page)	

AND

provided or request may be denied.

Member does not have a history of pheochromocytoma, paraganglioma, or other catecholamine-secreting neoplasms
For Inbrija® or Nourianz®, 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

☐ Member must have documentation of trial and failure of **TWO (2)** of the following:

- ☐ Monoamine oxidase type B inhibitors: rasagiline; **OR**
- Dopamine agonist: ropinirole/ropinirole ER, pramipexole/pramipexole ER; **OR**
- □ COMT inhibitor: generic entacapone, Ongentys® (requires prior authorization), tolcapone (requires prior authorization)

<u>AND</u>

- □ Member does not have a history of asthma, COPD, or other chronic underlying lung disease (for Inbrija[™] only)
- □ Reauthorization Approval: 12 months. 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.
 - ☐ Member must continue to meet initial approval criteria

<u>AND</u>

☐ Member has a documented positive clinical response to treatment (defined as: improvement and stabilization of "off episodes" associated with Parkinson's disease)

AND

☐ Medication is used in combination with carbidopa/levodopa (must have pharmacy paid claims)

AND

☐ Member must be absent of unacceptable toxicity from therapy

Medication being provided by Specialty Pharmacy - PropriumRx

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. **

Previous therapies will be verified through pharmacy paid claims or submitted chart notes.