## SENTARA HEALTH PLANS

## 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 1-844-668-1550</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>.

**For Medicare Members:** Medicare Coverage for outpatient (Part B) drugs is outlined in the Medicare Benefit Policy Manual (Pub. 100-2), Chapter 15, §50 Drugs and Biologicals. In addition, National Coverage Determination (NCD) and Local Coverage Determinations (LCDs) may exist and compliance with these policies is required where applicable. They can be found at: <a href="https://www.cms.gov/medicare-coverage-database/overview-and-quick-search.aspx">https://www.cms.gov/medicare-coverage-database/overview-and-quick-search.aspx</a>. Additional indications may be covered at the discretion of the health plan.

## **Botulinum Toxin Injections®**, Type A

<u>Drug Requested</u>: Botox<sup>®</sup> (onabotulinumtoxinA) (J0585) (Medical) (Chronic Migraine Headache Prophylaxis)

MEMBER & PRESCRIBER INFO	RMATION: Authorization may be delayed if incomplete.
Member Name:	
Member Sentara #:	Date of Birth:
Prescriber Name:	
Prescriber Signature:	
Phone Number:	
NPI #:	
DRUG INFORMATION: Authorization	on 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:
•	ne timeframe does not jeopardize the life or health of the member m function and would not subject the member to severe pain.

- Max quantity limits: 155 units once every 12 weeks
- Cosmetic indications are **EXCLUDED**

**NOTE:** In treating adult patients for one or more indications, the maximum cumulative dose should not exceed 400 units, in a 3-month interval. In pediatric patients, the total dose should not exceed the lower of 10 units/kg body weight or 340 units, in a 3-month interval

**CLINICAL CRITERIA:** Check below all that apply. <u>All criteria must be met for approval</u>. 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		
	Member experiences ≥ 15 headache days per month		
	Member experiences headaches which last $\geq 4$ hours per day		
	Member must have failed a <u>2-month</u> trial of a least one medication from <u>TWO</u> different migraine prophylactic classes supported by the American Headache Society/American Academy of Neurolo treatment guidelines 2012/2015/2021/2024, Level A and B evidence; ICSI 2013, high quality evide (verified by chart notes or pharmacy paid claims):		
	☐ Anticonvulsants (divalproex, valproate, topiramate)		
	☐ Beta blockers (atenolol, metoprolol, nadolol, propranolol, timolol)		
	☐ Antidepressants (amitriptyline, venlafaxine)		
	☐ Angiotensin II receptor blocker (candesartan) *requires prior authorization*		
	□ Injectable CGRP inhibitors (Aimovig <sup>®</sup> , Emgality <sup>®</sup> , Ajovy <sup>®</sup> ) or oral CGRP inhibitors indicated for migraine prevention (Qulipta <sup>™</sup> , Nurtec ODT <sup>®</sup> ) *requires prior authorization*		
	Member has been evaluated for medication overuse headache (MOH) (defined as headaches occurring greater than or equal to 15 days per month. It develops as a consequence of regular overuse of acute or symptomatic headache medication for more than 3 months)		
	Treatment will include a plan to taper off the offending medication if MOH is diagnosed		
	Requests for concurrent use of Calcitonin Gene-Related Peptide (CGRP) inhibitors with Botox® (onabotulinumtoxinA) for migraine headache prevention (if applicable): Member must meet <u>ALL</u> t following criteria (verified by chart notes and/or pharmacy paid claims):		
	□ Member must have a diagnosis of Chronic or Episodic Migraine Headache and is continuing to experience ≥ 4 migraine headache days per month after receiving therapy with <u>ALL</u> the following criteria:		
	☐ Member must have failed a <b>2-month</b> trial of at least one medication from <b>TWO</b> different migraine prophylactic classes supported by the American Headache Society/American Academy of Neurology treatment guidelines 2012/2015/2021/2024, Level A and B evidence: ICSI 2013, high quality evidence:		
	☐ Anticonvulsants (divalproex, valproate, topiramate)		
	☐ Beta blockers (atenolol, metoprolol, nadolol, propranolol, timolol)		
	☐ Antidepressants (amitriptyline, venlafaxine)		
	☐ Angiotensin II receptor blocker (candesartan) *requires prior authorization*		

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	u Memi	ber must meet ONE of the following:	
	in	lember has had an inadequate response to a <u>2-month</u> trial with an injectable CGRP hibitor (e.g., Aimovig <sup>®</sup> , Ajovy <sup>®</sup> , Emgality <sup>®</sup> ) or an oral CGRP inhibitor indicated for igraine prevention (e.g., Nurtec <sup>®</sup> ODT, Qulipta <sup>™</sup> ) *requires prior authorization*	
		Tember has had an inadequate response to a <u>6-month</u> trial (2 injection cycles) of Botox® onabotulinumtoxinA) *requires prior authorization*	
ıppo	ort each line ch	1: 12 months. Check below all that apply. All criteria must be met for approval. To necked, all documentation, including lab results, diagnostics, and/or chart notes, must be may be denied.	
	Member has experienced a positive response to therapy, demonstrated by a reduction in headache frequency		
	Use of acute	migraine medications (e.g., NSAIDs, triptans) has decreased since the start of Botox®	
	Member cont	tinues to be monitored for medication overuse headache (MOH)	
	Requests for continuation of concurrent use of Calcitonin Gene-Related Peptide (CGRP) inhibitor with Botox® (onabotulinumtoxinA) for migraine headache prevention (if applicable): Member must meet the following criteria:		
	number o onabotuli	has experienced further reduction in the overall number of migraine days or reduction in f severe migraine days per month compared to monotherapy with the initial agent numtoxinA (Botox) and a calcitonin gene-related peptide (CGRP) indicated for migraine n (submit documentation)	
<b>Sedication being provided by:</b> Please check applicable box below.			
	Location/site	e of drug administration:	
	NPI or DEA	# of administering location:	
		<u>OR</u>	
	Specialty Ph	armacy – Proprium Rx	

For urgent reviews: Practitioner should call Sentara Pre-Authorization Department if they believe a standard review would subject the member to adverse health consequences. Sentara's definition of urgent is a lack of treatment that could seriously jeopardize the life or health of the member or the member's ability to regain maximum function.

\*\*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. \*