

PHARMACY POLICY STATEMENT		
Arkansas PASSE		
DRUG NAME	Qbrexza (glycopyrronium) cloth, 2.4%	
BILLING CODE	Must use valid NDC code	
BENEFIT TYPE	Pharmacy	
SITE OF SERVICE ALLOWED	Home	
COVERAGE REQUIREMENTS	Prior Authorization Required (Non-Preferred Product) QUANTITY LIMIT— carton of 30 pouches for 30 days	
LIST OF DIAGNOSES CONSIDERED <b>NOT</b> MEDICALLY NECESSARY	Click Here	

Qbrexza (glycopyrronium) cloth, 2.4% will only be considered for coverage under the **pharmacy** benefit when the following criteria are met:

Members must be clinically diagnosed with one of the following disease states and meet their individual criteria as stated.

## PRIMARY AXILLARY HYPERHIDROSIS

For **initial** authorization:

- 1. Member must be 9 years of age or older; AND
- 2. Member has a diagnosis of severe axillary hyperhidrosis, including documentation in the chart notes of visible, excessive sweating of at least 6 months duration which significantly impairs daily activities; AND
- 3. Member has tried and failed topical prescription-strength aluminum chloride (e.g. Xerac) for at least 60 days.
- 4. **Dosage allowed:** Qbrexza cloth (one cloth per pouch) is used topically once daily to both axillae using a single cloth.

If member meets all the requirements listed above, the medication will be approved for 2 months. For reauthorization:

1. Chart notes must document clinically significant decreased severity of sweating.

If member meets all the reauthorization requirements above, the medication will be approved for an additional 12 months.

CareSource considers Qbrexza (glycopyrronium) cloth, 2.4% not medically necessary for the treatment of the following disease states based on a lack of robust clinical controlled trials showing superior efficacy compared to currently available treatments:

• Hyperhidrosis of palms/hands, soles (feet), forehead and other regions

DATE	ACTION/DESCRIPTION
11/27/2018	New policy for Qbrexza created.
09/22/2020	Reordered the criteria, matched wording to Botox policy, removed sweat quantification measure, removed HDSS score, summarized the list of exclusions, removed the

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		mental health issue piece, removed trial of Botox per IHS guideline, changed the reauth criteria, extended reauth duration per long term efficacy study, changed Drysol to Xerac and extended trial period to be 60 days, changed initial auth duration to 2 months instead of 1.
	12/21/2021	Removed prescriber specialty requirement, ruling out secondary causes of
		hyperhidrosis, and requirement that member does not have a medical condition that

## References:

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may be exacerbated by anticholinergic effects.

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- 5. Gelbard, Christina M. MD, et al. "Primary pediatric hyperhidrosis: a review of current treatment options". Pediatric Dermatology 25:6 (2008): 591-598.
- 6. Guidance for Industry: Patient-Reported Outcome Measures: Use in Medical Product Development to Support Labeling Claims. FDA, 2009.
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Effective date: 01/01/2022 Revised date: 12/21/2021