AETNA BETTER HEALTH® Coverage Policy/Guideline			<b>*ae</b>	etna <sup>™</sup>
Name:	Ponvory		Page:	1 of 2
Effective Date: 3/4/2024			Last Review Date:	01/12/2024
Applies to:	⊠Illinois ⊠Maryland □Michigan	□Florida ⊠Florida Kids □ Virginia	□New Jersey ⊠Pennsylvania Kids	

#### Intent:

The intent of this policy/guideline is to provide information to the prescribing practitioner outlining the coverage criteria for Ponvory under the patient's prescription drug benefit.

## **Description:**

### **FDA-Approved Indications**

Ponvory is indicated for the treatment of relapsing forms of multiple sclerosis, to include clinically isolated syndrome, relapsing-remitting disease, and active secondary progressive disease, in adults.

All other indications are considered experimental/investigational and not medically necessary.

# **Applicable Drug List:**

Ponvory

### Policy/Guideline:

#### . CRITERIA FOR INITIAL APPROVAL

### A. Relapsing forms of multiple sclerosis

- 1. The patient is unable to take the required number of preferred formulary alternatives (3) for the given diagnosis due to a trial and inadequate treatment response, intolerance, or a contraindication.
- 2. Authorization may be granted to members who have been diagnosed with a relapsing form of multiple sclerosis (including relapsing-remitting and secondary progressive disease for those who continue to experience relapse).
  - a. Pediatric members less than 18 years of age may be granted authorization when benefits outweigh risks
- 3. Ponvory must be prescribed by or in consultation with a neurologist.
- 4. Members will not use Ponvory concomitantly with other disease modifying multiple sclerosis agents

Note: Ampyra and Nuedexta are not disease modifying.

## **B.** Clinically isolated syndrome

- 1. The patient is unable to take the required number of preferred formulary alternatives (3) for the given diagnosis due to a trial and inadequate treatment response or intolerance, or a contraindication.
- 2. Authorization may be granted to members for the treatment of clinically isolated syndrome.

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- a. Pediatric members less than 18 years of age may be granted authorization when benefits outweigh risks
- 3. Ponvory must be prescribed by or in consultation with a neurologist.
- 4. Members will not use Ponvory concomitantly with other disease modifying multiple sclerosis agents
  - a. Ampyra and Nuedexta are not disease modifying.

### II. CRITERIA FOR CONTINUATION OF THERAPY

### A. For all indications:

- 1. Authorization may be granted to members who are experiencing disease stability or improvement while receiving Ponvory.
- 2. Ponvory must be prescribed by or in consultation with a neurologist.
- 3. Members will not use Ponvory concomitantly with other disease modifying multiple sclerosis agents

Note: Ampyra and Nuedexta are not disease modifying.

## **Approval Duration and Quantity Restrictions:**

## **Initial and Renewal Approval:**

12 months

#### **Quantity Level Limit:**

Starter Pack: 1 Starter Pack (14 tablets) per 14 days

Maintenance dose: 20 mg tablet, 30 tablets per 30 days

#### **References:**

1. Ponvory [package insert]. Titusville, NJ: Janssen Pharmaceuticals, Inc; August 2023.