# SENTARA COMMUNITY PLAN (MEDICAID)

#### 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, correct, or legible, the authorization process can be delayed.</u>

## Multiple Sclerosis Drugs – Mavenclad<sup>®</sup>, Mayzent<sup>®</sup>, Ponvory<sup>™</sup> or Zeposia<sup>®</sup> (Non-Preferred)

**Drug Requested:** (check box below that applies): ☐ Mavenclad® (cladribine) □ **Mayzent**® (siponimod) □ **Ponvory**<sup>™</sup> (ponesimod) □ **Zeposia**® (ozanimod) MEMBER & PRESCRIBER INFORMATION: 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 delayed if incomplete. Drug Form/Strength: Dosing Schedule: Length of Therapy: Diagnosis: ICD Code, if applicable: Weight: \_\_\_\_\_ Date: \_\_\_\_

#### **Recommended Dosage for Mavenclad:**

• Oral: 3.5 mg/kg over 2-year treatment course, administered as 1.75 mg/kg in each year. Divide the 1.75 mg/kg dose over 2 cycles, each cycle lasting 4 to 5 consecutive days; do not administer more than 2 tablets (20mg) /day. Following the administration of 2 treatment courses, do not administer additional Mavenclad treatment during the next 2 years.

Weight in kilograms: \_\_\_\_\_ Date:

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#### **Administration of First Treatment Course**

- First Course/First Cycle: start any time
- First Course/Second Cycle: administer 23 to 27 days after the last dose of First Course/First Cycle

#### **Administration of Second Treatment Course**

- Second Course/First Cycle: administer at least 43 weeks after the last dose of First Course/Second Cycle
- Second Course/Second Cycle: administer 23 to 27 days after the last dose of Second Course/First Cycle

Dose of MAVENCLAD per Cycle by Patient Weight in Each Treatment Course

Weight Range	Dose in mg (Number of 10 mg Tablets) per Cycle				
kg	First Cycle	Second Cycle			
40* to less than 50	40 mg (4 tablets)	40 mg (4 tablets)			
50 to less than 60	50 mg (5 tablets)	50 mg (5 tablets)			
60 to less than 70	60 mg (6 tablets)	60 mg (6 tablets)			
70 to less than 80	70 mg (7 tablets)	70 mg (7 tablets)			
80 to less than 90	80 mg (8 tablets)	70 mg (7 tablets)			
90 to less than 100	90 mg (9 tablets)	80 mg (8 tablets)			
100 to less than 110	100 mg (10 tablets)	90 mg (9 tablets)			
110 and above	100 mg (10 tablets)	100 mg (10 tablets)			

<sup>\*</sup>The use of MAVENCLAD in patients weighing less than 40 kg has not been investigated

**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.

		J			
1.	Is memb  ☐ Yes	at least 18 years of age?  No			
2. Has the member had a baseline magnetic resonance imaging (MRI) before initiating the first course (within 3 months prior to start of therapy)?					
	□ Yes	□ No			
		(Continued on next page)			

### PA Multiple Sclerosis Drugs (Non-Preferred) (Medicaid) (Continued from previous page)

3.	Ind	licate all that apply:						
		Relapsing-remitting Disease (RRMS)						
		Secondary Progressive Disease (SPMS) with relapses						
		Clinically Isolated Syndrome (CIS)						
		Member has had $\geq 1$ relapse within the previous two years						
		Member has new and unequivocally enlarging T2 contrast enhancing lesions as evidenced by MRI and has had $\geq 1$ relapse in the previous 12 months						
		Other:						
4.	Has the member had a treatment failure or contraindication to other agents used to treat multiple sclerosis (MS)? List previous medications (include drug name/dose):							
		Yes						
	Lis	st previous medications (include drug name/dose):						
5.	Wi	ill Mavenclad <sup>®</sup> , Mayzent <sup>®</sup> , Ponvory <sup>™</sup> <b>OR</b> Zeposia <sup>®</sup> be used as a single-agent therapy?						
		Yes   No						
5.	Has the member been tested for antibodies to the varicella zoster virus (VZV) or received immunizatio for VZV four weeks prior to beginning therapy?							
		Yes						
7.	На	s the member been screened for the presence of tuberculosis according to local guidelines?						
		Yes						
3.		s the member been evaluated and screened for the presence of hepatitis B and hepatitis C virus BV/HCV prior to initiating treatment?)						
		Yes □ No						
9.	Ma	avenclad® Specific						
		Is the lymphocyte count $\geq 800$ cells/mL prior to start of therapy?						
		□ Yes □ No						
	b.	Please attest that women of childbearing age are not pregnant and that members of reproductive potential must use effective contraception during treatment with therapy and for at least six months after the last dose.						
		□ Yes □ No						
	c.	Does the member have human immunodeficiency virus (HIV) infection?						
		□ Yes □ No						
10	Ma	ayzent® Specific						
	a.	Has the member been tested for CYP2C9 variant status to determine genotyping (required for dosing)?						
		□ Yes □ No						

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11. <b>M</b>	ayzent®, Ponvory™ OR Zeposia® Specific							
a.	Please attest that women of childbearing age are not pregnant and that members of reproductive potential must use effective contraception during treatment.							
	□ Yes □ No							
b.	Has the member obtained a baseline electrocardiogram (ECG)?							
	□ Yes □ No							
c	Has the member had a baseline ophthalmic evaluation of the fundus, including the macula, before starting treatment?							
	□ Yes □ No							
	efore using Mayzent <sup>®</sup> , Ponvory <sup>™</sup> <b>OR</b> Zeposia <sup>®</sup> , please attest that the member does <b>NOT</b> have any of e following:							
•	Recent myocardial infarction							
•	Unstable angina							
•	Stroke							
•	Transient Ischemic Attack							
•	Decompensated heart failure with hospitalization							
•	Class III/IV heart failure within the previous 6 months							
•	D 1 10T 1 1 1 1 (500 )							
•	CYP2C9*3/*3 genotype (Mayzent® ONLY)							
<ul> <li>History of Mobitz Type II second or third-degree atrioventricular block or sick sinus (unless treated with a functioning pacemaker)</li> </ul>								
	Yes   No							
13. Ca	in you confirm that Mayzent® will NOT be used in combination with the following?							
•	Moderate or strong CYP3A4 inducers (e.g., modafinil, efavirenz) in members with a CYP2C9*1/*3 and CYP2C9*2/*3 genotypes; <b>OR</b>							
•	• Drug regimens that contain CYP2C9/CY3A4 dual inhibitors (e.g., fluconazole); <b>OR</b>							
•	M. L. CVPACO: 171; J. L. L. L. CVPALA: 171; OB							
•	Other antineoplastic, immunosuppressive or immunomodulating drugs.							
	Yes  No							

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### PA Multiple Sclerosis Drugs (Non-Preferred) (Medicaid) (Continued from previous page)

- 14. Can you confirm **Zeposia**<sup>®</sup> will **NOT** be used in combination with the following?
  - Will **NOT** be initiating therapy after previous treatment with alemtuzaumab; **OR**
  - Monoamine oxidase inhibitor (MAOI) (e.g., selegiline, phenelzine, linezolid); **OR**
  - Drugs known to prolong the QT-interval (e.g., fluoroqinolone or macrolide antibiotics, venlafaxine, fluoxetine, quetiapine, ziprasidone, sumatriptan, zolmitriptan); **OR**
  - Strong cytochrome p450 2C8 (CYP2C8) inhibitors (e.g., gemfibrozil) or inducers (e.g., rifampin);
     OR
  - BCRP inhibitors (e.g., cyclosporine, eltombopag); **OR**
  - Adrenergic or serotonergic drugs which can increase norepinephrine or serotonin (e.g., opioids, selective serotonin reuptake inhibitors (SSRIs), selective norepinephrine reuptake inhibitors (SNRIs), tricyclics, tyramine; **OR**
  - Foods with large amounts of tyramine (e.g., >150mg), such as aged cheeses, cured meats, craft/unfiltered beers, beans); **OR**
  - Other antineoplastic, immunosuppressive or immunomodulating drugs (**Note**: if there is a history of prior use of these drugs, consider possible unintended additive immunosuppressive effects) **AND**
  - Patient will **NOT** receive live vaccines during and at least 4 weeks prior to and 12 weeks after treatment; **AND**

•	Patient do	es <b>NOT</b>	have an	active	infection,	including	clinically	important	localized	infections
	Yes	□ No								

Medication	being provided b	by Special	ty Pharmacy	- PropriumRx
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\*\*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. \*