Member Name: {{MEMFIRST}} {{MEMLAST}} DOB: {{MEMBERDOB}} PA Number: {{PANUMBER}}



Multiple Sclerosis

Prior Authorization Request

CVS Caremark administers the prescription benefit plan for the patient identified. This patient's benefit plan requires prior authorization for certain medications in order for the drug to be covered. To make an appropriate determination, providing the most accurate diagnosis for the use of the prescribed medication is necessary. Please respond below and fax this form to CVS Caremark toll-free at 1-866-249-6155. If you have questions regarding the prior authorization, please contact CVS Caremark at 1-866-814-5506. For inquiries or questions related to the patient's eligibility, drug copay or medication delivery; please contact the Specialty Customer Care Team: CaremarkConnect® 1-800-237-2767.

The recipient of this fax may make a request to opt-out of receiving telemarketing fax transmissions from CVS Caremark. There are numerous ways you may opt-out: The recipient may call the toll-free number at 877-265-2711, at any time, 24 hours a day/7 days a week. The recipient may also send an opt-out request via email to do not call@cvscaremark.com. An opt out request is only valid if it (1) identifies the number to which the request relates, and (2) if the person/entity making the request does not, subsequent to the request, provide express invitation or permission to CVS Caremark to send facsimile advertisements to such person/entity at that particular number. CVS Caremark is required by law to honor an opt-out request within thirty days of receipt.

	<u> FIENT INFORMATION</u>	PRESCRIBER INFORMATION
	te: {{TODAY}}	Name: {{PHYFIRST}} {{PHYLAST}}
	me: {{MEMFIRST}} {{MEMLAST}}	Office Telephone: {{PHYSICIANPHONE}}
	{{MEMBERID}}	Office Fax: {{PHYSICIANFAX}}
	te of Birth: {{MEMBERDOB}} quest Initiated For: {{DRUGNAME}}	Specialty:NPI#:
IXC	quest initiated Poil ((DROOMANIE))	111 III,
	glatiramer 20mg 🗖 glatiramer 40mg 🗖 Glatopa 🗖 Ke Plegridy 🗖 Rebif 🗖 dimethyl fumarate 🗖 teriflunomi	de □ Zeposia □ Vumerity
	n-preferred: □ Ampyra □ Aubagio □ Bafiertam □ Ponvory □ Briumvi □ Other	
	What is the patient's diagnosis? □ Relapsing form of multiple sclerosis (including relaps who continue to experience relapse) □ Primary progressive multiple sclerosis □ Other	sing-remitting and secondary progressive disease for those Clinically isolated syndrome of multiple sclerosis
2.	ICD-10:	te the PA electronically (ePA). You may sign up online
<u>PR</u> 1.		Avonex, Betaseron, Copaxone, dimethyl fumarate, crevus, Rebif, teriflunomide, Tysabri, Vumerity, and creferred product? If Yes, and drug is Ocrevus, please your office OR you may complete the PA electronically www.covermymeds.com/epa/caremark/ or call 1-866-452- at to the pharmacy and skip to next section.

Send completed form to: Case Review Unit, CVS Caremark Prior Authorization. Fax: 1-866-249-6155

Note: This fax may contain medical information that is privileged and confidential and is solely for the use of individuals named above. If you are not the intended recipient you hereby are advised that any dissemination, distribution, or copying of this communication is prohibited. If you have received the fax in error, please immediately notify the sender by telephone and destroy the original fax message. Multiple Sclerosis [non-Tysabri] PDPD CareFirst - 9/2023.

Me	mber Name: {{MEMFIRST}}	{{MEMLAST}} DOB: {{ME	EMBERDOB}} PA Number: {{PANUMBER}}			
2.			intolerable adverse event to any of the following importing chart note(s). Indicate ALL that apply. Intolerable adverse event			
<u>Aul</u> 3.	 If the patient had a documented intolerable adverse event to generic teriflunomide, was the documented intolerable adverse event an expected adverse event attributed to the active ingredient as described in the prescribing information? ACTION REQUIRED: If No, attach supporting chart note(s). □ Yes □ No □ N/A, the patient did NOT have a documented intolerable adverse event to generic teriflunomide 					
<u>Baj</u> 4.	afiertam Has the patient experienced a documented intolerable adverse event to dimethyl fumarate (including intolerable gastrointestinal adverse events from dimethyl fumarate) or Vumerity? <i>ACTION REQUIRED: If Yes, attach supporting chart note(s).</i> □ Yes □ No					
whi	ch are made in the same manuf Given that Betaseron and Exta	<i>facturing facility.</i> Evia are the same products, is the	there a documented clinical reason that the patient If Yes, attach supporting chart note(s).			
	 Filenya If the patient had a documented intolerable adverse event to generic fingolimod, was the documented intolerable adverse event an expected adverse event attributed to the active ingredient as described in the prescribing information? ACTION REQUIRED: If No, attach supporting chart note(s). □ Yes □ No □ N/A, the patient did NOT have a documented intolerable adverse event to generic fingolimod 					
	Is this request for continuation of therapy with the requested product? ☐ Yes ☐ No If No, skip to All Requests section.					
8.	Is the patient currently receiving the requested product through samples or a manufacturer's patient assistance program? \square Yes \square No					
<u>Tec.</u> 9.	intolerable adverse event an exprescribing information? AC	xpected adverse event attribute TION REQUIRED: If No, atta	generic dimethyl fumarate, was the documented of to the active ingredient as described in the ach supporting chart note(s). Intendiction of the support of			

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CVS Caremark Prior Authorization

1300 E. Campbell Road

Richardson, TX 75081

Me	ember Name: {{MEMFIRST}} {{MEMLAST}} DOB: {{MEMBERDOB}} PA Number: {{PANUMBER}}
	L REQUESTS (EXCEPT AMPYRA) If the patient is less than 18 years of age, has the prescriber evaluated the risks and benefits of treatment and attests the benefits outweigh the risks? Yes No N/A - Patient is not less than 18 years old
2.	If the prescribed drug is Zeposia, will the requested drug be used in combination with any other immunomodulator, biologic drug (e.g., Humira), targeted synthetic drug (e.g., Rinvoq, Xeljanz), or disease modifying multiple sclerosis (MS) agent? (Note: Ampyra and Nuedexta are not disease modifying.) ☐ Yes ☐ No ☐ N/A - Requested drug is not Zeposia
3.	Is the patient taking the requested medication with any other disease modifying multiple sclerosis (MS) agent? (Note: Ampyra and Nuedexta are not disease modifying). \square Yes \square No
4.	Will the requested drug be used in combination with any other disease modifying multiple sclerosis (MS) agent? (Note: Ampyra and Nuedexta are not disease modifying). \square Yes \square No
5.	Is the requested medication prescribed by or in consultation with a neurologist? ☐ Yes ☐ No If requested drug is Mavenclad, skip to drug specific questions.
6.	If requested drug is Kesimpta, is the patient currently receiving the requested medication? \square Yes \square No
7.	Is this a request for continuation of therapy? ☐ Yes ☐ No If No, skip to drug specific questions, if applicable.
8.	Is the patient experiencing disease stability or improvement while receiving the requested medication? ☐ Yes ☐ No
	RUG SPECIFIC QUESTIONS
	AVENCLAD Is this a request for continuation of therapy? If Yes, skip to #4 Yes No
2.	Has the patient received any cycles of Mavenclad previously? Note: Mavenclad is administered based on weight over 4 to 5 days. This 4 to 5 day administration period is a cycle. Two cycles administered three to four weeks apart are a course cycle(s) \text{None}
3.	Has the patient had an inadequate response or was unable to tolerate an alternative drug indicated for the treatment of multiple sclerosis (e.g., Rebif, Tecfidera, Copaxone, etc)? Yes No No further questions.
4.	How many cycles of Mavenclad has the patient received previously? Note: Mavenclad is administered based on weight over 4 to 5 days. This 4 to 5 day administration period is a cycle. Two cycles administered three to four weeks apart are a course cycle(s)
5.	Has the patient received a complete course (two 4-5 day cycles) of Mavenclad in the last 43 weeks? <i>Note: One course is two 4 to 5 day cycles administered 3 to 4 weeks apart).</i> □ Yes □ No
	IPYRA (dalfampridine ER) If brand Ampyra is being prescribed, is the prescriber willing to switch to the generic dalfampridine ER? If Yes, fax a new prescription to the pharmacy and skip to #5. □ Yes - generic dalfampridine ER □ No □ Generic dalfampridine ER is being requested, skip to #5
2.	Has the patient failed treatment with the generic medication due to an intolerable adverse event (e.g., rash, nausea, vomiting)? ☐ Yes ☐ No
3.	Was the intolerable adverse event an expected adverse event attributed to the <u>active</u> ingredient as described in the prescribing information (i.e., known adverse reaction for both the brand and generic medication)? Yes No
4.	Was this adverse event documented in the patient's chart? ACTION REQUIRED: Documentation is required for approval. Provide SPECIFIC and DETAILED chart documentation including description, date/time, and severity of the adverse event, dosage and duration of generic medication treatment, required intervention (if any), and relevant tests or laboratory data (if any) OR MedWatch form of this trial and failure including the adverse reaction. \square Yes \square No

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5.	Is this request for continuation of therapy with the requested medication? Yes No If No, skip to #7
6.	Has the patient experienced improvement in walking speed or other objective measure of walking ability since starting therapy with the requested medication? \square Yes \square No <i>No further questions</i> .
7.	Prior to initiation of therapy with the requested medication, does/did the patient have sustained walking impairment? Yes No
	LENYA (fingolimod hydrochloride), TASCENSO ODT Is the patient experiencing disease stability or improvement while receiving the requested medication? ☐ Yes ☐ No
LE	CMTRADA
1.	Is the patient currently receiving the requested product through samples or a manufacturer's patient assistance program? If unknown, answer Yes. \square Yes \square No
2.	How many courses of the requested medication has the patient previously received? courses If one course or more (5 doses or more), skip to #4
3.	Has the patient had an inadequate response to two or more drugs indicated for multiple sclerosis? ☐ Yes ☐ No No further questions.
4.	Has the patient received the last dose of the previous course of the requested medication at least 12 months prior to the planned date of the first dose of the subsequent treatment course of the requested medication? ☐ Yes ☐ No
ΑŪ	JTHORIZATION
\overline{Ia}	ttest that this information is accurate and true, and that documentation supporting this
inf	formation is available for review if requested by CVS Caremark or the benefit plan sponsor.
X_	Date (see that)
Pre	escriber or Authorized Signature Date (mm/dd/yy)

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