

Please complete all required sections to allow your request to be processed.

PATIENT INFORMATION				COVERAGE TYPE	
PATIENT LAST NAME	FIRST NAME	INITIAL	<input type="checkbox"/> Alberta Blue Cross <input type="checkbox"/> Alberta Human Services <input type="checkbox"/> Other		
BIRTH DATE (YYYY-MM-DD)	ALBERTA PERSONAL HEALTH NUMBER				
STREET ADDRESS	CITY	PROV	POSTAL CODE	ID/CLIENT/COVERAGE NUMBER	
PRESCRIBER INFORMATION					
PRESCRIBER LAST NAME	FIRST NAME	INITIAL	PRESCRIBER PROFESSIONAL ASSOCIATION REGISTRATION		
STREET ADDRESS			REGISTRATION NUMBER		
			<input type="checkbox"/> CPSA <input type="checkbox"/> ACO <input type="checkbox"/> CARNA <input type="checkbox"/> ADA+C <input type="checkbox"/> ACP <input type="checkbox"/> Other		
CITY, PROVINCE			PHONE	FAX	
POSTAL CODE			FAX NUMBER MUST BE PROVIDED WITH EACH REQUEST SUBMITTED		
Please provide the following information for ALL requests					
<input type="checkbox"/> NEW request (i.e. new to MS DMT and/or coverage) <input type="checkbox"/> MS disease modifying therapy (DMT) SWITCH					
For patients new to coverage and already on alemtuzumab , specify start date (YYYY-MM-DD) _____ and number of treatment courses and doses/course administered _____					
Diagnosis <input type="checkbox"/> Relapsing-remitting multiple sclerosis <input type="checkbox"/> Other (specify) _____			Current EDSS _____ . _____ Date (YYYY-MM-DD) _____		
NEW requests: Qualifying relapses					
Provide the dates of two relapses within the last two years OR the two years prior to starting MS DMT					
Date of relapse (YYYY-MM-DD)		Type of relapse (One MRI relapse may substitute for one clinical relapse)			
		<input type="checkbox"/> Clinical relapse		<input type="checkbox"/> MRI relapse (new T2 lesion or definite gadolinium-enhancing T1 lesion)	
		<input type="checkbox"/> Clinical relapse		<input type="checkbox"/> MRI relapse (new T2 lesion or definite gadolinium-enhancing T1 lesion)	
a) Has the patient been on MS DMT of any kind since the relapse(s)? <input type="checkbox"/> No <input type="checkbox"/> Yes → If yes, answer b) and c)					
b) Specify the MS DMT start date (YYYY-MM-DD) _____					
c) Indicate if there have been any interruptions in therapy since starting MS DMT <input type="checkbox"/> No <input type="checkbox"/> Yes → If yes, indicate					
i) Reason for the interruption in therapy _____					
ii) Specify time period of interruption: from (YYYY-MM-DD) _____ to (YYYY-MM-DD) _____					
iii) How many relapses did the patient experience while off therapy? _____					
ALL requests: Provide response to TWO of the following MS DMT					
DIMETHYL FUMARATE; GLATIRAMER ACETATE; INTERFERON BETA; OCRELIZUMAB; PEGINTERFERON BETA; TERIFLUNOMIDE					
Name of 1st MS DMT utilized _____ and date of treatment initiation (YYYY-MM-DD) _____					
<input type="checkbox"/> INTOLERANCE despite the use of symptom management techniques; OR <input type="checkbox"/> REFRACTORY → answer a) and b)					
a) Does the patient have clinically significant titres of neutralizing antibodies to interferon beta? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> N/A					
b) Within a consecutive 12-month period while on the MS DMT, did the patient experience at least two relapses of MS? <input type="checkbox"/> No <input type="checkbox"/> Yes → Provide the dates of either two clinical relapses OR one clinical relapse and one MRI relapse					
Date of relapse (YYYY-MM-DD)		Type of relapse (One MRI relapse may substitute for one clinical relapse)			
		<input type="checkbox"/> Moderate to very severe clinical relapse		<input type="checkbox"/> MRI relapse (new T2 lesion or definite gadolinium-enhancing T1 lesion)	
		<input type="checkbox"/> Moderate to very severe clinical relapse		<input type="checkbox"/> MRI relapse (new T2 lesion or definite gadolinium-enhancing T1 lesion)	
Name of 2nd MS DMT utilized _____ and date of treatment initiation (YYYY-MM-DD) _____					
<input type="checkbox"/> INTOLERANCE despite the use of symptom management techniques; OR <input type="checkbox"/> REFRACTORY → answer a) and b)					
a) Does the patient have clinically significant titres of neutralizing antibodies to interferon beta? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> N/A					
b) Within a consecutive 12-month period while on the MS DMT, did the patient experience at least two relapses of MS? <input type="checkbox"/> No <input type="checkbox"/> Yes → Provide the dates of either two clinical relapses OR one clinical relapse and one MRI relapse					
Date of relapse (YYYY-MM-DD)		Type of relapse (One MRI relapse may substitute for one clinical relapse)			
		<input type="checkbox"/> Moderate to very severe clinical relapse		<input type="checkbox"/> MRI relapse (new T2 lesion or definite gadolinium-enhancing T1 lesion)	
		<input type="checkbox"/> Moderate to very severe clinical relapse		<input type="checkbox"/> MRI relapse (new T2 lesion or definite gadolinium-enhancing T1 lesion)	
PRESCRIBER'S SIGNATURE		DATE (YYYY-MM-DD)		Please forward this request to Alberta Blue Cross, Clinical Drug Services 10009 108 Street NW, Edmonton, Alberta T5J 3C5 FAX 780-498-8384 in Edmonton • 1-877-828-4106 toll free all other areas	
ONCE YOUR REQUEST HAS SUCCESSFULLY TRANSMITTED, PLEASE DO NOT MAIL OR RE FAX YOUR REQUEST.					

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