

REPATHA (evolocumab)

Instructions

Plan Member Signature

Please complete Part A and have your physician complete Part B. Completion and submission is not a guarantee of approval. Any fees related to the completion of this form are the responsibility of the plan member. Drugs in the Prior Authorization Program may be eligible for reimbursement if the patient does not qualify for coverage under a primary plan or a government program. Drugs used for indications not approved by Health Canada may be denied. For Quebec plan members, RAMQ exception drug criteria may apply. The decision for approval versus denial is based on pre-defined clinical criteria, primarily based on Health Canada approved indication(s) and on supporting evidence-based clinical protocols. The plan member will be notified whether their request has been approved or denied. Please note that you have the right to appeal the decision made by Express Scripts Canada.

Part A - Patient Patient information First Name: Last Name: Insurance Carrier Name/Number: Group Number: Client ID: Relationship: Employee Spouse Dependent Date of Birth (YYYY/MM/DD): Gender: Male Female Language: | English | French Address: City: Province: Postal Code: Email address: Telephone (home): Telephone (cell): Telephone (work): Coordination of benefits **Patient** Is the patient enrolled in any patient assistance program? Yes No **Assistance Program** Contact Name: _ Has the patient applied for reimbursement under a provincial plan? Yes No N/A **Provincial** Coverage What is the coverage decision of the drug? Approved Denied *Attach decision letter* Has the patient applied for reimbursement under a primary plan? | Yes | No | N/A **Primary** Coverage What is the coverage decision of the drug? Approved Denied *Attach decision letter* **Authorization** On behalf of myself and my eligible dependents, I authorize my group benefit provider, and its agents, to exchange the personal information contained on this form. I give my consent on the understanding that the information will be used solely for purposes of administration and management of my group benefit plan. This consent shall continue so long as my dependents and I are covered by, or are claiming benefits under the present group contract, or any modification, renewal, or reinstatement thereof.

Date



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Part B - Prescriber

Please see instructions on page 1 and complete all sections below. <u>Incomplete forms may result in automatic denial</u>. Please do **not** provide genetic test information or results.

REPATHA (evolocumab)			New request Renewal request*			
Dose	Administration (ex:	oral, IV, etc)	Frequency	Duration		
Site of drug administration:						
☐ Home ☐ Physic	ian's office/Infusion clir	nic Hos	oital (outpatient)	Hospital (inpatient)		
* Please submit proof of price	or coverage if available					
ECTION 2 - ELIGIBILITY	CRITERIA					
Please indicate if the pa	tient satisfies the belov	/ criteria:				
Atherosclerotic Cardiovascul	ar Disease					
<u>INITIAL</u>						
following: ischemic		istory of heart atta		dult defined by one of the disease (stroke), and/or		
	g one moderate-to-high rapies in the chart belo		as a documented into	olerance to at least 2 statins		
The patient has had AND	an inadequate respons	se or has a docume	nted intolerance or c	ontraindication to ezetimibe,		
	ite taking a maximally t			L or greater, or Apo-B level is 0.7 east one of the patient's lipid		
Date (YYYY-MM-DD)	LDL-C (mmol/L)	non-HDL-C (mmol/L	Apo-B (g/L)			
RENEWAL						
	nonstrated LDL-C, non-F nd current lipid parame		uction to target. Pleas	se indicate at least one of the		
	BASE	LINE				
Date (YYYY-MM-DD)	LDL-C (mmol/L)	non-HDL-C (mmol/L	Apo-B (g/L)			
L			_1			
	CURR	ENT				
Date (YYY-MM-DD)	LDL-C (mmol/L)	non-HDI -C (mmol/l	Ano-R (g/L)			



REPATHA (evolocumab)

Heteroz	ygous Familial Hyperch	nolesterolemia					
INITIAL							
	For the treatment of heterozygous familial hypercholesterolemia (HeFH) in patients unable to reach target LDL-C levels, AND						
	The patient is 10 year	s of age or older, ANI)				
	The patient is currently receiving a maximally-tolerated dose of statin therapy (Please list prior therapies in the chart below), OR						
	The patient has a documented intolerance or contraindication to at least two different statins (Please list prior therapies in the chart below), AND						
	The patient has had an inadequate response or has a documented intolerance or contraindication to ezetimibe, AND						
	The patient's LDL-C le	vel is 2 mmol/L or gr	eater despite current t	therapy, OR			
	The patient has not ac	chieved a 50% reduct	tion in LDL-C from pre-	treatment levels desp	oite current therapy, OR		
	The patient's non-HDL-C level is 2.4 mmol/L or greater, or Apo-B level is 0.7 g/L or greater despite current therapy. Please indicate at least one of the patient's lipid parameter levels below:						
	Date (YYYY-MM-DD)	LDL-C (mmol/L)	non-HDL-C (mmol/L)	Apo-B (g/L)			
RENEW	RENEWAL The patient has demonstrated LDL-C, non-HDL-C, or Apo-B reduction to target. Please indicate at least one of the patient's baseline and current lipid parameter levels below:						
	BASELINE						
	Date (YYYY-MM-DD)	LDL-C (mmol/L)	non-HDL-C (mmol/L)	Apo-B (g/L)			
		CUR					
	Date (YYYY-MM-DD)	LDL-C (mmol/L)	non-HDL-C (mmol/L)	Apo-B (g/L)			



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omozy	gous Familial Hyperch	olesterolemia			
<u>IITIAL</u>					
	For the treatment of homozygous familial hypercholesterolemia (HoFH) defined by a family history of FH in both parents and/or premature atherosclerotic cardiovascular disease, AND				
	The patient is 10 years of age or older, AND				
	The patient has pre-treatment LDL-C levels greater than 12 mmol/L, AND				
	The patient had tendon xanthomas before the age of 10, AND				
	The patient is taking of	one high-intensity sta	tin therapy (<i>Please lis</i> i	t prior therapies in th	e chart below), OR
	The patient has a documented intolerance or contraindication to at least two different statins (<i>Please list prior therapies in the chart below</i>), AND				
	The patient has had an inadequate response or has a documented intolerance or contraindication to ezetimibe, AND				
	The patient's LDL-C le	evel is 2 mmol/L or gr	reater despite current	therapy, OR	
	The patient has not achieved a 50% reduction in LDL-C from pre-treatment level despite current therapy. Please indicate patient's current LDL-C level below:				
	Date (YYYY-MM-DD)	LDL-C (mmol/L)			
ENEW <i>E</i>		onstrated LDL-C reduc	ction to target. Please	indicate patient's ba	seline and current LDL-C level
	BASELINE		CURRENT		
	Date (YYYY-MM-DD)	LDL-C (mmol/L)	Date (YYYY-MM-DD)	LDL-C (mmol/L)	-
					-
					J
R □	Name of the all and a	taria anni			
Ш	None of the above cri	tena applies.			
Rele	evant additional inform	nation:			



REPATHA (evolocumab)

	Decede and	Duration of therapy		Reason for cessation	
Drug	Dosage and administration	From	То	Inadequate response	Allergy/ Intolerance

SECTION 3 - PRESCRIBER INFORMATION

Physician's Name:	
Address:	
Tel:	Fax:
License No.:	Specialty:
Physician Signature:	Date:

Please fax or mail the completed form to Express Scripts Canada®

Fax: Express Scripts Canada Clinical Services 1 (855) 712-6329

Mail: Express Scripts Canada Clinical Services 5770 Hurontario Street, 10th Floor Mississauga, ON L5R 3G5