

RINVOQ (upadacitinib)

#### **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: Date of Birth (YYYY/MM/DD): Relationship: | Employee | Spouse | Dependent Language: English French Gender: | | Male | | Female 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.

SECTION 1 - DRUG REQUESTED										
RINVOQ (upadacitinib)		New request	Renewal request*							
Dose	Administration (ex: oral, IV, etc)	Frequency	Duration							
Site of drug administration:		_								
Home Physician's office/Infusion clinic Hospital (outpatient) Hospital (inpatient)										
* Please submit proof of prior of	coverage if available									
SECTION 2 – ELIGIBILITY CRITERIA										
Please indicate if the patient satisfies the below criteria:										
Rheumatoid Arthritis										
For the treatment of moderately to severely active rheumatoid arthritis in an adult, AND										
	ninadequate response to a minim ying anti-rheumatic drug (DMARD									
Where combinations of non-biologic DMARDs are impossible, the patient has tried 3 consecutive non-biologic DMARDs, unless patient has a documented intolerance to DMARDs ( <i>Please list prior therapies in the chart below</i> )										
Psoriatic Arthritis										
For the treatment of ac	ctive psoriatic arthritis in an adult,	, AND								
The patient has had an inadequate response or has a documented intolerance to at least 2 disease modifying anti- rheumatic drugs (DMARDs), or to another biologic response modifier ( <i>Please list prior therapies in the chart below</i> )										
Ankylosing Spondylitis										
For the treatment of ar	nkylosing spondylitis in an adult, A	AND								
The patient has a Bath scale, AND	Ankylosing Spondylitis Disease A	ctivity Index (BASDAI) score	of 4 or greater on a 10-point							
	inadequate response or has a do SAIDs) for a minimum of 2 weeks		least 2 non-steroidal anti-							
The patient has had an (Please list prior therag	inadequate response or has a do pies in the chart below)	ocumented intolerance to a	biologic response modifier							



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ALODIC DE	rmotitio									
<u>INITIAL</u>	ermatitis									
	For the treatment of moderate-to-severe atopic dermatitis (AD), AND									
_	The patient is 12 years of age or older, AND									
	The patient has an affected body surface area (BSA) of 10% or greater, or there is involvement of the patient's face,									
_	hands, feet or genital region, AND  The patient has an Investigator's Global Assessment (IGA) score of 3 or greater, AND									
_	The patient has an Eczema Area and Severity Index (EASI) score of 16 or greater, AND									
_	The patient has an Eczema Area and Seventy Index (EASI) score of 16 or greater, AND  The patient has had an inadequate response or has a documented intolerance to at least 2 topical agents that are									
					the chart below), AND					
	The patient has had an inadequate response or has a documented intolerance to a systemic treatment, if an adult (Please list prior therapies in the chart below)									
RENEWA	I									
	– The patient has dem		ent defined as 75% o Irrent EASI score belo		ent from baseline in EASI score.					
	BASE	LINE	CURRENT							
-	Date (YYYY-MM-DD)									
	Date (1111-WIWI-DD)	EASI score	Date (YYYY-MM-DD)	EASI score						
	Date (TTT-WIW-DD)	EASI score	Date (YYYY-MM-DD)	EASI score						
_	iographic Axial Spond	dyloarthritis								
	iographic Axial Spond	<b>dyloarthritis</b> non-radiographic axi ctive signs of inflamr	al spondyloarthritis ir	n an adult, AND	e protein (CRP) and/or magnetic					
	iographic Axial Spond For the treatment of The patient has object resonance imaging (I	dyloarthritis non-radiographic axi ctive signs of inflamr MRI), AND an inadequate respo	al spondyloarthritis in mation as indicated bonse or has a docume	n an adult, AND y elevated C-reactive ented intolerance to	e protein (CRP) and/or magnetic at least 2 non-steroidal anti- apies in the chart below), AND					
	iographic Axial Spond For the treatment of The patient has object resonance imaging (I The patient has had a inflammatory drugs (	dyloarthritis non-radiographic axi ctive signs of inflamr MRI), AND an inadequate respo	al spondyloarthritis in mation as indicated bonse or has a docume um of 2 weeks each	n an adult, AND y elevated C-reactive ented intolerance to (Please list prior the	at least 2 non-steroidal anti-					
	iographic Axial Spond For the treatment of The patient has object resonance imaging (I The patient has had a inflammatory drugs ( The patient has had a rheumatic drug (DMA	dyloarthritis non-radiographic axi ctive signs of inflamr MRI), AND an inadequate respo	al spondyloarthritis in mation as indicated bonse or has a docume um of 2 weeks each conse or has a docume	n an adult, AND y elevated C-reactive ented intolerance to (Please list prior the	at least 2 non-steroidal anti- rapies in the chart below), AND					
Ulcerative	iographic Axial Spond For the treatment of The patient has object resonance imaging (I The patient has had a inflammatory drugs ( The patient has had a rheumatic drug (DMA	dyloarthritis non-radiographic axi ctive signs of inflamm MRI), AND an inadequate respo NSAIDs) for a minim an inadequate respo ARD) (Please list prio	al spondyloarthritis in mation as indicated bonse or has a docume um of 2 weeks each onse or has a docume or therapies in the charterists.	n an adult, AND y elevated C-reactive ented intolerance to (Please list prior the ented intolerance to art below)	at least 2 non-steroidal anti- capies in the chart below), AND a biologic disease modifying anti-					
JIcerative	iographic Axial Spond For the treatment of The patient has object resonance imaging (I The patient has had a inflammatory drugs ( The patient has had a rheumatic drug (DMA)	dyloarthritis non-radiographic axi ctive signs of inflammod inadequate responsion of a minimal and inadequate responsion (Please list priomoderately to severe	al spondyloarthritis in mation as indicated bonse or has a docume um of 2 weeks each conse or has a docume or therapies in the character dely active ulcerative conservative c	n an adult, AND y elevated C-reactive ented intolerance to (Please list prior the ented intolerance to art below)  olitis in an adult, AN	at least 2 non-steroidal anti- rapies in the chart below), AND a biologic disease modifying anti-					
Ulcerative	iographic Axial Spond For the treatment of The patient has object resonance imaging (I The patient has had a Inflammatory drugs ( The patient has had a Irheumatic drug (DMA  e Colitis  For the treatment of The patient has had a	dyloarthritis non-radiographic axi ctive signs of inflammoderately to severe an inadequate responsion inadequate responsion (Please list prior)	al spondyloarthritis in mation as indicated bonse or has a docume um of 2 weeks each conse or has a docume or therapies in the character dely active ulcerative conservative c	n an adult, AND y elevated C-reactive ented intolerance to (Please list prior the ented intolerance to art below)  olitis in an adult, AN ented intolerance to	at least 2 non-steroidal anti- rapies in the chart below), AND a biologic disease modifying anti- D corticosteroids and to either					



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Crohn's Disease									
_	rately to severely active Croh	ın's disease in a	n adult. AND						
_	For the treatment of moderately to severely active Crohn's disease in an adult, AND  The patient has had an inadequate response or has a documented intolerance to either aminosalicylates,								
	ticosteroids (Please list prior				,				
The patient has tried and f	ailed a biologic response mo	difier (Please lis	t prior therapie	es in the chart belo	ow)				
OR									
None of the above criteria	applies.								
Delevent additional information									
Relevant additional information	1:								
2. Please list previously tried ther	apies								
Drug	Dosage and administration	Duration	Duration of therapy		Reason for cessation				
		From	То	Inadequate response	Allergy/ Intolerance				
		110							
	'			•					
SECTION 3 - PRESCRIBER INF	FORMATION								
Physician's Name:									
A d d.v									
Address:									
el:		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, 10<sup>th</sup> Floor Mississauga, ON L5R 3G5