

SPECIAL AUTHORITY REQUEST ADALIMUMAB / INFLIXIMAB / VEDOLIZUMAB FOR MODERATE TO SEVERE ACTIVE CROHN'S / FISTULIZING CROHN'S DISEASE

HLTH 5368 Rev. 2023/05/03

INITIAL/SWITCH COVERAGE

For up-to-date criteria and forms, please check: www.gov.bc.ca/pharmacarespecialauthority

Fax requests to 1-800-609-4884 (toll free) OR mail requests to: PharmaCare, Box 9652 Stn Prov Govt, Victoria, BC V8W 9P4 This facsimile is doctor-patient privileged and contains confidential information intended only for PharmaCare. Any other distribution, copying or disclosure is strictly prohibited.

If you have received this fax in error, please write MISDIRECTED across the front of the form and fax toll-free to 1-800-609-4884, then destroy the pages received in error.

If PharmaCare approves this Special Authority request, approval is granted solely for the purpose of covering prescription costs. PharmaCare approval does not indicate that the requested medication is, or is not, suitable for any specific patient or condition.

Forms with information missing will be returned for completion. If no prescriber fax or mailing address is provided, PharmaCare will be unable to return a response

SECTION 1 – PRESCRIBING GASTROENTEROLOGIST'S INFO.	SECTION 2 - PATIENT INFORMATION		
Name and Mailing Address	Patient (Family) Name		
	Patient (Given) Name(s)		
College ID (use ONLY College ID number) Phone Number (include area code)	Date of Birth (YYYY / MM / DD) Date of Application (YYYY / MM / DD)		
CRITICAL FOR A TIMELY RESPONSE Prescriber's Fax Number	CRITICAL FOR PROCESSING Personal Health Number (PHN)		
SECTION 3 – MEDICATION REQUESTED			
ADALIMUMAB 12 week supply: 160 mg week 0, 80 mg week 2, then 40 mg weeks 4, 6, 8, and 10	INFLIXIMAB 3 doses: 5 mg/kg at 0, 2, and 6 weeks		
O ABRILADA® O AMGEVITA® O HADLIMA® O HULIO® O HYRIMOZ® O IDACIO® O SIMLANDI™ O YUFLYMA®	O AVSOLA™ INFLECTRA® RENFLEXIS® VEDOLIZUMAB 3 doses: 300 mg at 0, 2, and 6 weeks 12 weeks: 300 mg IV at 0, 2, and 6 weeks, plus 108 mg SC at weeks 8, 10, and 12		
ECTION 4 - PRE-TREATMENT CLINICAL INFORMATION			
Diagnosis Moderate to Severe Active Crohn's Active Fistulizing Crohn's	Year of Crohn's Diagnosis Current Weight in Kg		
Current Steroid Dose	Impact of Current Condition on Work/Social Life O None O Mild O Moderate O Severe		
FOR MODERATE TO SEVERE CROHN'S			
Site of Crohn's Olsolated Colonic Olsolated Colonic Small Bowel Other (specify)	Current Harvey Bradshaw Index (Hbi ≥ 8)		
FOR ACTIVELY FISTULIZING CROHN'S			
Site of Fistula(e) Perianal Enterocutaneous Recto-Vaginal Other (specify)	Number of Fistulae		
Fistula Drainage and Bleeding: Onone Omild Omoderate Osevere	Pain at Fistula Sites: None Mild Moderate Severe		
For consideration of off-criteria requests, additional information demonstrating mo assessment is required within 9 months of starting treatment to demonstrate medi Colonoscopy Fecal Calprotectin Level Dother (specify)			
SECTION 5 – CONCURRENT THERAPY INCLUDE ALL antidia	urrheals, narcotics, immunosuppressants, antibiotics		
DRUG, DOSE/ROUTE, FREQUENCY			
1			
2			
3			
PHARMACARE USE ONLY	Continued on page 2 >		
Status	ve Date (YYYY / MM / DD) Duration of Approval		

PATIENT NAME			PHN	N DATE (YYYY / MM / DD)	
ECTION 6 - PRIOR THERAPIE	S (INITIAL COVERAG	iE)			
OR PATIENTS WITH MODERATE TO	SEVERE ACTIVE CROHNS	- DETAILS OF	GLUCO	OCORTICOID TRIAL (REQUIRED)	
O Patient is steroid resistant , di	isplaying a lack of a sympto , unable to withdraw oral co	matic respons rticosteroid wit	e to then thin 3 m	nonths of initiation without a recurrence of symptoms; a symptomatic relap	
O Patient is unable to complete	a course of steroids eq	uivalent to o	ral pre	ednisone 40 mg or more daily for a minimum of 14 days.	
O Corticosteroid use is contrain	dicated (specify):				
O Intolerances/side effect(s) (spe	ecify):				
FOR PATIENTS WITH FISTULIZING C	ROHNS				
Drug Name, Dose,	Duration (required)			Details of Outcome (Failure, Contraindication, Intolerance, Other)	
Ciprofloxacin at maximally tolerate	d doses (min 3 week trial), w	vith/without	○ La	ack of Effect O Contraindication O Intolerance O Other	
☐ Metronidazole at maximally tole	e at maximally tolerated doses (min 3 week trial)		Specify:		
OR PATIENTS WITH MODERATE TO SURGICAL HISTORY AND PRIOR MEI		AND/OR FIST	ULIZIN	VG CROHNS	
MEDICATION	STARTING DATE	DURATION (OF USE	OUTCOME (specify intolerances, primary/secondary failures)	
PRIOR SURGERIES					
☐ None ☐ Ostomy* ☐ Pouch*	Prior resections*	*Please inclu with each co		ent's normal number of liquid bowel movements and HBI worksheet request	
ECTION 7 – ADDITIONAL INF	ORMATION, IF APPL	ICABLE			
ECTION 8 – PRESCRIBER SIG	NATURE				
Personal information on this form is collected with, the <i>British Columbia Pharmaceutical Serv.</i> Protection of Privacy Act 26 (a),(c),(e). The inform of (a) administering the PharmaCare program	ices Act 22(1) and Freedom of Info mation is being collected for the	rmation and purposes	infor	ve discussed with the patient that the purpose of releasing their rmation to PharmaCare is to obtain Special Authority for prescript erage and for the purposes set out here.	

PharmaCare may request additional documentation to support this Special Authority request. Actual reimbursement is subject to the rules of a patient's PharmaCare plan, including any annual deductible requirement, and to any other applicable PharmaCare pricing policy.

Prescriber's Signature (Mandatory)

Special Authority and other Ministry programs and (c) to manage and plan for the health system generally. If you have any questions about the collection of this information, call Health Insurance BC from Vancouver at 1-604-683-7151 or from elsewhere in BC toll free at 1-800-663-7100 and ask to consult a pharmacist concerning the Special Authority process.