

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

ADALIMUMAB for Hidradenitis Suppurativa SPECIAL AUTHORIZATION REQUEST FORM

Patients may or may not meet eligibility requirements as established by Alberta government-sponsored drug programs.

PATIENT INFORMATION				COVERAGE TYPE	
LAST NAME	FIRST NAME	INITIAL	<input type="checkbox"/> Alberta Blue Cross <input type="checkbox"/> Alberta Human Services <input type="checkbox"/> Other _____		
BIRTHDATE (YYYY/MM/DD)	ALBERTA PERSONAL HEALTH NUMBER				
ADDRESS	CITY	PROV.	POSTAL CODE	ID/CLIENT/COVERAGE NUMBER	

PRESCRIBER INFORMATION					
PRESCRIBER LAST NAME	FIRST NAME	INITIAL	PRESCRIBER PROFESSIONAL ASSOCIATION REGISTRATION		
ADDRESS			<input type="checkbox"/> CPSA	<input type="checkbox"/> ACO	REGISTRATION NUMBER
			<input type="checkbox"/> CARNA	<input type="checkbox"/> ADA+C	
CITY, PROVINCE			<input type="checkbox"/> ACP	<input type="checkbox"/> Other	
POSTAL CODE			PHONE	FAX	
			FAX NUMBER MUST BE PROVIDED WITH EACH REQUEST SUBMITTED		

Please provide the following information for ALL requests

Diagnosis Active moderate to severe Hidradenitis Suppurativa
 Other (specify) _____

Please provide the following information for INITIAL requests

- 1) Total abscess and nodule (AN) count at pre-treatment baseline _____ and date of count _____
- 2) Does the patient have lesions in at least two distinct anatomical areas? Yes No
- 3) Does the patient have Hurley Stage II or III lesions in at least one anatomical area? Yes No
- 4) Previous therapy
 - a) Have systemic antibiotics been tried for at least 90 days?
 - Yes → Specify antibiotics and response _____
 - No → Specify reason _____
 - b) Is there documented non-response to conventional therapy other than systemic antibiotics?
 - Yes → Specify which therapies have been tried, including dose and duration _____
 - No → Specify reason _____

Please provide the following information for RENEWAL requests

- 1) Current AN count _____ and date of count _____
- 2) Indicate the patient's response to treatment (check ALL that apply)
 - Fifty per cent or greater reduction in AN count from pre-treatment baseline.
 - No increase in abscess count or draining fistula count relative to pre-treatment baseline.

Note: Treatment with adalimumab should be discontinued if there is insufficient improvement after 12 weeks of treatment.

Additional information relating to request

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.

