

One Island Health System

PEI Pharmacare P.O. Box 2000 Charlottetown, PE C1A 7N8 www.healthpei.ca



www.healthpei.ca

Programmes provinciaux de medicaments C.P. 2000 Charlottetown, PE C1A 7N8

PEI Pharmacare Bulletin

Issue (2021 - 3) March 15, 2021

NEW PRODUCT(S) ADDED TO THE PEI PHARMACARE FORMULARY (EFFECTIVE DATE: MARCH 29, 2021)

Product (Generic name)	Product (Brand name)	Strength	Dosage Form	DIN	MFR
Benralizumab	Fasenra	30 mg/ml	Syringe	02473232	AZN
		30 mg/ml	Autoinjector	02496135	
Criteria	As add-on maintenance treatment for adult patients with severe eosinophilic asthma, if				sthma, if
	the following criteria are met:				
	Initiation Criteria:				
	 Patient must have 	e a documented d	iagnosis of asthma.		
	 Patient is inadequately controlled with high-dose inhaled corticosteroids, defined as greater or equal to 500 mcg of fluticasone propionate or equivalent daily, and one or more additional asthma controller(s) (e.g., long-acting beta agonists). Patient has one of the following: blood eosinophil count of ≥ 300 cells/μL within the past 12 months AND has experienced two or more clinically significant asthma exacerbations in the past 12 months, or blood eosinophil count of ≥ 150 cells/μL AND is receiving maintenance treatment with oral corticosteroids (OCS). 				
	Renewal Criteria:				
	 The effects of treatment should be assessed every 12 months to determine whether reimbursement should continue. 				mine
	Reimbursement of	of treatment shou	ld be discontinued if:		
	from bas the asthm months o	eline, when baseli na control questio of therapy has not	control questionnaire score has not improved paseline represents the initiation of treatment, or uestionnaire score achieved after the first 12 s not been maintained subsequently, or		
	 the number of clinically significant exacerbations has increased with the previous 12 months, or in patients on maintenance treatment with OCS, there has been not decrease in the OCS dose in the first 12 months of treatment, or in patients on maintenance treatment with OCS, the reduction in dose of OCS achieved after the first 12 months of treatment is not maintained subsequently. 				d within
					or

	Clinical Notes:
	 Benralizumab should not be used in combination with other biologics used to treat asthma.
	 A baseline assessment of asthma symptom control using a validated asthma control questionnaire must be completed prior to initiation of benralizumab treatment.
	 Patients should be managed by a physician with expertise in treating asthma.
Program Eligibility	High Cost Drug Program, Catastrophic Drug Program

0 0 ,	1 2 3 3 7		,			
24 1: 1		100	\" I	00440704	0014	
Mepolizumab	Nucala	100 mg	Vial	02449781	GSK	
		100 mg/ml	Autoinjector	02492989		
		100 mg/ml	Syringe	02492997		
Criteria	As add-on maintenance treatment for adult patients with severe eosinophilic asthma, if the following criteria are met: Initiation Criteria: Patient must have a documented diagnosis of asthma. Patient is inadequately controlled with high-dose inhaled corticosteroids, defined as greater or equal to 500 mcg of fluticasone propionate or equivalent daily, and one or more additional asthma controller(s) (e.g., long-acting beta agonists). Patient has one of the following: blood eosinophil count of ≥ 300 cells/µL within the past 12 months AND					
	has expering the passion of the pass	rienced two or mo st 12 months, or	ore clinically signific ≥ 150 cells/µL AND i	ant asthma exa	cerbations	
	Renewal Criteria:					
	 The effects of treatment should be assessed every 12 months to determine whether reimbursement should continue. 					
	Reimbursement of treatment should be discontinued if:					
	 the 12 month asthma control questionnaire score has not improved 					
	from baseline, when baseline represents the initiation of treatment, or					
	 the asthma control questionnaire score achieved after the first 12 months of therapy has not been maintained subsequently, or 					
	o the numb					
	o in patient decrease	 in patients on maintenance treatment with OCS, there has been no decrease in the OCS dose in the first 12 months of treatment, or 				
	dose of C		e treatment with OC the first 12 months			
	Clinical Notes:					
	 Mepolizumab should not be used in combination with other biologics used to treat asthma. 					
	 A baseline assessment of asthma symptom control using a validated asthma control questionnaire must be completed prior to initiation of mepolizumab treatment. 					
	 Patients should b 	e managed by a p	hysician with exper	tise in treating	asthma.	
Program Eligibility	High Cost Drug Program,	Catastrophic Drug	g Program			

Product (Generic name)	Product (Brand name)	Strength	Dosage Form	DIN	MFR	
Omalizumab	Xolair	150 mg	Vial	02260565	NVR	
Criteria	For the treatment of patients ≥ 12 years of age with moderate to severe chronic					
	idiopathic urticaria (CIU) who remain symptomatic (presence of hives and/or associated				ssociated	
	itching) despite optimum management with H1 antihistamines.					
	Initiation Criteria:					
	 Documentation of the most recent Urticaria Activity Score over 7 days (UAS7) to be 				S7) to be	
	provided on the submitte	•				
	 Approvals will be for a maximum dose of 300mg every four weeks. Initial approval period: 24 weeks. Renewal Criteria: Requests for renewal will be considered if the patient has achieved: complete symptom control for less than 12 consecutive weeks; or partial response to treatment, defined as at least a ≥ 9.5 point reduction in baseline UAS7 					
					auction in	
	Clinical Notes:					
	1. Moderate to severe CIU is defined as a UAS7 ≥16.					
	2. Treatment cessation could be considered for patients who experience complete symptom control for at least 12 consecutive weeks at the end of a 24 week treatment period.				lete	
	3. In patients who discontinue treatment due to temporary symptom control, re-					
	initiation can be considered if CIU symptoms reappear.					
	4. Optimal management	is defined as H1 ar	ntihistamines at up to	o 4 times the st	andard	
	daily dose					
Program Eligibility	High Cost Drug Program,	Catastrophic Drug	g Program			

BENEFIT STATUS/CRITERIA CHANGES (EFFECTIVE DATE: MARCH 15, 2021)

Product (Generic name)	Product (Brand name)	Strength	Dosage Form	DIN/PDIN	MFR
Bisacodyl	Magic Bullet	10 mg	Suppository	02241091	D&C
Criteria	Criteria The Special Authorization criteria has expanded to include: For use as part of a bowel program for neurogenic bowel dysfunction in patients with spinal cord injuries				
Program Eligibility	Family Health Benefit Drug Program, Financial Assistance Program, Nursing Home Dr				me Drug
	Program, Seniors Drug Pr	ogram	_	_	_