# Breztri (budesonide/glycopyrrolate/formoterol) Prior Authorization Request Form

Caterpillar Prescription Drug Benefit Phone: 877-228-7909 Fax: 800-424-7640

**Instructions:** Please fill out all applicable sections completely and legibly. Attach any additional documentation that is important for the review (e.g., chart notes or lab data, to support the authorization request). Information contained in this form is Protected Health Information under HIPAA.

MEMBER INFORMATION				
LAST NAME:	FIRST NAME:			
PHONE NUMBER:	DATE OF BIRTH:			
STREET ADDRESS:				
CITY:	STATE: ZIP CODE:			
PATIENT INSURANCE ID NUMBER:				
MALE FEMALE HEIGHT (IN/CM): WEIGH	HT (LB/KG): ALLERGIES:			

IF YOU ARE NOT THE PATIENT OR THE PRESCRIBER, YOU WILL NEED TO SUBMIT A PHI DISCLOSURE AUTHORIZATION FORM WITH THIS REQUEST WHICH CAN BE FOUND AT THE FOLLOWING LINK: PRIMETHERAPEUTICS.COM/NOPP

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PRESCRIBER INFORMATION				
LAST NAME:	FIRST NAME:			
PRESCRIBER SPECIALTY:	EMAIL ADDRESS:			
NPI NUMBER:	DEA NUMBER:			
PHONE NUMBER:	FAX NUMBER:			
STREET ADDRESS:				
CITY:	STATE: ZIP CODE:			
REQUESTOR (if different than prescriber):	OFFICE CONTACT PERSON:			

MEDICATION OR MEDICAL DISPENSING INFORMATION					
MEDICATION NAME:					
DOSE/STRENGTH:	FREQUENCY:	LENGTH OF THERAPY/REFILLS:	QUANTITY:		
NEW THERAPY	RENEWAL	IF RENEWAL: DATE THERAPY	INITIATED:		
DURATION OF THERAPY (SPECIFIC DATES):					

Continued on next page



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MEMBER'S LAST NAME:	MEMBER'S FIRST NAME:				
1. HAS THE PATIENT TRIED ANY OTHER	R MEDICATIONS FOR THIS CONDITION?	YES (if yes, complete below) NO			
MEDICATION/THERAPY (SPECIFY DRUG NAME AND DOSAGE):	<b>DURATION OF THERAPY</b> (SPECIFY DATES):	RESPONSE/REASON FOR FAILURE/ALLERGY:			
2. LIST DIAGNOSES:		ICD-10:			
Chronic obstructive pulmonary disease(COPD) Other diagnosis:ICD-10					
<b>3. REQUIRED CLINICAL INFORMATION:</b> PLEASE PROVIDE ALL RELEVANT CLINICAL INFORMATION TO SUPPORT A PRIOR AUTHORIZATION.					
Clinical Information: Is this drug being prescribed to this patient as part of a treatment regimen specified within a sponsored clinical trial? Pres No Has the patient previously been treated with a combination long-acting beta2-adrenergic agonist/long-acting muscarinic agonist (LABA/LAMA) such as Anoro Ellipta (umeclidinium/vilanterol), Stiolto (tiotropium/olodaterol), Bevespi (glycopyrrolate/formoterol), or Utibron (indacaterol/glycopyrrolate)? Yes No Has the patient previously been treated with a combination inhaled corticosteroid/ long-acting beta2-adrenergic agonist such as Advair(fluticasone/salmeterol), AirDuo(fluticasone/salmeterol),Wixela(fluticasone/salmeterol), Symbicort(budesonide/formoterol), or Dulera(mometasone/formoterol)? Yes No					
Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?					
<b>Please note:</b> Not all drugs/diagnosis are covered on all plans. This request may be denied unless all required information is received.					
<b>ATTESTATION:</b> I attest the information provided is true and accurate to the best of my knowledge. I understand that the Health Plan, insurer, Medical Group or its designees may perform a routine audit and request the medical information necessary to verify the accuracy of the information reported on this form.					
Prescriber Signature or Electronic I.D.	Verification:	Date:			
you are not the intended recipient, you are here	ompanying this transmission contain confidential eby notified that any disclosure, copying, distribut have received this information in error, please no se documents.	tion, or action taken in reliance on the contents			



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MEMBER'S LAST NAME: \_\_\_\_\_

MEMBER'S FIRST NAME: \_\_\_\_\_

### FAX THIS FORM TO: 800-424-7640

MAIL REQUESTS TO: Prime Therapeutics Management Prior Authorization Program

Attn: CP - 4201 P.O. Box 64811 St. Paul, MN 55164-0811

