

## BENEFIT INVESTIGATION REQUEST AND PRESCRIPTION FORM

Phone: 855-802-8746 Fax: 855-454-8746 MyQUTENZAConnect.com Hours: (M-F) 9 AM-7 PM ET

Case ID: \_\_\_\_

Received: \_\_\_\_\_

PATIENT INFORM	ATION												
Last Name			First Name			MI	Sex Assignment <sup>1</sup>		Date of Birth		Phone		
					State	Male Female							
Street Address Cir			City	City			ZIP	Email	Email				
Patient's Initial Treatment?  Yes No – If No, Date of Initial Treatme				nent:			Allergies Anticipated Treatment Date						
MEDICAL INSURANCE - PRIMARY					MEDICAL INSURANCE - SECON			SECON	IDARY				
Plan Name	Phone	Phone			Plan Name Member ID				Phone				
Member ID	Group #	Group #							Group #				
PHARMACY INSURANCE - PRIMARY						PHARMA	PHARMACY INSURANCE - SECONDARY						
Member ID		BIN				Member I	D			BIN			
PCN		Group #				PCN			Group #				
PRESCRIBER INFO	ORMATION												
Prescriber's Full Na		Practice Name					e Contact						
Address									State	ZIP			
Phone	Fax	Fax			NPI Number			TAX ID Number					
CLINICAL INFORM	ATION												
ICD-10-CM Code	CPT Cod				of codes that may be appropriate can be found in the QUTENZA Reimbursement Guide. he physician's responsibility to provide the correct indication and codes.								
Postherpetic po	N) 🗌 Diabo	Diabetic peripheral neuropathy of the fe				et (DPN) 🗌 Other:							
Physician Office	🗌 Hosp	Hospital Outpatient			Other Site of Service:								
PRESCRIPTION IN	NFORMATION												
(capsaicin) 8% topical system		# of Topi	Quantity # of Topical Systems (280 cm <sup>2</sup> billing units)		□ 1 □ 2	Ilment Options Kit (carton includes 1 topical system and Cleansing Gel) NDC #72512-928-01 Kit (carton includes 2 topical systems and Cleansing Gel) NDC #72512-929-01 Kit (carton includes 4 topical systems and Cleansing Gel) NDC #72512-930-01						12-929-01	
HA1C Levels Auto Transfer		 er				Shipping			g Address (if different fror				
By checking this box, if Rx coverage is found, your prescription will be automatically transferred to a specialty pharmacy for fulfillment ATTACH THE PATIENT CHART AND / OR CLINIC.			AL DATA WITH THE SUBMISSION OF THIS INTAKE FORM TO BEGIN THE PA PROCESS.							s.			
PRESCRIBER'S SI	GNATURE <sup>2</sup>												
Automatically r	e-investigate patie	nt for potential	retreatment in	91 days									
Prescriber's Signature:				Date:									
		<ol> <li>Authorizat necessary</li> </ol>	tion for Releas federal and st	ate authorizatio	ormation	n: By signin consents fr	om my patient t	to allow i	me to r	elease health ir	nformation	have obtained al to My QUTENZA es of transmitting	

this prescription to the appropriate pharmacy designated by the patient utilizing their benefit plan. My signature on this form also provides

consent to contact this patient's insurance provider for this prescription on the prescriber's behalf.





### BENEFIT INVESTIGATION REQUEST AND PRESCRIPTION FORM

# INDICATION

QUTENZA<sup>®</sup> (capsaicin) 8% topical system is indicated in adults for the treatment of neuropathic pain associated with postherpetic neuralgia (PHN) or associated with diabetic peripheral neuropathy (DPN) of the feet.

## IMPORTANT SAFETY INFORMATION

Do not dispense QUTENZA to patients for selfadministration or handling. Use only on dry, unbroken skin. Only physicians or healthcare professionals are to administer and handle QUTENZA, following the procedures in the label.

#### Warnings and Precautions

- Severe Irritation: Whether applied directly or transferred accidentally from other surfaces, capsaicin can cause severe irritation of eyes, mucous membranes, respiratory tract, and skin to the healthcare professional, patients, and others. Do not use near eyes or mucous membranes, including face and scalp. Take protective measures, including wearing nitrile gloves and not touching items or surfaces that the patient may also touch. Flush irritated mucous membranes or eyes with water and provide supportive medical care for shortness of breath. Remove affected individuals from the vicinity of QUTENZA. Do not re-expose affected individuals to QUTENZA if respiratory irritation worsens or does not resolve. If skin not intended to be treated comes into contact with QUTENZA, apply Cleansing Gel and then wipe off with dry gauze. Thoroughly clean all areas and items exposed to QUTENZA and dispose of properly. Because aerosolization of capsaicin can occur with rapid removal, administer QUTENZA in a wellventilated area, and remove gently and slowly, rolling the adhesive side inward.
- Application-Associated Pain: Patients may experience substantial procedural pain and burning upon application and following removal of QUTENZA. Prepare to treat acute pain during and following application with local cooling (e.g., ice pack) and/or appropriate analgesic medication.

- Increase in Blood Pressure: Transient increases in blood pressure may occur with QUTENZA treatment. Monitor blood pressure during and following treatment procedure and provide support for treatment-related pain. Patients with unstable or poorly controlled hypertension, or a recent history of cardiovascular or cerebrovascular events, may be at an increased risk of adverse cardiovascular effects. Consider these factors prior to initiating QUTENZA treatment.
- Sensory Function: Reductions in sensory function (generally minor and temporary) have been reported
- following administration of QUTENZA. All patients with sensory deficits should be assessed for signs of sensory deterioration or loss prior to each application of QUTENZA. If sensory loss occurs, treatment should be reconsidered.

#### Adverse Reactions

The most common adverse reactions ( $\geq$ 5% and > control group) in all controlled clinical trials are application site erythema, application site pain, and application site pruritus.

To report SUSPECTED ADVERSE REACTIONS, contact Averitas Pharma, Inc. at 1-877-900-6479 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

Please see accompanying full Prescribing Information.

