

Qutenza (capsaicin) Order Form

Patient Name:		DOB:) V	∕lale □ Female	
Diagnosis (please provide ICI	D10 code)						
Other:		□ NKDA Allergies:					
☐ New Start Therapy ☐ Continuation of Therapy Date of				of last dose (if applicable):			
Ordering Provider:							
Provider NPI:		Phone:		Fax:			
Practice Address:		City:		State:		Zip Code:	
Practice Contact:	Practice Contact: Email (f			tes):			
QUTENZA ORDERS							
DOSING:			REQUIRED TESTING/LABS:				
1 Topical System (1 patch - 280cm² billing units) 2 Topical Systems (2 patches- 560cm² billing units) 3 Topical Systems (3 patches- 840cm² billing units)			Ø	✓ Clinical/Progress Notes, Labs, Tests supporting primary diagnosis (please attach)			
☐ 4 Topical Systems (4 patch	nes-1120cm billing units)			FREQUENCY:			
LOCATION OF PATCH AND APPLICATION TIME:			V	Every 3 months			
☐ Left Foot (Dv: Diabetic Per	ipheral Neuropathy: 30 minute a	annlication)		Other:			
Right Foot (Dx: Diabetic Peripheral Neuropathy: 30 minute appl)	REFILLS:			
Right Side (Dx Post Herpetic Neuralgia : 60 minute application)			<i>_</i>	l			
☐ Left Side (Dx Post Herpetic Neuralgia : 60 minute application)			☐ One year from date signed				
Infuse One Standing Orde							
Provide treatment under In and Action Plan for Infusion	fuse One's Clinical Guidelines, Medi n Reactions.	cation Safety	Pro	tocol, Emergency Guidelin	es,		
Provider Name							
Provider Signature				Date		_	

Capsaicin can cause severe irritation of eyes, mucous membranes, respiratory tract, and skin to the healthcare professional, patients, and others.

Patients may experience substantial procedural pain and burning upon application of Qutenza and following removal of Qutenza. Prepare to treat acute pain during and following the application procedure with local cooling (such as an ice pack) and/or appropriate analgesic medication.

ransient increases in blood pressure may occur with Qutenza treatment. Monitor blood pressure during and following the treatment procedure.

Reductions in sensory function, which were 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.