# Otezla® Specialty Pharmacy (SP) START Form

Step 1: Please complete this form if you'd like an SP to provide prior authorization support or to process a prescription.





Preferred SP name			Fax #	
Please note that if th	e patient's insurance mandates the use of	a different SP than wha	t is preferred, your preferred SP ma	ry need to transfer the prescription to the mandated
Patient and Pres	criber Information			
ection 1: Patient	Information			
lame (First. Middle.	Last)		Date o	of birth/
				State ZIP
	7			
action 2: Insuran	ce Information *Include both sides of y	vour nationt's insuran	re and prescription benefit card	
	tached Prescription benefit card att	•		
_		_		Insurance phone
				narmacy insurance
				x PCN (if applicable)
	Rx BIN (if appli			х РСП (і) арріїсавіе)
x group 1D	RX BIN (IJ appu	<i>Cable)</i>		
ection 3: Prescrib	er Information			
lame (First, Last)	Facility name			
ddress			City	State ZIP
hone	Fax	NPI #*R6	equired Offi	ce contact
	on (PA) Information			
	oriatic arthropathy) ) (For PsO ONLY):	Medication Orals Topicals Biologics Other ADDITIONAL MED		Duration/Reason for discontinuation
Prescription Info	rmation for Otezla® (apremilast) FO	R ORAL USE		
Starting with in-	office sample			
	le was provided to patient: /	1	In-office 2-WEEK TITRATION S	AMPLE x14 days 27 tablets 0 refills
	start date if you directly provided the			ANTELEXIT days, 27 cablets, 0 ferrits
Starting with the	e Specialty Pharmacy			
Titration Starter Pack Rx is only for patients who did not receive a sample during their office visit. The SP will notify the patient via telephone prior to each shipment			Maintenance Dose: 30 mg o  ☐ Twice daily ☐ Once-daily r	f Otezla® renal dose 30 mg (For patients with severe renal impairme
ffice visit. The SP will	☐ Titration Dose: 4-WEEK STARTER PACK x28 days, 55 tablets, 0 refills			The state of the s
,	-WEEK STARTER PACK x28 days, 55 table	ets, 0 refills		lls:   11 or  Other (enter #)
Titration Dose: 4		ets, 0 refills		lls:
Titration Dose: 4	ure (dispense as written)	ets, 0 refills		
Titration Dose: 4				



Encourage commercially insured patients to enroll in the combined Co-Pay & Bridge Program by scanning the QR code, visiting otezla.com, or calling 1-844-4OTEZLA (1-844-468-3952).

Please see the back page for Indications and Important Safety Information. Please <u>click here</u> for the full Prescribing Information for Otezla.



## INDICATIONS AND IMPORTANT SAFETY INFORMATION

## **INDICATIONS**

Otezla® (apremilast) is indicated for the treatment of adult patients with plaque psoriasis who are candidates for phototherapy or systemic therapy.

Otezla is indicated for the treatment of adult patients with active psoriatic arthritis.

Otezla is indicated for the treatment of adult patients with oral ulcers associated with Behcet's Disease.

#### IMPORTANT SAFETY INFORMATION

#### **Contraindications**

 Otezla is contraindicated in patients with a known hypersensitivity to apremilast or to any of the excipients in the formulation

## **Warnings and Precautions**

- Hypersensitivity: Hypersensitivity reactions, including angioedema and anaphylaxis, have been reported during postmarketing surveillance. If signs or symptoms of serious hypersensitivity reactions occur, discontinue Otezla and institute appropriate therapy
- Diarrhea, Nausea, and Vomiting: Cases of severe diarrhea, nausea, and vomiting were associated with the use of Otezla. Most events occurred within the first few weeks of treatment. In some cases, patients were hospitalized. Patients 65 years of age or older and patients taking medications that can lead to volume depletion or hypotension may be at a higher risk of complications from severe diarrhea, nausea, or vomiting. Monitor patients who are more susceptible to complications of diarrhea or vomiting; advise patients to contact their healthcare provider. Consider Otezla dose reduction or suspension if patients develop severe diarrhea, nausea, or vomiting
- Depression: Carefully weigh the risks and benefits of treatment with Otezla for patients with a history of depression and/or suicidal thoughts/behavior, or in patients who develop such symptoms while on Otezla. Patients, caregivers, and families should be advised of the need to be alert for the emergence or worsening of depression, suicidal thoughts or other mood changes, and they should contact their healthcare provider if such changes occur
  - Plaque Psoriasis: Treatment with Otezla is associated with an increase in depression. During clinical trials in patients with moderate to severe plaque psoriasis, 1.3% (12/920) of patients reported depression compared to 0.4% (2/506) on placebo. Depression was reported as serious in 0.1% (1/1308) of patients exposed to Otezla, compared to none in placebo-treated patients (0/506). Suicidal behavior was observed in 0.1% (1/1308) of patients on Otezla, compared to 0.2% (1/506) on placebo. One patient treated with Otezla attempted suicide; one patient on placebo committed suicide
  - <u>Psoriatic Arthritis</u>: Treatment with Otezla is associated with an increase in depression. During clinical trials, 1.0% (10/998) reported depression or depressed mood compared to 0.8% (4/495) treated with placebo. Suicidal ideation and behavior was observed in 0.2% (3/1441) of patients on Otezla,

- compared to none in placebo-treated patients. Depression was reported as serious in 0.2% (3/1441) of patients exposed to Otezla, compared to none in placebo-treated patients (0/495). Two patients who received placebo committed suicide compared to none on Otezla
- Behçet's Disease: Treatment with Otezla is associated with an increase in depression. During the clinical trial, 1% (1/104) reported depression or depressed mood compared to 1% (1/103) treated with placebo. No instances of suicidal ideation or behavior were reported in patients treated with Otezla or treated with placebo
- Weight Decrease: Monitor body weight regularly; evaluate unexplained or clinically significant weight loss, and consider discontinuation of Otezla
- Plaque Psoriasis: Body weight loss of 5-10% occurred in 12% (96/784) of patients with moderate to severe plaque psoriasis treated with Otezla and in 5% (19/382) of patients treated with placebo. Body weight loss of ≥10% occurred in 2% (16/784) of patients treated with Otezla compared to 1% (3/382) of patients treated with placebo
- <u>Psoriatic Arthritis</u>: Body weight loss of 5-10% was reported in 10% (49/497) of patients taking Otezla and in 3.3% (16/495) of patients taking placebo
- Behçet's Disease: Body weight loss of >5% was reported in 4.9% (5/103) of patients taking Otezla and in 3.9% (4/102) of patients taking placebo
- Drug Interactions: Apremilast exposure was decreased when Otezla was co-administered with rifampin, a strong CYP450 enzyme inducer; loss of Otezla efficacy may occur. Concomitant use of Otezla with CYP450 enzyme inducers (e.g., rifampin, phenobarbital, carbamazepine, phenytoin) is not recommended

## **Adverse Reactions**

- <u>Plaque Psoriasis</u>: The most common adverse reactions (≥ 5%) are diarrhea, nausea, upper respiratory tract infection, and headache, including tension headache. Overall, the safety profile of Otezla in patients with mild to moderate plaque psoriasis was consistent with the safety profile previously established in adult patients with moderate to severe plaque psoriasis
- <u>Psoriatic Arthritis</u>: The most common adverse reactions (≥ 5%) are diarrhea, nausea, and headache
- <u>Behçet's Disease</u>: The most common adverse reactions (≥ 10%) are diarrhea, nausea, headache, and upper respiratory tract infection

## **Use in Specific Populations**

 Otezla has not been studied in pregnant women. Advise pregnant women of the potential risk of fetal loss

Please <u>click here</u> for the full Prescribing Information for Otezla.



