# **Free Trial and Dose Modification Program**



#### **CONTACT INFORMATION**

Phone: 502-805-3469 | Fax: 502-212-1107 | Email Address: ZejulaDE@pharmacord.com

#### **ENROLLMENT FORM**

The ZEJULA Free Trial and Dose Modification Program is available for new patients or patients who require mid-cycle dose changes of their current tablet strength. A 30-count bottle of ZEJULA will be provided to your patient at no cost.

#### INSTRUCTIONS FOR PRESCRIBER



### Complete

Review and complete this entire form



#### Sign

Prescriber must sign and date at the bottom of **section 4** 



## **Fax**

Fax this entire form to: 502–212–1107

# **ELIGIBILITY REQUIREMENTS**

To be eligible for the ZEJULA Free Trial and Dose Modification Program, a patient must:

- Be a legal resident of the United States (including Washington DC, Puerto Rico, US Virgin Islands, or US territories)
- Be prescribed ZEJULA for a US Food and Drug Administration (FDA)-approved indication
- Have remaining pills, from a current prescription for an FDA-approved indication, for those changing doses
- If using the first time, must not have already received a dose from the Free Trial Program
- Not have already had 2 separate dose adjustments from the Dose Modification Program if changing doses
- Not be receiving ZEJULA from the Patient Assistance Program

The program is available to all patients who meet eligibility requirements, including those enrolled in Medicare or other government-funded programs.

**Please note:** To provide your patient a new dose of ZEJULA at no charge, this program is dispensed by PharmaCord Pharmacy rather than your in-office dispensary or the specialty pharmacy that is currently dispensing your patient's prescription. ZEJULA can be shipped to the patient as early as 24 hours after the receipt of this form, if the completed form is received before 3 PM ET.

For ongoing refills, a new prescription will need to be submitted to the patient's existing specialty or in-office dispensing pharmacy. We will contact your office after your patient's dose has been shipped so that you may adjust the next month's prescription as needed.

Future prescriptions are not required to participate in the ZEJULA Free Trial and Dose Modification Program.

SECTION 1. PAT	IENT INFORMATION				
Name:		DOB (mm/dd/yyyy):	Phone	:	
Address:					
		Best time to contact:			
Alternative Contact Name:		Ph	Phone:		
SECTION 2. PRE	SCRIBER INFORMATION				
Name:		Practice:			
Address:					
		Preferred method of contact			
Phone:	Fax:	Email:			
NPI#:	State License #:	Tax ID#:		DEA#:	

# **Free Trial and Dose Modification Program**



Patient Name:	Patient DOB:MM /DD /YYYY
SECTION 3. PRESCRIPTION INFORMATION	
Provider Name: Provider	der NPI: Provider Phone:
<ul> <li>○ FREE TRIAL PROGRAM</li> <li>New Prescription (Select only 1)</li> <li>○ ZEJULA 200-mg tablets PO QD #30</li> <li>○ ZEJULA 300-mg tablets PO QD #30</li> </ul>	<ul> <li>DOSE MODIFICATION PROGRAM</li> <li>Current Prescription Dose (Select only 1)</li> <li>ZEJULA 100-mg tablets PO QD #30</li> <li>ZEJULA 200-mg tablets PO QD #30</li> <li>ZEJULA 300-mg tablets PO QD #30</li> </ul>
Diagnosis Code:	New Prescription Dose (Select only 1)  ZEJULA 100-mg tablets PO QD #30  ZEJULA 200-mg tablets PO QD #30  ZEJULA 300-mg tablets PO QD #30
"Dispense As Written" / Brand Medically Necessary / Do Not Substitute / No Substitution / DAW / May Not Substitute  Prescriber's Signature:	Date:MM /DD /YYYY
Known Drug Allergies:	ed to be submitted to the patient's existing specialty or
SECTION 4. TERMS AND CONDITIONS	
<ul> <li>The ZEJULA Free Trial and Dose Modification Program is available a indication.</li> <li>Patients using ZEJULA for the first time must have not already have.</li> <li>Patients who are changing doses must not have already had two (2) see.</li> <li>No one participating in the program may bill or seek payment or reparty payer, including any state or federal entity or any private or continuous.</li> <li>Those prohibited from billing or seeking payment or reimbursement institutions, pharmacies, pharmacists, or any other person or prescont Medicare Part D patients who participate in the Program must agree shall be counted towards their out-of-pocket costs, and no claim will</li> <li>Product provided pursuant to this Free Trial Program and Dose Mood</li> <li>GSK reserves the right to change or end the program at any time were continuous.</li> </ul>	re received a dose from the Free Trial Program. parate dose adjustments under the Dose Modification Program. imbursement for tablets received from the program from any third- other insurance plan. It for tablets received from the program include patients, prescribers, criber. It that no part of the costs of the drug provided as part of the Program be filed with a Part D plan for drug supplied by the Program. dification Progam may not be sold, traded, or distributed for sale.
complies with my state-specific prescribing requirements; 4)	cription medication identified in this form; (3) the prescription In my medical judgment, the starting or the new strength of nd its use is consistent with the FDA-approved indication; and 5) /e.
NO STANIF	S PLEASE Date: MM / DD / YYYY

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