Lecanemab-irmb (Legembi)

Provider Order Form rev. 10/04/2024

PATIENT INFORMATION		Referral Status:	□ New Referral		Updated Orde	er 🛛 Order Renewal
Patient Name:			DOB:		Patient Pho	ne:
Patient Address:		Patient Email:				
Allergies:			🗆 NKDA	Weight	t (lbs/kg):	Height (in/cm):
Sex: 🗆 M / 🗆 F	Date of Last Infusion:	Next Due Date	:	Pref	erred Location:	
DIAGNOSIS (PI	lease provide ICD-10 code i	n space provided)				
Alzheimer's Disea	ase:					
Other:		Description:				
REQUIRED INF	FORMATION FOR MED	ICARE F	RE-MED	CATIO	N ORDERS	
□ Z00.6: Encounter for examination for normal comparison and			🗆 Tylenol 🗆 500mg / 🗆 650mg PO			
control in clinical research program			Loratadine 10mg PO			
Medicare Trial Registry Number:			🗆 Pepcid 20mg 🗆 PO / 🗆 IVP			

THERAPY ADMINISTRATION & DOSING

🗹 Administer Leqembi 10n	ng/kg x	kg =	mg
IV every 2 weeks. Infuse in	250ml 0.9% NS o	ver 1 hour	
☑ Flush the IV line with no	rmal saline to ma	ke sure all	
medication is infused.			
☑ Dosing Weight:	kg		

ADDITIONAL ORDERS

- □ Benadryl □ 25mg / □ 50mg □ PO / □ IVP
- □ Solumedrol □ 40mg / □ 125mg IVP
- □ Other: ___

NURSING

☑ Hold infusion and notify provider for:

- Hold if amyloid beta pathology has not been confirmed. ٠
- Abnormal vital signs
- No brain MRI results in chart (need MRI within one year of starting treatment, and prior to 5th, 7th, and 14th infusion).
- Signs of Amyloid Related Imaging Abnormalities (ARIA) as reported on MRI results.
- New or worsening headache or altered mental status.

☑ Record vital signs before infusion, then every 30 minutes until patient discharge

☑ Provide nursing care per Nursing Procedure, including Hypersensitivity Reaction Management Protocol and postprocedure observation

☑ To report suspected adverse reactions, contact FDA at 1-800-FDA-1088 or www.fda.gov/medwatch

PROVIDER INFORMATION

Preferred Contact Name:	Prefe	Preferred Contact Email:			
Ordering Provider:	ing Provider: Provider NPI:				
Referring Practice Name:	Phone:	Fax:			
Practice Address:	City:	State:	Zip Code:		

REQUIRED DOCUMENTATION CHECKLIST (Additional documentation required for processing and insurance approval)

Required Documentation: Patient demos, copy of front and back of primary and secondary insurance, 2 most recent OVN including treatment failures or contraindications. Documentation confirming patient's enrollment in CMS National Patient Registry, MRI at initial and throughout treatment, PET or CSF analysis for amyloid bodies, cognitive function score

Provider Name (print)

Provider Signature

Date

Order valid for one year unless otherwise indicated. IV solutions/diluents may be substituted as allowed per manufacturer's instructions as necessitated by product availability.