Donanemab-azbt (Kisunla)

Provider Order Form rev. 10/04/2024

PATIENT INFO	RMATION	Referral Status:	🗆 New R	eferral	Updated Order	r 🛛 Order Renewal		
Patient Name:			DOB:		Patient Phone	e:		
Patient Address:			Patient Email:					
Allergies:			🗆 NKDA	Weight	t (lbs/kg):	Height (in/cm):		
Sex: 🗆 M / 🗆 F	Date of Last Infusion:	Next Due Date:	:	Prefe	erred Location:			
DIAGNOSIS (Please provide ICD-10 code in space provided)								
Alzheimer's Disea	ase:							
Other:		Description:						

REQUIRED INFORMATION FOR MEDICARE

□ Z00.6: Encounter for examination for normal comparison and control in clinical research program Medicare Trial Registry Number: _____

THERAPY ADMINISTRATION & DOSING

□ Administer Kisunla 700mg IV over 30 minutes every 4 weeks X 3 doses, then administer Kisunla 1400mg IV over 30 minutes every 4 weeks starting with the 4th dose.

Administer Kisunla 1400mg IV over 30 minutes every 4 weeks.

 \square Flush the IV line with normal saline to make sure all medication is infused.

☑ Monitor patient for at least 30mins after each infusion

ADDITIONAL ORDERS

PRE-MEDICATION ORDERS

- □ Tylenol □ 500mg / □ 650mg PO
- Loratadine 10mg PO
- 🗆 Pepcid 20mg 🗆 PO / 🗆 IVP
- □ Benadryl □ 25mg / □ 50mg □ PO / □ IVP
- □ Solumedrol □ 40mg / □ 125mg IVP
- □ Other:

NURSING

☑ Hold infusion and notify provider for:

- MRI not performed or read by radiologist. MRI must be done as a baseline before starting treatment and prior to 2nd, 3rd, 4th and 7th infusion.
- Signs of Amyloid Related Imaging Abnormalities (ARIA) as reported on MRI results.
- New neurological symptoms including headaches or altered mental status.

☑ 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:	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.