Vermont 28 Park Ave Suite #1A Williston, VT 05495





UDDED EUDM	Date:	

KISUNLA (donanemab-azbt)	O]	RDER FORM	Date:					
PATIENT INFORMATION								
Name:	DOB:		SEX: M □ F □					
Allergies:		Date of Referral:						
PHYSICIAN INFORMATION								
Physician Name*:		Practice Name:						
Address:		Office Contact*:						
Phone: Fax:		Email (for updates):						
REFERRAL STATUS								
□New Referral □Referral Renewal □Medication/Order Change □Benefits Verification Only □Discontinuation Order								
Kisunla: Kisunla is indicated for the treatment of Alzheimer's disease (AD). Treatment with Kisunla should be initiated in patients with mild cognitive impairment (MCI) or mild dementia stage of disease, the population in which treatment was initiated in the clinical trials.								
DOSAGE AND ADMINISTRATION:		Dosing Recomr	mendations for Patients With ARIA-E					
 Confirm the presence of amyloid beta pathology prior to initiating treatment. 	C	linical Symptom Severitya	ARIA	-E Severity on N	1RI			
Intravenous Infusion KISUNLA Dosage (every 4 weeks) (administered over approximately 30			Mild	Moderate	Severe			
(every 4 weeks) (administered over approximately 30 minutes)	ш—	Asymptomatic	May continue dosing at current dose and schedule	Suspend dosing ^b	Suspend dosing ^b			
Infusions 1, 2, and 3 700 mg	I I ⊨	Mild Moderate or Severe	May continue dosing based on clinical judgment	Suspend dosing ^b				
Infusion 4 and beyond 1400 mg	Moderate or Severe Suspend dosing ^b							
• The recommended dosage of KISUNLA is 700 mg administered as	Dosing Recommendations for Patients With ARIA-H							
an intravenous infusion over approximately 30 minutes every four weeks for the first three doses, followed by 1400 mg every four		linical Symptom Severitya		-E Severity on N				
weeks.		Asymptomatic	May continue dosing at current dose and schedule	Moderate Suspend dosinga	Severe			
 Consider stopping dosing with KISUNLA based on reduction of amyloid plaques to minimal levels on amyloid PET imaging. 	ΙΙ⊢		current dose and schedule Suspend dosing ^a	Suspend dosing ^a	Suspend dosing ^b			
Obtain a recent baseline brain MRI prior to initiating treatment. Obtain an MRI prior to the 2nd, 3rd, 4th, and 7th infusions. If radiographically observed ARIA occurs, treatment recommendations are based on type, severity, and presence of symptoms. Dilution to a final concentration of 4 mg/mL to 10 mg/mL with 0.9% Sodium Chloride Injection, is required prior to administration. DOSAGE FORMS AND STRENGTHS: Injection: 350 mg/20 mL (17.5 mg/mL) in a single-dose vial Diagnosis Code: Order/dosage: Signature: Signature:	WARNING: AMYLOID-RELATED IMAGING ABNORMALITIES Monoclonal antibodies directed against aggregated forms of beta amyloid, including Kisunla, can cause amyloid-related imaging abnormalities (ARIA), characterized as ARIA with edema (ARIA-E) and ARIA with hemosiderin deposition (ARIA-H). ARIA can be serious and life-threatening events can occur. Serious intracerebral hemorrhages >1 cm, some fatal, have been observed with this class of medications. Because ARIA-E can cause focal neurologic deficits that can mimic an ischemic stroke, treating clinicians should consider whether such symptoms could be due to ARIA-E before giving thrombolytic therapy. Apolipoprotein Ε ε4 (ApoE ε4) Homozygotes: Patients treated with this class of medications, including Kisunla, who are ApoE ε4 homozygotes have a higher incidence of ARIA, including symptomatic and serious ARIA, compared to heterozygotes and noncarriers. Testing for ApoE ε4 status should be performed prior to initiation of treatment to inform the risk of developing ARIA. Consider the benefit for the treatment of Alzheimer's disease and risk of ARIA when deciding to treat with Kisunla.							
ORDERING PROVIDER	-							
Signature X				Date				
Provider	_ Ph	one		_ Fax				