

Checklist for Kisunla (donanemab) Referral

Patien	nt Name:	DOB:	Date:	_
Referr	ring Physician:		NPI:	
Office	e Contact/Title/Email:			-
Office	e Address:			_
Office	e Phone:	Office Fax:		
Best co	contact number for physician in case of reac	Office Fax:		
Please	e return completed checklist and checklist ite	ems to initiate referral.	Use this form as fax cover sheet.	
	Patient demographic information			
	Insurance information and copy of insurar	nce card/s (front and b	ack). *Include primary and secondary insurance	ì
	Supporting clinical notes and office visits.	Two notes preferred.		
	 Note should include any therapies 	s tried/failed, and must	include discussion about Kisunla	
	 Medication list and allergies 			
	 Cognitive assessment and function Supporting lab reports/imaging for Kisunla 		ore and interpretation	
	 MRI within 1 year of treatment st 			
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	 ApoE testing to determine ARIA ri 			
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	Eav all information to our	r Infusion Coo	rdinator: 508-608-8671	_

Fax all information to our Infusion Coordinator: 508-698-8671

Call with any questions: 781-551-5812 option 4



Prescribing Order: Kisunla (donanemab)

Date of Order:	☐ New Start	☐ New Start ☐ Maintenance Date of last infusion:					
Patient Name:	DOB:	M/F:					
Diagnosis (include ICD-10 code/s):							
☐ NKDA Allergies:							
Patient Weight:							
Premedication:							
☐ Acetaminophen 1000mg PO	Dipher	enhydramine 25mg IV					
☐ Loratadine 10mg or Cetirizine 10mg PO	☐ Solu-m	medrol 125mg IV in 50ml over 15min					
☐ Diphenhydramine 25mg PO	☐ Other:	T					
ab Orders:							
☐ New start: Kisunla 700mg every 4 we	LA Medication Order: New start: Kisunla 700mg every 4 weeks for 3 infusions, then Kisunla 1400mg every 4 weeks over 30min Maintenance: Kisunla 1400mg every 4 weeks over 30min						
Other:	ther:						
Administration:							
✓ Hold infusion if no MRI Brain price	or to the 2^{nd} , 3^{rd} , 4^{th} and 7^{th} in	nfusion					
 Hold infusion and notify provider 	if patient experiencing any of	f the following signs of ARIA:					
	zziness, Nausea, Vision Change						
✓ In case of infusion reaction, STOF	infusion and follow ICNE infu	usion reaction protocol. Notify physician.					
Ordering Provider Name	NP	PI					
Signature	Da	ate					



Patient Name:	D	OB:	

Legembi/Kisunla Indication Checklist

	INFORMATION REQUIRED FO	OR TREATMENT INITIATION	
1)	Patient ICD-10 (select all that apply)	Clinical Diagnosis (select one)	
	G30.0 Alzheimer's disease, early onset	☐ Mild cognitive impairment due to AD	
	☐ G30.1 Alzheimer's disease, late onset	☐ Mild AD Dementia	
	G30.9 Alzheimer's disease, unspecified		
	G31.84 Mild cognitive impairment		
2)	Cognitive Screening: within 6 months Date:		
	Name of Test Used:	Score:	
	Interpretation:		
	Functional Screening: within 6 months Date:		
	Name of Test Used:	Score:	
3)	Confirmation of Amyloid-Beta Pathology: MUS	T provide supporting documentation	
	☐ Amyloid PET Scan Date:	Result:	
	☐ CSF Amyloid Confirmation Date:	Result:	
4)	Monitoring for Amyloid Related Imaging Abnormalities (ARIA) ***LEQEMBI/KISUNLA requires brain MRI within 1 year of treatment start date***		
	☐ Initial MRI Brain Date: Evidence of ARIA-E ☐ Negative	Positive	
	Evidence of ARIA-H Negative		
	Evidence of AlliA 11 — Negative	— 1 0510.00	
5)	Schedule for MRI Monitoring: the following is re	equired by Infusion Center of New England	
	☐ Leqembi: Obtain MRI prior to the 5 th , 7 th	, and 14 th infusions	
	☐ Kisunla: Obtain MRI prior to the 2 nd , 3 rd ,	4 th , and 7 th infusions	
6)	Is the patient on anticoagulation? Yes	□ No	
7)	Is the patient on antiplatelets?	□ No	
8)	Has ApoE testing been performed? ☐ Yes	□ No	
		Result:	
9)	CMS REGISTRATION NUMBER		
	(REQUIRED)	Date:	