



PATIENT INFORMATION (Complete or Fax Existing Chart)	PRESCRIBER INFORMATION
Name: _____ DOB: _____	Prescriber Name: _____
Address: _____	State License: _____
City, State, Zip: _____	NPI #: _____ Tax ID: _____
Phone: _____ Alt. Phone: _____	Address: _____
Email: _____ SS#: _____	City, State, Zip: _____
Gender: <input type="checkbox"/> M <input type="checkbox"/> F Weight: _____ (lbs) Ht: _____	Phone: _____ Fax: _____
Allergies: _____	Office Contact: _____ Phone: _____
Is patient the main point of contact? <input type="checkbox"/> Yes <input type="checkbox"/> No If not, who?	
Name/Relationship: _____ Phone	
Number: _____	

INSURANCE INFORMATION – AND – Send a copy of the patient's prescription/insurance cards (front & back)	
Primary Insurance: _____	Secondary Insurance (If Applicable): _____
Plan #: _____	Plan #: _____
Group #: _____	Group #: _____
RX Card (PBM): _____	RX Card (PBM): _____
BIN: _____ PCN: _____	BIN: _____ PCN: _____
MEDICARE/MEDICARE ADVANTAGE PATIENTS ONLY:	
CMS Registry # (ALZH#): _____	CMS Registry Enrollment Date: _____

CLINICAL INFORMATION
Please Select Diagnosis:
<input type="checkbox"/> G30.0 Alzheimer's disease with early onset <input type="checkbox"/> G30.1 Alzheimer's disease with late onset <input type="checkbox"/> G30.8 Other Alzheimer's disease <input type="checkbox"/> G30.9 Alzheimer's disease, unspecified <input type="checkbox"/> G31.84 Mild cognitive impairment, so stated <input type="checkbox"/> Other: _____
Prescriber must indicate the following requirements have been met to confirm diagnosis and that Patient has evidence of AD neuropathology and has been assessed for baseline ARIA risk via MRI:
<input type="checkbox"/> Amyloid pathology confirmed via: <input type="checkbox"/> Amyloid PET Scan <input type="checkbox"/> CSF analysis <input type="checkbox"/> Blood plasma Date: _____ Resu It: <input type="checkbox"/> Amyloid Positive <input type="checkbox"/> Amyloid Negative <input type="checkbox"/> Recent MRI obtained prior to initiating Leqembi® (including FLAIR and T2/GRE or SWI) to assess ARIA risk <input type="checkbox"/> Prescriber has verified that this Patient does not have evidence of prior ARIA-H Date: _____ <input type="checkbox"/> Completion of cognitive assessment type: <input type="checkbox"/> MMSE <input type="checkbox"/> MoCA <input type="checkbox"/> CDR <input type="checkbox"/> Other: _____ Date: _____ <input type="checkbox"/> Completion of functional assessment type: <input type="checkbox"/> FAQ <input type="checkbox"/> FAST <input type="checkbox"/> Other: _____ Date: _____
**Note: MRIs must be obtained prior to initial infusion to assess ARIA risk. During treatment, conduct an ARIA monitoring MRI before Infusions 3, 5, 7, and 14 and if symptoms consistent with ARIA occur.

LEQEMBI® ORDERS		
Prescription type: <input type="checkbox"/> New start <input type="checkbox"/> Restart <input type="checkbox"/> Continued therapy Total Doses Received: _____ Date of Last Injection/Infusion: _____		
Medication	Dose/Frequency	Refills
<input type="checkbox"/> Leqembi® (lecanemab-irmb) 500 mg/5 mL (100 mg/mL) <input type="checkbox"/> Leqembi® (lecanemab-irmb) 200 mg/2 mL (100 mg/mL)	<input type="checkbox"/> 10 mg/kg intravenous infusion over approximately one hour, once every two weeks. <input type="checkbox"/> Other: _____	_____
Pre-Medication	Dose/Strength	Directions
<input type="checkbox"/> Acetaminophen	<input type="checkbox"/> 500mg	<input type="checkbox"/> Take 1-2 tablets PO prior to infusion or post-infusion as directed

CONFIDENTIALITY STATEMENT: This facsimile and documents accompanying this transmission contain confidential health information that is legally privileged. This information is intended only for the use of the individual or entity named above. The authorized recipient of this information is prohibited from disclosing this information to any other party unless required to do so by law or regulation. If you are not the intended recipient, you are hereby notified that any disclosure, copying, distribution, or action taken in reliance on the contents of these documents is strictly prohibited. If you have received this information in error, please notify the sender at the address and telephone number set forth herein and arrange for return or destruction of the material. In no event should such material be read by anyone other than the named addressee, except by express authority of the sender to the named addressee.



Please Fax Completed Form To: 888-340-3699
Please Send a Copy of The Patient's Insurance Cards (Front & Back)

Please Send a Copy of The Patient's Up-to-Date Clinical Notes

Table with 3 columns for medication options: Diphenhydramine, Methylprednisolone, and an empty row. Includes dosage and administration instructions.

INFUSION REACTION ORDERS

Mild reaction protocol:
Moderate reaction protocol:
Severe reaction protocol: (Call 911 if initiated):

FLUSHING & LOCKING ORDERS

Flushing Protocol (>66lbs/33kg)
PIV and Midline:
Implanted Port, PICC, Tunneled Catheter, and Non-tunneled Catheter:

Locking Protocol (>66lbs/33kg)
PIV and Midline:
PICC:
Implanted Port, Tunneled Catheter, and Non-tunneled Catheter:

** May substitute Dextrose 5% in Water, or alternative, for 0.9% Sodium Chloride, when indicated due to incompatibility with medications bring infused

SIGNATURE

We hereby authorize Infuse IQ LLC to provide all supplies and additional services (nursing/patient training) required to provide and deliver the medicine as prescribed in this referral.
X _____ Date: _____
Prescriber Signature

To ensure payment by insurance carrier, please include supporting clinical documentation for specified ICD 10 Code, demographic, and insurance information along with faxed order. Initial appointment will be verified upon insurance approval.

CONFIDENTIALITY STATEMENT: This facsimile and documents accompanying this transmission contain confidential health information that is legally privileged. This information is intended only for the use of the individual or entity named above.