

MEDICARE FORM

Leqvio® (inclisiran) Medication **Precertification Request**

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(All fields must be completed and legible for precertification review.)

For Michigan MMP: FAX: 1-844-241-2495 **PHONE**: 1-855-676-5772 For other lines of business:

Please use other form.

Note: For MAPD plans, Leqvio is nonpreferred. Praluent is preferred through the Part D benefit. Repatha is also preferred for MAPD plans with open formularies. Leqvio is not subject to step therapy on MA only plans.

Please indicate: Start of treatment: start date/				☐ Continuation of therapy, date of last treatment//					
Precertification Requ	ested By:			Phone	e:	Fax: _			
A. PATIENT INFORMA									
First Name:			Last Name:			DOB:			
Address:				City:		State:	ZIP:		
Home Phone:		Work Phone:		Cell Phone:		Email:	_		
Patient Current Weight:	lbs or	kgs Patien	t Height: inche	s orcms	Allergies:				
B. INSURANCE INFO									
Aetna Member ID #: _			Does patient have o	ther coverage?	☐ Yes ☐ No				
Group #:			If yes, provide ID#:Carrier Name:						
Insured:			Insured:						
Medicare: ☐ Yes ☐	No If yes, provi	de ID #:	N	Medicaid: 🗌 Yes	☐ No If yes, prov	vide ID #:			
C. PRESCRIBER INFO	ORMATION								
First Name:			Last Name:		(Check On	ne): 🔲 M.D. 🗀] D.O. 🗌 N.P. 🗌 P.A.		
Address:				City:		State:	ZIP:		
Phone:	Fax:		St Lic #:	NPI #:	DEA #:		UPIN:		
Provider Email:	•		Office Contact Name	e:	<u> </u>	Phone:			
Specialty (Check one):	☐ Cardiologis	st	1			-1			
D. DISPENSING PRO									
Place of Administration				Dispensing	Provider/Pharmac	v: Patient Se	lected choice		
☐ Self-administered	☐ Physi	cian's Office			Dispensing Provider/Pharmacy: Patient Selected choice ☐ Physician's Office ☐ Retail Pharmacy				
Outpatient Infusion Center Phone:			· · · · · · · · · · · · · · · · · · ·						
Center Name:		-		- ' '	, ,				
☐ Home Infusion Cen	ter Pl	none:							
Agency Name									
☐ Administration code									
Address:				_		PIN: _			
NPI:				NPI:					
E. PRODUCT INFORM			_						
Request is for: Leqvio						HCPCS C	ode:		
F. DIAGNOSIS INFOR	MATION - Pleas	se indicate primai	ry ICD code and spec	ify any other where	e applicable.				
Primary ICD Code:			=						
G. CLINICAL INFORM	IATION - Requir	ed clinical informa	ation must be complet	ted in its <u>entirety</u> fo	or all precertification	requests.			
Please indicate the curre	ent LDL-C level in	mg/dL:							
For Initiation Requests									
Note: Lequio is non-preferred on MAPD plans. Praluent is preferred through the Part D benefit. Repatha is also preferred for MAPD plans with									
open formularies. Leqvio is not subject to step therapy on MA only plans. Yes No Has the patient had prior therapy with Leqvio (inclisiran) within the last 365 days?									
Yes No Has the patient had a trial and failure, intolerance, or contraindication to any of the following? (select all that apply)									
☐ Praluent (alirocumab) ☐ Repatha (evolocumab)									
Please explain if there are any other medical reason(s) that the patient cannot use any of the following preferred products when indicated for the patient's									
diagnosis (select all that apply) □ Praluent (alirocumab) □ Repatha (evolocumab)									
∐ Pra	luent (alirocumab)	olocumab)						
☐ Ves ☐ No. Will the	natient continue t	o receive concom	itant statin therany?						
☐ Yes ☐ No Will the patient continue to receive concomitant statin therapy? ☐ Yes ☐ No Does the patient have intolerance or contraindication to high-intensity statin therapy?									
Please indicate the prior therapy the patient has previously received (select all that applies to the patient):									
☐ The patient is receivi	The patient is receiving a high-intensity statin dose daily, such as rosuvastatin (Crestor) 20 mg daily or atorvastatin (Lipitor) 40 mg daily								
Please indicate			for at locat 2 m = = th = 0						
☐ Yes ☐ No Has the patient received this dose for at least 3 months? ☐ Yes ☐ No Was the patient unable to tolerate a high-intensity statin due to adverse effects?									
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Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (continued	Required clinical information must be or	completed in its entirety for all pre-	certification requests					
For Initiation Requests (clinical document	•	ompleted in its <u>entirety</u> for all proc	portinioation requests.					
The patient is receiving a moderate-intensity statin dose daily, such as atorvastatin (Lipitor) 20 mg or equivalent Please indicate the start date: / /								
Yes No Has the patient received this dose for at least 3 months?								
The patient has intolerance to a high-intensity statin therapy								
Yes No Did the patient score a 7 or higher on the Statin-Associated Muscle Symptom Clinical Index (SAMS-CI)?								
☐ Yes ☐ No Did the patient experience a statin-associated increase in creatine kinase (CK) level of greater than or equal to 10 times the upper limit of normal (ULN) during previous treatment with a statin?								
The patient has contraindication to a high-intensity statin therapy								
Please indicate which of the following applies to the patient:								
Active liver disease, including <u>unexplained</u> persistent elevations in hepatic transaminase levels (e.g., ALT greater than or equal to 3 times the upper limit of normal)								
☐ Currently pregnant ☐ Planning pregnancy ☐ Breastfeeding ☐ None of the above								
Clinical atherosclerotic cardiovascular disease (ASCVD)								
Please indicate which of the following manifestations of clinical atherosclerotic cardiovascular disease (ASCVD) the patient has experienced: Acute coronary syndrome								
☐ Coronary Artery Calcium (CAC) score of greater than or equal to 1000								
Coronary or other arterial revascularization procedure (e.g., percutaneous coronary intervention [PCI], coronary artery bypass graft [CABG] surgery)								
Myocardial infarction								
☐ Non-cardiac peripheral arterial disease (F	PAD) of presumed atherosclerotic origin (e.g.,	carotid artery stenosis, lower extre	mity PAD)					
□ Obstructive coronary artery disease (defined as fifty percent or greater stenosis on cardiac computed tomography angiogram or catheterization)								
☐ Stable or unstable angina								
☐ Stroke of presumed atherosclerotic origin								
☐ Transient ischemic attack (TIA)								
☐ Other								
Heterozygous familial hypercholesterolemia (HeFH)								
Yes No Does the patient possess an	·							
Please indicate the patient's untreated (before any lipid-lowering therapy) LDL-C level in mg/dL:								
Please select which of the following applies to the patient: ☐ Family history of myocardial infarction (MI) at less than 60 years of age in a first degree relative or less than 50 years of age in a second								
degree relative	, ,	,	,					
	☐ Family history of total cholesterol (TC) greater than 290 mg/dL in a first/second degree relative							
☐ Presence of tendon xanthoma(s) in the patient or first/second-degree relative								
☐ None of the above- the patient does not meet any of the criteria listed above								
For Continuation Requests (clinical docum								
Yes No Has the patient achieved or the requested medication th	erapy?	is now at goal, robust lowering of LI	DL-C) as the result of					
Please indicate which of the following applies to the patient:								
The patient is currently receiving concomitant statin therapy								
☐ The patient has intolerance to a high-intensity statin therapy								
Yes No Did the patient score a 7 or higher on the Statin-Associated Muscle Symptom Clinical Index (SAMS-CI)?								
Yes No Did the patient experience a statin-associated increase in creatine kinase (CK) level of greater than or equal to 10 times								
the upper limit of normal (ULN) during previous treatment with a statin?								
The patient has contraindication to a high-intensity statin therapy								
Please indicate which of the following applies to the patient:								
Active liver disease, including <u>unexplained</u> persistent elevations in hepatic transaminase levels (e.g., ALT greater than or equal to 3 times upper limit of normal)								
☐ Currently pregnant ☐ Planning	pregnancy	ie above						
H. ACKNOWLEDGEMENT								
Request Completed By (Signature Requ	uired):		Date: /					
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive								