

INDIANA HEALTH COVERAGE PROGRAMS (IHCP) PCSK9 INHIBITORS AND SELECT LIPOTROPICS PRIOR AUTHORIZATION (PA) REQUEST FORM



CareSource Pharmacy Prior Authorization Form P.O. Box 8738 Dayton, OH 45401-8738 Fax: (866) 930-0019

(Holps					
Today's Date			Non-Urgent □ Urgent □		
Note: This form must be completed by the prescribing provider. *All sections must be completed or the request will be returned**					
Patient's CareSource #		Date of Birth			
Patient's Name		Prescriber's Na	ame		
Prescriber's Indiana		Specialty	Specialty		
Prescriber's NPI #		Prescriber's Si	Prescriber's Signature		
Prescriber Fax		Prescriber Phone	Prescriber Phone		
Prescriber Address:		Date(s) of Serv	vice:		
		Start Date:			
Diagnosis:		Diagnosis Cod	de:		
Lattest that the informatio	on on this form is accurate:				
Takos and and anomina	71 011 allo 10 12 222.222				
Physician Signature:			Date:		
Requested Medica	ation Strength	Quantity	Dosage Regimen		
DA Boquiroments for F	Evkeeza (evinacumab-dgr	h).			
-	sis of homozygous familial hyp		a (HoFH)? □ Yes □ No		
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 Medication prescribed by, or in consultation with, a cardiologist or endocrinologist? ☐ Yes ☐ No Select one of the following: 					
☐ Member is 5 years of age or older and less than 7 years of age.					
☐ Member is 7 years of age or older and less than 10 years of age and one of the following:					

PA Requirements for Evkeeza (evinacumab-dgnb) continued:
i. Member has trial and failure history of at least 90 days of therapy with rosuvastatin 20 mg?
□ Yes □ No
ii. Provider has submitted documentation of intolerance/contraindication to rosuvastatin?
□ Yes □ No
\square Member is 10 years of age or older and less than 18 years of age and one of the following:
i. Member has trial and failure history with Repatha (evolocumab)? ☐ Yes ☐ No
Drug/dose/date(s):
ii. Member has trial and failure history of at least 90 days of high dose rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy concurrently with ezetimibe (or documented intolerance/contraindication to statins/ezetimibe) AND provider has submitted medical justification for use of Evkeeza (evinacumab-dgnb) over Repatha
(evolocumab)? ☐ Yes ☐ No
Drug/dose/date(s):
☐ Member is 18 years of age or older and one of the following:
i. Member has trial and failure history with Praluent (alirocumab) OR Repatha (evolocumab)?
□ Yes □ No
Drug/dose/date(s):
 ii. Member has trial and failure history of at least 90 days of high dose rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy concurrently with ezetimibe (or documented intolerance/contraindication to statins/ezetimibe) AND provider has submitted medical justification for use of Evkeeza (evinacumab-dgnb) over Praluent
(alirocumab) and Repatha (evolocumab)? ☐ Yes ☐ No
Drug/dose/date(s):
4. Select one of the following:
☐ Member will utilize maximally tolerated statin therapy with or without ezetimibe concurrently with Evkeeza (for those 7 years of age and older).
☐ Provider has submitted documented intolerance to statin and/or ezetimibe therapy or medical rationale against use of statin or ezetimibe therapy.
5. Requested dose is 15 mg/kg every 4 weeks or less? Yes No Member weight: LB / KG (circle one)
DA Paguiraments for Juytanid (Iomitanida magylata):
PA Requirements for Juxtapid (lomitapide mesylate):
1. Member is enrolled in the Juxtapid/lomitapide REMS program and prescriber is monitoring in accordance with
REMS requirements? ☐ Yes ☐ No
2. Member is 18 years of age or older? ☐ Yes ☐ No
3. Medication prescribed by, or in consultation with, a cardiologist or endocrinologist? ☐ Yes ☐ No
4. Select one of the following:
☐ Member has trial and failure history of Praluent (alirocumab) or Repatha (evolocumab).
Drug/dose/date(s):

PA Requirements for Juxtapid (lomitapide mesylate) continued:		
	□ Member has trial and failure history of at least 90 days of high dose rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy concurrently with ezetimibe (or documented intolerance/contraindication to statins/ezetimibe) AND provider has submitted medical justification for use of Juxtapid (lomitapide mesylate) over Praluent (alirocumab) and Repatha (evolocumab). Drug/dose/date(s):	
5.	For those of childbearing potential, documentation of a negative pregnancy test obtained in the past 30 days is attached and prescriber has counseled member on risks associated with conceiving while utilizing Juxtapid and appropriate methods of contraception? □ Yes □ No	
	Prescriber Name and Signature:	
6.	Select one of the following: Member will utilize maximally tolerated statin therapy with or without ezetimibe concurrently with Juxtapid. Provider has submitted documented intolerance to statin and/or ezetimibe therapy or medical rationale against use of statin or ezetimibe therapy.	
7. F	Requested dose is 60 mg/day or less? □ Yes □ No	
-D/		
	A Requirements for Leqvio (inclisiran):	
1.	Select one of the following: ☐ Member has a diagnosis of primary hyperlipidemia with clinical atherosclerotic cardiovascular disease (ASCVD) or is at increased risk for ASCVD with a baseline LDL-C level of ≥55 mg/dL (documentation required). ☐ Member has diagnosis of heterozygous familial hypercholesterolemia (HeFH) with a baseline LDL-C level of ≥70 mg/dL (documentation required).	
2.	Member is 18 years of age or older? ☐ Yes ☐ No	
3.	Prescribed by, or in consultation with, a cardiologist or endocrinologist? ☐ Yes ☐ No	
4.	Select one of the following:	
	☐ Member has trial and failure history of Praluent (alirocumab) or Repatha (evolocumab). Drug/dose/date(s):	
	☐ Member has trial and failure history of at least 90 days of high dose rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy concurrently with ezetimibe (or documented intolerance/contraindication to statins/ezetimibe) AND provider has submitted medical justification for use of Leqvio (inclisiran) over Praluent (alirocumab) and Repatha (evolocumab). Drug/dose/date(s):	
5.	Select one of the following:	
	Member will utilize maximally tolerated statin therapy with or without ezetimibe concurrently with Leqvio.	
	Provider has submitted documented intolerance to statin and/or ezetimibe therapy or medical rationale against use of statin or ezetimibe therapy.	
6.	Select one of the following:	
	☐ Member is initiating therapy and requested dose does not exceed 284 mg every three months.	
	$\hfill \square$ Member is established on therapy and requested dose does not exceed 284 mg every six months.	

PA	Requirements for Niacin ER
1.	Diagnosis of severe hypertriglyceridemia (baseline triglycerides ≥500 mg/dL)? ☐ Yes ☐ No
	If <u>Yes</u> , then select one of the following:
	 ☐ Member is on concurrent therapy with all of the following for at least 90 days: omega-3 fatty acid (omega-3-acid ethyl esters or icosapent ethyl), fibric acid derivative and statin therapy. ☐ Drug/dose/date(s):
	☐ Member has a documented intolerance of omega-3 fatty acid, fibric acid derivative AND statin therapy OR medical justification for use of Niacin ER over omega-3 fatty acid, fibric acid derivative AND statin therapy. Please explain:
2.	Member is 17 years of age or older? ☐ Yes ☐ No
PA	A Requirements for Praluent (alirocumab):
	Select one of the following:
	☐ Member has a diagnosis of clinical ASCVD, is at Very High Risk requiring therapy for secondary prevention, AND has persistently elevated LDL-C (≥55 mg/dL) despite treatment with 90 days of therapy with high intensity rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy or has documented intolerance of both rosuvastatin and atorvastatin OR medical rationale against the use of statin therapy.*
	☐ Member has a diagnosis of clinical ASCVD, is NOT at Very High Risk requiring therapy for secondary prevention, AND has persistently elevated LDL-C (≥70 mg/dL) despite treatment with 90 days of therapy with high intensity rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy WITH ezetimibe or has documented intolerance of rosuvastatin and atorvastatin and/or ezetimibe OR medical rationale against the use of statin therapy and/or ezetimibe therapy.
	☐ Member has a diagnosis of clinical ASCVD, with a baseline LDL-C ≥190 mg/dL, not due to secondary causes, without clinical or genetic diagnosis of familial hypercholesterolemia, requiring therapy for secondary prevention AND has persistently elevated LDL-C (≥70 mg/dL) despite treatment with 90 days of therapy with high intensity rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy or has documented intolerance of both rosuvastatin and atorvastatin OR medical rationale against the use of statin therapy.*
	☐ Member has a diagnosis of clinical ASCVD, is at Very High Risk with a baseline LDL-C ≥190 mg/dL not due to secondary causes, a diagnosis of familial hypercholesterolemia, requiring therapy for secondary prevention, AND has persistently elevated LDL-C (≥55 mg/dL) despite treatment with 90 days of therapy with high intensity rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy or has documented intolerance of both rosuvastatin and atorvastatin OR medical rationale against the use of statin therapy.*

Ρ,	A Requirements for Praluent (alirocumab) continued:
	☐ Member has a diagnosis of primary hyperlipidemia, without clinical ASCVD, with a baseline LDL-C ≥190 mg/dL not due to secondary causes, with or without concomitant ASCVD risk factors, requiring therapy for primary prevention AND persistently elevated LDL-C (≥100 mg/dL) despite treatment with 90 days of therapy with high intensity rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy or has documented intolerance of both rosuvastatin and atorvastatin OR medical rationale against the use of statin therapy.*
	☐ Member has a diagnosis of homozygous familial hypercholesterolemia (HoFH) or heterozygous familial hypercholesterolemia (HeFH) AND persistently elevated LDL-C (≥70 mg/dL) despite treatment with 90 days of therapy with high intensity rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy WITH ezetimibe or has documented intolerance of rosuvastatin and atorvastatin and/or ezetimibe OR medical rationale against the use of statin therapy and/or ezetimibe therapy.
	* For members requiring >25% additional lowering of LDL-C ONLY (≤ 25% LDL-C lowering must utilize high intensity statin therapy WITH ezetimibe as first line).
	For above diagnoses that require medical justification for use of Praluent over statin and/or ezetimibe therapy, please provide justification here:
2.	Member is 18 years of age or older? \Box Yes \Box No
3.	Select one of the following:
	Member will utilize maximally tolerated statin therapy with or without ezetimibe concurrently with Praluent.
	 Provider has submitted documented intolerance to statin and/or ezetimibe therapy or medical rationale against use of statin or ezetimibe therapy.
4.	Select one of the following:
	☐ Requested dose is 75 mg every two weeks.
	☐ Requested dose is 300 mg every four weeks.
	 □ Requested dose is 150 mg every two weeks AND the member has one of the following: □ Diagnosis of homozygous familial hypercholesterolemia.
	Diagnosis of heterozygous familial hypercholesterolemia and member is undergoing LDL apheresis.
	Member has not achieved clinically meaningful response after at least 4 weeks of dosing at 75 mg every 2 weeks or 300 mg every 4 weeks.

PF	Requirements for Repatha (evolocumab):
1.	Select one of the following:
	☐ Member has a diagnosis of clinical ASCVD, is at Very High Risk requiring therapy for secondary prevention, AND has persistently elevated LDL-C (≥55 mg/dL) despite treatment with 90 days of therapy with high intensity rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy or has documented intolerance of both rosuvastatin and atorvastatin OR medical rationale against the use of statin therapy.*
	☐ Member has a diagnosis of clinical ASCVD, is NOT at Very High Risk requiring therapy for secondary prevention, AND has persistently elevated LDL-C (≥70 mg/dL) despite treatment with 90 days of therapy with high intensity rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy WITH ezetimibe or has documented intolerance of rosuvastatin and atorvastatin and/or ezetimibe OR medical rationale against the use of statin therapy and/or ezetimibe.
	☐ Member has a diagnosis of clinical ASCVD, with a baseline LDL-C ≥190 mg/dL, not due to secondary causes, without clinical or genetic diagnosis of familial hypercholesterolemia, requiring therapy for secondary prevention AND has persistently elevated LDL-C (≥70 mg/dL) despite treatment with 90 days of therapy with high intensity rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy or has documented intolerance of both rosuvastatin and atorvastatin OR medical rationale against the use of statin therapy.*
	☐ Member has a diagnosis of clinical ASCVD, is at Very High Risk with a baseline LDL-C ≥190 mg/dL not due to secondary causes, a diagnosis of familial hypercholesterolemia, requiring therapy for secondary prevention, AND has persistently elevated LDL-C (≥55 mg/dL) despite treatment with 90 days of therapy with high intensity rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy or has documented intolerance of both rosuvastatin and atorvastatin OR medical rationale against the use of statin therapy.*
	☐ Member has a diagnosis of primary hyperlipidemia, without clinical ASCVD, with a baseline LDL-C ≥190 mg/dL not due to secondary causes, with or without concomitant ASCVD risk factors, requiring therapy for primary prevention AND persistently elevated LDL-C (≥100 mg/dL) despite treatment with 90 days of therapy with high intensity rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy or has documented intolerance of both rosuvastatin and atorvastatin OR medical rationale against the use of statin therapy.*
	☐ Member has a diagnosis of homozygous familial hypercholesterolemia (HoFH) or heterozygous familial hypercholesterolemia (HeFH) AND persistently elevated LDL-C (≥70 mg/dL) despite treatment with 90 days of therapy with high intensity rosuvastatin (20 mg/40 mg) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy WITH ezetimibe or has documented intolerance of rosuvastatin and atorvastatin and/or ezetimibe OR medical rationale against the use of statin therapy and/or ezetimibe.
2.	For members requiring >25% additional lowering of LDL-C ONLY (≤ 25% LDL-C lowering must utilize high intensity statin therapy WITH ezetimibe as first line).
	Note: Documentation of any and all intolerances to statins and/or ezetimibe must be provided.
	For any of the above diagnoses that have medical rationale against the use of statin and/or ezetimibe therapy please provide here:

PA	Requirements for Repatha (evolocumab) continued:
2.	Select one of the following:
	☐ Member is 18 years of age or older.
	\square Member is 10 years of age or older and has a diagnosis of either HoFH or HeFH.
3.	Select one of the following: Member will utilize maximally tolerated statin therapy with or without ezetimibe concurrently with Repatha. Provider has submitted documented intolerance to statin and/or ezetimibe therapy or medical rationale against use of statin or ezetimibe therapy.
4.	Select one of the following:
	☐ Requested dose is 140 mg every two weeks.
	☐ Requested dose is 420 mg once monthly.
	 Requested dose is 420 mg every two weeks AND the member has one of the following: Diagnosis of HoFH and has not achieved clinically meaningful response after at least 12 weeks at 420 mg once monthly dosing.
	☐ Member is receiving lipid apheresis.

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