## MHS PHARMACY BENEFIT PCSK9 INHIBITORS AND SELECT LIPOTROPICS PRIOR AUTHORIZATION REQUEST FORM

## MHS 550 N. Meridian St. Suite 101 Indianapolis, IN, 46204-1208 Phone: (877) 647-4848 Fax: (866) 399-0929



Today's Date			
Note: This form must be completed by th  **All sections		ler. d or the request wil	ll be returned**
Patient's Medicaid #		Date of Birth	
Patient's Name		Prescriber's Name	
Prescriber's IN License #		Specialty	
Prescriber's NPI #		Prescriber's Signature	e
Return Fax #		Return Phone #	
Check box if requesting retro-active PA		Date(s) of service requested for retro-active eligibility (if applicable):	
		or to eligibility determi	nation, but within established eligibility timelines) quests (dates of service 30 calendar days or less
Requested Medication	Strength	Quantity	Dosage Regimen
PA Requirements for Evkeeza (ev	vinacumab-dgn	b):	
			(HoFH) □ Yes □ No
Member has a diagnosis of homozy	ygous familial hyp	ercholesterolemia (	· ,
<ol> <li>Member has a diagnosis of homozy</li> <li>Medication prescribed by, or in con</li> <li>Select one of the following:</li> </ol>	ygous familial hypessultation with, a ca	ercholesterolemia	crinologist □ Yes □ No
<ol> <li>Member has a diagnosis of homozy</li> <li>Medication prescribed by, or in con</li> <li>Select one of the following:</li> <li>Member is 5 years of a</li> </ol>	ygous familial hypesultation with, a ca	ercholesterolemia of ardiologist or endo	crinologist □ Yes □ No
<ol> <li>Member has a diagnosis of homozy</li> <li>Medication prescribed by, or in con</li> <li>Select one of the following:         <ul> <li>Member is 5 years of a</li> <li>Member is 7 years of a</li> </ul> </li> </ol>	ygous familial hypesultation with, a can age or older and leage or older and leage or older and leage	ercholesterolemia of ardiologist or endocess than 7 years of ess than 10 years of	crinologist □ Yes □ No
<ol> <li>Member has a diagnosis of homozy</li> <li>Medication prescribed by, or in con</li> <li>Select one of the following:         <ul> <li>Member is 5 years of a</li> <li>Member is 7 years of a</li> </ul> </li> </ol>	ygous familial hypesultation with, a can age or older and leage or older and leage or older and leage	ercholesterolemia of ardiologist or endocess than 7 years of ess than 10 years of	crinologist □ Yes □ No age of age and one of the following:
1. Member has a diagnosis of homozy  2. Medication prescribed by, or in con  3. Select one of the following:  ☐ Member is 5 years of a  ☐ Member is 7 years of a  i. Member has to	ygous familial hypessultation with, a canage or older and leage or older and leage and failure hist	ercholesterolemia of ardiologist or endocess than 7 years of ess than 10 years of ory of at least 90 d	crinologist □ Yes □ No age of age and one of the following:
1. Member has a diagnosis of homozy  2. Medication prescribed by, or in con  3. Select one of the following:  ☐ Member is 5 years of a  ☐ Member is 7 years of a  i. Member has to	ygous familial hypessultation with, a canage or older and leage or older and leage and failure hist	ercholesterolemia of ardiologist or endocess than 7 years of ess than 10 years of ory of at least 90 d	crinologist
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1. Member has a diagnosis of homozy  2. Medication prescribed by, or in conditions  3. Select one of the following:	ygous familial hyposultation with, a canage or older and leage or older and leage and failure hist submitted document	ercholesterolemia of ardiologist or endocess than 7 years of ess than 10 years of ory of at least 90 dentation of intolerancess than 18 years	crinologist
1. Member has a diagnosis of homozy  2. Medication prescribed by, or in conditions  3. Select one of the following:	ygous familial hypesultation with, a cauge or older and leage or older and leage or older and leage and failure histerial	ercholesterolemia of ardiologist or endocess than 7 years of ess than 10 years of ory of at least 90 dentation of intolerancess than 18 years	age of age and one of the following: lays of therapy with rosuvastatin 20 mg ce/contraindication to rosuvastatin of age and one of the following:
2. Medication prescribed by, or in con 3. Select one of the following:  Member is 5 years of a  Member is 7 years of a  i. Member has tr  Yes No  ii. Provider has s  Yes No  Member is 10 years of  i. Member has tr  Drug/dose/dat  ii. Member has tr  mg) or atorvas ezetimibe (or o	ygous familial hypesultation with, a cauge or older and leage or older and leage or older and failure histerial and failure histeria	ercholesterolemia of ardiologist or endocases than 7 years of ess than 10 years of ory of at least 90 dentation of intolerance ory with Repatha (cory of at least 90 dentation of at least 90 dentat	age of age and one of the following: lays of therapy with rosuvastatin 20 mg ce/contraindication to rosuvastatin of age and one of the following:

	Dr	ug/dose/date(s):
		18 years of age or older and one of the following: ember has trial and failure history with Praluent (alirocumab) OR Repatha (evolocumab)
		Yes □ No
	Dr	ug/dose/date(s):
	m( ez	ember has trial and failure history of at least 90 days of high dose rosuvastatin (20 mg/40 g) or atorvastatin (40 mg/80 mg, if rosuvastatin intolerant) therapy concurrently with etimibe (or documented intolerance/contraindication to statins/ezetimibe) AND provider has bmitted medical justification for use of Evkeeza (evinacumab-dgnb) over Praluent
	(a	lirocumab) and Repatha (evolocumab) □ Yes □ No
	Dr	ug/dose/date(s):
4.	Select one of the follow	ring:
		ill utilize maximally tolerated statin therapy with or without ezetimibe concurrently with for those 7 years of age and older)
		as submitted documented intolerance to statin and/or ezetimibe therapy or medical rationale e of statin or ezetimibe therapy
5.	Requested dose is 15 r	mg/kg every 4 weeks or less ☐ Yes ☐ No
	Member weight:	LB / KG (circle one)
PA	A Requirements for J	uxtapid (lomitapide mesylate):
1.	Member is enrolled in t	he Juxtapid/lomitapide REMS program and prescriber is monitoring in accordance with
	REMS requirements [	
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 follow	ring:
		s trial and failure history of Praluent (alirocumab) or Repatha (evolocumab)  date(s):
	atorvastatin documented	s trial and failure history of at least 90 days of high dose rosuvastatin (20 mg/40 mg) or (40 mg/80 mg, if rosuvastatin intolerant) therapy concurrently with ezetimibe (or d intolerance/contraindication to statins/ezetimibe) AND provider has submitted medical for use of Juxtapid (lomitapide mesylate) over Praluent (alirocumab) and Repatha b)
	Drug/dose/	date(s):
5.		ng potential, documentation of a negative pregnancy test obtained in the past 30 days is r has counseled member on risks associated with conceiving while utilizing Juxtapid and
		contraception □ Yes □ No Signature:
6.	Select one of the follow	ving:
	☐ Member w Juxtapid	ill utilize maximally tolerated statin therapy with or without ezetimibe concurrently with
	☐ Provider h	as submitted documented intolerance to statin and/or ezetimibe therapy or medical rationale e of statin or ezetimibe therapy
7.	Requested dose is 60 r	ng/day or less □ Yes □ No

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PΑ	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)
	<ul> <li>Member has diagnosis of heterozygous familial hypercholesterolemia (HeFH) with a baseline LDL-C level of ≥70 mg/dL (documentation required)</li> </ul>
2.	Member is 18 years of age or older □ Yes □ No
3.	Prescribed by, or in consultation with, a cardiologist or endocrinologist $\ \square$ Yes $\ \square$ 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:
	<ul> <li>Member will utilize maximally tolerated statin therapy with or without ezetimibe concurrently with Leqvio</li> </ul>
c	Provider has submitted documented intolerance to statin and/or ezetimibe therapy or medical rationale against use of statin or ezetimibe therapy
О.	Select one of the following:
	☐ Member is initiating therapy and requested dose does not exceed 284 mg every 3 months
	☐ Member is established on therapy and requested dose does not exceed 284 mg every 6 months
PΑ	Requirements for Niacin ER
1.	Diagnosis of severe hypertriglyceridemia (baseline triglycerides ≥500 mg/dL) ☐ Yes ☐ No
	If Yes, 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, 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

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PA Requirements for Praluent (alirocumab):
PA Requirements for Praluent (alirocumab):  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 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*
☐ 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)
Note: documentation of any and all intolerances to statins and/or ezetimibe must be provided
For any of the above diagnoses that require medical justification for use of Praluent over statin and/or ezetimibe therapy, please provide justification here:

Se	elect one of the following:
	☐ Member is 18 years of age or older
	$\square$ Member is 8 years of age or older and has a diagnosis of HeFH
Se	elect one of the following:
	<ul> <li>Member will utilize maximally tolerated statin therapy with or without ezetimibe concurrently with Praluent</li> </ul>
	<ul> <li>Provider has submitted documented intolerance to statin and/or ezetimibe therapy or medical ration against use of statin or ezetimibe therapy</li> </ul>
Se	elect one of the following:
	☐ Requested dose is 75 mg every 2 weeks
	☐ Requested dose is 300 mg every 4 weeks
	$\ \square$ Requested dose is 150 mg every 2 weeks <b>AND the member has one of the following:</b>
	☐ Diagnosis of homozygous familial hypercholesterolemia
	<ul> <li>Diagnosis of heterozygous familial hypercholesterolemia and member is undergoing LDL apheresis</li> </ul>
	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
	$\square$ Requested dose is 150 mg every 4 weeks <b>AND all of the following:</b>
	<ul> <li>Diagnosis of heterozygous familial hypercholesterolemia</li> <li>Member is under 18 years of age and weighs less than 50 kg</li> </ul>
\ Re	equirements for Repatha (evolocumab):
or P	Praluent (alirocumab): elect 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 again the use of statin therapy*
	☐ Member has a diagnosis of clinical ASCVD, is NOT at Very High Risk requiring therapy for secondar 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 ezetimile 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 secondar
	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 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*

	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
	* 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:
_	
2.	Select one of the following:
2.	Select one of the following:
2.	
	☐ Member is 18 years of age or older
3.	<ul> <li>☐ Member is 18 years of age or older</li> <li>☐ Member is 10 years of age or older and has a diagnosis of either HoFH or HeFH</li> <li>Select one of the following:         <ul> <li>a. Member will utilize maximally tolerated statin therapy with or without ezetimibe concurrently with Repatha</li> <li>b. Provider has submitted documented intolerance to statin and/or ezetimibe therapy or medical rationale</li> </ul> </li> </ul>
3.	<ul> <li>□ Member is 18 years of age or older</li> <li>□ Member is 10 years of age or older and has a diagnosis of either HoFH or HeFH</li> <li>Select one of the following:         <ul> <li>a. Member will utilize maximally tolerated statin therapy with or without ezetimibe concurrently with Repatha</li> <li>b. Provider has submitted documented intolerance to statin and/or ezetimibe therapy or medical rationale against use of statin or ezetimibe therapy</li> </ul> </li> </ul>
3.	<ul> <li>□ Member is 18 years of age or older</li> <li>□ Member is 10 years of age or older and has a diagnosis of either HoFH or HeFH</li> <li>Select one of the following:         <ul> <li>a. Member will utilize maximally tolerated statin therapy with or without ezetimibe concurrently with Repatha</li> <li>b. Provider has submitted documented intolerance to statin and/or ezetimibe therapy or medical rationale against use of statin or ezetimibe therapy</li> </ul> </li> <li>Select one of the following:</li> </ul>
3.	<ul> <li>□ Member is 18 years of age or older</li> <li>□ Member is 10 years of age or older and has a diagnosis of either HoFH or HeFH</li> <li>Select one of the following:         <ul> <li>a. Member will utilize maximally tolerated statin therapy with or without ezetimibe concurrently with Repatha</li> <li>b. Provider has submitted documented intolerance to statin and/or ezetimibe therapy or medical rationale against use of statin or ezetimibe therapy</li> </ul> </li> <li>Select one of the following:         <ul> <li>□ Requested dose is 140 mg every 2 weeks</li> </ul> </li> </ul>

## CONFIDENTIAL INFORMATION

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