

State of Oklahoma SoonerCare **PCSK9 Inhibitor Prior Authorization Form**

Sooner Select	♥ aetna
_	

Pharmacy Section									
Mem	ber N	ame:		Date	of Birth:		Member ID#	t:	
Phar	тасу	NPI:		Pharmacy Phon	ne:		Pharmacy Fa	ax:	
Phar	тасу	Name:		P	harmacist Na	ame:			
Pres	criber	· NPI:		Prescriber Name):		_ Specialty:		
Pres	criber	Phone:	<i>I</i>	Prescriber Fax:_		Drug N	lame/Strengt	th:	
NDC	:		Regimen:_		_ Fill Date:		Quantity:	Day Sup	oply:
Has	memb	er been trained	on proper ad	ministration and	storage of th	nis medicati	on? Yes	_No	
Phar	macis	st Signature:				Date:			
				Prescri	iber Section				
Page 1 of 2—Please complete and return all pages. Failure to complete all pages will result in processing delays. All information must be provided and SoonerCare may verify through further requested documentation. The member's prescription claim history will be reviewed prior to approval. For Initial Authorization (Initial approval will be for the duration of 3 months):									
		•		will be for the al	uration of 3 n	nontns):			
	Het	Documented fureceptor function Pre-treatment to History of tendor Dutch Lipid Clir mozygous familiar Documented furectionality via Untreated LDL- Documented Documented Documented LDL- Presence	al hypercholesinctional mutational mutational mutationality via generated cholestero an xanthomas in Network Critical hypercholestero and mutational mutational mutational mutational mutational mutational mutational mutational et al.	terolemia (HeFH) ion(s) in low-densitic testing <i>(result</i> : 1 >290mg/dL or LI in either the membiteria score of >8 erolemia (HoFH) of ion(s) in both LDL of <i>(results of gene</i> and at least 1 of the cutaneous xantho	ity lipoprotein s of genetic to DL-cholestero per, first degree confirmed by receptor alleletic testing managements to 10 managements of the street of the s	(LDL) recept testing must I (LDL-C) >1 tee relative, on 1 or more of the es or alleles thust be subtances by years of ag	tor alleles or a t be submitte 90mg/dL r second degr the following: known to affe mitted)	e d) ree relative ect LDL recepto	or
	hos con Dia Dia	pitalization in additions and dates agnosis/condition	ults with estables of occurrences:	farction, stroke, co ished cardiovascu e signifying establi	ular disease (0 shed CVD: Da	CVD). Please	e provide suppence:	porting diagno	ses/
2. H	ow wil	I this medication	be used? \square N	Monotherapy \square	Adjunct to sta	atin therapy,	diet, and exer	rcise	
3. P		specify the mem		• •					
a)				Dosir					
b)			ū	h-dose statin thera					
c)				DL-C level following conducted to ver			py:		

Page 1 of 2

Fax completed prior authorization request form to 888-601-8461 or submit Electronic Prior Authorization through CoverMyMeds® or SureScripts. All requested data must be provided. Incomplete forms or forms without the chart notes will be returned. Pharmacy Coverage Guidelines are available at AetnaBetterHealth.com/Oklahoma.

CONFIDENTIALITY NOTICE
This document, including any attachments, contains information which is confidential or privileged. If you are not the intended recipient, be aware that any disclosure, copying, distribution, or use of the contents of this information is prohibited. If you have received this document in error, please notify the sender immediately by telephone to arrange for the return of the transmitted documents or to verify their destruction.

Pharm - 38 6/17/2024



State of Oklahoma SoonerCare **PCSK9 Inhibitor Prior Authorization Form**

Sooner Select	⇔ aetna°
2001 lei Select	Vacuia

Weilib	r Name: Date of Birth: Wember ID#:					
	Prescriber Section					
	Page 2 of 2—Please complete and return all pages. Failure to complete all pages will result in processing delays. For Initial Authorization, Continued:					
into	e member has <u>not</u> been adherent to high-dose statin therapy for at least 12 continuous weeks, is the member erant to statin therapy? Yes No If yes, please indicate 1 of the following: Rhabdomyolysis - creatine kinase (CK) labs verifying this diagnosis must be provided. An FDA labeled contraindication to all statins. Provide contraindication: Documented intolerance to at least 2 different statins at lower doses or at intermittent dosing: Please provide all of the following: 1) Medication/strength: Dosing regimen: Duration of treatment: Reason for discontinuation:					
	2) Medication/strength: Dosing regimen:					
	Duration of treatment: Reason for discontinuation:					
a) 6. If th a) 7. Ple 8. If e. why 9. Me 10. Has For Co 1. Has 2. Has	the member had a recent trial of a statin with ezetimibe? Yes No If yes, please provide statin tried with ezetimibe: trial dates: e member is intolerant to statin therapy, has the member had a recent trial of ezetimibe alone? Yes No If yes, please provide ezetimibe trial dates: se provide member's LDL-C level following ezetimibe therapy with statin therapy or without statin therapy: etimibe has not been tried either with or without a statin, please provide a patient-specific, clinically significant reason ezetimibe is not appropriate for the member: her's baseline LDL-C: Goal LDL-C: Goal LDL-C: the member been counseled on proper administration and storage of PCSK9 therapy? Yes No tinued Authorization: member been compliant with PCSK9 Inhibitor treatment? Yes No PCSK9 Inhibitor treatment been effective for this member? Yes No se provide a recent LDL-C level for this member: Date taken:					
Presci	(Page 2 of 2) ber Signature: Date:					

Fax completed prior authorization request form to 888-601-8461 or submit Electronic Prior Authorization through CoverMyMeds® or SureScripts. All requested data must be provided. Incomplete forms or forms without the chart notes will be returned. Pharmacy Coverage Guidelines are available at

AetnaBetterHealth.com/Oklahoma.

CONFIDENTIALITY NOTICE
This document, including any attachments, contains information which is confidential or privileged. If you are not the intended recipient, be aware that any disclosure, copying, distribution, or use of the contents of this information is prohibited. If you have received this document in error, please notify the sender immediately by telephone to arrange for the return of the transmitted documents or to verify their destruction.

By signature, the physician confirms the criteria information above is accurate and verifiable in patient records. Please do not send in chart

notes. Specific information will be requested if necessary. Failure to complete this form in full will result in processing delays.