## SENTARA HEALTH PLANS

## PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST\*

<u>Directions</u>: The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; fax to 1-800-750-9692. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. If information provided is not complete, correct, or legible, authorization may be delayed.

## CNS Stimulants for Adults Age 19 and Above

 A review of written documentation to substantiate a complete, appropriate, and covered diagnosis for both new starts and members currently receiving any CNS stimulant listed below will be required before Prior Authorization approval. <u>Prescribing history alone WILL NOT meet criteria for approval.</u>

MEMBER & PRESCRIBER INFOR	MATION: Authorization may be delayed if incomplete.			
Member Name:				
Member Sentara #:				
Prescriber Name:				
Prescriber Signature:				
Office Contact Name:				
Phone Number:				
NPI #:				
DRUG INFORMATION: Authorization				
Drug Form/Strength:				
Dosing Schedule:	Length of Therapy:			
Diagnosis: ICD Code:				
Weight (if applicable):	Date weight obtained:			
• Will the member be discontinuing a previously prescribed central nervous system (CNS) stimulant medication if approved for requested medication?				
-	□ Yes <b>OR</b> □ No			
<ul> <li>If yes, please list the medication that will be approval along with the corresponding effe</li> </ul>	e discontinued and the medication that will be initiated upon ctive date.			
Medication to be discontinued:	Effective date:			
Medication to be initiated:	Effective date:			

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<b>DRUG(S) REQUESTED:</b>	Check applicable drug(s) below	v. Box(es) <b>must</b> be checked to qua	alify, or
authorization process will be del	ayed.		

	Adhansia XR®		Adzenys XR-ODT®  Adzenys ER®  Suspension		amphetamine/ dextroamphetamine (Adderall®)		amphetamine/ dextroamphetamine ER (Adderall XR®)
	amphetamine sulfate (Evekeo®)		Azstarys®		Cotempla XR/ODT®		dexmethylphenidate (Focalin®)
	dexmethylphenidate ER (Focalin XR®)		dextroamphetamine (Dextrostat®)		dextroamphetamine (ProCentra®)		dextroamphetamine (Zenzedi®)
	dextroamphetamine ER (Dexedrine Spansule®)		Dyanavel® XR Suspension Dyanavel® XR Chewable Tablets	_	Evekeo ODT®		Jornay PM®
	methamphetamine (Desoxyn®)		methylphenidate ER (Aptensio XR®)		methylphenidate ER (Concerta®)		methylphenidate TD Patch (Daytrana®)
	methylphenidate ER (Metadate ER®/ Ritalin SR®)		methylphenidate (Methylin®/Ritalin®)		methylphenidate LA (Ritalin LA®)		methylphenidate CD (Metadate CD®)
	Mydayis <sup>®</sup>		Quillichew® ER		Quillivant XR®		Vyvanse <sup>®</sup>
	Xelstrym <sup>™</sup> (dextroamphetamine)						
<b>DIAGNOSES:</b> Check applicable diagnosis below with ICD Code and description. For **BINGE EATING DISORDER, obtain BED specific form, found under "Vyvanse (Binge Eating Disorder). **							
	□ ADHD/ADD: ICD-9/10: Description:						
,	*please complete table below and attach/fax any documentation as requested						
	□ Narcolepsy: ICD-9/10: Description:						
	*please attach and fax documentation (polysomnogram and MSLT results) to support diagnosis						
	☐ Other*: ICD-9/10: Description:						
	*please attach and fax documentation (i.e. chart notes, previous therapies tried) to support diagnosis						
*NON-FDA approved indications - submit two (2) peer reviewed clinical studies documenting the safety							
and efficacy of the specified drug for that particular indication.							

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**CLINICAL CRITERIA:** Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.

Na	ıme	e of Diagnosing Prescriber:		Date of Diagnosis:			
of syl	the mpt	prescriber, the date of diagnosis, and cop	oies of tes	either a child or an adult, please submit the name ting and chart notes detailing signs and e prescribing physician in the table below or as a			
☐ Existence of at least 5 symptoms for a minimum of 6 months. (indicate symptoms below)							
		<b>Inattentive Symptoms</b> : 5 or more					
		<b>Hyperactive-Impulsive Symptoms</b> : 5 or :	more				
		Combined Symptoms: 10 or more ADHE or more hyperactive-impulsive symptoms	) symptoi	ms including 5 or more inattentive symptoms <b>AND</b> 5			
	Documentation that symptoms impair or compromise normal functioning.						
	Do	ocumentation that symptoms are present in t	wo (2) oi	more settings/environments (indicate settings):			
	1.		2				
		Medical Chart/Progress Notes documenting	g childho	od diagnosis and/or symptoms			
		School Records					
		Corroborated by a relative/friend					
		Not Available					
		ymptoms are not better explained by another isorder, Substance Abuse, Dissociative Diso		(e.g., Schizophrenia, Mood Disorder, Anxiety ersonality Disorder)			
		ne diagnosis has been verified using a standaraluation	ırdized ra	ting scale, patient interview, or psychological			
		Adult Self-Report Scale- V1.1		Member Interview			
		8		Psychological Evaluation			
	Αľ		LE USE	ITERIA, PSYCHOLOGICAL EVALUATION, D TO MAKE OR VERIFY THE DIAGNOSIS APPROVAL.			

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If requesting Brand or generic when applicable for Adhansia XR<sup>®</sup>, Adzenys<sup>®</sup>, Aptensio XR<sup>®</sup>, Azstarys<sup>®</sup>, Cotempla XR ODT<sup>®</sup>, Daytrana<sup>®</sup>, Dyanavel<sup>®</sup> XR, Evekeo<sup>®</sup>/Evekeo ODT<sup>®</sup>, Jornay PM<sup>®</sup>, Mydayis<sup>®</sup>, Quillichew<sup>®</sup> ER, Quillivant<sup>®</sup> XR or Xelstrym<sup>™</sup>, BOTH of the following criteria MUST be met:

☐ Member must have tried and failed 30 days of therapy with two (2) of the following:

amphetamine-dextroamphetamine IR/ER (generic Adderall/Adderall XR®)	dexmethylphenidate IR/ER (generic Focalin®/Focalin XR®)
dextroamphetamine IR/SR (generic Dextrostat®/Procentra®/Zenzedi®/Dexedrine® IR/ER)	methylphenidate IR/ER (generic Ritalin®/Methylin®/Ritalin SR®/Ritalin LA®/Concerta®/ Metadate CD®/Metadate ER®

☐ Member must have tried and failed <u>30 days of therapy</u> with Vyvanse<sup>®</sup> (<u>NOT</u> required for amphetamine sulfate (Evekeo<sup>®</sup>) or Evekeo ODT<sup>®</sup> requests)

Please be aware if this request is for a dose that <u>EXCEEDS</u> Sentara Health's Maximum Daily Dosage Limits, a second prior authorization request will need to be submitted for dosage approval. The correct form can be downloaded from <a href="http://providers.sentarahealthplans.com/">http://providers.sentarahealthplans.com/</a>

Not all drugs may be covered under every Plan.

If a drug is non-formulary on a Plan, documentation of medical necessity will be required.

\*\*Use of samples to initiate therapy does not meet step edit/ preauthorization criteria. \*\*

\*Previous therapies will be verified through pharmacy paid claims or submitted chart notes. \*