NARCOLEPSY AGENTS PRIOR AUTHORIZATION REQUEST FORM

and the second	<i>OptumRx</i> P.O. Box 25184 Santa Ana, CA, 92799 Phone: (800) 310-6826 Fax: (866) 940-7328	Optum Rx [®] United Healthcare
		Community Plan
Today's Date		
/ /		

Note: This form must be completed by the prescribing provider.

All sections must be completed or the request will 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
Return Fax # - - -	Return Phone # - - - -
Check box if requesting retro-active PA	Date(s) of service requested for retro-active eligibility (if applicable):

Note: Submit PA requests for retroactive claims (dates of service prior to eligibility determination, but within established eligibility timelines) with dates of service prior to 30 calendar days of submission separately from current PA requests (dates of service 30 calendar days or less and going forward).

Requested Medication	Quantity	Dosing

PA Requirements for Nuvigil (armodafinil):		
The member is 18 years of age or older and has one of the following diagnoses:		
 Bipolar depression in conjunction with appropriate medical intervention(s) List any other medical intervention(s) being utilized for bipolar depression (e.g., mood stabilizers): 		
□ Narcolepsy with excessive daytime sleepiness		
 Obstructive sleep apnea/hypopnea syndrome with residual excessive daytime sleepiness in conjunction with appropriate medical intervention(s) List any other medical intervention(s) being utilized for obstructive sleep apnea (e.g., PAP, OPT, etc.)? 		
□ Shift work sleep disorder		

PA Requirements for Provigil (modafinil):			
Select ONE of	the following:		
1)	The member is 6 years of age or older and has one of the following diagnoses:		
	Attention deficit hyperactivity disorder (ADHD)		
2)	 Narcolepsy with excessive daytime sleepiness The member is 18 years of age or older and has one of the following diagnoses: 		
	Depression-related fatigue in conjunction with appropriate medical intervention(s)		
	 List any other medical intervention(s) being utilized for depression (e.g., antidepressants): 		
	□ Idiopathic hypersomnia		
	 Obstructive sleep apnea/hypopnea syndrome with residual excessive daytime sleepiness in conjunction with appropriate medical intervention(s) List any other medical intervention(s) being utilized for obstructive sleep apnea (e.g., PAP, OPT, etc.)? 		
	□ Shift work sleep disorder		
	□ Sleep deprivation		
	Steinert myotonic dystrophy syndrome		
	 Unipolar or bipolar depression in conjunction with appropriate medical intervention(s) List any other medical intervention(s) being utilized for unipolar/bipolar depression (e.g., antidepressants/mood stabilizers): 		
PA Require	ments for Sunosi (solriamfetrol):		

The member is 18 years of age or older and has one of the following diagnoses:

□ Narcolepsy with excessive daytime sleepiness

□ Obstructive sleep apnea/hypopnea syndrome with residual excessive daytime sleepiness in conjunction with appropriate medical intervention(s)

- List any other medical intervention(s) being utilized for obstructive sleep apnea (e.g., PAP, OPT, etc.)?
- Has the member had a previous trial and failure with any of the following in the past year:
 - Modafinil
 Dates of use:
 - Armodafinil Dates of use: ______

If no, please provide any other medical justification for use: _____

PA Require	ments for Wakix (pito	lisant):
The member is	18 years of age or older an	nd has one of the following diagnoses:
Narcole	psy with cataplexy or exces	sive daytime sleepiness
	tive sleep apnea/hypopnea priate medical intervention(syndrome with residual excessive daytime sleepiness in conjunction
•	List any other medical inte OPT, etc.)?	ervention(s) being utilized for obstructive sleep apnea (e.g., PAP,
•	Has the member had a pr	evious trial and failure with any of the following in the past year:
	🗆 Modafinil 🛛 Da	tes of use:
		tes of use:
		ny other medical justification for use:
A Require	ments for Xyrem (soc	lium oxybate):
nitial Authoriz		
	ONE of the following:	
1)	The member is 7 years of	age or older and has narcolepsy with cataplexy or excessive
	daytime sleepiness diagne	osis □ Yes □ No
		sted dose per day:
	Please provide memb	er's weight (include date of collection):
2)	 2) The member is 18 years of age or older and has fibromyalgia diagnosis □ Yes □ No Has the member had a previous trial and failure with ONE of the following? 	
		Dates of use:
		Medication name and dates of use:
		Medication name and dates of use:
	 Anticonvulsan (gabapentin, preg 	Medication name and dates of use:
	\Box NSAIDs and A	PAP Dates of use:
		t trialed all of the above agents, please provide medical justification ent or an agent in that class was not trialed.

Please provide requested dose per day: _____

Reauthorization

- 1) Please provide the following information showing an attempt to decrease dose or trial and failure of an alternative therapy within the past year:
 - Date of dose reduction attempt: ______
 - Original dose member was prescribed:______

 - Outcomes of dose reduction: ______
 - Trial and failure of an alternative therapy (name of medication, date of trial, and an explanation as to how the member failed):
- 2) Please provide documentation showing continued benefit from the medication (i.e., reduction in frequency of cataplexy, reduction in symptoms of excessive daytime sleepiness, etc.) without significant adverse events (documentation must include most recent chart notes)

PA Requirements for Xywav (calcium/magnesium/potassium/sodium oxybates solution):			
Initial Authorization			
Select ONE of the following:			
1) The member is 7 years of age or older and has narcolepsy with cataplexy or excessive			
daytime sleepiness diagnosis □ Yes □ No			
Please provide requested dose per day:			
Please provide member's weight (include date of collection):			
2) The member is 18 years of age or older has idiopathic hypersomnia \Box Yes \Box No			
Please provide requested dose per day:			
Reauthorization			
 Please provide the following information showing an attempt to decrease dose or trial and failure of an alternative therapy within the past year: Date of dose reduction attempt: 			
Original dose member was prescribed:	_		
Dose member was reduced to:			
Outcomes of dose reduction:	_		
• Trial and failure of an alternative therapy (name of medication, date of trial, and an			
explanation as to how the member failed):			
 Please provide documentation showing continued benefit from the medication (i.e., reduction in frequency of cataplexy, reduction in symptoms of excessive daytime sleepiness, etc.) without significant adverse events (documentation must include most recent chart notes) 			

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