

Pharmacy Prior Approval Request for Antinarcolepsy: Wakix

Beneficiary Information			
1. Beneficiary Last Name:	2. First Na	ıme:	
3. Beneficiary ID #:	4. Beneficiary Date of Birth		5. Beneficiary Gender:
Prescriber Information			
6 Prescribing Provider NPI #:			
7. Requester Contact Information - Nam	 ie:	Phone #:	Ext
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Drug Information			
8. Drug Name:	9. Strength:		10. Quantity Per 30 Days:
11. Length of Therapy (in days): up to 3	30 Days □ 60 Days □ 90 Days	☐ 120 Days [☐ 180 Days ☐ 365 Days ☐ Other
Clinical Information			
1. Is the beneficiary 18 years of age or older? ☐ Yes ☐ No 2. Does the beneficiary have daily periods of irrepressible need to sleep or daytime lapses into sleep occurring for at least three (3) months? ☐ Yes ☐ No 3. Is the beneficiary receiving treatment with sedative hypnotic agents (e.g., zolpidem, eszopiclone, zaleplon, benzodiazepines, barbiturates)? ☐ Yes ☐ No 4. Will the beneficiary use drugs that prolong the QT interval (e.g., quinidine, procainamide, disopyramide, amiodarone, sotalol, ziprasidone, chlorpromazine, thioridazine, moxifloxacin) concomitantly? ☐ Yes ☐ No 5. Will the beneficiary use histamine-1 (H1) receptor antagonists (e.g., pheniramine maleate, diphenhydramine, promethazine, imipramine, clomipramine, mirtazapine) concomitantly? ☐ Yes ☐ No 6. Does the beneficiary have a history of prolonged QTc interval (e.g., QTc interval > 450 milliseconds)? ☐ Yes ☐ No 7. Does the beneficiary have end-stage renal disease (estimated glomerular filtration rate [eGFR] < 15 mL/min/1.73 m2)? ☐ Yes ☐ No 8. Does the beneficiary have severe hepatic impairment? ☐ Yes ☐ No 9. Does the beneficiary have a diagnosis of cataplexy with narcolepsy? ☐ Yes ☐ No 10. Does the beneficiary have a diagnosis of narcolepsy? ☐ Yes ☐ No 11. Does the beneficiary have an adequate documented trial and failure of, or contraindication to, modafinil and armodafinil? ☐ Yes ☐ No 11. Does ☐ No 12. Please explain if contraindicated: ☐ Yes ☐ No 13. Please explain if contraindicated: ☐ Yes ☐ No			
For continuation of therapy, please and 12. If treating narcolepsy, has the benefit treatment baseline as measured by Sleepiness Scale, Cleveland Adolesce 13. If treating cataplexy with narcolepsy baseline? ☐ Yes ☐ No 14. Has the beneficiary experienced any dreams or nightmares, anhedonia, a prolongation, sleep disorder, suicide	iciary reported a documented re a validated scale (e.g., Epworth ent Sleepiness Questionnaire, or t, has the beneficiary had reduce treatment-restricting adverse e anxiety, bipolar disorder, depres	Sleepiness Scal a Visual Analord d frequency of ffects (e.g., ab sion or depress	le, Stanford Sleepiness Scale, Karolinska og Scale)? Yes No f cataplexy attacks from pretreatment normal behavior, abnormal
Signature of Prescriber:(Prescri	riber Signature Mandatory)	Date: _	

I certify that the information provided is accurate and complete to the best of my knowledge, and I understand that any falsification, omission, or concealment of material fact may subject me to civil or criminal liability.

Fax this form to: (833) 404-2393

Pharmacy PA Call Center: (833) 585-4309