

NC Medicaid and NC Health Choice Pharmacy Prior Approval Request for Amondys 45

Beneficiary Information		
1. Beneficiary Last Name:	2. First Name:	
3. Beneficiary ID #:	4. Beneficiary Date of Birth:	5. Beneficiary Gender:
Prescriber Information		
6. Prescribing Provider NPI #:		:Ext
7. Requester Contact Information - Name:	Phone #	:Ext
Drug Information		
8. Drug Name:	9. Strength:	10. Quantity Per 30 Days:
11. Length of Therapy (in days): 🗆 ເ		
Clinical Information		
For initial authorization requests: 1. What is the beneficiary's weight? 2. Does the beneficiary have a diagnosis of	Duchenne Muscular Dystrophy? ☐ Yes ☐	
3. Are medical records attached to this request that confirm the mutation of the Duchenne Muscular Dystrophy gene is amenable to		
exon 45 skipping? □ Yes □ No 4. Is Amondys 45 being prescribed by or in consultation with a neurologist? □ Yes □ No		
5. Does the beneficiary retain meaningful voluntary motor function (beneficiary is able to speak, manipulate objects using upper		
extremities, ambulate, etc? Yes No		
6. Has the beneficiary has been assessed for any physical therapy and/or occupational therapy needs? ☐ Yes ☐ No 7. Has the beneficiary's serum cystatin C, urine dipstick, and urine protein-to-creatinine ratio (UPCR) have been measured prior to		
starting therapy? ☐ Yes ☐ No		
8. Does the prescriber attest that serum cystatin C, urine dipstick, and urine protein-to-creatinine ratio will be measured during treatment (monthly urine dipstick with serum cystatin C and urine protein-to-creatinine ratio every 3 months)? Yes No		
9. Has baseline documentation of at least 1 of the following been performed: Dystrophin level, 6-minute walk test (6WMT)		
or other timed function tests, Upper limb function (ULM) test, North Star Ambulatory Assessment (NSAA),Forced Vital		
Capacity (FVC) % predicted, of Performance of Upper Limb (PUL)? Yes No List 10. Is the beneficiary taking any other RNA antisense agent or any other gene therapy? Yes No		
11. 12. Is the beneficiary receiving a dose that does not exceed 30mg/kg once per week? Yes No		
For reauthorization (answer 1-12): 13. Please attach documentation that she to pretreatment baseline in at least 1 of t slowed rate of decline in 6MWT or other tim	the following: Increase in dystrophin level;	OR Stability, improvement, or
ULM test; OR Stability, improvement, or slot decline in FVC% predicted; OR Improvement treatment-restricting adverse effects (e.g. re	wed rate of decline in NSAA; OR Stability, int in quality of life; and that the beneficiary	mprovement, or slowed rate of
Signature of Prescriber:		Date:
(Presc	criber Signature Mandatory)	

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 1-866-940-7328

06/23/2023