POLICY NAME Spinraza (nusinersen) POLICY # 2609P

Criteria

Coverage Criteria	
	Diagnosis of Spinal Muscular Atrophy (SMA) types I, II, or III
	Documentation of 5q SMA homozygous gene mutation, homozygous gene deletion, or compound heterozygote
	Ordered by a Geneticist or provider specializing in the treatment of SMA
	Member is 15 years of age or younger at initiation of treatment
	Documented baseline motor milestone scores according to one of the following age-appropriate assessments:
	Hammersmith Infant Neurologic Exam (HINE)
	Modified Hammersmith Functional Motor-Scale
	Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders (CHOP –INTEND)
	Bayley Scales of Infant and Toddler Development (BSID-III)
	Motor Function Measure 32 (MFM-32)
	Review of chart notes and labs documenting diagnosis and confirming that patient has met all of the above requirements for treatment with Spinraza by both a pharmacist and medical director
Evaluation Oritaria - Any of the following provents according	
EXCI	usion Criteria – Any of the following prevents coverage
	Spinraza will not be covered after treatment with Zolgensma because its use following Zolgensma infusion has not been studied and is considered experimental/investigational Note: Patients in the clinical trials that received Zolgensma before the age of 2 were followed up to 5 years post-treatment and did not require additional medications
	Spinraza will not be covered in combination with Evrysdi because the concomitant use of these two drugs has not been studied and is considered experimental/investigational