

Evrysdi® (Risdiplam) Prior Authorization Form

Member Name: _____ Date of Birth: _____ Member ID#: _____

Drug Information

Pharmacy billing (NDC: _____) Start Date (or date of next dose): _____

Member's Weight: _____ Date Taken: _____ Dose: _____ Regimen: _____

Billing Provider Information

Pharmacy NPI: _____ Pharmacy Name: _____

Pharmacy Phone: _____ Pharmacy Fax: _____

Will Evrysdi® be constituted to an oral solution by a pharmacist prior to dispensing? Yes ___ No ___

Will Evrysdi® be shipped via cold chain supply to adhere to the storage and handling requirements? Yes ___ No ___

Pharmacist signature: _____ Date: _____

Prescriber Information

Prescriber NPI: _____ Prescriber Name: _____

Prescriber Phone: _____ Prescriber Fax: _____ Specialty: _____

Criteria

*Page 1 of 2—Please complete and return all pages. Failure to complete all pages will result in processing delays.

For Initial Authorization (Initial approval will be for the duration of 6 months):

1. What is the member's diagnosis?
 - Spinal Muscular Atrophy (SMA)
 - A. What type of SMA does the member have (0-4)? _____
 - B. Does member currently have symptoms consistent with SMA? Yes ___ No ___
 - C. Has the diagnosis been confirmed by molecular genetic testing? Yes ___ No ___
 - D. Does member have biallelic pathogenic variants in the survival motor neuron gene 1 (SMN1)? Yes ___ No ___
 - Other: _____
2. Is member currently dependent on permanent ventilation? Yes ___ No ___
 - A. If member is currently dependent on permanent ventilation, please specify number of hours per day member requires ventilator support: _____
3. Is Evrysdi® being prescribed by a neurologist, specialist with expertise in the treatment of SMA, or an advanced care practitioner with a supervising physician who is a neurologist or specialist with expertise in the treatment of SMA? Yes ___ No ___
4. Does prescriber agree to evaluate the member's liver function prior to initiating Evrysdi® and verify the member does not have severe hepatic impairment (Child-Pugh C)? Yes ___ No ___
5. Has the member or caregiver been instructed on the proper storage of Evrysdi® and how to prepare the prescribed daily dose of Evrysdi® prior to administration of the first dose? Yes ___ No ___
6. For female members of reproductive potential, please answer all of the following:
 - A. Is the member pregnant? Yes ___ No ___
 - B. Does the member have a negative pregnancy test prior to initiation of Evrysdi® treatment? Yes ___ No ___
 - C. Is the member willing to use effective contraception during treatment with Evrysdi® and for at least 1 month after the last dose? Yes ___ No ___
7. For male members of reproductive potential, has the member been counseled on the potential effects of Evrysdi® on fertility and is the potential of compromised male fertility acceptable? Yes ___ No ___
8. Has member previously received treatment with Zolgensma® (onasemnogene abeparvovec-xioi)? Yes ___ No ___
9. Has the member previously been treated with Spinraza® (nusinersen)? Yes ___ No ___
 - A. If yes, will the member discontinue treatment with Spinraza® upon approval of Evrysdi®? Yes ___ No ___
10. Has a baseline assessment been performed and documented using a functionally appropriate exam [e.g., Hammersmith Infant Neurological Exam (HINE), Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders (CHOP-INTEND), Upper Limb Module (ULM) Test, or Hammersmith Functional Motor Scale Expanded (HFMSE)]? Yes ___ No ___
 - A. If yes, please indicate the exam performed: _____
 - B. Please provide member's baseline score to exam listed above: _____

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Fax completed prior authorization request form to **888-601-8461** or submit Electronic Prior Authorization through CoverMyMeds® or SureScripts. All requested data must be provided. Incomplete forms or forms without the chart notes will be returned. Pharmacy Coverage Guidelines are available at **AetnaBetterHealth.com/OKlahoma.**

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*Page 2 of 2—Please complete and return all pages. Failure to complete all pages will result in processing delays.

For Continued Authorization:

1. Has the member previously been approved through the SoonerCare prior authorization process? Yes ___ No ___
A. If no, please complete the initial authorization section above.
2. Is member responding to the medication as demonstrated by a clinically significant improvement or maintenance of function from pretreatment baseline status using the same exam as performed at baseline assessment?
Yes ___ No ___
3. Please indicate exam used to perform assessment: _____
A. Please provide member's baseline score to exam listed above: _____
B. Please provide member's current score to exam listed above: _____
4. If member is currently dependent on permanent ventilation, please specify number of hours per day member requires ventilator support: _____

Additional Information: _____

Please complete and return all pages. Failure to complete all pages will result in processing delays.

Prescriber Signature: _____ **Date:** _____

I certify that the indicated treatment is medically necessary and all information is true and correct to the best of my knowledge. Please do not send in chart notes. Specific information will be requested if necessary. Failure to complete this form in full will result in processing delays.

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