





Fax Completed Form To 1.833.404.2392

Prescriber Help Desk 1.833.587.2012

Online

covermymeds.com/main/

Request for Prior Authorization Ensifentrine (Ohtuvayre)

(PLEASE PRINT - ACCURACY IS IMPORTANT)

| | • | , | prior-authorization-iorms/ | | | |
|--|--|------------------------|-------------------------------|--|--|--|
| IA Medicaid Member ID # | Patient name | DOB | | | | |
| Patient address | | | | | | |
| Provider NPI | Prescriber name | | Phone | | | |
| Prescriber address | | | Fax | | | |
| Pharmacy name Address | | | Phone | | | |
| Prescriber must complete all informa | । ation above. It must be legible, corre | ct, and complete or fo | rm will be returned. | | | |
| Pharmacy NPI | Pharmacy fax | NDC | | | | |
| | | | | | | |
| Prior authorization (PA) is required for ensifentrine (Ohtuvayre). Requests for non-preferred agents may be considered when documented evidence is provided that the use of the preferred agent(s) would be medically contraindicated. Payment will be considered for an FDA approved or compendia indicated diagnosis for the requested drug when the following conditions are met: 1. Request adheres to all FDA approved labeling for requested drug and indication, including age, dosing, contraindications, warnings and precautions, drug interactions, and use in specific populations; and | | | | | | |
| 2. Patient has a diagnosis of moderate to severe COPD when all of the following are met: | | | | | | |
| a. FEV1/FVC ratio < 0.7; and | | | | | | |
| b. Post-bronchodilato | b. Post-bronchodilator FEV1 % predicted of 30% to 79%; and | | | | | |
| c. Modified Medical Ro 10; and | esearch Council (mMRC) dyspnea so | core of ≥ 2 or a COPD | Assessment Test (CAT) score ≥ | | | |
| 3. Patient is adherent with COPD treatments, meeting one of the following criteria: | | | | | | |
| The patient has a blood eosinophil of ≥ 100 and has experienced an exacerbation while adherent to a current 60-day trial of a triple combination regimen consisting of a long-acting beta agonist (LABA), a long-acting muscarinic antagonist (LAMA), and an inhaled corticosteroid (ICS); or | | | | | | |
| | The patient has a blood eosinophil of < 100 and has experienced an exacerbation while adherent to a current 60-day trial of a dual combination regimen consisting of a LABA and LAMA; and | | | | | |
| 4. Dual or triple combination regimen will be continued in combination with ensifentrine (Ohtuvayre). | | | | | | |
| The required trials may be overridden when documented evidence is provided that the use of these agents would be medically contraindicated. | | | | | | |
| If the criteria for coverage are met, initial authorization will be given for 6 months to assess the response to treatment. Additional authorizations will be considered upon documentation of a response to treatment (e.g. improved dyspnea, decrease exacerbations) and patient continues their dual or triple combination regimen. | | | | | | |
| Non-Preferred | | | | | | |
| ☐ Ohtuvayre | | | | | | |
| Strength | Usage Instructions | Quantity | Day's Supply | | | |
| | | | | | | |
| Diagnosis: | | | | | | |
| Provide all of the following inform | nation for a diagnosis of modera | te to severe COPD: | | | | |
| a. FEV1/FVC ratio: | Date obtained: | | | | | |
| Date obtained: | | | | | | |

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| c. | mMRC dyspnea score: | OR CAT score : | Date obt | tained: | | |
|-------------------------|--|-----------------------------|----------------|---|--|--|
| Blood eosinophil count: | | Date obtain | Date obtained: | | | |
| | atient has a blood eosinophil o triple combination regimen: | f ≥ 100 and has experienced | exacerbation v | vhile adherent to a current 60 day trial of | | |
| L | ABA Trial: | | | | | |
| Na | ame/dose: | | Trial da | ates: | | |
| Fa | ailure reason/medical contraindica | ation: | | | | |
| L | AMA Trial: | | | | | |
| Na | ame/dose: | Trial dates: | | | | |
| Fa | ailure reason/medical contraindica | ation: | | | | |
| IC | S Trial: | | | | | |
| Na | ame/dose: | | Trial da | ates: | | |
| | ailure reason/medical contraindica | | | | | |
| а | dual combination regimen: | . Too and not experienced | exacerbation (| vhile adherent to a current 60 day trial of | | |
| | | | Trial dates: | | | |
| | me/dose:Trial dates: lure reason/medical contraindication: | | | | | |
| L | AMA Trial: | | | | | |
| Na | ame/dose: | | Trial dates: | | | |
| | ailure reason/medical contraindica | | | | | |
| Re | enewal: | | | | | |
| Do | ocument response to treatmen | t: | | | | |
| At | patient currently on dual or trip | umentation as necessary. |] Yes | No | | |
| P | rescriber signature (Must match pi | rescriber listed above.) | | Date of submission | | |
| \Box | | | | | | |

IMPORTANT NOTE: In evaluating requests for prior authorization the consultant will consider the treatment from the standpoint of medical necessity only. If approval of this request is granted, this does not indicate that the member continues to be eligible for Medicaid. It is the responsibility of the provider who initiates the request for prior authorization to establish by inspection of the member's Medicaid eligibility card and, if necessary by contact with the county Department of Health and Human Services, that the member continues to be eligible for Medicaid.

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