

Prior Authorization Criteria

ALYFTREK™ (vanzacaftor, tezacaftor, and deutivacaftor) PA:

ALYFTREKTM (*vanzacaftor, tezacaftor, and deutivacaftor*) is indicated for the treatment of cystic fibrosis (CF) in patients aged 6 years and older who have at least one F508del mutation or another responsive mutation in the CFTR gene.

Prior authorization is required for ALYFTREKTM. Prior authorization approval will be considered when the following criteria are met. Along with the Universal PA Form, please submit any supporting clinical documentation.

Initial Authorization: 6 months

- 1. Age of the patient is within the age range as recommended by the FDA label; AND
- 2. Diagnosis of CF; **AND**
- 3. Prescribed by or in consultation with a CF specialist/pulmonologist specializing in treating CF patients. Please submit the name of the CF-treating or consulting specialist/pulmonologist on the request. Please also provide chart documentation from the consulting provider, including the name, strength, and dosing instructions of CF drug; **AND**
- 4. Patient has a diagnosis of CF with a CFTR gene mutation responsive to ALYFTREK™ as outlined in the FDA labeling.
 - If the patient's genotype is unknown, an FDA-cleared CF mutation test should be used to confirm the presence of at least one indicated mutation.
 - Please submit laboratory results documenting the patient's CFTR mutation upon request; AND
- 5. ALYFTREK™ is not prescribed concurrently with other CFTR modulators (e.g., Trikafta®, Orkambi®, Kalydeco®, Symdeko®); **AND**
- 6. Baseline measures submitted by provider of **ALL** of the following:
 - a. For age-appropriate patients, percent predicted expiratory volume in 1 second (ppFEV1)
 - b. Body mass index (BMI)
 - c. Pulmonary exacerbations- number in preceding 6 months
 - d. Liver function tests (e.g., ALT, AST, alkaline phosphatase, bilirubin); AND
- 7. Dose does not exceed vanzacaftor 20 mg, tezacaftor 100 mg, and deutivacaftor 250 mg (two tablets of vanzacaftor 10 mg, tezacaftor 50 mg, and deutivacaftor 125 mg) per day.

<u>Reauthorization</u>: 12 months with evidence of appropriate clinical response to therapy

- MISSISSIPPI DIVISION OF
- 1. Prescribed by or in consultation with a CF specialist/pulmonologist specializing in treating CF patients.

 Please submit the name of the CF-treating or consulting specialist/pulmonologist on the request. Please also provide chart documentation from the consulting provider, including the name, strength, and dosing instructions of CF drug; AND
- 2. ALYFTREK™ is not prescribed concurrently with other CFTR modulators (e.g., Trikafta®, Orkambi®, Kalydeco®, Symdeko®); **AND**
- 3. Provider attests that the patient has achieve a clinically meaningful response and displayed tolerance while on ALYFTREKTM based on ALL the following:
 - a. For age-appropriate patients, improved or stable lung function as demonstrated by percent predicted expiratory volume in 1 second (ppFEV1)
 - b. Body mass index (BMI)
 - c. Pulmonary exacerbations number of exacerbations compared to the number of exacerbations prior to medication initiation
 - d. Liver function tests; AND
- 4. Dose does not exceed vanzacaftor 20 mg, tezacaftor 100 mg, and deutivacaftor 250 mg (two tablets of vanzacaftor 10 mg, tezacaftor 50 mg, and deutivacaftor 125 mg) per day.

ALYFTREKTM **Dosing**:

The recommended dosage of ALYFTREKTM in patients aged 6 years and older is listed in the table below. Please see the full prescribing information for dose modifications when ALYFTREKTM is used concomitantly with strong or moderate CYP3A inhibitors.

Age	Weight	Once Daily Oral Dosage
6 to Less than	Less than 40 kg	Three tablets of vanzacaftor 4 mg/tezacaftor 20 mg/
12 Years Old		deutivacaftor 50 mg
	Greater than or	Two tablets of vanzacaftor 10 mg/tezacaftor 50 mg/
	equal to 40 kg	deutivacaftor 125 mg
12 Years and	Any Weight	Two tablets of vanzacaftor 10 mg/tezacaftor 50 mg/
Older		deutivacaftor 125 mg

Formulations: Available as a fixed-dose combination containing vanzacaftor 4 mg, tezacaftor 20 mg, and deutivacaftor 50 mg and a fixed-dose combination containing vanzacaftor 10 mg, tezacaftor 50 mg, and deutivacaftor 125 mg.