

## Oral Cystic Fibrosis Modulators

### Goals:

- To ensure appropriate drug use and limit to patient populations in which they have demonstrated to be effective and safe.
- To monitor for clinical response for appropriate continuation of therapy.

### Length of Authorization:

- 90 days to 6 months

### Requires PA:

- Ivacaftor (Kalydeco®)
- Lumacaftor/Ivacaftor (Orkambi®)
- Tezacaftor/Ivacaftor (Symdeko®)

### Preferred Alternatives:

- Current PMPDP preferred drug list per OAR 410-121-0030 at [www.orpdl.org](http://www.orpdl.org)
- Searchable site for Oregon FFS Drug Class listed at [www.orpdl.org/drugs/](http://www.orpdl.org/drugs/)

Approval Criteria		
1. Is this a request for continuation of therapy previously approved by the FFS program (patient already on ivacaftor, lumacaftor/ivacaftor, or tezacaftor/ivacaftor)?	<b>Yes:</b> Go to <b>Renewal Criteria</b>	<b>No:</b> Go to #2
2. What diagnosis is being treated?	Record ICD10 code. Go to #3	
3. Is the request from a practitioner at an accredited Cystic Fibrosis Center or a pulmonologist?	<b>Yes:</b> Go to #4	<b>No:</b> Pass to RPh. Deny; medical appropriateness
4. How many exacerbations and/or hospitalizations in the past 12 months has the patient had?	Prescriber must provide documentation before approval. Document baseline value. Go to #5	
5. Is the request for ivacaftor?	<b>Yes:</b> Go to #6	<b>No:</b> Go to #10
6. What is the patient's baseline sweat chloride level?	Prescriber must provide documentation before approval. Document baseline value. Go to #7	
7. Does the patient have a diagnosis of cystic fibrosis and is 12 months of age or older?	<b>Yes:</b> Go to #8	<b>No:</b> Pass to RPh. Deny; medical appropriateness

## Approval Criteria

<p>8. Does the patient have a documented mutation in the CFTR gene that ivacaftor is FDA approved for (see below)?</p> <p>FDA approved CFTR mutations include: E56K, G178R, S549R K1060T, G1244E, P67L, E193K, G551D, A1067T, S1251N R74W, L206W, G551S, G1069R, S1255P, D110E, R347H, D579G, R1070Q, D1270N, D110H, R352Q, S945L, R1070W G1349D, R117C, A455E, S977F, F1074L, R117H, S549N, F1052V, D1152H 3849 + 10kbc -T, 2789 +5G&gt;A, 3272-26A-G, 711+3A-G, E831X</p>	<p><b>Yes:</b> Go to #17</p>	<p><b>No:</b> Go to #9</p> <p>If unknown, there needs to be a CF mutation test to detect the presence of the CFTR mutation prior to use.</p> <p>CF due to other CFTR gene mutations are not approved indications (including the F508del mutation).</p>
<p>9. Does the patient have a documented R117H mutation in the CFTR gene detected by a CF mutation test?</p>	<p><b>Yes:</b> Pass to RPh. Refer request to Medical Director for manual review and assessment of clinical severity of disease for approval.</p>	<p><b>No:</b> Pass to RPh. Deny; medical appropriateness.</p> <p>If unknown, there needs to be a CF mutation test to detect the presence of the CFTR mutation prior to use.</p> <p>CF due to other CFTR gene mutations are not approved indications (including the F508del mutation).</p>
<p>10. Is the request for lumacaftor/ivacaftor?</p>	<p><b>Yes:</b> Go to #11</p>	<p><b>No:</b> Go to #13</p>
<p>11. Does the patient have a diagnosis of cystic fibrosis and is 2 years of age or older?</p>	<p><b>Yes:</b> Go to #12</p>	<p><b>No:</b> Pass to RPh. Deny; medical appropriateness</p>

## Approval Criteria

<p>12. Does the patient have a documented homozygous Phe508del mutation in the CFTR gene detected by a CF mutation test?</p>	<p><b>Yes:</b> If the patient is younger than 12 years of age, refer case to <u>OHP Medical Director</u>; otherwise, Go to #17</p>	<p><b>No:</b> Pass to RPh. Deny; medical appropriateness</p> <p>If unknown, there needs to be a CF mutation test to detect the presence of the CFTR mutation prior to use.</p> <p>CF due to other CFTR gene mutations are not approved indications (including those who are heterozygous for the F508del mutation)</p>
<p>13. Is the request for tezacaftor/ivacaftor?</p>	<p><b>Yes:</b> Go to #14</p>	<p><b>No:</b> Pass to RPh. Deny; medical appropriateness</p>
<p>14. Does the patient have a diagnosis of cystic fibrosis and is 12 years of age or older?</p>	<p><b>Yes:</b> Go to #15</p>	<p><b>No:</b> Pass to RPh. Deny; medical appropriateness</p>
<p>15. Does the patient have a documented homozygous Phe508del mutation in the CFTR gene detected by a CF mutation test?</p>	<p><b>Yes:</b> Go to #17</p>	<p><b>No:</b> Go to #16</p> <p>If unknown, there needs to be a CF mutation test to detect the presence of the CFTR mutation prior to use.</p>
<p>16. Does the patient have at least one mutation that is responsive to tezacaftor/ivacaftor based on in vitro data and FDA labeling?</p> <p>Note: A list of CFTR gene mutations that produce CFTR protein and are responsive to tezacaftor/ivacaftor include: A455E, A1067T, D110E, D110H, D579G, D1152H, D1270N, E56K, E193K, E831X, F1052V, F1074L, K1060T, L206W, P67L, R74W, R1070W, R117C, R347H, R352Q, S945L, S977F, 711+3A→G, 2789+5G→A, 3272-26A→G, 3849+10kbC→T</p>	<p><b>Yes:</b> Go to #17</p>	<p><b>No:</b> Pass to RPh. Deny; medical appropriateness.</p> <p>If unknown, there needs to be a CF mutation test to detect the presence of the CFTR mutation prior to use.</p>

## Approval Criteria

<p>17. Is the patient on ALL the following drugs, or has had an adequate trial of each drug, unless contraindicated or not appropriate based on age &lt;6 years and normal lung function:</p> <ul style="list-style-type: none"> <li>• Dornase alfa; AND</li> <li>• Hypertonic saline; AND</li> <li>• Inhaled or oral antibiotics (if appropriate)?</li> </ul>	<p><b>Yes:</b> Go to #18</p>	<p><b>No:</b> Pass to RPh. Deny; medical appropriateness</p>
<p>18. Is the patient on concomitant therapy with a strong CYP3A4 inducer (see Table 1)?</p>	<p><b>Yes:</b> Pass to RPh. Deny; medical appropriateness</p>	<p><b>No:</b> Go to #19</p>
<p>19. What are the baseline liver function (AST/ALT) and bilirubin levels (within previous 3 months)?</p>	<p>Document labs. Go to #20</p> <p>If unknown, these labs need to be collected prior to approval.</p>	
<p>20. Is medication dosed appropriately based on age, weight, and co-administered drugs (see dosing and administration below)?</p>	<p><b>Yes:</b> Approve for 90 days.</p> <p>Note: Approve for 90 days to allow time for patient to have a sweat chloride test done after 30 days of treatment if on IVA (see <b>Renewal Criteria</b>).</p> <p>If approved, a referral will be made to case management by the Oregon Health Authority.</p>	<p><b>No:</b> Pass to RPh. Deny; medical appropriateness</p>

## Renewal Criteria

<p>1. Is this the first time the patient is requesting a renewal (after 90 days of initial approval)?</p>	<p><b>Yes:</b> Go to #2</p>	<p><b>No:</b> Go to #4</p>
<p>2. If prescription is for ivacaftor: Does the patient have a documented physiological response to therapy and evidence of adherence after 30 days of treatment, as defined by a sweat chloride test that has decreased by at least 20 mmol/L from baseline?</p>	<p><b>Yes:</b> Go to #7</p>	<p><b>No:</b> Go to #3 Consider patient's adherence to therapy and repeat test in 2 weeks to 45 days to allow for variability in test. If sodium chloride has still not decreased by 20 mmol/L, deny therapy for medical appropriateness</p>
<p>3. If the prescription is for lumacaftor/ivacaftor or tezacaftor/ivacaftor: Is there evidence of adherence and tolerance to therapy through pharmacy claims/refill history and provider assessment?</p>	<p><b>Yes:</b> Go to #7</p>	<p><b>No:</b> Pass to RPh; Deny (medical appropriateness)</p>
<p>4. Does the patient have documented response to therapy as defined as below :</p> <p>For patients age <math>\geq 6</math> years:</p> <ul style="list-style-type: none"> <li>• An improvement or lack of decline in lung function as measured by the FEV1 when the patient is clinically stable; OR</li> <li>• A reduction in the incidence of pulmonary exacerbations; OR</li> <li>• A significant improvement in BMI by 10% from baseline?</li> </ul> <p>For patients age 2-5 years (cannot complete lung function tests)</p> <ul style="list-style-type: none"> <li>• Significant improvement in BMI by 10% from baseline; OR</li> <li>• Improvement in exacerbation frequency or severity; OR</li> <li>• Sweat chloride test has decreased from baseline by 20 mmol/L from baseline?</li> </ul>	<p><b>Yes:</b> Go to #5</p>	<p><b>No:</b> Pass to RPh. Deny; medical appropriateness</p>

Renewal Criteria		
5. Has the patient been compliant with therapy, as determined by refill claims history?	<b>Yes:</b> Go to #6	<b>No:</b> Pass to RPh. Deny; medical appropriateness
6. Have liver function tests been appropriately monitored? What are the most recent liver function tests (AST, ALT, and bilirubin)?  Note: Monitoring LFTs is recommended every 3 months for the first year, followed by once a year.	Document. Go to #7  Note: Therapy should be interrupted in patients with AST or ALT >5x the upper limit of normal (ULN), or ALT or AST >3x ULN with bilirubin >2x ULN.	
7. Is the CFTR modulator dosed appropriately based on age, weight, and co-administered drugs (see dosing and administration below)?	<b>Yes:</b> Approve for additional 3 months (total of 6 months since start of therapy)	<b>No:</b> Pass to RPh. Deny; medical appropriateness

### Dosage and Administration:

#### Ivacaftor:

- Adults and pediatrics age ≥6 years: 150 mg orally every 12 hours with fat-containing foods
- Children age 1 to <6 years:
  - < 14 kg: 50 mg packet every 12 hours
  - ≥ 14 kg: 75 mg packet every 12 hours
- Hepatic Impairment
  - Moderate Impairment (Child-Pugh class B):
    - Age ≥6 years: one 150 mg tablet once daily
    - Age 1 to < 6 years with body weight < 14 kg: 50 mg packet once daily; with body weight ≥ 14 kg : 75 mg packet of granules once daily
  - Severe impairment (Child-Pugh class C): Use with caution at a dose of 1 tablet or 1 packet of oral granules once daily or less frequently.
- Dose adjustment with concomitant medications:

Table 1. Examples of CYP3A4 inhibitors and inducers.

Drug co-administered with IVA	Co-administered drug category	Recommended dosage adjustment for IVA
Ketoconazole Itraconazole Posaconazole Voriconazole Clarithromycin Telithromycin	CYP3A4 strong inhibitors	Reduce IVA dose to 1 tablet or 1 packet of oral granules <b>twice weekly</b> (one-seventh of normal initial dose)
Fluconazole Erythromycin Clofazimine	CYP3A4 moderate inhibitors	Reduce IVA dose to 1 tablet or 1 packet of oral granules <b>once daily</b> (half of normal dose)

Rifampin Rifabutin Phenobarbital Phenytoin Carbamazepine St. John's wort Grapefruit Juice	CYP3A4 strong inducers	Concurrent use is <b>NOT</b> recommended
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Lumacaftor/ivacaftor

- Adults and pediatrics age ≥12 years: 2 tablets (LUM 200 mg/IVA 125 mg) every 12 hours
- Pediatric patients age 6 through 11 years: 2 tablets (LUM 100mg/IVA 125 mg) every 12 hours
- Children age 2 to <6 years:
  - < 14 kg: 1 packet (LUM 100mg/IVA125mg) every 12 hours
  - ≥ 14 kg: 1 packet (LUM 150mg/IVA 188mg) every 12 hours
- Hepatic impairment
  - Moderate impairment (Child-Pugh class B):
    - Age ≥ 6 years: 2 tablets in the morning and 1 tablet in the evening
    - Age 2 to <6 years: 1 packet in the morning and 1 packet every other day in the evening
  - Severe impairment (Child-Pugh class C): Use with caution after weighing the risks and benefits of treatment.
    - Age ≥ 6 years: 1 tablet twice daily, or less
    - Age 2 to <6 years: 1 packet once daily, or less
- Dose adjustment with concomitant medications:
  - When initiating therapy in patients taking strong CYP3A inhibitors (see table above), reduce dose to 1 tablet daily for the first week of treatment. Following this period, continue with the recommended daily dose.

Tezacaftor/ivacaftor:

- Adults and pediatrics age ≥12 years: 1 tablet (TEZ 100 mg/IVA 150 mg) in the morning and IVA 150 mg in the evening
- Hepatic impairment
  - Moderate impairment (Child-Pugh class B):
    - 1 tablet (TEZ 100 mg/IVA 150 mg) in the morning. The evening IVA dose should not be administered.
  - Severe impairment (Child-Pugh class C):
    - 1 tablet (TEZ 100 mg/IVA 150 mg) in the morning (or less frequently). The evening IVA dose should not be administered.
- Dose adjustment with concomitant medications:
  - When initiating therapy in patients taking moderate CYP3A inhibitors (see table above), reduce dose to:
    - On day 1, TEZ 100/IVA 150 once daily in the morning, and on day 2, IVA 150 mg once daily in the morning; continue this dosing schedule.
  - When initiating therapy in patients taking strong CYP3A4 inhibitors (See table above), reduce dose to:
    - TEZ 100 mg/IVA 150 mg twice a week, administered 3 to 4 days apart. The evening dose of IVA 150 mg should not be administered.