HIGHLIGHTS OF PRESCRIBING INFORMATION
These highlights do not include all the information needed to use OSELTAMIVIR PHOSPHATE
FOR ORAL SUSPENSION safely and effectively. See full prescribing information for
OSELTAMIVIR PHOSPHATE FOR ORAL SUSPENSION.

OSELTAMIVIR PHOSPHATE for oral suspension Initial U.S. Approval: 1999

INDICATIONS AND USAGE

Oseltamivir phosphate for oral suspension is an influenza neuraminidase inhibitor (NAI) indicated for:

- Treatment of acute, uncomplicated influenza A and B in patients 2 weeks of age and older who have been symptomatic for no more than 48 hours. (1.1)
 Prophylaxis of influenza A and B in patients 1 year and older. (1.2)

- Not a substitute for annual influenza vaccination. (1.3) Consider available information on influenza drug susceptibility patterns and treatment effects when deciding whether to use. (1.3) Not recommended for patients with end-stage renal disease not undergoing dialysis. (1.3)

... DOSAGE AND ADMINISTRATION

Treatment of influenza

- Treatment of Influenza

 Adults and adolescents (13 years and older): 75 mg twice daily for 5 days (2,2)

 Pediatric patients 1 to 12 years of age: Based on weight twice daily for 5 days (2,2)

 Pediatric patients 2 weeks to less than 1 year of age: Based on weight twice daily for 5 days (2,2)

 Renally impaired adult patients (creatinine clearance >30 to 80 mL/min): Reduce to 30 mg twice daily

 Renally impaired adult patients (creatinine clearance >10 to 80 mL/min): Reduce to 30 mg one daily

 for 5 days (2,4)

 Renally impaired adult patients (creatinine clearance >10 to 80 mL/min): Reduce to 30 mg once daily

 for 5 days (2,4)

 ESRO patients on Intracallalysis: Reduce to 30 mg immediately and then 30 mg after every

 ESRO patients on CAPD: Reduce to a single 30 mg dose immediately (2,4)

- ESHU patients of Live A resource of a large may Prophylaxis of Influenza
 Adults and adolescents (13 years and older): 75 mg once daily for at least 10 days (2.3)
 Community outbreak: 75 mg once daily for up to 6 weels (2.3)
 Pediatric patients 1 to 12 years of age: Based on weight once daily for 10 days (2.3)
 Community outbreak: Based on weight once daily for up to 6 weels (2.3)
 Renally impaired adult patients (creatinine clearance > 30 to 60 mL/min): Reduce to 30 mg once daily
 Renally impaired adult patients (creatinine clearance > 30 to 60 mL/min): Reduce to 30 mg once even
- other day (2.4).

 © ESRD patients on hemodialysis: Reduce to 30 mg immediately and then 30 mg after alternate hemodialysis cycles for the recommended duration of prophylaxis (2.4).

 © ESRD patients on ACPD: Reduce to 30 mg immediately and then 30 mg once weekly for the recommended duration of prophylaxis (2.4).

 DOSAGE FORMS AND STRENGTHS.

For oral suspension: 360 mg oseltamivir base supplied as powder (constituted to a final concentration of 6 mg/mL) (3)

Patients with known serious hypersensitivity to oseltamivir or any of the components of oseltamivir phosphate (4)

.... WARNINGS AND PRECAUTIONS

- Serious skin/hypersensitivity reactions such as Stevens-Johnson Syndrome, toxic epidermal necrolysis
 and eylythema mulditorme: Discontinue oselatinuto phosphate and initiate appropriate treatment if
 Neuropsychiatric events: Patients with influenza, including those receiving osetamitir phosphate,
 particularly pediatric patients, may be at an increased risk of confusion or abnormal behavior early in
 their illness. Monitor for signs of adnormal behavior, (5.2)

Most common adverse reactions (>1% and more common than with placebo)

- Treatment studies Nausea, vomiting, headache. (6.1)
 Prophylaxis studies Nausea, vomiting, headache, pain. (6.1)

Avoid admittiscretors to cave where a medically indicated. (7)

See 17 for PATIENT COUNSELING INFORMATION and FDA-approved patient labeling.

Revised: 9/2022

FULL PRESCRIBING INFORMATION: CONTENTS* 1 INDICATIONS AND USAGE 1.1 Treatment of Influenza 1.2 Prophylaxis of Influenza

- 1. 2 Prophylaxs of Influenza
 1. 3 Limitations of Use
 2 DOSAGE AND ADMINISTRATION
 2. 1 Dosage and Administration Overview
 2. Recommended Dosage for Treatment of Influenza
 2. 3 Recommended Dosage for Prophylaxis of Influenza
 2. 4 Dosage in Patients with Renal Impairment
 2. 5 Preparation and Storage of Constituted Osetamivir Phosphate Oral Suspension
 2.6 Emergency Preparation of Oral Suspension from 75 mg Osetamivir Phosphate
 Capsules
- Capsules
 3 DOSAGE FORMS AND STRENGTHS

- 3 DOSAGE FORMS AND STRENG INS
 4 CONTRAINDICATIONS
 5 WARNINGS AND PRECAUTIONS
 5.1 Serious skin/Hypersensithity Reactions
 5.2 Neuropsychiatric Events
 5.3 Risk of Bacterial Infections
 5.4 Fructose Intolerance in Patients with Hereditary Fructose Intolerance

- 6.1 Clinical Trials Experience 6.2 Postmarketing Experienc 7 DRUG INTERACTIONS 7.1 Influenza Vaccines

- 7.2 Drugs Without Clinically Significant Drug Interaction with Oseltamivir Phosphate 8 USE IN SPECIFIC POPULATIONS

- 8.1 Pregnancy 8.2 Lactation 8.4 Pediatric Use 8.5 Geriatric Use 8.6 Renal Impairment 8.7 Henatic Impairment

- 8.7 Hepatic Impairment 8.8 Use in Patients with Chronic Conditions 8.9 Immunocompromised Patients 10 OVERDOSAGE
- 11 DESCRIPTION 12 CLINICAL PHARMACOLOGY
- 12.4 Microbiology
 13 NONCLINICAL TOXICOLOGY
 23 Corcinogenesis, Mutagenesis, Impairment of Fertility

- 13 NONCLINICAL TOXICOLOGY
 13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility
 14 CLINICAL STUDIES
 14.1 Treatment of Influenza
 14.2 Prophylaxis of Influenza
 16 HOW SUPPLIED/STORAGE AND HANDLING
 17 PATIENT COUNSELING INFORMATION

 * Sections or subsections omitted from the full prescribing information are not listed.

FULL PRESCRIBING INFORMATION

1 INDICATIONS AND USAGE

1.1 Treatment of Influenza

Oseltamivir phosphate for oral suspension is indicated for the treatment of acute, uncomplicated illness due to influenza A and B infection in patients 2 weeks of age and older who have been symptomatic for no more than 48 hours.

1.2 Prophylaxis of Influenza

Oseltamivir phosphate for oral suspension is indicated for the prophylaxis of influenza A and B in patients $\bf 1$ year and older.

1.3 Limitations of Use

- Oseltamivir phosphate for oral suspension is not a substitute for early influenza vaccination on an annual basis as recommended by the Centers for Disease Control and Prevention Advisory Committee on Immunization Practice
- might also diminish clinical benefit of antiviral drugs. Prescribers should consider available information on influenza drug susceptibility patterns and treatment effects

when deciding whether to use oseltamivir phosphate for oral suspension [see Microbiology (12.4)].

Microbiology (12.4)].

Oseltamivir phosphate for oral suspension is not recommended for patients with end-stage renal disease not undergoing dialysis [see Dosage and Administration (2.4) and Use in Specific Populations (8.6)].

2.1 Dosage and Administration Overview

Administer osekamivir phosphate for oral suspension for the treatment of influenza in patients 2 weeks of age or older [see Dosage and Administration (2.2)] or for prophylaxis of influenza in patients 1 year and older [see Dosage and Administration (2.3)] using:

O Sekamivir phosphate for oral suspension (supplied as a powder). This is the preferred formulation (6 mg per ml.) for patients who cannot swallow capsules. Prior to use, the supplied osetamivir phosphate for or real suspension powder must be constituted with water by the pharmackit to produce the oral suspension (see Dosage and Administration (2.5)).

An oral suspension may be taken with or without food; however, tolerability may be enhanced if oseltamivir phosphate for oral suspension is taken with food.

Adjust the oseltamivir phosphate for oral suspension dosage in patients with moderate or severe renal impairment [see Dosage and Administration (2.4)].

or severe renal impairment [see Dosage and Administration (2.4)]. For patients who cannot swallow capsules, osekamivir phosphate for oral suspension is the preferred formulation. When osekamivir phosphate for oral suspension is not available from wholesaler or the manufacturer, osekamivir phosphate capsules may be opened and mixed with sweetened liquids such as regular or sugar-free chocolate syrup, corn syrup, caramel topping, or light brown sugar (dissolved in water). During emergency stuations and when neither the oral suspension for the age-appropriate strengths of osekamivir phosphate capsules to mix with sweetened liquids are available, then a pharmacst thay prepare an emergency supply of oral suspension from osekamivir phosphate 75 mg capsules (see Dosage and Administration (2.6)).

2.2 Recommended Dosage for Treatment of Influenza

Initiate treatment with oseltamivir phosphate for oral suspension within 48 hours of influenza symptom onset.

Adults and Adolescents (13 Years of Age and Older)

The recommended oral dosage of oseltamivir phosphate for oral suspension for treatment of influenza in adults and adolescents 13 years and older is 75 mg twice daily (one 75 mg capsule or 12.5 mL of oral suspension twice daily) for 5 days.

Pediatric Patients (2 Weeks of Age Through 12 Years of Age)

Table 1 displays the recommended oral dosage of oseltamivir phosphate for oral suspension for treatment of influenza in pediatric patients 2 weeks of age through years of age and provides information about prescribing the capsule or the formula for oral suspension.

2.3 Recommended Dosage for Prophylaxis of Influenza

Initiate post-exposure prophylaxis with oseltamivir phosphate for oral suspension within 48 hours following close contact with an infected individual. Initiate seasonal prophylaxis with oseltamivir phosphate for oral suspension during a community outbreak.

Adults and Adolescents (13 Years of Age and Older)

Adults and Adolescents (13 Years of Age and Older)

The recommended dosage of oseltamivir phosphate for oral suspension for prophylaxis of influenza in adults and adolescents 13 years and older is 75 mg orally once daily (one 75 mg capsule or 12.5 mt. of oral suspension once daily) for at least 10 days following close contact with an infected individual and up to 6 weeks during a community outbreak. In immunocompromised patients, oselaminir phosphate for oral suspension may be confinited for up to 12 weeks feet by 8 in Specific Populations (6.9). The during may be continued for up to 12 weeks feet by 8 in Specific Populations (6.9). The during may be continued for the state of the stat

Pediatric Patients (1 Year to 12 Years of Age)

Table 1 display the recommended or all dosage of oseltamivir phosphate for oral suspension for prophylaxis of influenza in pediatric patients 1 year to 12 years of age based on body weight and provides information about prescribing the formulation for oral suspension. Prophylaxis in pediatric patients is recommended for 10 days following close contact with an infected individual and up to 6 weeks during a community outbreak (see Use in Specific Populations (8.4) and Clinical Studies (14.2)).

Table 1 Oseltamivir Phosphate Dosage Recommendations in Pediatric

| | Patient | s for Treatm | ent and Prophy | ylaxis of Influen | za |
|------------|-------------|---------------------------------------|---|--|--|
| - | Dosage | Prophylaxis Dosage for 10 days* | Volume of <u>Oral</u> <u>Suspension</u> (6 mg/mL) for <u>each Dose</u> [†] | Number of Bottles of <u>Oral</u> <u>Suspension</u> to Dispense | Number of <u>Capsules</u> to Dispense (Strength) [‡] |
| Pat ients | from 2 We | eks to less t | han 1 Year of | Age | |
| Any | 3 mg/kg | Not applicable | 0.5 mL/kg§ | 1 bottle | Not applicable |
| weight | twice daily | | _ | | |
| Pat ients | 1 to 12 Ye | ars of Age B | ased on Body \ | Weight | |
| 15 kg or | 30 mg | 30 mg once | 5 mL | 1 bottle | 10 capsules (30 |
| less | twice daily | daily | | | mg) |
| 15.1 kg to | 45 mg | 45 mg once | 7.5 mL | 2 bottles | 10 capsules (45 |
| 23 kg | twice daily | daily | | | mg) |
| 23.1 kg to | 60 mg | 60 mg once | 10 mL | 2 bottles | 20 capsules (30 |
| 40 kg | twice daily | daily | | | mg) |
| 40.1 kg or | 75 mg | 75 mg once | 12.5 mL | 3 bottles | 10 capsules (75 |
| more | twice daily | daily | | | mg) |

more | twice daily | daily | mg|

**The recommended duration for post-exposure prophylaxis is 10 days and the recommended
duration for community outbreak (seasonal/pre-exposure) prophylaxis is up to 6 weeks (or up to
12 weeks in immunocompromised patients). The amount supplied (e.g., number of bottles or
capsules) for seasonal prophylaxis may be greater than for post-exposure prophylaxis.
14 Use an oral dosing dispensing device that measures the appropriate volume in mit with the oral
suspension.

5 Osetamiker phosphate for oral suspension is the preferred formulation for patients who cannot

- swallow capsules.
 § For patients less than 1 year of age, provide an appropriate dosing device that can accurately measure and administer small volumes.

2.4 Dosage in Patients with Renal Impairment

Table 2 displays the dosage recommendations for the treatment and prophylaxis of influenza in adults with various stages of renal impairment (estimated creatinine clearance of less than or equal to 90 mL per minute). Dosage modifications are recommended in adults with an estimated creatinine clearance less than or equal to 60 mL per minute [see Use in Specific Populations (8.6) and Clinical Pharmacology (12.3)].

Table 2 Recommended Dosage Modifications for Treatment and Prophylaxis of Influenza in Adults with Renal Impairment or End Stage Renal Disease (ESRD) on Dialysis

| R enal Impairment (C reatinine Clearance) | R ecommended Treatment Regimen* | R ecommended Prophylaxis Regimen*† |
|--|---|--|
| Mild (>60 to 90 mL/minute) | 75 mg twice daily for 5 days | 75 mg once daily |
| Moderate (>30 to 60 mL/minute) | 30 mg twice daily for 5 days | 30 mg once daily |
| Severe (>10 to 30 mL/minute) | 30 mg once daily for 5 days | 30 mg every other day |
| ESRD Patients on Hemodialysis (≤10 mL/minute) | 30 mg immediately and then 30 mg after every hemodialysis cycle (treatment duration not to exceed 5 days) | 30 mg immediately and then 30 mg after alternate hemodialysis cycles |
| ESRD Patients on Continuous Ambulatory Peritoneal Dialysis† (≤10 mL/minute) | A single 30 mg dose administered immediately | 30 mg immediately and then 30 mg once weekly |
| ESRD Patients not on Dialysis | Oseltamivir phosphate is not recommended | Oseltamivir phosphate is not recommended |

* Oral suspension can be used for 30 mg dosing.
 + Data derived from studies in continuous ambulatory peritoneal dialysis (CAPD) patients

2.5 Preparation and Storage of Constituted Oseltamivir Phosphate Oral Suspension

Prior to dispensing to the patient, constitute oseltamivir phosphate for oral suspension (supplied as powder):

- a. Tap the closed bottle containing the supplied oseltamivir phosphate for oral suspension white to light brown powder several times to loosen the powder.
 b. Measure 55 m. of water in a graduated cylinder.
 c. Add the total amount of water for constitution to the bottle.
 d. Close bottle with child-resistant cap tightly and shake the closed bottle well for 15

- seconds.

 e. Label the bottle with instructions to "Shake Well Before Use".

 f. The constituted oral suspension contains 360 mg of oseltamivir base per 60 mL of volume (6 mg per mL) and is white to light brown, tutti-frutti-flavored). Use the constituted oral suspension with 17 days of preparation when stored under refrigeration, 2° to 8°C (36° to 46°F), or within 10 days if stored at controlled room temperature, 25°C (77°F). Write the expiration date of the constituted oral suspension on the bottle label.

 g. Ensure patients have an oral dosing dispenser that measures the appropriate volume in militers. Counsel patients on how to utilize the oral dosing dispenser and correctly measure the oral suspension as prescribed (see Tables 1 and 2).

2.6 Emergency Preparation of Oral Suspension from 75 mg Oseltamivir Phosphate Capsules

The following directions are provided for use only during emergency situations and when FDA-approved, commercially manufactured oseltamivir phosphate for oral suspension is not available from wholesalers or the manufacturer.

The following emergency preparation instructions will provide one patient with enough oseltamivir phosphate for a 5-day course of treatment of influenza or a 10-day course of prophylaxis of influenza:

Step #1: Determine the dosage of oseltamivir phosphate for the patient (see Dosage and Administration (2.2, 2.3, and 2.4)) then determine the total volume of oral suspension needed to be prepared (see Table 3).

Table 3 Emergency Preparation: Volume of Prepared Oral Suspension (6 mg per mL) Based Upon Oseltamivir Phosphate Dose

| Oseltamivir Phosphate Dose* | Total Volume to Prepare p e r Patient |
|-----------------------------|---------------------------------------|
| 15 mg or less | 37.5 mL |
| 30 mg | 75 mL |
| 45 mg | 100 mL |
| 60 mg | 125 mL |
| 75 mg | 150 mL |

If the oseltamivir phosphate dose is between the doses listed, use the greater listed dose to determine the total volume of prepared oral suspension.

Step.#2: Preparation must be performed with only one of the following vehicles (other vehicles have not been studied): Cherry Syrup (Humco®), Ora-Sweet® SF (sugar-free) (Paddock Laboratories), or simple syrup. Determine the number of capsules and the amount of water and vehicle needed to prepare the total volume (see Table 3) of prepared oral suspension (6 mg per mL) for a complete treatment or prophylaxis course (see Table 4).

Table 4 Emergency Preparation: Number of Oseltamivir Phosphate 75 mg Capsules and Amount of Water and Vehicle Needed to Prepare the Total Volume of a Prepared Oral Suspension (6 mg per mL)

| T otal Volume of Prepared | 37.5 mL | 75 mL | 100 mL | 125 mL | 150 mL |
|----------------------------|----------|----------|----------|----------|----------|
| Oral Suspension | | | | | |
| Number of Oseltamivir | 3 | 6 | 8 | 10 | 12 |
| Phosphate 75 mg | (225 mg) | (450 mg) | (600 mg) | (750 mg) | (900 mg) |
| Capsules (Total Strength)* | | | | _ | |
| A m ount of Water | 2.5 mL | 5 mL | 7 mL | 8 mL | 10 mL |
| Volume of Vehicle | 34.5 mL | 69 mL | 91 mL | 115 mL | 137 mL |
| Cherry Syrup (Humco®) OR | | | | | |
| Ora-Sweet® SF (Paddock | | | | | |
| Laboratories) OR | | | | | |
| simple syrup | | | | | |

* Includes overage to ensure all doses can be delivered

Step #3: Follow the instructions below for preparing the 75 mg oseltamivir phosphate capsules to produce the oral suspension (6 mg per mL):

- a. Place the specified amount of water into a polyethyleneterephthalate (PET) or glass bottle (see Table 4). Constitution in other bottle types is not recommended because there is no stability data with other bottle types.
- b. Carefully separate the capsule body and cap and pour the contents of the required number of oseltamivir phosphate 75 mg capsules into the PET or glass bottle.
- c. Gently swirl the suspension to ensure adequate wetting of the oseltamivir phosphate powder for at least 2 minutes.
- d. Slowly add the specified amount of vehicle to the bottle.
- e. Close the bottle using a child-resistant cap and shake well for 30 seconds to completely dissolve the active drug auspension. The active drug, oselamivir phosphate, dissolved drug inter resulting suspension. The active drug, oselamivir phosphate, read oselamivir phosphate capsules which are insoluble in these vehicles.
- f. Put an ancillary label on the bottle indicating "Shake Well Before Use."
- g. Instruct the parent or caregiver that any unused suspension remaining in the bottle following completion of therapy must be discarded by either affixing an ancillary label to the bottle or adding a statement to the pharmacy label instructions.
- h. Place a pharmacy label on the bottle that includes the patient's name, dosing instructions, drug name and any other required information to be in compliance wit State and Federal Pharmacy Regulations. Place an appropriate expiration date on the label according to storage conditions below.
- i. Include the recommended dosage on the pharmacy label as per Tables 1 and 2 [see Dosage and Administration (2.2, 2.3, and 2.4)].
- j. Store the prepared oral suspension in glass or PET bottles either:
- In a refrigerator [2° to 8°C (36° to 46°F)]: Stable for 5 weeks when stored in a refrigerator.
- At room temperature [25°C (77°F)]: Stable for 5 days when stored at room temperature.

3 DOSAGE FORMS AND STRENGTHS

Oseltamivir Phosphate for Oral Suspension: 6 mg per mL (final concentration when constituted)

White to light brown colored granular powder blend for constitution

4 CONTRAINDICATIONS

Oseltamivir phosphate is contraindicated in patients with known serious hypersensitivity to oseltamivir or any component of the product. Severe altergic reactions have included anaphylaxis and serious skin reactions including toxic epidermal necroyiss, Stevens-Johnson Syndrome, and erythema multiform [see Warnings and Precautions (5.1)].

5 WARNINGS AND PRECAUTIONS

5.1 Serious Skin/Hypersensitivity Reactions

5.1 Serious Skini/rypersensitivity reactions
Cases of anaphykais and serious skin reactions including toxic epidermal necrolysis,
Stevens-Johnson Syndrome, and erythema multiforme have been reported in postmarketing experience with osetamivir phosphate. Stop Osetamivir phosphate and institute appropriate treatment if an alterjicit he reaction occurs or is suspected. The use of osetamivir phosphate for oral suspension is contraindicated in patients with known serious hypersensitivity to osetamivir phosphate (see Contraindications (4) and Adverse Reactions (6.2)].

5.2 Neuropsychiatric Events

There have been postmarketing reports of delirum and abnormal behavior leading to injury, and in some cases resulting in fatal outcomes, in patients with influenza who were receiving oseitamivir phosphate [see Adverse Reactions (6.2)]. Because these events were reported voluntarily during clinical practice, estimates of frequency cannot be made but they appear to be uncommon based on oseitamivir phosphate usage data. These events were reported primarily among pediatric patients and often had an abrupt onset and rapid resolution. The contribution of oseitamivir phosphate to these events has not been established. Influenza can be associated with a variety of neurologic and abnormal behavior, in some cases resulting in fatal outcomes. These events may occur in the setting of encephalish or encephalopathy but can occur without obvious severe disease. Closely monitor oseitamivir phosphate-treated patients with influenza for signs of abnormal behavior. If neuropycychiatric symptoms occur, evaluate the risks and benefits of continuing oseitamivir phosphate for each patient. There have been postmarketing reports of delirium and abnormal behavior leading to

5.3 Risk of Bacterial Infections

There is no evidence for efficacy of oseltamivir phosphate in any illness caused by pathogens other than influenza viruses. Serious bacterial infections may begin with influenza-like symptoms or may coexist with or occur as complications during the course of influenza. Osetlamity phosphate has not been shown to prevent such complications. Prescribers should be alert to the potential for secondary bacterial infections and treat them as appropriate.

5.4 Fructose Intolerance in Patients with Hereditary Fructose Intolerance

Fructose can be harmful to patients with hereditary fructose intolerance. One dose of 75 mg osetlamivir phosphate for oral suspension delivers 2 grams of sorbibol. This is above the daily maximum limit of sorbibol for patients with hereditary fructose intolerance and may cause dyspepsia and diarrhea.

The following serious adverse reactions are discussed below and elsewhere in the labeling:

- Serious skin and hypersensitivity reactions [see Warnings and Precautions (5.1)]
 Neuropsychiatric events [see Warnings and Precautions (5.2)]

6.1 Clinical Trials Experience

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be directly compared to rates in the clinical trials of another drug and may not reflect the rates observed in practice.

Adverse Reactions from Treatment and Prophylaxis Trials in Adult and Adolescent Subjects (13 years of age and older)

The overall safety profile of oseltamivir phosphate is based on data from 2,646 adult and adolescent subjects that received the recommended dosage of 75 mg orally twice daily for 5 days for treatment of influenza and 1,943 adult and adolescent subjects that received the recommended dosage of 75 mg orally once daily for up to 6 weeks for prophylaxis of influenza in clinical.

The most common adverse reactions in the pooled treatment and pooled prophylaxis trials in adults and adolescents are displayed in Table 5. The majority of these adverse reactions were reported on a single occasion, occurred on either the first to second treatment day and resolved spontaneously within 1 to 2 days. This summary includes otherwise healthy adults/adolescents and subjects art is first subjects at higher risk of developing complications associated with influenza, e.g., etlerly patients and patients with chronic cardiac or respiratory disease), in general, the safety profile in the subjects "at risk" was qualitatively similar to that in otherwise healthy adults/adolescents.

Table 5 Adverse Reactions Occurring in ≥1% of Adults and Adolescents (13 years of age and older) in Treatment and Prophylaxis Trials*

| System Organ Class | Treatment Trials | | Prophylaxis Trials | |
|----------------------------|------------------|-----------------------|--|-----------------------|
| Adverse Reaction | | Placebo (n = 1977) | Oseltamivir Phosphate 75 mg once daily (n = 1943) | Placebo (n = 1586) |
| Gastrointestinal Disorders | | | | |
| Nausea | 10% | 6% | 8% | 4% |
| Vomiting | 8% | 3% | 2% | 1% |
| Nervous System Disorders | | | | |
| Headache | 2% | 1% | 17% | 16% |
| General Disorders | | | | |
| Pain | <1% | <1% | 4% | 3% |

compared to placebo-treated subjects in either the treatment or prophylaxis trials

Adverse Reactions from Treatment and Prophylaxis Trials in Pediatric Subjects (1 year to 12 years of age)

A total of 1,481 pediatric subjects (including otherwise healthy pediatric subjects aged 1 A rota or 1.481 pediatric subjects (including otherwise healthy pediatric subjects aged to 12 years and asthmatic pediatric subjects aged 6 to 12 years) participated in clinical trials of oseltamivir phosphate for the treatment of influenza. A total of 859 pediatric subjects received treatment with oseltamivir phosphate for or als suspension either at a 2 mg per kgt wice daily for 5 days or weight-band dosing. Vomiting was the only adverse reaction reported at a frequency of ${\,\cong\,}1\%$ in subjects receiving oseltamivir phosphate (16%) compared to placebo (8%).

Amongst the 148 pediatric subjects aged 1 year to 12 years who received oseltamivir phosphate at doses of 30 to 60 mg once daily for 10 days in a post-exposure prophylaxis study in household contacts (n = 99), and in a separate 6-week seasonal influenza prophylaxis safety study (n = 49), vomiting was the most frequent adverse reaction (8% on oseltamivir phosphate versus 2% in the no prophylaxis groups).

Adverse Reactions from Treatment Trials in Pediatric Subjects (2 weeks to less than 1 year of age)

less than 1 year of age)

Assessment of adverse reactions in pediatric subjects 2 weeks to less than 1 year of age was based on two open-label studies that included safety data on 135 influenza-infected subjects 2 weeks to less than 1 year of age (including premature infants at least 36 weeks post conceptional age) exposed to osetamivir phosphate at doses ranging from 2 to 3.5 mg per kg of the formulation for oral suspension twice daily orally for 5 dosy. The safety profile of osetamivir phosphate was similar across the age range studied, with vomiting (9%), diarrhea (7%) and diaper rash (7%) being the most frequently reported adverse reactions, and was generally comparable to that observed in older pediatric and adult subjects.

Adverse Reactions from the Prophylaxis Trial in Immunocompromised Subjects

In a 12-week seasonal prophylaxis study in 475 immunocompromised subjects, including 18 pediatric subjects 1 year to 12 years of age, the safety profile in the 238 subjects receiving osetaminivi phosphate 75 mg once daily was consistent with that previously observed in other osetaminir phosphate prophylaxis clinical trials [see Clinical Studies (14-2)].

6.2 Postmarketing Experience

The following adverse reactions have been identified during post-approval use of osetamivir phosphate. Because these reactions are reported voluntarily from a population of uncertain size, it is not possible to reliably estimate their frequency or establish a causal relationship to osetamivir phosphate exposure.

General Disorders and Administration Site Conditions

Swelling of the face or tongue, allergy, anaphylactic/anaphylactoid reactions,

Skin and Subcutaneous Tissue Disorders

Rash, dermatitis, urticaria, eczema, toxic epidermal necrolysis, Stevens-Johnson Syndrome, erythema multiforme [see Warnings and Precautions (5.1)]

Gastrointestinal Disorders

Gastrointestinal bleeding, hemorrhagic colitis

Cardiac Disorders

Arrhythmia

Hepatobiliary Disorders

Hepatitis, abnormal liver function tests Nervous System Disorders

Seizure

Metabolism and Nutrition Disorders

Aggravation of diabetes Psychiatric Disorders

Annormal behavior, delirium, including symptoms such as hallucinations, agitation, anxiety, altered level of consciousness, confusion, nightmares, delusions [see Warnings and Precautions (5.2)]

7 DRUG INTERACTIONS

7.1 Influenza Vaccines

Live Attenuated Influenza Vaccine

The concurrent use of oseltamivir phosphate with live attenuated influenza vaccine (LAIV) intranasal has not been evaluated. However, because of the potential for oseltamivir phosphate to inhibit replication of five vaccine virus and possibly reduce the efficacy of LAIV, avoid administration of LAIV within 2 weeks before or 48 hours after oseltamivir phosphate administration, unless medically indicated.

Inactivated Influenza Vaccine

Inactivated influenza vaccine can be administered at any time relative to use of oseltamivir phosphate.

7.2 Drugs Without Clinically Significant Drug Interaction with Oseltamivir Phosphate

No dose adjustments are needed for either oseltamivir or the concomitant drug when coadministering oseltamivir with amoxicilin, acetaminophen, aspirin, cimetidine, antacids (magnesium and aluminum Hydroxides and calcium carbonates), rimantadine, amantadine, or warfarin [see Clinical Pharmacology (12.3)].

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Risk Summary

Risk Summary

There are no adequate and well-controlled studies with oseltamivir phosphate in pregnant women to inform a drug-associated risk of adverse developmental outcomes. Available published epidemiological data suggest that oseltamivir phosphate, taken in an timester, is not associated with an increased risk of brith defects. However, these studies individually are limited by small sample sizes, use of different comparison groups, and some lacked information on dose, which preclude a definitive assessment of the risk (see Data and Clinical Pharmacology (12.3)). In animal reproduction studies with oseltamivir, no adverse developmental effects were observed at clinically relevant exposures (see Data).

The background risk of major birth defects and miscarriage for the indicated populations is unknown. All pregnancies have a background risk of birth defect, loss, or other adverse outcomes. In the U.S. general population, the estimated background risk of major birth defects and miscarriage is 2 to 4% and 15 to 20%, respectively.

Clinical Considerations:

Disease-Associated Maternal and/or Embryo/Fetal Risk:

Pregnant women are at higher risk of severe complications from influenza, which may lead to adverse pregnancy and/or fetal outcomes including maternal death, still births, birth defects, pretern delivery, low birth weight and small for gestational age.

Data:

Published prospective and retrospective observational studies of more than 5,000 women exposed to osekamivir phosphate during pregnancy, including more than 1,000 women exposed in the first trimester, suggest that the observed rate of congenital mailrormations was not increased above the rate in the general comparison population, regardless of when therapy was administered during the gestational period. However, individually, none of these studies had adequate sample sizes and some lacked information on dose, which preclude a definitive assessment of the risk.

Animal Data:

Osetamivir was administered orally during organogenesis to pregnant rats (at 50, 250, or 1500 mg/kg/day on gestation days 6 to 17) and rabbits (at 50, 150, or 500 mg/kg/day on gestation days 6 to 18). In rats, embryo-fetal effects consisting of an increased incidence of minor skeletal malformations were observed at a maternaly toxic dose (1500 mg/kg/day), resulting in systemic drug exposures (based on AUC modes (1500 mg/kg/day). The systemic ratio of the maximum recommend to when dose (1500 mg/kg/day) resulting in systemic (75 mg twice a day). In the rabbit study, embryo-fetal effects consisting of an increased incidence of minor skeletal abnormalities and variants of the maternally toxic doses (1510 mg/kg/day) resulting in systemic exposures after observed at maternally toxic doses (1510 mg/kg/day) resulting in systemic exposures at the MHHD of osetamivir phosphate.

In prenatal and postnatal development studies in rats, oseltamivir was administered orally (at 50, 250, 500, or 1500 mg/kg/day) from organogenesis through late gestation, delivery, and lactation (gestation day 6 to postpartum/lactation day 20). Prolonged parturition duration and reduced offspring viability were observed at a maternally toxic dose (1500 mg/kg/day). No adverse maternal or offspring effects were observed. doses \$500 mg/kg/day). No adverse maternal or offspring effects were observed doses \$500 mg/kg/day, resulting in systemic drug exposures (based on AUC for oseltamivir carboxylate) 44 times human exposures at the MRHD of oseltamivir phosphate.

8.2 Lactation

Risk Summary

Based on limited published data, oseltamivir and oseltamivir carboxylate have been shown to be present in human milk at low levels considered unlikely to lead to toxicity in the breastfed infant. Postmarketing experience has not reported any information to suggest serious adverse effects of oseltamivir exposure via breast milk in infants. It is not known if oseltamivir affects human milk production. The developmental and heabth benefits of breastfeding should be considered along with the mother's clinical need for oseltamivir phosphate and any potential adverse effects on the breastfed child from the drug or from the underlying maternal condition.

8.4 Pediatric Use

Treatment of Influenza

The safety and efficacy of oseltamivir phosphate for the treatment of influenza in pediatric patients 2 weeks old to 17 years of age has been established [see Dosage and Administration (2.2), Clinical Pharmacology (12.3), and Clinical Studies (14.1)] and is been don:

- a3 to 17 years of age: Safety and efficacy in adolescent patients 13 to 17 years of age was supported by adequate and well-controlled trials in adults and adolescents and younger pediatric patients and safety data in adolescents treated with osetamin'r phosphate in a study of treatment and prophylaxis.
 1 year to 12 years of age: Safety and efficacy in pediatric patients 1 year to 12 years of age was supported by results of one double-blind, placebo-controlled trial in 452 pediatric patients with influenza in whom osetamin'r phosphate 2 mg per kg twice daily or placebo was administered with 48 hours of symptom onset [see Clinical Studies (14.11)]. Additionals afety information was provided in a double-blind, placebo-controlled trial in pediatric patients 6 to 12 years of age with known asthma. Efficacy could not be established in pediatric patients with asthma.
 2 weeks to less than 1 year of age: Safety and efficacy in pediatric patients 2 weeks to less than 1 year of age. In these two trials, the osetamin'r plasma concentrations in these subjects were similar to or higher than the osetamin'r plasma concentrations oserved in older pediatric administration of the pediatric patients and two open-label trials of osetamin'r plasma concentrations in these subjects were similar to or higher than the osetamin'r plasma concentrations of served in older pediatric administration of the pediatric patients and two trials. The osetamin'r plasma concentrations of served in older pediatric administration of the pediatric patients and the pediatric patients and the pediatric patients and the pediatric patients and pediatric patients.

The safety and efficacy of oseltamivir phosphate for treatment of influenza in pediatric patients less than 2 weeks of age have not been established.

Prophylaxis of Influenza

The safety and efficacy of oseltamivir phosphate for the prophylaxis of influenza in pediatric patients 1 year to 17 years old has been established *(see Dosage and Administration (2.3), Clinical Pharmacology (12.3), and Clinical Studies (14.2)* and is based on:

- saeet oil:

 13 to 17 years of age: Prophylaxis in adolescent patients 13 to 17 years of age is supported by one randomized, placebo-controlled post-exposure household prophylaxis trial of osetlamivir phosphate 75 mg taken orally once daily for 7 days in household contacts including 207 adolescents [see Clinical Studies (14-2)].

 1 years to 12 years of age: Osetlamivir phosphate for prophylaxis in pediatric patients 1 year to 12 years of age is supported by one randomized, open-label, post-exposure household prophylaxis trial including pediatric subjects 1 year to 12 years of age who received 30 to 60 mg of osetlamivir phosphate for oral suspension (supplied as powder) taken orally once daily for 10 days [see Clinical Studies (14-2)]. Additional safety information was provided in a 6-week seasonal prophylaxis (community outbreak) safety study in 49 patients 1 year to 12 years of age.

The safety and efficacy of oseltamivir phosphate for prophylaxis of influenza have not been established for pediatric patients less than $1\ \text{year}$ of age.

8.5 Geriatric Use

Treatment of Influenza

Of the 4,765 adults in clinical trials of oselamivir phosphate for the treatment of influenza, 948 (20%) were 65 years and older, while 329 (7%) were 75 years and older, in three double-blind, placebo-controlled trials in the treatment of influenza in patients at least 65 years old, that enrolled 741 subjects (374 received placebo and 362 received oselamivir phosphate), no overall differences in safety or effectiveness were observed between these subjects and younger subjects, and other reported clinical experience has not identified differences in responses between the elderly and younger subjects (see Clinical Studies (14.1)).

Prophylaxis of Influenza

Of the 4,603 adults in clinical trials of oseltamivir phosphate for the prophylaxis of Of the 4,603 adults in clinical trial ocstaminary phosphate for the prophylaxs of influenza, 1,046 (23%) were 75 years and older, while 719 (16%) were 75 years and older. In a randomized, placebo-controlled trial in eleterly residents of nursing homewho took oselaminy phosphate for up to 42 days for the prophylaxs of influenza (oselaminar phosphate = 276, placebo = 272), no overall differences in safety of effectiveness where the event between these subjects and younger subjects, and other reported clinical experience has not identified differences in responses between the eleterly and younger subjects and younger subject be earlied of studies (14.2.1).

8.6 Renal Impairment Patients with renal impairment had higher blood levels of oseltamivir carboxylate

Fracetis with retaining an internal right group of the compared to possible with compared to possible six with normal renal function which may surpress either fisk of osekamity between executions. Therefore, devise a digital recommended for patients with a serum creations clearance between each group and to multiminate and for patients with a serum creations clearance less (ESRD) undergoing routine hemodules and for continuous personnel dislayes treatment (see Dosage and Administration and Compared Continuous personnel dislayes the creatment (see Dosage and Administration and Continuous personnel dislayes the c

(2.4)]. Oseltamivir phosphate is not recommended for patients with ESRD not undergoing dialysis [see Indications and Usage (1.3) and Clinical Pharmacology (12.3)].

No dosage adjustment is required in patients with mild to moderate hepatic impairment. The safety and pharmacokinetics in patients with severe hepatic impairment have not been evaluated [see Clinical Pharmacology (12.3)].

8.8 Use in Patients with Chronic Conditions

Efficacy of oseltamivir phosphate in the treatment of influenza in patients with chronic cardiac disease and/or respiratory disease was evaluated in one randomized, placebo-controlled clinical trial. Efficacy in this population, as measured by time to alleviation of all symptoms, was not established, but no new safety signals were identified [see Clinical Studies (14.1)].

No clinical trial data are available regarding treatment of influenza in patients with any medical condition sufficiently severe or unstable to be considered at imminent risk of requiring hospitalization.

8.9 Immunocompromised Patients

Efficacy of oseltamivir phosphate for the treatment or prophylaxis of influenza has not been established in immunocompromised patients [see Clinical Studies (14.2)]. Safety of oseltamivir phosphate has been demonstrated for up to 12 weeks for prophylaxis of influenza in immunocompromised patients [see Adverse Reactions (6.1)].

10 OVERDOSAGE

Reports of overdoses with oseltamivir phosphate have been received from clinical trials and during postmarketing experience. In the majority of cases reporting overdose, no adverse reactions were reported. Adverse reactions reported following overdose were similar in nature to those observed with therapeutic doses of oseltamivir phosphate (see Adverse Reactions (6)1.

11 DESCRIPTION

Oseltamivir phosphate, an influenza neuraminidase inhibitor (NAI), is available as:

A powder for oral suspension, which when constituted with water as directed contains 6 mg per mL oseltamivir base.

In addition to the active ingredient, the powder for oral suspension contains monosodium citrate, saccharin sodium, sodium benzoate, sorbitol, titanium dioxide, tutti-frutti flavor and xanthan gum.

Oseltamivir phosphate is a white crystalline solid with the chemical name (3R,4R,5S)-4-accylamino-5-amino 31-ethylpropoxy)-1-cyclohecene-1-carboxylic acid, ethyl ester, phosphate (1:1). The chemical formula is C₃(H₂B₃V₂O₄ (free base). The molecular weight is 312.4 for oseltamivir free base and 410.40 for oseltamivir phosphate salt. The structural formula is as follows:

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Oseltamivir is an antiviral drug with activity against influenza virus [see Microbiology (12.4)].

12.3 Pharmacokinetics

Absorption and Bioavailability

Oseltamivir is absorbed from the gastrointestinal tract after oral administration of oseltamivir phosphate and is extensively converted predominantly by hepatic esterases to oseltamivir carboxylate. All least 75% of an oral dose reaches the systemic circulation as oseltamivir carboxylate and less than 5% of the oral dose reaches the systemic circulation as oseltamivir (see Table 6).

Table 6 Mean (% CV) Pharmacokinetic Parameters of Oseltamivir and Oseltamivir Carboxylate Following Multiple Dosing of 75 mg Capsules Twice Daily (n=20)

| Parameter | Oseltamivir | Oseltamivir Carboxylate | | | | |
|--------------------------------|-------------|-------------------------|--|--|--|--|
| C _{max} (ng/mL) | 65 (26) | 348 (18) | | | | |
| AUC _{0-12h} (ng·h/mL) | 112 (25) | 2719 (20) | | | | |

Plasma concentrations of oselkamivir carboxylate are proportional to doses up to 500 mg given twice daily (about 6.7 times the maximum recommended oselkamivir phosphate dosage) [see Dosage and Administration (2)].

Coadministration with food had no significant effect on the peak plasma concentration (551 ng/mL under fasted conditions and 441 ng/mL under fed conditions) and the area under the plasma concentration time curve (6218 ng-l/mL under fasted conditions and 6069 ng h/mL under fed conditions) of osekamivir carboxylate.

The volume of distribution (V_{ss}) of oseltamivir carboxylate, following intravenous administration in 24 subjects (oseltamivir phosphate is not available as an IV formulation), ranged between 23 and 26 liters.

The binding of osettamivir carboxylate to human plasma protein is low (3%). The binding of osettamivir to human plasma protein is 42%, which is insufficient to cause significant displacement-based drug interactions.

Flimination

Absorbed oseltamivir is primarily (>90%) eliminated by conversion to the active metabolite, oseltamivir carboxylate. Plasma concentrations of oseltamivir declined with a half-life of 10 al hours in most subjects after oral administration. Oseltamivir carboxylate is not further metabolized and is eliminated unchanged in urine. Plasma concentrations of oseltamivir carboxylate declined with a half-life of 6 to 10 hours in most subjects after oral administration.

Oseltamivir is extensively converted to the active metabolite, oseltamivir carboxylate, by esterases located predominantly in the liver. Oseltamivir carboxylate is not further metabolized. Neither oseltamivir nor oseltamivir carboxylate is a substrate for, or inhibitor of, cytochrome P450 Soforms.

Excretion:

Osetamivir carboxylate is eliminated entirely (>99%) by renal excretion. Renal clearance (18.8 L/h) exceeds glomerular filtration rate (7.5 L/h), indicating that tubular secretion (via organic anion transporter) occurs in addition to glomerular filtration. Less than 20% of an oral radiolabeled dose is eliminated in feces.

Specific Populations

Renal Impairment:

Administration of 100 mg of oseltamivir phosphate twice daily (about 1.3 times the maximum recommended dosage) for 5 days to subjects with various degrees of renal impairment showed that exposure to oseltamivir carboxylate is inversely proportional to declining renal function.

Decuming tenal unicuon.

Population -derived pharmacokinetic parameters were determined for patients with varying degrees of renal function including ESRD patients on hemodialysis. Median simulated exposures of oselamivir carboxysite for recommended treatment and prophylaxis regimens are provided in Table 7. The pharmacokinetics of oselamivir have not been studied in ESRD patients not undergoing dialysis [see Indications and Usage (1.3), and Use in Specific Populations (8.6)].

| Renal Function/ Impairment | Normal Creatinine Clearance 90 to 140 mL/min | Mild Creatinine Clearance 60 to 90 mL/min | Moderate Creatinine Clearance 30 to 60 mL/min | Severe Creatinine Clearance 10 to 30 mL/min | ESRD Creatinine Clearance |
|---------------------------------|--|---|---|---|-----------------------------------|
| Renal Function/ Impairment | (n=57) | (n=45) | (n=13) | (n=11) | <10 mL/min on Hemodialysis (n=24) |
| Recommended Treatment | Regimens | | | | |
| PK exposure parameter | 75 mg twice daily | 75 mg twice daily | 30 mg twice daily | 30 mg once daily | 30 mg every HD cycle |
| C _{min} (ng/mL) | 145 | 253 | 180 | 219 | 221 |
| C _{max} (ng/mL) | 298 | 464 | 306 | 477 | 1170 |
| AUC ₄₈ (ng • h/mL)* | 11224 | 18476 | 12008 | 16818 | 23200 |
| Recommended Prophylaxis | Regimens | | | | |
| PK exposure parameter | 75 mg once daily | 75 mg once daily | 30 mg once daily | 30 mg every other day | 30 mg alternate HD cycle |
| C _{min} (ng/mL) | 39 | 62 | 57 | 70 | 42 |
| C _{max} (ng/mL) | 213 | 311 | 209 | 377 | 903 |
| AUC ₄₈ (ng • hr/mL)* | 5294 | 8336 | 6262 | 9317 | 11200 |
| * AUC normalized to 48 hours. | | • | | • | |

In continuous ambulatory peritoneal dialysis (CAPD) patients, the peak concentration of oselamivir carboxylate following a single 30 mg dose of oselamivir or once weekly oselamivir was approximately 3-fold higher than in patients with normal renal function who received 75 mg twice daily. The plasma concentration of oselamivir carboxylate on Day 5 (147 ngml) following a single 30 mg dose in CAPD patients is similar to the predicted C_{min} (160 ng/mL) in patients with normal renal function following 75 mg twice daily. Administration of 30 mg once weekly to CAPD patients resulted in plasma concentrations of oselamivir carboxylate at the 168-hour blood sample of 63 ng/mL, which were comparable to the C_{min} in patients with normal renal function receiving the approved regimen of 75 mg once daily (40 ng/mL).

Hepatic Impairment:

In clinical studies oseltamivir carboxylate exposure was not altered in subjects with mild or moderate hepatic impairment [see Use in Specific Populations (8.7)].

Pregnant Women:

A pooled population pharmacokinetic analysis indicates that the oseltamivir phosphate dosage regimen resulted in lower exposure to the active metabolite in pregnant women (n=99) compared to non-pregnant women (n=39). However, this predicted exposure is expected to have activity against susceptible influenza virus strains and there are insufficient pharmacokinetics and safety data to recommend a dose adjustment for pregnant women (see Use in Specific Populations (6.1)).

Pediatric Subjects (1 year to 12 Years of Age):

The pharmacokinetics of osetamivir and osetlamivir carboxylate have been evaluated in a single-dose pharmacokinetic study in pediatric subjects aged 5 to 16 years (n=18) and in a small number of pediatric subjects aged 3 to 12 years (n=26) enrolled in a clinical trial. Younger pediatric subjects cleared both the prodrug and the active metabolite faster than adult subjects resulting in a lower exposure for a given molky dose. For osetamivir carboxylate, apparent total clearance decreases linearly with increasing age (up to 12 years). The pharmacokinetics of osetamivir neglectivic subjects over 12 years of age are similar to those in adult subjects (see Use in Specific Populations (8.4)).

Pediatric Subjects (2 Weeks to Less Than 1 Year of Age):

The pharmac chinetics of oselamivia and oselamiviar carboxylate have been evaluated in two open-label studies of pediatric subjects less than one year of age (n=122) infected with influenza. Apparent clearance of the active metabolite decreases with decreasing age in subjects less than 1 year of age; however the oselamivir and oselamivir carboxylate exposure following a 3 mg/kg dose in subjects under 1 year of age is expected to be within the observed exposures in adults and adolescents receiving 75 mg twice daily and 150 mg twice daily [see Use in Specific Populations (8.4)].

Geriatric Patients:

Exposure to osekamiwir carboxylate at steady-state was 25 to 35% higher in geriatric subjects (age range 65 to 78 years) compared to young adults given comparable doses of osekamiwir. Half-lives observed in the geriatric subjects were simal to those seen in young adults. Based on drug exposure and tolerability, dose adjustments are not required for geriatric patients for either treatment or prophylaxis [see Use in Specific Populations (8.51)].

Drug Interaction Studies

Oselamivir is extensively converted to oseltamivir carboxylate by esterases, located predominantly in the liver. Drug interactions involving competition for esterases have not been extensively reported in iterature. Low protein binding of oseltamivir and oseltamivir carboxylate suggests that the probability of drug displacement interactions is low.

In vitro studies demonstrate that neither oseltamivir nor oseltamivir carboxylate is a good substrate for P450 mixed-function oxidases or for glucuronyl transferases.

Coadministration of probenecid results in an approximate two-fold increase in exposure to oselamivir carboxylate due to a decrease in active anionit tubular secretion in the kidney. However, due to the safety margin of oselamivir carboxylate, no dose adjustments are required when coadministering with probenecid.

No clinically relevant pharmacokinetic interactions have been observed when coadmistering oseltamivir with amoxicilin, acetaminophen, aspirin, cimetitine, antacids (magnesium and aluminum hydroxides and calcium carbonates), rimantadine, amantadine, or warfarin.

Mechanism of Action

Oseltamivir phosphate is an ethyl ester prodrug requiring ester hydrolysis for conversion to the active form, oseltamivir carboxylate. Oseltamivir carboxylate is an inhibitor of influenza virus neuraminidase affecting release of viral particles. The median IC5₉ values of oseltamivir against influenza AH1N1, influenza AH3N2, and influenza BIC5₁₀ values of oseltamivir against influenza AH1N1, influenza AH3N2, and influenza BIC10 values of oseltamivir against influenza AH1N1, oseltamivirus objective values of oseltamivirus values of oseltamivirus values values of oseltamivirus values value

Antiviral Activity

Antiviral Activity

The antiviral activity of oseltamivir carboxylate against laboratory strains and clinical isolates of influenza virus was determined in cell culture. The concentrations of oseltamivir carboxylate required for inhibition of influenza virus in cell culture were highly variable depending on the assay method used and the virus tested. The 50% and 90% effective concentrations (EC₅₀ and EC₅₀) were in the range of 0.0008 micromolar to greater than 35 micromolar and 0.004 micromolar to greater than 100 micromolar, respectively (1 micromolar=0.284 microgram per mL). The relationship between the antiviral activity in cell culture, inhibitory activity in the neuramindsae assay, and the inhibition of influenza virus replication in humans has not been established.

Resistance

Cell Culture Studies:

Influenza A virus isolates with reduced susceptibility to oseltamivir carboxylate have been recovered by serial passage of virus in cell culture in the presence of increasing concentrations of oseltamivir carboxylate. Reduced susceptibility of influenza virus to inhibition by oseltamivir carboxylate may be conferred by amino acid substitutions in the viral neuramindase and/or hemagglithin proteins.

Clinical Studies:

Reduced susceptibility isolates have been obtained during treatment with oseitamivir and from sampling during community surveillance studies. Changes in the viral neuraminidase that have been associated with reduced susceptibility to oseitamivir carboxylate are summarized in Table 8. The clinical impact of this reduced susceptibility is unknown.

Hemagglutini (HA) substitutions selected in cell culture and associated with reduced susceptibility to oseltamivir include (influenza virus subtype-specific numbering) A11T, K173E, and R453M in H3N2: and H99Q in influenza B virus (Yamagata lineage). In some cases, HA substitutions were selected in conjunction with known NA resistance substitutions and may contribute to reduced susceptibility to oseltamivir; however, the impact of HA substitutions on antiviral activity of oseltamivir in humans is unknown and likely to be strain-dependent.

Table 8 Neuraminidase Amino Acid Substitutions Associated with Reduced Susceptibility to Oseltamivir

A m ino Acid Substitution'
Influenza A NI. (NI numbering in brackets)
117V (117V), E119V (E119V), R152K (R152K), Y155H (Y155H), F173V (F174V),
D198CM (D199CM), 1222K(R7IV) (1232K(R7IV), S246CM (S247CM), G248R+1266V
(G249R+1267V), H274V (H275V), N294S (N295S), Q312R+1427T (Q313R+1427T),
N325K (N325K), R371K (R368K)

Influenza A N2 E41G, E119I/V, D151V, I222L/V, Q226H, SASG245-248 deletion, S247P, R292K, N294S

Participant (1974) (197

Selection of influenza A viruses resistant to oseltamivir can occur at higher frequencies in children. Oseltamivir treatment-associated resistance in pediatric treatment studies has been detected at frequencies of 27 to 37% and 3 to 18% (3/11 to 7/19 and 1/34 to 9/50 post-treatment isolates, respectively) for influenza Alri NIN virus and influenza A/H3N2 virus, respectively

The frequency of resistance selection to oseltamivir and the prevalence of such resistant virus vary seasonally and geographically

vrus vary seasonaly and geographicasy. Circulating seasonal influenza strains expressing neuraminidase resistance-associated substitutions have been observed in individuals who have not received seelamivir treatment. The seelamivir resistance-associated substitution HZ75Y was found in more than 99% of US-circulating 2008 H1N1 influenza virus loaltes. The 2009 H1N1 influenza virus ("swine fill") was almost uniformly susceptible to seelamivir: however, the frequency of circulating resistant variants can change from season to season. Prescribers should consider available information from the CDC on influenza virus drug susceptibility patterns and treatment effects when deciding whether to use oselamivir phososhate.

Cross-resistance

Cross-resistance between oseltamivir and zanamivir has been observed in neuraminidase biochemical assays. The H275Y (N1 numbering) or N2945 (N2 numbering) oseltamivir resistance-associated substitutions observed in the N1 neuraminidase subtype, and the E119V or N2945 oseltamivir resistance-associated neuramindase subtype, and the E119V or N2945 osebarnivir resistance-associated substitutions observed in the N2 subtype (N2 numbering), are associated with reduced susceptibility to osebarnivir but not zanamivir. The Q136K and K150T zanamivir resistance-associated substitutions observed in N1 neuraminidase, or the S250G zanamivir resistance-associated substitutions observed in influenza B virus neuraminidase, confer reduced susceptibility to zanamivir but not osebarnivir. The R292K osebarnivir resistance-associated substitution observed in N2, and the I227T, D198E/N, A871K, or G4025 osebarnivir resistance-associated substitutions observed in influenza B virus neuraminidase, confer reduced susceptibility to both osebarnivir and zanamivir. These examples do not represent an exhaustive list of cross resistance-associated substitutions and prescribers should consider available information from the CDC on influenza drug susceptibility patterns and treatment effects when deciding whether to use osebarnivir phosphate.

No single amino acid substitution has been identified that could confer cross-resistance between the neuraminidase inhibitor class (oseltamivir, zanamivir) and the M2 ion channel inhibitor class (amantadine, rimantadine). However, a virus may carry a neuraminidase inhibitor-associated substitution in neuraminidase and an M2 ion channel inhibitor associated substitution in M2 and may therefore be resistant to both classes of inhibitors. The clinical relevance of phenotypic cross-resistance evaluations has not been established.

Immune Response

No influenza vaccine/oseltamivir interaction study has been conducted. In studies of naturally acquired and experimental influenza, treatment with oseltamivir phosphate did not impair normal humoral antibody response to infection.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

In 2-year carcinogenicity studies in mice and rats given daily oral doses of the prodrug osetamive prosphate up to 400 mg/kg and 500 mg/kg, respectively, the prodrug and the active from setamivir carboyate induced no statistically significant increases in tumors over control. The mean maximum daily exposures to the prodrug in mice and rats were approximately 130- and 320-fodi, respectively, greater than those in humans at the recommended clinical dose based on AUC comparisons. The respective specified margins of the active osetamivir carboxistic were 15- and 50-fodi.

or the exposures to the active osetamive carboxylate were 15- and 50-fold. Osetamivit was found to be non-mulagenic in the Ames test and the human hymphocyte chromosome assay with and without enzymatic activation and negative in the mouse introducelus test. It was found to be positive in a Syrian Hamster Embryo (SHE) cell transformation liest. Osetamivi carboxylate was non-mulagenic in the Ames test and the LS178Y mouse lymphoma assay with and without enzymatic activation and negative in the SHE cell transformation test.

In a fertility and early embryonic development study in rats, doses of oseltamivir at 50, 250, and 1500 mg/kg/day were administered to females for 2 weeks before mating, during mating and until day 6 of pregnancy. Males were dosed for 4 weeks before mating, during mating, and for 2 weeks after mating. There were no effects on fertility, mating performance or early embryonic development at any dose level. The highest dose in this study was approximately 115 times the human systemic exposure (AUC_{0-24h}) of oseltamivir carboxylate that occurs after administration of the maximum recommended human dose.

14 CLINICAL STUDIES

14.1 Treatment of Influenza

Adults

Two randomized, placebo-controlled, double-blind clinical trials of oseltamivir phosphate were conducted in adults between 18 and 65 years old, one in the U.S. and one outside U.S. and fever of at least 100°F, accompanied by at least one respiratory symptoms (ocuph, nasal symptoms, or sore throat) and at least one systemic symptom (myalgia, chilk/sweats, mabiae, fatigue, or headache), and influenza virus was known to be circulating in the community. Subjects were randomized to receive oral oselaminir phosphate or placeb for 5 days. All enrolled subjects were allowed to take fever- reducing medications.

Of 1,355 subjects enrolled in these two trials, 849 (63%) subjects were influenza-infected (median age 34 years; 52% male; 90% Caucasian; 31% smokers). Of the 849 influenza-infected subjects, 95% were infected with influenza A, 3% with influenza B, and 2% with influenza of unknown type.

2% with influenza of unknown type.

Study medication was started within 40 hours of onset of symptoms and administered twice daily for 5 days. Subjects were required to self-assess the influenza-associated symptoms finals congestion, sore throat, cough, aches, fatigue, headaches, and chillis/sweats) twice daily as "none." mild. "moderate," or "severe". Time to improvement was calculated from the time of treatment initiation to the time when all symptoms were assessed as "none" or "mild". In both trials, there was a 1.3-day reduction in the median time to improvement in influenza-infected subjects who received osetamivir phosphate 75 mg twice a day for 5 days compared to subjects who received placebo. Subgroup analyses by gender showed no differences in the treatment effect of osetamivir phosphate in men and women.

. h. the treatment of influenza, no increased efficacy was demonstrated in subjects who received higher doses of oseltamivir phosphate.

Adolescents and Adults with Chronic Cardiac or Respiratory Disease

A double-blind, placebo-controlled, multicenter trial was unable to demonstrate efficacy of osetamivir phosphate (75 mg kwc daily for 5 days) in the treatment of influenza adult and adolescent subjects (13 years or older) with chronic cardiac (excluding chronic idopathic hypertension) or respiratory diseases, as measured by time to alleviation of all symptoms. However, in patients treated with osetamivir phosphate there was a more rapid cessation of febrile liness. No difference in the incidence of influenza complications was observed between the treatment and placebo groups in this population.

Geriatric Subjects

Three double-blind placebo-controlled treatment trials were conducted in subjects who were at least 65 years of age in three consecutive seasons. The enrollment criteria were similar to that of adult trials with the exception of fever being defined as higher than 97.5°F. Of 741 subjects enrolled, 476 (65%) subjects were influenza-infected; of these, 95% were influenza-infected; of these, 95% were influenza type A and 5% with influenza type B.

In the pooled analysis, there was a 1-day reduction in the median time to improvement influenza-infected subjects who received oseltamivir phosphate 75 mg twice daily for 5 days compared to those who received placebo (p=NS) (see Use in Specific Populations (8.5)). Some seasonal variability was noted in the clinical efficacy outcomes.

Pediatric Subjects (1 year to 12 years of age)

One double-blind placebo-controlled treatment trial was conducted in pediatric subject aged 1 year to 12 years (median age 5 years) who had fever (at least 100°F) plus one respiratory symptom (cough or coryza) when filtenza virus was known to be circulating in the community. Of 698 subjects enrolled in this trial, 452 (65%) were influenza-infected (50% male; 68% Caucasian). Of the 452 influenza-infected subjects, 67% were infected with influenza A and 33% with influenza B.

Efficacy in this trial was determined by the time to alleviation or resolution of influenza signs and symptoms, measured by a composte endpoint that required the following four individual conditions be met:) alleviation of cough, ii) alleviation of coryas, iii) resolution of fever, and iv) parental opinion of a return to normal health and activity. Osetamivir phosphate treatment of 2 mp per kg twice daily, started within 48 hours of onset of symptoms, reduced the total composite time to freedom from liness by 1.5 days compared to placebo. Subgroup analyses by gender showed no differences in the treatment effect of osetamivir phosphate in male and female pediatric subjects.

Pediatric Subjects (2 weeks to less than 1 year of age)

Two open-bled trisk evaluated the safety and pharmacokinetics of oseltamivir and oseltamivir carboxylate in influenza-infected pediatric subjects 2 weeks to less than 1 year of age (including premature infants at least 36 weeks post conceptional age). Subjects received oseltamivir phosphate at doses ranging from 2 to 3.5 mg per kg tw daily for 5 days depending on subject age. These clinical trials were not designed to evaluate clinical efficacy or virologic response.

Of the 136 subjects under the age of 1 year enrolled and dosed in the trials, the majority of the subjects were male (55%), white (79%), non-Hispanic (74%), full term (76%) and infected with influenza A (80%). Pharmacokinetic data indicated that a dose of 3 mg per

kg twice daily in pediatric subjects 2 weeks to less than 1 year of age provided oseltamivir phosphate concentrations similar to or higher than those observed in older pediatric subjects and adults receiving the approved dose and provided the basis for approval [see Adverse Reactions (6.1) and Use in Specific Populations (8.4)].

14.2 Prophylaxis of Influenza

Adult and Adolescent Subjects (13 years of age and older)

The efficacy of osekamivir phosphate in preventing naturally occurring influenza illness has been demonstrated in three seasonal prophylaxis (community outbreak) clinical trials and one post-exposure prophylaxis trial in household contacts. The efficacy endpoint for all of these trials was the incidence of laboratory-confirmed clinical influenza defined as meeting all the following criteria (all signs and symptoms must have been recorded within 24 hours):

- oral temperature greater than or equal to 99.0°F (37.2°C).

- una usripur auure greater than or equal to 99.0°F (37.2°C), at least one respiratory symptom (cough, sore throat, nasal congestion), at least one constitutional symptom (aches and pain, fatigue, headache, chills/sweats), and either a postive virus isolation or a four-fold increase in virus antibody titers from baseline.

In a pooled analysis of two seasonal prophylaxis trials in healthy unvaccinated adults (aged 18 to 65 years), oseltamivir phosphate 75 mg once dally taken for 42 days dut a community outbreak reduced the incidence of laboratory-confirmed clinical influence from 5% (25/519) for the placebo group to 1% (6/520) for the oseltamivir phosphate

in the seasonal (community outbreak) prophylaxis trial in elderly residents of skilled nursing homes, about 80%, 43%, and 14% of the subjects were vaccinated, had cardiac disorders, and had chronic airway obstructive disorders, respectively. In this trial, subjects were randomized to oselaminir phosphate 75 mg once daily or placebo taken or ally for 42 days. The incidence of laboratory-confirmed clinical influenza was 4% (12/277) in the placebo- treated subjects compared to less than 1% (1/276) in the oselaminir phosphate -treated subjects.

oseralming prospirate—treated subjects. In the post-exposure prophylays trial in household contacts (aged 13 years or older) of an index influenza case, osertamivir phosphate 75 mg once daily or placebo taken orally was administered within 48 hours of onset of symptoms in the index case and continued for 7 days (index cases did not receive osertamivir phosphate treatment). The incidence of laboratory-confirmed clinical influenza was 12% (24/200) in the placebo-treated subjects compared to 1% (2/205) in the osertamivir phosphate -treated subjects.

Pediatric Subjects (1 year to 12 years of age)

The efficacy of oselamivir phosphate in preventing naturally occurring influenza illness was demonstrated in a randomized, open-label post-exposure prophylaxic trial in household contacts that included pediatric subjects aged 1 year on 12 years, both as index cases and as family contacts. All index cases in this trial received oselamiving phosphate for oral supersion 30 to 60 mg taken orally once daily for 10 days. The effect of the properties of the properties of the object of the properties of the object of the o household. Labor following criteria:

- oral temperature at least 100F (37.8C), cough and/or coryza recorded within 48 hours, and either a positive virus isolation or a four-fold or greater increase in virus antibody titers from baseline or at illness visits.

Among household contacts 1 year to 12 years of age not already shedding virus at baseline, the incidence of laboratory-confirmed clinical influenza was lower in the gro who received osekamivir phosphate prophylaxis [3% (3/95)] compared to the group who did not receive osekamivir phosphate prophylaxis [17% (18/106)].

Immunocompromised Subjects

Immunocompromised Subjects

A double-blind, placebo-controlled trial was conducted for seasonal prophylaxis of influenza in 475 immunocompromised subjects (including 18 pediatric subjects 1 year to 12 years of age) who had received solid organ (n=388; liver, kidney, liver and kidney) or hematopoietic stem cell transplants (n=87). Median time since transplant for solid organ transplant receipients was 1,105 days for the placebo group and 1,379 days for the oselarmivir phosphate group. Median time since transplant for hematopoietic stem cell transplant receipients was 424 days for the placebo group and 367 days for the oselarmivir phosphate group. Approximately 40% of subjects received influenza vaccine prior to entering the study. The primary efficacy endpoint was the incidence of confirmed clinical influenza, defined as oral temperature higher than 99.0 f (37.2 C) plus cough and/or coryza, all recorded within 24 hours, plus either a positive virus culture or a four-fold increase in virus antibody itters from baseline. Subjects received treatment with oseltamivir phosphate 75 ing or placebo once daily by mouth for 12 weeks. The with 3% (57.237) in the oseltamivir phosphate group; this difference was not statisticated with 3% (57.237) in the oseltamivir phosphate group; this difference was not statisticated with 3% (57.237) in the oseltamivir phosphate group; this difference was not statisticated with 3% (57.237) in the oseltamivir phosphate group; this difference was not statisticated with 3% (57.237) in the oseltamivir phosphate group; this difference was not statisticated with 3% (57.237) in the oseltamivir phosphate group; this difference was not statisticated with 3% (57.237) in the oseltamivir phosphate group; this difference was not statisticated with 3% (57.237) in the oseltamivir phosphate group; this difference was not statisticated with 3% (57.237) in the oseltamivir phosphate group; this difference was not statisticated with 3% (57.237) in the oseltamivir phosphate group; this difference was not stati

16 HOW SUPPLIED/STORAGE AND HANDLING

Oseltamivir Phosphate for Oral Suspension (Supplied as Powder)

Supplied as a white to light brown colored granular powder in a glass bottle. After constitution, the powder blend produces a white to light brown tuttl-frutti-flavored oral suspension. After constitution with 55 mL of water, each bottle delivers a usable volume of 60 mL of oral suspension equivalent to 360 mg osefamiwir base (6 mg/mL) (see Dosage and Administration (2.5)) (INDC 71205-489-60).

Storage

Store dry powder at 25°C (77°F); excursions permitted between 15° to 30°C (59° to 86°F) [See USP Controlled Room Temperature].

Store constituted oral suspension under refrigeration for up to 17 days at 2° to 8°C (36° to 46°F). Do not freeze. Alternatively, store constituted oral suspension for up to days at 25°c (77°F); excursion permitted to 15° to 30°C (59° to 86°F) [See USP Controlled Room Temperature].

17 PATIENT COUNSELING INFORMATION

Advise the patient to read the FDA-approved patient labeling (Patient Information and Instructions for Use).

Serious Skin/Hypersensitivity Reactions

Advise patients and/or caregivers of the risk of severe allergic reactions (including anaphylaxis) or serious skin reactions. Instruct patients and/or caregiver to stop osetlamivir phosphate and seek immediate medical attention if an allergic-like reaction occurs or is suspected [see Warnings and Precautions (5.1)].

Neuropsychiatric Events

Advise patients and/or caregivers of the risk of neuropsychiatric events in oseltamivir phosphate-treated patients with influenza and instruct patients to contact their physicia if they experience signs of abnormal behavior while receiving oseltamivir phosphate (see Warnings and Precautions (5.2)].

Important Dosing Information

Instruct patients to begin treatment with oseltamivir phosphate as soon as possible from the first appearance of flu symptoms, within 48 hours of onset of symptoms. Similarly, instruct patients to start taking oseltamivir phosphate for prevention as soon as possible after exposure [see Dosage and Administration (2)]. Instruct patients to take any missed doses as soon as they remember, except if it is near the next scheduled dose (within 2 hours), and then continue to take oseltamivir phosphate at the usual times.

Instruct patients that osetamivir phosphate is not a substitute for receiving an annual flu vaccination. Patients should continue receiving an annual flu vaccination according to guidelines on immunization practices. Because of the potential for osetamivir phosphate to inhibit replication of live attenuated influence vaccine (LANV) and possibly reduce efficacy of LAIV, avoid administration of LAIV within 2 weeks or 48 hours after osetamivir phosphate administration, unless medically necessary (see Drug Interactions (7.1)).

Fructose Intolerance

Inform patients with hereditary fructose intolerance that one dose of 75 mg oseltamivir phosphate oral suspension (supplied as powder) delivers 2 grams of sorbitol. Inform patients with hereditary fructose intolerance that this is above the daily maximum limit of sorbitol and may cause dyspepsia and dierrhea [see Warnings and Precautions (5.4)].

The brands listed are trademarks of their respective owners and are not trademarks of Lupin Pharmaceuticals, Inc. The makers of these brands are not affiliated with and do not endorse Lupin Pharmaceuticals, Inc. or its products.

Lupin Pharmaceuticals, Inc. Baltimore, Maryland 21202

United States

Manufactured by:

Lupin Limited

Aurangabad - 431 210

Relabeled by

Proficient Rx LP

Thousand Oaks, CA 91320 Revised: June 2020

PATIENT INFORMATION

Oseltamivir Phosphate for Oral Suspension

(OH-sel-TAM-i-vir FOS-fate)

What is oseltamivir phosphate for oral suspension?

Oseltamivir phosphate for oral suspension is a prescription medicine used to:

- treat the flu (influenza) in people 2 weeks of age and older who have had flu symptoms for no more than two days.
 prevent the flu in people who are 1 year of age and older.

It is not known if oseltamivir phosphate for oral suspension is:

- effective in people who start treatment after 2 days of developing flu symptoms. effective for the treatment of the flu in people with long-time (chronic) heart problems or breathing problems. effective for the treatment or prevention of flu in people who have weakened immune systems (immunocompromised) safe and effective for the treatment of the flu in children less than 2 weeks of age. safe and effective for the treatment of the flu in children less than 1 year of age.

Oseltamivir phosphate for oral suspension does not treat or prevent illness that is caused by infections other than the influenza virus.

Oseltamivir phosphate for oral suspension does not prevent bacterial infections that may happen with the flu.

Oseltamivir phosphate for oral suspension is not recommended for people with end-stage renal disease (ESRD) who are not receiving dialysis.

Soseitamis of the state of receiving a flu vaccination. Talk to your healthcare provider about when you should receive an annual flu vaccination.

Who should not take oseltamivir phosphate for oral suspension?

Do not take oseltamivir phosphate for oral suspension if you are allergic to oseltamivir phosphate or any of the ingredients in oseltamivir phosphate for oral suspension. See the end of this leaflet for a complete list of ingredients in oseltamivir phosphate for oral suspension.

What should I tell my healthcare provider before taking oseltamiving phosphate for oral suspension?

Before you take oseltamivir phosphate for oral suspension, tell your healthcare provider if you:

- have problems swallowing oseltamivir phosphate capsules.
- have kindey problems have kindey problems have a history of frustee (fruit sugar) intolerance. Oselkamivir phosphate for oral suspension contains sorbitol and may cause stomach upset and diarrhea in people who are fructose intolerant. have any other medical conditions
- are pregnant or plan to become pregnant. Available information indicate that oselamivir phosphate for oral suspension does not increase the risk of birth defects.

 are breastfeeding or plan to breastfeed. Oseltamivir phosphate can pass into
- breast milk in small amounts.

Tell your healthcare provider about all the medicines you take, including prescription or over-the-counter medicines, vitamins, and herbal supplements.

Know the medicines you take. Keep a list of them to show your healthcare provider and pharmacist when you get a new medicine.

How should I take oseltamivir phosphate for oral suspens

- Take oseltamivir phosphate for oral suspension exactly as your healthcare
- The continuer principle for oral suspension exactly as your healthcare provider tells you to.

 Take osekamivir phosphate for oral suspension with food or without food. There is less chance of stomach upset if you take osekamivir phosphate for oral suspension with food.
- If you miss a dose of oseltamivir phosphate for oral suspension, take it as soon as
- If you miss a dose of osetamivir phosphate for oral suspension, take it as soon as you remember, if it is 2 hours or less before your next dose, do not take the missed dose. Take your next dose, dose of osetamivir phosphate for oral suspension at your scheduled time. Do not take 2 doses at the same time. If osetamivir phosphate for oral suspension is not available or you cannot swallow osetamivir phosphate capsules, your healthcare provider or pharmacist may instruct you to open osetamivir phosphate capsules and mix the capsules contents with sweetened fliguids such as chocolate syrup (regular or sugar-free), corn syrup, caramel topping, or light brown sugar (dissolved in water).

If your healthcare provider or pharmacist has instructed you to take oseltamivir phosphate for oral suspension, read the detailed instructions for Use at the end of this leaflet. Ask your pharmacist if you have any questions.

What are the possible side effects of oseltamivir phosphate for oral suspension?

Oseltamivir phosphate for oral suspension may cause serious side effects, including

- Serious skin and allergic reactions. Oseltamivir phosphate for oral suspension
 can cause serious skin and allergic reactions. Stop taking oseltamivir phosphate for
 oral suspension and get medical help right away if you get any of the following
- symptoms: skin rash or hives your skin blisters and peels blisters or sores in your mouth
- itching

- Riching of your face, eyes, lips, tongue, or throat trouble breathing chest pain or tightness (Change in behavior. People, especially children, who have the flu can develop nervous system problems and abnormal behavior that can lead to death. During treatment with oseltamivin phosphate for or pal suspension, tell your healthcare provider right away if you or your child have confusion, speech problems, shaky movements, seizures, or start hearing voices or seeing things that are not really there (hallucinations).

The most common side effects of oseltamivir phosphate for oral suspension when used for treatment of the flu include nausea, vomiting, and headache.

The most common side effect of oseltamivir phosphate for oral suspension when used for prevention of the flu include nausea, vomiting, headache, and pain.

Tell your healthcare provider if you have any side effect that bothers you or that does not go away.

These are not all of the possible side effects of oseltamivir phosphate for oral suspension.

Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088. You may also report side effect to Lupin Pharmaceuticals, Inc. at 1-800-399-2561.

How should I store oseltamivir phosphate for oral suspension?

- Store osetamivir phosphate for oral suspension in the refrigerator for up to 17 days between 36°F to 46°F (2°C to 8°C). Do not freeze. Store osetamivir phosphate for oral suspension for up to 10 days at room temperature between 68°F to 77°F (20°C to 25°C). Safely throw away any urused osetamivir phosphate for oral suspension that is out of date or no longer needed.

Keep oseltamivir phosphate for oral suspension and all medicines out of the reach of children.

General information about the safe and effective use of oseltamivir phosphate for oral suspension.

Medicines are sometimes prescribed for purposes other than those listed in a Patient Information leaflet. Do not use oseltamivir phosphate for oral suspension for a condition for which it was not prescribed. Do not give oseltamivir phosphate for oral suspension

What are the ingredients in oseltamivir phosphate for oral suspension?

Active ingredient: oseltamivir phosphate

Inactive ingredients:

Oseltamivir phosphate for oral suspension: monosodium citrate, saccharin sodium, sodium benzoate, sorbitol, titanium dioxide, tutti-frutti flavor and xanthan gum. This Patient Information has been approved by the U.S. Food and Drug Administration.

Manufactured for:

Lupin Pharmaceuticals, Inc.

Baltimore, Maryland 21202

United States

Manufactured by:

Lupin Limited

Aurangabad - 431 210

India

Relabeled by:

Proficient Rx LP

Thousand Oaks, CA 91320

Revised: June 2020

INSTRUCTIONS FOR USE

Oseltamivir Phosphate for Oral Suspension

(OH-sel-TAM-i-vir FOS-fate)

How do I give a dose of oseltamivir phosphate for oral suspension?

Step 1. each use. Shake the oseltamivir phosphate for oral suspension bottle well before

Step 3. Measure the oral suspension with an appropriate oral dosing dispenser to be sure you get the correct dose. Contact your pharmacist if you do not have an appropriate oral dosing dispenser.

Step 4. Give the full contents of oral dosing dispenser directly into the mouth.

Close the bottle with the child-resistant bottle cap after each use. Step 6. Rinse oral dosing dispenser under running tap water and allow to air dry after each use.

How do I mix the contents of oseltamivir phosphate capsules with sweetened liquids, if directed by my healthcare provider or pharmacist?

You will need:

- the prescribed dose of oseltamivir phosphate capsules
 a small bowl
 sweetened liquid, such as chocolate syrup (regular or sugar-free), corn syrup, caramel topping, or light brown sugar (dissolved in water)

Step 1. Open the contents of the prescribed dose of oseltamivir phosphate capsules into a small bowl.

Step 2. Add a small amount of the sweetened liquid to the capsule contents.

Step 3. Stir the mixture and give the entire dose of oseltamivir phosphate capsules.

This Instructions for Use has been approved by the U.S. Food and Drug Administration. Manufactured for:

Lupin Pharmaceuticals, Inc.

Baltimore. Maryland 21202

United States

Manufactured by: Lupin Limited

Aurangabad - 431 210

India

Relabeled by: Proficient Rx LP

Thousand Oaks, CA 91320

Revised: June 2020 264977

ID#:

PACKAGE LABEL.PRINCIPAL DISPLAY PANEL

NDC 71205-489-60

Oseltamivir Phosphate for Oral Suspension

6 mg/mL

Rx only

Glass Bottle of 60 mL



| Product Informa | tion | | | | | | |
|--|---|--|------------|-------|----------------------------------|------|-----------------|
| Product Type HUMAN PRESCRIPTION DRUG (Source) | | | | | NDC:71205-489(NDC:68180- 678) | | |
| Route of Administra | | | | | | | |
| Active Ingredient | /Active | Moiety | | | | | |
| | Ing | redient Name | | | Basis Streno | | Strength |
| OSELTAMIVIR PHOSPH UNII:K6106LV5Q8) | ATE (UNII: | 4A3O49NGEZ) (OSELTAMI | VIR ACID - | 09 | ELTAMIVIR | ACID | 6 mg in 1 mL |
| | | | | | | | |
| Inactive Ingredie | nts | | | | | | |
| Inactive Ingredie | nts | Ingredient Name | | | | St | renath |
| | | • | | | | St | rength |
| MONOSODIUM CITRAT | re (UNII: 68 | 538UP9SE) | | | | St | rength |
| MONOSODIUM CITRAT SACCHARIN SODIUM (| FE (UNII: 68 UNII: SB8Z | 8538UP9SE) UX40TY) | | | | St | rength |
| MONOSODIUM CITRAT SACCHARIN SODIUM (U SODIUM BENZOATE (U | FE (UNII: 68 UNII: SB8Z INII: OJ2458 | 8538UP9SE) UX40TY) | | | | St | rength |
| MONOSODIUM CITRAT SACCHARIN SODIUM (I SODIUM BENZOATE (U SORBITOL (UNII: 506T6) TITANIUM DIOXIDE (UN | FE (UNII: 68 UNII: SB8Z INII: OJ2458 0A25R) NII: 15FIX9\ | 1538UP9SE) UX40TY) FESEU) (2JP) | | | | St | rength |
| Inactive Ingredie | FE (UNII: 68 UNII: SB8Z INII: OJ2458 0A25R) NII: 15FIX9\ | 1538UP9SE) UX40TY) FESEU) (2JP) | | | | St | rength |
| MONOSODIUM CITRAT SACCHARIN SODIUM (I SODIUM BENZOATE (U SORBITOL (UNII: 506T6) TITANIUM DIOXIDE (UN | FE (UNII: 68 UNII: SB8Z INII: OJ2458 0A25R) NII: 15FIX9\ | 1538UP9SE) UX40TY) FESEU) (2JP) | | | | St | rength |
| MONOSODIUM CITRAT SACCHARIN SODIUM (I SODIUM BENZOATE (I) SORBITOL (UNII: 506T6 TITANIUM DIOXIDE (UN XANTHAN GUM (UNII: T | FE (UNII: 68 UNII: SB8Z INII: OJ245I 0A25R) NII: 15FIX9A TV12P4NEI | 1538UP9SE) UX40TY) FESEU) (2JP) | | | | St | rength |
| MONOSODIUM CITRAT SACCHARIN SODIUM (I SODIUM BENZOATE (U SORBITOL (UNIE: 506766 TITANIUM DIOXIDE (UN XANTHAN GUM (UNIE: T | FE (UNII: 68 UNII: SB8Z UNII: OJ245I OA25R) NII: 15FIX9V TV12P4NEI | 1538UP9SE) UX40TY) FESEU) (2JP) | į. | icore | | St | rength |
| MONOSODIUM CITRAT SACCHARIN SODIUM (II SODIUM BENZOATE (II SORBITOL (UNIE: SOOTE) TITTANIUM DIOXIDE (UN XANTHAN GUM (UNIE: T | FE (UNII: 68 UNII: SB8Z UNII: OJ245I OA25R) NII: 15FIX9V TV12P4NEI | 1538UP9SE) UX40TY) FESEU) (2JP) | | icore | | St | rength |
| MONOSODIUM CITRAT SACCHARIN SODIUM (II SODIUM BENZOATE (II SORBITOL (UNIE: SO676 TITANIUM DIOXIDE (UNIE: TITANIUM DIOXIDE (UNIE: TITANIUM CUNIE: TITANIUM CUNIE: TITANIUM COLOR XANTHAN GUM (UNIE: TITANIUM COLOR Product Characte Color Shape | FE (UNII: 68 UNII: SB8Z UNII: OJ245I 0A25R) NII: 15FIX9V TV12P4NEI | 1538UP9SE) UX40TY) FESEU) ZJP) E) te to light brown) | 9 | ize | nt Code | St | rength |

| P | ackaging | | | |
|---|------------------------------------|---|-------------------------|-----------------------|
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:71205- 489-60 | 1 in 1 CARTON | 10/19/2020 | |
| 1 | | 60 mL in 1 BOTTLE, GLASS; Type 0: Not a | | |
| - | | Combination Product | | |
| _ | | Combination Product | | |
| | larketing | Information | | |
| | larketing Marketing Category | | Marketing Start | Marketing End |

Labeler - Proficient Rx LP (079196022)

| Establishment | | | | | | |
|------------------|---------|-----------|---------------------|--|--|--|
| Name | Address | ID/FEI | Business Operations | | | |
| Proficient Rx LP | | 079196022 | RELABEL (71205-489) | | | |

Revised: 9/2022 Proficient Rx LP