HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use PRIFTIN® safely and effectively. See full prescribing information for PRIFTIN.

PRIFTIN® (rifapentine) Tablets

Initial U.S. Approval: 1998

INDICATIONS AND USAGE

PRIFTIN is indicated for the treatment of latent tuberculosis infection (LTBI) caused by M. tuberculosis in combination with isoniazid in patients 2 years of age and older at high risk of progression to TB disease. (1.2)

See Limitations of Use. (1.1, 1.2)

DOSE AND ADMINISTRATION

- Active pulmonary tuberculosis: PRIFTIN should be used in regimens consisting of an initial 2 month phase followed by a 4 month continuation phase. (2.1)
- Initial phase (2 Months): 600 mg twice weekly for two months as directly observed therapy (DOT), with no less than 72 hours between doses, in combination with other anti-tuberculosis drugs. (2.1)
- Continuation phase (4 Months): 600 mg once-weekly for 4 months as directly observed therapy with isoniazid or another appropriate anti-tuberculosis agent. (2.1)
- Latent tuberculosis infection: PRIFTIN should be administered in combination with isoniazid once-weekly for 12 weeks as directly observed therapy. Adults and children ≥ 12 years: PRIFTIN (based on weight, see table below) and isoniazid 15 mg/kg (900 mg maximum) Children 2–11 years: PRIFTIN (based on weight, see table below) and isoniazid 25 mg/kg (900 mg maximum)

Weight range PRIFTIN dose Number of PRIFTIN tablets
10–14 kg 300 mg 2
14.1–25 kg 450 mg 3
25.1–32 kg 600 mg 4
32.1–50 kg 750 mg 5
> 50 kg 900 mg 6

For Latent Tuberculosis Infection, the maximum recommended dose of PRIFTIN is 900 mg once-weekly for 12 weeks.
- Take with food. Tablets may be crushed and added to semi-solid food. (2.3)

DOSE FORMS AND STRENGTHS

150 mg tablets (3)

CONTRAINDICATIONS

Known hypersensitivity to any rifamycin (4.1)

WARNINGS AND PRECAUTIONS

- Hepatotoxicity: Monitor for symptoms of liver injury and discontinue PRIFTIN if signs or symptoms of liver injury occur. (5.1)
- Hypersensitivity: Discontinue PRIFTIN if signs or symptoms of hypersensitivity reaction occur. (5.2)
- Relapse in the treatment of active pulmonary tuberculosis: Do not use as a once-weekly continuation phase regimen with isoniazid in HIV-infected patients. Monitor for signs or symptoms of relapse in patients with cavitary lesions or bilateral disease. (5.3, 14.1)
- Drug Interactions: May interact with drugs metabolized by CYP450. (5.4, 7.1, 7.4)
- Discoloration of body fluids: May permanently stain contact lenses or dentures red-orange. (5.5)
- Clostridium difficile-associated colitis: Evaluate if diarrhea occurs. (5.6)
- Porphyria: Avoid use in patients with porphyria. (5.7)

ADVERSE REACTIONS

The most common adverse reactions with regimen for active pulmonary tuberculosis (1% and greater) are anemia, lymphopenia, neutropenia, increased ALT, arthralgia, conjunctivitis, headache, vomiting, nausea, diarrhea, rash, pruritus, anorexia and lymphadenopathy. The most common adverse reaction (1% and greater) with the regimen for latent tuberculosis infection is hypersensitivity reaction. (6.1)

To report SUSPECTED ADVERSE REACTIONS, contact sanofi-aventis U.S. LLC at 1-800-633-1610 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch

DRUG INTERACTIONS

- Protease Inhibitors and Reverse Transcriptase Inhibitors. (5.2, 7.1)
- Hormonal Contraceptives: Use another means of birth control. (7.3)
- May increase metabolism and decrease the activity of drugs metabolized by CYP450. (5.4, 7.1, 7.4)
- Hepatotoxicity: Monitor for symptoms of liver injury and discontinue PRIFTIN if signs or symptoms of liver injury occur. (5.1)

USE IN SPECIFIC POPULATIONS

- Pregnancy: Based on animal data, may cause fetal harm. (8.1)
- Nursing Mothers: Discontinue drug or nursing taking into consideration importance of drug to mother. (8.3)
- Pediatrics: Safety and effectiveness in treating active pulmonary tuberculosis in children under the age of 12 years have not been established. (8.4)

See 17 for PATIENT COUNSELING INFORMATION and Medication Guide

Revised: 12/2014

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FULL PRESCRIBING INFORMATION

1 INDICATIONS AND USAGE

1.1 Active Pulmonary Tuberculosis

PRIFTIN® (rifapentine) is indicated in adults and children 12 years and older for the treatment of active pulmonary tuberculosis (TB) caused by Mycobacterium tuberculosis. PRIFTIN must always be used in combination with one or more antituberculosis (anti-TB) drugs to which the isolate is susceptible.[see Dosage and Administration (2.1) and Clinical Studies (14.1)].

Limitations of Use

Do not use PRIFTIN monotherapy in either the initial or the continuation phases of active antituberculosis treatment.

PRIFTIN should not be used once-weekly in the continuation phase regimen in combination with isoniazid (INH) in HIV-infected patients with active pulmonary tuberculosis because of a higher rate of failure and/or relapse with rifampin (rifampicin)-resistant organisms [see Warnings and Precautions (5.3)] and Clinical Studies (14.2).

PRIFTIN has not been studied as part of the initial phase treatment regimen in HIV-infected patients with active pulmonary tuberculosis.

1.2 Latent Tuberculosis Infection

PRIFTIN is indicated in adults and children 2 years and older for the treatment of latent tuberculosis infection caused by Mycobacterium tuberculosis in patients at high risk of progression to tuberculosis disease (including those in close contact with active tuberculosis patients, recent conversion to a positive tuberculin skin test, HIV-infected patients, or those with pulmonary fibrosis on radiograph) [see Clinical Studies (14.2)].

Limitations of Use

Active tuberculosis disease should be ruled out before initiating treatment for latent tuberculosis infection.

PRIFTIN must always be used in combination with isoniazid as a 12-week once-weekly regimen for the treatment of latent tuberculosis infection. [see Dosage and Administration (2.2) and Clinical Studies (14.2)].

• PRIFTIN in combination with isoniazid is not recommended for Individuals presumed to be exposed to rifampin- or isoniazid-resistant M. tuberculosis.

2 DOSE ADMINISTRATION

2.1 Dosage in Active Pulmonary Tuberculosis

PRIFTIN is only recommended for the treatment of active pulmonary tuberculosis caused by drug-susceptible organisms as part of regimens consisting of a 2 month initial phase followed by a 4 month continuation phase.

Initial phase (2 Months): PRIFTIN should be administered at a dose of 600 mg twice weekly for 2 months as directly observed therapy (DOT), with an interval of no less than 3 consecutive days (72 hours) between doses, in combination with other anti-tuberculosis drugs as part of an appropriate regimen which includes daily companion drugs such as isoniazid (INH), ethambutol (EMB) and pyrazinamide (PZA).

Continuation phase (4 Months): Following the initial phase (2 phases), continuation phase (4 months) treatment consists of PRIFTIN 600 mg once-weekly for 4 months in combination with isoniazid or another appropriate anti-tuberculosis agent for susceptible organisms administered as directly observed therapy.

2.2 Dosage in Latent Tuberculosis Infection

PRIFTIN should be administered once-weekly in combination with isoniazid for 12 weeks as directly observed therapy.

Adults and children 12 years and older: The recommended dose of PRIFTIN should be determined based on weight of the patient up to a maximum of 900 mg once-weekly (see Table 1). The recommended dose of isoniazid is 15 mg/kg (rounded to the nearest 50 mg or 100mg) up to a maximum of 900 mg once-weekly for 12 weeks.

Children 2–11 years: The recommended dose of PRIFTIN should be determined based on weight of the patient up to a maximum of 900 mg once-weekly (see Table 1). The recommended dose of isoniazid is 25 mg/kg (rounded to the nearest 50 mg or 100mg) up to a maximum of 900 mg once-weekly for 12 weeks.

Table 1: Weight based dose of PRIFTIN in the treatment of latent tuberculosis infection

<table>
<thead>
<tr>
<th>Weight range</th>
<th>PRIFTIN dose</th>
<th>Number of PRIFTIN tablets</th>
</tr>
</thead>
<tbody>
<tr>
<td>10–14 kg</td>
<td>300 mg</td>
<td>2</td>
</tr>
<tr>
<td>14.1–25 kg</td>
<td>450 mg</td>
<td>3</td>
</tr>
<tr>
<td>25.1–32 kg</td>
<td>600 mg</td>
<td>4</td>
</tr>
<tr>
<td>32.1–50 kg</td>
<td>750 mg</td>
<td>5</td>
</tr>
<tr>
<td>&gt; 50 kg</td>
<td>900 mg</td>
<td>6</td>
</tr>
</tbody>
</table>

2.3 Administration

Take PRIFTIN with meals. Administration of PRIFTIN with a meal increases oral bioavailability and may improve the rate of absorption and compliance with therapy [see Patient Counseling Information (17)].

For patients who cannot swallow tablets, the tablets may be crushed and added to a small amount of semi-solid food, all of which should be consumed immediately [see Clinical Pharmacology (12.3)].

3 DOSAGE FORMS AND STRENGTHS

PRIFTIN is supplied as 150 mg round normal convex dark-pink film-coated tablets debossed “Prifitin” on top and “150” on the bottom.

4 CONTRAINDICATIONS

4.1 Hypersensitivity

PRIFTIN is contraindicated in patients with a history of hypersensitivity to rifamycins.

5 WARNINGS AND PRECAUTIONS

5.1 Hepatotoxicity

Elevations of liver transaminases may occur in patients receiving PRIFTIN [see Adverse Reactions (6.1)]. Patients on PRIFTIN should be monitored for symptoms of liver injury.

Patients with abnormal liver tests and/or liver disease or patients initiating treatment for active pulmonary tuberculosis should only be given PRIFTIN in cases of necessity and under strict medical supervision. In such patients, obtain serum transaminase levels prior to therapy and every 2–4 weeks while on therapy. Discontinue PRIFTIN if evidence of liver injury occurs.

5.2 Hypersensitivity and Related Reactions

Hypersensitivity reactions may occur in patients receiving PRIFTIN. Signs and symptoms of these reactions may include hypotension, urticaria, angioedema, acute bronchospasm, conjunctivitis, thrombocytopenia, neutropenia or flu-like syndrome (weakness, fatigue, muscle pain, nausea, vomiting, headache, fever, chills, aches, rash, itching, sweats, dizziness, shortness of breath, chest pain, cough, syncope, palpitations). There have been reports of anaphylaxis [see Patient Counseling Information (17)].

Monitor patients receiving PRIFTIN therapy for signs and/or symptoms of hypersensitivity reactions. If these symptoms occur, administer supportive measures and discontinue PRIFTIN.

5.3 Relapse in the Treatment of Active Pulmonary Tuberculosis

PRIFTIN has not been evaluated as part of the initial phase treatment regimen in HIV-infected patients with active pulmonary TB.

Do not use PRIFTIN as a once-weekly continuation phase regimen in HIV-infected patients with active pulmonary tuberculosis because of a higher rate of failure and/or relapse with rifampin-resistant organisms [see Clinical Studies (14.1)].

Higher relapse rates may occur in patients with cavitary pulmonary lesions and/or positive sputum cultures after the initial phase of active tuberculosis treatment and in patients with evidence of bilateral pulmonary disease. Monitor for signs and symptoms of TB relapse in these patients [see Clinical Studies (14.1)].

Poor adherence to therapy is associated with high relapse rate. Emphasize the importance of compliance with therapy [see Patient Counseling Information (17)].

5.4 Drug Interactions

Rifapentine is an inducer of CYP450 enzymes. Concomitant use of rifapentine with other drugs metabolized by these enzymes, such as protease inhibitors, certain reverse transcriptase inhibitors, and hormonal contraception may cause a significant decrease in plasma concentrations and loss of therapeutic effect [see Drug Interactions (7.1, 7.2, 7.3 and 7.4) and Clinical Pharmacology (12.3)].

5.5 Discoloration of Body Fluids

PRIFTIN may produce a red-orange discoloration of body tissues and/or fluids (e.g., skin, teeth, tongue, urine, feces, saliva, sputum, tears, sweat, and cerebrospinal fluid). Contact lenses or dentures may become permanently stained.

5.6 Clostridium difficile-Associated Diarrhea

Clostridium difficile-associated diarrhea (CDAD) has been reported with the use of nearly all systemic antibacterial agents, including PRIFTIN, with severity ranging from mild diarrhea to fatal colitis.

Treatment with antibacterial agents can alter the normal flora of the colon and permit overgrowth of C. difficile.

C. difficile produces toxins A and B which contribute to the development of CDAD. Hypotoxin producing strains of C. difficile cause increased morbidity and mortality, as these infections can be refractory to antimicrobial therapy and may require colectomy. CDAD must be considered in all patients who present with diarrhea following antibacterial use. Careful medical history is necessary because CDAD has been reported to occur over two months after the administration of antibacterial agents.

If CDAD is suspected or confirmed, discontinue antibacterial use not directed against C. difficile if possible. Institute appropriate measures such as fluid and electrolyte management, protein supplementation, antibacterial treatment of C. difficile, and surgical evaluation as clinically indicated.

5.7 Porphyrin

Porphyria has been reported in patients receiving rifampin, attributed to induction of delta amino levulinic acid synthetase. Because PRIFTIN may have similar enzyme induction properties, avoid the use of PRIFTIN in patients with porphyria.

6 ADVERSE REACTIONS

The following serious and otherwise important adverse drug reactions are discussed in greater detail in other sections of labeling:

• Hepatotoxicity [see Warnings and Precautions (5.1)]

• Hypersensitivity [see Contraindications (4.1) and Warnings and Precautions (5.2)]

• Discoloration of Body Fluids [see Warnings and Precautions (5.5)]

• Clostridium difficile-Associated Diarrhea [see Warnings and Precautions (5.6)]

• Porphyria [see Warnings and Precautions (5.7)]

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.

Active Pulmonary Tuberculosis

PRIFTIN was studied in a randomized, open label, active-controlled trial of HIV-negative patients with active pulmonary tuberculosis. The population consisted of primarily of male subjects with a mean age of 37 ± 11 years. In the initial 2 month phase of treatment, 361 patients received PRIFTIN 600 mg twice a week in combination with daily isoniazid, pyrazinamide, and ethambutol and 361 subjects received...
rifampin in combination with isoniazid, pyrazinamide and ethambutol all administered daily. Ethambutol was discontinued when drug susceptibly testing was known. During the 4 month continuation phase, 317 patients in the PRIFTIN group were randomized to receive PRIFTIN 600 mg twice-weekly with isoniazid and 304 patients in the rifampin group received twice weekly rifampin and isoniazid. Both treatment groups received pyridoxine (Vitamin B6) over the 6 month treatment period. Because PRIFTIN was administered as part of a combination regimen, the adverse reaction profile reflects the entire regimen.

Twenty-two deaths occurred in the study, eleven in the rifampin combination therapy group and eleven in the PRIFTIN combination therapy group. Three patients (two rifampin combination therapy patients and one PRIFTIN combination therapy patient) were discontinued in the initial phase due to hepatotoxicity. Concomitant medications for all three patients included isoniazid, pyrazinamide, ethambutol, and pyridoxine. All three recovered without sequelae.

Five patients had adverse reactions associated with PRIFTIN overdose. These reactions included hematuria, neutropenia, hyperglycemia, ALT increased, hyperuricemia, pruritus, and arthritis.

Table 2 presents selected treatment-emergent adverse reactions associated with the treatment regimens which occurred in at least 1% of patients during treatment and post-treatment through the first three months of follow-up.

A total of 4040 patients received at least one dose of the 3RPT/INH regimen, including 348 children 1–17 years of age and 3692 adults over 17 years of age. The death rate was 6.5% for the PRIFTIN combination regimen compared to 6.7% for the rifampin combination regimen.

The following selected treatment-emergent adverse reactions were reported in less than 1% of the PRIFTIN combination therapy patients during treatment and post-treatment through the first three months of follow-up.

<table>
<thead>
<tr>
<th>System Organ Class Preferred Term</th>
<th>Initial Phase</th>
<th>Continuation Phase</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PRIFTIN</td>
<td>Rifampin</td>
</tr>
<tr>
<td></td>
<td>Combination</td>
<td>Combination</td>
</tr>
<tr>
<td>(N=361) N (%)</td>
<td>(N=361) N (%)</td>
<td>(N=361) N (%)</td>
</tr>
<tr>
<td><strong>BLOOD AND LYMPHATICS</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anemia</td>
<td>41 (11.4)</td>
<td>41 (11.4)</td>
</tr>
<tr>
<td>Lymphopenia</td>
<td>38 (10.5)</td>
<td>37 (10.2)</td>
</tr>
<tr>
<td>Neutropenia</td>
<td>22 (6.1)</td>
<td>21 (5.8)</td>
</tr>
<tr>
<td>Leukocytosis</td>
<td>6 (1.7)</td>
<td>13 (3.6)</td>
</tr>
<tr>
<td>Thrombocytosis</td>
<td>20 (5.5)</td>
<td>13 (3.8)</td>
</tr>
<tr>
<td>Thrombocytopenia</td>
<td>6 (1.7)</td>
<td>6 (1.7)</td>
</tr>
<tr>
<td>Lymphadenopathy</td>
<td>4 (1.1)</td>
<td>6 (1.8)</td>
</tr>
<tr>
<td>Nonprotein Nitrogen Increased</td>
<td>4 (1.1)</td>
<td>3 (0.8)</td>
</tr>
<tr>
<td><strong>EYE</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Conjunctivitis</td>
<td>6 (1.7)</td>
<td>11 (3)</td>
</tr>
<tr>
<td>Vomiting</td>
<td>6 (1.7)</td>
<td>14 (3.9)</td>
</tr>
<tr>
<td>Nausea</td>
<td>7 (1.9)</td>
<td>3 (0.8)</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>5 (1.4)</td>
<td>2 (0.6)</td>
</tr>
<tr>
<td><strong>GENERAL</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Back Pain</td>
<td>15 (4.2)</td>
<td>11 (3)</td>
</tr>
<tr>
<td>Abdominal Pain</td>
<td>3 (0.8)</td>
<td>3 (0.8)</td>
</tr>
<tr>
<td>Fever</td>
<td>5 (1.4)</td>
<td>7 (1.9)</td>
</tr>
<tr>
<td>Anorexia</td>
<td>14 (3.9)</td>
<td>18 (5)</td>
</tr>
<tr>
<td><strong>HEPATIC &amp; BILIARY</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ALT Increased</td>
<td>18 (5)</td>
<td>23 (6.4)</td>
</tr>
<tr>
<td>AST Increased</td>
<td>15 (4.2)</td>
<td>18 (5)</td>
</tr>
<tr>
<td><strong>MUSCULOSKELETAL</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Arthralgia</td>
<td>13 (3.6)</td>
<td>13 (3.6)</td>
</tr>
<tr>
<td><strong>NEUROLOGIC</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Headache</td>
<td>11 (3)</td>
<td>13 (3.6)</td>
</tr>
<tr>
<td>Dizziness</td>
<td>5 (1.4)</td>
<td>5 (1.4)</td>
</tr>
<tr>
<td><strong>RESPIRATORY</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hemoptysis</td>
<td>27 (7.5)</td>
<td>20 (5.5)</td>
</tr>
<tr>
<td>Coughing</td>
<td>21 (5.8)</td>
<td>8 (2.2)</td>
</tr>
<tr>
<td><strong>SKIN</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rash</td>
<td>15 (4.2)</td>
<td>26 (7.2)</td>
</tr>
</tbody>
</table>
Table 3 Select Adverse Reactions occurring in 0.5% or greater of patients’ in the Latent Tuberculosis Infection Main Study

<table>
<thead>
<tr>
<th>System Organ Class Preferred Term</th>
<th>3RPT/INH (N=4040) N (%)</th>
<th>9INH (N=3579) N (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Immune system disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypersensitivity</td>
<td>161 (4)</td>
<td>18 (0.5)</td>
</tr>
<tr>
<td>Hepatobiliary disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hepatitis</td>
<td>24 (0.6)</td>
<td>113 (3)</td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Headache</td>
<td>26 (0.6)</td>
<td>17 (0.5)</td>
</tr>
<tr>
<td>Skin and subcutaneous tissue disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Skin reaction</td>
<td>31 (0.8)</td>
<td>21 (0.6)</td>
</tr>
</tbody>
</table>

*Includes events reported through 60 days after last dose of study drug.

Pediatric Substudy
Six-hundred and ninety children 2 years–17 years of age received at least one dose of study drugs in the main study. An additional 342 children 2 years–17 years of age received at least one dose in the pediatric extension study (total 1032 children; 539 received 3RPT/INH and 493 received 9INH). No children in either treatment arm developed hepatotoxicity. Using the same definition for rifamycin hypersensitivity reaction as in the main study, 7 (1.5%) of children in the 3RPT/INH group experienced a rifamycin hypersensitivity reaction. Adverse reactions in children 2 years–11 years of age and 12 years–17 years of age were similar.

HIV Substudy
Two-hundred HIV-infected patients with latent tuberculosis infection received at least one dose of study drugs in the main study and an additional 193 patients received at least one dose in the extension study (total of 393; 207 received 3RPT/INH and 186 received 9INH). Compared to the HIV-negative patients enrolled in the main study, a higher proportion of HIV-infected patients in each treatment arm experienced a treatment emergent adverse reaction, including a higher incidence of hepatotoxicity. Hepatotoxicity occurred in 3/207 (1.5%) patients in the 3RPT/INH arm and in 14/186 (7.5%) in the 9INH arm. Rifamycin hypersensitivity occurred in only one HIV-infected patient. Eleven deaths occurred during the 33 month follow up period (6/207 in the 3RPT/INH group and 5/186 in the 9INH group) including one death in the 9INH arm during the treatment emergent period. None of the reported deaths were considered related to treatment with study drugs or tuberculosis disease. Selected treatment-emergent adverse reactions reported during treatment and 60 days post-treatment in less 0.5% of the 3RPT/INH combination-therapy group in the main study are presented below by body system.

Eye Disorders: conjunctivitis.
Blood and Lymphatic System Disorders: leukopenia, anemia, lymphadenopathy, neutropenia.
Gastrointestinal Disorders: nausea, diarrhea, vomiting, abdominal pain constipation, dry mouth, dyspepsia, esophageal irritation, gastritis, pancreatitis.
General Disorders and Administration Site Conditions: fatigue, pyrexia, asthenia, chest pain, chills, feeling jittery.
Infections and Infestations: pharyngitis, viral infection, vulvovaginal candidiasis.
Metabolism and Nutrition Disorders: hyperglycemia, gout, hyperkalemia, decreased appetite, hyperlipidemia.
Musculoskeletal and Connective Tissue Disorders: arthralgia, myalgia, back pain, rhabdomyolysis.
Nervous system disorders: dizziness, convulsion, paresthesia, headache, neuropathy peripheral, syncope.
Psychiatric Disorders: depression, anxiety, disorientation, suicidal ideation.
Renal and Urinary Disorders: azotemia.
Reproductive System and Breast Disorders: vulvovaginal pruritus.
Respiratory, Thoracic and Mediastinal Disorders: cough, dyspnea, oropharyngeal pain, asthma, bronchial hyperactivity, epistaxis.
Skin and Subcutaneous Tissue Disorders: rash, hyperhidrosis, pruritus, urticaria.

7 DRUG INTERACTIONS
7.1 Protease Inhibitors and Reverse Transcriptase Inhibitors
Ritonavir is an inducer of CYP3A4 enzymes. Concomitant use of PRIFTIN with other drugs metabolized by these enzymes, such as protease inhibitors and certain reverse transcriptase inhibitors, may cause a significant decrease in plasma concentrations and loss of therapeutic effect of the protease inhibitor or reverse transcriptase inhibitor [see Warnings and Precautions (5.4) and Clinical Pharmacology (12.2)].

7.2 Fixed Dose Combination of Efavirenz, Emtricitabine and Tenofovir
Once-weekly co-administration of 900 mg PRIFTIN with the antiretroviral fixed dose combination of efavirenz 600 mg, emtricitabine 200 mg and tenofovir disoproxil fumarate 300 mg in HIV-infected patients did not result in any substantial change in steady state exposures of efavirenz, emtricitabine, and tenofovir. No clinically significant change in CD4 cell counts or viral loads were noted [see Clinical Pharmacology (12.3)].

7.3 Hormonal Contraceptives
PRIFTIN may reduce the effectiveness of hormonal contraceptives. Therefore, patients using oral, transdermal patch, or other systemic hormonal contraceptives should be advised to change to non-hormonal methods of birth control.

7.4 Cytochrome P450 3A4 and 2C8/9
Rifapentine is an inducer of cytochromes P450 3A4 and P4502C8/9. Therefore, PRIFTIN may increase the metabolism of other coadministered drugs that are metabolized by these enzymes. Induction of enzyme activities by PRIFTIN occurred within 4 days after the first dose. Enzyme activities returned to baseline levels 14 days after discontinuing PRIFTIN.

Rifapentine has been reported to accelerate the metabolism and may reduce the activity of the following drugs; hence, PRIFTIN may also increase the metabolism and decrease the activity of these drugs. Dosage adjustments of the drugs in Table 4 or of other drugs metabolized by cytochrome P450 3A4 or P4502C8/9 may be necessary if they are given concurrently with PRIFTIN.

Table 4. Drug Interactions with PRIFTIN: Dosage Adjustment may be Necessary

<table>
<thead>
<tr>
<th>Drug Class</th>
<th>Examples of Drugs Within Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antiarrhythmics</td>
<td>Disopyramide, mexiletine, quinidine, tocainide</td>
</tr>
<tr>
<td>Antibiotics</td>
<td>Chloramphenicol, clarithromycin, dapson, doxycycline, Fluoroquinolones (such as ciprofloxacin)</td>
</tr>
<tr>
<td>Oral Anticoagulants</td>
<td>Warfarin</td>
</tr>
<tr>
<td>Anticonvulsants</td>
<td>Phenytoin</td>
</tr>
<tr>
<td>Antimarial</td>
<td>Quinine</td>
</tr>
<tr>
<td>Azole Antifungals</td>
<td>Fluconazole, itraconazole, ketoconazole</td>
</tr>
<tr>
<td>Antipsychotics</td>
<td>Haloperidol</td>
</tr>
<tr>
<td>Barbiturates</td>
<td>Phenobarbital</td>
</tr>
<tr>
<td>Benzodiazepines</td>
<td>Diazepam</td>
</tr>
<tr>
<td>Beta-Blockers</td>
<td>Propanolol</td>
</tr>
<tr>
<td>Calcium Channel Blockers</td>
<td>Diltiazem, nifedipine, verapamil</td>
</tr>
<tr>
<td>Cardiac Glycoside Preparations</td>
<td>Digoxin</td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>Prednisone</td>
</tr>
<tr>
<td>Fibrates</td>
<td>Clofibrate</td>
</tr>
<tr>
<td>Oral Hypoglycemics</td>
<td>Sulfonylureas (e.g., glyburide, glipizide)</td>
</tr>
<tr>
<td>Hormonal Contraceptives/ Progestins</td>
<td>Ethinyl estradiol, levonorgestrel</td>
</tr>
<tr>
<td>Immunosuppressants</td>
<td>Cyclosporine, tacrolimus</td>
</tr>
<tr>
<td>Methylxanthines</td>
<td>Theophylline</td>
</tr>
<tr>
<td>Narcotic analgesics</td>
<td>Methadone</td>
</tr>
<tr>
<td>Phosphodiesterase-5 (PDE-5) Inhibitors</td>
<td>Sildenafil</td>
</tr>
<tr>
<td>Thyroid preparations</td>
<td>Levothyroxine</td>
</tr>
<tr>
<td>Tricyclic antidepressants</td>
<td>Amitriptyline, nortriptyline</td>
</tr>
</tbody>
</table>

7.5 Other Interactions
The conversion of PRIFTIN to 25-desacetyl rifapentine is mediated by an esterase enzyme. There is minimal potential for PRIFTIN metabolism to be inhibited or induced by another drug, based upon the characteristics of the esterase enzymes. Since PRIFTIN is highly bound to albumin, drug displacement interactions may also occur [see Clinical Pharmacology (12.5)].

7.6 Interactions with Laboratory Tests
Therapeutic concentrations of rifampin have been shown to inhibit standard microbiological assays for serum folate and Vitamin B12. Similar drug-laboratory interactions should be considered for PRIFTIN; thus, alternative assay methods should be considered.

8 USE IN SPECIFIC POPULATIONS
8.1 Pregnancy
Pregnancy Category: C. Risk Summary
There are no adequate and well controlled trials of PRIFTIN in pregnant women; however, there are limited pregnancy outcome data reported from women enrolled in clinical trials of various PRIFTIN treatment regimens for active tuberculosis and latent tuberculosis infection. The reported rate of spontaneous abortion following PRIFTIN exposure did not represent an increase over the background rate of spontaneous abortion reported in the general population. Further interpretation of these data is limited by the quality of clinical trial adverse event reporting. In animal reproduction and developmental toxicity studies, rifapentine produced fetal harm and was teratogenic at doses less than and similar to the recommended human dose. Because animal studies are not always predictive of human response, PRIFTIN should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

Clinical Considerations
Labor or Delivery
When administered during the last few weeks of pregnancy, rifampin, another rifamycin product, may increase the risk for maternal postpartum hemorrhage and bleeding in the exposed neonate. Monitor prothrombin time of pregnant women and neonates who are exposed to PRIFTIN during the last few weeks of pregnancy. Treatment with Vitamin K may be indicated.
Human Data

Fourteen patients with active tuberculosis treated with multiple anti-tuberculosis drugs including PRIFTIN became pregnant during clinical studies. Six delivered normal infants; four had first trimester spontaneous abortions (of these, one patient abused ethanol and another patient was HIV-infected); one had an elective abortion; and outcome was unknown in three patients. These data are, however, limited by the quality of reporting and confounded by co-morbid medical conditions and multiple anti-tuberculosis drug exposures.

In the trial that compared the safety and effectiveness of PRIFTIN in combination with isoniazid once-weekly regimen has been established in pediatric patients under the age of 12.

8.4 Pediatric Use

Ingestion of rifapentine during organogenesis at 40 mg/kg/day (0.6 times the human dose of 600 mg based on body surface area), produced pups with cleft palates, rachitic arch, increased incidence of delayed ossification, and increased numbers of ribs. When rifapentine was administered orally to mated female rats late in gestation, at 20 mg/kg/day (0.3 times the human dose based on body surface area), pup weights and gestational survival (live pups born/pups born) were reduced compared to controls. Increased resorptions and post-implantation loss, decreased mean fetal weights, increased numbers of stillborn pups, and slightly increased pup mortality during lactation were also noted. When pregnant rabbits received oral rifapentine at 10 mg/kg to 40 mg/kg (0.3 times to 1.3 times the human dose based on body surface area), major fetal malformations occurred including: ovarian agenesis, pes varus, arthritis, microphthalmia and irregularities of the ossified facial tissues. At the higher dose, there were increases in post-implantation loss and the incidence of stillbirth pups.

8.3 Nursing Mothers

It is not known whether PRIFTIN is present in human milk. Because many drugs are excreted in human milk and because of the potential for serious adverse reactions in nursing infants, a decision should be made whether to discontinue nursing or discontinue the drug, taking into account the importance of the drug to the mother [see Nonclinical Toxicology (13.1)]. Since PRIFTIN may produce a red-orange discoloration of body fluids, there is a potential for discoloration of breast milk. A slight increase in rat pup mortality was observed during lactation when dams were dosed late in gestation through lactation.

8.4 Pediatric Use

The safety and effectiveness of PRIFTIN in the treatment of active pulmonary tuberculosis have not been established in pediatric patients under the age of 12.

The safety and effectiveness of PRIFTIN in combination with isoniazid once-weekly regimen has been evaluated in pediatric patients (2–17 years of age) for the treatment of latent tuberculosis infection. In clinical studies, the safety profile in children was similar to that observed in adult patients [see Adverse Reactions (6.1) and Clinical Studies (14.2)].

In a pharmacokinetic study conducted in 2 to 11-year-old pediatric patients with latent tuberculosis infection, PRIFTIN was administered once-weekly based on weight (15mg/kg to 30 mg/kg, up to a maximum of 900 mg). Exposures (AUC) in children 2 years–11 years with latent tuberculosis infection were higher (average 31%) than those observed in adults receiving PRIFTIN 900mg once-weekly [see Dosage and Administration (2.2) and Clinical Pharmacology (12.3)].

8.5 Geriatric Use

Clinical studies with PRIFTIN did not include sufficient numbers of subjects aged 65 years and over to determine whether they respond differently from younger subjects. In a pharmacokinetic study with PRIFTIN, no substantial differences in the pharmacokinetics of rifapentine and 25-desacetyl rifapentine metabolite were observed in the elderly compared to younger adults [see Clinical Pharmacology (12.3)].

10 OVERDOSAGE

While there is no experience with the treatment of acute overdose with PRIFTIN, clinical experience with rifampicin suggests that gastric lavage to evacuate gastric contents (within a few hours of overdose), followed by instillation of an activated charcoal slurry into the stomach, may help absorb any remaining drug from the gastrointestinal tract.

Rifapentine and 25-desacetyl rifapentine are 91.7% and 93.2% plasma protein bound, respectively. Rifapentine and related compounds excreted in urine account for only 17% of the administered dose; therefore, neither hemodialysis nor forced diuresis is expected to enhance the systemic elimination of unchanged rifapentine from the body of a patient with PRIFTIN overdose.

11 DESCRIPTION

PRIFTIN (rifapentine) for oral administration contains 150 mg of the active ingredient rifapentine per tablet.

The 150 mg tablets also contain, as inactive ingredients: calcium stearate, disodium EDTA, FD&C Blue No. 2 aluminum lake, hydroxypropyl cellulose, hypromellose USP, microcrystalline cellulose, polyethylene glycol, pregelatinized starch, propylene glycol, sodium ascorbate, sodium lauryl sulfate, sodium starch glycolate, synthetic red iron oxide, and titanium dioxide.

Rifapentine is a rifamycin derivative antimicrobial and has a similar profile of microbiological activity to rifampicin. The molecular weight is 877.04.

The molecular formula is C_{57}H_{86}N_{10}O_{12}.

The chemical name for rifapentine is rifamycin, 3-{[4-cyclopropyl-1-piperazinyl][liminomethyl]-or 3-[N-(4-Cyclopropyl-1-piperazinyl)formimidoyl]liminomethyl}-or 5,6,9,17,19,21-hexahydroxy-23-methoxy-2,4,12,16,18,20,22-heptamethyl-8-N-[4-cyclopentyl-l-piperazinyl]-formimidoyl]-2,7-(epoxypentadeca [11,13]triminomethyl)napththo[2,1-b]uracil-1,11(2H)-dione 21-acetate. It has the following structure:

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Rifapentine, a cyclopentyl rifamycin, is an antimycobacterial agent [see Clinical Pharmacology, Microbiology (12.4)].

12.3 Pharmacokinetics

When oral doses of PRIFTIN were administered once daily or once every 72 hours to healthy volunteers for 10 days, single dose AUC of rifapentine was similar to its steady-state AUC (0–24h) or AUC(0–72h) values, suggesting no significant auto-induction effect on steady-state pharmacokinetics of rifapentine. Steady-state conditions were achieved by day 10 following daily administration of PRIFTIN 600 mg. No plasma accumulation of rifapentine and 25-desacetyl rifapentine (active metabolite) is expected after once weekly administration of PRIFTIN.

The pharmacokinetic parameters of rifapentine and 25-desacetyl rifapentine on day 10 following oral administration of 600 mg PRIFTIN every 72 hours to healthy volunteers are described in Table 5.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Rifapentine</th>
<th>25-Desacetyl Rifapentine</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean ± SD</td>
<td></td>
</tr>
<tr>
<td>C_{max} (µg/mL)</td>
<td>15.06 ± 4.62</td>
<td>6.26 ± 2.06</td>
</tr>
<tr>
<td>AUC (0–72h) (µg*h/mL)</td>
<td>319.54 ± 91.52</td>
<td>215.88 ± 85.96</td>
</tr>
<tr>
<td>T_{max} (h)</td>
<td>13.19 ± 1.38</td>
<td>13.35 ± 2.67</td>
</tr>
<tr>
<td>T_{1/2} (h)</td>
<td>4.83 ± 1.80</td>
<td>11.25 ± 2.73</td>
</tr>
<tr>
<td>C_{INF} (L/h)</td>
<td>2.03 ± 0.60</td>
<td>--</td>
</tr>
</tbody>
</table>

The pharmacokinetic parameters of rifapentine and 25-desacetyl rifapentine following single-dose oral administration of 900 mg PRIFTIN in combination with 900 mg isoniazid in fed conditions are described in Table 6.

12.6 Clinical Pharmacology, Microbiology

In a population pharmacokinetic analysis in 351 tuberculosis patients who received 600 mg PRIFTIN every 72 hours to healthy adult volunteers was 70%. The maximum concentrations were achieved from 5 hours to 6 hours after administration of the 600 mg PRIFTIN dose.

The administration of PRIFTIN with a high fat meal increased rifapentine C_{max} and AUC by 40% to 50% over that observed when PRIFTIN was administered under fasting conditions. The administration of PRIFTIN (900 mg single dose) and isoniazid (900 mg single dose) with a low fat, high carbohydrate breakfast, led to a 47% and 51% increase in rifapentine C_{max} and AUC, respectively. In contrast, the ingestion of the same meal decreased isoniazid C_{max} and AUC by 46% and of 23%, respectively.

Distribution

In a population pharmacokinetic analysis in 351 tuberculosis patients who received 600 mg PRIFTIN in combination with isoniazid, pyrazinamide and ethambutol, the estimated apparent volume of distribution was 70.2 ± 9.1 L. In healthy volunteers, rifapentine and 25-desacetyl rifapentine were 97.7%
Following a single 600 mg oral dose of radiolabeled rifapentine to healthy volunteers (n=15), 87% of the total 14C-rifapentine was recovered in the urine (17%) and feces (70%). Greater than 80% of the total 14C-rifapentine dose was excreted from the body within 7 days. Rifapentine was hydrolyzed by an esterase enzyme to form a microbiologically active 25-desacetyl rifapentine. Rifapentine and 25-desacetylrifapentine were recovered from 99% of the total radioactivity in plasma. Plasma AUC0–∞ and Cmax values of the 25-desacetyl rifapentine metabolite were one-half and one-third those of the rifapentine, respectively. Based upon relative in vitro activities and AUC0–∞ values, rifapentine and 25-desacetyl rifapentine potentially contributes 62% and 38% to the clinical activities against M. tuberculosis, respectively.

Specific Populations

Gender: In a population pharmacokinetics analysis of sparse blood samples obtained from 361 tuberculosis patients who received 600 mg PRIFTIN in combination with isoniazid, pyrazinamide and ethambutol, the estimated apparent oral clearance of PRIFTIN for males and females was 2.51 ± 0.14 L/h, respectively. The clinical significance of the difference in the estimated apparent oral clearance is not known.

Elderly: Following oral administration of a single 600 mg dose of PRIFTIN to elderly (65 years and older) male healthy volunteers (n=14), the pharmacokinetics of rifapentine and 25-desacetyl metabolite were similar to that observed for young (18 to 45 years) healthy male volunteers (n=20).

Pediatric: In a pharmacokinetic study in pediatric patients (age 2 to 12 years), a single oral dose of 150 mg PRIFTIN was administered to those weighing less than 30 kg (n=11) and a single oral dose of 300 mg was administered to those weighing greater than 30 kg (n=12). The mean estimates of AUC and Cmax were approximately 33% to 52% lower in these pediatric patients than those observed in healthy adults administered single oral doses of 600 mg and 900 mg.

A study compared the pharmacokinetics of rifapentine in pediatric patients (age 2 years to 11 years) with latent tuberculosis infection (n=80) receiving PRIFTIN once weekly based on weight (15 mg/kg–30 mg/kg, up to a maximum of 900 mg, see Table 7) to that of adults (n=77) receiving PRIFTIN 900 mg once weekly. Children who could not swallow whole tablets were administered crushed tablets. Rifapentine was administered to patients in soft food. Overall, the mean AUC of rifapentine in this age group was 31% higher compared to adult patients receiving 900 mg PRIFTIN once weekly (720 versus 551 mcg*h/mL). The geometric mean AUC of rifapentine was 60% higher in children administered whole tablets (884 versus 551 mcg*h/mL) and 19% higher in children administered crushed tablets (656 versus 551 mcg*h/mL), as compared to exposures in adults. Pediatric patients administered crushed PRIFTIN tablets had 26% lower rifapentine exposures compared to those pediatric patients who were given whole tablets.

Population pharmacokinetic analysis showed that rifapentine clearance adjusted to body weight decreased with increasing age of pediatric patients (2–18 years).

In another pharmacokinetics study of PRIFTIN in healthy adolescents (age 12 to 15 years), 600 mg PRIFTIN was administered to those weighing ≥45 kg (n=10) and 450 mg was administered to those weighing less than 45 kg (n=2). The pharmacokinetics of rifapentine were similar to those observed in healthy adult subjects.

Renal Impaired Patients: The pharmacokinetics of rifapentine have not been evaluated in renal impaired patients. Although only about 17% of an administered dose is excreted via the kidneys, the clinical significance of impaired renal function on the disposition of rifapentine and its 25-desacetyl metabolite is not known.

Hepatic Impaired Patients: Following oral administration of a single 600 mg dose of PRIFTIN to mild to severe hepatic impaired patients (n=15), the pharmacokinetics of rifapentine and 25-desacetyl metabolite were similar in patients with various degrees of hepatic impairment and to that observed in another study for healthy volunteers (n=12).

Asymptomatic HIV-Infected Volunteers: Following oral administration of a single 600 mg dose of PRIFTIN to asymptomatic HIV-infected volunteers (n=15) under fasting conditions, mean Cmax and AUC values of rifapentine were reduced by 52%–58% compared to those observed in healthy subjects (n=55). In a cross-study comparison, mean Cmax and AUC values of the 25-desacetyl rifapentine, when compared to healthy volunteers were higher (6%–21%) in one study (n=20), but lower (15%–16%) in a different study (n=40). The clinical significance of this observation is not known. Food (850 total calories: 33 g protein, 55 g fat, and 58 g carbohydrate) increases the mean AUC and Cmax by 6%–21% in one study and 15%–16% in another study compared to exposures in healthy volunteers.

12.4 Microbiology

Mechanism of Action

Rifapentine, a cyclopentyl rifamycin, inhibits DNA-dependent RNA polymerase in susceptible strains of Mycobacterium tuberculosis but does not affect mammalian cells at concentrations that are active against these bacteria. At therapeutic levels, rifapentine inhibits RNA transcription by preventing the initiation of RNA chain formation. It forms a stable complex with bacterial DNA-dependent RNA polymerase, leading to repression of RNA synthesis and cell death. Rifapentine and its 25-desacetyl metabolite accumulate in human monocyte-derived macrophages and are bactericidal to both intracellular and extracellular M. tuberculosis bacilli.

Mechanism of Resistance

The mechanism of resistance to rifapentine appears to be similar to that of rifampin. Bacterial resistance to rifapentine is caused by an alteration in the target site, the beta subunit of the DNA-dependent RNA polymerase. Resistance to rifapentine is conferred by the rifampin-resistant mutants in an otherwise susceptible population of M. tuberculosis strains is approximately one in 105 to 106 bacilli. Rifapentine resistance appears to be associated with monotherapy. Therefore, rifapentine should always be used in combination with other antibacterial drugs.

Cross Resistance

Mycobacterial species resistant to other rifamycins are likely to be resistant to rifapentine. A high level of cross-resistance between rifampin and rifapentine has been demonstrated with M. tuberculosis strains. Cross-resistance between rifapentine and non-rifamycin antitubercular agents has not been identified in clinical isolates.

Susceptibility Test Methods

In vitro susceptibility tests should be performed according to published methods1. Susceptibility test interpretive criteria and quality control ranges for in vitro susceptibility testing of Rifapentine have not been established.

13. NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

Hepatocellular carcinomas were increased in male NMRI mice (Harlan Winkelmann) which were treated orally with rifapentine for two years at or above doses of 5 mg/kg/day (0.04 times the recommended human dose based on body surface area conversions). In a two year rat study, there was an increase in nasal cavity adenomas in Wistar rats treated orally with rifapentine at 40 mg/kg/day (0.6 times human dose based on body surface area conversions). Rifapentine was negative in the following genotoxicity tests: in vitro gene mutation assay in bacteria (Ames test); in vitro point mutation test in Aspergillus nidulans; in vitro gene conversion assay in Saccharomyces cerevisiae; in vitro CHO cell transferase (CHO/HGPRT) forward mutation assay; in vivo chromosomal aberration assay utilizing rat lymphocytes; and in vivo mouse bone marrow micronucleus assay. The 25-desacetyl metabolite of rifapentine was positive in the in vitro mammalian chromosome aberration test in V79 Chinese Hamster cells, but was negative in the in vitro gene mutation assay in bacteria (Ames test), in the in vitro Chinese hamster ovary cell (CHO/HGPRT) forward mutation assay, and in the in vivo mouse bone marrow micronucleus assay. Fertility and reproductive performance were not affected by oral administration of rifapentine to male and female rats at doses of up to 20 mg/kg/day (one-third of the human dose based on body surface area conversions).

14. CLINICAL STUDIES

14.1 Active Pulmonary Tuberculosis

PRIFTIN was studied in two randomized, open-label controlled clinical trials in the treatment of active pulmonary tuberculosis.

The first trial was an open-label, prospective, parallel group, active controlled trial in HIV-negative patients with active pulmonary tuberculosis. The population comprised Black (approximately 60%) or multiracial (approximately 31%) patients. Treatment groups were comparable for age and sex and consisted primarily of male subjects with a mean age of 37 ± 11 years. In the initial 2 month phase of treatment, 361 patients received PRIFTIN 600 mg twice a week in combination with daily isoniazid, pyrazinamide, and ethambutol and 361 subjects received rifampin600 mg in combination with isoniazid, pyrazinamide and ethambutol all administered daily. The dosages of the combination drugs were the same in both treatment groups during the initial phase: isoniazid 300 mg, pyrazinamide 2000 mg, and ethambutol 1200 mg. For patients weighing less than 50 kg, the doses of rifampin (450 mg), pyrazinamide (1500 mg) and ethambutol (800 mg) were reduced. Ethambutol was discontinued when isoniazid and rifampin susceptibility testing results were confirmed. During the 4 month continuation phase of treatment, 17 patients in the PRIFTIN group continued to receive PRIFTIN 600 mg once weekly and isoniazid 300 mg and 304 patients in the rifampin group received twice weekly rifampin and isoniazid 900 mg. For patients weighing less than 50 kg, the doses of rifampin (450 mg) and isoniazid (600 mg) were reduced. Both treatment groups received pyridoxine (Vitamin B6) over the 6 month treatment period. Treatment was directly observed. 65/361 (18%) of patients in the PRIFTIN group and 143/361 (40%) in the rifampin group received overdoses of one or more of the administered study medications during the initial or continuation phase of treatment. Seven of these patients had adverse reactions reported with the overdose (5 in the PRIFTIN group and 2 in the rifampin group).

Table 7 below contains assessments of sputum conversion at end of treatment (6 months) and relapse rates at the end of follow-up (24 months).
**Table 8. Clinical Outcome in HIV Negative Patients with Active Pulmonary Tuberculosis**

<table>
<thead>
<tr>
<th>Status at End of 6 months of Treatment</th>
<th>PRIFTIN Combination Treatment % and (n/N)</th>
<th>Rifampin Combination Treatment % and (n/N)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Converted</td>
<td>87% (248/286)</td>
<td>80% (226/283)</td>
</tr>
<tr>
<td>Not Converted</td>
<td>1% (4/286)</td>
<td>3% (8/283)</td>
</tr>
<tr>
<td>Lost to Follow-up</td>
<td>12% (34/286)</td>
<td>17% (49/283)</td>
</tr>
<tr>
<td>Status Through 24 Month Follow-up*:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Relapsed</td>
<td>12% (29/246)</td>
<td>7% (15/226)</td>
</tr>
<tr>
<td>Sputum Negative</td>
<td>57% (142/248)</td>
<td>64% (145/226)</td>
</tr>
<tr>
<td>Lost to Follow-Up</td>
<td>31% (77/246)</td>
<td>29% (66/226)</td>
</tr>
</tbody>
</table>

All data for patients with confirmed susceptible M. tuberculosis (PRIFTIN combination treatment, N=286; rifampin combination treatment, N=283).

†Twenty-two (22) deaths occurred during the study; 11 in each treatment group.

Risk of relapse was greater in the group treated with the PRIFTIN combination. Higher relapse rates were associated with a lower rate of compliance as well as a failure to convert sputum cultures at the end of the initial 2 month treatment phase. Relapse rates were also higher for males in both regimens. Relapse in the PRIFTIN group was not associated with development of mono-resistance to rifampin.

The second trial was randomized, open-label performed in 1075 HIV-negative and positive patients with active pulmonary tuberculosis. Patients with culture-positive, drug-susceptible pulmonary tuberculosis who had completed the initial 2-month phase of treatment with 4 drugs (rifampin, isoniazid, pyrazinamide, and either ethambutol or streptomycin) under direct observation were randomly assigned to receive either PRIFTIN 600 mg and isoniazid 15 mg/kg (max 600 mg) once-weekly or rifampin 10 mg/kg (max 600 mg) and isoniazid 15 mg/kg (max 900 mg) twice weekly for the 4 month continuation phase. Study drugs were given under direct observation therapy in both groups.

In the PRIFTIN group, 502 HIV-negative and 36 HIV-positive patients were randomized and in the rifampin group 502 HIV-negative and 35 HIV-positive patients were randomized to treatment. Enrollment of HIV-infected patients was stopped when 4 of 36 patients in the PRIFTIN combination group relapsed with isolates that were rifampin resistant.

Table 9 below contains assessments of sputum conversion at the end of treatment (6 months total: 2 months of initial and 4 months of randomized continuation treatment) and relapse rates at the end of follow-up (24 months) in all HIV-negative patients randomized to treatment. Positive culture was based on either one sputum sample with >10 colonies on solid media or at least 2 positive sputum samples on liquid or solid media. However, only one sputum sample was collected at each visit in a majority of patients.

**Table 9. Clinical Outcome in HIV Negative Patients with Active Pulmonary Tuberculosis**

<table>
<thead>
<tr>
<th>Status at End of 4 Months Continuation Phase</th>
<th>PRIFTIN Combination Treatment % (n/N)</th>
<th>Rifampin Combination Treatment % (n/N)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment Response *</td>
<td>93.8% (471/502)</td>
<td>91% (457/502)</td>
</tr>
<tr>
<td>Not Converted</td>
<td>1% (5/502)</td>
<td>1.2% (6/502)</td>
</tr>
<tr>
<td>Did Not Complete Treatment†</td>
<td>4.2% (21/502)</td>
<td>7% (35/502)</td>
</tr>
<tr>
<td>Deaths</td>
<td>1% (5/502)</td>
<td>0.8% (4/502)</td>
</tr>
<tr>
<td>Status Through 24 Month Follow-up:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Relapsed</td>
<td>8.7% (41/471)</td>
<td>4.8% (22/477)</td>
</tr>
<tr>
<td>Sputum Negative</td>
<td>79.4% (374/471)</td>
<td>80.1% (368/457)</td>
</tr>
<tr>
<td>Lost to Follow-up</td>
<td>7.9% (37/471)</td>
<td>9.8% (45/457)</td>
</tr>
<tr>
<td>Deaths</td>
<td>4% (19/471)</td>
<td>5.3% (24/457)</td>
</tr>
</tbody>
</table>

*Treatment response was defined as subjects who had two negative sputum cultures after 16 doses of rifampin and isoniazid or after 8 doses of PRIFTIN and isoniazid, and remained sputum negative through the end of continuation phase therapy.

†Due to drug toxic effects, non-adherence, withdrawal of consent, receipt of non-study regimen, other.

In HIV-negative patients, higher relapse rates were seen in patients with a positive sputum culture at 2 months (i.e., at the time of study randomization), cavitary on chest x-ray, and bilateral pulmonary involvement.

Sixty-one HIV-positive patients were assessed for relapse. The rates of relapse were 16.7% (5/30) in the PRIFTIN group and 9.7% (3/31) in the rifampin group. In HIV-positive patients, 4 of the 5 relapses in the PRIFTIN combination group involved M. tuberculosis strains with rifampin monoresistance. No relapse strain in the twice weekly rifampin / isoniazid group acquired drug resistance.

The death rate among all study participants did not differ between the two treatment groups.

**Table 10. Outcomes in randomized patients at 33-months post-enrollment**

<table>
<thead>
<tr>
<th>Outcome</th>
<th>3RPT/INH (n=374)</th>
<th>9INH (n=374)</th>
<th>Difference*, 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tuberculosis n (%)</td>
<td>5 (0.16)</td>
<td>10 (0.32)</td>
<td>-0.16 (-0.42, 0.01)</td>
</tr>
<tr>
<td>Cumulative TB Rate (%)</td>
<td>0.17</td>
<td>0.35</td>
<td>-0.17 (-0.43, 0.09)</td>
</tr>
<tr>
<td>Deaths</td>
<td>22 (0.72)</td>
<td>35 (1.14)</td>
<td>-0.42 (-0.91, 0.06)</td>
</tr>
<tr>
<td>Lost to Follow-Up</td>
<td>320 (10.41)</td>
<td>357 (11.61)</td>
<td>-1.20 (-2.77, -0.36)</td>
</tr>
</tbody>
</table>

*Similar results were observed when all enrolled patients were included in the analysis.

†Rate in the 3RPT/INH group minus the rate in the 9INH group.

The proportion of patients completing treatment was 81.2% in the 3RPT/INH group and 68.3% in the 9INH group for a difference (3RPT/INH/9INH) of 12.8%, 95% CI (10.7, 15.0).

In the 9INH treatment group, two of the thirteen culture-confirmed cases were found to be isoniazid -monoresistant. In the 3RPT/INH treatment group, one of the seven cases was rifampin -resistant, isoniazid -susceptible M. bovis infection.

Pediatric Sub-study

Enrollment of children was extended after the overall target number of patients was attained in the main study. Data from both the main study and the extension were pooled resulting in an eligible population for analysis of 375 children in the 3RPT/INH arm and 367 in the 9INH arm.

One child in the 9INH group developed tuberculosis (1/367, cumulative rate 0.32%) versus zero tuberculosis cases in the 3RPT/INH group (0/375) at 33 months post-enrollment. The proportion of patients completing treatment in the 3RPT/INH and the 9INH groups was 87.5% and 79.6% respectively for a difference of 7.9%, 95% CI (2.5, 13.2).

HIV Sub-study

Enrollment of HIV-positive patients was extended after the overall target number of patients was attained in the main study. Data from both the main study and the extension were pooled resulting in an eligible population for analysis of 206 patients in the 3RPT/INH group and 193 in the 9INH group.

Tuberculosis disease developed in 1/206 patients in the 3RPT/INH group (cumulative rate, 0.32%) versus zero tuberculosis cases in the 3RPT/INH group at 33 months post-enrollment. The proportion of patients completing treatment in the 3RPT/INH and the 9INH groups was 97.7% and 96.2% respectively for a difference of 1.5%, 95% CI (0.8, 3.2).

**15 REFERENCES**


HOW SUPPLIED/STORAGE AND HANDLING

**How Supplied**

Priftin is supplied as 150 mg round normal convex dark-pink film-coated tablets debossed “Priftin” on top and “150” on the bottom, packaged in aluminum formable foil blister strips inserted into an aluminum foil laminated pouch.

Carton of 32 tablets (4 strips of 8 tablets) NDC 0088-2100-32

Carton of 24 tablets (3 strips of 8 tablets) NDC 0088-2100-24

**Storage**

Store at 25°C (77°F); excursions permitted 15–30°C (59–86°F) (see USP Controlled Room Temperature). Protect from excessive heat and humidity.

**17 PATIENT COUNSELING INFORMATION**

**17.1 Treatment Adherence**

Emphasize the importance of compliance with the full course of therapy, and the importance of not missing any doses of PRIFTIN or companion medications in the treatment of active pulmonary tuberculosis or the treatment of latent tuberculosis infection.

**17.2 Hypersensitivity Reactions**

Inform patients that PRIFTIN may cause hypersensitivity reactions. Signs and symptoms of this reaction may include a febrile illness, hypotension, urticaria, angioedema, bronchospasm, conjunctivitis, thrombocytopenia or neutropenia. Anaphylaxis may also occur [see Warnings and Precautions (5.2)]. Inform patients of signs and symptoms of hypersensitivity reactions and advise them to stop the medication and contact their healthcare provider if they experience any of these symptoms.

**17.3 Hepatitis**

Instruct patients to stop the medication and notify their physician promptly if they experience any of the following: fever, loss of appetite, malaise, nausea and vomiting, darkening urine, yellowish discoloration of the skin and eyes, and pain or swelling of the joints [see Warnings and Precautions (5.1)].

**17.4 Drug Interactions**

Rifapentine may increase the metabolism and decrease the activity of other drugs that are metabolized by the P4503A4 and 2C8/9 pathways. Dosage adjustments of the co-administered drugs may be
necessary. Advise patients to discuss with their physician any other medications they are taking before starting treatment with PRIFTIN. [see Warnings and Precautions (5.4), Drug Interactions 7.1 and 7.4]

Concomitant use of PRIFTIN with protease inhibitors or reverse transcriptase inhibitors may cause a significant decrease in plasma concentrations and loss of therapeutic effect of the protease inhibitor or reverse transcriptase inhibitor. [see Warnings and Precautions (5.4) and Drug Interactions 7.3]

Rifapentine may reduce the effectiveness of hormonal contraceptives. Advise patients using oral,transdermal patch, or other systemic hormonal contraceptives to change to non-hormonal methods of birth control. [see Drug Interactions 7.3]

17.5 Discoloration of Body Fluids
Inform the patient that PRIFTIN produces a reddish coloration of the urine, sweat, spu tum, tears, and breast milk. Contact lenses or dentures may be permanently stained. [see Warnings and Precautions (5.5), see Warnings and Precautions (5.4) and Drug Interactions 7.4]

17.6 Administration with Food
Advise patients to take PRIFTIN with food.

17.7 Nursing Mothers
Advise nursing mothers that breastfeeding is not recommended with PRIFTIN use [see Use in Specific Populations (8.3)].

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sanofi-aventis U.S. LLC
Bridgewater, NJ 08807
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Medication Guide
PRIFTIN (prif-tin) (rifapentine)
Tablets
Read this Medication Guide before you start taking PRIFTIN and each time you get a refill. There may be new information. This information does not take the place of talking with your doctor about your medical condition or your treatment.

What is the most important information I should know about PRIFTIN?
PRIFTIN may cause serious side effects, including:
- Liver problems. PRIFTIN may cause serious liver problems. Your doctor may do a blood test to check your liver function before and while you take PRIFTIN. Stop taking PRIFTIN and call your doctor right away if you have any of the following signs and symptoms of liver problems:
  - nausea
  - stomach pain
  - tiredness, yellowing skin or whites of your eyes
- Allergic reactions and flu-like symptoms. Allergic reactions and flu-like symptoms have happened in some people taking PRIFTIN. Signs and symptoms of an allergic reaction may include:
  - low blood pressure (hypotension)
  - hives
  - cough with wheezing
  - difficulty breathing
  - red eyes (conjunctivitis)
  - lower blood platelet levels
- signs and symptoms of a flu-like reaction may include:
  - weakness
  - nausea and vomiting
  - chills
  - itching
  - shortness of breath
  - fainting

What is PRIFTIN?
PRIFTIN is a prescription medicine used with other anti-tuberculosis (TB) medicines to:
- treat active tuberculosis disease of the lung in people age 12 years and older.
- prevent progression of inactive (latent) tuberculosis infection to active tuberculosis disease in people age 2 years and older.

PRIFTIN should not be used:
- alone to treat people with active or latent TB
- in people with active TB who had taken the medicines rifampin or isoniazid in the past and did not respond (resistant)
- in people who had been exposed to patients with TB that cannot be treated with isoniazid or rifampin

PRIFTIN is safe and effective in children older than 2 years of age who have inactive (latent TB), but it is not known if PRIFTIN is safe and effective for use in the treatment of active TB in children under 12 years of age.

Who should not take PRIFTIN?
- Do not take PRIFTIN if you are allergic to a group of medicines called rifamycins.

What should I tell my doctor before taking PRIFTIN?
Before you take PRIFTIN, tell your doctor if you:
- have active TB disease
- know that you have TB that is resistant to treatment with some medicines
- have HIV infection or taking medicines to treat HIV infection
- have liver problems
- have a condition called porphyria
- are pregnant or planning to become pregnant. It is not known if PRIFTIN will harm your unborn baby.
- are breastfeeding or plan to breastfeed. It is not known if PRIFTIN passes into your breast milk. You and your doctor should decide if you will take PRIFTIN or breastfeed.

Tell your doctor about all the medicines you take, including prescription and over-the-counter medicines, vitamins, and herbal supplements. Using PRIFTIN with other medicines may affect each other causing serious side effects. PRIFTIN may affect the way other medicines work, and other medicines may affect how PRIFTIN works. Especially tell your doctor if you take medicines to treat HIV infection or oral contraceptives.

Ask your doctor or pharmacist for a list of these medicines if you are not sure.

Know the medicines you take. Keep a list of them to show your doctor or pharmacist when you get a new medicine.

How should I take PRIFTIN?
- Take PRIFTIN exactly as your doctor tells you to take it.
- It is important to take all of your PRIFTIN and your other TB medicines. Do not skip doses. Skipping doses may cause PRIFTIN to not work as well and may increase the chance that your TB will not be treatable by PRIFTIN or other medicines.
- Take PRIFTIN with food.
- If you cannot swallow PRIFTIN tablets whole, they can be crushed and mixed with small amount of semisolid food. Be sure to take all of the semisolid food with PRIFTIN in it right away.

What are possible side effects of PRIFTIN?
PRIFTIN may cause serious side effects, including:
- See “What is the most important information I should know about PRIFTIN?”
- Relapse of your TB symptoms. Active TB disease may return after improvement (relapse) in some people, especially people who do not take PRIFTIN exactly as their doctor tells them to. It is important that you take PRIFTIN exactly as your doctor tells you to. Your doctor should check you for worsening signs and symptoms of your TB while you take PRIFTIN.
- change in the normal color of your skin, mouth and body fluids. PRIFTIN may cause your skin, teeth, tongue, urine, feces, saliva, sputum, tears, sweat, and breast milk to turn a red-orange color. Contact lenses or dentures may become permanently stained.
- diarrhea. A type of diarrhea called Clostridium difficile-associated diarrhea (CDAD) may occur during or after taking antibiotics, including PRIFTIN. The severity of CDAD can range from mild
diarrhea to severe diarrhea that may cause death (fatal colitis). Tell your doctor right if you have diarrhea while you take or after you stop taking PRIFTIN.

- worsening of a condition called porphyria.

The most common side effects of PRIFTIN include change in the color of body fluids to orange-red, allergic reactions and flu-like symptoms, abnormalities in liver tests, decrease in white blood cell and red blood cell count, decreased appetite, skin rash or itching, and red eyes. Tell your doctor if you have any side effect that bothers you or that does not go away. These are not all the possible side effects of PRIFTIN. For more information, ask your doctor or pharmacist.

Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

How should I store PRIFTIN?

- Store PRIFTIN at room temperature between 68°F to 77°F (20°C to 25°C).
- Keep PRIFTIN dry and away from heat.
- Keep PRIFTIN and all medicines out of reach of children.

General information about the safe and effective use of PRIFTIN.

Medicines are sometimes prescribed for purposes other than those listed in a Medication Guide. Do not use PRIFTIN for a condition for which it was not prescribed. Do not give PRIFTIN to other people, even if they have the same symptoms you have. It may harm them.

This Medication Guide summarizes the most important information about PRIFTIN. If you would like more information, talk with your doctor. You can ask your doctor or pharmacist for information about PRIFTIN that is written for healthcare professionals.

For more information, go to www.sanofi.us or call 1-800-633-1610, and select option 1.

What are the ingredients in PRIFTIN?

Active ingredient: rifapentine

Inactive ingredients: calcium stearate, disodium EDTA, FD&C Blue No. 2 aluminum lake, hydroxypropyl cellulose, hypromellose USP, microcrystalline cellulose, polyethylene glycol, pregelatinized starch, propylene glycol, sodium ascorbate, sodium lauryl sulfate, sodium starch glycolate, synthetic red iron oxide, and titanium dioxide

Manufactured by: sanofi-aventis U.S. LLC, Bridgewater, NJ 08807

This Medication Guide has been approved by the U.S. Food and Drug Administration

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RIE-FPLR-SL-DEC14 Rx Only