PRODUCT MONOGRAPH

PrMYCOBUTIN®

(rifabutin capsules USP)

150 mg Capsules

Antibacterial Agent

Pfizer Canada ULC
17,300 Trans-Canada Highway
Kirkland, Quebec H9J 2M5

Date of Preparation: 24 September 2003

Date of revision: February 11, 2021

Control No. 244143

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Pfizer Canada ULC, licensee
Pfizer Canada ULC, 2021
PRODUCT MONOGRAPH

NAME OF DRUG

PrMYCOBUTIN®
(rifabutin capsules USP)
150 mg Capsules

THERAPEUTIC CLASSIFICATION
Antibacterial Agent

ACTION AND CLINICAL PHARMACOLOGY

MYCOBUTIN® (rifabutin) is a derivative of rifamycin S, belonging to the class of ansamycins. The rifamycins owe their antimycobacterial efficacy to their ability to penetrate the cell wall and to their ability to complex with and to inhibit DNA-dependent RNA polymerase. Rifabutin has been found to interact with and to penetrate the outer layers of the mycobacterial envelope. Rifabutin inhibits DNA-dependent RNA polymerase in susceptible strains of Escherichia coli and Bacillus subtilis but not in mammalian cells. In resistant strains of E. coli, rifabutin, like rifampin, did not inhibit this enzyme.

It is not known whether rifabutin inhibits DNA-dependent RNA polymerase in Mycobacterium avium or in M. intracellulare which constitutes M. avium complex (MAC). Rifabutin inhibited incorporation of thymidine into DNA of rifampin-resistant M. tuberculosis suggesting that rifabutin may also inhibit DNA synthesis which may explain its activity against rifampin-resistant organisms.

Following oral administration, at least 53% of MYCOBUTIN dose is rapidly absorbed with rifabutin peak plasma concentrations attained in 2 to 4 hours. High-fat meals slow the rate without influencing the extent of absorption of rifabutin from the capsule dosage form.

The mean (± SD) absolute bioavailability assessed in HIV positive patients in a multiple dose study was 20% (±16%, n=5) on day 1 and 12% (± 5%, n=7) on day 28.

In healthy adult volunteers administered a single oral dose of 300 mg of rifabutin, the mean (± SD) peak plasma concentration (Cmax) was 375 (± 267) ng/mL (range: 141 to 1033 ng/mL).
Mean rifabutin steady-state trough levels (C\textsubscript{p, min}\textsuperscript{8h}, 24-hour post dose) ranged from 50 to 65 ng/mL in HIV positive patients and in healthy normal volunteers. Pharmacokinetic dose-proportionality over the 300 to 900 mg single dose range has been demonstrated in early symptomatic HIV positive patients and in healthy normal volunteers over the 300 to 600 mg single dose range.

Rifabutin appears to be widely distributed throughout the body and has been detected in all tissues and body fluids examined. Several times higher concentrations than those achieved in plasma have been observed in lung parenchyma, gall bladder and the small intestinal wall. The apparent volume of distribution at steady-state (V\textsubscript{ss}) estimated in early symptomatic HIV positive male patients following intravenous dosing was large (8 to 9 liters/kg), suggesting extensive distribution of rifabutin into the tissues. About 85% of the drug is bound to plasma proteins over a concentration range of 50 to 1000 ng/mL. Binding is predominantly to human serum albumin, is concentration independent and does not appear to be influenced by renal or hepatic dysfunction.

Rifabutin undergoes extensive oxidative metabolism. Of the five metabolites that have been identified, 25-O-deacetyl and 31-hydroxy are the most predominant and show a plasma metabolite:parent area under the curve ratio of 0.10 for 25-O-deacetyl and 0.07 for 31-hydroxy metabolite. The 25-O-deacetyl metabolite has antimycobacterial activity equal to the parent drug and contributes up to 10% to the total antimicrobial activity. The 31-hydroxy metabolite has some antimicrobial activity (1/16 that of parent drug), but, considering its concentration in plasma, it is probably not contributing significantly to the therapeutic activity of rifabutin. Rifabutin can induce its own metabolism on multiple dosing. The area under the plasma concentration-time curve (AUC) following multiple dosing decreased by 38%, but its terminal half-life remained unchanged.

The plasma elimination profile of rifabutin is biphasic with an initial half-life of approximately 4 hours followed by a mean terminal half-life of 45 (± 17) hours (range: 16 to 69 hours). Mean systemic clearance in healthy adult volunteers following a single oral dose was 0.69 (± 0.32) L/hr/kg (range: 0.46 to 1.34 L/hr/kg). Rifabutin is mainly excreted in the urine, primarily as metabolites and to a lesser extent in the faeces. Fifty-three percent (53%) of the oral dose of \textsuperscript{14}C-labelled drug was recovered in the urine by five days post-dose and 30% was recovered in the faeces over the same period. Renal and biliary excretion of unchanged drug each contribute approximately 5% to the systemic clearance.

The pharmacokinetic profile of rifabutin is not significantly modified by age or by hepatic dysfunction, although the inter-individual variability in elderly subjects (71-80 years) was slightly higher. Renal insufficiency was correlated to a decrease in urinary excretion with AUC and C\textsubscript{max} increases most apparent in severe disease. Caution and dose reductions may be required when treating patients with severe renal or severe hepatic impairment.

Rifabutin steady-state pharmacokinetics in early symptomatic HIV positive patients are similar to those in healthy normal volunteers but the variability between individuals is higher in the HIV positive patients. No rifabutin disposition information is currently available in children or adolescents under 18 years of age.
INDICATIONS AND CLINICAL USAGE

MYCOBUTIN® (rifabutin) is indicated for the prevention of disseminated *Mycobacterium avium* complex (MAC) disease in patients with advanced HIV infection (CD4+ cell count #200/mm³ with an AIDS defining diagnosis, or CD4+ cell count # 100/mm³ without an AIDS defining diagnosis).

To reduce the development of drug-resistant bacteria and maintain the effectiveness of MYCOBUTIN and other antibacterial drugs, MYCOBUTIN should be used only to treat infections that are proven or strongly suspected to be caused by susceptible bacteria. When culture and susceptibility information are available, they should be considered in selecting or modifying antibacterial therapy. In the absence of such data, local epidemiology and susceptibility patterns may contribute to the empiric selection of therapy.

CONTRAINDICATIONS

MYCOBUTIN® (rifabutin) is contraindicated in patients who have had clinically significant hypersensitivity to this drug, or to any other rifamycins.

WARNINGS

General:

MYCOBUTIN® (rifabutin) prophylaxis must not be administered to patients with active tuberculosis. Among HIV positive patients, tuberculosis is common and may present with atypical or extrapulmonary findings. Patients are likely to have a nonreactive purified protein derivative (PPD) test despite active disease. In addition to chest X-ray and sputum culture, the following studies may be useful in the diagnosis of tuberculosis in the HIV positive patient: blood culture, urine culture, or biopsy of a suspicious lymph node.

Patients who develop signs and symptoms consistent with active tuberculosis while on MYCOBUTIN prophylaxis should be evaluated immediately, so that those with active disease may be given an effective combination regimen of anti-tuberculosis medications. Administration of MYCOBUTIN, as a single-agent, to patients with active tuberculosis is likely to lead to the development of tuberculosis which is resistant both to MYCOBUTIN and to rifampin.

There is no evidence that MYCOBUTIN provides effective prophylaxis against *M. tuberculosis* infections. Patients requiring prophylaxis against both *M. tuberculosis* and *Mycobacterium avium* complex may be given isoniazid and MYCOBUTIN concurrently.
Clostridium difficile-associated disease (CDAD) has been reported with use of many antibacterial agents, including MYCOBUTIN (rifabutin). CDAD may range in severity from mild diarrhea to fatal colitis. It is important to consider this diagnosis in patients who present with diarrhea, or symptoms of colitis, pseudomembranous colitis, toxic megacolon, or perforation of colon subsequent to the administration of any antibacterial agent. CDAD has been reported to occur over 2 months after the administration of antibacterial agents.

Treatment with antibacterial agents may alter the normal flora of the colon and may permit overgrowth of Clostridium difficile. C. difficile produces toxins A and B, which contribute to the development of CDAD. CDAD may cause significant morbidity and mortality. CDAD can be refractory to antimicrobial therapy.

If the diagnosis of CDAD is suspected or confirmed, appropriate therapeutic measures should be initiated. Mild cases of CDAD usually respond to discontinuation of antibacterial agents not directed against Clostridium difficile. In moderate to severe cases, consideration should be given to management with fluids and electrolytes, protein supplementation, and treatment with an antibacterial agent clinically effective against Clostridium difficile. Surgical evaluation should be instituted as clinically indicated, as surgical intervention may be required in certain severe cases.

Due to increased plasma concentrations and other pharmacokinetic considerations, a decrease in dosage may be considered in patients with severe renal impairment or severe liver insufficiency or because of potential CYP450 3A-related interactions in patients co-administered certain other drugs (see PRECAUTIONS, DOSAGE AND ADMINISTRATION).

Significant drug-drug interactions between rifabutin and protease inhibitors, among many other drugs, requires careful consideration based upon the overall assessment of the patient and patient’s specific drug profile since drug safety and efficacy can be impacted. Patients should be carefully monitored to avoid uveitis. If uveitis is suspected, the patient should be referred to an ophthalmologist and, if considered necessary, treatment with MYCOBUTIN should be suspended (see PRECAUTIONS, Drug Interactions, ADVERSE REACTIONS).

Rifabutin is a CYP450 3A inducer. Therefore, co-administration with antiretroviral products including but not limited to bictegravir, rilpivirine, or doravirine is not recommended due to the expected decrease in plasma concentrations of the antiretrovirals which may lead to loss of virologic response and possible development of resistance (see PRECAUTIONS, Rifabutin Interaction Studies).

For further recommendations, please refer to the most recent Product Monograph of the antiretrovirals or contact the specific manufacturer.

Anaphylactic shock has occurred with other antibiotics of the same class.

There have been reports of severe cutaneous adverse reactions (SCARs), such as Stevens-Johnson syndrome (SJS), toxic epidermal necrolysis (TEN), drug reaction with eosinophilia and systemic symptoms (DRESS), and acute generalized exanthematous pustulosis (AGEP) with anti-
tuberculosis drugs (see ADVERSE REACTIONS). If patients develop a skin rash they should be monitored closely and suspect drug(s) discontinued if lesions progress. Identifying the specific drug is difficult, as multiple anti-tuberculosis drugs are prescribed in association concurrently. Specifically, for DRESS, a multi-system potential life-threatening SCAR, time to onset of the first symptoms may be prolonged. DRESS is a clinical diagnosis, and its clinical presentation remains the basis for decision making. An early withdrawal of the suspect drug is essential because of the syndrome’s mortality and visceral involvement (e.g., liver, bone marrow or kidney).

Susceptibility/Resistance:

Development of Drug Resistant Bacteria

Prescribing MYCOBUTIN in the absence of a proven or strongly suspected bacterial infection is unlikely to provide benefit to the patient and risks the development of drug-resistant bacteria.

PRECAUTIONS

General
Because MYCOBUTIN® (rifabutin) may be associated with neutropenia, and more rarely thrombocytopenia, physicians should consider obtaining hematologic studies periodically in patients receiving MYCOBUTIN prophylaxis.

Use in the Elderly
MYCOBUTIN administered as a single dose has been evaluated in 24 healthy, elderly (71-80 years) volunteers. The pharmacokinetic profile of MYCOBUTIN is not significantly modified by age, although the inter-individual variability in this age group was slightly higher when compared to younger (25-37 years) volunteers.

Use in Children
The safety and effectiveness of MYCOBUTIN for prophylaxis of MAC disease in children and adolescents under 18 years of age have not been established. However, limited safety data are available from 22 HIV positive children who received MYCOBUTIN as treatment for disseminated MAC disease, in combination with at least two other antimycobacterials for periods ranging from 1 to 183 weeks.

The mean daily doses (mg/kg) for these children were: infants one year of age, 18.5 (range 15.0 to 25.0); children 2 to 10 years, 8.6 (range 4.4 to 18.8); adolescents 14 to 16 years, 4.0 (range 2.8 to 5.4). MYCOBUTIN was generally safe in this treatment group. Adverse experiences were similar to those observed in the adult population, and included leukopenia, neutropenia and skin rash. Doses of MYCOBUTIN may be administered mixed with foods such as applesauce.

Usage in Pregnancy
There are no adequate and well-controlled studies of MYCOBUTIN use in pregnant women. No teratogenic effects were observed in reproduction studies carried out in rats and rabbits. Because animal reproduction studies are not always predictive of human response, MYCOBUTIN should
be used in pregnant women only if the potential benefit justifies the potential risk to the fetus.

**Nursing Mothers**
It is not known whether MYCOBUTIN is excreted in human milk. Because many drugs are excreted in human milk and given 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 nursing mother.

**Use In Renal Impairment**
Caution is recommended when treating patients with severe renal insufficiency.

The disposition of rifabutin (300 mg) was studied in 18 patients with varying degrees of renal function. Area under plasma concentration time curve (AUC) increased by about 71% in patients with severe renal insufficiency (creatinine clearance below 30 mL/min) compared to patients with creatinine clearance (Cr\(_{\text{cl}}\)) between 61–74 mL/min. In patients with mild to moderate renal insufficiency (Cr\(_{\text{cl}}\) between 30–61 mL/min), the AUC increased by about 41%. A 50% reduction in the dosage of rifabutin is recommended for patients with Cr\(_{\text{cl}}\) < 30 mL/min. No dosage adjustment is recommended in mild to moderate renal impairment (see **DOSAGE AND ADMINISTRATION**).

**Use In Hepatic Impairment**
The pharmacokinetics of rifabutin were studied in 40 patients with mild (n=30), moderate (n=6) and severe (n=4) hepatic impairment. Significant variability was noted. Caution should be exercised in treating patients with severe hepatic disease. For patients with severe liver insufficiency a dose reduction should be considered. Mild and moderate hepatic impairment does not require a dose modification.

**Drug Interactions**
Multiple dosing of rifabutin has been associated with induction of hepatic metabolic enzymes of the CYP450 3A subfamily. Rifabutin’s predominant metabolite (25-desacetyl rifabutin; LM 565), may also contribute to this effect. Metabolic induction due to rifabutin is likely to produce a decrease in circulating levels of concomitantly administered drugs (especially those metabolized by the CYP450 3A pathway). Kinetic data suggest that enzymatic induction by rifabutin is complete within 5 days and is dose-independent over the 300 to 600 mg dose-range. Similarly, concomitant medications that competitively inhibit the CYP450 3A activity may increase circulating levels of rifabutin.

**Malabsorption**
Gastric pH alteration due to progressing HIV disease has been linked with malabsorption of some drugs used in HIV-positive patients (e.g., rifampin, isoniazid). Drug serum concentration data from AIDS patients with varying disease severity (based on CD4+ counts) suggest that rifabutin absorption is not influenced by progressing HIV disease. However, when stomach pH is increased with drug co-administration, rifabutin absorption may be impaired.
Effects on Other Drugs
Rifabutin induces CYP450 3A enzymes and therefore may reduce the plasma concentrations of drugs metabolized by those enzymes. This effect may reduce the efficacy of standard doses of such drugs (see below).

Effects on Rifabutin
Some drugs that inhibit CYP450 3A may significantly increase the plasma concentration of rifabutin. Because high plasma levels of rifabutin may increase the risk of adverse reactions, patients coadministered such drugs should be carefully monitored and in some cases the doses of MYCOBUTIN may need to be reduced (see below).

The following table summarizes the results and magnitude of the pertinent drug interactions assessed with rifabutin as reported in selected not all-inclusive publications from the scientific literature. The clinical relevance of these interactions and subsequent dose modifications, which are largely based upon pharmacokinetic data extrapolations, should be judged in light of the population studied, severity of the disease, patient drug profile, and the likely impact on the risk/benefit ratio.

Rifabutin Interaction Studies

<table>
<thead>
<tr>
<th>Coadministered Drugs</th>
<th>Effect on Rifabutin</th>
<th>Effect on Coadministered Drug</th>
<th>Comments</th>
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</thead>
<tbody>
<tr>
<td><strong>ANTIRETROVIRALS</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Amprenavir</td>
<td>2.9-fold ↑ AUC, 2.2-fold ↑ Cmax</td>
<td>No significant change in kinetics.</td>
<td>A 50% reduction in the rifabutin dose is recommended when combined with amprenavir. Increased monitoring for adverse reactions is warranted.</td>
</tr>
<tr>
<td>Bictegravir</td>
<td>ND</td>
<td>AUC ↓38% Cmin ↓56% Cmax ↓20%</td>
<td>Although not studied, co-administration of rifabutin with Biktarvy (bictegravir/entricitabine/tenofovir alafenamide) is not recommended due to an expected decrease in tenofovir alafenamide in addition to the reported reduction in bictegravir.</td>
</tr>
<tr>
<td>Delavirdine</td>
<td>ND</td>
<td>Oral clearance ↑ 5-fold resulting in significantly lower mean trough plasma concentrations (18±15 to 1.0±0.7 µM)</td>
<td>Study conducted in HIV-1 infected patients Rifabutin is not recommended for patients dosed with delavirdine mesylate 400 mg q8h.</td>
</tr>
<tr>
<td>Didanosine</td>
<td>No significant change in kinetics.</td>
<td>No significant change in kinetics at steady state.</td>
<td>Didanosine administration with pH increasing buffers may decrease rifabutin absorption</td>
</tr>
<tr>
<td>Doravirine</td>
<td>ND</td>
<td>50% ↓ in AUC 68% ↓ in C24 ↔ in Cmax</td>
<td>If concomitant use is necessary, increase the doravirine dosage as instructed in doravirine-containing product prescribing information.</td>
</tr>
<tr>
<td>Fosamprenavir/ritonavir *</td>
<td>64% ↑ AUC **</td>
<td>35% ↑ AUC and 36% ↑ Cmax, no effect Ctrough (amprenavir)</td>
<td>Dosage reduction of rifabutin by at least 75% (to 150 mg every other day or three times per week) is recommended when combined with fosamprenavir</td>
</tr>
<tr>
<td>Coadministered Drugs</td>
<td>Effect on Rifabutin</td>
<td>Effect on Coadministered Drug</td>
<td>Comments</td>
</tr>
<tr>
<td>----------------------</td>
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</tr>
<tr>
<td>Indinavir</td>
<td>173% ↑ in AUC, 134% ↑ Cmax</td>
<td>34%↓ in AUC, 25%↓ in Cmax</td>
<td>Dose reduction of rifabutin to half the standard dose and increase of indinavir to 1000 mg every 8 hours are recommended when rifabutin and indinavir are coadministered.</td>
</tr>
<tr>
<td>Lopinavir/ritonavir*</td>
<td>5.7-fold ↑ AUC, 3.4 fold ↑ Cmax**</td>
<td>No significant change in lopinavir kinetics.</td>
<td>Dosage reduction of rifabutin by at least 75% of the usual dose of 300 mg/day is recommended (i.e., a maximum dose of 150 mg every other day or three times per week). Increased monitoring for adverse reactions is warranted. Further dosage reduction of rifabutin may be necessary.</td>
</tr>
<tr>
<td>Saquinavir</td>
<td>ND</td>
<td>40% ↓ in AUC</td>
<td></td>
</tr>
<tr>
<td>Rilpivirine</td>
<td>ND</td>
<td>42% ↓ in AUC, 48% ↓ in Cmin, 31% ↓ in Cmax</td>
<td>Although not studied, co-administration of rifabutin with Odefsey (rilpivirine/tenofovir alafenamide/emtricitabine) is not recommended due to an expected decrease in tenofovir alafenamide in addition to the reported reduction in rilpivirine.</td>
</tr>
<tr>
<td>Ritonavir *</td>
<td>4 fold increase in AUC, 2.5 fold increase in Cmax</td>
<td>ND</td>
<td>In the presence of ritonavir the subsequent risk of side effects, including uveitis may be increased. If a protease inhibitor is required in a patient treated with rifabutin, agents other than ritonavir should be considered.</td>
</tr>
<tr>
<td>Tipranavir/ritonavir *</td>
<td>2.9-fold ↑ AUC, 1.7-fold ↑ Cmax</td>
<td>No significant change in tipranavir kinetics.</td>
<td>Therapeutic drug monitoring of rifabutin is recommended.</td>
</tr>
<tr>
<td>Zidovudine</td>
<td>No significant change in kinetics.</td>
<td>Approximately 32%↓ in Cmax and AUC</td>
<td>A large controlled clinical study has shown that these changes are of no clinical relevance.</td>
</tr>
</tbody>
</table>

**ANTIFUNGALS**

<table>
<thead>
<tr>
<th></th>
<th>Effect on Rifabutin</th>
<th>Effect on Steady-State Plasma Concentrations</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluconazole</td>
<td>82% ↑ in AUC</td>
<td>No significant change in</td>
<td>One case report suggests a kinetic interaction resulting in an increase in serum rifabutin levels and a risk for developing uveitis in the presence of itraconazole.</td>
</tr>
<tr>
<td>Itraconazole</td>
<td>ND</td>
<td>70% to 75% ↓ in Cmax and AUC</td>
<td></td>
</tr>
<tr>
<td>Posaconazole</td>
<td>31%↑ Cmax, 72%↑ AUC</td>
<td>43%↓ Cmax, 49%↓ AUC</td>
<td>If the drugs are coadministered, patients should be monitored for adverse events associated with rifabutin administration.</td>
</tr>
<tr>
<td>Coadministered Drugs</td>
<td>Effect on Rifabutin</td>
<td>Effect on Coadministered Drug</td>
<td>Comments</td>
</tr>
<tr>
<td>----------------------</td>
<td>---------------------</td>
<td>-------------------------------</td>
<td>----------</td>
</tr>
<tr>
<td>Voriconazole</td>
<td>195%↑ Cmax, 331%↑ AUC ***</td>
<td>Rifabutin (300 mg once daily) decreased the Cmax and AUC of voriconazole at 200 mg twice daily by 69% and 78%, respectively. During co-administration with rifabutin, the Cmax and AUC of voriconazole at 350 mg twice daily were 96% and 68% of the levels when administered alone at 200 mg twice daily. At a voriconazole dose of 400 mg twice daily Cmax and AUC were 104% and 87% higher, respectively, compared with voriconazole alone at 200 mg twice daily.</td>
<td>Concurrent administration of voriconazole and rifabutin is not recommended.</td>
</tr>
</tbody>
</table>

**ANTI-PCP (Pneumocystis jirovecii pneumonia †)**

<table>
<thead>
<tr>
<th>Dapsone</th>
<th>ND</th>
<th>Approximately 27% to 40% ↓ in AUC</th>
<th>Study conducted in HIV infected patients (rapid and slow acetylators).</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulfamethoxazole-Trimethoprim</td>
<td>No significant change in Cmax and AUC</td>
<td>Approximately 15% to 20% ↓ in AUC</td>
<td>In another study, only trimethoprim (not sulfamethoxazole) had 14% ↓ in AUC and 6%↓ in Cmax but were not considered clinically significant.</td>
</tr>
</tbody>
</table>

**ANTI-MAC (Mycobacterium avium intracellulare complex)**

<table>
<thead>
<tr>
<th>Azithromycin</th>
<th>No PK interaction</th>
<th>No PK interaction</th>
<th>Study conducted in HIV infected patients. Dose of rifabutin should be adjusted in the presence of clarithromycin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clarithromycin</td>
<td>Approximately 77% ↑ in AUC</td>
<td>Approximately 50%↓ in AUC</td>
<td></td>
</tr>
</tbody>
</table>

**ANTI-TB (Tuberculosis)**

<table>
<thead>
<tr>
<th>Ethambutol</th>
<th>ND</th>
<th>No significant change in AUC or Cmax</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Isoniazid</td>
<td>ND</td>
<td>Pharmacokinetics not affected</td>
<td>Study data being evaluated.</td>
</tr>
<tr>
<td>Pyrazinamide</td>
<td>ND</td>
<td>ND</td>
<td></td>
</tr>
</tbody>
</table>

**OTHER**

| Methadone | ND | No significant effect | No apparent effect of rifabutin on either peak levels of methadone or systemic exposure based upon AUC. Rifabutin kinetics not evaluated. |
| Ethinylestradiol | ND | 35%↓ AUC 20%↓ Cmax | Patients should be advised to use other methods of contraception. |
| Norethindrone | ND | 46%↓ AUC | Patients should be advised to use other methods of contraception. |
| Tacrolimus | ND | ND | Authors report that rifabutin decreases tacrolimus trough blood levels. |
### Coadministered Drugs

<table>
<thead>
<tr>
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<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Theophylline</td>
<td>ND</td>
<td>No significant change in AUC or Cmax compared with baseline.</td>
<td></td>
</tr>
</tbody>
</table>

ND: Not done; AUC: Area under the Concentration vs. Time Curve; Cmax: Maximum serum concentration
* A lower dose of ritonavir was used when combined with fosamprenavir, lopinavir or tipranavir than when used alone; the PK effects upon ritonavir used alone or in combination with these antivirals were not studied.
** - Drug plus active metabolite
*** - Voriconazole dosed at 400 mg twice daily
† Formerly known as Pneumocystis carinii pneumonia

### Other drugs

MYCOBUTIN has liver enzyme-inducing properties. The related drug rifampin is known to reduce the activity of a number of drugs, including dapsone, narcotics (including methadone), anticoagulants, corticosteroids, cyclosporine, cardiac glycoside preparations, quinidine, oral contraceptives, oral hypoglycemic agents (sulfonylureas), and analgesics. Rifampin has also been reported to decrease the effects of concurrently administered ketoconazole, barbiturates, diazepam, verapamil, beta-adrenergic blockers, clofibrate, progestins, disopyramide, mexiletine, theophylline, chloramphenicol, and anticonvulsants. Because of the structural similarity of rifabutin and rifampin, MYCOBUTIN may be expected to have some effect on these drugs as well. However, unlike rifampin, MYCOBUTIN appears not to affect the acetylation of isoniazid. When the effects of rifabutin on hepatic microsomal enzyme activity were compared to those of rifampin in a study with 8 healthy normal volunteers, rifabutin appeared to be a less potent enzyme inducer than rifampin. The significance of this finding for clinical drug interactions is not known. Dosage adjustment of drugs listed above may be necessary if they are given concurrently with MYCOBUTIN.

Patients using oral contraceptives should consider changing to nonhormonal methods of birth control.

### ADVERSE REACTIONS

MYCOBUTIN® (rifabutin) was generally well tolerated in the controlled clinical trials involving 566 patients treated with MYCOBUTIN and 580 patients treated with placebo. The most serious adverse reaction to MYCOBUTIN was neutropenia.

The most common adverse events, reported more frequently in the MYCOBUTIN treated patients than in the placebo group were: urine discoloration, neutropenia, skin rash, nausea and/or vomiting, and abdominal pain (see tables). The incidence of urine discoloration and neutropenia in patients treated with MYCOBUTIN were significantly greater than in patients treated with placebo (Fisher's Test, p< 0.01 and p=0.03 respectively).

Sixteen percent (16%) of MYCOBUTIN treated patients discontinued therapy due to an adverse event as compared to 8% of placebo-treated patients. The primary reasons for discontinuation of MYCOBUTIN were: skin rash (4%), gastrointestinal intolerance (3%) and neutropenia (2%).
The following table enumerates adverse experiences that occurred at a frequency of 1% or greater among the patients treated with MYCOBUTIN and those treated with placebo in the Phase III clinical trials.

<table>
<thead>
<tr>
<th>ADVERSE EVENT</th>
<th>MYCOBUTIN (n=566) %</th>
<th>PLACEBO (n=580) %</th>
</tr>
</thead>
<tbody>
<tr>
<td>BODY AS A WHOLE</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abdominal Pain</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>Headache</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>Fever</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Asthenia</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Chest Pain</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Pain</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>DIGESTIVE SYSTEM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nausea</td>
<td>6</td>
<td>5</td>
</tr>
<tr>
<td>Nausea and vomiting</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>Vomiting</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Dyspepsia</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Eructation</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Anorexia</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Flatulence</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>MUSCULOSKELETAL SYSTEM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myalgia</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>NERVOUS SYSTEM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Insomnia</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>SKIN AND APPENDAGES</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rash</td>
<td>11</td>
<td>8</td>
</tr>
<tr>
<td>Pruritus</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>SPECIAL SENSES</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Taste Perversion</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>UROGENITAL SYSTEM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Discoloured urine</td>
<td>30</td>
<td>6</td>
</tr>
</tbody>
</table>

Considering data from the Phase III clinical trials, and from other clinical studies, MYCOBUTIN appears to be a likely cause of the following adverse events which occurred in less than 1% of the treated patients: arthralgia, chest pressure or pain with dyspnea, hemolysis, hepatitis, myositis, and skin discoloration.
The following adverse events have occurred in more than one patient receiving MYCOBUTIN, but an etiologic role for MYCOBUTIN has not been established: aphasia, confusion, non-specific T wave changes on the electrocardiogram, and seizures. The following adverse event has occurred in one patient receiving MYCOBUTIN, but an etiologic role for MYCOBUTIN has not been established: Pseudomembranous Colitis

When MYCOBUTIN was administered at doses from 1050 mg/day to 2400 mg/day, generalized arthralgia and uveitis were reported. These adverse experiences abated when MYCOBUTIN was discontinued.

**Laboratory Test Abnormalities**
The following table enumerates the changes in laboratory values that were considered as laboratory test abnormalities in the Phase III clinical trials.

<table>
<thead>
<tr>
<th>Laboratory Abnormality</th>
<th>MYCOBUTIN (n=566)%</th>
<th>PLACEBO (n=580)%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemistry:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Increased ALT (&gt; 150 U/L)</td>
<td>9</td>
<td>11</td>
</tr>
<tr>
<td>Increased AST (&gt; 150 U/L)</td>
<td>7</td>
<td>12</td>
</tr>
<tr>
<td>Increased Alkaline Phosphatase (&gt; 450 U/L)</td>
<td>&lt;1</td>
<td>3</td>
</tr>
<tr>
<td>Hematology:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neutropenia (ANC &lt; 750/mm$^3$)</td>
<td>25</td>
<td>20</td>
</tr>
<tr>
<td>Leukopenia (WBC &lt; 1500/mm$^3$)</td>
<td>17</td>
<td>16</td>
</tr>
<tr>
<td>Anemia (Hemoglobin &lt; 8.0 g/dL)</td>
<td>6</td>
<td>7</td>
</tr>
<tr>
<td>Thrombocytopenia</td>
<td>5</td>
<td>4</td>
</tr>
<tr>
<td>(Platelet count &lt; 50,000/mm$^3$)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Eosinophilia</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

The incidence of neutropenia in patients treated with MYCOBUTIN was significantly greater than in patients treated with placebo (p = 0.03). Although thrombocytopenia was not significantly more common among patients treated with MYCOBUTIN in these trials, MYCOBUTIN has been clearly linked to thrombocytopenia in rare cases. One patient developed thrombotic thrombocytopenic purpura, which was attributed to MYCOBUTIN.

Uveitis is rare when MYCOBUTIN is used as a single agent at 300 mg/day for prophylaxis of MAC in HIV-infected persons, even with the concomitant use of fluconazole and/or macrolide antibiotics. However, if higher doses of MYCOBUTIN are administered in combination with these agents, the incidence of uveitis is higher.

Patients who developed uveitis had mild to severe symptoms that resolved after treatment with corticosteroids and/or mydriatic eye drops; in some severe cases, however, resolution of symptoms occurred after several weeks.

When uveitis occurs, temporary discontinuance of MYCOBUTIN and ophthalmologic evaluation are recommended. In most mild cases, MYCOBUTIN may be restarted; however, if signs or symptoms recur, use of MYCOBUTIN should be discontinued.
Post-Marketing Adverse Reactions

Adverse reactions identified through post-marketing surveillance by system organ class (SOC) are listed below:

**Blood and lymphatic system disorders:** Pancytopenia, white blood cell disorders (including agranulocytosis, lymphopenia, granulocytopenia.

**Immune system disorders:** Hypersensitivity, bronchospasm.

**Eye disorders:** Corneal deposits.

**Hepato-biliary disorders:** Jaundice, hepatic enzyme increased.

Corneal deposits have been reported during routine ophthalmologic surveillance of some HIV-positive pediatric patients receiving MYCOBUTIN as part of a multiple drug regimen for MAC prophylaxis. The deposits are tiny, almost transparent, asymptomatic peripheral and central corneal deposits, and do not impair vision.

Anti-tuberculosis drug use may lead to the occurrence of drug reaction with eosinophilia and systemic symptoms (DRESS) as well as other SCARs such as SJS, TEN, and AGEP (see **WARNINGS, General**).

**SYMPTOMS AND TREATMENT OF OVERDOSAGE**

**Symptoms:**
No information is available on accidental overdose in humans.

**Treatment:**
While there is no experience in the treatment of overdose with MYCOBUTIN® (rifabutin), clinical experience with rifamycins suggest 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.

MYCOBUTIN is 85% protein bound, and distributed extensively into tissues ($V_{ss}$: 8 to 9 L/kg). As unchanged drug, MYCOBUTIN is not primarily excreted via the urinary route (less than 10%), therefore, neither hemodialysis nor forced diuresis is expected to enhance the systemic elimination of unchanged MYCOBUTIN from the body in a patient with MYCOBUTIN overdose.

For management of suspected overdosage, please contact your regional Poison Control Centre.
DOSAGE AND ADMINISTRATION

It is recommended that MYCOBUTIN (rifabutin) 300 mg be administered once daily with or without food. For those patients who experience nausea, vomiting or other gastrointestinal upsets, it may be useful to split the MYCOBUTIN dose in half (one 150 mg capsule) twice a day with food.

Limited pharmacokinetic data suggests that dose reductions may be required in patients with severe renal or hepatic impairment and in patients receiving concomitant treatment with certain drugs (see PRECAUTIONS).
PHARMACEUTICAL INFORMATION

DRUG SUBSTANCE

Proper Name: Rifabutin capsules USP


Structural Formula:

![Structural Formula Image]

Molecular Formula: C_{46}H_{62}N_{4}O_{11}

Molecular Weight: 847.02

Description:
Rifabutin is a red-violet powder soluble in chloroform and methanol, sparingly soluble in ethanol, and very slightly soluble in water (0.19 mg/mL).

Melting Point = 148°C - 156°C (with decomposition)
pKa Value = 6.9 in methanol/water (1/1, v/v)
Partition Coefficient = The partition coefficient of rifabutin between n-octanol and pH 6.8 buffer was determined to be > 100.
COMPOSITION

MYCOBUTIN® (rifabutin) is supplied as hard gelatin capsules having an opaque red-brown cap and body, imprinted with PHARMACIA & UPJOHN/ MYCOBUTIN, in white ink, each containing 150 mg rifabutin. The capsules also contain, as inactive ingredients, microcrystalline cellulose, magnesium stearate, red iron oxide, silica gel, sodium lauryl sulfate, titanium dioxide, and edible white ink.

STABILITY AND STORAGE RECOMMENDATIONS

Store at controlled room temperature, 15º-30ºC. Keep container tightly closed.

AVAILABILITY OF DOSAGE FORMS

MYCOBUTIN® (rifabutin) 150 mg
Bottles of 100 capsules

MICROBIOLOGY

Action
MYCOBUTIN® (rifabutin) inhibits DNA-dependant RNA polymerase in susceptible strains of Escherichia coli and Bacillus subtilis, but not in mammalian cells. In resistant strains of E. coli, rifabutin, like rifampin, did not inhibit this enzyme. It is not known whether rifabutin inhibits DNA-dependant RNA polymerase in Mycobacterium avium or in M. intracellulare which constitutes M. avium complex (MAC). Rifabutin inhibited incorporation of thymidine into DNA of rifampin-resistant M. tuberculosis suggesting that rifabutin may also inhibit DNA synthesis which may explain its activity against rifampin-resistant organisms.

In vitro Susceptibility Studies
There are no standardized methods for identification or susceptibility testing for M. avium complex. Various in vitro methodologies employ radiometric broth (7H12) or solid media. In general the radiometric broth (7H12) procedure has been used for isolation of mycobacteria with specific identification by DNA probes. Susceptibility testing is often performed using a radiometric broth method. However, neither method has been optimized for M. avium complex.

The MIC of rifabutin against M. avium in broth media were 2 to 4 times lower when polysorbate 80 (Tween 80) was present. This effect was attributed to an increase in the permeability of the cell envelope caused by Tween 80. It is recommended that susceptibility testing of rifabutin against M. avium be carried out in the absence of Tween 80 or other detergents.
**Mycobacterium avium Complex (MAC)**

Rifabutin has demonstrated *in vitro* activity against *M. avium* Complex (MAC) organisms isolated from both HIV positive and HIV negative people. These studies predated the widespread use of gene probe techniques to distinguish between the two organisms, although it is now known that the vast majority of isolates from MAC-infected, HIV positive people are *M. avium*, whereas in HIV negative people, about 40% of the isolates are *M. intracellulare*.

In one reported study on 100 MAC isolates from AIDS patients summarized in the table below, 83% had MIC\textsubscript{99} values of \(\leq 0.25\) \(\mu\text{g/mL}\), and 96% had MIC\textsubscript{99} values of \(\leq 0.5\) \(\mu\text{g/mL}\) when evaluated by a radiometric method 7H12 broth. In comparison, higher MICs were observed when the same isolates were tested in 7H11 by the agar proportion method. Twenty-three percent (23%) and 46%, of isolates had MIC\textsubscript{99} values of \(\leq 0.25\) and \(\leq 0.5\), respectively.

| In Vitro Minimum Inhibitory Concentrations of Rifabutin Against *Mycobacterium avium-intracellulare* Isolated from AIDS Patients |
|---|---|---|---|---|---|
| No. Isolates Tested (Method) | MIC (\(\mu\text{g/mL}\)) | \(\leq 0.25\) | 0.5 | 1.0 | 2.0 |
| 100(agar) | | 23(23) | 23(46) | 25(71) | 29(100) |
| 100(broth) | | 83(83) | 13(96) | 2(98) | 2(100) |

Other studies to determine the susceptibilities of MAC isolates from AIDS patients have yielded similar results.

The susceptibility to rifabutin of the initial positive MAC isolates from AIDS patients who developed MAC bacteremia during two large, rifabutin, multicenter placebo-controlled trials for prevention of MAC is as follows:

| Susceptibility to Rifabutin of the First Blood Isolates of MAC Recovered from Patients Receiving Prophylaxis with Rifabutin or Placebo |
|---|---|---|---|---|---|---|---|
| Prophylaxis | No. of Isolates Tested | Number of Isolates (cumulative %) ||
|  |  | MIC (\(\mu\text{g/mL}\)) | \(\leq 0.12\) | 0.25 | 0.5 | 1.0 | 2.0 | >2.0 |
| Placebo | 59 | | 4 (6.8) | 8 (20.3) | 20 (54.2) | 15 (79.7) | 11 (98.3) | 1 (100) |
| Rifabutin | 29 | | 3 (10.3) | 3 (20.7) | 16 (75.9) | 4 (89.7) | 3 (100) | 0 (100) |
For people who received placebo and rifabutin prophylaxis, 20.3% and 20.7% of the MAC isolates had MIC\textsubscript{99} values of \( \leq 0.25 \) \( \mu \)g/mL, 54.2% and 75.9% had MIC\textsubscript{99} values of \( \leq 0.5 \) \( \mu \)g/mL, and 79.7% and 89.7% had MIC\textsubscript{99} values of \( \leq 1.0 \) \( \mu \)g/mL, respectively, when evaluated by the radiometric 7H12 broth method. The distribution of MICs in the two groups did not differ significantly (Kruskal-Wallis p-value = 0.143).

In these studies there was no significant difference between the results of the broth and agar methods; the geometric means of MIC\textsubscript{99} values for all isolates were 0.64 and 0.78 \( \mu \)g/mL, respectively.

**Mycobacterium tuberculosis**

Rifabutin has *in vitro* activity against many strains of *M. tuberculosis* including some which are rifampin-resistant. In one study, utilizing the 7H12 broth dilution method, each of 20 rifampin-naive clinical isolates tested from Taiwan, had an MIC\textsubscript{99} value of \( \leq 0.125 \) \( \mu \)g/mL. The table below shows susceptibilities of 122 strains of *M. tuberculosis* isolates from rifampin-treated patients to rifampin and rifabutin in another study.

| Susceptibility of Rifampin-Resistant Strains of *M. tuberculosis* to Rifabutin+ |
|-------------------------------|-------------------------------|
| **No. Strains** | **RIFAMPIN MIC (µg/mL)** | **RIFABUTIN MIC (µg/mL)** |
| 1.0 | 5.0 | 10.0 | 0.5 | 1.0 | 2.0 |
| 122 | 0 | 12.3 | 24.6 | 27.0 | 41.8 | 63.9 |

+ Values are cumulative percentages

**Other Mycobacterial Species**

Rifabutin has shown activity against other mycobacterial species. Most strains of the following organisms were susceptible to concentrations of rifabutin of 0.5 \( \mu \)g/mL using the agar dilution method:

<table>
<thead>
<tr>
<th>Isolates Tested</th>
<th>No. of Isolates (Cumulative %)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MIC\textsubscript{99} (µg/mL)</td>
</tr>
<tr>
<td><em>Mycobacterium gordonae</em></td>
<td>100</td>
</tr>
<tr>
<td><em>Mycobacterium kansasii</em></td>
<td>100</td>
</tr>
<tr>
<td><em>Mycobacterium marinum</em></td>
<td>100</td>
</tr>
<tr>
<td><em>Mycobacterium terrae</em></td>
<td>71.4</td>
</tr>
<tr>
<td><em>Mycobacterium xenopi</em></td>
<td>100</td>
</tr>
</tbody>
</table>
Rifabutin was active against *Mycobacterium leprae* with MIC<sub>99</sub> values of 3.1 to 12.5 ng/mL when tested by the broth dilution method (BACTEC 460). It has an MIC<sub>99</sub> value of 0.016 to 2 μg/mL against *Mycobacterium phlei* when tested by the agar dilution method.

Rifabutin is rapidly taken up by macrophages and monocytes and has been shown to be active against phagocytized *M. avium intracellulare* and *M. tuberculosis*.

**Resistance**

*Mycobacterium avium* Complex:  
The cross-resistance relationship between rifampin and rifabutin appears to be partial. Rifampin and rifabutin MIC<sub>99</sub> values against 523 isolates of *M. avium* complex were determined utilizing the agar dilution method.

| SUSCEPTIBILITY OF *M. AVIUM* COMPLEX STRAINS TO RIFAMPIN AND RIFABUTIN | % of Strains Susceptible/Resistant to Different Concentrations of Rifabutin (μg/mL) |
|---|---|---|---|---|---|
| Susceptibility to Rifampin (μg/mL) | Number of Strains | Susceptible to 0.5 | Resistant to 0.5 only | Resistant to 1.0 | Resistant to 2.0 |
| Susceptible to 1.0 | 30 | 100.0 | 0.0 | 0.0 | 0.0 |
| Resistant to 1.0 only | 163 | 88.3 | 11.7 | 0.0 | 0.0 |
| Resistant to 5.0 | 105 | 38.0 | 57.1 | 2.9 | 2.0 |
| Resistant to 10.0 | 225 | 20.0 | 50.2 | 19.6 | 10.2 |
| TOTAL | 523 | 49.5 | 36.7 | 9.0 | 4.8 |

The majority of the strains (94%) were "naturally" resistant to rifampin (1.0 μg/mL or higher). Approximately 13% of those resistant strains were resistant to rifabutin in a concentration of ≥ 1.0 μg/mL.

*Mycobacterium tuberculosis*  
The cross-resistance between rifampin and rifabutin is frequently observed with *M. tuberculosis* isolates but it is not complete. Analysis of cross-resistance between rifampin and rifabutin among 302 *M. tuberculosis* strains indicate that all strains that were susceptible to 1.0 μg/mL of rifampin were also susceptible to 0.5 μg/mL of rifabutin. 46.7% of the strains that were resistant to 5.0 μg/mL of rifampin and 12.0% of the strains that were resistant to 10.0 μg/mL of rifampin were susceptible to rifabutin at concentrations of 0.5 μg/mL.
SUSCEPTIBILITY OF M. TUBERCULOSIS STRAINS TO RIFAMPIN AND RIFABUTIN

<table>
<thead>
<tr>
<th>Susceptibility to Rifampin (μg/mL)</th>
<th>Number of Strains</th>
<th>Susceptible to 0.5</th>
<th>Resistant to 0.5 only</th>
<th>Resistant to 1.0</th>
<th>Resistant to 2.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Susceptible to 1.0</td>
<td>180</td>
<td>100.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>Resistant to 1.0 only</td>
<td>15</td>
<td>100.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>Resistant to 5.0</td>
<td>15</td>
<td>46.7</td>
<td>33.3</td>
<td>13.3</td>
<td>6.7</td>
</tr>
<tr>
<td>Resistant to 10.0</td>
<td>92</td>
<td>12.0</td>
<td>14.1</td>
<td>27.2</td>
<td>45.7</td>
</tr>
<tr>
<td>TOTAL</td>
<td>302</td>
<td>70.5</td>
<td>6.0</td>
<td>8.9</td>
<td>14.6</td>
</tr>
</tbody>
</table>

In Vivo Studies

*Mycobacterium avium* Complex
Rifabutin and rifampin, each at doses of 6.25, 12.5 and 25 mg/kg orally, given six days per week for 12 weeks starting immediately after infection, were efficacious in ddY mice infected I.V. with *Mycobacterium avium-intracellulare* (MAI or MAC). Rifabutin was more effective than rifampin. In control mice, 96% developed macroscopic lung lesions. In mice treated with rifampin, 79% showed lung lesions, and in mice treated with rifabutin, 70% showed lung lesions. Advanced disease developed in 67% of control mice, 42% of rifampin-treated mice, and 31% of rifabutin-treated mice.

In another study, rifabutin and rifampin at doses of 20 and 80 mg/kg orally, given 6 days per week for 4 weeks, exhibited comparable efficacy in ddY mice infected with two different strains of MAI (N-260 and N-276).

Treatment, via the drinking water, was started 24 hours post-infection. Effectiveness was determined by counting the number of colony-forming units (CCU) in lungs and spleen homogenate plated on agar.

Rifabutin, at doses of 20 mg/kg three days per week, prevented *M. avium* infection in cyclosporine-treated rats. In a treatment study, at doses of 20 mg/kg five days per week, rifabutin reduced the number of mycobacteria in the spleen and liver by 200-fold in cyclosporine rats infected with *M. avium*.

Rifabutin was also effective in reducing the number of mycobacteria in tissues of thymectomized T-cell deficient mice infected with *M. avium*. Enhanced activity was observed in this animal
model when rifabutin was combined with either ethambutol or a combination of ethambutol, clofazimine, amikacin and ciprofloxacin.

Rifabutin was also effective in T-cell deficient mice infected with various strains of *M. intracellulare* or *M. tuberculosis* but not with the severely virulent *M. avium* 724. In this study, rifabutin was given in the drinking water at a dose of 40 mg/kg for 2 months beginning 20-40 days after infection.

Rifabutin and rifampin were also compared at a dose of 40 mg/kg daily for 120 days, alone and in combination with other antibiotics with anti-mycobacterial activity (all via drinking water) in T-cell deficient mice infected with AIDS-associated strains of *M. avium*.

Rifabutin was more active than the other drugs against most of the strains, and the combinations containing rifabutin exhibited greater efficacy than those containing rifampin.

When rifabutin at a dose of 10 mg/kg was added to a combination of amikacin (50 mg/kg intramuscularly) and clofazimine (20 mg/kg orally) and administered by gavage to MAI-infected beige mice, no additional improvement to the efficacy was observed.

The activity of rifabutin alone or in combination with clofazimine or ethambutol or both against acute and chronic experimental MAI infections were studied. Groups of male beige mice were infected intravenously with *M. intracellulare*, Strain 571-8 (serotype 8) isolated from a hospital patient. Drug treatment was rifabutin, given at 5, 10, or 20 mg/kg day alone, or at 10 mg/kg/day in combination with clofazimine (20 mg/kg/day) and/or ethambutol (125 mg/kg/day), via oral gavage. Administration of drugs was begun three weeks before infection, immediately after infection, or three weeks after infection, and continued seven days a week for 4-8 weeks following infection.

Given immediately after infection, rifabutin at 5 mg/kg/day alone showed no antibacterial effect. At 10 mg/kg/day, lung tissue, 6 weeks after infection, showed a significant (p<0.05) reduction in CCU. At 20 mg/kg/day, both lung and spleen showed significant reductions in CCU. Given three weeks after infection, 5 mg/kg/day of rifabutin showed no significant protection. At 10 mg/kg/day, significant reductions in CCU of the spleen and complete elimination of CCU in the lungs were observed.

When treatment at 10 mg/kg/day was started three weeks before infection, CCU counts were eliminated in the lungs at 4 weeks and in the spleen at 8 weeks after infection. The lungs of untreated animals were cleared spontaneously 8 weeks after infection. Thus, the effectiveness of rifabutin was especially pronounced when the drug was given prophylactically, three weeks before infection.

When combined therapy with rifabutin and clofazimine was initiated immediately after infection, complete sterilization of the tissues was achieved within 6 to 8 hours. Addition of ethambutol did not improve the efficacy. The same combinations given 3 weeks after infection was less effective.
The combination of rifabutin (10 mg/kg p.o.) with either clofazimine (20 mg/kg p.o.) or kanamycin (20 mg/kg s.c.) or both was more effective than any of these drugs given alone to *M. intracellulare*-infected mice (C57BL/6) strain for six days/week for 12 weeks starting 24 hours post-infection. The combination of rifabutin and clofazimine plus kanamycin was the most effective of the treatments tested giving complete protection against pulmonary lesions and significant reductions in bacterial counts in the lungs, liver and spleen.

Rifabutin in combination with clarithromycin was significantly more efficacious than clarithromycin alone in Beige mice infected with *M. avium* complex.

*Mycobacterium tuberculosis*

The efficacy of rifabutin (up to 10 mg/kg/day x three days) and rifampin (up to 25 mg/kg/day x ten days) was evaluated in CD1 (COBS) mice infected I.V. with *M. tuberculosis* H37 Rv. Drug treatment was started at either three days or ten days post-infection. Rifabutin was more effective than rifampin, although plasma levels of drug were much higher in rifampin-treated mice. In COBS rats, plasma levels of rifampin were higher than those of rifabutin but, except for liver, tissue levels of rifabutin were much higher than those observed with rifampin. This was especially true in the lung.

*Mycobacterium fortuitous*

In CD1 mice infected intravenously with *M. fortuitous*, rifampin was marginally active at 30 mg/kg/day, 5 days a week for 6 weeks, starting from 24 hours post-infection. Rifabutin exhibited significant activity at doses of 4 and 10 mg/kg/day.

*Mycobacterium leprae*

In a study in which rifabutin (0.00003 - 0.001%) or rifampin (0.01%) were administered in the diet to BALL/C mice infected with a rifampin-resistant of *M. leprae*, for approximately 6 months, rifabutin was found to be highly effective even at the lowest dose, while rifampin had no effect.

**Antibacterial**

Several non-Mycobacterial organisms were tested for *in vivo* susceptibility to rifabutin. In CD1 (COBS) mice infected with *Staphylococcus aureus*, single oral doses of rifabutin, administered 1-4 hours after infection, showed good antibiotic activity. The ED$_{50}$ value (on the basis of mortality) for rifabutin given to mice two hours after infection with *S. aureus* strain PV-1 was 0.4 mg/kg. Gram-negative bacterial infections were more susceptible to rifampin than to rifabutin, while *Streptococci* and *Diplococci* infections were more susceptible to rifabutin.

In the majority of the above *in vivo* studies, where doses equivalent to human doses were given, rifabutin was bacteriostatic, not bactericidal. Bactericidal activity was achieved when additional drugs were given.
PHARMACOLOGY

Clinical Pharmacology

Pharmacodynamics:
Over a dose range of 300 to 900 mg/day, administration of multiple, daily rifabutin doses did not elicit any serious/unexpected adverse events in 34 HIV positive patients. Doses higher than 900 mg/day produced chest discomfort, flu-like syndromes, lower back pain, skin discoloration, and gastrointestinal symptoms. However, a steep dose-related increase in the incidence of arthralgia, from 0% at 900 mg/day to 100% at 1200 mg/day, and the occurrence of uveitis at doses higher than 1800 mg/day was seen. No apparent effects on hematological or hepatic parameters were seen for doses up to 1200 mg/day, except mild leukopenia. Rifabutin in HIV positive patients was well tolerated up to 900 mg/day. The minimum effective dose and the optimal dose of rifabutin for prophylaxis, are not known.

Pharmacokinetics:
Rifabutin pharmacokinetics have been studied using daily doses ranging from 150 mg to 1200 mg (600 mg bid) in the pivotal pharmacokinetics and dose-tolerance studies involving both healthy normal volunteers and HIV positive patients, for periods of up to 28 days.

Absorption: In normal volunteers, a nominal therapeutic dose of 300 mg produces a mean $C_{\text{max}}$ of 375 ng/mL which is attained at approximately 3 hours post dose. At least 53% of the oral dose is absorbed.

Absolute Bioavailability: The mean (± SD) absolute bioavailability (F) of rifabutin from the capsule formulation in 15 early symptomatic HIV positive patients was found to be 20% (± 16%, n=5) following a single dose (first day), and was 12% (± 5%, n=7), after daily dosing for 4 weeks. This change was not statistically significant. The possible decrease in absolute bioavailability on day 28 may be due to autoinduction of rifabutin metabolism.

Effect of Food on Oral Absorption: A single study simultaneously examined the absorption of rifabutin from the capsule dosage form relative to a solution and also assessed the effect of food. A single 150 mg dose was administered to 12 male volunteers in a crossover manner. The administration of rifabutin with a high fat content meal decreased mean $C_{\text{max}}$ by 17% (156.2 vs. 187.9 ng/mL), increased mean $T_{\text{max}}$ from 3.0 to 5.4 hours and increased mean percent of dose excreted unchanged in urine by 26% (11.4% vs 9.1%). Only the changes in mean $T_{\text{max}}$ and the percent of dose excreted unchanged in urine were statistically significant. The data indicate that a high-fat content meal decreased the rate of absorption and increased urinary excretion of rifabutin with no change in extent of absorption. The relative bioavailability of the capsule with respect to the solution was estimated to be 85%.

Distribution: In a radio-tracer intravenous study for unchanged rifabutin, a triexponential concentration versus time disposition profile was observed. The fastest distributive phase ($\lambda_1$) has a half-life of 9-15 minutes. The half-lives associated with the slow distributive phase ($\lambda_2$) and terminal elimination ($\lambda_z$) seem comparable to those following oral dosing, after which absorption,
distribution and elimination phases are clearly observed. The half-life of the slow distributive phase (t1/2,λ2) was estimated to be 3.5 to 4.5 hours in healthy males, and 1.7-3.3 hours in early symptomatic HIV patients, thus suggesting a lack of disease effects on the slow distributive phase. The volume of distribution at steady-state (Vss) was determined to be 8-9 L/kg in early symptomatic HIV patients. This estimate is approximately 13 to 15-fold larger than the total body water (TBW: 0.6 L/kg).

**Tissue Distribution**: A study in 4 surgical patients provided limited but useful information regarding tissue uptake. Rifabutin (measured as total antimicrobial activity) was found in all tissues studied, e.g. lung, gall bladder, ileum, jejunum, and muscle. Tissue concentrations were several times higher than those found in plasma. The lung to plasma ratio ranged from 1.4 to 8.6 at about 6 hours and 5.6 to 6.8 at 12 hours post-oral dosing. Even when plasma levels were undetectable, measurable levels of rifabutin persisted in lung up to 48 hours. The partitioning of rifabutin into ileum, jejunum, and bile appeared higher than the lung, and supports biliary excretion as a significant route of elimination. Distribution of rifabutin was lowest in the muscle, with a muscle:plasma ratio of < 1.

**Plasma Protein Binding**: The extent of *in vitro* protein binding of 14C-rifabutin in fresh human plasma was assessed by equilibrium dialysis. Approximately 90% of rifabutin was bound to plasma proteins over a concentration range of 0.1-10.0 μg/mL. The binding decreased to approximately 85% at higher concentrations of 20-100 μg/mL. At a concentration of 100 ng/mL, rifabutin was 68.3 ± 1.9% bound to human serum albumin (HSA) and 19.6 ± 3.1% bound to α-acid glycoprotein (AAG). These results suggest that rifabutin is predominantly bound to HSA. The free fraction (fu) in plasma is approximately 0.15 (95% CI: 0.138-0.158) and is independent of drug concentration within the concentration range observed following the standard 300 mg dose. Protein binding was assessed at 20 μg/mL by equilibrium dialysis in healthy subjects, the elderly, patients with alcoholic liver disease and renal insufficiency. Mean % bound averaged 95% in healthy subjects, 91% in the elderly, 90% in patients with alcoholic liver disease, and 92-94% in patients with renal insufficiency. Protein binding was therefore > 90% for all cases.

**Metabolism**: Investigations elucidating the metabolic profile of rifabutin suggest extensive biotransformation. Urinary metabolism studies, using mass spectrometry, 1H-NMR spectrometry, and HPLC, have shown that rifabutin is metabolized to more than 20 different metabolites. Five among them have been identified: 25-O-deacetyl rifabutin (M1), 31-OH-rifabutin (M2), 32-OH rifabutin, 32-OH-25-O-deacetyl rifabutin, and 25-O-deacetyl rifabutin-N-oxide. The M1 derivative was determined to be as equally active as rifabutin microbiologically. The M2 derivative seems to possess approximately 1/16 of the activity of rifabutin. The evidence of rifabutin autoinduction comes from estimates of CLT in early symptomatic HIV positive patients administrated tracer intravenous 14C-rifabutin doses. The mean CLT was estimated to be 10.2 L/h on day 1 and 18.5 L/h on day 28, following 4 weeks of daily dosing, an increase of 80%. This estimate of CLT was not confounded by absolute bioavailability (F).

**Excretion**: Urinary and faecal excretion are the two major routes of elimination for rifabutin. The excretion characteristics of rifabutin and its metabolites have been assessed in a radiolabelled study. Following an oral dose (270 mg) as 14C-rifabutin solution (100 μCi) to 3 healthy
volunteers, 53% of the dose was recovered in the urine over a 120 hour period and 30% in the faeces. Approximately 8% of the administered radioactive dose was recovered in urine as unchanged parent drug. After administration of rifabutin in the capsule dosage form, 5.17% was eliminated through bile, and 5.8% was eliminated in urine as unchanged drug.

**Animal Safety Pharmacology**

**Neuropharmacology**
Oral doses of 200 mg/kg rifabutin in the mouse and 100 mg/kg in the rat caused CNS depression lasting up to six hours post dose. Rifabutin (50 mg/kg) did not antagonize amphetamine, pentylentetrazol or reserpine in the mouse. Rifabutin did not affect neuromuscular coordination (rotorod) or conditioned avoidance response in the rat at 50 mg/kg. There were no consistent effects on body temperature in mice, rats or dogs treated with rifabutin.

**Cardiovascular**
In the rat, single intravenous doses of up to 36 mg/kg or oral doses of up to 200 mg/kg daily for four days did not affect blood pressure or heart rate or the responses of these parameters to several autocoids. There was an increase of about 50% in respiratory rate beginning about 100 minutes following an intraduodenal dose of 50 mg/kg in the anaesthetized dog. There were, however, no other significant changes in cardiovascular or respiratory system parameters.

**Gastrointestinal**
Oral doses of up to 20 mg/kg rifabutin did not affect gastric emptying rate in the rat.

**Genitourinary**
Rifabutin, at oral doses of up to 100 mg/kg, did not affect urinary volume, pH or electrolytes in the rat.

**Immunopharmacology**
The effects of rifabutin and rifampin on humoral and cell-mediated immunity were determined in mice and guinea pigs. Neither rifabutin nor rifampin affected humoral antibody response to SRBC in the mouse at intraperitoneal doses of up to 300 mg/kg and 150 mg/kg, respectively. Although there was no effect on delayed hypersensitivity (DH) when give oral doses of 50 mg/kg in the mouse, both drugs did reduce DH when given intraperitoneal dose of 150 mg/kg. Also, both drugs reduced DH to tuberculin in the mouse but in guinea pigs, rifampin decreased DH to tuberculin whereas rifabutin had no effect. These studies, however, were not extensive and it is difficult to assess their significance.

A study was also conducted to determine the effect of rifabutin on the phagocytic and bactericidal activities of splenic macrophages in the mouse. The mice were given 300 mg/kg rifabutin daily for 30 days and macrophage function was determined for up to 30 days following cessation of treatment.

The results indicated that rifabutin and/or its active metabolites persist in the lungs, liver, and spleen at measurable concentrations for up to 15 days post-treatment and that the phagocytic and
bactericidal activities of splenic macrophages as measured using *L. monocytogenes* as a target were not affected despite the lipidotic effect which rifabutin has on macrophages.

Rifabutin and rifampin were found to inhibit random migration but not chemotaxis of human PMNL. In a study comparing the effects of rifabutin and rifampin on rabbit PMNL, it was found that rifabutin reduced cellular respiration at concentrations of 50 and 100 μg/mL but had no effect on their phagocytic or bactericidal functions. Rifampin inhibited phagocytic function at 100 μg/mL. Since the concentrations of rifabutin used in this study were much higher than those found in plasma of patients treated with rifabutin, it is unlikely that rifabutin will affect PMNL function clinically.

Also, rifabutin did not affect the ability of human PMNL to phagocytize *S. aureus in vitro* and did not interfere with superoxide formation by human neutrophils *in vitro*.

**TOXICOLOGY**

The acute oral toxicity of rifabutin in rats, given single oral doses up to 5 g/kg, or in beagle dogs and cynomolgus monkeys, given 2 and 4 g/kg rifabutin was low with no mortality. The oral LD_{50} in mice was 4.8 g/kg for males and 3.3 g/kg for females. The pharmacotoxic effects observed were relatively minor and consisted of decreased spontaneous activity in mice and rats and gastrointestinal disturbances, i.e. emesis and/or diarrhea in dogs and monkeys. These results are consistent with the antimicrobial properties of the drug.

Subchronic 13-week studies were performed in mice (0, 12.5, 25, 50 and 100 mg/kg/day; 0, 50, 100 and 200 mg/kg/day), rats (0, 25, 50, 100 and 200 mg/kg/day; 0, 25, 50 and 100 mg/kg/day on alternate days; 0, 50, 100 and 200 mg/kg/day), cynomolgus monkeys (0, 10, 20, 40 and 80 mg/kg/day) and baboons (0, 10, 40, and 80 mg/kg/day). No drug-related mortality occurred in all species.

In mice, only slight functional hepatic changes were seen. Body weight gain was reduced at all doses in male mice in one study, and relative spleen weights were increased in both sexes at all doses in another mouse study. Some liver inflammation was seen in high dose female mice.

In rats, multinucleated hepatocytes were observed both following daily administration of rifabutin or administration on alternate days. Ultrastructurally, the multiple nuclei and nucleoli closely resemble those in mononucleated cells. This was not seen in any other species. Rifabutin induced slight dose-related decreases in RBC and related parameters, hyperplasia of the gastric mucosa, increased spleen and liver weights, reduced body weight gains in high-dose rats, decreased spermatogenesis in male rats, and uterine hyperplasia in female rats.

In cynomolgus monkeys, the daily oral administration of rifabutin at doses up to 80 mg/kg/day for 13 weeks had minimal effects, the most important of which was an increased bilirubin and decreased LAP at the 80 mg/kg/day dose. Slight fatty infiltration of the liver was seen in cynomolgus monkeys and in baboons at all doses. Decreased testes weight, beginning at a dose of 10 mg/kg/day, was also noted in baboons.
Twelve month toxicity studies were carried out in mice (0,8,32 and 128 mg/kg/day), rats (0,10,28 and 80 mg/kg/day) and cynomolgus monkeys (0,8,24 and 72 mg/kg/day). These results were in agreement with the previous 13-week studies. No drug-related mortality was noted in all species. In mice, body weight gains were increased in high dose females. In males there was a dose-related increase in plasma cholesterol, with bilirubin and reticulocytes increased at the high dose. High dose females had increased ALT. At 128 mg/kg/day in males, testes weight decreased while liver and lung weights increased. In high-dose females, adrenal, lung, liver and spleen weights increased.

There was a dose-related increase at the 32 and 128 mg/kg/day dose in the incidence of Heinz bodies in reticulocytes.

Both male and female rats showed increased plasma bilirubin at the high dose, and high dose males had increased ALT and AST. Multinucleated hepatocytes were seen in males at all dose groups and in females at 28 and 80 mg/kg/day. Liver and spleen weights increased in mid-dose males and in both sexes at the high dose. High dose males also had decreased testes weights and atrophy of seminiferous tubules. Hyperplasia of the gastric mucosa occurred in both sexes given 80 mg/kg/day.

Male monkeys experienced occasional episodes of emesis at 24 and 72 mg/kg/day. High dose animals showed an increase in plasma bilirubin and triglycerides with lipid deposits in hepatocytes and increased liver weight.

Multinucleated hepatocytes were seen in rats only; liver hypertrophy with functional effects (increased ALT and/or AST, cholesterol, triglycerides and bilirubin values) was observed in all species. Minor changes included slight decreases in RBC and related parameters, sclerosis/hyperplasia of the glandular mucosa of the stomach, and atrophy of the testes in rats.

The significance of the multinucleated hepatocytes found in rats given repeated oral doses of rifabutin is not clear. They have been reported in association with various physiological, nutritional and disease states, and after administration of several structurally and pharmacologically diverse chemical and drug substances. In any case, the presence of multinucleated hepatocytes must be considered rat specific, as they were not found in the other species.

There was no increase of the multinucleated hepatocytes with time and there was no effect in their life span. As expected, this finding is not preneoplastic, as confirmed by the oncogenicity study in rats.

Reproductive Toxicology
Fertility studies have been performed in male and female rats given rifabutin orally at 0,10,40 and 160 mg/kg/day. Fertility was impaired in male rats given 160 mg/kg/day (32 times the recommended human daily dose). The 160 mg/kg/day dose resulted in a reduced number of
implants and offspring. In the 160 mg/kg/day treated females, there was an 18% decrease in the number of fetuses per dam associated with a 5-fold increase in post implantation losses. At 40 mg/kg/day (8 times the human dose), rifabutin caused an increase in skeletal variants. Peri- and post-natal studies have been performed in rats given rifabutin orally at 0, 12.5, 50 and 200 mg/kg/day. At 200 mg/kg/day (40 times the human dose), there was a decrease in fetal viability.

**Teratogenicity**
Embryotoxicity studies have been performed in rats and rabbits given rifabutin orally at 0, 20, 40 and 80 mg/kg/day. At all doses and in both species no teratogenic effects were seen. In rats, at 40 and 80 mg/kg/day, and in rabbits, at 80 mg/kg/day (16 times the human dose), rifabutin caused maternotoxicity and an increase in fetal minor skeletal anomalies.

**Mutagenicity**
Rifabutin (0.125 - 1000 μg/mL) was not mutagenic in the bacterial point mutation assay (Ames Test) using both rifabutin-susceptible and resistant strains of *Salmonella typhimurium*. Rifabutin was not mutagenic in *Schizosaccharomyces pombe P*1 and was not genotoxic in V-79 Chinese hamster cells, human lymphocytes *in vitro*, or mouse bone marrow cells *in vivo*.

**Carcinogenicity**
Long term carcinogenicity studies were conducted with rifabutin in mice (0, 20, 60 and 180/100 mg/kg/day) and rats (0, 15, 30, and 60 mg/kg/day). Rifabutin was not carcinogenic in mice at doses up to 180 mg/kg/day, or approximately 36 times the recommended human daily dose. The high dose was reduced from 180 mg/kg/day to 100 mg/kg/day on Day 561, due to excessive mortality in males.

This high-dose treatment group was terminated after 21 months of treatment. Necropsy and histopathological findings indicated that deaths were caused by myocardial vacuolation and fibrosis. Rifabutin was not carcinogenic in the rat at doses up to 60 mg/kg/day, about 12 times the recommended human daily dose. There was a significant increase in the incidence of incidental tumours in the liver of the high dose females but most of these were adenomas. In summary, rifabutin was not carcinogenic in either mice or rats when administered in the diet for up to two years at the MTD.
REFERENCES


Read this carefully before you start taking MYCOBUTIN. Read it again every time you get a refill. This leaflet is a summary. It will not tell you everything about this drug. Talk to your healthcare professional about your medical condition and treatment. Ask whether there is any new information about MYCOBUTIN.

What is MYCOBUTIN used for?
- MYCOBUTIN is given to patients with advanced HIV infection.
- It is used to prevent serious disease caused by germs a group of (bacteria) called Mycobacterium avium complex (MAC).
- MAC may cause a lung disease like tuberculosis (TB).

Antibacterial drugs like MYCOBUTIN treat only bacterial infections. They do not treat viral infections such as the common cold. Although you may feel better early in treatment, MYCOBUTIN should be used exactly as directed. Misuse or overuse of MYCOBUTIN could lead to the growth of bacteria that will not be killed by MYCOBUTIN (resistance). This means that MYCOBUTIN may not work for you in the future. Do not share your medicine.

How does MYCOBUTIN work?
- MYCOBUTIN is an antibiotic drug that treats infections caused by germs (bacteria) called mycobacteria.
- Mycobacteria are germs (bacteria) that cannot be killed using other antibiotics.

What are the ingredients in MYCOBUTIN?

Medicinal ingredients: Rifabutin

Non-medicinal ingredients: Edible white ink, magnesium stearate, microcrystalline cellulose, red iron oxide, silica gel, sodium lauryl sulfate and titanium dioxide.

MYCOBUTIN comes in the following dosage form:
It comes as a 150 mg capsule (pill). The capsule is dark red with the name of the drug ‘PHARMACIA & UPJOHN/ MYCOBUTIN’ written in white letters.

Do not use MYCOBUTIN if:
- If you are allergic (hypersensitive) to this drug.
- If you are allergic to any of the other ingredients in MYCOBUTIN (Read also “What are the ingredients in MYCOBUTIN” above.)
- If you are allergic to any drug containing rifamycin.
To help avoid side effects and make sure you take the drug properly, talk to your healthcare professional before you take MYCOBUTIN. Talk about any health conditions or problems you may have, including if you:

- Have severe kidney or liver problems. (You may need a lower dose of the drug.)
- Get mild to severe diarrhea. (This could be a serious infection called *C. difficile*.)
- Get eye problems like
  - pain
  - blurred vision
  - floating dark spots in your vision
  - redness
  - sensitivity to light.
  These could be signs of a problem called uveitis (*inflammation in the inside of the eye.*)
- Are pregnant or thinking of getting pregnant.
- If you are breast feeding your baby or thinking of breast feeding your baby.
- Have ever had an allergic reaction to any medicines.

Your health care professional will test to be sure you do not have active TB or another infection.

If you are taking MYCOBUTIN with other medicines that treat infection, you may also be asked to have regular eye exams.

**Other warnings you should know about**

Taking MYCOBUTIN with anti-tuberculosis drugs can increase the risk of side effects to your skin. You may develop a serious skin condition called Severe Cutaneous Adverse Reactions (SCARs). **Speak to your doctor immediately** if you notice any skin rashes. The following skin problems may develop:

- **Stevens Johnson Syndrome (SJS) and Toxic Epidermal Necrolysis (TEN).** SJS and TEN are severe skin rashes. Typical symptoms include peeling skin, fever, body aches, blisters, sores, and a flat red rash
- **Drug Reaction with Eosinophilia and Systemic Symptoms (DRESS).** DRESS is serious skin reaction that may affect more than one or more organs. Typical symptoms include fever, severe rash, peeling skin, flu-like feeling, yellow skin or eyes, shortness of breath
- **Acute Generalized Exanthematous Pustulosis (AGEP).** AGEP is a serious skin reaction that appears suddenly. Typical symptoms include a rash of small red and white bumps and fever.

Tell your healthcare professional about all the medicines you take, even the ones you get without a prescription (example vitamins).

**The following drugs may interact with MYCOBUTIN:**

- Drugs to treat diabetes
- Painkillers like acetylsalicylic acid (ASA) and others
- Narcotics, including methadone
- Blood thinners (anticoagulants) like warfarin
- Steroids (to treat inflammation or allergy) like prednisolone
- Drugs to suppress the immune system like cyclosporine (ciclosporin) and tacrolimus
• Drugs for heart conditions like quinidine or digitalis (digoxin is okay)
• Drugs to treat skin infections or pneumonia like dapsone
• Drugs to treat epilepsy or seizures like phenytoin (Dilantin)
• Drugs to treat fungal infections like fluconazole, itraconazole, posaconazole, voriconazole, ketoconazole and miconazole
• Drugs to treat viral infections like indinavir, saquinavir, ritonavir or amprenavir, fosamprenavir/ritonavir, lopinavir/ritonavir, tipranavir/ritonavir, bictegravir, doravirine and rilpivirine
• The anti-HIV drug called delviradine
• The antibiotic drug called clarithromycin
• Birth control pills that have ethinyl estradiol and/or norethindrone. (These drugs may not be as effective if you are on MYCOBUTIN and your healthcare professional may recommend a different kind or an additional kind of birth control.)

**How to take MYCOBUTIN:**

• It is very important to take MYCOBUTIN exactly as your healthcare professional has told you to. Never change the dose by yourself. Do not stop taking MYCOBUTIN unless your healthcare professional tells you to because your infection could return.
• Take this drug at the same time every day.
• Swallow the MYCOBUTIN capsules (pills) whole with a drink of water.

**Usual dose:**

• Take two MYCOBUTIN capsules (pills) by mouth once a day.
• If you have nausea, feel sick to your stomach, or throw up you can take one capsule (pill) by mouth twice a day. In this case, take with food.

**Overdose:**

If you think you have taken too much MYCOBUTIN, contact your healthcare professional, hospital emergency department or regional Poison Control Centre immediately, even if there are no symptoms.

**Missed Dose:**

If you forget to take a dose, take it as soon as you remember, unless it is time for your next dose. Do not take a double dose to make up for a missed dose.

**What are the possible side effects from using MYCOBUTIN?**

Some of the possible side effects are listed below. These are not all the possible side effects you may get when taking MYCOBUTIN. If you have side effects not listed here, contact your healthcare professional.

The most common side effect of MYCOBUTIN is a change in the colour of your urine from yellow to brown-orange. Colour changes like this may also affect bowel movements, spit, sweat, tears or skin. If you wear contact lenses, they may be permanently stained.
Other side effects include:

- A drop in the number of white blood cells (these fight infections).
- Skin rashes
- Stomach problems like upset stomach, burping, passing gas, nausea (feeling sick to your stomach), vomiting (throwing up) and belly pain.
- Ask your healthcare professional about how to tell if you have *Mycobacterium avium* complex (MAC) disease and/or TB.
- Let your healthcare professional know if you get any of these side effects and symptoms.

If you notice symptoms such as severe diarrhea (bloody or watery) with or without fever, abdominal (belly) pain, or tenderness, you may have *Clostridium difficile* colitis (C. difficile – bowel inflammation). If this occurs, stop taking MYCOBUTIN and contact your healthcare professional immediately.

<table>
<thead>
<tr>
<th>Serious side effects and what to do about them</th>
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<tbody>
<tr>
<td><strong>Symptom / effect</strong></td>
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<tr>
<td></td>
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<tr>
<td><strong>COMMON</strong></td>
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<tr>
<td>Diarrhea</td>
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<tr>
<td><strong>UNCOMMON</strong></td>
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<tr>
<td>● Sudden wheeziness or problems breathing</td>
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<tr>
<td>● Pain in your chest</td>
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<tr>
<td>● Swelling of your eyelids, face or lips</td>
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<tr>
<td>● Rash or itching (especially over your whole body)</td>
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<tr>
<td><strong>RARE</strong></td>
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<tr>
<td>● Muscle aches</td>
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<tr>
<td>● Uveitis: symptoms can be eye pain, blurred vision, floating dark spots, eye redness and sensitivity to light</td>
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<tr>
<td>● Pain in a number of joints</td>
</tr>
<tr>
<td>● Severe cutaneous adverse reactions (SCAR) (severe)</td>
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skin reactions that may also affect other organs):
- skin peeling, scaling, or blistering (with or without pus) which may also affect your eyes, mouth, nose or genitals, itching, severe rash, bumps under the skin, skin pain, skin color changes (redness, yellowing, purplish)
- swelling and redness of eyes or face
- flu-like feeling, fever, chills, body aches, swollen glands, cough
- shortness of breath, chest pain or discomfort

If you have a symptom or side effect that is not listed here and that bothers you or becomes bad enough to interfere with your daily activities, talk with your healthcare professional.

**Reporting Side Effects**
You can report any suspected side effects associated with the use of health products to Health Canada by:

- Visiting the Web page on Adverse Reaction Reporting ([https://www.canada.ca/en/health-canada/services/drugs-health-products/medeffect-canada/adverse-reaction-reporting.html](https://www.canada.ca/en/health-canada/services/drugs-health-products/medeffect-canada/adverse-reaction-reporting.html) for information on how to report online, by mail or by fax; or
- Calling toll-free at 1-866-234-2345.

*NOTE: Contact your health professional if you need information about how to manage your side effects. The Canada Vigilance Program does not provide medical advice.*

**Storage:**

- Store your bottle of capsules at room temperature (15º to 30ºC).
- Keep the container tightly closed.
- Keep this medication where children cannot reach it or see it.

**If you want more information about MYCOBUTIN:**

- Talk to your healthcare professional.
- Find the full product monograph that is prepared for healthcare professionals and includes this Patient Medication Information by visiting the Health Canada website, the manufacturer’s website http://www.pfizer.ca or by calling 1-800-463-6001.