

HIGHLIGHTS OF PRESCRIBING INFORMATION

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

BOSENTAN tablets, for oral use

Initial U.S. Approval: 2001

WARNING: RISKS OF HEPATOTOXICITY and EMBRYO-FETAL TOXICITY
See full prescribing information for complete boxed warning.
Bosentan tablets are available only through a restricted distribution program called the Bosentan REMS Program because of these risks (5.3):
Elevations of liver aminotransferases (ALT, AST) and liver failure have been reported with bosentan (5.1).
• Measure liver aminotransferases prior to initiation of treatment and then monthly (2.1, 5.1).
• Discontinue bosentan if aminotransferase elevations are accompanied by signs or symptoms of liver dysfunction or injury or increases in bilirubin greater than or equal to 2 x ULN (2.4, 5.1).
Based on animal data, bosentan is likely to cause major birth defects if used during pregnancy (4.1, 5.2, 8.1).
• Must exclude pregnancy before and during treatment (2.1, 4.1, 8.1).
• To prevent pregnancy, females of reproductive potential must use two reliable forms of contraception during treatment and for one month after stopping bosentan (4.1, 5.2, 8.1).

----- INDICATIONS AND USAGE -----
Bosentan is an endothelin receptor antagonist indicated for the treatment of pulmonary arterial hypertension (PAH) (WHO Group 1):
• in adults to improve exercise ability and to decrease clinical worsening. Studies establishing effectiveness included predominantly patients with WHO Functional Class II-IV symptoms and etiologies of idiopathic or heritable PAH (60%), PAH associated with connective tissue diseases (21%), and PAH associated with congenital heart disease with left-to-right shunts (18%) (1).
----- DOSAGE AND ADMINISTRATION -----
• Patients older than 12 years of age, initiate at 62.5 mg orally twice daily; if tolerated, increase to 125 mg twice daily; if tolerated, increase to 125 mg orally twice daily after 4 weeks (2,3).

FULL PRESCRIBING INFORMATION: CONTENTS
WARNING: RISKS OF HEPATOTOXICITY AND EMBRYO-FETAL TOXICITY
INDICATIONS AND USAGE
2 DOSAGE AND ADMINISTRATION
2.1 Required Monitoring
2.2 Recommended Dosage
2.3 Administration
2.4 Dosage Adjustments for Aminotransferase Elevations
2.5 Use with Ritonavir
2.6 Use in Patients with Pre-existing Hepatic Impairment
3 DOSAGE FORMS AND STRENGTHS
CONTRAINDICATIONS
4.1 Pregnancy
4.2 Use with Cyclosporine A
4.3 Use with Glyburide
4.4 Hypersensitivity
5 WARNINGS AND PRECAUTIONS
5.1 Hepatotoxicity
5.2 Embryo-fetal Toxicity
5.3 Prescribing and Distribution Program for Bosentan Tablets
5.4 Fluid Retention
5.5 Pulmonary Venous-Occlusive Disease
5.6 Decreased Sperm Counts
5.7 Decreases in Hemoglobin and Hematocrit
6 ADVERSE REACTIONS
6.1 Clinical Trials Experience
6.2 Postmarketing Experience

7 DRUG INTERACTIONS
7.1 Cyclochrome P450 Drug Interactions
7.2 Hormonal Contraceptives
8 USE IN SPECIFIC POPULATIONS
8.1 Pregnancy
8.2 Lactation
8.3 Females and Males of Reproductive Potential
8.4 Pediatric Use
8.5 Geriatric Use
8.6 Hepatic Impairment
8.7 Renal Impairment
10 OVERDOSAGE
11 DESCRIPTION
12 CLINICAL PHARMACOLOGY
12.1 Mechanism of Action
12.2 Pharmacokinetics
12.3 Pharmacokinetics
13 NONCLINICAL TOXICOLOGY
13.1 Carcinogenicity, Mutagenesis, Impairment of Fertility
14 CLINICAL STUDIES
14.1 Pulmonary Arterial Hypertension
14.2 Lack of Benefit in Congestive Heart Failure
16 HOW SUPPLIED/STORAGE AND HANDLING
17 PATIENT COUNSELING INFORMATION
*Sections or subsections omitted from the full prescribing information are not listed.

18 REFERENCES
6.2 Postmarketing Experience

FULL PRESCRIBING INFORMATION
WARNING: RISKS OF HEPATOTOXICITY AND EMBRYO-FETAL TOXICITY
Because of the risks of hepatotoxicity and birth defects, bosentan tablets are available only through a restricted program called the Bosentan REMS Program. The Bosentan REMS Program is a component of the Bosentan Risk Evaluation and Mitigation Strategy (REMS). Under the Bosentan REMS, prescribers, patients, and pharmacies must enroll in the program (see *Warnings and Precautions* (5.3)).
Hepatotoxicity
In clinical studies, bosentan caused at least 3-fold upper limit of normal (ULN) elevation of liver aminotransferases (ALT and AST) in about 11% of patients, accompanied by elevated bilirubin in a small number of cases. Because these changes are a marker for potential serious hepatotoxicity, serum aminotransferase levels must be measured prior to initiation of treatment and then monthly (see *Dosage and Administration* (2.4), *Warnings and Precautions* (5.1)). In the postmarketing period, in the setting of close monitoring, rare cases of unexplained hepatic cirrhosis were reported after prolonged (greater than 12 months) therapy with bosentan in patients with multiple comorbidities and drug therapies. There have also been reports of liver failure. The contribution of bosentan in these cases could not be excluded.
In at least one case, the initial presentation (after greater than 20 months of treatment) included pronounced elevations in aminotransferases and bilirubin levels accompanied by non-specific symptoms, all of which resolved slowly over time after discontinuation of bosentan tablets. This case reinforces the importance of strict adherence to the monthly monitoring schedule for the duration of treatment and the treatment algorithm, which includes stopping bosentan with a rise of aminotransferases accompanied by signs or symptoms of liver dysfunction (see *Dosage and Administration* (2.4)). Elevations in aminotransferases require close attention (see *Dosage and Administration* (2.4)). Bosentan should generally be avoided in patients with elevated aminotransferases (greater than 3 x ULN) at baseline because monitoring for hepatotoxicity may be more difficult. If liver aminotransferase elevations are accompanied by clinical symptoms of hepatotoxicity (such as nausea, vomiting, fever, abdominal pain, jaundice, or unusual lethargy or fatigue) or increases in bilirubin greater than or equal to 2 x ULN, treatment with bosentan should be stopped. There is no experience with the reintroduction of bosentan in these circumstances.

Embryo-fetal Toxicity
Bosentan is likely to cause major birth defects if used by pregnant females based on animal data (see *Warnings and Precautions* (4.1, 5.2, 8.1)). Because of these risks, bosentan tablets should be excluded before the start of treatment with bosentan tablets. Throughout treatment and for one month after stopping bosentan tablets, females of reproductive potential must use two reliable methods of contraception unless the patient has an intrauterine device (IUD) or tubal sterilization, in which case no other contraception is needed. Hormonal contraceptives, including oral, injectable, transdermal, and implantable contraceptives should not be used as the sole means of contraception because these may not be effective in patients receiving bosentan tablets (see *Drug Interactions* (7.2)). Obtain monthly pregnancy tests.

1. INDICATIONS AND USAGE
Bosentan tablets are indicated for the treatment of pulmonary arterial hypertension (PAH) (WHO Group 1):
• in adults to improve exercise ability and to decrease clinical worsening. Studies establishing effectiveness included predominantly patients with WHO Functional Class II-IV symptoms and etiologies of idiopathic or heritable PAH (60%), PAH associated with connective tissue diseases (21%), and PAH associated with congenital heart disease with left-to-right shunts (18%) (see *Clinical Studies* (14.1)).
2. DOSAGE AND ADMINISTRATION
2.1 Required Monitoring
Healthcare professionals who prescribe bosentan tablets must enroll in the Bosentan REMS Program and must comply with the required monitoring to minimize the risks associated with bosentan tablets (see *Warnings and Precautions* (5.3)). Obtain a pregnancy test in females of reproductive potential prior to bosentan treatment, monthly during treatment and one month after stopping bosentan. Initiate treatment with bosentan in females of reproductive potential only after a negative pregnancy test (see *Boxed Warning, Contraindications* (4.1), *Warnings and Precautions* (5.3), *Use in Specific Populations* (8.1, 8.3)). Measure liver aminotransferase levels prior to initiation of treatment and then monthly (see *Warnings and Precautions* (5.1)).
2.2 Recommended Dosage
Administer bosentan tablets orally following the dosing recommendations in Table 1. Doses above 125 mg twice daily did not appear to confer additional benefit sufficient to offset the increased risk of hepatotoxicity.

Table 1. Dosing Recommendations

	Initial 4 weeks	Maintenance (after 4 weeks)
Patients >12 years of age and >40 kg	62.5 mg twice daily	125 mg twice daily
Patients >12 years of age and <40 kg	62.5 mg twice daily	62.5 mg twice daily

2.3 Administration
Bosentan film-coated tablets should be administered orally twice daily.
2.4 Dosage Adjustments for Aminotransferase Elevations
If aminotransferase levels increase, adjust monitoring and treatment plan according to Table 2.
Discontinue bosentan if liver aminotransferase elevations are accompanied by clinical symptoms of hepatotoxicity (such as nausea, vomiting, fever, abdominal pain, jaundice, or unusual lethargy or fatigue) or bilirubin greater than or equal to 2 x Upper Limit of Normal (ULN). There is no experience with the reintroduction of bosentan in these circumstances.

Table 2. Dosage Adjustment and Monitoring in Patients Developing Aminotransferase Elevations > 3 x ULN

ALT/AST levels	Treatment and monitoring recommendations
> 3 and ≤ 5 x ULN	Confirm by another aminotransferase test. If confirmed, - in patients >12 years and >40 kg, reduce the daily dose to 62.5 mg twice daily or interrupt treatment, and monitor aminotransferase levels at least every 2 weeks. If the aminotransferase levels return to pretreatment values, treatment may continue or be reintroduced at 62.5 mg twice daily, with reassessment of aminotransferase levels within 3 days. - in patients >12 years and <40 kg, interrupt treatment with no prior dose reduction. If the aminotransferase levels return to pretreatment values, reintroduce at the dose used prior to treatment interruption, with reassessment of aminotransferase levels within 3 days.
> 5 and ≤ 8 x ULN	Confirm by another aminotransferase test. If confirmed, stop treatment and monitor aminotransferase levels at least every 2 weeks. Once the aminotransferase levels return to pretreatment values, - in patients >12 years, consider reintroduction of treatment at 62.5 mg twice daily, with reassessment of aminotransferase levels within 3 days. - > 8 x ULN Stop treatment permanently. There is no experience with reintroduction of Bosentan in these circumstances.

2.5 Use with Ritonavir
Coadministration of Bosentan Tablets in Patients on Ritonavir
In patients who have been receiving ritonavir for at least 10 days, start bosentan tablets at the recommended initial dose once daily or every other day based upon individual tolerability (see *Cytochrome P450 Drug Interactions* (7.1)).
Coadministration of Ritonavir in Patients on Bosentan Tablets
Discontinue use of bosentan tablets at least 36 hours prior to initiation of ritonavir. After at least 10 days following the initiation of ritonavir, resume bosentan tablets at the recommended initial dose once daily or every other day based upon individual tolerability (see *Cytochrome P450 Drug Interactions* (7.1)).
2.6 Use in Patients with Pre-existing Hepatic Impairment
Avoid initiation of bosentan tablets in patients with aminotransferases greater than 3 x ULN. No dose adjustment is required in patients with mildly impaired liver function (see *Warnings and Precautions* (5.3), *Use in Specific Populations* (8.6), *Clinical Pharmacology* (12.3)).

3. DOSAGE FORMS AND STRENGTHS
Bosentan tablets for oral administration are available as follows:
62.5 mg - Light orange, oval, biconvex, film-coated tablets, debossed with "WPI" on one side and "62.5" on the other side contains 64.541 mg of bosentan monohydrate, equivalent to 62.5 mg of bosentan base.
125 mg - Light orange, oval, biconvex, film-coated tablets, debossed with "WPI" on one side and "125" on the other side contains 129.082 mg of bosentan monohydrate, equivalent to 125 mg of bosentan base.

• Reduce the dose and closely monitor patients developing aminotransferase elevations more than 3 x Upper Limit of Normal (ULN) (2.1).
----- DOSAGE FORMS AND STRENGTHS -----
• Film-coated tablet: 62.5 mg and 125 mg (3)
----- CONTRAINDICATIONS -----
• Pregnancy (4.1)
• Use with Cyclosporine A (4.2)
• Use with Glyburide (4.3)
• Hypersensitivity (4.4)
----- WARNINGS AND PRECAUTIONS -----
• Fluid retention. (May require intervention) (5.4).
• Pulmonary veno-occlusive disease (PVOD): If signs of pulmonary edema occur, consider the diagnosis of associated PVOD and consider discontinuing bosentan (5.5).
• Decreased sperm counts (5.6).
• Decreases in hemoglobin and hematocrit: Monitor hemoglobin levels after 1 and 3 months of treatment, then every 3 months thereafter (5.7).
----- ADVERSE REACTIONS -----
• Common adverse reactions (greater than or equal to 3% more than placebo) for the film-coated tablet are respiratory tract infection and anemia (6.1).
----- DRUG INTERACTIONS -----
• Cyclochrome P450: Coadministration of bosentan with drugs metabolized by CYP2C9 and CYP3A can increase exposure to bosentan and/or the coadministered drug (4.2, 4.3, 7.1).
• Hormonal contraceptives: bosentan use decreases contraceptive exposure and reduces effectiveness (7.2).
----- USE IN SPECIFIC POPULATIONS -----
• Nursing mothers: Choose breastfeeding or bosentan (8.2).
See 17 for PATIENT COUNSELING INFORMATION and Medication Guide

To report SUSPECTED ADVERSE REACTIONS, contact Actavis at 1-800-272-5525 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.
----- DRUG INTERACTIONS -----
• Cyclochrome P450: Coadministration of bosentan with drugs metabolized by CYP2C9 and CYP3A can increase exposure to bosentan and/or the coadministered drug (4.2, 4.3, 7.1).
• Hormonal contraceptives: bosentan use decreases contraceptive exposure and reduces effectiveness (7.2).
----- USE IN SPECIFIC POPULATIONS -----
• Nursing mothers: Choose breastfeeding or bosentan (8.2).
See 17 for PATIENT COUNSELING INFORMATION and Medication Guide

Revised: 2/2019

4. CONTRAINDICATIONS
4.1 Pregnancy
Use of bosentan is contraindicated in females who are or may become pregnant. To prevent pregnancy, females of reproductive potential must use two reliable forms of contraception during treatment and for one month after stopping bosentan (see *Boxed Warning, Warnings and Precautions* (5.2), *Drug Interactions* (7.2), *Use in Specific Populations* (8.1)).
4.2 Use with Cyclosporine A
Coadministration of cyclosporine A and bosentan resulted in markedly increased plasma concentrations of bosentan. Therefore, concomitant use of bosentan and cyclosporine A is contraindicated (see *Cytochrome P450 Drug Interactions* (7.1)).
4.3 Use with Glyburide
An increased risk of liver enzyme elevations was observed in patients receiving glyburide concomitantly with bosentan. Therefore coadministration of glyburide and bosentan is contraindicated (see *Cytochrome P450 Drug Interactions* (7.1)).
4.4 Hypersensitivity
Bosentan is contraindicated in patients who are hypersensitive to bosentan or any component of the product. Adverse reactions to these cyclosporine A and bosentan resulted in markedly increased plasma concentrations of bosentan. Therefore, concomitant use of bosentan and cyclosporine A is contraindicated (see *Cytochrome P450 Drug Interactions* (7.1)).

5. WARNINGS AND PRECAUTIONS
5.1 Hepatotoxicity
ALT or AST greater than 3 x ULN were observed in 11% of bosentan-treated patients (n = 658) compared to 2% of placebo-treated patients (n = 288). Three-fold increases were seen in 12% of 95 pulmonary arterial hypertension (PAH) patients on 125 mg twice daily and 14% of 70 PAH patients on 250 mg twice daily. Eight-fold increases were seen in 2% of PAH patients on 125 mg twice daily and 7% of PAH patients on 250 mg twice daily. Bilirubin increases to greater than or equal to 3 x ULN were associated with aminotransferase increases in 2 of 658 (0.3%) of patients treated with bosentan. The combination of hepatocellular injury (increases in aminotransferases of greater than 3 x ULN) and increases in total bilirubin (greater than or equal to 2 x ULN) is a marker for potential serious hepatotoxicity.
Elevations of AST or ALT associated with bosentan are dose-dependent, occur both early and late in treatment, usually progress slowly, are typically asymptomatic, and usually have been reversible after treatment interruption or cessation. Aminotransferase elevations also may reverse spontaneously while continuing treatment with bosentan.
Liver aminotransferase levels must be measured prior to initiation of treatment and then monthly and therapy adjusted accordingly (see *Dosage and Administration* (2.1, 2.4)). Discontinue bosentan if liver aminotransferase elevations are accompanied by clinical symptoms of hepatotoxicity (such as nausea, vomiting, fever, abdominal pain, jaundice, or unusual lethargy or fatigue) or increases in bilirubin greater than or equal to 2 x ULN.
Avoid initiation of bosentan in patients with elevated aminotransferases (greater than 3 x ULN) prior to drug initiation because monitoring hepatotoxicity in these patients may be more difficult (see *Boxed Warning, Dosage and Administration* (2.6), *Use in Specific Populations* (8.6)).

In WHO Functional Class II patients, consider whether the benefits of bosentan are sufficient to offset the risk of hepatotoxicity, which may preclude future use as their disease progresses.
5.2 Embryo-fetal Toxicity
Based on data from animal reproduction studies, bosentan may cause fetal harm when administered to a pregnant female and is contraindicated in females who are pregnant. Advise females of reproductive potential of the potential risk to a fetus. Obtain a pregnancy test prior to bosentan treatment, monthly during treatment and for one month after stopping treatment. Advise females of reproductive potential to use two reliable forms of contraception during treatment with bosentan and for at least one month after the last dose (see *Dosage and Administration* (2), *Use in Specific Populations* (8.1, 8.3)). Bosentan is only available for females through a restricted program under REMS (see *Warnings and Precautions* (5.3)).

5.3 Prescribing and Distribution Program for Bosentan Tablets
Because of the risks of hepatotoxicity and birth defects, bosentan tablets are available only through a restricted program called the Bosentan REMS Program. The Bosentan REMS Program is a component of the Bosentan Risk Evaluation and Mitigation Strategy (REMS). Under the Bosentan REMS, prescribers, patients, and pharmacies must enroll in the program (see *Boxed Warning, Warnings and Precautions* (5.1, 5.2), *Contraindications* (4.1)).
Required components of the Bosentan REMS are:
• Healthcare professionals who prescribe bosentan tablets must review the prescriber educational materials, enroll in the Bosentan REMS Program and comply with its requirements.
• Healthcare professionals must (1) review serum aminotransferases (ALT/AST) and bilirubin, and agree to order and monitor these laboratory tests (2) for females of reproductive potential, confirm that the patient is not pregnant, and agree to order and monitor pregnancy tests monthly.
• To receive bosentan tablets, all patients must understand the risks and benefits, complete a patient enrollment form.
• Pharmacies that dispense bosentan tablets must enroll in the program and agree to comply with the Bosentan REMS Program requirements.

Further information about bosentan tablets and the Bosentan REMS Program is available at www.BosentanREMSProgram.com or 1-866-359-2612.
5.4 Fluid Retention
13.1 Carcinogenicity, Mutagenesis, Impairment of Fertility
14 CLINICAL STUDIES
14.1 Pulmonary Arterial Hypertension
14.2 Lack of Benefit in Congestive Heart Failure
16 HOW SUPPLIED/STORAGE AND HANDLING
17 PATIENT COUNSELING INFORMATION
*Sections or subsections omitted from the full prescribing information are not listed.

5.5 Pulmonary Veno-Occlusive Disease
If signs of pulmonary edema occur, consider the possibility of associated pulmonary veno-occlusive disease and consider whether bosentan should be discontinued.
5.6 Decreased Sperm Counts
Decreased sperm counts have been observed in patients receiving bosentan. Preclinical data also suggest that bosentan, similar to other endothelin receptor antagonists, may have an adverse effect on spermatogenesis (see *Adverse Reactions* (6.1), *Nonclinical Toxicology* (13.1)).
5.7 Decreases in Hemoglobin and Hematocrit
Treatment with bosentan can cause a dose-related decrease in hemoglobin and hematocrit. There have been postmarketing reports of decreases in hemoglobin concentration and hematocrit that have resulted in anemia requiring transfusion. It is recommended that hemoglobin concentrations be checked after 1 and 3 months, and every 3 months thereafter. If a marked decrease in hemoglobin concentration occurs, further evaluation should be undertaken to determine the cause and need for specific treatment (see *Adverse Reactions* (6.1)).

6. ADVERSE REACTIONS
The following table lists adverse reactions as described elsewhere in the labeling:
• Hepatotoxicity (see *Boxed Warning, Warnings and Precautions* (5.1))
• Embryo-fetal Toxicity (see *Boxed Warning, Warnings and Precautions* (5.2))
• Fluid Retention (see *Warnings and Precautions* (5.4))
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.
Safety data on bosentan were obtained from 13 clinical studies (9 placebo-controlled and 4 open-label) in 870 adult patients with WHO Functional Class II-IV PAH who were more frequent on bosentan (6% vs 15/288 patients) than on placebo (3% vs 1/122 patients). In this database the only cause of discontinuations greater than 1% and occurring more often on bosentan was abnormal liver function. The adverse drug events that occurred in greater than or equal to 3% of the bosentan-treated patients and were more common on bosentan in placebo-controlled trials in PAH at doses of 125 mg or 250 mg twice daily are shown in Table 3.
Table 3. Adverse Events* Occurring in ≥3% of Patients Treated with Bosentan 125 mg to 250 mg Twice Daily and More Common on Bosentan in Placebo-Controlled Studies in Pulmonary Arterial Hypertension

Adverse Event	Bosentan n = 258		Placebo n = 172	
	No.	%	No.	%
Respiratory Tract Infection**	56	22%	30	17%
Headache	39	15%	25	14%
Edema	28	11%	16	9%
Chest Pain	13	5%	8	5%
Syncope	12	5%	7	4%
Flushing	10	4%	5	3%
Hypotension	10	4%	4	2%
Sinusitis	9	4%	4	2%
Arthralgia	9	4%	3	2%
Serum Aminotransferases, abnormal	9	4%	3	2%
Painful legs	9	4%	4	2%
Anemia	8	3%	3	2%
	8	3%		

*Note: only AEs with onset from start of treatment to 1 calendar day after end of treatment are included. All reported events (at least 3%) are included except those too general to be informative, and those not reasonably associated with the use of the drug because they were associated with the condition being treated or are very common in the treated population.
** Respiratory Tract Infection combines the terms "Nasopharyngitis", "Upper Respiratory Tract Infection" and "Respiratory Tract Infection". Combined data from Study 351, BREATHE-1 and EARLY.
Decreased Sperm Counts
An open-label, single-arm, multicenter, safety study evaluated the effect on testicular function of bosentan 62.5 mg twice daily for 4 weeks, followed by 125 mg twice daily for 5 months. Twenty-five male patients with WHO functional class III and IV PAH and normal baseline sperm count were enrolled. Twenty-three completed the study and 2 discontinued due to adverse events not related to testicular function. There was a decline in sperm count of at least 50% in 25% of the patients after 3 or 6 months of treatment with bosentan. Sperm count remained within the normal range in all 22 patients with data after 6 months and no changes in sperm morphology, sperm motility, or hormone levels were observed. One patient developed marked oligospermia at 3 months and the sperm count remained low with 2 follow-up measurements over the subsequent 6 weeks. Bosentan was discontinued and after 2 months the sperm count had returned to baseline levels. Based on these findings and preclinical data from endothelin receptor antagonists, it cannot be excluded that endothelin receptor antagonists such as bosentan may have an adverse effect on spermatogenesis.
Decreases in Hemoglobin and Hematocrit
Treatment with bosentan can cause a dose-related decrease in hemoglobin and hematocrit. It is recommended that hemoglobin concentrations be checked after 1 and 3 months, and every 3 months thereafter. If a marked decrease in hemoglobin concentration occurs, further evaluation should be undertaken to determine the cause and need for specific treatment.
The overall mean decrease in hemoglobin concentration for adult bosentan-treated patients was 0.9 g/dL (change to end of treatment). Most of this decrease of hemoglobin concentration was detected during the first few weeks of bosentan treatment and hemoglobin levels stabilized by 4 to 12 weeks of bosentan treatment. In placebo-controlled studies of all uses of bosentan, marked decreases in hemoglobin (greater than 15% decrease from baseline resulting in values less than 11 g/dL) were observed in 6% of bosentan-treated patients and 3% of placebo-treated patients. In patients with PAH treated with doses of 125 mg and 250 mg twice daily, marked decreases in hemoglobin occurred in 2% compared to 1% in placebo-treated patients. A decrease in hemoglobin concentration by at least 1 g/dL was observed in 57% of bosentan-treated patients as compared to 29% of placebo-treated patients. In 80% of those patients whose hemoglobin decreased by at least 1 g/dL, the decrease occurred during the first 6 weeks of bosentan treatment.
During the course of treatment, the hemoglobin concentration remained within normal limits in 68% of bosentan-treated patients compared to 76% of placebo patients. The explanation for the change in hemoglobin is not known, but it does not appear to be hemolytic in nature.

6.2 Postmarketing Experience
There have been several postmarketing reports of angioedema associated with the use of bosentan. The onset of the reported cases occurred within a range of 8 hours to 21 days after starting therapy. Some patients were treated with an antihistamine and their signs of angioedema resolved without discontinuing bosentan.
The following additional adverse reactions have been reported during the postapproval use of bosentan. Because these adverse reactions are reported from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to bosentan exposure:
Unexplained hepatic cirrhosis (see *Boxed Warning*)
Liver failure (see *Boxed Warning*)
Hypersensitivity, DRESS, and anaphylaxis (see *Contraindications* (4.4))
Thrombocytopenia
Rash
Jaundice
Anemia requiring transfusion
Neutropenia and leukopenia
Nasal congestion

7. DRUG INTERACTIONS
7.1 Cyclochrome P450 Drug Interactions
Bosentan is metabolized by CYP2C9 and CYP3A. Inhibition of these enzymes may increase the plasma concentration of bosentan (see *Pharmacokinetics* (12.3)). Concomitant administration of both a CYP2C9 inhibitor (such as fluconazole or amiodarone) and a strong CYP3A inhibitor (e.g., ketoconazole, itraconazole) or a moderate CYP3A inhibitor (e.g., ampravir, erythromycin, fluconazole, delamanid) with bosentan will likely lead to large increases in plasma concentrations of bosentan. Coadministration of such combinations of a CYP2C9 inhibitor plus a strong or moderate CYP3A inhibitor with bosentan is not recommended.
Bosentan is an inducer of CYP3A and CYP2C9. Consequently plasma concentrations of drugs metabolized by these two isozymes will be decreased when bosentan is coadministered. Bosentan had no relevant inhibitory effect on any CYP isozyme *in vitro* (CYP1A2, CYP2C9, CYP2C19, CYP2D6, CYP3A). Consequently, bosentan is not expected to increase the plasma concentrations of drugs metabolized by these enzymes.

Figure 1. CYP3A Induction-Mediated Effect of Bosentan on Other Drugs

Interacting drug	Cmax	AUC	Recommendation
Cyclosporin A	↔	↔	contraindicated
Glyburide	↔	↔	contraindicated
Simvastatin	↔	↔	monitor cholesterol
β-Hydroxy acid simvastatin	↔	↔	monitor cholesterol
Norethindrone	↔	↔	use additional contraception
Ethinyl estradiol	↔	↔	use additional contraception

Figure 2. Effect of Other Drugs on Bosentan

Interacting drug	Ctrough Day 2	Ctrough (ss)	AUC	Recommendation
Cyclosporin A	↔	↔	↔	contraindicated
DATP inhibition	↔	↔	↔	
Lopinavir/ritonavir	↔	↔	↔	dose reduction
DATP and CYP3A inhibition	↔	↔	↔	
Rifampin	↔	↔	↔	AST/ALT weekly
CYP3A and CYP2C9 induction and DATP inhibition	↔	↔	↔	

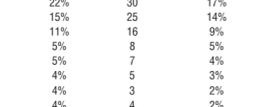
7.2 Hormonal Contraceptives
Hormonal contraceptives, including oral, injectable, transdermal, and implantable forms, may not be reliable when bosentan is coadministered. Females should practice additional methods of contraception and not rely on hormonal contraception alone when taking bosentan (see *Use in Specific Populations* (8.3)).
An interaction study demonstrated that coadministration of bosentan and a combination oral hormonal contraceptive produced average decreases of norethindrone and ethinyl estradiol levels of 14% and 31%, respectively. However, decreases in exposure were as small as 56% and 66%, respectively, in individual subjects.
8. USE IN SPECIFIC POPULATIONS
8.1 Pregnancy
Risk Summary
Based on data from animal reproduction studies, bosentan may cause fetal harm, including birth defects and fetal death, when administered to a pregnant female and is contraindicated during pregnancy (see *Contraindications* (4.1)). There are limited data on bosentan use in pregnant women. In animal reproduction studies, oral administration of bosentan to pregnant rats at 2-times the maximum recommended human dose (MRHD) on a mg/m² basis caused teratogenic effects in rats, including malformations of the head, mouth, face, and large blood vessels (see *Animal Data*). Advise pregnant women of the potential risk to a fetus.
The estimated background risk of major birth defects and miscarriage for the indicated population is unknown. All pregnancies have a background risk of birth defect, loss, or miscarriage. In the U.S. general population, the estimated background risk of major birth defects and miscarriage in clinically recognized pregnancies is 2% to 4% and 15% to 20%, respectively.
Data
Animal Data
Bosentan was teratogenic in rats given oral doses two times the MRHD (on a mg/m² basis). In an embryo-fetal toxicity study in rats, bosentan showed dose-dependent teratogenic effects, including malformations of the head, mouth, face and large blood vessels. Bosentan increased stillbirths and pup mortality at oral doses 2 and 10 times the MRHD (on a mg/m² basis). Although birth defects were not observed in rabbits given oral doses of up to the equivalent of 10.5/day in a 70 kg person, plasma concentrations of bosentan in rabbits were lower than those reached in the rat. The similarity of malformations induced by bosentan and those observed in endothelin-1 knockout mice and in animals treated with other endothelin receptor antagonists indicates that embryo-fetal toxicity is a class effect of these drugs.

8.2 Lactation
Risk Summary
There are no data on the presence of bosentan in human milk, the effects on the breastfed infant, or the effect on milk production. Because of the potential for serious adverse reactions, such as fluid retention and hepatotoxicity, in breastfed infants from bosentan, advise women not to breastfeed during treatment with bosentan.
8.3 Females and Males of Reproductive Potential
Pregnancy Testing
Verify the pregnancy status of females of reproductive potential prior to initiating bosentan, monthly during treatment and one month after stopping treatment with bosentan. The patient should contact her physician immediately for pregnancy testing if onset of menses is delayed or pregnancy is suspected. If the pregnancy test is positive, the physician and patient must discuss the risks to her, the pregnancy, and the fetus.
Contraception
Drug interaction studies show that bosentan reduces serum levels of the estrogen and progestin in oral contraceptives. Based on these findings, hormonal contraceptives (including oral, injectable, transdermal, and implantable contraceptives) may be less effective for preventing pregnancy in patients using bosentan and should not be used as a patient's only contraceptive method (see *Drug Interactions* (7.2)). Females of reproductive potential using bosentan must use two acceptable methods of contraception during treatment and for 1 month after treatment with bosentan. Patients may choose one highly effective form of contraception (intrauterine device (IUD) or tubal sterilization) or a combination of methods (hormone method with a barrier method or two barrier devices). If a partner's vasectomy is the chosen method of contraception, a hormone or barrier method must be used along with the vasectomy until the partner has had a physician's counseling plan on pregnancy planning and prevention, or designate counseling by another healthcare provider trained in contraceptive counseling (see *Boxed Warning*).

Infertility
Males
Decreased sperm counts have been observed in patients receiving bosentan. Based on these findings and findings in animals, bosentan may impair fertility in males of reproductive potential.
It is not known whether effects on fertility would be reversible (see *Warnings and Precautions* (5.6), *Adverse Reactions* (6.1), *Nonclinical Toxicology* (13.1)).
8.4 Pediatric Use
Juvenile Animal Toxicity Data
In a juvenile rat toxicity study, rats were treated from Day 4 postpartum to adulthood (Day 69 postpartum). Decreased body weights, absolute weights of testes and epididymides, and reduced number of sperm in epididymides were observed after weaning. No effect on testis histology or sperm morphology and function was seen. The NOAEL was 4 times at (Day 4 postpartum) and 2 times (Day 69 postpartum) the human therapeutic exposure, respectively.
No effects on general development, sensory, cognitive function and reproductive performance were detected at the highest dose tested in juvenile rats, 7 times the therapeutic exposure in children with PAH.

8.5 Geriatric Use
Clinical studies of bosentan did not include sufficient numbers of subjects aged 65 and older to determine whether they respond differently from younger subjects.
8.6 Hepatic Impairment
Because there is *in vitro* and *in vivo* evidence that the main route of excretion of bosentan is biliary, liver impairment could be expected to increase exposure (C_{max} and AUC) of bosentan. The pharmacokinetics of bosentan have been evaluated in patients with severe liver impairment (Child-Pugh Class C). In patients with moderate hepatic impairment (Child-Pugh Class B), the systemic exposures to bosentan and its active metabolite increased significantly. Bosentan should generally be avoided in patients with moderate or severe liver impairment. Pharmacokinetics of bosentan were not altered in patients with mild impairment of hepatic function (Child-Pugh Class A) (see *Dosage and Administration* (2.6), *Warnings and Precautions* (5.1), *Pharmacokinetics* (12.3)).
8.7 Renal Impairment
The effect of renal impairment on the pharmacokinetics of bosentan is small and does not require dosing adjustment (see *Pharmacokinetics* (12.3)).
10. OVERDOSAGE
Bosentan has been given as a single dose of up to 2,400 mg in normal volunteers, or up to 2,000 mg/day for 2 months in patients, without any major clinical consequences. The most common side effect was headache of mild to moderate intensity. In the cyclosporine A interaction study, in which doses of 500 and 1,000 mg twice daily of bosentan were given concomitantly with cyclosporine A, trough plasma concentrations of bosentan increased 30-fold, resulting in severe headache, nausea, and vomiting, but no serious adverse events. Mild decreases in blood pressure and increases in heart rate were observed. In the postmarketing period, there was one reported overdose of 10,000 mg of bosentan taken by an adolescent male patient. He had symptoms of nausea, vomiting, hypotension, dizziness, sweating, and blurred vision. He recovered within 24 hours with blood pressure support. Bosentan is unlikely to be effectively removed by dialysis due to the high molecular weight and extensive plasma protein binding.

11. DESCRIPTION
Bosentan, an endothelin receptor antagonist that belongs to a class of highly substituted pyrimidine derivatives, with no chiral centers, is designated chemically as 4-tert-butyl-N-[6-(2-hydroxy-ethoxy)-5-(2-methoxy-ethoxy)-1,2,3,4-tetrahydro-1H-benzene-sulfonamide] monohydrate and has the following structural formula:



Who should not take bosentan tablets?

Do not take bosentan tablets if you:

- are pregnant, plan to become pregnant, or become pregnant during bosentan treatment. Bosentan tablets can cause serious birth defects. All females should read the birth defects section of “What is the most important information I should know about bosentan tablets?”
- take any of these medicines:

- cyclosporine A used to treat psoriasis and rheumatoid arthritis, and to prevent rejection of heart, liver, and kidney transplants
- glyburide used to treat diabetes

- are allergic to bosentan or any of the ingredients in bosentan tablets. See the end of this Medication Guide for a complete list of the ingredients in bosentan tablets. If you have a rash, hives or your lips swell after taking bosentan tablets, it may be a sign of allergy. You should stop taking your bosentan tablets and talk to your healthcare provider.

What should I tell my healthcare provider before taking bosentan tablets?

Bosentan tablets may not be right for you. Tell your healthcare provider about all your medical conditions, including if you:

- have liver problems.
- are breast-feeding or plan to breast feed. It is not known if bosentan passes into your milk. You and your healthcare provider should decide if you will take bosentan tablets or breastfeed. You should not do both.
- Tell your healthcare provider about all the medicines you take, including prescription and over-the-counter medicines, vitamins, and herbal supplements. Bosentan tablets and other medicines may affect how each other works and cause side effects. Especially tell your healthcare provider if you take:
 - hormone-based birth control, such as pills, shots, patches, and implants. These birth control methods may not work as well when taken with bosentan tablets.
 - simvastatin or other “-statin” medicines used to lower cholesterol
 - rifampin used for tuberculosis
 - ketoconazole, fluconazole, itraconazole, or voriconazole used for fungal infections
 - warfarin sodium used to prevent blood clots
 - ritonavir used to treat HIV

There may be more than one brand name medicine. Ask your healthcare provider if you are not sure if your medicine is one that is listed above.

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

How should I take bosentan tablets?

Your healthcare provider will give you detailed information about the Bosentan REMS Program.

- Bosentan tablets will be mailed to you by a specialty pharmacy. You will only receive a 30-day supply of bosentan tablets at one time.
- Take bosentan tablets exactly as prescribed.
- Your healthcare provider will tell you how many bosentan tablets to take and when to take it.
- In most cases, you will take 1 tablet in the morning and 1 in the evening.
- You can take bosentan tablets orally with or without food.
- If you take more than the prescribed dose of bosentan tablets, call your healthcare provider right away.
- If you miss a dose of bosentan tablets, take your tablet as soon as you remember. Do not take 2 doses at the same time. If it is almost time for your next dose, skip the missed dose. Just take the next dose at your regular time.
- Do not stop taking bosentan tablets unless your healthcare provider tells you to. Suddenly stopping your treatment may cause your symptoms to get worse. If you need to stop taking bosentan tablets, speak with your healthcare provider about the right way to stop.

What are the possible side effects of bosentan tablets?

Bosentan tablets can cause serious side effects, including:

- See “What is the most important information I should know about bosentan tablets?”
- Fluid retention and swelling of your ankles and legs. Bosentan tablets can cause your body to hold too much water, and you may get swelling of your ankles and legs. Tell your healthcare provider if you have swelling of your ankles and legs that happens either with or without weight gain, or if you have more trouble with your breathing than normal. Your healthcare provider will look for the cause of this.
- Lower Sperm Count. Some men who take bosentan tablets may have lower sperm counts. This may affect your ability to father a child. Tell your healthcare provider if fertility is a concern for you.
- Low red blood cell levels (anemia). Your healthcare provider will do blood tests to check your red blood cells during treatment with bosentan tablets.

The most common side effects of bosentan tablets include:

- respiratory tract infection
- headache
- fainting
- flushing
- low blood pressure
- inflamed nose passages (sinusitis)
- joint pain
- irregular heart beats

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 bosentan tablets. 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 bosentan tablets?

- Store bosentan tablets at room temperature between 68°F to 77°F (20°C to 25°C).

Keep bosentan tablets and all medicines out of the reach of children.

General information about bosentan tablets

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

This Medication Guide summarizes the most important information about bosentan tablets. If you would like more information, talk with your healthcare provider. You can ask your pharmacist or healthcare provider for information about bosentan tablets that is written for health professionals. For more information call Actavis at 1-800-272-5525.

What are the ingredients in bosentan tablets?

Active ingredient: bosentan

Inactive ingredients: corn starch, ethyl cellulose, glyceryl behenate, hypromellose, iron oxide red, iron oxide yellow, magnesium stearate, pregelatinized corn starch, povidone K90, sodium starch glycolate, titanium dioxide and triacetin.

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

Manufactured by:
Watson Pharma Private Ltd.
Verna, Salcette Goa 403 722 India

Distributed by:
Actavis Pharma, Inc.
Parsippany, NJ 07054 USA

12.3 Pharmacokinetics

General

After oral administration, maximum plasma concentrations of bosentan are attained within 3 to 5 hours and the terminal elimination half-life is about 5 hours in healthy adult subjects. The exposure to bosentan after intravenous and oral administration is about twice as high in adult patients with PAH as it is in healthy adult subjects.

Absorption

The absolute bioavailability of bosentan in normal volunteers is about 50% and is unaffected by food.

Bosentan is highly bound (greater than 98%) to plasma proteins, mainly albumin. Bosentan does not penetrate into erythrocytes. The volume of distribution is about 18 L.

Elimination

Metabolism
Bosentan has three metabolites, one of which is pharmacologically active and may contribute 10% to 20% of the effect of bosentan. Bosentan is an inducer of CYP2C9 and CYP3A and possibly also of CYP2C19. Upon multiple oral dosing, plasma concentrations in healthy adults decrease gradually to 50% to 65% of those seen after single dose administration, probably the effect of auto-induction of the metabolizing liver enzymes. Steady-state is reached within 3 to 5 days.

Excretion

Bosentan is eliminated by biliary excretion following metabolism in the liver. Less than 3% of an administered oral dose is recovered in urine. Total clearance after a single intravenous dose is about 4 L/h in patients with PAH.

Specific Populations

Hepatic Impairment

In vitro and *in vivo* evidence showing extensive hepatic metabolism of bosentan suggests that liver impairment could significantly increase exposure to bosentan. In a study comparing 8 patients with mild liver impairment (Child-Pugh Class A) to 8 controls, the single- and multiple-dose pharmacokinetics of bosentan were not altered in patients with mild hepatic impairment.

In another small (N=8) pharmacokinetic study, the steady-state AUC of bosentan was on average 4.7 times higher and the active metabolite Ro 48-5033 was 12.4 times higher in 5 patients with moderately impaired liver function (Child-Pugh Class B) and PAH associated with portal hypertension than in 3 patients with normal liver function and PAH of other etiologies.

The pharmacokinetics of bosentan have not been evaluated in patients with severe liver impairment (Child-Pugh Class C) [see Dosage and Administration (2.2), Warnings and Precautions (5.1), Use in Specific Populations (8.6)].

Renal Impairment

In patients with severe renal impairment (creatinine clearance 15 to 30 mL/min), plasma concentrations of bosentan were essentially unchanged and plasma concentrations of the three metabolites were increased about 2-fold compared to subjects with normal renal function. These differences do not appear to be clinically important.

Drug Interactions

Ketoconazole

Coadministration of bosentan 125 mg twice daily and ketoconazole, a potent CYP3A inhibitor, increased the plasma concentrations of bosentan by approximately 100% in normal volunteers. No dose adjustment of bosentan is necessary, but increased effects of bosentan should be considered.

Warfarin

Coadministration of bosentan 500 mg twice daily for 6 days in normal volunteers decreased the plasma concentrations of both S-warfarin (a CYP2C9 substrate) and R-warfarin (a CYP3A substrate) by 29 and 38%, respectively. Clinical experience with concomitant administration of bosentan and warfarin in patients with PAH did not show clinically relevant changes in INR or warfarin dose (baseline vs. end of the clinical studies), and the need to change the warfarin dose during the trials due to changes in INR or due to adverse events was similar among bosentan- and placebo-treated patients.

Digoxin, Nimodipine, and Losartan

Bosentan has no significant pharmacokinetic interactions with digoxin and nimodipine, and losartan has no significant effect on plasma levels of bosentan.

Sildenafil

In normal volunteers, coadministration of multiple doses of 125 mg twice daily bosentan and 80 mg three times daily sildenafil resulted in a reduction of sildenafil plasma concentrations by 63% and increased bosentan plasma concentrations by 50%. This change in plasma concentrations were not considered clinically relevant and dose adjustments are not necessary. This recommendation holds true when sildenafil is used for the treatment of PAH or erectile dysfunction.

Tadalafil

Bosentan (125 mg twice daily) reduced tadalafil (40 mg once per day) systemic exposure (AUC) by 42% and *C*_{max} by 27% following multiple dose coadministration. Tadalafil did not affect the exposure (AUC and *C*_{max}) of bosentan or its metabolites.

Figure 3. CYP Induction-Mediated Effect of Bosentan on Other Drugs

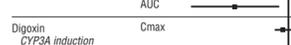
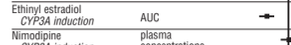
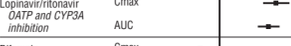
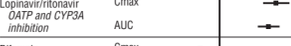
Interacting drug		Recommendation
Cyclosporin A <i>CYP3A induction</i>		contraindicated
Glyburide <i>CYP3A induction</i>		contraindicated
Simvastatin <i>CYP3A induction</i>		monitor cholesterol
8-Hydroxy acid simvastatin <i>CYP3A induction</i>		monitor cholesterol
Digoxin <i>CYP3A induction</i>		no adjustment
Sildenafil <i>CYP3A induction</i>		no adjustment
S-Warfarin <i>CYP2C9 induction</i>		no adjustment
R-Warfarin <i>CYP3A induction</i>		no adjustment
Norethindrone <i>CYP3A induction</i>		use additional contraception
Ethinyl estradiol <i>CYP3A induction</i>		use additional contraception
Nimodipine <i>CYP3A induction</i>		no adjustment
Tadalafil <i>CYP3A induction</i>		no adjustment

Figure 4. Effects of Other Drugs on Bosentan

Interacting drug		Recommendation
Cyclosporin A <i>OATP inhibition</i>		contraindicated
Lopinavir/ritonavir <i>OATP and CYP3A inhibition</i>		dose reduction
Rifampin <i>CYP3A and CYP2C9 induction and OATP inhibition</i>		AST/ALT weekly
Ketoconazole <i>CYP3A inhibition</i>		no adjustment
Losartan <i>CYP3A4 and CYP2C9 substrate</i>		no adjustment
Sildenafil <i>OATP inhibition</i>		no adjustment
Tadalafil <i>CYP3A4 and CYP2C9 substrate</i>		no adjustment

13. NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis and Mutagenesis, Impairment of Fertility

Carcinogenesis and Mutagenesis

Two years of dietary administration of bosentan to mice produced an increased incidence of hepatocellular adenomas and carcinomas in males at doses as low as 450 mg/kg/day (about 8 times the maximum recommended human dose (MRHD) of 125 mg twice daily, on a mg/m² basis). In the same study, doses greater than 2,000 mg/kg/day (about 32 times the MRHD) were associated with an increased incidence of colon adenomas in both males and females. In rats, dietary administration of bosentan for two years was associated with an increased incidence of brain astrocytomas in males at doses as low as 500 mg/kg/day (about 16 times the MRHD). In a comprehensive battery of *in vitro* tests (the microbial mutagenesis assay, the unscheduled DNA synthesis assay, the V-79 mammalian cell mutagenesis assay, and human lymphocyte assay) and an *in vivo* mouse micronucleus assay, there was no evidence for any mutagenic or clastogenic activity of bosentan.

Impairment of Fertility/Testicular Function

The development of testicular tubular atrophy and impaired fertility has been linked with the chronic administration of certain endothelin receptor antagonists in rodents.

Treatment with bosentan at oral doses of up to 1,500 mg/kg/day (50 times the MRHD on a mg/m² basis) or intravenous doses up to 40 mg/kg/day had no effects on sperm count, sperm motility, mating performance or fertility in male and female rats. An increased incidence of testicular tubular atrophy was observed in rats given bosentan orally at doses as low as 125 mg/kg/day (about 4 times the MRHD) and the lowest doses tested) for two years but not at doses as high as 1,500 mg/kg/day (about 50 times the MRHD) for 6 months. Effects on sperm count and motility were evaluated only in the much shorter duration fertility studies in which males had been exposed to the drug for 4 to 6 weeks. An increased incidence of tubular atrophy was not observed in mice treated for 2 years at doses up to 4,500 mg/kg/day (about 75 times the MRHD) or in dogs treated up to 12 months at doses up to 500 mg/kg/day (about 50 times the MRHD).

14. CLINICAL STUDIES

14.1 Pulmonary Arterial Hypertension

WHO Functional Class III-IV

Two randomized, double-blind, multi-center, placebo-controlled trials were conducted in 32 and 213 patients. The larger study (BREATHE-1) compared 2 doses (125 mg twice daily and 250 mg twice daily) of bosentan with placebo. The smaller study (Study 351) compared 125 mg twice daily with placebo. Patients had severe (WHO functional Class III-IV) PAH: idiopathic or heritable PAH (72%) or PAH associated with scleroderma or other connective tissue diseases (21%), or to autoimmune diseases (7%). There were no patients with PAH associated with other conditions such as HIV disease or recurrent pulmonary emboli.

In both studies, bosentan or placebo was added to patients' current therapy, which could have included a combination of digoxin, anticoagulants, diuretics, and vasodilators (e.g., calcium channel blockers, ACE inhibitors), but not epoprostenol.

Bosentan was given at a dose of 62.5 mg twice daily for 4 weeks and then at 125 mg twice daily or 250 mg twice daily for either 12 (BREATHE-1) or 8 (Study 351) additional weeks. The primary study endpoint was 6-minute walk distance. In addition, symptoms and functional status were assessed. Hemodynamic measurements were made at 12 weeks in Study 351.

The mean age was about 49 years. About 80% of patients were female, and about 80% were Caucasian. Patients had been diagnosed with pulmonary hypertension for a mean of 2.4 years.

Submaximal Exercise Ability

Results of the 6-minute walk distance at 3 months (Study 351) or 4 months (BREATHE-1) are shown in Table 4.

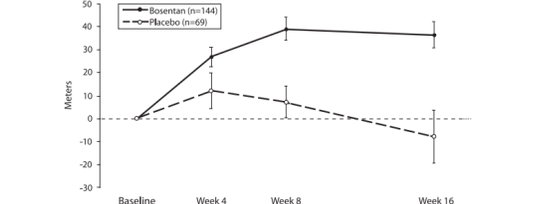
Table 4. Effects of Bosentan on 6-minute walk distance

	BREATHE-1			Study 351	
	Bosentan 125 mg twice daily (n = 74)	Bosentan 250 mg twice daily (n = 70)	Placebo (n = 69)	Bosentan 125 mg twice daily (n = 21)	Placebo (n = 11)
Baseline	326 ± 73	333 ± 75	344 ± 76	360 ± 86	355 ± 82
End point	353 ± 115	379 ± 101	336 ± 129	431 ± 66	350 ± 147
Change from baseline	27 ± 75	46 ± 62	-8 ± 96	70 ± 56	-6 ± 121
Placebo – subtracted	35 (a)	54 (b)		76 (c)	

Distance in meters; mean ± standard deviation. Changes are to week 16 for BREATHE-1 and to week 12 for Study 351. (a)*p*≤0.01; by Wilcoxon; (b)*p*≤0.0001; by Wilcoxon; (c)*p*≤0.02; by Student's t-test.

In both trials, treatment with bosentan resulted in a significant increase in exercise ability. The improvement in walk distance was apparent after 1 month of treatment (with 62.5 mg twice daily) and fully developed by about 2 months of treatment (Figure 5). It was maintained for up to 7 months of double-blind treatment. Walking distance was somewhat greater with 250 mg twice daily, but the potential for increased hepatotoxicity causes this dose not to be recommended [see Dosage and Administration (2.1)]. There were no apparent differences in treatment effects on walk distance among subgroups analyzed by demographic factors, baseline disease severity, or disease etiology, but the studies had little power to detect such differences.

Figure 5. Mean Change in 6-min Walk Distance (BREATHE-1)



Change from baseline in 6-minute walking distance from start of therapy to week 16 in the placebo and combined bosentan (125 mg twice daily and 250 mg twice daily) groups. Values are expressed as mean ± standard error of the mean.

Hemodynamic Changes

Invasive hemodynamic parameters were assessed in Study 351. Treatment with bosentan led to a significant increase in cardiac index (CI) associated with a significant reduction in pulmonary artery pressure (PAP), pulmonary vascular resistance (PVR), and mean right atrial pressure (RAP) (Table 5).

The relationship between hemodynamic effects and improvements in 6-minute walk distance is unknown.

Table 5. Change from Baseline to Week 12: Hemodynamic Parameters

	Bosentan 125 mg twice daily	Placebo
CI (L/min/m²)	n=20	n=10
Baseline	2.35±0.73	2.48±1.03
Absolute Change	0.50±0.46	-0.52±0.48
Treatment Effect		1.02(a)
Mean PAP (mmHg)	n=20	n=10
Baseline	53.7±13.4	55.7±10.5
Absolute Change	-1.6±5.1	5.1±8.8
Treatment Effect		-6.7(b)
PVR (dyn·sec·cm⁻⁵)	n=19	n=10
Baseline	896±425	942±430
Absolute Change	-223±245	191±235
Treatment Effect		-415(a)
Mean RAP (mmHg)	n=19	n=10
Baseline	9.7±5.6	9.9±4.1
Absolute Change	-1.3±4.1	4.9±4.6
Treatment Effect		-6.2(a)

Values shown are means ± SD

(a)*p*≤0.001; (b)*p*<0.02

Symptoms and Functional Status

Symptoms of PAH were assessed by Borg dyspnea score, WHO functional class, and rate of “clinical worsening.” Clinical worsening was assessed as the sum of death, hospitalizations for PAH, discontinuation of therapy because of PAH, and need for epoprostenol. There was a significant reduction in dyspnea during walk tests (Borg dyspnea score), and significant improvement in WHO functional class in bosentan-treated patients. There was a significant reduction in the rate of clinical worsening (Table 6 and Figure 6). Figure 6 shows the log-rank test reflecting clinical worsening over 28 weeks.

Table 6. Incidence of Clinical Worsening, Intent To Treat Population

	BREATHE-1		Study 351	
	Bosentan 125/250 mg twice daily (n = 144)	Placebo (n = 69)	Bosentan 125 mg twice daily (n = 21)	Placebo (n = 11)
Patients with clinical worsening	9 (6%) ^(a)	14 (20%)	0 (0%) ^(b)	3 (27%)
Death	1 (1%)	2 (3%)	0 (0%)	0 (0%)
Hospitalization for PAH	6 (4%)	9 (13%)	0 (0%)	3 (27%)
Discontinuation due to worsening of PAH	5 (3%)	6 (9%)	0 (0%)	3 (27%)
Receipt of epoprostenol ^(c)	4 (3%)	3 (4%)	0 (0%)	3 (27%)

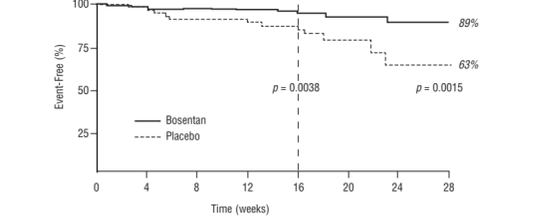
Note: Patients may have had more than one reason for clinical worsening.

(a)*p*≤0.0015 vs. placebo by log-rank test. There was no relevant difference between the 125 mg and 250 mg twice daily groups.

(b)*p*=0.033 vs. placebo by Fisher's exact test.

(c)Receipt of epoprostenol was always a consequence of clinical worsening.

Figure 6. Time to Clinical Worsening (BREATHE-1)



Time from randomization to clinical worsening with Kaplan-Meier estimate of the proportions of failures in BREATHE-1. All patients (n=144 in the bosentan group and n=69 in the placebo group) participated in the first 16 weeks of the study. A subset of this population (n=35 in the bosentan group and 13 in the placebo group) continued double-blind therapy for up to 28 weeks.

WHO Functional Class II

In a randomized, double-blind, multicenter, placebo-controlled trial, 185 mildly symptomatic PAH patients with WHO Functional Class II (mean baseline 6-minute walk distance of 443 meters) received bosentan 62.5 mg twice daily for 4 weeks followed by 125 mg twice daily (n = 93), or placebo (n = 92) for 6 months. Enrolled patients were treatment-naïve (n = 156) or on a stable dose of sildenafil (n = 29). The coprimary endpoints were change from baseline to month 6 in PVR and 6-minute walk distance. Time to clinical worsening (assessed as the sum of death, hospitalization due to PAH complications, or symptomatic progression of PAH), Borg dyspnea index, change in WHO functional class and hemodynamics were assessed as secondary endpoints.

Compared with placebo, bosentan treatment was associated with a reduced incidence of worsening of at least one functional class (3% bosentan vs. 13% placebo, *p* = 0.03), and improvement in hemodynamic variables (PVR, mPAP, TPR, cardiac index, and SVD₂; *p* < 0.05). The +19 m mean (+14 m median) increase in 6-minute walk distance with bosentan vs. placebo was not significant (*p* = 0.06). There was a significant delay in time to clinical worsening (first seen primarily as symptomatic progression of PAH) with bosentan compared with placebo (hazard ratio 0.2, *p* = 0.01). Findings were consistent in strata with or without treatment with sildenafil at baseline.

Long-term Treatment of PAH

Long-term follow-up of patients with Class III and IV PAH who were treated with bosentan in open-label extensions of trials (N=235) showed that 93% and 84% of patients were still alive at 1 and 2 years, respectively, after the start of treatment.

These uncontrolled observations do not allow comparison with a group not given bosentan and cannot be used to determine the long-term effect of bosentan on mortality.

Pulmonary Arterial Hypertension in Adults related to Congenital Heart Disease with Left-to-Right Shunts

A small study (N=54) and its open-label extension (N=37) of up to 40 weeks in adult patients with Eisenmenger physiology demonstrated effects of bosentan on exercise and safety that were similar to those seen in other trials in patients with PAH (WHO Group 1).

14.2 Lack of Benefit in Congestive Heart Failure

Bosentan is not effective in the treatment of congestive heart failure with left ventricular dysfunction. In a pair of studies, 1,613 subjects with NYHA Class III-IV heart failure, left ventricular ejection fraction less than 35%, on diuretics, ACE inhibitor, and other therapies, were randomized to placebo or bosentan (62.5 mg twice daily titrated as tolerated to 125 mg twice daily) and followed for up to 70 weeks. Use of bosentan was associated with no benefit on patient global assessment (the primary end point) or mortality. However, hospitalizations for heart failure were more common during the first 4 to 8 weeks after bosentan was initiated. In a placebo-controlled trial of patients with severe chronic heart failure, there was an increased incidence of hospitalization for CHF associated with weight gain and increased leg edema during the first 4 to 8 weeks of treatment with bosentan. Patients required intervention with a diuretic, fluid management, or hospitalization for decompensating heart failure.

16. HOW SUPPLIED/STORAGE AND HANDLING

Bosentan tablets