THE ALOSETRON REMS PROGRAM

Prescriber Education Slide Deck

Understanding the Benefits and Risks of Alosetron

The Alosetron REMS Program - Please see complete Prescribing Information for Alosetron Tablets for full details.
# Table of Contents

<table>
<thead>
<tr>
<th>Section</th>
<th>Title</th>
<th>Slide</th>
</tr>
</thead>
<tbody>
<tr>
<td>Section 1</td>
<td>Purpose</td>
<td>3</td>
</tr>
<tr>
<td>Section 2</td>
<td>Indication and Usage</td>
<td>8</td>
</tr>
<tr>
<td>Section 3</td>
<td>Important Safety Information</td>
<td>11</td>
</tr>
<tr>
<td>Section 4</td>
<td>How to Prescribe and Dispense Alosetron Tablets</td>
<td>27</td>
</tr>
<tr>
<td>Section 5</td>
<td>Alosetron REMS Program</td>
<td>31</td>
</tr>
</tbody>
</table>
Section 1:

Purpose
Purpose of the Prescriber Educational Slide Deck for Alosetron

• By reviewing the information provided in this presentation, prescribers who prescribe alosetron hydrochloride (alosetron) will better understand the:

  - Restricted distribution process for this product;
  - Risks and benefits of alosetron;
  - Etiology of irritable bowel syndrome;
  - The Alosetron REMS Program
Risk Evaluation and Mitigation Strategy (REMS)

The Food and Drug Administration (FDA) has determined that a Risk Evaluation and Mitigation Strategy (REMS) is necessary for alosetron to ensure the benefits of the drug outweigh the risk of serious gastrointestinal adverse reactions. A REMS is a strategy to address the serious risks associated with a drug. The REMS can range from periodic assessment of a product’s postmarketing safety profile to strict limitations on the way a drug is prescribed, distributed, or dispensed.
Goal of The Alosetron REMS Program and Key Elements

The Alosetron REMS Program was implemented to help reduce the risks of serious GI adverse events.

The Goals of the Alosetron REMS Program are:

• To mitigate the risk of ischemic colitis (IC) and serious complications of constipation (CoC) associated with alosetron hydrochloride (hereinafter, referred to as alosetron) by ensuring that alosetron is used in only severely affected patients in whom benefits exceed the risks.

• To ensure that the risk of IC and serious CoC with the use of alosetron are communicated to patients, pharmacists, and prescribers.
Goal of The Alosetron REMS Program and Key Elements (cont’d)

The Key Elements of the Alosetron REMS are:

- only prescribers who have enrolled in the Alosetron REMS Program, based on their understanding of the benefits and risks, can prescribe alosetron.

- pharmacists may only dispense alosetron from prescriptions with a sticker and written by prescribers participating in the Alosetron REMS Program.¹

¹ In the alternative, a prescription for alosetron may be dispensed in the presence of a Prescribing Program for LOTRONEX™ sticker, if generic substitution of alosetron is permitted pursuant to State substitution laws.

Goal of The Alosetron REMS Program and Key Elements (cont’d)

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Section 2: Indication and Usage
Indication and Usage

Alosetron is indicated ONLY for women with severe diarrhea-predominant IBS who have:

- chronic IBS symptoms (generally lasting 6 months or longer),
- had anatomic or biochemical abnormalities of the GI tract excluded, and
- not responded adequately to conventional therapy.
Indication and Usage (cont’d)

• Diarrhea-predominant IBS is severe if it includes diarrhea and one or more of the following:
  - frequent and severe abdominal pain/discomfort,
  - frequent bowel urgency or fecal incontinence,
  - disability or restriction of daily activities due to IBS.

• Because of infrequent but serious GI adverse reactions associated with alosetron, the indication is restricted to those patients for whom the benefit-to-risk balance is most favorable.

• Clinical studies have not been performed to adequately confirm the benefits of alosetron in men.
Section 3:
Important Safety Information

The Alosetron REMS Program - Please see complete Prescribing Information for Alosetron Tablets for full details.
Boxed Warning

WARNING: SERIOUS GASTROINTESTINAL ADVERSE REACTIONS

• Infrequent but serious gastrointestinal adverse reactions have been reported with the use of alosetron. These events, including ischemic colitis and serious complications of constipation, have resulted in hospitalization, and rarely, blood transfusion, surgery, and death.

• The Alosetron REMS Program was implemented to help reduce risks of serious gastrointestinal adverse reactions. Only prescribers who have enrolled in the Alosetron REMS Program, based on their understanding of the benefits and risks, should prescribe alosetron.

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Boxed Warning (cont’d)

• Alosetron is indicated only for women with severe diarrhea-predominant IBS who have not responded adequately to conventional therapy. Before receiving the initial prescription for alosetron, the patient must read and sign the Patient Acknowledgement Form for alosetron.

• Alosetron should be discontinued immediately in patients who develop constipation or symptoms of ischemic colitis. Patients should immediately report constipation or symptoms of ischemic colitis to their prescriber. Alosetron should not be resumed in patients who develop ischemic colitis. Patients who have constipation should immediately contact their prescriber if the constipation does not resolve after alosetron is discontinued. Patients with resolved constipation should resume alosetron only on the advice of their treating prescriber.
Warnings and Precautions

Serious Complications of Constipation

- Some patients have experienced serious complications of constipation without warning. Examples include:
  - obstruction, ileus, impaction, toxic megacolon, and secondary bowel ischemia have been reported with use of alosetron during clinical trials.
  - in addition, rare cases of intestinal perforation and death have been reported from postmarketing clinical practice.
  - in some cases, complications of constipation required intestinal surgery, including colectomy.

The Alosetron REMS Program - Please see complete Prescribing Information for Alosetron Tablets for full details.
Warnings and Precautions (cont’d)

Serious Complications of Constipation (cont’d)

• The incidence of serious complications of constipation was ~0.1%, or 1 per 1,000 patients, in women receiving either alosetron or placebo.

• Patients who are elderly, debilitated, or taking additional medications that decrease GI motility may be at greater risk for complications of constipation.

• Alosetron should be discontinued immediately in patients who develop constipation.

The Alosetron REMS Program – Please see complete Prescribing Information for Alosetron Tablets for full details.
Warnings and Precautions (cont’d)

Ischemic Colitis

- Some patients have experienced symptoms of ischemic colitis without warning.

- Ischemic colitis has been reported in patients receiving alosetron in clinical trials as well as during marketed use of the drug.

- In IBS clinical trials:
  - cumulative incidence of ischemic colitis in women receiving alosetron was:
    0.2%, or 2 per 1,000 patients (95% CI 1 to 3), over 3 months
    0.3%, or 3 per 1,000 patients (95% CI 1 to 4), over 6 months
  - patient experience in controlled clinical trials is insufficient to estimate the incidence of ischemic colitis in patients taking alosetron for longer than 6 months
Warnings and Precautions (cont’d)

**Ischemic Colitis (cont’d)**

- Alosetron should be discontinued immediately in patients with signs of ischemic colitis, e.g., rectal bleeding, bloody diarrhea, or new or worsening abdominal pain.

- Because ischemic colitis can be life threatening, patients with signs or symptoms of ischemic colitis should be evaluated promptly and have appropriate diagnostic testing performed.

- Treatment with alosetron should not be resumed in patients who develop ischemic colitis.
Contraindications

• Alosetron should not be initiated in patients with constipation.

• Alosetron is contraindicated in patients with a history of:
  – chronic or severe constipation or sequelae from constipation;
  – intestinal obstruction, stricture, toxic megacolon, gastrointestinal perforation, and/or adhesions;
  – ischemic colitis, impaired intestinal circulation, thrombophlebitis, or hypercoagulable state;
  – Crohn’s disease or ulcerative colitis;
  – diverticulitis;
  – severe hepatic impairment.

The Alosetron REMS Program - Please see complete Prescribing Information for Alosetron Tablets for full details.
Contraindications (cont’d)

- Alosetron should not be used by patients who are unable to understand or comply with the Patient Acknowledgement Form.

- Concomitant administration of alosetron with fluvoxamine is contraindicated.
In vivo data suggest that alosetron is primarily metabolized by cytochrome P450 (CYP) 1A2, with minor contributions from CYP3A4 and CYP2C9. Therefore, inducers or inhibitors of these enzymes may change the clearance of alosetron.

- Concomitant administration of alosetron and fluvoxamine is contraindicated.
- Concomitant administration of alosetron and moderate CYP1A2 inhibitors, including quinolone antibiotics and cimetidine, has not been evaluated, but should be avoided unless clinically necessary because of similar potential drug interactions.
Drug Interactions (cont’d)

• Caution should be used when alosetron and ketoconazole are administered concomitantly.

• Coadministration of alosetron and strong CYP3A4 inhibitors, such as clarithromycin, telithromycin, protease inhibitors, voriconazole, and itraconazole has not been evaluated but should be undertaken with caution because of similar potential drug interactions.

• The effect of induction or inhibition of other pathways on exposure to alosetron and its metabolites is not known.

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Use in Specific populations

• Pregnancy Category B.

• It is not known whether alosetron is excreted in human milk; caution should be exercised when alosetron is administered to a nursing woman.

• Safety and effectiveness in pediatric patients have not been established.

• Postmarketing experience suggests that elderly patients may be at greater risk for complications of constipation; therefore, appropriate caution and follow-up should be exercised if alosetron is prescribed for these patients.
Use in Specific populations (cont’d)

• Increased exposure to alosetron and/or its metabolites is likely to occur in patients with hepatic impairment. Alosetron should not be used in patients with severe hepatic impairment and should be used with caution in patients with mild or moderate hepatic impairment.
### Adverse Reactions Reported in ≥ 1% of IBS Patients

<table>
<thead>
<tr>
<th>Gastrointestinal Adverse Reactions</th>
<th>Alosetron 1 mg BID (n=8,328b)</th>
<th>Placebo (n=2,363)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Constipation</td>
<td>29%</td>
<td>6%</td>
</tr>
<tr>
<td>Abdominal discomfort and pain</td>
<td>7%</td>
<td>4%</td>
</tr>
<tr>
<td>Nausea</td>
<td>6%</td>
<td>5%</td>
</tr>
<tr>
<td>GI discomfort and pain</td>
<td>5%</td>
<td>3%</td>
</tr>
<tr>
<td>Abdominal distention</td>
<td>2%</td>
<td>1%</td>
</tr>
<tr>
<td>Regurgitation and reflux</td>
<td>2%</td>
<td>2%</td>
</tr>
<tr>
<td>Hemorrhoids</td>
<td>2%</td>
<td>1%</td>
</tr>
</tbody>
</table>

a Reported in ≥1% of alosetron patients and occurring more frequently on alosetron 1 mg twice-a-day than on placebo.
b Data reported from 22 repeat-dose studies in patients with IBS treated for 8 to 24 weeks.
c P<0.0001 vs placebo.
Adverse Reactions

Constipation is a frequent and dose-related side effect of treatment with alosetron.

• In clinical studies constipation was reported in ~29% of patients with IBS treated with alosetron 1 mg twice daily (n=9,316).
  – The effect was statistically significant compared with placebo (P<0.0001);
  – 11% of patients treated with alosetron 1 mg twice daily withdrew from the studies due to constipation.

• Although the number of IBS patients treated with alosetron 0.5 mg twice daily is relatively small (n=243), 11% of patients reported constipation and 4% of patients withdrew from clinical studies due to constipation.
Overdosage

• No specific antidote available for overdose of alosetron.

• Patients should be managed with appropriate supportive therapy.
Section 4:
How to Prescribe and Dispense Alosetron Tablets
Dosage and Administration

For safety reasons, only prescribers who enroll in the Alosetron REMS Program can prescribe alosetron.

- **Usual Dose in Adults**
  - To lower the risk of constipation, alosetron should be started at 0.5 mg twice-a-day.
  - Patients well controlled on 0.5 mg twice-a-day may be maintained on this regimen.
  - If, after 4 weeks, the 0.5 mg twice-a-day dosage is tolerated but does not adequately control IBS symptoms, increase dose to 1 mg twice-a-day, the dose used in controlled clinical trials.
Dosage and Administration (cont’d)

• Usual Dose in Adults
  – **Alosetron should be started at a dosage of 0.5 mg twice-a-day. Patients controlled on this dose may be maintained on this regimen.**
  – If after 4 weeks, the 0.5 mg twice-a-day dosage is well tolerated but does not adequately control the IBS symptoms, then the dosage can be increased up to 1 mg twice-a-day.
  – **Alosetron should be discontinued in patients who have not had adequate control of IBS symptoms after 4 weeks of treatment with 1 mg twice-a-day.**
  – Alosetron should be discontinued immediately in patients who develop constipation or signs of ischemic colitis.
  – Alosetron should not be restarted in patients who develop ischemic colitis.
Dosage and Administration (cont’d)

• Clinical trial and postmarketing experience suggest that debilitated patients or patients taking additional medications that decrease GI motility may be at greater risk of serious complications of constipation.

• Therefore, appropriate caution and follow-up should be exercised if alosetron is prescribed for these patients.

• Alosetron can be taken with or without food.

The Alosetron REMS Program - Please see complete Prescribing Information for Alosetron Tablets for full details.
Section 5:
Alosetron
REMS Program
Enrolling in the Alosetron REMS Program

• Prescribers must read the full PI and understand the benefits and risks of treatment with alosetron for severe diarrhea-predominant IBS.

• Prescribers then complete the Prescriber Enrollment Form at www.AlosetronREMS.com or return it by mail or by fax.

• **The form must be returned to the sponsor before a prescriber can be considered enrolled in the Alosetron REMS Program.**

• Alosetron REMS Program Kits, including Medication Guides and stickers that need to be affixed to every prescription, are provided after enrollment.

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Overview of Prescriber Responsibilities

Step 1: Review and provide Medication Guide to the patient.

Step 2: Have the patient complete the Patient Acknowledgment Form.
Place the original in the patient’s medical record and give a copy to the patient.

Step 3: Provide patient with written prescription with affixed Alosetron REMS sticker.
Refills are permitted on written prescription.

The Alosetron REMS Program - Please see complete Prescribing Information for Alosetron Tablets for full details.
Informing the Patients about the Requirements of the Alosetron REMS Program

- Once an appropriate patient has been selected for therapy, review the Medication Guide and explain the risks of alosetron therapy to the patient.

- Answer any questions the patient may have.

- Instruct the patient to complete the Patient Acknowledgement Form. Place the original signed form in the patient’s medical record and give a copy to the patient.

Informing the Patients about the Requirements of the Alosetron REMS Program

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- Answer any questions the patient may have.

- Instruct the patient to complete the Patient Acknowledgement Form. Place the original signed form in the patient’s medical record and give a copy to the patient.
The Alosetron REMS Sticker

• The Alosetron REMS sticker must be affixed to every prescription:

  Alosetron Tablets
  The sticker indicates that this prescription is in compliance with the
  Alosetron REMS
  REFILL PERMITTED

• Refills may be written on prescriptions.

• No telephone, facsimile, or computerized prescriptions are permitted with this program.
Patient Responsibilities

Patients should be instructed to:

• read the Medication Guide before starting alosetron and each time they refill their prescription.

• not take alosetron if they are constipated.

• immediately discontinue alosetron and contact their prescriber if they become constipated or have symptoms of ischemic colitis such as new or worsening abdominal pain, bloody diarrhea, or blood in the stool.

• immediately contact their prescriber again if their constipation does not resolve after discontinuation of alosetron.
Patient Responsibilities (cont’d)

Patients should be instructed to:

• resume alosetron only if their constipation has resolved and after discussion with and the agreement of their treating prescriber.

• stop taking alosetron and contact their prescriber if alosetron does not adequately control IBS symptoms after 4 weeks of taking 1 mg twice-a-day.

The Alosetron REMS Program - Please see complete Prescribing Information for Alosetron Tablets for full details.
Encourage the Patients to Enroll in the Voluntary Follow-Up Survey for Alosetron

- Sponsors offer a patient survey to monitor the process of prescribing alosetron in clinical practice.

- The survey is important for monitoring the Alosetron REMS Program.

- The Alosetron REMS Program Patient Follow-up Survey Pre-Enrollment Form will be sent to prescribers after enrollment and is available upon request.

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• You have now reached the end of this Education Slide Deck.

• If you have questions or want to enroll in the Alosetron REMS Program, please call 1-844-267-8675 or visit www.AlosetronREMS.com.