

## Oxycodone Hydrochloride Oral Solution, USP 5 mg/5 mL

On 01/18/2017, VistaPharm recalled (Lot Numbers: 427900, 426700, 424800, 423600, 420800, 416300, 407700, 407300, 405900, 403900, 426900, 404700, 390200) of Oxycodone Hydrochloride Oral Solution, USP 5 mg/5 mL (NDC# 66689-401-50, 66689-403-16), the purified water used to manufacture the drug products may have been contaminated with Burkholderia cepacia. The U.S. Food and Drug Administration (FDA) has issued a Class II of the affected medications. More information about the recall is at: <http://www.fda.gov/Safety/Recalls/EnforcementReports/default.htm>.

The detailed information of the recalled product is listed below:

**Recalled Drug:** Oxycodone Hydrochloride Oral Solution, USP 5 mg/5 mL  
**NDC Number:** 66689-401-50, 66689-403-16  
**Lot Numbers:** 427900, 426700, 424800, 423600, 420800, 416300, 407700, 407300, 405900, 403900, 426900, 404700, 390200  
**Expiration Date:** 11/17, 10/17, 09/17, 08/17, 06/17, 05/17, 02/17

### What you should do:

- **Do not continue to use Oxycodone Hydrochloride Oral Solution, USP 5 mg/5 mL if it has been recalled**
- Check your prescription label to see if you have Oxycodone Hydrochloride Oral Solution, USP 5 mg/5 mL.
- If you have the medication, contact the pharmacy that you received it from.
- Adverse events from the use of prescription drugs should be reported to FDA by visiting [www.fda.gov/medwatch/report.htm](http://www.fda.gov/medwatch/report.htm) or calling 1-800-FDA-0178.

## GlucaGen HypoKit (glucagon [rDNA origin] for injection).

On 09/08/2016, Novo Nordisk recalled (FS6X270, FS6X296, FS6X538, FS6X597, FS6X797, FS6X875) of GlucaGen HypoKit (glucagon [rDNA origin] for injection) (NDC# 0169-7065-15) Defective delivery system: detached needles on the syringe in the kit. The U.S. Food and Drug Administration (FDA) has issued a Class I of the affected medications. More information about the recall is at: <http://www.fda.gov/Safety/Recalls/EnforcementReports/default.htm>.

The detailed information of the recalled product is listed below:

**Recalled Drug:** GlucaGen HypoKit (glucagon [rDNA origin] for injection).  
**NDC Number:** 0169-7065-15  
**Lot Numbers:** FS6X270, FS6X296, FS6X538, FS6X597, FS6X797, FS6X875  
**Expiration Date:** 9/30/2017

### What you should do:

- **Do not continue to use GlucaGen HypoKit (glucagon [rDNA origin] for injection) if it has been recalled.**
- Check your prescription label to see if you have GlucaGen HypoKit (glucagon [rDNA origin] for injection).
- If you have the medication, contact the pharmacy that you received it from.
- Adverse events from the use of prescription drugs should be reported to FDA by visiting [www.fda.gov/medwatch/report.htm](http://www.fda.gov/medwatch/report.htm) or calling 1-800-FDA-0178.

## Senna Syrup (sennosides) 8.8 mg, 8 fl. oz. (237 mL) bottle

On 8/8/2016, Major Pharmaceuticals recalled Senna Syrup (sennosides) Syrup (NDC# 0904-6289-09). This recall was initiated as a precautionary measure due to a potential risk of product contamination with the bacteria B. cepacia. The U.S. Food and Drug Administration (FDA) has issued a Class II of the affected medications. More information about the recall is at: <http://www.fda.gov/Safety/Recalls/EnforcementReports/default.htm>.

The detailed information of the recalled product is listed below:

**Recalled Drug:** Senna Syrup (sennosides) 8.8 mg  
**NDC Number:** 0904-6289-09  
**Lot Number:** 20391517, 20391518, 20391519, 20391601, 20391602, 20391604, 20391605, 20391608.  
**Expiration Date:** 09/17, 10/17, 11/17, 01/18, 02/18, 03/18, 06/18.

**What you should do:**

- **Do not continue to use Senna Syrup (sennosides) if it has been recalled**
- Check your prescription label to see if you have Senna Syrup (sennosides).
- If you have the medication, contact the pharmacy that you received it from.
- Adverse events from the use of prescription drugs should be reported to FDA by visiting [www.fda.gov/medwatch/report.htm](http://www.fda.gov/medwatch/report.htm) or calling 1-800-FDA-0178.

## Senexon Liquid (sennosides) 8.8 mg, 8 fl oz. (237 mL)

On 8/8/2016, Rugby Laboratories recalled Senexon Liquid (sennosides) (NDC# 0536-1000-59). This recall was initiated as a precautionary measure due to a potential risk of product contamination with the bacteria *B. cepacia*. The U.S. Food and Drug Administration (FDA) has issued a Class II of the affected medications. More information about the recall is at: <http://www.fda.gov/Safety/Recalls/EnforcementReports/default.htm>.

The detailed information of the recalled product is listed below:

**Recalled Drug:** Senexon Liquid (sennosides) 8.8 mg  
**NDC Number:** 0536-1000-59

**What you should do:**

- **Do not continue to use Senexon Liquid (sennosides) 8.8 mg if it has been recalled**
- Check your prescription label to see if you have < Senexon Liquid (sennosides) 8.8 mg.
- If you have the medication, contact the pharmacy that you received it from.
- Adverse events from the use of prescription drugs should be reported to FDA by visiting [www.fda.gov/medwatch/report.htm](http://www.fda.gov/medwatch/report.htm) or calling 1-800-FDA-0178.

## Diocto Syrup (docusate sodium), 60 mg/15 mL

On 8/8/2016, Rugby Laboratories recalled Diocto Syrup (docusate sodium), 60 mg/15 mL (NDC# 0536-1001-85). This recall was initiated as a precautionary measure due to a potential risk of product contamination with the bacteria *B. cepacia*. The U.S. Food and Drug Administration (FDA) has issued a Class II of the affected medications. More information about the recall is at: <http://www.fda.gov/Safety/Recalls/EnforcementReports/default.htm>.

The detailed information of the recalled product is listed below:

**Recalled Drug:** Diocto Syrup (docusate sodium), 60 mg/15 mL  
**NDC Number:** 0536-1001-85

**What you should do:**

- **Do not continue to use Diocto Syrup (docusate sodium), 60 mg/15 mL if it has been recalled**
- Check your prescription label to see if you have Diocto Syrup (docusate sodium), 60 mg/15 mL.
- If you have the medication, contact the pharmacy that you received it from.
- Adverse events from the use of prescription drugs should be reported to FDA by visiting [www.fda.gov/medwatch/report.htm](http://www.fda.gov/medwatch/report.htm) or calling 1-800-FDA-0178.

## Sennazon (sennosides) Syrup, 8.8 mg, 8 fl. oz. (237 mL) bottle

On 8/8/2016, Bayshore Pharmaceuticals recalled Sennazon (sennosides) Syrup (NDC# 76518-100-08). This recall was initiated as a precautionary measure due to a potential risk of product contamination with the bacteria *B. cepacia*. The U.S. Food and Drug Administration (FDA) has issued a Class II of the affected medications. More information about the recall is at: <http://www.fda.gov/Safety/Recalls/EnforcementReports/default.htm>.

The detailed information of the recalled product is listed below:

<b>Recalled Drug:</b> Sennazon (sennosides) Syrup <b>NDC Number:</b> 76518-100-08
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### What you should do:

- **Do not continue to use Sennazon (sennosides) Syrup if it has been recalled**
- Check your prescription label to see if you have Sennazon (sennosides) Syrup.
- If you have the medication, contact the pharmacy that you received it from.
- Adverse events from the use of prescription drugs should be reported to FDA by visiting [www.fda.gov/medwatch/report.htm](http://www.fda.gov/medwatch/report.htm) or calling 1-800-FDA-0178.

## Albuterol Sulfate Syrup, 2 mg/ 5 mL, 473 mL bottle

On 06/22/2016, Teva Pharmaceuticals recalled Lot # 95113 of Albuterol Sulfate Syrup, 2 mg/ 5 mL, 473 mL bottle (NDC# 0093-0661-16) due to the presence of foreign substances; presence of black particles described generically as cellulose-based bundles of brown fibrous material. The U.S. Food and Drug Administration (FDA) has issued a Recall Class II of the affected medications. More information about the recall can be found at:

<http://www.fda.gov/Safety/Recalls/EnforcementReports/default.htm>.

The detailed information of the recalled product is listed below:

<b>Recalled Drug:</b> Albuterol Sulfate Syrup, 2 mg/ 5 mL <b>NDC Number:</b> 0093-0661-16 <b>Lot Numbers:</b> 95113 <b>Expiration Date:</b> 01/2017
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### What you should do:

- **Do not continue to use Albuterol Sulfate Syrup, 2 mg/ 5 mL if it has been recalled**
- Check your prescription label to see if you have Albuterol Sulfate Syrup, 2 mg/ 5 mL.
- If you have the medication, contact the pharmacy that you received it from.
- Adverse events from the use of prescription drugs should be reported to FDA by visiting [www.fda.gov/medwatch/report.htm](http://www.fda.gov/medwatch/report.htm) or calling 1-800-FDA-0178.



