

Vesitirim® 10 mg film-coated tablets

(solifenacin succinate)

Package Leaflet: Information for the user

Read all of this leaflet carefully before you start taking this medicine because it contains important information for you.

- * Keep this leaflet. You may need to read it again.
- * If you have any further questions, ask your doctor or pharmacist.
- * This medicine has been prescribed for you only. Do not pass it on to others. It may harm them, even if their signs of illness are the same as yours.
- * If you get any side effects, talk to your doctor or pharmacist. This includes any possible side effects not listed in this leaflet. See section 4.

What is in this leaflet

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1 What Vesitirim is and what it is used for

The active substance of Vesitirim belongs to the group of anticholinergics. These medicines are used to reduce the activity of an overactive bladder. This enables you to wait longer before having to go to the bathroom and increases the amount of urine that can be held by your bladder.

Vesitirim is used to treat the symptoms of a condition called overactive bladder. These symptoms include: having a strong, sudden urge to urinate without prior warning, having to urinate frequently or wetting yourself because you could not get to the bathroom in time.

2 What you need to know before you take Vesitirim

Do not take Vesitirim

- * if you have an inability to pass water or to empty your bladder completely (urinary retention)
- * if you have a severe stomach or bowel condition (including toxic megacolon, a complication associated with ulcerative colitis)
- * if you suffer from the muscle disease called myasthenia gravis, which can cause an extreme weakness of certain muscles
- * if you suffer from increased pressure in the eyes, with gradual loss of eye sight (glaucoma)
- * if you are allergic to solifenacin or any of the other ingredients of this medicine (listed in section 6)
- * if you are undergoing kidney dialysis
- * if you have severe liver disease
- * if you suffer from severe kidney disease or moderate liver disease AND at the same time are being treated with medicines that may decrease the removal of Vesitirim from the body (for example, ketoconazole). Your doctor or pharmacist will have informed you if this is the case.

Inform your doctor if you have or ever had any of the above mentioned conditions before treatment with Vesitirim starts.

Warnings and precautions

Talk to your doctor or pharmacist before taking Vesitirim

- * if you have trouble emptying your bladder (= bladder obstruction) or have difficulty in passing urine (e.g. a thin urine flow). Risk of accumulation of urine in the bladder (urinary retention) is much higher.
- * if you have some obstruction of the digestive system (constipation).
- * if you are at risk of your digestive system slowing down (stomach and bowel movements). Your doctor will have informed you if this is the case.
- * if you suffer from severe kidney disease.
- * if you have moderate liver disease.

- * if you have a stomach tear (hiatus hernia) or heartburn.
- * if you have a nervous disorder (autonomic neuropathy).

Children and adolescents

Vesitirim is not to be used in children or adolescents under 18 years.

Inform your doctor if you have or ever had any of the above mentioned conditions before treatment with Vesitirim starts.

Before starting Vesitirim, your doctor will assess whether there are other causes for your need to pass urine frequently (for example heart failure (insufficient pumping power of the heart) or kidney disease). If you have a urinary tract infection, your doctor will prescribe you an antibiotic (a treatment against particular bacterial infections).

Other medicines and Vesitirim

Please tell your doctor or pharmacist if you are taking or have recently taken or might take other medicines.

It is especially important to inform your doctor if you are taking:

- * other anticholinergic medicines, effects and side effects of both medications can be enhanced.
- * cholinergics as they can reduce the effect of Vesitirim.
- * medicines, like metoclopramide and cisapride, which make the digestive system work faster. Vesitirim can reduce their effect.
- * medicines, like ketoconazole, ritonavir, nelfinavir, intraconazole, verapamil and diltiazem, which decrease the rate at which Vesitirim is broken down by the body
- * medicines like rifampicin, phenytoin and carbamazepine, as they may increase the rate at which Vesitirim is broken down by the body.
- * medicines such as bisphosphonates, that can cause or exacerbate inflammation of the gullet (oesophagitis).

Vesitirim with food and drink

Vesitirim can be taken with or without food, depending on your preference.

Pregnancy and breast-feeding

You should not use Vesitirim if you are pregnant unless clearly necessary. Do not use Vesitirim if you are breast-feeding as solifenacin may get into your breast milk.

Ask your doctor or pharmacist for advice before taking this medicine.

Driving and using machines

Vesitirim may cause blurred vision and sometimes sleepiness or tiredness. If you suffer from any of these side effects, do not drive or operate machinery.

Vesitirim contains lactose. If you have been told by your doctor that you have a rare hereditary problem of galactose intolerance, Lapp lactase deficiency or glucose-galactose malabsorption you should not use this medicine.

3 How to take Vesitirim

Instructions for proper use

Always take this medicine exactly as your doctor has told you. Check with your doctor or pharmacist if you are not sure.

You should swallow the whole tablet with some liquid. It can be taken with or without food, according to your preference. Do not crush the tablets.

The usual dose is 5 mg per day, unless your doctor told you to take 10 mg per day.

If you take more Vesitirim than you should

If you have taken too much Vesitirim or if a child has accidentally taken Vesitirim, contact your doctor or pharmacist immediately.

Symptoms of overdose may include: headache, dry mouth, dizziness, drowsiness and blurred vision, perceiving things that are not there (hallucinations), over- excitability, seizures (convulsions), difficulty breathing, elevated heart rate (tachycardia), accumulation of urine in the bladder (urinary retention) and dilated pupils (mydriasis).

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(solifenacin succinate)

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If you forget to take Vesitirim

If you forget to take a dose at the usual time, take it as soon as you remember, unless it is time to take your next dose. Never take more than one dose per day. If you are in doubt, always consult your doctor or pharmacist.

If you stop taking Vesitirim

If you stop taking Vesitirim, your symptoms of overactive bladder may return or worsen. Always consult your doctor, if you are considering stopping the treatment.

If you have any further questions on the use of this product, ask your doctor or pharmacist.

4 Possible side effects

Like all medicines, this medicine can cause side effects, although not everybody gets them.

If you experience an allergic attack, or a severe skin reaction (e.g. blistering and peeling of the skin), you must inform your doctor or pharmacist immediately

Angioedema (skin allergy that results in the swelling that occurs in the tissue just below the surface of the skin) with airway obstruction (difficulty in breathing) has been reported in some patients on solifenacin succinate (Vesitirim). If angioedema occurs, solifenacin succinate (Vesitirim) should be discontinued immediately and appropriate therapy and/or measures should be taken.

Vesitirim may cause the following other side effects:

Very common (may affect more than 1 in 10 people)

- * dry mouth

Common (may affect up to 1 in 10 people)

- * blurred vision
- * constipation, nausea, indigestion with symptoms such as abdominal fullness, abdominal pain, burping, nausea, and heartburn (dyspepsia), stomach discomfort

Uncommon (may affect up to 1 in 100 people)

- * urinary tract infection, bladder infection
- * sleepiness, impaired sense of taste (dysgeusia)
- * dry (irritated) eyes
- * dry nasal passages
- * reflux disease (gastro-oesophageal reflux), dry throat
- * dry skin
- * difficulty in passing urine
- * tiredness, accumulation of fluid in the lower legs (oedema)

Rare (may affect up to 1 in 1,000 people)

- * lodging of a large amount of hardened stool in the large intestine (faecal impaction)
- * build up of urine in the bladder due to inability to empty the bladder (urinary retention)
- * dizziness, headache
- * vomiting
- * itching, rash,

Very rare (may affect up to 1 in 10,000 people)

- * hallucinations, confusion
- * allergic rash

Not known (frequency cannot be estimated from the available data)

- * decreased appetite, high levels of blood potassium which can cause abnormal heart rhythm
- * increased pressure in the eyes
- * changes in the electrical activity of the heart (ECG), irregular heartbeat feeling your heartbeat, faster heart beat
- * voice disorder
- * liver disorder
- * muscle weakness
- * renal disorder

If you get any side effects, talk to your doctor or pharmacist. This includes any side effects not listed in this leaflet.

Reporting of side effects

If you get any side effects, talk to your doctor, pharmacist or nurse.

This includes any possible side effects not listed in this leaflet. You can also report side effects directly via HPRA Pharmacovigilance, Earlsfort Terrace, IRL - Dublin 2; Tel: +353 1 6764971; Fax: +353 1 6762517. Website: www.hpra.ie; E-mail: medsafety@hpra.ie. By reporting side effects you can help provide more information on the safety of this medicine.

5 How to store Vesitirim

Keep out of the reach and sight of children.

Do not use Vesitirim after the expiry date which is stated on the carton after EXP. The expiry date refers to the last day of that month. This medicinal product does not require any special storage conditions.

Medicines should not be disposed of via wastewater or household waste.

Ask your pharmacist how to dispose of medicines no longer required.

These measures will help to protect the environment.

6 Contents of the pack and other information

What Vesitirim contains

- * The active substance is solifenacin succinate 10 mg
- * The other ingredients are maize starch, lactose, hypromellose (E464) magnesium stearate, macrogol, talc, titanium dioxide (E171), iron oxide (E172)

What Vesitirim looks like and contents of the pack

Vesitirim 10 mg tablets are round, light pink tablets marked with the company logo and the code "151" on the same side.

Vesitirim tablets are supplied in blister packs of 30 tablets.

The Parallel Product Authorisation Holder is: LTT Pharma Limited, Unit 18, Oxleasow Road, East Moons Moat, Redditch, Worcestershire, B98 0RE.

Repackaged by Lexon (UK) Limited, Unit 18, Oxleasow Road, East Moons Moat, Redditch, Worcestershire, B98 0RE.

Manufactured by ASTELLAS PHARMA EUROPE BV, SYLVIVSWEG 62, 2333 BE LEIDEN, THE NETHERLANDS.

Vesitirim is a registered trademark of Astellas Pharma Europe B.V

Revision date: 07/07/16

POM PPA1562/086/004

Blind or partially sighted?

Is this leaflet hard to see or read?

Phone LTT Pharma Limited,

Tel: +44 (0)1527 505414

for help.

This medicinal product is authorised in the Member States of the EEA under the following names:

Austria, Belgium, Cyprus, Czech Republic, Denmark, Estonia, Finland, France, Greece, Hungary, Iceland, Latvia, Lithuania, Luxemburg, Malta, Netherlands, Norway, Poland, Portugal, Sweden, Slovenia, Slovakia, Spain and United Kingdom: Vesitirim

Italy: Vesiker

Germany: Vesikur

Ireland: Vesitirim