Package leaflet: Information for the user

Ezetator 10 mg/10 mg, filmomhulde tabletten Ezetator 10 mg/20 mg, filmomhulde tabletten Ezetator 10 mg/40 mg, filmomhulde tabletten Ezetator 10 mg/80 mg, filmomhulde tabletten

ezetimibe/atorvastatin

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

- 1. What < Product name > is and what it is used for
- 2. What you need to know before you take < Product name >
- 3. How to take < Product name >
- 4. Possible side effects
- 5. How to store < Product name >
- 6. Contents of the pack and other information

1. What <Product name> is and what it is used for

<Product name> is a medicine to lower increased levels of cholesterol. <Product name> contains ezetimibe and atorvastatin.

<Product name> is used in adults to lower levels of total cholesterol, "bad" cholesterol (LDL cholesterol), and fatty substances called triglycerides in the blood. In addition, <Product name> raises levels of "good" cholesterol (HDL cholesterol).

<Product name> works to reduce your cholesterol in two ways. It reduces the cholesterol absorbed in your digestive tract, as well as the cholesterol your body makes by itself.

Cholesterol is one of several fatty substances found in the bloodstream. Your total cholesterol is made up mainly of LDL and HDL cholesterol.

LDL cholesterol is often called "bad" cholesterol because it can build up in the walls of your arteries forming plaque. Eventually this plaque build-up can lead to a narrowing of the arteries. This narrowing can slow or block blood flow to vital organs such as the heart and brain. This blocking of blood flow can result in a heart attack or stroke.

HDL cholesterol is often called "good" cholesterol because it helps keep the bad cholesterol from building up in the arteries and protects against heart disease.

Triglycerides are another form of fat in your blood that may increase your risk for heart disease.

<Product name> is used for patients who cannot control their cholesterol levels by diet alone. You should stay on your cholesterol lowering diet while taking this medicine.

<Product name> is used in addition to your cholesterol-lowering diet if you have:

- a raised cholesterol level in your blood (primary hypercholesterolaemia [heterozygous familial and non-familial]) or elevated fat levels in your blood (mixed hyperlipidaemia)
 - that is not well controlled with a statin alone;
 - for which you have used a statin and ezetimibe as separate tablets.
- a hereditary illness (homozygous familial hypercholesterolaemia) that increases the cholesterol level in your blood. You may also receive other treatments.
- heart disease. <Product name> reduces the risk of heart attack, stroke, surgery to increase heart blood flow, or hospitalisation for chest pain.

<Product name> does not help you lose weight.

2. What you need to know before you take <Product name>

Do not take < Product name > if

- you are allergic to atorvastatin, ezetimibe or any of the other ingredients of this medicine (listed in section 6),
- you have or have ever had a disease that affects the liver,
- you have had any unexplained abnormal blood tests for liver function,
- you are a woman able to have children and are not using reliable contraception,
- you are pregnant, trying to become pregnant or are breast-feeding,
- you use the combination of glecaprevir/pibrentasvir in the treatment of hepatitis C.

Warnings and precautions

Talk to your doctor or pharmacist before taking <Product name> if:

- you have had a previous stroke with bleeding into the brain, or have small pockets of fluid in the brain from previous strokes,
- you have kidney problems,
- you have an under-active thyroid gland (hypothyroidism),
- you have had repeated or unexplained muscle aches or pains, a personal history or family history of muscle problems,
- you have had previous muscular problems during treatment with other lipid-lowering medicines (e.g. other "statin" or "fibrate" medicines),
- you regularly drink a large amount of alcohol,
- you have a history of liver disease,
- you are older than 70 years,
- you have been told by your doctor that you have an intolerance to some sugars, contact your doctor before taking this product,
- you are taking or have taken in the last 7 days a medicine called fusidic acid, (a medicine for bacterial infection) orally or by injection. The combination of fusidic acid and <Product name> can lead to serious muscle problems (rhabdomyolysis),
- you have or have had myasthenia (a disease with general muscle weakness including in some cases muscles used when breathing), or ocular myasthenia (a disease causing eye muscle weakness) as statins may sometimes aggravate the condition or lead to the occurrence of myasthenia (see section 4).

Contact your doctor promptly if you experience unexplained muscle pain, tenderness, or weakness while taking <Product name>. This is because on rare occasions, muscle problems can be serious, including muscle breakdown resulting in kidney damage. Atorvastatin is known to cause muscle problems, and cases of muscle problems have also been reported with ezetimibe.

Also tell your doctor or pharmacist if you have a muscle weakness that is constant. Additional tests and medicines may be needed to diagnose and treat this.

Check with your doctor or pharmacist before taking <Product name>:

• if you have severe respiratory failure.

If any of these apply to you (or you are not sure), talk to your doctor or pharmacist before taking <Product name> because your doctor will need to carry out a blood test before and possibly during your <Product name> treatment to predict your risk of muscle-related side effects. The risk of muscle-related side effects, e.g. rhabdomyolysis (breakdown of damaged skeletal muscle), is known to increase when certain medicines are taken at the same time (see section 2 "Other medicines and <Product name>").

While you are on this medicine your doctor will monitor you closely if you have diabetes or are at risk of developing diabetes. You are likely to be at risk of developing diabetes if you have high levels of sugars and fats in your blood, are overweight and have high blood pressure.

Tell your doctor about all medical conditions including allergies.

The combined use of <Product name> and fibrates (medicines for lowering cholesterol) should be avoided since the combined use of <Product name> and fibrates has not been studied.

Children

<Product name> is not recommended for children and adolescents.

Other medicines and <Product name>

Tell your doctor or pharmacist if you are taking, have recently taken, or might take any other medicines.

There are some medicines that may change the effect of <Product name> or their effect may be changed by <Product name> (see section 3). This type of interaction could make one or both of the medicines less effective. Alternatively it could increase the risk or severity of side effects, including the important muscle wasting condition known as "rhabdomyolysis" described in section 4:

- ciclosporin (a medicine often used in organ transplant patients),
- erythromycin, clarithromycin, telithromycin, fusidic acid**, rifampicin (medicines for bacterial infections),
- ketoconazole, itraconazole, voriconazole, fluconazole, posaconazole (medicines for fungal infections),
- gemfibrozil, other fibrates, nicotinic acid, derivatives, colestipol, cholestyramine (medicines for regulating lipid levels),
- some calcium channel blockers used for angina or high blood pressure, e.g. amlodipine, diltiazem.
- digoxin, verapamil, amiodarone (medicines to regulate your heart rhythm),
- medicines used in the treatment of HIV, e.g. ritonavir, lopinavir, atazanavir, indinavir, darunavir, the combination of tipranavir/ritonavir, etc. (medicines for AIDS),
- some medicines used in the treatment of hepatitis C, e.g. telaprevir, boceprevir and the combination of elbasvir/grazoprevir,
- daptomycin (a medicine used to treat complicated skin and skin structure infections and bacteraemia).

**If you need to take oral fusidic acid to treat a bacterial infection you will need to temporarily stop using this medicine. Your doctor will tell you when it is safe to restart <Product name>. Taking <Product name> with fusidic acid may rarely lead to muscle weakness, tenderness or pain (rhabdomyolysis). See more information regarding rhabdomyolysis in section 4.

- Other medicines known to interact with the <Product name>:
 - oral contraceptives (medicines for preventing pregnancy)
 - stiripentol (an anticonvulsant medicine for epilepsy)
 - cimetidine (a medicine used for heartburn and peptic ulcers)
 - phenazone (a painkiller)
 - antacids (indigestion products containing aluminium or magnesium)
 - warfarin, phenprocoumon, acenocoumarol or fluindione (medicines to prevent blood clots)
 - colchicine (used to treat gout)
 - St John's Wort (a medicine to treat depression)

<Pre><Product name> with food and alcohol

See section 3 for instructions on how to take <Product name>. Please note the following:

Grapefruit juice

Do not take more than one or two small glasses of grapefruit juice per day because large quantities of grapefruit juice can change the effects of <Product name>.

Alcohol

Avoid drinking too much alcohol while taking this medicine. See section 2 "Warnings and precautions" for details.

Pregnancy and breast-feeding

Do not take <Product name> if you are pregnant, are trying to become pregnant or think you may be pregnant.

Do not take <Product name> if you are able to become pregnant unless you use reliable contraceptive measures. If you get pregnant while taking <Product name>, stop taking it immediately and tell your doctor.

Do not take <Product name> if you are breast-feeding.

The safety of <Product name> during pregnancy and breast-feeding has not yet been proven. Ask your doctor or pharmacist for advice before taking this medicine.

Driving and using machines

<Product name> is not expected to interfere with your ability to drive or to use machinery. However, it should be taken into account that some people may get dizzy after taking <Product name>.

<Product name> contains lactose

<Product name> 10 mg/10 mg, 10 mg/20 mg and 10 mg/40 mg film-coated tablets contain a sugar called lactose. If you have been told by your doctor that you have an intolerance to some sugars, contact your doctor before taking this medicinal product.

<Product name> contains sodium

<Product name> contains less than 1 mmol sodium (23 mg) per tablet, that is to say essentially 'sodium-free'.

3. How to take <Product name>

Always take this medicine exactly as your doctor has told you. Your doctor will determine the appropriate tablet strength for you, depending on your current treatment and your personal risk status. Check with your doctor or pharmacist if you are not sure.

- Before starting <Product name>, you should be on a diet to lower your cholesterol.
- You should keep on this cholesterol-lowering diet while taking < Product name >.

How much to take

The recommended dose is one <Product name> tablet once a day.

When to take

Take <Product name> at any time of the day. You can take it with or without food.

If your doctor has prescribed <Product name> along with cholestyramine or any other bile acid sequestrant (medicines for lowering cholesterol), you should take <Product name> at least 2 hours before or 4 hours after taking the bile acid sequestrant.

If you take more <Product name> than you should

Please contact your doctor or pharmacist.

If you forget to take <Product name>

Do not take an extra dose; just take your normal amount of <Product name> at the usual time the next day.

If you have any further questions on the use of this medicine, 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 any of the following serious side effects or symptoms, stop taking your tablets and tell your doctor immediately or go to the nearest hospital accident and emergency department and take your tablets with you.

- serious allergic reaction which causes swelling of the face, tongue and throat that can cause great difficulty in breathing
- serious illness with severe peeling and swelling of the skin, blistering of the skin, mouth, eyes, genitals and fever; skin rash with pink-red blotches especially on palms of hands or soles of feet, which may blister
- muscle weakness, tenderness, pain, rupture or red-brown discolouration of urine and particularly, if at the same time, you feel unwell or have a high temperature it may be caused by an abnormal muscle breakdown which can be life-threatening and lead to kidney problems
- lupus-like disease syndrome (including rash, joint disorders and effects on blood cells).

You should consult your doctor as soon as possible if you experience problems with unexpected or unusual bleeding or bruising, because this may be suggestive of a liver complaint.

The following common side effects were reported (may affect up to 1 in 10 people):

- diarrhoea,
- muscle aches.

The following uncommon side effects were reported (may affect up to 1 in 100 people):

- the flu,
- depression; trouble sleeping; sleep disorder,
- dizziness; headache; tingling sensation,
- slow heartbeat,
- hot flush,
- shortness of breath,
- abdominal pain; abdominal bloating; constipation; indigestion; flatulence; frequent bowel movements; inflammation of the stomach; nausea; stomach discomfort; upset stomach,
- acne: hives.
- joint pain; back pain; leg cramps; muscle fatigue, spasms, or weakness; pain in arms and legs,

- unusual weakness; feeling tired or unwell; swelling, especially in the ankles (oedema),
- elevations in some laboratory blood tests of liver or muscle (CK) function,
- weight gain.

Additionally, the following side effects have been reported in people taking <Product name>, or ezetimibe or atorvastatin tablets:

- allergic reactions including swelling of the face, lips, tongue, and/or throat that may cause difficulty in breathing or swallowing (which requires treatment immediately),
- raised red rash, sometimes with target-shaped lesions,
- liver problems,
- cough,
- heartburn,
- decreased appetite; loss of appetite,
- high blood pressure,
- skin rash and itching; allergic reactions including rash and hives,
- tendon injury,
- gallstones or inflammation of the gallbladder (which may cause abdominal pain, nausea, vomiting),
- inflammation of the pancreas often with severe abdominal pain,
- reduction in blood cell counts, which may cause bruising/bleeding (thrombocytopenia),
- inflammation of the nasal passages; nose bleed,
- neck pain; pain; chest pain; pain in the throat,
- increases and decreases in blood sugar levels (if you have diabetes you should continue careful monitoring of your blood sugar levels),
- having nightmares,
- numbness or tingling in the fingers and toes,
- reduction of sensation to pain or touch,
- change in sense of taste; dry mouth,
- loss of memory,
- ringing in the ears and/or head; hearing loss,
- vomiting,
- belching,
- hair loss,
- raised temperature,
- urine tests that are positive for white blood cells,
- blurred vision; visual disturbances,
- gynaecomastia (breast enlargement in men),
- myasthenia gravis (a disease causing general muscle weakness including in some cases muscles used when breathing) *,
- ocular myasthenia (a disease causing eye muscle weakness) *,

Possible side effects reported with some statins

- sexual difficulties,
- depression,
- breathing problems including persistent cough and/or shortness of breath or fever,
- diabetes. This is more likely if you have high levels of sugars and fats in your blood, are overweight and have high blood pressure. Your doctor will monitor you while you are taking this medicine,
- muscle pain, tenderness, or weakness that is constant and particularly if, at the same time, you feel unwell or have a high temperature that may not go away after stopping <Product name> (frequency not known).

^{*} talk to your doctor if you experience weakness in your arms or legs that worsens after periods of activity, double vision or drooping of your eyelids, difficulty swallowing, or shortness of breath.

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 the national reporting system listed in <u>Appendix V</u>. By reporting side effects you can help provide more information on the safety of this medicine.

5. How to store < Product name>

- Keep this medicine out of the sight and reach of children.
- Do not take this medicine after the expiry date which is stated on the blister and carton, after EXP. The expiry date refers to the last day of that month.
- This medicine does not require any special storage conditions.

Do not throw away any medicines via wastewater or household waste. Ask your pharmacist how to throw away medicines you no longer use. These measures will help protect the environment.

6. Contents of the pack and other information

What < Product name > contains

The active substances are ezetimibe and atorvastatin.

<Product name> 10 mg/10 mg: Each tablet contains 10 mg of ezetimibe and 10 mg atorvastatin (as atorvastatin calcium trihydrate).

<Product name> 10 mg/20 mg: Each tablet contains 10 mg of ezetimibe and 20 mg atorvastatin (as atorvastatin calcium trihydrate).

<Product name> 10 mg/40 mg: Each tablet contains 10 mg of ezetimibe and 40 mg atorvastatin (as atorvastatin calcium trihydrate).

<Product name> 10 mg/80 mg: Each tablet contains 10 mg of ezetimibe and 80 mg atorvastatin (as atorvastatin calcium trihydrate).

The other ingredients are:

Tablet core:

Cellulose, microcrystalline 101; mannitol; calcium carbonate; croscamellose sodium; hydroxypropylcellulose; polysorbate 80; iron oxide yellow (E172); magnesium stearate; povidone K29/32; sodium laurilsulfate (see section 2 "<Product name> contains sodium").

Tablet Coating

<Product name> 10 mg/10 mg, 10 mg/20 mg, 10 mg/40 mg:

Lactose monohydrate (see section 2 "<Product name> contains lactose"); hypromellose 2910 (E464); titanium dioxide (E171); macrogol 4000 (E1521).

<Product name> 10 mg/80 mg:

Hypromellose 2910 (E464); titanium dioxide (E171); talc (E553b); macrogol 400 (E1521); iron oxide yellow (E172).

What <Product name> looks like and contents of the pack

<Product name> 10 mg/10 mg tablets: White, round, biconvex film coated tablets, with diameter 8.1 mm approximately.

<Product name> 10 mg/20 mg tablets: White, ovaloid, biconvex film coated tablets, with dimensions 11.6 x 7.1 mm approximately.

<Product name> 10 mg/40 mg tablets: White, capsule shape, biconvex film coated tablets, with dimensions 16.1 x 6.1 mm approximately.

<Product name> 10 mg/80 mg tablets: Yellow, oblong, biconvex film coated tablets, with dimensions 19.1 x 7.6 mm approximately.

OPA/Al/PVC//Al blisters and perforated unit dose blisters packed into carton boxes.

For <Product Name> 10 mg/10 mg, 10 mg/20 mg, 10 mg/40 mg film-coated tablets: Pack sizes of 10, 30, 30 x 1, 45 x 1, 90, 90 x 1 and 100 film-coated tablets.

For <Product Name> 10 mg/80 mg film-coated tablets:

Pack sizes of 10, 30, 30 x 1, 45 x 1, 90, 90 x 1 and 100 film-coated tablets and multipacks containing 90 (2 packs of 45), 90 x 1 (2 packs of 45 x 1), 100 (2 packs of 50), 100 x 1 (2 packs of 50 x 1) film-coated tablets.

Not all pack sizes may be marketed.

Houder van de vergunning voor het in de handel brengen en fabrikant

Houder van de vergunning voor het in de handel brengen Teva B.V. Swensweg 5 2031 GA Haarlem

Nederland

Fahrikant

ELPEN Pharmaceutical Co. Inc. Marathonos Avenue 95 190 09 Pikermi Attiki

Griekenland

In het register ingeschreven onder

RVG 130564, filmomhulde tabletten, 10 mg/10 mg RVG 130565, filmomhulde tabletten, 10 mg/20 mg RVG 130566, filmomhulde tabletten, 10 mg/40 mg RVG 130567, filmomhulde tabletten, 10 mg/80 mg

Dit medicijn is geregistreerd in lidstaten van de Europese Economische Ruimte onder de volgende namen

Bulgarije: Avanor Plus 40 mg/10 mg film-coated tablets

Avanor Plus 80 mg/10 mg film-coated tablets

Estland: Atorvastatin/Ezetimibe Teva

Frankrijk: EZETIMIBE / ATORVASTATINE TEVA 10mg/10mg, comprimé pelliculé

EZETIMIBE / ATORVASTATINE TEVA 10mg/20mg, comprimé pelliculé EZETIMIBE / ATORVASTATINE TEVA 10mg/40mg, comprimé pelliculé EZETIMIBE / ATORVASTATINE TEVA 10mg/80mg, comprimé pelliculé

Italië: EZETIMIBE E ATORVASTATINA TEVA

Litouwen: Atorvastatin/Ezetimibe Teva 10mg/10mg plèvele dengtos tabletès

Atorvastatin/Ezetimibe Teva 20mg/10mg plèvele dengtos tabletės Atorvastatin/Ezetimibe Teva 40mg/10mg plèvele dengtos tabletės Atorvastatin/Ezetimibe Teva 80mg/10mg plèvele dengtos tabletės

Letland: Atorvastatin/Ezetimibe Teva 10mg/10mg apvalkotās tabletes

Atorvastatin/Ezetimibe Teva 20mg/10mg apvalkotās tabletes Atorvastatin/Ezetimibe Teva 40mg/10mg apvalkotās tabletes Atorvastatin/Ezetimibe Teva 80mg/10mg apvalkotās tablets

Nederland: Ezetator 10 mg/10 mg, filmomhulde tabletten

Ezetator 10 mg/20 mg, filmomhulde tabletten Ezetator 10 mg/40 mg, filmomhulde tabletten Ezetator 10 mg/80 mg, filmomhulde tabletten

Portugal: Atorvastatina + Ezetimiba ratiopharm

Deze bijsluiter is voor het laatst goedgekeurd in januari 2024.