

COSYREL

Bisoprolol fumarate/perindopril arginine

5/5 mg, 5/10 mg, 10/5 mg, 10/10 mg

NAME OF THE MEDICINAL PRODUCT

Cosyrel 5 mg/5 mg film-coated tablets
Cosyrel 5 mg/10 mg film-coated tablets
Cosyrel 10 mg/5 mg film-coated tablets
Cosyrel 10 mg/10 mg film-coated tablets

QUALITATIVE AND QUANTITATIVE COMPOSITION

One film-coated tablet contains 5 mg of bisoprolol fumarate (equivalent to 4.24 mg bisoprolol) and 5 mg of perindopril arginine (equivalent to 3.395 mg perindopril).

One film-coated tablet contains 5 mg of bisoprolol fumarate (equivalent to 4.24 mg bisoprolol) and 10 mg of perindopril arginine (equivalent to 6.790 mg perindopril).

One film-coated tablet contains 10 mg of bisoprolol fumarate (equivalent to 8.49 mg bisoprolol) and 5 mg of perindopril arginine (equivalent to 3.395 mg perindopril).

One film-coated tablet contains 10 mg of bisoprolol fumarate (equivalent to 8.49 mg bisoprolol) and 10 mg of perindopril arginine (equivalent to 6.790 mg perindopril).

PHARMACEUTICAL FORM

Film-coated tablet

Pink beige, oblong, bilayer scored film-coated tablet of 8.3 mm length and 4.5 mm width, engraved with '  ' on one face and '5/5' on the other face.

Cosyrel 5 mg/5 mg scored tablet can be divided into equal doses.

Pink beige, oblong, bilayer scored film-coated tablet of 9.8 mm length and 5.4 mm width,

engraved with '  ' on one face and '5/10' on the other face.

Cosyrel 5 mg/10 mg scored tablet can be divided into equal doses.

Pink beige, round, bilayer film-coated tablet with a diameter of 7 mm and a curvature radius of 12.7 mm, engraved with '  ' on one face and '10/5' on the other face.

Pink beige, oblong, bilayer film-coated tablet of 10 mm length and 5.7 mm width, engraved with '  ' on one face and '10/10' on the other face.

CLINICAL PARTICULARS

Therapeutic indications

Cosyrel 5 mg/5 mg and Cosyrel 10 mg/5 mg

Cosyrel 5 mg/5 mg and Cosyrel 10 mg/5 mg are indicated as substitution therapy in adult patients who adequately controlled with bisoprolol and perindopril given concurrently at the same dose level, for treatment of:

- Hypertension
- Stable coronary artery disease (in patients with a history of myocardial infarction and/or revascularization)
- Stable chronic heart failure with reduced systolic left ventricular function

Cosyrel 5 mg/10 mg and Cosyrel 10 mg/10 mg

Cosyrel 5 mg/10 mg and Cosyrel 10 mg/10 mg are indicated as substitution therapy in adult patients who adequately controlled with bisoprolol and perindopril given concurrently at the same dose level, for treatment of:

- Hypertension
- Stable coronary artery disease (in patients with a history of myocardial infarction and/or revascularization)

Posology and method of administration

Posology

The usual posology is one tablet once daily.

Patients should be stabilized with bisoprolol and perindopril at the same dose level for at least 4 weeks. The fixed dose combination is not suitable for initial therapy.

If a change of posology is required, titration should be done with the individual components.

Special population

Renal impairment (see section [Special warnings and precautions for use](#) and [Pharmacokinetic properties](#))

Cosyrel is not suitable for patients with renal impairment.

Individual dose titration with the monocomponent is recommended.

Hepatic impairment (see section [Special warnings and precautions for use](#) and [Pharmacokinetic properties](#))

No dosage adjustment is necessary in patients with hepatic impairment.

Elderly

Cosyrel should be administered according to the renal function.

Paediatric population

The safety and efficacy of Cosyrel in children and adolescents have not been established. No data are available. Therefore, use in children and adolescents is not recommended.

Method of administration

Cosyrel tablet should be taken as a single dose once daily in the morning before a meal.

Contraindications

- Hypersensitivity to the active substances, or to any of the excipients, or to any other

angiotensin converting enzyme (ACE) inhibitor

- Acute heart failure or during episodes of heart failure decompensation requiring *i.v.* inotropic therapy
- Cardiogenic shock
- Second or third degree AV block (without pacemaker)
- Sick sinus syndrome
- Sinoatrial block
- Symptomatic bradycardia
- Symptomatic hypotension
- Severe bronchial asthma or severe chronic obstructive pulmonary disease
- Severe forms of peripheral arterial occlusive disease or severe forms of Raynaud's syndrome
- Untreated phaeochromocytoma (see section [Special warnings and precautions for use](#))
- Metabolic acidosis
- History of angioedema associated with previous ACE inhibitor therapy (see section [Special warnings and precautions for use](#))
- Hereditary or idiopathic angioedema
- Second and third trimesters of pregnancy (see section [Special warnings and precautions for use](#) and [Fertility, pregnancy and lactation](#))
- Concomitant use of Cosyrel with aliskiren-containing products in patients with diabetes mellitus or renal impairment (GFR < 60 ml/min/1.73m²) (see section [Special warnings and precautions for use](#), [Interaction with other medicinal products and other forms of interaction](#) and [Pharmacodynamic properties](#)).
- Concomitant use with sacubitril/valsartan therapy. [Cosyrel must not be initiated earlier than 36 hours after the last dose of sacubitril/valsartan](#) (see section [Special warnings and precautions for use](#) and [Interaction with other medicinal products and other forms of interaction](#))
- Extracorporeal treatments leading to contact of blood with negatively charged surfaces (see section [Interaction with other medicinal products and other forms of interaction](#))

- Significant bilateral renal artery stenosis or stenosis of the artery to a single functioning kidney (see section *Special warnings and precautions for use*)

Special warnings and precautions for use

All warnings and precautions for use related to each component are applicable to Cosyrel

Hypotension:

ACE inhibitors may cause a fall in blood pressure. Symptomatic hypotension is seen rarely in uncomplicated hypertensive patients and is more likely to occur in patients who have been volume-depleted e.g. by diuretic therapy, dietary salt restriction, dialysis, diarrhoea or vomiting, or who have severe renin-dependent hypertension (see sections *Interaction with other medicinal products and other forms of interaction* and *undesirable effects*). In patients with symptomatic heart failure, with or without associated renal insufficiency, symptomatic hypotension has been observed. This is most likely to occur in those patients with more severe degrees of heart failure, as reflected by the use of high doses of loop diuretics, hyponatraemia or functional renal impairment. In patients at increased risk of symptomatic hypotension, initiation of therapy and dose adjustment should be closely monitored. Similar considerations apply to patients with ischaemic heart or cerebrovascular disease in whom an excessive fall in blood pressure could result in a myocardial infarction or cerebrovascular accident.

If hypotension occurs, the patient should be placed in the supine position and, if necessary, should receive an intravenous infusion of sodium chloride 9 mg/ml (0.9%) solution. A transient hypotensive response is not a contraindication to further doses, which can be given usually without difficulty once the blood pressure has increased after volume expansion. In some patients with congestive heart failure who have normal or low blood pressure, additional lowering of systemic blood pressure may occur with perindopril. This effect is anticipated and is usually not a reason to

discontinue treatment. If hypotension becomes symptomatic, a reduction of dose or gradual discontinuation of treatment, using the individual components, may be necessary.

Hypersensitivity/Angioedema:

Angioedema of the face, extremities, lips, mucous membranes, tongue, glottis and/or larynx has been reported rarely in patients treated with ACE inhibitors, including perindopril (see section *undesirable effects*). This may occur at any time during therapy. In such cases, Cosyrel should promptly be discontinued. Therapy with beta-blocker must be continued. Appropriate monitoring should be initiated and continued until complete resolution of symptoms has occurred. In those instances where swelling was confined to the face and lips the condition generally resolved without treatment, although antihistamines have been useful in relieving symptoms.

Angioedema associated with laryngeal oedema may be fatal. Where there is involvement of the tongue, glottis or larynx, likely to cause airway obstruction, emergency therapy should be administered promptly. This may include the administration of adrenaline and/or the maintenance of a patent airway. The patient should be under close medical supervision until complete and sustained resolution of symptoms has occurred.

Patients with a history of angioedema unrelated to ACE inhibitor therapy may be at increased risk of angioedema while receiving an ACE inhibitor (see section *Contraindications*).

Intestinal angioedema has been reported rarely in patients treated with ACE inhibitors. These patients presented with abdominal pain (with or without nausea or vomiting); in some cases there was no prior facial angioedema and C-1 esterase levels were normal. The angioedema was diagnosed by procedures including abdominal CT scan, or ultrasound or at surgery and symptoms resolved after stopping the ACE inhibitor. Intestinal angioedema should be included in the differential diagnosis of patients on ACE inhibitors presenting with abdominal pain.

The combination of perindopril with sacubitril/valsartan is contraindicated due to the increased risk of angioedema (see section *Contraindications*). Sacubitril/valsartan must not be initiated until 36 hours after taking the last dose of perindopril therapy. If treatment with sacubitril/valsartan is stopped, perindopril therapy must not be initiated until 36 hours after the last dose of sacubitril/valsartan (see sections *Contraindications* and *Interaction with other medicinal products and other forms of interaction*). Concomitant use of ACE inhibitors with NEP inhibitors (e.g. racecadotril), mTOR inhibitors (e.g. sirolimus, everolimus, temsirolimus) and gliptins (e.g. linagliptin, saxagliptin, sitagliptin, vildagliptin) may lead to an increased risk of angioedema (e.g. swelling of the airways or tongue, with or without respiratory impairment) (see *Interaction with other medicinal products and other forms of interaction*). Caution should be used when starting racecadotril, mTOR inhibitors (e.g. sirolimus, everolimus, temsirolimus) and gliptins (e.g. linagliptin, saxagliptin, sitagliptin, vildagliptin) in a patient already taking an ACE inhibitor.

Hepatic failure:

Rarely, ACE inhibitors have been associated with a syndrome that starts with cholestatic jaundice and progresses to fulminant hepatic necrosis and (sometimes) death. The mechanism of this syndrome is not understood. Patients receiving ACE inhibitors who develop jaundice or marked elevations of hepatic enzymes should discontinue the ACE inhibitor and receive appropriate medical follow-up (see section *undesirable effects*).

Race:

ACE inhibitors cause a higher rate of angioedema in black patients than in non-black patients.

As with other ACE inhibitors, perindopril may be less effective in lowering blood pressure in black people than in non-blacks, possibly

because of a higher prevalence of low-renin states in the black hypertensive population.

Cough:

Cough has been reported with the use of ACE inhibitors. Characteristically, the cough is non-productive, persistent and resolves after discontinuation of therapy. ACE inhibitor induced cough should be considered as part of the differential diagnosis of cough.

Hyperkalaemia:

Elevations in serum potassium have been observed in some patients treated with ACE inhibitors, including perindopril, ACE inhibitors can cause hyperkalaemia because they inhibit the release of aldosterone. The effect is usually not significant in patients with normal renal function. Risk factors for the development of hyperkalemia include those with renal insufficiency, worsening of renal function, age (> 70 years), diabetes mellitus, intercurrent events, in particular dehydration, acute cardiac decompensation, metabolic acidosis and concomitant use of potassium-sparing diuretics (e.g. spironolactone, eplerenone, triamterene, or amiloride), potassium supplements or potassium-containing salt substitutes; or those patients taking other drugs associated with increases in serum potassium (e.g. heparin, co-trimoxazole also known as trimethoprim/sulfamethoxazole) and especially aldosterone antagonists or angiotensin-receptor blockers. The use of potassium supplements, potassium-sparing diuretics, or potassium-containing salt substitutes particularly in patients with impaired renal function may lead to a significant increase in serum potassium. Hyperkalemia can cause serious, sometimes fatal arrhythmias. Potassium-sparing diuretics and angiotensin-receptor blockers should be used with caution in patients receiving ACE inhibitors, and serum potassium and renal function should be monitored. If concomitant use of the above-mentioned agents is deemed appropriate, they should be used with caution and with frequent monitoring of serum potassium (see section

Interaction with other medicinal products and other forms of interaction).

Combination with lithium:

The combination of lithium and perindopril is generally not recommended (see section *Interaction with other medicinal products and other forms of interaction*).

Combination with potassium sparing drugs, potassium supplements or potassium-containing salt substitutes:

The combination of perindopril and potassium sparing drugs, potassium supplements or potassium-containing salt substitutes is generally not recommended (see section *Interaction with other medicinal products and other forms of interaction*).

Dual blockade of the renin-angiotensin-aldosterone system (RAAS):

There is evidence that the concomitant use of ACE-inhibitors, angiotensin II receptor blockers or aliskiren increases the risk of hypotension, hyperkalaemia and decreased renal function (including acute renal failure). Dual blockade of RAAS through the combined use of ACE-inhibitors, angiotensin II receptor blockers or aliskiren is therefore not recommended (see sections *Interaction with other medicinal products and other forms of interaction* and *Pharmacodynamic properties*).

If dual blockade therapy is considered absolutely necessary, this should only occur under specialist supervision and subject to frequent close monitoring of renal function, electrolytes and blood pressure.

ACE-inhibitors and angiotensin II receptor blockers should not be used concomitantly in patients with diabetic nephropathy.

Combination with calcium antagonists, Class I antiarrhythmic drugs and centrally acting antihypertensive drugs:

Combination of bisoprolol with calcium antagonists of the verapamil or diltiazem type, with Class I antiarrhythmic drugs and with centrally acting antihypertensive drugs is generally not recommended see section

Interaction with other medicinal products and other forms of interaction)

Stopping treatment:

Abrupt cessation of therapy with a beta-blocker should be avoided, especially in patients with ischaemic heart disease, because this may lead to transitional worsening of heart condition. The posology should be decreased gradually, using the individual components, ideally over a period of two weeks while at the same time starting the replacement therapy if necessary.

Bradycardia:

If, during treatment, resting heart rate drops below 50-55 beats per minute and the patient experiences symptoms related to bradycardia, Cosyrel dose should be downtitrated using the individual components with an appropriate dose of bisoprolol.

First degree AV block:

Given their negative dromotropic effect, beta-blockers should be administered with caution to patients with first degree AV block.

Aortic and mitral valve stenosis / hypertrophic cardiomyopathy:

As with other ACE inhibitors, perindopril should be given with caution to patients with mitral valve stenosis and obstruction in the outflow of the left ventricle such as aortic stenosis or hypertrophic cardiomyopathy.

Prinzmetal's angina:

Beta-blockers may increase the number and the duration of angina episodes in patients with Prinzmetal's angina. The use of selective blockers of beta-1 adrenergic receptors is possible in mild cases and only in combination with vasodilators.

Renal impairment:

In case of renal impairment, the daily dose of Cosyrel should be adjusted according to creatinine clearance (see section *Posology and method of administration*). Routine monitoring of potassium and creatinine are part of normal

medical practice for these patients (see section *Undesirable effects*).

In patients with symptomatic heart failure, hypotension following the initiation of therapy with ACE inhibitors may lead to some further impairment in renal function. Acute renal failure, usually reversible, has been reported in this situation.

In some patients with bilateral renal artery stenosis or stenosis of the artery to a solitary kidney, who have been treated with ACE inhibitors, increases in blood urea and serum creatinine, usually reversible upon discontinuation of therapy, have been seen. This is especially likely in patients with renal insufficiency. If renovascular hypertension is also present there is an increased risk of severe hypotension and renal insufficiency. In these patients, treatment should be started under close medical supervision with low doses and careful dose titration. Since treatment with diuretics may be a contributory factor to the above, they should be discontinued and renal function should be monitored during the first weeks of treatment therapy.

Some hypertensive patients with no apparent pre-existing renal vascular disease have developed increases in blood urea and serum creatinine, usually minor and transient, especially when perindopril has been given concomitantly with a diuretic. This is more likely to occur in patients with pre-existing renal impairment. Dosage reduction and/or discontinuation of the diuretic and/or perindopril may be required.

Renovascular hypertension:

There is an increased risk of hypotension and renal insufficiency when patient with bilateral renal artery stenosis or stenosis of the artery to a single functioning kidney are treated with ACE inhibitors (see *Contraindications*). Treatment with diuretics may be a contributory factor. Loss of renal function may occur with only minor changes in serum creatinine even in patients with unilateral renal artery stenosis.

Kidney transplantation:

There is no experience regarding the administration of perindopril arginine in patients with recent kidney transplantation.

Haemodialysis patients:

Anaphylactoid reactions have been reported in patients dialysed with high flux membranes, and treated concomitantly with an ACE inhibitor. In these patients consideration should be given to using a different type of dialysis membrane or different class of antihypertensive agent.

Anaphylactoid reactions during low-density lipoproteins (LDL) apheresis:

Rarely, patients receiving ACE inhibitors during LDL apheresis with dextran sulphate have experienced life-threatening anaphylactoid reactions. These reactions were avoided by temporarily withholding ACE inhibitor therapy prior to each apheresis.

Anaphylactoid reactions during desensitisation:

Patients receiving ACE inhibitors during desensitisation treatment (e.g. hymenoptera venom) have experienced anaphylactoid reactions. In the same patients, these reactions have been avoided when the ACE inhibitors were temporarily withheld, but they reappeared upon inadvertent rechallenge. As with other beta-blockers, bisoprolol may increase both the sensitivity towards allergens and the severity of anaphylactoid reactions. Epinephrine treatment does not always yield the expected therapeutic effect.

Neutropenia/Agranulocytosis/Thrombocytopenia/Anaemia:

Neutropenia/agranulocytosis, thrombocytopenia and anaemia have been reported in patients receiving ACE inhibitors. In patients with normal renal function and no other complicating factors, neutropenia occurs rarely. Perindopril should be used with extreme caution in patients with collagen vascular disease, immunosuppressant therapy, treatment with allopurinol or procainamide, or a combination of these complicating factors,

especially if there is pre-existing impaired renal function. Some of these patients developed serious infections, which in a few instances did not respond to intensive antibiotic therapy. If perindopril is used in such patients, periodic monitoring of white blood cell counts is advised and patients should be instructed to report any sign of infection (e.g. sore throat, fever).

Bronchospasm (Bronchial asthma, obstructive airways diseases):

In bronchial asthma or other chronic obstructive lung diseases, which may cause symptoms, bronchodilating therapy should be given concomitantly. Occasionally an increase of the airway resistance may occur when beta-blockers are used in patients with asthma, therefore the dose of beta2-stimulants may have to be increased.

Diabetic patients:

Caution is advised when Cosyrel is used in patients with diabetes mellitus with large fluctuations in blood glucose values. Symptoms of hypoglycaemia can be masked by beta-blockers.

Strict fasting:

Caution is advised in patients with strict fasting.

Peripheral arterial occlusive disease:

Aggravation of symptoms may occur with beta-blockers, especially when starting therapy.

Anaesthesia:

In patients undergoing general anaesthesia beta-blockade reduces the incidence of arrhythmias and myocardial ischemia during induction and intubation, and the post-operative period. It is currently recommended that maintenance beta-blockade be continued peri-operatively. The anaesthetist must be aware of beta-blockade because of the potential for interactions with other drugs, resulting in bradyarrhythmias, attenuation of the reflex tachycardia and the decreased reflex ability to compensate for blood loss. If it is thought necessary to withdraw beta-blocker therapy before surgery, this should be done

gradually and completed about 48 hours before anaesthesia.

In patients undergoing major surgery or during anaesthesia with agents that produce hypotension, perindopril may block angiotensin II formation secondary to compensatory renin release. The treatment should be discontinued one day prior to the surgery. If hypotension occurs and is considered to be due to this mechanism, it can be corrected by volume expansion.

Psoriasis:

Patients with psoriasis or with a history of psoriasis should only be given beta-blockers after carefully balancing the benefits against the risks.

Phaeochromocytoma:

In patients with known or suspected to have phaeochromocytoma bisoprolol should always be given in combination with an alpha-receptor blocker.

Thyreotoxicosis:

Under treatment with bisoprolol the symptoms of a thyreotoxicosis may be masked.

Primary aldosteronism:

Patients with primary hyperaldosteronism generally will not respond to anti-hypertensive drugs acting through inhibition of the renin-angiotensin system. Therefore, the use of this product is not recommended.

Pregnancy:

Unless continued ACE inhibitor therapy is considered essential, patients planning pregnancy should be changed to alternative anti-hypertensive treatments which have an established safety profile for use in pregnancy. When pregnancy is diagnosed, treatment with ACE inhibitors should be stopped immediately, and, if appropriate, alternative therapy should be started (see sections *Contraindications* and *Fertility, pregnancy and lactation*).

Heart failure:

There is no therapeutic experience of bisoprolol treatment of heart failure in patients with the following diseases and conditions:

- insulin dependent diabetes mellitus (type I),
- severely impaired renal function,
- severely impaired hepatic function,
- restrictive cardiomyopathy,
- congenital heart disease,
- haemodynamically significant organic valvular disease,
- myocardial infarction within the last 3 months.

Excipients:

Level of sodium

Cosyrel contains less than 1 mmol sodium (23 mg) per tablet, i.e. essentially 'sodium-free'.

Interaction with other medicinal products and other forms of interaction

No interactions between bisoprolol and perindopril have been observed in an interaction study conducted in healthy volunteers. Only information on interactions with other products that are known for the individual active substances is provided below.

Drugs increasing the risk of angioedema:

Concomitant use of ACE inhibitors with sacubitril/valsartan is contraindicated as this increases the risk of angioedema (see *Contraindications* and *Special warnings and precautions for use*). Sacubitril/valsartan must not be started until 36 hours after taking the last dose of perindopril therapy. Perindopril therapy must not be started until 36 hours after the last dose of sacubitril/valsartan (see *Contraindications* and *Special warnings and precautions for use*).

Concomitant use of ACE inhibitors with racecadotril, mTOR inhibitors (e.g. sirolimus, everolimus, temsirolimus) and gliptins (e.g. linagliptin, saxagliptin, sitagliptin, vildagliptin) may lead to an increased risk for angioedema (see *Special warnings and precautions for use*).

Drugs inducing hyperkalaemia:

Although serum potassium usually remains within normal limits, hyperkalaemia may occur in some patients treated with Cosyrel. Some drugs or therapeutic classes may increase the occurrence of hyperkalaemia: aliskiren, potassium salts, potassium-sparing diuretics (e.g. spironolactone, triamterene or amiloride), ACE inhibitors, angiotensin-II receptors antagonists, NSAIDs, heparins, immunosuppressant agents such as ciclosporin or tacrolimus, trimethoprim and cotrimoxazole (trimethoprim/sulfamethoxazole), as trimethoprim is known to act as a potassium-sparing diuretic like amiloride. The combination of these drugs increases the risk of hyperkalaemia. Therefore, the combination of Cosyrel with the above-mentioned drugs is not recommended. If concomitant use is indicated, they should be used with caution and with frequent monitoring of serum potassium.

Concomitant use contraindicated (see section Contraindications)

Aliskiren:

The concomitant therapy with Cosyrel and aliskiren is contra-indicated in diabetic or impaired renal patients, due to the risk of hyperkalaemia, worsening of renal function and cardiovascular morbidity and mortality increase.

Extracorporeal treatments:

Extracorporeal treatments leading to contact of blood with negatively charged surfaces such as dialysis or haemofiltration with certain high-flux membranes (e.g. polyacrylonitril membranes) and low density lipoprotein apheresis with dextran sulphate due to increased risk of severe anaphylactoid reactions (see section *Contraindications*). If such treatment is required, consideration should be given to using a different type of dialysis membrane or a different class of antihypertensive agent.

Concomitant use not recommended

Linked to bisoprolol

Centrally acting antihypertensives such as clonidine and others (e.g. methyldopa, moxonidine, rilmenidine):

Concomitant use of centrally acting antihypertensives may worsen heart failure by lowering the central sympathetic tonus (reduced heart rate and cardiac output, vasodilation). Abrupt termination, particularly before down-titration of beta-blocker therapy, may increase the risk of rebound hypertension.

Class I antiarrhythmic drugs (e.g. quinidine, disopyramide; lidocaine, phenytoin; flecainide, propafenone):

Effect on atrio-ventricular conduction time may be potentiated and negative inotropic effect increased.

Calcium antagonists of the verapamil type and to a lesser extent of the diltiazem type:

Negative influence on contractility and atrio-ventricular conduction. Intravenous administration of verapamil in patients on beta-blocker treatment may lead to profound hypotension and atrio-ventricular block.

Linked to perindopril

Aliskiren:

In patients other than diabetic or impaired renal patients, risk of hyperkalaemia, worsening of renal function and cardiovascular morbidity and mortality increase.

Concomitant therapy with ACE inhibitor and angiotensin-receptor blocker:

Clinical trial data has shown that dual blockade of the renin-angiotensin-aldosterone-system (RAAS) through the combined use of ACE-inhibitors, angiotensin II receptor blockers or aliskiren is associated with a higher frequency of adverse events such as hypotension, hyperkalaemia and decreased renal function (including acute renal failure) compared to the use of a single RAAS-acting agent (see sections *Contraindications*, *Special warnings and precautions for use* and *Pharmacodynamic properties*).

It has been reported in the literature that in patients with established atherosclerotic

disease, heart failure, or with diabetes with end organ damage, concomitant therapy with ACE inhibitor and angiotensin-receptor blocker is associated with a higher frequency of hypotension, syncope, hyperkalaemia, and worsening renal function (including acute renal failure) as compared to use of a single renin-angiotensin-aldosterone system agent. Dual blockade (e.g. by combining an ACE inhibitor with an angiotensin II receptor antagonist) should be limited to individually defined cases with close monitoring of renal function, potassium levels, and blood pressure.

Estramustine:

Risk of increased adverse effects such as angioneurotic oedema (angioedema).

Potassium sparing diuretics (e.g. triamterene, amiloride...), potassium (salts):

Hyperkalaemia (potentially lethal), especially in conjunction with renal impairment (additive hyperkalaemic effects).

The combination of perindopril with the above-mentioned drugs is not recommended (see section *Special warnings and precautions for use*). If concomitant use is nonetheless indicated they should be used with caution and with frequent monitoring of serum potassium. For use of spironolactone in heart failure, see below.

Lithium:

Reversible increases in serum lithium concentrations and toxicity have been reported during concomitant administration of lithium with ACE inhibitors. Use of perindopril with lithium is not recommended, but if the combination proves necessary, careful monitoring of serum lithium levels should be performed (see section *Special warnings and precautions for use*).

Concomitant use which require special care

Linked to bisoprolol and perindopril

Antidiabetic agents (insulins, oral hypoglycaemic agents):

Epidemiological studies have suggested that concomitant administration of ACE inhibitors and antidiabetic medicines (insulins, oral hypoglycaemic agents) may cause an increased blood-glucose lowering effect with risk of hypoglycaemia. This phenomenon appeared to be more likely to occur during the first weeks of combined treatment and in patients with renal impairment.

Concomitant administration of bisoprolol with insulin and oral antidiabetic drugs may increase blood sugar lowering effect. Blockade of beta-adrenoreceptors may mask symptoms of hypoglycaemia.

Non-steroidal anti-inflammatory medicinal products (NSAIDs) (including acetylsalicylic acid ≥ 3 g/day):

The administration of Cosyrel simultaneously with non-steroidal anti-inflammatory drugs (i.e. acetylsalicylic acid at anti-inflammatory dosage regimens, COX-2 inhibitors and non-selective NSAIDs) may attenuate the antihypertensive effect of bisoprolol and perindopril.

In addition, concomitant use of ACE inhibitors and NSAIDs may lead to an increased risk of worsening of renal function, including possible acute renal failure, and an increase in serum potassium, especially in patients with poor pre-existing renal function. The combination should be administered with caution, especially in the elderly. Patients should be adequately hydrated and consideration should be given to monitoring renal function after initiation of concomitant therapy, and periodically thereafter.

Antihypertensive agents and vasodilators:

Concomitant use with antihypertensive agents, vasodilators (such as nitroglycerin, other nitrates or other vasodilators) or with other medications which have a blood-pressure-reducing potential (e.g. tricyclic antidepressants, barbiturates, phenothiazines) may increase the risk of hypotensive effects of perindopril and bisoprolol.

Tricyclic

antidepressants/Antipsychotics/Anesthetics:

Concomitant use of ACE inhibitors with certain anaesthetic medicinal products, tricyclic antidepressants and antipsychotics may result in further reduction of blood pressure.

Concomitant use of bisoprolol with anaesthetics may lead to reduced reflex tachycardia and increased risk of hypotension.

Sympathomimetics:

Beta-sympathomimetics (e.g. isoprenaline, dobutamine): combination with bisoprolol may reduce the effects of both agents.

Sympathomimetics that activate both beta- and alpha-adrenoceptors (e.g. norepinephrine, epinephrine): combination with bisoprolol may unmask the alpha-adrenoceptor-mediated vasoconstrictor effects of these agents, leading to blood pressure increase and exacerbated intermittent claudication. Such interactions are considered to be more likely with nonselective beta-blockers.

Sympathomimetics may reduce the antihypertensive effects of ACE inhibitors.

Linked to bisoprolol

Calcium antagonists of the dihydropyridine type such as felodipine and amlodipine:

Concomitant use may increase the risk of hypotension, and an increase in the risk of a further deterioration of the ventricular pump function in patients with heart failure cannot be excluded.

Class-III antiarrhythmic drugs (e.g. amiodarone):
Effect on atrio-ventricular conduction time may be potentiated.

Parasympathomimetic drugs:

Concomitant use may increase atrio-ventricular conduction time and the risk of bradycardia.

Topical beta-blockers (e.g. eye drops for glaucoma treatment):

Concomitant use may add to the systemic effects of bisoprolol.

Digitalis glycosides:

Reduction of heart rate, increase of atrio-ventricular conduction time.

Linked to perindopril

Baclofen:

Increased antihypertensive effect. Monitor blood pressure and adapt antihypertensive dosage if necessary.

Non-potassium-sparing diuretics:

Patients on diuretics, and especially those who are volume and/or salt depleted, may experience excessive reduction in blood pressure after initiation of therapy with an ACE inhibitor. The possibility of hypotensive effects can be reduced by discontinuation of the diuretic, by increasing volume or salt intake prior to initiating therapy with low and progressive doses of perindopril.

In arterial hypertension, when prior diuretic therapy can have caused salt/volume depletion, either the diuretic must be discontinued before initiating the ACE inhibitor, in which case a non-potassium-sparing diuretic can be thereafter reintroduced or the ACE inhibitor must be initiated with a low dosage and progressively increased.

In diuretic-treated congestive heart failure, the ACE inhibitor should be initiated at a very low dosage, possibly after reducing the dosage of the associated non-potassium-sparing diuretic. In all cases, renal function (creatinine levels) must be monitored during the first few weeks of ACE inhibitor therapy.

Potassium-sparing diuretics (eplerenone, spironolactone):

With eplerenone or spironolactone at doses between 12,5 mg to 50 mg by day and with low doses of ACE inhibitors:

In the treatment of class II-IV heart failure (NYHA) with an ejection fraction < 40%, and previously treated with ACE inhibitors and loop diuretics, risk of hyperkalaemia, potentially lethal, especially in case of non-observance of the prescription recommendations on this combination.

Before initiating the combination, check the absence of hyperkalaemia and renal impairment.

A close monitoring of the kalaemia and creatininemia is recommended in the first month of the treatment once a week at the beginning and, monthly thereafter.

Combination use to be taken into consideration

Linked to bisoprolol

Mefloquine:

Increased risk of bradycardia.

Monoamine oxidase inhibitors (except MAO-B inhibitors):

Enhanced hypotensive effect of the beta-blockers but also risk for hypertensive crisis.

Linked to perindopril

Gold:

Nitritoid reactions (symptoms include facial flushing, nausea, vomiting and hypotension) have been reported rarely in patients on therapy with injectable gold (sodium aurothiomalate) and concomitant ACE inhibitor therapy including perindopril.

Fertility, pregnancy and lactation

Pregnancy:

Based on existing data on monocomponents, Cosyrel is not recommended during the first trimester of pregnancy and is contraindicated during the second and third trimesters of pregnancy.

Bisoprolol

Bisoprolol has pharmacological effects that may cause harmful effects on pregnancy and/or the foetus/newborn (reduce placental perfusion associated with growth retardation, intrauterine death, abortion or early labour and adverse effects (e.g. hypoglycaemia and bradycardia) may occur in the foetus and newborn infant). If treatment with beta-adrenoceptor blockers is necessary, beta-1-selective adrenoceptor blockers are preferable.

Bisoprolol should not be used during pregnancy unless clearly necessary. If treatment with bisoprolol is considered necessary, the uteroplacental blood flow and the foetal growth should be monitored. In case of harmful effects on pregnancy or the foetus alternative treatment should be considered. The newborn infant must be closely monitored. Symptoms of hypoglycaemia and bradycardia are generally to be expected within the first 3 days.

Perindopril

Epidemiological evidence regarding the risk of teratogenicity following exposure to ACE inhibitors during the first trimester of pregnancy has not been conclusive; however a small increase in risk cannot be excluded. Unless continued ACE inhibitor therapy is considered essential, patients planning pregnancy should be changed to alternative anti-hypertensive treatments which have an established safety profile for use in pregnancy. When pregnancy is diagnosed, treatment with ACE inhibitors should be stopped immediately, and, if appropriate, alternative therapy should be started.

Exposure to ACE inhibitor therapy during the second and third trimesters is known to induce human foetotoxicity (decreased renal function, oligohydramnios, skull ossification retardation) and neonatal toxicity (renal failure, hypotension, hyperkalaemia) (see section *Preclinical safety data*). Should exposure to ACE inhibitor have occurred from the second trimester of pregnancy, ultrasound check of renal function and skull is recommended. Infants whose mothers have taken ACE inhibitors should be closely observed for hypotension (see also sections *Contraindications* and *Special warnings and precautions for use*).

Breast-feeding:

Cosyrel is not recommended during lactation. It is not known whether bisoprolol is excreted in human milk. Therefore, breastfeeding is not

recommended during administration of bisoprolol.

Because no information is available regarding the use of perindopril during breastfeeding, perindopril is not recommended and alternative treatments with better established safety profiles during breast-feeding are preferable, especially while nursing a newborn or preterm infant.

Fertility:

There are no clinical data on fertility with the use of Cosyrel.

Effects on ability to drive and use machines

Cosyrel has no direct influence on the ability to drive and use machines but individual reactions related to low blood pressure may occur in some patients, particularly at the start of treatment or upon change of medication as well as in conjunction with alcohol.

As a result the ability to drive or operate machinery may be impaired.

Undesirable effects

Summary of the safety profile:

The most common adverse reactions to bisoprolol include headache, dizziness, worsening of heart failure, hypotension, cold extremities, nausea, vomiting, abdominal pain, diarrhoea, constipation, asthenia and fatigue.

The most common adverse reactions reported in clinical trials and observed with perindopril include headache, dizziness, vertigo, paraesthesia, visual disturbance, tinnitus, hypotension, cough, dyspnoea, nausea, vomiting, abdominal pain, diarrhoea, constipation, dysgeusia, dyspepsia, rash, pruritus, muscle cramps and asthenia.

Tabulated list of adverse reactions:

The following undesirable effects have been observed during clinical trials and/or post-marketing use with bisoprolol or perindopril given separately and ranked under the MedDRA classification by body system and under the following frequency:

Very common ($\geq 1/10$) ; common ($\geq 1/100$ to $< 1/10$) ; uncommon ($\geq 1/1000$ to $< 1/100$) ; rare ($\geq 1/10000$ to $< 1/1000$) ; very rare ($< 1/10000$) ; not known (cannot be estimated from the available data).

MedDRA System Organ Class	Undesirable Effects	Frequency	
		Bisoprolol	Perindopril
Infections and infestations	Rhinitis	Rare	Very rare
Blood and lymphatic System Disorders	Eosinophilia	-	Uncommon*
	Agranulocytosis (see section <i>Special warnings and precautions for use</i>)	-	Very rare
	Pancytopenia	-	Very rare
	Leukopenia	-	Very rare
	Neutropenia (see section <i>Special warnings and precautions for use</i>)	-	Very rare
	Thrombocytopenia (see section <i>Special warnings and precautions for use</i>)	-	Very rare
Endocrine disorders	Haemolytic anaemia in patients with a congenital deficiency of G-6PDH	-	Very rare
	Syndrome of inappropriate antidiuretic hormone secretion (SIADH)	-	Rare
Metabolism and nutrition disorders	Hypoglycaemia (see sections <i>Special warnings and precautions for use</i> and <i>Interaction with other medicinal products and other forms of interaction</i>)	-	Uncommon*
	Hyperkalaemia, reversible on discontinuation	-	Uncommon*
	Hyponatraemia	-	Uncommon*
Psychiatric disorders	Mood altered	-	Uncommon
	Sleep disorder	Uncommon	Uncommon

	Depression	Uncommon	Uncommon*
	Nightmares, Hallucinations	Rare	-
	Confusion	-	Very rare
Nervous system disorders	Headache**	Common	Common
	Dizziness**	Common	Common
	Vertigo	-	Common
	Dysgeusia	-	Common
	Paraesthesia	-	Common
	Somnolence	-	Uncommon*
	Syncope	Rare	Uncommon*
Eye disorders	Visual impairment	-	Common
	Reduced tear flow (to be considered if the patient uses lenses)	Rare	-
	Conjunctivitis	Very rare	-
Ear and labyrinth disorders	Tinnitus	-	Common
	Hearing disorders	Rare	-
Cardiac disorders	Palpitations	-	Uncommon*
	Tachycardia	-	Uncommon*
	Bradycardia	Very common	-
	Worsening of heart failure	Common	-
	AV-conduction disturbances	Uncommon	-
	Arrhythmia	-	Very rare
	Angina pectoris	-	Very rare
	Myocardial infarction possibly secondary to excessive hypotension in high-risk patients (see section <i>Special warnings and precautions for use</i>)	-	Very rare
Vascular disorders	Hypotension and effects related to hypotension	Common	Common
	Feeling of coldness or numbness in the extremities	Common	-
	Orthostatic hypotension	Uncommon	-
	Vasculitis	-	Uncommon*
	Flushing	-	Rare*

	Stroke possibly secondary to excessive hypotension in high-risk patients (see section <i>Special warnings and precautions for use</i>)	-	Very rare
	Raynaud's phenomenon	-	Not known
Respiratory, thoracic and mediastinal disorders	Cough	-	Common
	Dyspnoea	-	Common
	Bronchospasm	Uncommon	Uncommon
	Eosinophilic pneumonia	-	Very rare
Gastro-intestinal disorders	Abdominal pain	Common	Common
	Constipation	Common	Common
	Diarrhoea	Common	Common
	Nausea	Common	Common
	Vomiting	Common	Common
	Dyspepsia	-	Common
	Dry mouth	-	Uncommon
	Pancreatitis	-	Very rare
Hepato-biliary disorders	Hepatitis either cytolytic or cholestatic (see section <i>Special warnings and precautions for use</i>)	Rare	Very rare
Skin and subcutaneous tissue disorders	Rash	-	Common
	Pruritus	-	Common
	Angioedema of face, extremities, lips, mucous membranes, tongue, glottis and/or larynx (see section <i>Special warnings and precautions for use</i>)	-	Uncommon
	Urticaria	-	Uncommon
	Photosensitivity reactions	-	Uncommon*
	Pemphigoid	-	Uncommon*
	Hyperhidrosis	-	Uncommon
	Hypersensitivity reactions (itching, flush, rash)	Rare	-
	Psoriasis aggravation	-	Rare*
	Erythema multiform	-	Very rare
	Alopecia	Very rare	-
	Beta-blockers may provoke or worsen psoriasis or induce psoriasis-like rash	Very rare	-
Musculoskeletal and connective tissue disorders	Muscle cramps	Uncommon	Common
	Muscular weakness	Uncommon	-
	Arthralgia	-	Uncommon*
	Myalgia	-	Uncommon*
Renal and urinary disorders	Renal insufficiency	-	Uncommon
	Acute renal failure	-	Rare
	Anuria/Oliguria	-	Rare*
	Erectile dysfunction	-	Uncommon
Reproductive system and breast disorders	Potency disorders	Rare	-
	Asthenia	Common	Common
General disorders and administration site conditions	Fatigue	Common	-
	Chest pain	-	Uncommon*
	Malaise	-	Uncommon*
	Oedema peripheral	-	Uncommon*
	Pyrexia	-	Uncommon*
	Blood urea increased	-	Uncommon*
Investigations	Blood creatinine increased	-	Uncommon*
	Hepatic enzyme increased	Rare	Rare
	Blood bilirubin increased	-	Rare
	Increased triglycerides	Rare	-
	Haemoglobin decreased and haematocrit decreased (see section <i>Special warnings and precautions for use</i>)	-	Very rare
	Fall	-	Uncommon*
	Injury, poisoning and procedural complications	-	Uncommon*

** Frequency calculated from clinical trials for adverse events detected from spontaneous report*

***These symptoms especially occur at the beginning of the therapy. They are generally mild and often disappear within 1-2 weeks.*

Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via e-meso sub-site at <https://e-meso.pom.go.id>, or reporting to Pusat Farmakovigilans/ MESO Nasional, Badan Pengawas Obat dan Makanan: Phone: (+62 21) 4244691 Ext.1079 and Email: pv-center@pom.go.id.

Overdose

There is no information on overdose with Cosyrel in humans.

Bisoprolol

Symptoms:

In general the most common signs expected with overdosage of a beta-blocker are bradycardia, hypotension, bronchospasm, acute cardiac insufficiency and hypoglycaemia. To date a few cases of overdose (maximum: 2000 mg) with bisoprolol have been reported in patients suffering from hypertension and/or coronary heart disease showing bradycardia and/or hypotension; all patients recovered. There is a wide interindividual variation in sensitivity to one single high dose of bisoprolol and patients with heart failure are probably very sensitive.

Management:

If overdose occurs, bisoprolol treatment should be stopped and supportive and symptomatic treatment should be provided. Limited data suggest that bisoprolol is hardly dialysable. Based on the expected pharmacologic actions and recommendations for other beta-blockers, the following general measures should be considered when clinically warranted.

Bradycardia: Administer intravenous atropine. If the response is inadequate, isoprenaline or another agent with positive chronotropic properties may be given cautiously. Under some circumstances, transvenous pacemaker insertion may be necessary.

Hypotension: Intravenous fluids and vasopressors should be administered. Intravenous glucagon may be useful.

AV block (second or third degree): Patients should be carefully monitored and treated with isoprenaline infusion or transvenous cardiac pacemaker insertion.

Acute worsening of heart failure: Administer i.v. diuretics, inotropic agents, vasodilating agents.

Bronchospasm: Administer bronchodilator therapy such as isoprenaline, beta2-sympathomimetic drugs and/or aminophylline.

Hypoglycaemia: Administer i.v. glucose.

Perindopril

Symptoms:

Limited data are available for overdosage in humans. Symptoms associated with overdosage of ACE inhibitors may include hypotension, circulatory shock, electrolyte disturbances, renal failure, hyperventilation, tachycardia, palpitations, bradycardia, dizziness, anxiety, and cough.

Management:

The recommended treatment of overdosage is intravenous infusion of sodium chloride 9 mg/ml (0.9%) solution. If hypotension occurs, the patient should be placed in the shock position. If available, treatment with angiotensin II infusion and/or intravenous catecholamines may also be considered. Perindopril may be removed from the general circulation by haemodialysis (see section *Special warnings and precautions for use*). Pacemaker therapy is indicated for therapy-resistant bradycardia. Vital signs, serum electrolytes and creatinine concentrations should be monitored continuously.

PHARMACOLOGICAL PROPERTIES

Pharmacodynamic properties

Pharmacotherapeutic group: ACE inhibitors, other combinations.

ATC code: C09BX02

Mechanism of action

Bisoprolol

Bisoprolol is a highly beta1-selective-adrenoceptor blocking agent, lacking intrinsic stimulating and relevant membrane stabilising activity. It only shows low affinity to the beta2-receptor of the smooth muscles of bronchi and vessels as well as to the beta2-receptors concerned with metabolic regulation. Therefore, bisoprolol is generally not to be expected to influence the airway resistance and beta2-mediated metabolic effects. Its beta1-selectivity extends beyond the therapeutic dose range.

Perindopril

Perindopril is an inhibitor of the enzyme that converts angiotensin I into angiotensin II (ACE). The converting enzyme, or kininase, is an exopeptidase that allows conversion of angiotensin I into the vasoconstrictor angiotensin II as well as causing the degradation of the vasodilator bradykinin into an inactive heptapeptide. Inhibition of ACE results in a reduction of angiotensin II in the plasma, which leads to increased plasma renin activity (by inhibition of the negative feedback of renin release) and reduced secretion of aldosterone. Since ACE inactivates bradykinin, inhibition of ACE also results in an increased activity of circulating and local kallikrein-kinin systems (and thus also activation of the prostaglandin system). It is possible that this mechanism contributes to the blood pressure-lowering action of ACE inhibitors and is partially responsible for certain of their side effects (e.g. cough).

Perindopril acts through its active metabolite, perindoprilat. The other metabolites show no inhibition of ACE activity in vitro.

Pharmacodynamic effects

Bisoprolol

Bisoprolol has no significant negative inotropic effects.

Bisoprolol reaches its maximum effects 3-4 hours after administration. Due to the half-life of 10-12 hours, bisoprolol acts for 24 hours.

The maximum blood-pressure-lowering effects of bisoprolol are generally reached after 2 weeks.

In acute administration in patients with coronary heart disease without chronic heart failure bisoprolol reduces the heart rate and stroke volume and thus the cardiac output and oxygen consumption. In chronic administration the initially elevated peripheral resistance decreases. The decrease in plasma renin activity is proposed as a mechanism of action underlying the antihypertensive effect of beta-blockers.

Bisoprolol reduces the sympatho-adrenergic response by blocking cardiac beta-adrenergic receptors. This results in a decrease in heart rate and contractility, causing a reduction in oxygen consumption by the myocardium, which is the desired effect in the case of angina associated with underlying coronary heart disease.

Perindopril

Hypertension:

Perindopril is active in all grades of hypertension: mild, moderate, severe; a reduction in systolic and diastolic blood pressures in both supine and standing positions is observed.

Perindopril reduces peripheral vascular resistance, leading to blood pressure reduction. As a consequence, peripheral blood flow increases, with no effect on heart rate.

Renal blood flow increases as a rule, while the glomerular filtration rate (GFR) is usually unchanged.

For Cosyrel 5 mg/5 mg and Coyrel 10 mg/5 mg

Heart failure:

Perindopril reduces cardiac work by a decrease in pre-load and after-load.

Clinical efficacy and safety
Bisoprolol

In total 2647 patients were included in the CIBIS II trial. 83% (n = 2202) were in NYHA class III and 17% (n = 445) were in NYHA class IV. They had stable symptomatic systolic heart failure (ejection fraction <35%, based on echocardiography). Total mortality was reduced from 17.3% to 11.8% (relative reduction 34%). A decrease in sudden death (3.6% vs 6.3%, relative reduction 44%) and a reduced number of heart failure episodes requiring hospital admission (12% vs 17.6%, relative reduction 36%) was observed. Finally, a significant improvement of the functional status according to NYHA classification has been shown. During the initiation and titration of bisoprolol hospital admission due to bradycardia (0.53%), hypotension (0.23%), and acute decompensation (4.97%) were observed, but they were not more frequent than in the placebo-group (0%, 0.3% and 6.74%). The numbers of fatal and disabling strokes during the total study period were 20 in the bisoprolol group and 15 in the placebo group.

The CIBIS III trial investigated 1010 patients aged ≥65 years with mild to moderate chronic heart failure (CHF; NYHA class II or III) and left ventricular ejection fraction ≤35%, who had not been treated previously with ACE inhibitors, beta-blockers, or angiotensin receptor blockers. Patients were treated with a combination of bisoprolol and enalapril for 6 to 24 months after an initial 6 months treatment with either bisoprolol or enalapril.

There was a trend toward higher frequency of chronic heart failure worsening when bisoprolol was used as the initial 6 months treatment. Non inferiority of bisoprolol-first versus enalapril-first treatment was not proven in the per-protocol analysis, although the two strategies for initiation of CHF treatment showed a similar rate of the primary combined endpoint death and hospitalization at study end (32.4% in the bisoprolol-first group vs. 33.1 % in the enalapril-first group, per-protocol population). The study shows that bisoprolol can also be used in

elderly chronic heart failure patients with mild to moderate disease.

Perindopril

Hypertension:

Perindopril is active in all grades of hypertension: mild, moderate, severe; a reduction in systolic and diastolic blood pressures in both supine and standing positions is observed.

Perindopril reduces peripheral vascular resistance, leading to blood pressure reduction. As a consequence, peripheral blood flow increases, with no effect on heart rate.

Renal blood flow increases as a rule, while the GFR is usually unchanged.

The antihypertensive activity is maximal between 4 and 6 hours after a single dose and is sustained for at least 24 hours: trough effects are about 87- 100 % of peak effects.

The decrease in blood pressure occurs rapidly. In responding patients, normalisation is achieved within a month and persists without the occurrence of tachyphylaxis.

Discontinuation of treatment does not lead to a rebound effect.

Perindopril reduces left ventricular hypertrophy.

In man, perindopril has been confirmed to demonstrate vasodilatory properties. It improves large artery elasticity and decreases the media: lumen ratio of small arteries.

An adjunctive therapy with a thiazide diuretic produces an additive-type of synergy. The combination of an ACE inhibitor and a thiazide also decreases the risk of hypokalaemia induced by the diuretic treatment.

For Cosyrel 5 mg/5 mg and Coyrel 10 mg/5 mg

Heart failure:

Studies in patients with heart failure have demonstrated:

- decreased left and right ventricular filling pressures,
- reduced total peripheral vascular resistance,
- increased cardiac output and improved cardiac index.

In comparative studies, the first administration of 2.5 mg of perindopril arginine to patients with mild to moderate heart failure was not associated with any significant reduction of blood pressure as compared to placebo.

Patients with stable coronary artery disease:

The EUROPA study was a multicentre, international, randomised, double-blind, placebo-controlled clinical trial lasting 4 years.

Twelve thousand two hundred and eighteen (12 218) patients aged over 18 were randomised to 8 mg perindopril tert-butylamine (equivalent to 10 mg perindopril arginine) (n = 6 110) or placebo (n = 6 108).

The trial population had evidence of coronary artery disease with no evidence of clinical signs of heart failure. Overall, 90% of the patients had a previous myocardial infarction and/or a previous coronary revascularisation. Most of the patients received the study medication on top of conventional therapy including platelet inhibitors, lipid lowering agents and beta-blockers.

The main efficacy criterion was the composite of cardiovascular mortality, non-fatal myocardial infarction and/or cardiac arrest with successful resuscitation. The treatment with 8 mg perindopril tert-butylamine (equivalent to 10 mg perindopril arginine) once daily resulted in a significant absolute reduction in the primary endpoint of 1.9% (relative risk reduction of 20%, 95%CI [9.4; 28.6] – p<0.001).

In patients with a history of myocardial infarction and/or revascularisation, an absolute reduction of 2.2% corresponding to a RRR of 22.4% (95%CI [12.0; 31.6] – p<0.001) in the primary endpoint was observed by comparison to placebo.

In a subgroup of patients treated with beta-blockers from EUROPA study defined in a post-hoc analysis, the addition of perindopril to beta-blockers (n=3789) showed a significant absolute reduction of 2.2% (relative risk reduction of 24%, 95%CI [9.5; 36.4]) compared to beta-blockers without perindopril (n=3745) in the composite of cardiovascular mortality, non-fatal

myocardial infarction and/or cardiac arrest with successful resuscitation.

Dual blockade of the renin-angiotensin-aldosterone system (RAAS) clinical trial data:

Two large randomised, controlled trials (ONTARGET (ONgoing Telmisartan Alone and in combination with Ramipril Global Endpoint Trial) and VA NEPHRON-D (The Veterans Affairs Nephropathy in Diabetes) have examined the use of combination of an ACE-inhibitor with an angiotensin II receptor blocker.

ONTARGET was a study conducted in patients with a history of cardiovascular or cerebrovascular disease, or type 2 diabetes mellitus accompanied by evidence of end-organ damage. VA NEPHRON-D was a study in patients with type 2 diabetes mellitus and diabetic nephropathy.

These studies have shown no significant beneficial effect on renal and/or cardiovascular outcomes and mortality, while an increased risk of hyperkalaemia, acute kidney injury and/or hypotension as compared to monotherapy was observed. Given their similar pharmacodynamic properties, these results are also relevant for other ACE-inhibitors and angiotensin II receptor blockers.

ACE-inhibitors and angiotensin II receptor blockers should therefore not be used concomitantly in patients with diabetic nephropathy.

ALTITUDE (Aliskiren Trial in Type 2 Diabetes Using Cardiovascular and Renal Disease Endpoints) was a study designed to test the benefit of adding aliskiren to a standard therapy of an ACE-inhibitor or an angiotensin II receptor blocker in patients with type 2 diabetes mellitus and chronic kidney disease, cardiovascular disease, or both. The study was terminated early because of an increased risk of adverse outcomes. Cardiovascular death and stroke were both numerically more frequent in the aliskiren group than in the placebo group and adverse events and serious adverse events of interest (hyperkalaemia, hypotension and renal dysfunction) were more frequently reported in the aliskiren group than in the placebo group.

Paediatric population:

No data are available with Cosyrel in children. The European Medicines Agency has granted a product-specific waiver for Cosyrel in all subsets of the paediatric population in the treatment of hypertension, ischaemic coronary artery disease stable and chronic heart failure (see section 4.2 for information on paediatric use).

Pharmacokinetic properties

The rate and extent of absorption of bisoprolol and perindopril from Cosyrel are not significantly different, respectively, from the rate and extent of absorption of bisoprolol and perindopril when taken alone as monotherapy.

Bisoprolol

Absorption

Bisoprolol is almost completely (>90%) absorbed from the gastrointestinal tract and, because of its small hepatic first-pass metabolism (approximately 10%), it has a bioavailability of approximately 90% after oral administration.

Distribution

The distribution volume is 3.5 l/kg. The plasma protein binding of bisoprolol is about 30%.

Biotransformation and elimination

Bisoprolol is excreted from the body by two routes. 50% is metabolised by the liver to inactive metabolites which are then excreted by the kidneys. The remaining 50% is excreted by the kidneys in an unmetabolised form. Total clearance is approximately 15 l/h. The half-life in plasma of 10-12 hours gives a 24 hour effect after dosing once daily.

Special populations

The kinetics of bisoprolol are linear and independent of age.

Since the elimination takes place in the kidneys and the liver to the same extent a dosage adjustment is not required for patients with impaired liver function or renal insufficiency. The pharmacokinetics in patients with chronic

heart failure and with impaired liver or renal function has not been studied. In patients with chronic heart failure (NYHA stage III) the plasma levels of bisoprolol are higher and the half-life is prolonged compared to healthy volunteers. Maximum plasma concentration at steady state is 64 ± 21 ng/ml at a daily dose of 10 mg and the half-life is 17 ± 5 hours.

Perindopril

Absorption

After oral administration, the absorption of perindopril is rapid and the peak concentration is achieved within 1 hour. The plasma half-life of perindopril is equal to 1 hour.

Distribution

The volume of distribution is approximately 0.2 l/kg for unbound perindoprilat. Protein binding of perindoprilat to plasma proteins is 20%, principally to ACE, but is concentration-dependent.

Biotransformation

Perindopril is a prodrug. Twenty seven percent of the administered perindopril dose reaches the bloodstream as the active metabolite perindoprilat. In addition to active perindoprilat, perindopril yields five metabolites, all inactive. The peak plasma concentration of perindoprilat is achieved within 3 to 4 hours.

As ingestion of food decreases conversion to perindoprilat, hence bioavailability, perindopril arginine should be administered orally in a single daily dose in the morning before a meal.

Elimination

Perindoprilat is eliminated in the urine and the terminal half-life of the unbound fraction is approximately 17 hours, resulting in steady-state within 4 days.

Linearity

It has been demonstrated a linear relationship between the dose of perindopril and its plasma exposure.

Special population:

Elimination of perindoprilat is decreased in the elderly, and also in patients with heart or renal failure. Dosage adjustment in renal insufficiency is desirable depending on the degree of impairment (creatinine clearance).

Dialysis clearance of perindoprilat is equal to 70 ml/min.

Perindopril kinetics are modified in patients with cirrhosis: hepatic clearance of the parent molecule is reduced by half. However, the quantity of perindoprilat formed is not reduced and therefore no dosage adjustment is required (see sections *Posology and method of administration* and *Special warnings and precautions for use*).

Preclinical safety data

Bisoprolol

Preclinical data reveal no special hazard for humans based on conventional studies of safety pharmacology, repeated dose toxicity, genotoxicity or carcinogenicity.

In reproductive toxicology studies, bisoprolol had no effect on fertility or other general results of reproduction.

Like other beta-blockers, bisoprolol caused maternal (decreased food intake and decreased body weight) and embryo/fetal toxicity (increased incidence of resorptions, reduced birth weight of the offspring, retarded physical development) at high doses but was not teratogenic.

Perindopril

In the chronic oral toxicity studies (rats and monkeys), the target organ is the kidney, with reversible damage.

No mutagenicity has been observed in in vitro or in vivo studies.

Reproduction toxicology studies (rats, mice, rabbits and monkeys) showed no sign of embryotoxicity or teratogenicity. However, ACE inhibitors, as a class, have been shown to induce adverse effects on late foetal development, resulting in foetal death and

congenital effects in rodents and rabbits: renal lesions and an increase in peri- and postnatal mortality have been observed. Fertility was not impaired either in male or in female rats

No carcinogenicity has been observed in long term studies in rats and mice.

Environmental Risk Assessment:

Cosyrel contains known active substances, bisoprolol and perindopril. Cosyrel will be prescribed as a direct replacement for individual doses of bisoprolol and perindopril, so there will be no increase in the environmental exposure.

STORAGE CONDITIONS

Store below 30°C

Shelf life: 2.5 years

The in-use stability after first opening is 30 days.

PACK SIZE

Cosyrel 5 mg/5 mg

Reg.No.: DK12068601817A1

Box of 1 container of 30 tablets.

Cosyrel 5 mg/10 mg

Reg.No.: DK12068601817B1

Box of 1 container of 30 tablets.

Cosyrel 10 mg/5 mg

Reg.No.: DK12068601817C1

Box of 1 container of 30 tablets.

Cosyrel 10 mg/10 mg

Reg.No.: DK12068601817D1

Box of 1 container of 30 tablets.

HARUS DENGAN RESEP DOKTER

Les Laboratoires Servier - France

Manufactured by:



Les Laboratoires Servier Industrie
Gidy – France

Registered by:

PT. Genero Pharmaceuticals
Bekasi - Indonesia

Imported and Marketed by:

PT. Servier Indonesia
Jakarta – Indonesia

DDMMYY

Informasi untuk pasien

Cosyrel 5 mg/5 mg, tablet salut selaput

Cosyrel 5 mg/10 mg, tablet salut selaput

Cosyrel 10 mg/5 mg, tablet salut selaput

Cosyrel 10 mg/10 mg, tablet salut selaput

bisoprolol fumarat / perindopril arginin

Bacalah seluruh isi dari leaflet ini dengan seksama sebelum Anda mulai menggunakan obat ini karena berisi informasi penting untuk Anda.

- Simpan lembar informasi obat ini. Anda mungkin perlu membacanya kembali.
- Jika Anda memiliki pertanyaan lebih lanjut, tanyakan kepada dokter, apoteker atau perawat Anda.
- Obat ini diresepkan untuk Anda saja. Jangan memberikannya kepada orang lain karena dapat membahayakan mereka, bahkan jika gejala penyakit mereka sama seperti Anda.
- Jika Anda mengalami efek samping obat, konsultasikan dengan dokter atau apoteker atau perawat Anda. Efek samping ini termasuk yang mungkin tidak tercantum dalam leaflet ini. Lihat bagian 4.

Apa yang terdapat dalam leaflet ini

1. Apakah Cosyrel dan digunakan untuk apa
2. Apakah yang perlu Anda ketahui sebelum Anda menggunakan Cosyrel
3. Bagaimana cara menggunakan Cosyrel
4. Kemungkinan efek samping
5. Bagaimana cara menyimpan Cosyrel
6. Isi dari kemasan obat ini dan informasi lainnya

1. Apakah Cosyrel dan digunakan untuk apa

Cosyrel mengandung dua bahan aktif yaitu bisoprolol fumarate dan perindopril arginine dalam satu tablet:

- Bisoprolol fumarate termasuk kelompok obat yang disebut penghambat beta / *beta-blockers*. Beta-blockers memperlambat denyut jantung dan membuat jantung lebih efisien dalam memompa darah ke seluruh tubuh.
- Perindopril arginin merupakan inhibitor *angiotensin converting enzyme* (ACE) yang melebarkan pembuluh darah, yang membuatnya lebih mudah bagi jantung Anda untuk memompa darah melalui pembuluh darah.

Cosyrel 5 mg/5 mg dan Cosyrel 10 mg/5 mg

Cosyrel 5 mg/5 mg dan Cosyrel 10 mg/5 mg digunakan untuk mengobati:

- tekanan darah tinggi (hipertensi)
- gagal jantung kronik yang stabil (suatu kondisi dimana jantung tidak dapat memompa cukup darah untuk memenuhi kebutuhan tubuh sehingga mengakibatkan sesak napas dan pembengkakan)
- untuk mengurangi risiko sakit jantung, seperti serangan jantung pada pasien dengan penyakit arteri koroner yang stabil (suatu kondisi dimana pasokan darah ke jantung berkurang atau terhambat) dan yang telah mengalami serangan jantung dan / atau operasi untuk meningkatkan pasokan darah ke jantung dengan melebarkan pembuluh darah yang mengalirkan darah ke jantung.

Cosyrel 5 mg/10 mg dan Cosyrel 10 mg/10 mg

Cosyrel 5 mg/10 mg dan Cosyrel 10 mg/10 mg digunakan untuk mengobati:

- tekanan darah tinggi (hipertensi)
- untuk mengurangi risiko sakit jantung, seperti serangan jantung pada pasien dengan penyakit arteri koroner yang stabil (suatu kondisi dimana pasokan darah ke jantung berkurang atau terhambat) dan yang telah mengalami serangan jantung dan / atau operasi untuk meningkatkan pasokan darah ke jantung dengan melebarkan pembuluh darah yang mengalirkan darah ke jantung.

Daripada menggunakan bisoprolol fumarate dan perindopril arginine sebagai tablet terpisah, Anda lebih baik menggunakan hanya satu tablet Cosyrel yang mengandung kedua bahan aktif dalam kekuatan yang sama.

2. Apa yang perlu Anda ketahui sebelum Anda menggunakan Cosyrel

Jangan menggunakan Cosyrel jika Anda:

- alergi terhadap bisoprol atau *beta-blocker* lainnya, perindopril atau inhibitor ACE lainnya, atau salah satu bahan lain dari obat ini (tercantum dalam bagian 6),
- memiliki gagal jantung yang tiba-tiba menjadi buruk dan / atau yang memerlukan perawatan di rumah sakit,
- mengalami syok kardiogenik (kondisi jantung serius yang disebabkan oleh tekanan darah sangat rendah),
- memiliki penyakit jantung yang ditandai dengan detak jantung yang lambat atau tidak teratur (derajat dua atau tiga blok atrioventrikular, blok sinoatrial, sindrom *sick sinus*),
- memiliki detak jantung yang lambat,
- memiliki tekanan darah yang sangat rendah,
- memiliki asma berat atau penyakit paru kronik berat,
- memiliki masalah sirkulasi darah yang berat di kaki Anda (seperti sindrom Raynaud), yang dapat menyebabkan jari-jari dan kaki Anda menjadi gatal, atau menjadi pucat atau biru,
- memiliki feokromositoma yang tidak diobati, yang merupakan tumor yang jarang dari kelenjar adrenal (medula),
- memiliki asidosis metabolik, suatu kondisi dimana darah Anda mengandung terlalu banyak asam,
- pernah mengalami gejala seperti mengi, pembengkakan wajah, lidah atau tenggorokan, rasa gatal atau ruam pada kulit dengan pengobatan inhibitor ACE sebelumnya atau jika Anda atau anggota keluarga Anda memiliki berbagai gejala ini dalam keadaan lain (kondisi yang disebut angioedema),
- hamil lebih dari 3 bulan (lebih baik menghindari Cosyrel pada awal kehamilan - lihat bab kehamilan),
- memiliki diabetes atau fungsi ginjal terganggu dan Anda diobati dengan obat penurun tekanan darah yang mengandung aliskiren.
- sedang menjalani terapi dialisis atau jenis filtrasi darah lainnya. Tergantung pada alat yang digunakan. Cosyrel mungkin tidak cocok untuk Anda.
- memiliki masalah pada ginjal, di mana terdapat penurunan aliran darah ke ginjal Anda (stenosis arteri renalis).

- sudah meminum atau sedang menjalani terapi dengan sacubitril/valsartan, obat yang digunakan untuk gagal jantung, karena meningkatkan resiko angioedema (pembengkakan yang cepat di bawah kulit di area seperti tenggorokan) (lihat bagian “Peringatan dan tindakan pencegahan” dan bagian “Obat lain dan Cosyrel”)

Peringatan dan tindakan pencegahan

Konsultasikan dengan dokter atau apoteker Anda sebelum menggunakan Cosyrel, jika Anda:

- memiliki diabetes,
- memiliki penyakit ginjal (termasuk transplantasi ginjal) atau jika Anda menjalani dialisis,
- memiliki penyakit hati,
- memiliki stenosis aorta dan mitral (penyempitan pembuluh darah utama dari jantung) atau kardiomiopati hipetrofik (penyakit otot jantung) atau stenosis arteri ginjal (penyempitan arteri yang memasok darah ke ginjal),
- mengalami peningkatan abnormal kadar hormone yang disebut aldosteron dalam darah Anda (aldosteronisme primer),
- memiliki gagal jantung atau masalah jantung lainnya seperti gangguan kecil dalam irama jantung atau nyeri dada yang parah saat istirahat (angina varian),
- menderita penyakit vaskular kolagen (penyakit jaringan ikat) seperti lupus eritematosus sistemik atau skleroderma,
- dalam diet pembatasan garam atau menggunakan pengganti garam yang mengandung kalium (terlalu banyak kalium dalam darah dapat menyebabkan perubahan denyut jantung),
- menderita diare atau muntah belakangan ini, atau mengalami dehidrasi (Cosyrel dapat menyebabkan penurunan tekanan darah),
- yang menjalani aferesis LDL (pembuangan kolesterol dari darah dengan mesin),
- menjalani pengobatan anti-alergi saat ini atau akan menjalani pengobatan desensitisasi untuk mengurangi efek dari alergi terhadap sengatan lebah atau tawon,
- dalam puasa yang ketat atau diet,
- yang menjalani anestesi dan / atau operasi besar,
- memiliki masalah sirkulasi di kaki Anda,
- memiliki asma atau penyakit paru kronik,

- telah (atau memiliki) psoriasis,
- memiliki tumor pada kelenjar adrenal (feokromositoma),
- memiliki gangguan tiroid (Cosyrel dapat menyembunyikan gejala tiroid yang terlalu aktif),
- memiliki angioedema (reaksi alergi yang parah dengan pembengkakan wajah, bibir, lidah atau tenggorokan dengan kesulitan menelan atau bernapas). Hal ini dapat terjadi setiap saat selama pengobatan. Jika Anda mengalami gejala tersebut, Anda harus berhenti minum Cosyrel dan segera periksa ke dokter,
- berasal dari kulit hitam karena Anda mungkin memiliki risiko angioedema lebih tinggi dan obat-obatan ini mungkin kurang efektif dalam menurunkan tekanan darah Anda dibandingkan pasien non-kulit hitam,
- menggunakan salah satu obat berikut yang digunakan untuk mengobati tekanan darah tinggi:

- "angiotensin II receptor blocker" (ARBs) (juga dikenal sebagai sartans - misalnya valsartan, telmisartan, irbesartan), khususnya jika Anda memiliki masalah ginjal terkait diabetes.
- Aliskiren.

Dokter Anda mungkin memeriksa fungsi ginjal, tekanan darah, dan jumlah elektrolit (misalnya kalium) dalam darah Anda secara rutin. Lihat juga informasi dengan judul "Jangan menggunakan Cosyrel".

- menggunakan salah satu obat berikut, yang dapat meningkatkan risiko angioedema:
 - racecadotril (digunakan untuk mengobati diare)
 - sirolimus, everolimus, temsirolimus, dan obat lainnya yang termasuk Kelas Inhibitor mTOR (digunakan untuk menghindari penolakan pada transplantasi organ)
 - sacubitril (tersedia dalam bentuk kombinasi tetap dengan valsartan), digunakan untuk mengobati gagal jantung jangka panjang.
 - [linagliptin](#), [saxagliptin](#), [sitagliptin](#), [vildagliptin](#) dan obat lain yang termasuk dalam golongan [gliptin](#) (digunakan untuk mengobati [diabetes](#)).

Jangan tiba-tiba berhenti menggunakan Cosyrel karena dapat menyebabkan perburukan berat kondisi jantung. Pengobatan tidak boleh dihentikan tiba-tiba, terutama pada pasien dengan penyakit arteri koroner.

Anda harus memberitahu dokter Anda jika Anda berpikir Anda (atau mungkin menjadi) hamil. Cosyrel tidak dianjurkan di awal kehamilan, dan tidak boleh digunakan lebih dari 3 bulan kehamilan, karena dapat menyebabkan bahaya serius untuk bayi Anda jika digunakan pada tahap itu (lihat bab kehamilan).

Anak dan remaja

Cosyrel tidak dianjurkan untuk digunakan pada anak dan remaja berusia di bawah 18 tahun.

Obat lain dan Cosyrel

Tolong beritahu dokter atau apoteker Anda jika Anda sedang menggunakan, baru-baru ini menggunakan atau mungkin menggunakan obat-obatan lainnya.

Ada beberapa obat yang dapat mengubah efek dari Cosyrel atau efek obat lain yang dapat diubah oleh Cosyrel. Jenis interaksi ini bisa membuat satu atau kedua obat kurang efektif. Atau bisa meningkatkan risiko atau keparahan efek samping.

Pastikan untuk memberitahu dokter Anda jika Anda sedang mengonsumsi obat berikut:

- obat yang digunakan untuk mengontrol tekanan darah atau obat untuk masalah jantung (seperti amiodarone, amlodipine, clonidine, glikosida digitalis, diltiazem, Disopiramid, felodipine, flekainid, lidocaine, metildopa, moxonidine, procainamide, propafenone, quinidine, rilmenidine, verapamil),
- obat lain yang digunakan untuk mengobati tekanan darah tinggi, termasuk angiotensin II receptor blocker (ARB), aliskiren (lihat juga informasi dengan judul "Jangan mengambil Cosyrel" dan "Peringatan dan tindakan pencegahan") atau diuretik (obat yang meningkatkan jumlah urin yang diproduksi oleh ginjal),
- obat hemat kalium (misalnya triamterene, amilorid), suplemen kalium atau pengganti garam yang mengandung kalium, obat lain yang dapat meningkatkan kalium dalam tubuh Anda (seperti heparin _obat yang digunakan untuk mengencerkan darah_ untuk mencegah pembekuan; _—trimethoprim_ dan kotrimoksazol juga dikenal sebagai as-trimetoprim/sulfametoksazol _untuk infeksi yang disebabkan oleh bakteri_),

- obat hemat kalium yang digunakan dalam pengobatan gagal jantung: eplerenon dan spironolakton dengan dosis antara 12,5 mg sampai 50 mg per hari,
- agen simpatomimetik untuk mengobati syok klinis (adrenalin, noradrenalin, dobutamin, isoprenalin, efedrin),
- estramustine yang digunakan dalam terapi kanker,
- obat yang sering digunakan untuk mengobati diare (racecadotril) atau menghindari penolakan pada transplantasi organ (sirolimus, everolimus, temsirolimus, dan obat lainnya yang termasuk Kelas Inhibitor mTOR). Lihat bagian "Peringatan dan tindakan pencegahan".
- sacubitril/valsartan (digunakan untuk mengobati gagal jantung jangka panjang). Lihat bagian "Jangan menggunakan Cosyrel" dan "Peringatan dan tindakan pencegahan".
- lithium yang digunakan untuk mengobati manik atau depresi,
- obat tertentu yang digunakan untuk mengobati depresi seperti imipramine, amitriptilin, inhibitor monoamine oxidase (MAO) (kecuali inhibitor MAO-B),
- obat tertentu yang digunakan untuk mengobati skizofrenia (antipsikotik),
- obat tertentu yang digunakan untuk mengobati epilepsi (fenitoin, barbiturat seperti fenobarbital),
- agen anestesi yang digunakan untuk operasi,
- vasodilator termasuk nitrat (obat yang membuat pembuluh darah menjadi lebih lebar),
- trimetoprim yang digunakan untuk mengobati infeksi,
- heparin yang digunakan untuk mengobati darah yang encer,
- imunosupresan (obat yang mengurangi mekanisme pertahanan tubuh) seperti siklosporin, tacrolimus, yang digunakan untuk pengobatan gangguan auto-imun atau setelah operasi transplantasi,
- allopurinol yang digunakan untuk mengobati asam urat,
- obat parasimpatomimetik yang digunakan untuk mengobati kondisi seperti penyakit Alzheimer atau glaukoma,
- beta-blocker topikal yang digunakan untuk mengobati glaukoma (peningkatan tekanan dalam mata),
- mefloquine yang digunakan untuk mencegah atau mengobati malaria,

- baclofen yang digunakan untuk mengobati kekakuan otot pada penyakit seperti sklerosis multipel,
- garam emas, terutama dengan pemberian intravena (digunakan untuk mengobati gejala artritis reumatoid),
- obat untuk mengobati diabetes seperti insulin, metformin, linagliptin, saxagliptin, sitagliptin, vildagliptin,
- obat anti-inflamasi non-steroid (AINS) seperti ibuprofen atau diklofenak atau [acetylsalicylic acid](#) dosis tinggi yang digunakan untuk mengobati radang sendi, sakit kepala, nyeri atau peradangan.

Cosyrel dengan makanan, minuman dan alkohol

Lebih baik menggunakan Cosyrel sebelum makan.

Kehamilan dan menyusui

Jika Anda sedang hamil atau menyusui, kemungkinan hamil atau berencana untuk hamil, konsultasikan kepada dokter atau apoteker Anda sebelum menggunakan obat ini.

Kehamilan

Anda harus memberitahu dokter Anda jika Anda berpikir Anda (atau mungkin menjadi) hamil. Dokter biasanya akan menyarankan Anda untuk berhenti menggunakan Cosyrel sebelum Anda hamil atau segera setelah Anda tahu Anda sedang hamil dan akan menyarankan Anda untuk menggunakan obat lain selain Cosyrel. Cosyrel tidak dianjurkan di awal kehamilan, dan tidak boleh digunakan lebih dari 3 bulan kehamilan, karena dapat menyebabkan bahaya serius untuk bayi Anda jika digunakan setelah bulan ketiga kehamilan.

Menyusui

Katakan kepada dokter Anda jika Anda menyusui atau akan mulai menyusui. Cosyrel tidak dianjurkan untuk ibu yang sedang menyusui, dan dokter Anda dapat memilih obat lain untuk Anda jika Anda ingin menyusui, terutama jika bayi Anda baru lahir, atau lahir prematur.

Mengemudi dan menggunakan mesin

Cosyrel biasanya tidak mempengaruhi kewaspadaan tetapi pusing atau lemas karena tekanan darah rendah dapat terjadi pada beberapa pasien, terutama pada awal pengobatan atau setelah penggantian pengobatan serta terkait dengan alkohol. Jika Anda mengalami hal tersebut maka kemampuan Anda untuk mengemudi atau mengoperasikan mesin mungkin terganggu.

Cosyrel mengandung kurang dari 1 mmol natrium (23 mg) per tablet, sehingga pada dasarnya ‘bebas-natrium’.

3. Bagaimana menggunakan Cosyrel

Selalu minum obat ini sesuai anjuran dokter atau apoteker Anda. Konsultasikan dengan dokter atau apoteker Anda jika Anda tidak yakin.

Dosis yang dianjurkan adalah satu tablet sekali sehari. Minum tablet dengan segelas air pada pagi hari sebelum makan.

Dalam beberapa kasus, dokter mungkin meresepkan satu setengah tablet Cosyrel sekali sehari pada pagi hari sebelum makan.

Pasien dengan penyakit ginjal

Cosyrel tidak dianjurkan jika Anda menderita penyakit ginjal.

Penggunaan pada anak-anak dan remaja

Penggunaan pada anak dan remaja tidak dianjurkan.

Jika Anda menggunakan Cosyrel lebih dari yang seharusnya

Jika Anda menggunakan lebih dari yang disarankan, hubungi dokter atau apoteker Anda segera.

Yang paling mungkin berpengaruh dalam kasus overdosis adalah tekanan darah rendah yang dapat membuat Anda merasa pusing atau pingsan (jika hal ini terjadi, berbaring

dengan kaki diangkat dapat membantu), kesulitan bernafas berat, tremor (karena gula darah menurun) dan denyut jantung lambat.

Jika Anda lupa menggunakan Cosyrel

Penting untuk menggunakan obat setiap hari secara teratur supaya pengobatan bekerja lebih baik. Namun, jika Anda terlupa menggunakan dosis Cosyrel, gunakan dosis berikutnya pada waktu yang biasa. Jangan menggunakan dosis ganda untuk mengganti dosis yang terlewat.

Jika Anda berhenti menggunakan Cosyrel

Jangan tiba-tiba berhenti menggunakan Cosyrel atau mengubah dosis tanpa berkonsultasi dengan dokter Anda karena ini dapat menyebabkan memburuknya kondisi jantung yang berat. Pengobatan tidak boleh dihentikan tiba-tiba, terutama pada pasien dengan penyakit arteri koroner.

Jika Anda memiliki pertanyaan lebih lanjut tentang penggunaan obat ini, tanyakan kepada dokter, apoteker atau perawat Anda.

4. Kemungkinan efek samping

Seperti semua obat lainnya, obat ini dapat menyebabkan efek samping, meskipun tidak semua orang mengalaminya.

Berhenti menggunakan produk obat dan hubungi dokter segera, jika Anda mengalami salah satu dari efek samping berikut:

- pusing berat atau pingsan karena tekanan darah rendah (sering terjadi - dapat terjadi hingga 1 dari 10 orang),
- memburuknya gagal jantung dapat menyebabkan peningkatan sesak napas dan / atau retensi cairan (sering terjadi - dapat terjadi hingga 1 dari 10 orang),
- pembengkakan wajah, bibir, mulut, lidah atau tenggorokan, kesulitan bernapas (angioedema) (tidak sering terjadi - dapat terjadi hingga 1 dari 100 orang),
- mengi tiba-tiba, nyeri dada, sesak napas, atau kesulitan bernafas (bronkospasme) (tidak sering terjadi - dapat terjadi hingga 1 dari 100 orang),

- jantung berdetak tidak biasa, cepat atau tidak teratur, nyeri dada (angina) atau serangan jantung (sangat jarang terjadi - dapat terjadi hingga 1 dari 10.000 orang),
- kelemahan pada lengan atau kaki, atau gangguan berbicara yang dapat menjadi tanda stroke (sangat jarang terjadi - dapat terjadi hingga 1 dari 10.000 orang),
- radang pankreas yang dapat menyebabkan nyeri perut dan punggung yang berat disertai dengan perasaan sangat tidak enak (sangat jarang terjadi - dapat terjadi hingga 1 dari 10.000 orang),
- menguningnya kulit atau mata (jaundis) yang bisa menjadi tanda hepatitis (sangat jarang terjadi - dapat terjadi hingga 1 dari 10.000 orang),
- ruam kulit yang sering dimulai dengan bercak merah gatal di wajah, lengan atau kaki (eritema multiformis) (sangat jarang terjadi - dapat terjadi hingga 1 dari 10.000 orang).

Cosyrel biasanya dapat ditoleransi tetapi seperti obat lainnya, setiap orang dapat mengalami efek samping yang berbeda, terutama ketika pengobatan pertama dimulai.

Jika Anda mengalami salah satu efek samping seperti yang tercantum di bawah atau yang tidak tercantum, silahkan beritahu dokter atau apoteker Anda segera:

Sangat sering terjadi (dapat terjadi lebih dari 1 pada 10 orang):

- jantung berdetak lambat.

Sering terjadi (dapat terjadi hingga 1 dari 10 orang):

- sakit kepala,
- pusing,
- vertigo,
- gangguan nafsu makan,
- kesemutan
- mati rasa pada tangan dan kaki,
- gangguan penglihatan,
- berdengung (sensasi suara di telinga),
- perasaan dingin di tangan atau kaki,
- batuk,

- sesak napas,
- gangguan gastrointestinal seperti mual, muntah, sakit perut, kesulitan pencernaan atau dispepsia, diare, sembelit,
- reaksi alergi seperti ruam kulit, gatal,
- kram otot,
- perasaan lelah,
- kelelahan.

Tidak sering terjadi (dapat terjadi hingga 1 dari 100 orang):

- **depresi**
- perubahan suasana hati,
- gangguan tidur,
- depresi,
- mulut kering,
- rasa gatal atau ruam kulit yang berat,
- gambaran melepuh pada kulit,
- peningkatan sensitivitas kulit terhadap sinar matahari (reaksi fotosensitivitas),
- berkeringat,
- gangguan ginjal,
- impotensi,
- eosinofil bertambah (sejenis sel darah putih),
- mengantuk,
- pingsan,
- palpitasi,
- takikardia,
- denyut jantung tidak teratur (gangguan konduksi AV, peradangan pembuluh darah (vaskulitis),
- pusing ketika berdiri,
- kelemahan otot,
- artralgia (nyeri sendi),
- mialgia (nyeri otot),
- sakit dada,

- malaise,
- pembengkakan terlokalisir (edema perifer),
- demam,
- perasaan jatuh,
- perubahan parameter laboratorium: kalium darah meningkat reversibel ketika obat dihentikan, natrium rendah, gula darah yang sangat rendah (hipoglikemia) pada kasus pasien diabetes, peningkatan urea darah, peningkatan kreatinin darah.

Jarang terjadi (dapat terjadi hingga 1 dari 1000 orang):

- gagal ginjal akut,
- urin berwarna gelap, merasa tidak enak badan (mual) atau merasa sakit (muntah-muntah), kram otot, bingung dan kejang. Ini mungkin gejala dari suatu kondisi yang disebut SIADH (sekresi hormon antidiuretik yang tidak tepat).
- Keluar urin menurun atau tidak keluar sama sekali,
- *flushing*,
- mimpi buruk, halusinasi,
- berkurangnya air mata (mata kering),
- masalah pendengaran,
- masalah ereksi,
- peradangan hati yang dapat menyebabkan menguningnya kulit atau putih mata,
- hidung berair, alergi, bersin,
- reaksi alergi seperti gatal, memerah, ruam,
- psoriasis memburuk
- perubahan parameter laboratorium: peningkatan enzim hati, bilirubin serum meningkat, kadar lemak yang berbeda dari normal.

Sangat jarang (mungkin mempengaruhi hingga 1 dari 10.000 orang):

- kebingungan,
- iritasi dan kemerahan pada mata (konjungtivitis),
- eosinophilic pneumonia (jenis pneumonia langka),
- radang pankreas (yang menyebabkan sakit parah di perut dan punggung),
- rambut rontok,
- ruam kulit bersisik (psoriasis), ruam seperti psoriasis,

- perubahan jumlah sel darah seperti jumlah sel darah putih dan merah yang berkurang, hemoglobin berkurang, jumlah trombosit yang berkurang.

Frekuensi tidak diketahui (tidak dapat diperkirakan dari data yang tersedia)

- perubahan warna, mati rasa dan nyeri di jari tangan atau kaki (gejala Raynaud)

Jika anda mengalami gejala-gejala tersebut, hubungi dokter Anda segera.

Pelaporan efek samping obat

Jika Anda mengalami efek samping, konsultasikan dengan dokter atau apoteker Anda. Hal ini termasuk efek samping yang mungkin tidak tercantum dalam leaflet ini. Anda juga dapat melaporkan efek samping langsung melalui e-meso subsite di <https://e-meso.pom.go.id>, atau melaporkan ke Pusat Farmakovigilans/MESO Nasional, Badan Pengawas Obat dan Makanan: Tlp. 021-4244691 Ext.1079, dan Email: pv-center@pom.go.id.

Dengan melaporkan efek samping obat Anda dapat membantu dalam memberikan informasi lebih lanjut tentang keamanan obat ini.

5. Bagaimana cara menyimpan Cosyrel

Jauhkan obat ini dari jangkauan anak-anak.

Masa kadaluwarsa obat ini adalah 2,5 tahun.

Jangan menggunakan obat ini setelah tanggal kadaluwarsa yang tertera pada kemasan.

Tanggal kadaluwarsa mengacu pada hari terakhir pada bulan tersebut.

Obat ini tidak memerlukan kondisi penyimpanan khusus.

Setelah dibuka, Cosyrel harus digunakan dalam waktu 30 hari untuk kemasan berisi 30 tablet salut selaput

Jangan membuang obat melalui saluran air limbah atau sampah rumah tangga.

Tanyakan apoteker Anda bagaimana membuang obat yang tidak lagi digunakan.

Langkah ini akan membantu dalam melindungi lingkungan.

6. Isi dari kemasan dan informasi lainnya

Apa kandungan Cosyrel

- Cosyrel 5 mg/5 mg: Zat aktif bisoprolol fumarate dan perindopril arginine. Tiap tablet Cosyrel mengandung 5 mg bisoprolol fumarate yang setara dengan 4,24 bisoprolol mg dan 5 mg perindopril arginine yang setara dengan 3,395 mg perindopril.
- Cosyrel 5m g/10 mg: Zat aktif bisoprolol fumarate dan perindopril arginine. Tiap tablet Cosyrel mengandung 5 mg bisoprolol fumarate yang setara dengan 4,24 bisoprolol mg dan 10 mg perindopril arginine yang setara dengan 6,790 mg perindopril.
- Cosyrel 10 mg/5 mg: Zat aktif bisoprolol fumarate dan perindopril arginine. Tiap tablet Cosyrel mengandung 10 mg bisoprolol fumarate yang setara dengan 8,49 bisoprolol mg dan 5 mg perindopril arginine yang setara dengan 3,395 mg perindopril.
- Cosyrel 10 mg/10 mg: Zat aktif bisoprolol fumarate dan perindopril arginine. Tiap tablet Cosyrel mengandung 10 mg bisoprolol fumarate yang setara dengan 8,49 bisoprolol mg dan 10 mg perindopril arginine yang setara dengan 6,790 mg perindopril.
- Bahan lainnya adalah mikrokristalin selulosa PH 102 (E460), kalsium karbonat (E170), pati jagung pregelatinisasi, natrium pati glikolat tipe A (E468), silica colloidal anhydrous (E551), magnesium stearate (Liga E572), natrium kroskarmelosa (E468), gliserol (E422), Hypromellose (E464), Macrogol 6000, titanium dioxyde (E171), iron oxide yellow (E172), iron oxide red (E172) dan air murni.

Apa sediaan Cosyrel dan isi kemasan

Cosyrel merupakan tablet salut selaput dua lapis berwarna krem merah muda, lonjong, dengan gambar '  ' pada satu sisi dan '5/5' di sisi lain. Tablet ini dapat dibagi menjadi dua bagian yang sama.

Cosyrel merupakan tablet salut selaput dua lapis berwarna krem merah muda, lonjong, dengan gambar '  ' pada satu sisi dan '5/10' di sisi lain. Tablet ini dapat dibagi menjadi dua bagian yang sama.

Cosyrel merupakan tablet salut selaput dua lapis berwarna krem merah muda, bulat, dengan gambar '☞' pada satu sisi dan '10/5' di sisi lain.

Cosyrel merupakan tablet salut selaput dua lapis berwarna krem merah muda, lonjong, dengan gambar '☞' pada satu sisi dan '10/10' di sisi lain.

Tablet tersedia dalam kemasan kotak berisi 30 tablet salut selaput.

KEMASAN

- Cosyrel 5mg/5mg No. Reg: DKI2068601817A1
Dus, botol plastik @ 30 tablet
- Cosyrel 5mg/10mg No. Reg: DKI2068601817B1
Dus, botol plastik @ 30 tablet
- Cosyrel 10mg/5mg No. Reg: DKI2068601817C1
Dus, botol plastik @ 30 tablet
- Cosyrel 10mg/10mg No. Reg: DKI2068601817D1
Dus, botol plastik @ 30 tablet

HARUS DENGAN RESEP DOKTER

Les Laboratoires Servier - France

Diproduksi oleh:



Les Laboratoires Servier Industrie
Gidy – France

Didaftarkan oleh:

PT. Genero Pharmaceuticals
Bekasi - Indonesia

Diimpor dan dipasarkan oleh:

PT. Servier Indonesia
Jakarta – Indonesia

DDMMYY