Summary of the risk management plan (RMP) for Budesonide-Formoterol Teva Pharma B.V. (budesonide / formoterol)

This is a summary of the risk management plan (RMP) for Budesonide-Formoterol Teva Pharma B.V., which details the measures to be taken in order to ensure that Budesonide-Formoterol Teva Pharma B.V. is used as safely as possible. For more information on RMP summaries, see here.

This RMP summary should be read in conjunction with the EPAR summary and the product information for Budesonide-Formoterol Teva Pharma B.V., which can be found on <u>Budesonide-Formoterol Teva Pharma B.V.'s EPAR page</u>.

Overview of disease epidemiology

Budesonide-Formoterol Teva Pharma B.V. is used in adults to treat asthma-

Asthma

Asthma is a common, life-long inflammatory disease of the airways that affects children and adults of all ages. It is one of the most common chronic diseases worldwide, and can be life-threatening. Symptoms come and go and include shortness of breath, wheezing, chest tightness and cough. The cause of asthma is unknown; however, a family history of asthma, eczema or allergy makes it more likely that an individual will develop asthma.

Estimates of the prevalence in European countries range from around 10 to 13% in the UK to 0.28% in Georgia. There is evidence that its prevalence has considerably increased in recent years, especially in children. Across the world, the number of deaths related to asthma is estimated at around 250,000 per year.

Summary of treatment benefits

Budesonide-Formoterol Teva Pharma B.V. contains two active substances, budesonide and formoterol:

- budesonide belongs to a group of medicines called 'corticosteroids'. It works by reducing and preventing swelling and inflammation in the lungs.
- formoterol belongs to a group of medicines called 'long-acting beta₂-adrenoceptor agonists'
 (LABAs) or 'bronchodilators'. It works by relaxing the muscles in the airways. This helps to breathe
 more easily.

Budesonide-Formoterol Teva Pharma B.V. is a 'hybrid medicine'. This means that it is similar to a 'reference medicine' containing the same active substances, but Budesonide-Formoterol Teva Pharma B.V. is given using a different inhaler. Because Budesonide-Formoterol Teva Pharma B.V. is a hybrid medicine, its benefits and risks are taken as being the same as those of Symbicort Turbohaler, the

reference medicine. Studies in patients have been limited to tests to determine that Budesonide-Formoterol Teva Pharma B.V. is bioequivalent to the reference medicine (two medicines are bioequivalent when they produce the same levels of the active substance in the body). Clinical studies have shown that the addition of formoterol to budesonide improved symptoms and lung function, and reduced exacerbations with asthma.

Unknowns relating to treatment benefits

Budesonide-Formoterol Teva Pharma B.V. has not been studied in children or adolescents under the age of 18 years. There are no data available for use of budesonide/formoterol in patients with reduced liver and kidney function.

Summary of safety concerns

Important identified risks

Risks	What is known	Preventability	
Systemic glucocorticosteroid effects (systemic means that the medicine is carried throughout the body in the bloodstream from the site of application and have general rather than only local effects)	Budesonide is a glucocorticosteroid and shares the actions of this class of hormones. Corticosteroids used at high doses for a long time can lead to the following: changes in bone mineral density (thinning of the	Budesonide/formoterol should be used exactly as instructed by a doctor or pharmacist. Patients should check with their doctor or pharmacist if they are not sure. Patients should talk to their doctor or pharmacist before using budesonide/formoterol if they have	
	bones);cataract (clouding of the lens in the eye);	problems with the thyroid or adrenal glands.	
	 glaucoma (high pressure in the eye); a slowing of the rate of growth 	Patients should tell their doctor or pharmacist if they are taking steroid medicines taken by mouth (such as prednisolone).	
	of children and adolescents; an effect on the adrenal gland (a small gland next to the kidney); restlessness, nervousness or agitation; disturbed sleep;	If patients have been taking steroid tablets for asthma, the doctor may reduce the number of tablets being taken when treatment with budesonide/formoterol is started. If patients have been taking oral steroid tablets for a long time, the doctor may carry out blood tests from time to time.	
	bruising of the skin;depression;changes in behaviour, especially in children.	The doctor may consider adding steroid tablets to the usual treatment during periods of stress (for example, during a chest infection or before an operation).	
	These effects are much less likely to happen with inhaled corticosteroids than with		

corticosteroid tablets. Formoterol is a LABA and shares Heart problems due to Patients should talk to their doctor or formoterol (which the actions of this class of pharmacist before using belongs to a class of 'bronchodilators'. budesonide/formoterol if they have: medicines called long-Palpitations (awareness of the high blood pressure, or have ever acting adrenergic beta heart beating) have been reported had a heart problem (including an 2 receptor agonists or commonly with uneven heartbeat, a very fast LABAs) budesonide/formoterol (affecting pulse, narrowing of the arteries or less than 1 in 10 people), fast heart failure); heartbeat uncommonly (affecting problems with the thyroid or less than 1 in 100 people), chest adrenal glands; pain or tightness in the chest low levels of potassium in the (angina pectoris) very rarely (affecting less than 1 in 10,000 blood. people), and uneven heart beat Patients should tell their doctor or rarely (may affect up to 1 in 1,000 pharmacist if they are taking any of the people). following medicines: medicines for a fast or uneven Mediciral product no lor heart beat (such as quinidine, disopyramide and procainamide); medicines like digoxin, often used to treat heart failure; tricyclic antidepressants (such as amitriptyline) and the antidepressant nefazodone; medicines called monoamine oxidase inhibitors (such as phenelzine, furazolidone and procarbizine); phenothiazine medicines (such as chlorpromazine); medicines for Parkinson's disease (such as levodopa); medicines for thyroid problems (such as levothyroxine); medicines for allergies or antihistamines (such as terfenadine); diuretics, also known as 'water tablets' (such as furosemide), used to treat high blood pressure; steroid medicines taken by mouth

		(such as prednisolone);
		xanthine medicines (such as theophylline or aminophylline),often used to treat asthma;
		other bronchodilators (such as salbutamol).
		The levels of potassium in the blood may be monitored if patients suffer from unstable asthma with variable use of rescue bronchodilators, from acute severe asthma or from other conditions associated with low potassium levels.
Life-threatening and	Serious asthma-related adverse	If asthma symptoms remain
fatal asthma events	events and exacerbations may	uncontrolled or worsen after initiating
with long-acting	occur during treatment.	therapy with budesonide/formoterol,
adrenergic beta 2		patients should continue treatment and
receptor agonists		see their doctor as soon as possible.
(LABAs)		
A sudden temporary	As with other inhalation therapies,	If symptoms occur, patients should
narrowing of the	a paradoxical bronchospasm can	stop using budesonide/formoterol
airways (paradoxical	occur with budesonide/formoterol,	straight away and use their "reliever"
bronchospasm)	causing an immediate increase in	inhaler. They should also contact their
	wheezing, shortness of breath and	doctor immediately as they may need
	cough after dosing.	to have their treatment changed.
	Paradoxical bronchospasm was	
	reported very rarely in patients	
	taking budesonide/formoterol	
	(may affect up to 1 in 10,000	
	people).	
Laurente de la contra del la contra de la contra de la contra del la contra del la contra de la contra de la contra del		Detients should bell be their destance
Low blood levels of	Low levels of potassium in the	Patients should talk to their doctor or
potassium (hypokalaemia) which	blood have been reported rarely (may affect up to 1 in 1,000	pharmacist before using budesonide/formoterol if they have low
can cause muscle	people) in patients taking	levels of potassium in the blood.
weakness, twitching or	budesonide/formoterol.	·
abnormal heart rhythm	Successing of the certain	Patients should tell their doctor or
		pharmacist if they are taking any
		medicines that can lower the amount of
		potassium in the blood:
		diuretics, also known as 'water
		tablets' (such as furosemide).
		These are used to treat high blood
		pressure.
		steroid medicines that are taken by
		mouth (such as prednisolone).
		· , , , , , , , , , , , , , , , , , , ,

 xanthine medicines (such as theophylline or aminophylline). These are often used to treat asthma. 	
 other bronchodilators (such as salbutamol). 	
Patients should tell their doctor if they experience muscle weakness, twitching or abnormal heart rhythm.	
The levels of potassium in the blood may be monitored if patients suffer from unstable asthma with variable use of rescue bronchodilators, from acute	
severe asthma or from other conditions associated with low potassium levels.	

Important potential risks

Risks	What is known
Off-label use in children and adolescents under 18 years (unlicensed use of medicine)	Budesonide-Formoterol Teva Pharma B.V. should not be used in children and adolescents under the age of 18 years. The risk of off-label use arises from the fact that the reference product Symbicort Turbohaler is approved for treatment of asthma as a maintenance therapy in children 6 years and older.
Potential for off-label use of the highest strength (320/9.0) of budesonide/formoterol in the "maintenance and reliever therapy regimen"	Budesonide-Formoterol Teva Pharma B.V. 320/9 micrograms (mcg) should not be used as a reliever inhaler. There is a potential safety concern if the 320/9.0 mcg strength is used to substituted directly for the 160/4.5 mcg strength, given the recommended maximum number of daily inhalations (12) with the maintenance and reliever approach. If the highest strength were used this way, this would expose patients to a high dose of budesonide and formoterol.
Simultaneous use with medicines called 'beta-adrenergic blockers' and 'strong inhibitors of CYP3A4'	 Budesonide/formoterol may interact with certain medicines that are potent inhibitors of CYP3A4: medicines to treat infections (e.g. ketoconazole, itraconazole, voriconazole, posaconazole, clarithromycin and telithromycin); medicines called 'HIV-protease inhibitors' (e.g. ritonavir) to treat HIV infection; the anti-depressant nefazodone. These medicines are likely to markedly increase blood levels of budesonide and potentially induce systemic glucocorticosteroid effects of budesonide/formoterol. Beta-blocker medicines (such as atenolol or propranolol used to treat high blood pressure), including eye drops (such as timolol for

glaucoma), may weaken or inhibit the effect of formoterol and therefore of budesonide/formoterol as well.
Patients should tell their doctor or pharmacist if they are taking, have recently taken or might take any other medicines.

Missing information

Risk	What is known
Use in pregnant or breastfeeding women	There are no data from studies with budesonide/formoterol in pregnant women. During pregnancy, budesonide/formoterol should only be used when the benefits outweigh the potential risks. Budesonide passes into breast milk. It is not known whether formoterol passes into human breast milk. Use of budesonide/formoterol in women who are breastfeeding should only be considered if the expected benefit to the mother is greater than any possible risk to the child.
Use in patients with reduced kidney function	There are no data available on the use of Budesonide-Formoterol Teva Pharma B.V. in patients with reduced kidney function.
Use in patients with reduced liver function	There are no data available on the use of Budesonide-Formoterol Teva Pharma B.V. in patients with reduced liver function. As budesonide and formoterol are mostly broken down in the liver, patients with severe liver problems (e.g. severe liver cirrhosis) may be exposed to higher levels of these medicines.
Use in children and adolescents	The safety and efficacy of Budesonide-Formoterol Teva Pharma B.V. in children and adolescents has not yet been established.

Summary of risk minimisation measures by safety concern

All medicines have a summary of product characteristics (SmPC) which provides physicians, pharmacists and other healthcare professionals with details on how to use the medicine, and also describes the risks and recommendations for minimising them. Information for patients is available in lay language in the package leaflet. The measures listed in these documents are known as 'routine risk minimisation measures'.

The SmPC and the package leaflet are part of the medicine's product information. The product information for Budesonide-Formoterol Teva Pharma B.V. can be found on <u>Budesonide-Formoterol Teva Pharma B.V.'s EPAR page</u>.

This medicine has no additional risk minimisation measures.

Planned post-authorisation development plan

List of studies in post-authorisation development plan

Study/activity (including study number)	Objectives	Safety concerns /efficacy issue addressed	Status	Planned date for submission of (interim and) final results
A paediatric / adolescent pharmacokinetic study (a study of what happens to the drug in the body, including its movement and its metabolism)	To demonstrate that the pharmacokinetic and safety profile of budesonide/formote rol (80/4.5 mcg) is comparable with the reference product Symbicort Turbohaler (100/6 mcg)	Pharmacokinetic and safety profiles of budesonide/formoter ol (80/4.5 mcg) in mild asthmatic children and adolescents is not yet determined	Planned (start of the study is anticipated for Q4 2014)	Not available

Studies which are a condition of the marketing authorisation

None

Summary of changes to the risk management plan over time

Not applicable

This summary was last updated in 10-2014.