ANNEX I SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE MEDICINAL PRODUCT

Daxas 250 micrograms tablets

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each tablet contains 250 micrograms of roflumilast.

Excipient with known effect:

Each tablet contains 49.7 mg lactose monohydrate.

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Tablet.

White to off-white, round tablet, 5 mm in diameter, embossed with "D" on one side and "250" on the other side.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Daxas is indicated for maintenance treatment of severe chronic obstructive pulmonary disease (COPD) (FEV₁ post-bronchodilator less than 50% predicted) associated with chronic bronchitis in adult patients with a history of frequent exacerbations as add on to bronchodilator treatment.

4.2 Posology and method of administration

Posology

Starting dose

The recommended starting dose is one tablet of 250 micrograms roflumilast to be taken once daily, for 28 days.

This starting dose is intended to reduce adverse reactions and patient discontinuation when initiating therapy, but it is a sub-therapeutic dose. Therefore, the 250 micrograms dose should be used only as a starting dose (see sections 5.1 and 5.2).

Maintenance dose

After 28 days of treatment with the 250 micrograms starting dose, patients must be up-titrated to one tablet of 500 micrograms roflumilast, to be taken once daily.

Roflumilast 500 micrograms may need to be taken for several weeks to achieve its full effect (see sections 5.1 and 5.2). Roflumilast 500 micrograms has been studied in clinical trials for up to one year, and is intended for maintenance treatment.

Special populations

Elderly

No dose adjustment is necessary.

Renal impairment

No dose adjustment is necessary.

Hepatic impairment

The clinical data with roflumilast in patients with mild hepatic impairment classified as Child-Pugh A are insufficient to recommend a dose adjustment (see section 5.2) and therefore Daxas should be used with caution in these patients.

Patients with moderate or severe hepatic impairment classified as Child-Pugh B or C must not take Daxas (see section 4.3).

Paediatric population

There is no relevant use of Daxas in the paediatric population (under 18 years) for the indication of COPD.

Method of administration

For oral use.

The tablet should be swallowed with water and taken at the same time every day. The tablet can be taken with or without food.

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients listed in section 6.1. Moderate or severe hepatic impairment (Child-Pugh B or C).

4.4 Special warnings and precautions for use

All patients should be informed about the risks of Daxas and the precautions for safe use before starting treatment.

Rescue medicinal products

Daxas is not indicated as rescue medicinal product for the relief of acute bronchospasms.

Weight decrease

In 1-year studies (M2-124, M2-125), a decrease of body weight occurred more frequently in patients treated with roflumilast compared to placebo-treated patients. After discontinuation of roflumilast, the majority of patients had regained body weight after 3 months.

Body weight of underweight patients should be checked at each visit. Patients should be advised to check their body weight on a regular basis. In the event of an unexplained and clinically concerning weight decrease, the intake of roflumilast should be stopped and body weight should be further followed-up.

Special clinical conditions

Due to lack of relevant experience, treatment with roflumilast should not be initiated or existing treatment with roflumilast should be stopped in patients with severe immunological diseases (e.g. HIV infection, multiple sclerosis, lupus erythematosus, progressive multifocal leukoencephalopathy), severe acute infectious diseases, cancers (except basal cell carcinoma), or patients being treated with immunosuppressive medicinal products (i.e. methotrexate, azathioprine, infliximab, etanercept, or oral corticosteroids to be taken long-term; except short-term systemic corticosteroids). Experience in patients with latent infections such as tuberculosis, viral hepatitis, herpes viral infection and herpes zoster is limited.

Patients with congestive heart failure (NYHA grades 3 and 4) have not been studied and therefore treatment of these patients is not recommended.

Psychiatric disorders

Roflumilast is associated with an increased risk of psychiatric disorders such as insomnia, anxiety, nervousness and depression. Rare instances of suicidal ideation and behaviour, including suicide, have been observed in patients with or without history of depression, usually within the first weeks of treatment (see section 4.8). The risks and benefits of starting or continuing treatment with roflumilast should be carefully assessed if patients report previous or existing psychiatric symptoms or if concomitant treatment with other medicinal products likely to cause psychiatric events is intended. Roflumilast is not recommended in patients with a history of depression associated with suicidal ideation or behaviour. Patients and caregivers should be instructed to notify the prescriber of any changes in behaviour or mood and of any suicidal ideation. If patients suffered from new or worsening psychiatric symptoms, or suicidal ideation or suicidal attempt is identified, it is recommended to discontinue treatment with roflumilast.

Persistent intolerability

While adverse reactions like diarrhoea, nausea, abdominal pain and headache mainly occur within the first weeks of therapy and mostly resolve on continued treatment, roflumilast treatment should be reassessed in case of persistent intolerability. This might be the case in special populations that may have higher exposure, such as in black, non-smoking females (see section 5.2) or in patients concomitantly treated with CYP1A2/ 2C19/3A4 inhibitors (such as fluvoxamine and cimetidine) or the CYP1A2/3A4 inhibitor enoxacin (see section 4.5).

Body weight < 60 kg

Treatment with roflumilast may lead to a higher risk of sleep disorders (mainly insomnia) in patients with a baseline body weight of <60 kg, due to a higher total PDE4 inhibitory activity found in these patients (see section 4.8).

Theophylline

There are no clinical data to support the concomitant treatment with the ophylline for maintenance therapy. Therefore, the concomitant treatment with the ophylline is not recommended.

Lactose content

This medicinal product contains lactose. Patients with rare hereditary problems of galactose intolerance, total lactase deficiency or glucose-galactose malabsorption should not take this medicinal product.

4.5 Interaction with other medicinal products and other forms of interaction

Interaction studies have only been performed in adults.

A major step in roflumilast metabolism is the N-oxidation of roflumilast to roflumilast N-oxide by CYP3A4 and CYP1A2. Both roflumilast and roflumilast N-oxide have intrinsic phosphodiesterase-4 (PDE4) inhibitory activity. Therefore, following administration of roflumilast, the total PDE4 inhibition is considered to be the combined effect of both roflumilast and roflumilast N-oxide. Interaction studies with CYP1A2/3A4 inhibitor enoxacin and the CYP1A2/2C19/3A4 inhibitors cimetidine and fluvoxamine, resulted in increases of the total PDE4 inhibitory activity of 25%, 47% and 59%, respectively. The tested dose of fluvoxamine was 50 mg. A combination of roflumilast with these active substances might lead to an increase of exposure and persistent intolerability. In this case, roflumilast treatment should be reassessed (see section 4.4).

Administration of the cytochrome P450 enzyme inducer rifampicin resulted in a reduction in total PDE4 inhibitory activity by about 60%. Therefore, the use of strong cytochrome P450 enzyme inducers (e.g. phenobarbital, carbamazepine, phenytoin) may reduce the therapeutic efficacy of roflumilast. Thus, roflumilast treatment is not recommended in patients receiving strong cytochrome P450 enzyme inducers.

Clinical interaction studies with CYP3A4 inhibitors erythromycin and ketoconazole showed increases of 9% of the total PDE4 inhibitory activity. Co-administration with theophylline resulted in an increase of 8% of the total PDE4 inhibitory activity (see section 4.4). In an interaction study with an oral contraceptive containing gestodene and ethinyl oestradiol, the total PDE4 inhibitory activity was increased by 17%. No dose adjustment is necessary in patients receiving these active substances.

No interactions were observed with inhaled salbutamol, formoterol, budesonide and oral montelukast, digoxin, warfarin, sildenafil and midazolam.

Co-administration with an antacid (combination of aluminium hydroxide and magnesium hydroxide) did not alter the absorption or pharmacokinetics of roflumilast or its N-oxide.

4.6 Fertility, pregnancy and lactation

Women of childbearing potential

Women of childbearing age should be advised to use an effective method of contraception during treatment. Roflumilast is not recommended in women of childbearing potential not using contraception.

Pregnancy

There are limited amount of data from the use of roflumilast in pregnant women.

Studies in animals have shown reproductive toxicity (see section 5.3). Roflumilast is not recommended during pregnancy.

Roflumilast has been demonstrated to cross the placenta in pregnant rats.

Breastfeeding

Available pharmacokinetic data in animals have shown excretion of roflumilast or its metabolites in milk. A risk to the breastfed infant cannot be excluded. Roflumilast should not be used during breast-feeding.

Fertility

In a human spermatogenesis study, roflumilast 500 micrograms had no effects on semen parameters or reproductive hormones during the 3-month treatment period and the following 3-month off-treatment period.

4.7 Effects on ability to drive and use machines

Daxas has no influence on the ability to drive and use machines.

4.8 Undesirable effects

Summary of the safety profile

The most commonly reported adverse reactions are diarrhoea (5.9%), weight decreased (3.4%), nausea (2.9%), abdominal pain (1.9%) and headache (1.7%). These adverse reactions mainly occurred within the first weeks of therapy and mostly resolved on continued treatment.

Tabulated list of adverse reactions

Within the following table, adverse reactions are ranked under the MedDRA frequency classification:

Very common ($\geq 1/10$); common ($\geq 1/100$ to <1/10); uncommon ($\geq 1/1,000$ to <1/100); rare ($\geq 1/10,000$ to <1/1,000); very rare (<1/10,000), not known (cannot be estimated from the available data).

Within each frequency grouping, adverse reactions are presented in order of decreasing seriousness.

Table 1. Adverse reactions with roflumilast in clinical COPD studies and post-marketing experience

Frequency	Common	Uncommon	Rare
System			
System Organ Class			
Immune system		Hypersensitivity	Angioedema
disorders		Trypersensitivity	ringioedema
Endocrine disorders			Gynaecomastia
Metabolism and	Weight decreased		
nutrition disorders	Decreased		
	appetite		
Psychiatric disorders	Insomnia	Anxiety	Suicidal ideation and
			behaviour
			Depression
			Nervousness
			Panic attack
Nervous system	Headache	Tremor	Dysgeusia
disorders		Vertigo	
		Dizziness	
Cardiac disorders		Palpitations	
Respiratory, thoracic			Respiratory tract
and mediastinal			infections (excluding
disorders	D: 1		Pneumonia)
Gastrointestinal	Diarrhoea	Gastritis	Haematochezia
disorders	Nausea	Vomiting	Constipation
	Abdominal pain	Gastro-esophageal reflux disease	
II		Dyspepsia	Gamma-GT increased
Hepatobiliary disorders			
			Aspartate aminotransferase (AST)
			increased
Skin and subcutaneous		Rash	Urticaria
tissue disorders		1/4511	Officaria
Musculoskeletal and		Muscle spasms and	Blood creatine
connective tissue		weakness	phosphokinase (CPK)
disorders		Myalgia	increased
MINJI MOI U		Back pain	1131 0400 4
General disorders and		Malaise	
administration site		Asthenia	
conditions		Fatigue	

Description of selected adverse reactions

In clinical studies and post-marketing experience, rare instances of suicidal ideation and behaviour, including suicide, were reported. Patients and caregivers should be instructed to notify the prescriber of any suicidal ideation (see also section 4.4).

Other special populations

Elderly

A higher incidence of sleep disorders (mainly insomnia) in patients ≥75 years or older was observed in Study RO-2455-404-RD for patients treated with roflumilast when compared to those treated with placebo (3.9% vs 2.3%). The incidence observed was also higher in patients less than 75 years old, treated with roflumilast when compared to those treated with placebo (3.1% vs 2.0%).

Body weight <60 kg

A higher incidence of sleep disorders (mainly insomnia) in patients with a baseline body weight <60 kg was observed in Study RO-2455-404-RD for patients treated with roflumilast when compared to those treated with placebo (6.0% vs 1.7%). The incidence was 2.5% vs 2.2% in patients with a baseline body weight $\ge 60 \text{ kg}$, treated with roflumilast when compared to those treated with placebo.

Concomitant treatment with long acting muscarinic antagonists (LAMA)

A higher incidence of weight decrease, decreased appetite, headache and depression was observed during Study RO-2455-404-RD in patients receiving concomitant roflumilast and long-acting muscarinic antagonists (LAMA) plus concomitant inhaled corticosteroids (ICS) and long acting B₂ agonists (LABA) compared to those treated only with concomitant roflumilast, ICS and LABA. Difference of incidence between roflumilast and placebo was quantitatively greater with concomitant LAMA for weight decreased (7.2% vs 4.2%), decreased appetite (3.7% vs 2.0%), headache (2.4% vs 1.1%) and depression (1.4% vs -0.3%).

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 the national reporting system listed in Appendix V.

4.9 Overdose

Symptoms

In Phase I studies, the following symptoms were observed at an increased rate after single oral doses of 2,500 micrograms and one single dose of 5,000 micrograms (ten times the recommended dose): headache, gastrointestinal disorders, dizziness, palpitations, light-headedness, clamminess and arterial hypotension.

Management

In case of overdose, it is recommended that the appropriate supportive medical care is provided. Since roflumilast is highly protein bound, haemodialysis is not likely to be an efficient method of its removal. It is not known whether roflumilast is dialysable by peritoneal dialysis.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Drugs for obstructive airway diseases, other systemic drugs for obstructive airway diseases, ATC code: R03DX07

Mechanism of action

Roflumilast is a PDE4 inhibitor, a non-steroid, anti-inflammatory active substance designed to target both the systemic and pulmonary inflammation associated with COPD. The mechanism of action is the inhibition of PDE4, a major cyclic adenosine monophosphate (cAMP)-metabolizing enzyme found in structural and inflammatory cells important to the pathogenesis of COPD. Roflumilast targets the PDE4A, 4B and 4D splicing variants with similar potency in the nanomolar range. The affinity to the PDE4C splicing variants is 5 to 10-fold lower. This mechanism of action and the selectivity also apply to roflumilast N-oxide, which is the major active metabolite of roflumilast.

Pharmacodynamic effects

Inhibition of PDE4 leads to elevated intracellular cAMP levels and mitigates COPD-related malfunctions of leukocytes, airway and pulmonary vascular smooth muscle cells, endothelial and airway epithelial cells and fibroblasts in experimental models. Upon *in vitro* stimulation of human neutrophils, monocytes, macrophages or lymphocytes, roflumilast and roflumilast N-oxide suppress the release of inflammatory mediators e.g. leukotriene B4, reactive oxygen species, tumour necrosis factor α , interferon γ and granzyme B.

In patients with COPD, roflumilast reduced sputum neutrophils. Furthermore, roflumilast attenuated influx of neutrophils and eosinophils into the airways of endotoxin challenged healthy volunteers.

Clinical efficacy and safety

In two confirmative replicate one-year studies (M2-124 and M2-125) and two supplementary six-month studies (M2-127 and M2-128), a total number of 4,768 patients were randomised and treated, of whom 2,374 were treated with roflumilast. The design of the studies was parallel-group, double-blind and placebo-controlled.

The one-year studies included patients with a history of severe to very severe COPD [FEV1 (forced expiratory volume in one second) $\leq 50\%$ of predicted] associated with chronic bronchitis, with at least one documented exacerbation in the previous year and with symptoms at baseline as determined by cough and sputum score. Long-acting beta-agonists (LABAs) were allowed in the studies and were used in approximately 50% of the study population. Short-acting anticholinergics (SAMAs) were allowed for those patients not taking LABAs. Rescue medicinal products (salbutamol or albuterol) were allowed on an as-needed basis. The use of inhaled corticosteroids and theophylline was prohibited during the studies. Patients with no history of exacerbations were excluded.

In a pooled analysis of the one-year studies M2-124 and M2-125, roflumilast 500 micrograms once daily significantly improved lung function compared to placebo, on average by 48 ml (pre-bronchodilator FEV₁, primary endpoint, p<0.0001), and by 55 ml (post-bronchodilator FEV₁, p<0.0001). The improvement in lung function was apparent at the first visit after 4 weeks and was maintained up to one year (end of treatment period). The rate (per patient per year) of moderate exacerbations (requiring intervention with systemic glucocorticosteroids) or severe exacerbations (resulting in hospitalisation and/or leading to death) after 1 year was 1.142 with roflumilast and 1.374 with placebo corresponding to a relative risk reduction of 16.9% (95% CI: 8.2% to 24.8%) (primary endpoint, p=0.0003). Effects were similar, independent of previous treatment with inhaled corticosteroids or underlying treatment with LABAs. In the subgroup of patients with history of frequent exacerbations (at least 2 exacerbations during the last year), the rate of exacerbations was 1.526 with roflumilast and 1.941 with placebo corresponding to a relative risk reduction of 21.3% (95% CI: 7.5% to 33.1%). Roflumilast did not significantly reduce the rate of exacerbations compared with placebo in the subgroup of patients with moderate COPD.

The reduction of moderate or severe exacerbations with roflumilast and LABA compared to placebo and LABA was on average 21% (p=0.0011). The respective reduction in exacerbations seen in patients without concomitant LABAs was on average 15% (p=0.0387). The numbers of patients who died due to any reason were equal for those treated with placebo or roflumilast (42 deaths each group; 2.7% each group; pooled analysis).

A total of 2,690 patients were included and randomised in two supportive 1-year studies (M2-111 and M2-112). In contrast to the two confirmative studies, a history of chronic bronchitis and of COPD exacerbations was not requested for patients' inclusion. Inhaled corticosteroids were used in 809 (61%) of the roflumilast treated patients, whereas the use of LABAs and theophylline was prohibited. Roflumilast 500 micrograms once daily significantly improved lung function compared to placebo, on average by 51 ml (pre-bronchodilator FEV₁, p<0.0001), and by 53 ml (post-bronchodilator FEV₁, p<0.0001). The rate of exacerbations (as defined in the protocols) were not significantly reduced by roflumilast in the individual studies (relative risk reduction: 13.5% in Study M2-111 and 6.6% in Study M2-112; p= not significant). Adverse events rates were independent of concomitant treatment with inhaled corticosteroids.

Two six-month supportive studies (M2-127 and M2-128) included patients with a history of COPD for at least 12 months prior to baseline. Both studies included moderate to severe patients with a non-reversible airway obstruction and a FEV $_1$ of 40% to 70% of predicted. Roflumilast or placebo treatment was added to continuous treatment with a long-acting bronchodilator, in particular salmeterol in Study M2-127 or tiotropium in Study M2-128. In the two six-month studies, pre-bronchodilator FEV $_1$ was significantly improved by 49 ml (primary endpoint, p<0.0001) beyond the bronchodilator effect of concomitant treatment with salmeterol in Study M2-127 and by 80 ml (primary endpoint, p<0.0001) incremental to concomitant treatment with tiotropium in Study M2-128.

Study RO-2455-404-RD was a one-year study in COPD patients with a baseline (pre-bronchodilator) FEV $_1$ <50% of predicted normal and a history of frequent exacerbations. The study assessed the effect of roflumilast on COPD exacerbation rate in patients treated with fixed combinations of LABA and inhaled corticosteroids, compared to placebo. A total of 1935 patients were randomised to double-blind medication and approximately 70% were also using a long-acting muscarinic antagonist (LAMA) through the course of the trial. The primary endpoint was reduction in rate of moderate or severe COPD exacerbations per patient per year. The rate of severe COPD exacerbations and changes in FEV $_1$ were evaluated as key secondary endpoints.

Table 2. Summar	v of	f COPD exaceri	bation endn	oints in Stud	v RO-2455-404-RD

		Roflumilast	Placebo	Ratio	Roflumilas	t/Placebo	2-Sided
Exacerbation Category	Analysis model	(N=969) Rate (n)	(N=966) Rate (n)	Rate Ratio	Change (%)	95% CI	p- value
Moderate or severe	Poisson regression	0.805 (380)	0.927 (432)	0.868	-13.2	0.753, 1.002	0.0529
Moderate	Poisson regression	0.574 (287)	0.627 (333)	0.914	-8.6	0.775, 1.078	0.2875
Severe	Negative binomial regression	0.239 (151)	0.315 (192)	0.757	-24.3	0.601, 0.952	0.0175

There was a trend towards a reduction in moderate or severe exacerbations in subjects treated with roflumilast compared with placebo over 52 weeks, which did not achieve statistical significance (Table 2). A pre-specified sensitivity analysis using the negative binomial regression model treatment showed a statistically significant difference of -14.2% (rate ratio: 0.86; 95% CI: 0.74 to 0.99).

The per-protocol Poisson regression analysis and the non-significant sensitivity to drop-out Poisson regression intention-to-treat analysis rate ratios were 0.81 (95% CI: 0.69 to 0.94) and 0.89 (95% CI: 0.77 to 1.02), respectively.

Reductions were achieved in the subgroup of patients concomitantly treated with LAMA (rate ratio: 0.88; 95% CI: 0.75 to 1.04) and in the subgroup not treated with LAMA (rate ratio: 0.83; 95% CI: 0.62 to 1.12).

The rate of severe exacerbations was reduced in the overall patient group (rate ratio: 0.76; 95% CI: 0.60 to 0.95) with a rate of 0.24 per patient/year compared to a rate of 0.32 per patient/year in patients treated with placebo. A similar reduction was achieved in the subgroup of patients concomitantly treated with LAMA (rate ratio: 0.77; 95% CI: 0.60 to 0.99) and in the subgroup not treated with LAMA (rate ratio: 0.71; 95% CI: 0.42 to 1.20).

Roflumilast improved lung function after 4 weeks (sustained over 52 weeks). Post-bronchodilator FEV_1 increased for the roflumilast group by 52 mL (95% CI: 40, 65 mL) and decreased for the placebo group by 4 mL (95% CI: -16, 9 mL). Post-bronchodilator FEV_1 showed a clinically significant improvement in favour of roflumilast by 56 mL over placebo (95% CI: 38, 73 mL).

Seventeen (1.8%) patients in the roflumilast group and 18 (1.9%) patients in the placebo group died during the double-blind treatment period due to any reason and 7 (0.7%) patients in each group due to a COPD exacerbation. The proportion of patients who experienced at least 1 adverse event during the double-blind treatment period were 648 (66.9%) patients and 572 (59.2%) patients in the roflumilast and placebo groups, respectively. The observed adverse reactions for roflumilast in Study RO-2455-404-RD were in line with those already included in section 4.8.

More patients in the roflumilast group (27.6%) than placebo (19.8%) withdrew study medication due to any reason (risk ratio: 1.40; 95% CI: 1.19 to 1.65). The major reasons for trial discontinuation were withdrawal of consent and reported adverse events.

Starting dose titration trial

The tolerability of roflumilast was evaluated in a 12-week randomised, double-blind, parallel group trial (RO-2455-302-RD) in patients with severe COPD associated with chronic bronchitis. At screening, patients were required to have had at least one exacerbation in the previous year and on standard of care COPD maintenance treatment for at least 12 weeks. A total of 1323 patients were randomised to receive roflumilast 500 micrograms once a day for 12 weeks (n=443), roflumilast 500 micrograms every other day for 4 weeks followed by roflumilast 500 micrograms once a day for 8 weeks (n=439), or roflumilast 250 micrograms once a day for 4 weeks followed by roflumilast 500 micrograms once a day for 8 weeks (n=441).

Over the entire study period of 12 weeks, the percentage of patients discontinuing treatment due to any reason was statistically significantly lower in patients initially receiving roflumilast 250 micrograms once a day for 4 weeks followed by roflumilast 500 micrograms once a day for 8 weeks (18.4%) compared to those receiving roflumilast 500 micrograms once a day for 12 weeks (24.6%; Odds Ratio 0.66, 95% CI [0.47, 0.93], p=0.017). The discontinuation rate for those receiving 500 micrograms every other day for 4 weeks followed by 500 micrograms once a day for 8 weeks was not statistically significantly different to those receiving 500 micrograms once a day for 12 weeks. The percentage of patients experiencing a Treatment Emergent Adverse Event (TEAE) of interest, defined as diarrhoea, nausea, headache, decreased appetite, insomnia, and abdominal pain (secondary endpoint), was nominally statistically significantly lower in patients initially receiving roflumilast 250 micrograms once a day for 4 weeks followed by roflumilast 500 micrograms once a day for 8 weeks (45.4%) compared to those receiving roflumilast 500 micrograms once a day for 12 weeks (54.2%, Odds Ratio 0.63, 95% CI [0.47, 0.83], p=0.001). The rate of experiencing a TEAE of interest for those receiving 500 micrograms every other day for 4 weeks followed by 500 micrograms once a day for 8 weeks was not statistically significantly different to those receiving 500 micrograms once a day for 8 weeks was not statistically significantly different to those receiving 500 micrograms once a day for 12 weeks.

Patients receiving a 500 micrograms dose once a day had a median PDE4 inhibitory activity of 1.2 (0.35, 2.03) and those receiving a 250 micrograms dose once a day had a median PDE4 inhibitory activity of 0.6 (0.20, 1.24). Long-term administration at the 250 micrograms dose level may not induce

sufficient PDE4 inhibition to exert clinical efficacy. 250 micrograms once a day is a sub-therapeutic dose, and should be used only as a starting dose for the first 28 days (see sections 4.2 and 5.2).

Paediatric population

The European Medicines Agency has waived the obligation to submit the results of studies with roflumilast in all subsets of the paediatric population in chronic obstructive pulmonary disease (see section 4.2 for information on paediatric use).

5.2 Pharmacokinetic properties

Roflumilast is extensively metabolised in humans, with the formation of a major pharmacodynamically active metabolite, roflumilast N-oxide. Since both roflumilast and roflumilast N-oxide contribute to PDE4 inhibitory activity *in vivo*, pharmacokinetic considerations are based on total PDE4 inhibitory activity (i.e. total exposure to roflumilast and roflumilast N-oxide).

Absorption

The absolute bioavailability of roflumilast following a 500 micrograms oral dose is approximately 80%. Maximum plasma concentrations of roflumilast typically occur approximately one hour after dosing (ranging from 0.5 to 2 hours) in the fasted state. Maximum concentrations of the N-oxide metabolite are reached after about eight hours (ranging from 4 to 13 hours). Food intake does not affect the total PDE4 inhibitory activity, but delays time to maximum concentration (t_{max}) of roflumilast by one hour and reduces C_{max} by approximately 40%. However, C_{max} and t_{max} of roflumilast N-oxide are unaffected.

Distribution

Plasma protein binding of roflumilast and its N-oxide metabolite is approximately 99% and 97%, respectively. Volume of distribution for single dose of 500 micrograms roflumilast is about 2.9 l/kg. Due to the physico-chemical properties, roflumilast is readily distributed to organs and tissues including fatty tissue of mice, hamster and rat. An early distribution phase with marked penetration into tissues is followed by a marked elimination phase out of fatty tissue most probably due to pronounced break-down of parent compound to roflumilast N-oxide. These studies in rats with radiolabelled roflumilast also indicate low penetration across the blood-brain barrier. There is no evidence for a specific accumulation or retention of roflumilast or its metabolites in organs and fatty tissue.

Biotransformation

Roflumilast is extensively metabolised via Phase I (cytochrome P450) and Phase II (conjugation) reactions. The N-oxide metabolite is the major metabolite observed in the plasma of humans. The plasma AUC of the N-oxide metabolite on average is about 10-fold greater than the plasma AUC of roflumilast. Thus, the N-oxide metabolite is considered to be the main contributor to the total PDE4 inhibitory activity *in vivo*.

In vitro studies and clinical interaction studies suggest that the metabolism of roflumilast to its N-oxide metabolite is mediated by CYP1A2 and 3A4. Based on further *in vitro* results in human hepatic microsomes, therapeutic plasma concentrations of roflumilast and roflumilast N-oxide do not inhibit CYP1A2, 2A6, 2B6, 2C8, 2C9, 2C19, 2D6, 2E1, 3A4/5, or 4A9/11. Therefore, there is a low probability of relevant interactions with substances metabolised by these P450 enzymes. In addition, *in vitro* studies demonstrated no induction of the CYP1A2, 2A6, 2C9, 2C19, or 3A4/5 and only a weak induction of CYP2B6 by roflumilast.

Elimination

The plasma clearance after short-term intravenous infusion of roflumilast is about 9.6 l/h. Following an oral dose, the median plasma effective half-life of roflumilast and its N-oxide metabolite are approximately 17 and 30 hours, respectively. Steady state plasma concentrations of roflumilast and its N-oxide metabolite are reached after approximately 4 days for roflumilast and 6 days for roflumilast N-oxide following once-daily dosing. Following intravenous or oral administration of radiolabelled roflumilast, about 20% of the radioactivity was recovered in the faeces and 70% in urine as inactive metabolites.

Linearity/non-linearity

The pharmacokinetics of roflumilast and its N-oxide metabolite are dose-proportional over a range of doses from 250 micrograms to 1,000 micrograms.

Special populations

In older people, females and in non-Caucasians, total PDE4 inhibitory activity was increased. Total PDE4 inhibitory activity was slightly decreased in smokers. None of these changes were considered to be clinically meaningful. No dose adjustment is recommended in these patients. A combination of factors, such as in black, non-smoking females, might lead to an increase of exposure and persistent intolerability. In this case, roflumilast treatment should be reassessed (see section 4.4).

In Study RO-2455-404-RD when compared with the overall population, the total PDE4 inhibitory activity determined from $ex\ vivo$ unbound fractions was found to be 15% higher in patients \geq 75 years of age, and 11% higher in patients with baseline body weight <60 kg (refer to section 4.4).

Renal impairment

Total PDE4 inhibitory activity decreased by 9% in patients with severe renal impairment (creatinine clearance 10-30 ml/min). No dose adjustment is necessary.

Hepatic impairment

The pharmacokinetics of roflumilast 250 micrograms once-daily was tested in 16 patients with mild to moderate hepatic impairment classified as Child-Pugh A and B. In these patients, the total PDE4 inhibitory activity was increased by about 20% in patients with Child-Pugh A and about 90% in patients with Child-Pugh B. Simulations suggest dose proportionality between roflumilast 250 and 500 micrograms in patients with mild and moderate hepatic impairment. Caution is necessary in Child-Pugh A patients (see section 4.2). Patients with moderate or severe hepatic impairment classified as Child-Pugh B or C should not take roflumilast (see section 4.3).

5.3 Preclinical safety data

There is no evidence for an immunotoxic, skin sensitising or phototoxic potential.

A slight reduction in male fertility was seen in conjunction with epididymal toxicity in rats. No epididymal toxicity or changes in semen parameters were present in any other rodent or non-rodent species including monkeys in spite of higher exposures.

In one of two rat embryofetal development studies, a higher incidence of incomplete skull bone ossification was seen at a dose producing maternal toxicity. In one of three rat studies on fertility and embryofetal development, post-implantation losses were observed. Post-implantation losses were not seen in rabbits. Prolongation of gestation was seen in mice.

The relevance of these findings to humans is unknown.

Most relevant findings in safety pharmacology and toxicology studies occurred at higher doses and exposure than that intended for clinical use. These findings consisted mainly of gastrointestinal

findings (i.e. vomiting, increased gastric secretion, gastric erosions, intestine inflammation) and cardiac findings (i.e. focal haemorrhages, haemosiderin deposits and lympho-histiocytic cell infiltration in the right atria in dogs, and decreased blood pressure and increased heart rate in rats, guinea pigs and dogs).

Rodent-specific toxicity in the nasal mucosa was observed in repeat-dose toxicity and carcinogenicity studies. This effect seems to be due to an ADCP (4-Amino-3,5-dichloro-pyridine) N-oxide intermediate specifically formed in rodent olfactory mucosa, with special binding affinity in these species (i.e. mouse, rat and hamster).

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Lactose monohydrate Maize starch Povidone Magnesium stearate

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

4 years.

6.4 Special precautions for storage

This medicinal product does not require any special storage conditions.

6.5 Nature and contents of container

PVC/PVDC aluminium blisters in packs of 28 tablets.

6.6 Special precautions for disposal

No special requirements.

7. MARKETING AUTHORISATION HOLDER

AstraZeneca AB SE-151 85 Södertälje Sweden

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/10/636/008 28 tablets

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 05 July 2010 Date of latest renewal: 20 May 2020

10. DATE OF REVISION OF THE TEXT

Detailed information on this medicinal product is available on the website of the European Medicines Agency http://www.ema.europa.eu

1. NAME OF THE MEDICINAL PRODUCT

Daxas 500 micrograms film-coated tablets

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each tablet contains 500 micrograms of roflumilast.

Excipient with known effect:

Each film-coated tablet contains 198.64 mg lactose monohydrate.

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Film-coated tablet (tablet).

Yellow, D-shaped film-coated tablet of 9 mm, embossed with "D" on one side.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Daxas is indicated for maintenance treatment of severe chronic obstructive pulmonary disease (COPD) (FEV₁ post-bronchodilator less than 50% predicted) associated with chronic bronchitis in adult patients with a history of frequent exacerbations as add on to bronchodilator treatment.

4.2 Posology and method of administration

Posology

Starting dose

The recommended starting dose is one tablet of 250 micrograms roflumilast to be taken once daily, for 28 days.

This starting dose is intended to reduce adverse reactions and patient discontinuation when initiating therapy, but it is a sub-therapeutic dose. Therefore, the 250 micrograms dose should be used only as a starting dose (see sections 5.1 and 5.2).

Maintenance dose

After 28 days of treatment with the 250 micrograms starting dose, patients must be up-titrated to one tablet of 500 micrograms roflumilast, to be taken once daily.

Roflumilast 500 micrograms may need to be taken for several weeks to achieve its full effect (see sections 5.1 and 5.2). Roflumilast 500 micrograms has been studied in clinical trials for up to one year, and is intended for maintenance treatment.

Special populations

Elderly

No dose adjustment is necessary.

Renal impairment

No dose adjustment is necessary.

Hepatic impairment

The clinical data with roflumilast in patients with mild hepatic impairment classified as Child-Pugh A are insufficient to recommend a dose adjustment (see section 5.2) and therefore Daxas should be used with caution in these patients.

Patients with moderate or severe hepatic impairment classified as Child-Pugh B or C must not take Daxas (see section 4.3).

Paediatric population

There is no relevant use of Daxas in the paediatric population (under 18 years) for the indication of COPD.

Method of administration

For oral use.

The tablet should be swallowed with water and taken at the same time every day. The tablet can be taken with or without food.

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients listed in section 6.1. Moderate or severe hepatic impairment (Child-Pugh B or C).

4.4 Special warnings and precautions for use

All patients should be informed about the risks of Daxas and the precautions for safe use before starting treatment.

Rescue medicinal products

Daxas is not indicated as rescue medicinal product for the relief of acute bronchospasms.

Weight decrease

In 1-year studies (M2-124, M2-125), a decrease of body weight occurred more frequently in patients treated with roflumilast compared to placebo-treated patients. After discontinuation of roflumilast, the majority of patients had regained body weight after 3 months.

Body weight of underweight patients should be checked at each visit. Patients should be advised to check their body weight on a regular basis. In the event of an unexplained and clinically concerning weight decrease, the intake of roflumilast should be stopped and body weight should be further followed-up.

Special clinical conditions

Due to lack of relevant experience, treatment with roflumilast should not be initiated or existing treatment with roflumilast should be stopped in patients with severe immunological diseases (e.g. HIV infection, multiple sclerosis, lupus erythematosus, progressive multifocal leukoencephalopathy), severe acute infectious diseases, cancers (except basal cell carcinoma), or patients being treated with immunosuppressive medicinal products (i.e. methotrexate, azathioprine, infliximab, etanercept, or oral corticosteroids to be taken long-term; except short-term systemic corticosteroids). Experience in patients with latent infections such as tuberculosis, viral hepatitis, herpes viral infection and herpes zoster is limited.

Patients with congestive heart failure (NYHA grades 3 and 4) have not been studied and therefore treatment of these patients is not recommended.

Psychiatric disorders

Roflumilast is associated with an increased risk of psychiatric disorders such as insomnia, anxiety, nervousness and depression. Rare instances of suicidal ideation and behaviour, including suicide, have been observed in patients with or without history of depression, usually within the first weeks of treatment (see section 4.8). The risks and benefits of starting or continuing treatment with roflumilast should be carefully assessed if patients report previous or existing psychiatric symptoms or if concomitant treatment with other medicinal products likely to cause psychiatric events is intended. Roflumilast is not recommended in patients with a history of depression associated with suicidal ideation or behaviour. Patients and caregivers should be instructed to notify the prescriber of any changes in behaviour or mood and of any suicidal ideation. If patients suffered from new or worsening psychiatric symptoms, or suicidal ideation or suicidal attempt is identified, it is recommended to discontinue treatment with roflumilast.

Persistent intolerability

While adverse reactions like diarrhoea, nausea, abdominal pain and headache mainly occur within the first weeks of therapy and mostly resolve on continued treatment, roflumilast treatment should be reassessed in case of persistent intolerability. This might be the case in special populations that may have higher exposure, such as in black, non-smoking females (see section 5.2) or in patients concomitantly treated with CYP1A2/2C19/3A4 inhibitors (such as fluvoxamine and cimetidine) or the CYP1A2/3A4 inhibitor enoxacin (see section 4.5).

Body weight < 60 kg

Treatment with roflumilast may lead to a higher risk of sleep disorders (mainly insomnia) in patients with a baseline body weight of <60 kg, due to a higher total PDE4 inhibitory activity found in these patients (see section 4.8).

Theophylline

There are no clinical data to support the concomitant treatment with the ophylline for maintenance therapy. Therefore, the concomitant treatment with the ophylline is not recommended.

Lactose content

This medicinal product contains lactose. Patients with rare hereditary problems of galactose intolerance, total lactase deficiency or glucose-galactose malabsorption should not take this medicinal product.

4.5 Interaction with other medicinal products and other forms of interaction

Interaction studies have only been performed in adults.

A major step in roflumilast metabolism is the N-oxidation of roflumilast to roflumilast N-oxide by CYP3A4 and CYP1A2. Both roflumilast and roflumilast N-oxide have intrinsic phosphodiesterase-4 (PDE4) inhibitory activity. Therefore, following administration of roflumilast, the total PDE4 inhibition is considered to be the combined effect of both roflumilast and roflumilast N-oxide. Interaction studies with CYP1A2/3A4 inhibitor enoxacin and the CYP1A2/2C19/3A4 inhibitors cimetidine and fluvoxamine, resulted in increases of the total PDE4 inhibitory activity of 25%, 47% and 59%, respectively. The tested dose of fluvoxamine was 50 mg. A combination of roflumilast with these active substances might lead to an increase of exposure and persistent intolerability. In this case, roflumilast treatment should be reassessed (see section 4.4).

Administration of the cytochrome P450 enzyme inducer rifampicin resulted in a reduction in total PDE4 inhibitory activity by about 60%. Therefore, the use of strong cytochrome P450 enzyme inducers (e.g. phenobarbital, carbamazepine, phenytoin) may reduce the therapeutic efficacy of roflumilast. Thus, roflumilast treatment is not recommended in patients receiving strong cytochrome P450 enzyme inducers.

Clinical interaction studies with CYP3A4 inhibitors erythromycin and ketoconazole showed increases of 9% of the total PDE4 inhibitory activity. Co-administration with theophylline resulted in an increase of 8% of the total PDE4 inhibitory activity (see section 4.4). In an interaction study with an oral contraceptive containing gestodene and ethinyl oestradiol, the total PDE4 inhibitory activity was increased by 17%. No dose adjustment is necessary in patients receiving these active substances.

No interactions were observed with inhaled salbutamol, formoterol, budesonide and oral montelukast, digoxin, warfarin, sildenafil and midazolam.

Co-administration with an antacid (combination of aluminium hydroxide and magnesium hydroxide) did not alter the absorption or pharmacokinetics of roflumilast or its N-oxide.

4.6 Fertility, pregnancy and lactation

Women of childbearing potential

Women of childbearing age should be advised to use an effective method of contraception during treatment. Roflumilast is not recommended in women of childbearing potential not using contraception.

Pregnancy

There are limited amount of data from the use of roflumilast in pregnant women.

Studies in animals have shown reproductive toxicity (see section 5.3). Roflumilast is not recommended during pregnancy.

Roflumilast has been demonstrated to cross the placenta in pregnant rats.

Breastfeeding

Available pharmacokinetic data in animals have shown excretion of roflumilast or its metabolites in milk. A risk to the breastfed infant cannot be excluded. Roflumilast should not be used during breast-feeding.

Fertility

In a human spermatogenesis study, roflumilast 500 micrograms had no effects on semen parameters or reproductive hormones during the 3-month treatment period and the following 3-month off-treatment period.

4.7 Effects on ability to drive and use machines

Daxas has no influence on the ability to drive and use machines.

4.8 Undesirable effects

Summary of the safety profile

The most commonly reported adverse reactions are diarrhoea (5.9%), weight decreased (3.4%), nausea (2.9%), abdominal pain (1.9%) and headache (1.7%). These adverse reactions mainly occurred within the first weeks of therapy and mostly resolved on continued treatment.

Tabulated list of adverse reactions

Within the following table, adverse reactions are ranked under the MedDRA frequency classification:

Very common ($\geq 1/10$); common ($\geq 1/100$ to <1/10); uncommon ($\geq 1/1,000$ to <1/100); rare ($\geq 1/10,000$ to <1/1,000); very rare (<1/10,000), not known (cannot be estimated from the available data).

Within each frequency grouping, adverse reactions are presented in order of decreasing seriousness.

Table 1. Adverse reactions with roflumilast in clinical COPD studies and post-marketing experience

Frequency	Common	Uncommon	Rare
System			
Organ Class			
Immune system		Hypersensitivity	Angioedema
disorders			8
Endocrine disorders			Gynaecomastia
Metabolism and	Weight decreased		
nutrition disorders	Decreased		
	appetite		
Psychiatric disorders	Insomnia	Anxiety	Suicidal ideation and
			behaviour
			Depression
			Nervousness
			Panic attack
Nervous system	Headache	Tremor	Dysgeusia
disorders		Vertigo	
		Dizziness	
Cardiac disorders		Palpitations	
Respiratory, thoracic			Respiratory tract
and mediastinal			infections (excluding
disorders			Pneumonia)
Gastrointestinal	Diarrhoea	Gastritis	Haematochezia
disorders	Nausea	Vomiting	Constipation
	Abdominal pain	Gastro-esophageal	
		reflux disease	
		Dyspepsia	
Hepatobiliary disorders			Gamma-GT increased
			Aspartate
			aminotransferase (AST)
		- 1	increased
Skin and subcutaneous		Rash	Urticaria
tissue disorders		26 1	D1 1 1
Musculoskeletal and		Muscle spasms and	Blood creatine
connective tissue		weakness	phosphokinase (CPK)
disorders		Myalgia	increased
G 111 1 1		Back pain	
General disorders and		Malaise	
administration site		Asthenia	
conditions		Fatigue	

Description of selected adverse reactions

In clinical studies and post-marketing experience, rare instances of suicidal ideation and behaviour, including suicide, were reported. Patients and caregivers should be instructed to notify the prescriber of any suicidal ideation (see also section 4.4).

Other special populations

Elderly

A higher incidence of sleep disorders (mainly insomnia) in patients ≥75 years or older was observed in Study RO-2455-404-RD for patients treated with roflumilast when compared to those treated with placebo (3.9% vs 2.3%). The incidence observed was also higher in patients less than 75 years old, treated with roflumilast when compared to those treated with placebo (3.1% vs 2.0%).

Body weight <60 kg

A higher incidence of sleep disorders (mainly insomnia) in patients with a baseline body weight <60 kg was observed in Study RO-2455-404-RD for patients treated with roflumilast when compared to those treated with placebo (6.0% vs 1.7%). The incidence was 2.5% vs 2.2% in patients with a baseline body weight $\ge60 \text{ kg}$, treated with roflumilast when compared to those treated with placebo.

Concomitant treatment with long acting muscarinic antagonists (LAMA)

A higher incidence of weight decrease, decreased appetite, headache and depression was observed during Study RO-2455-404-RD in patients receiving concomitant roflumilast and long-acting muscarinic antagonists (LAMA) plus concomitant inhaled corticosteroids (ICS) and long acting B₂ agonists (LABA) compared to those treated only with concomitant roflumilast, ICS and LABA. Difference of incidence between roflumilast and placebo was quantitatively greater with concomitant LAMA for weight decreased (7.2% vs 4.2%), decreased appetite (3.7% vs 2.0%), headache (2.4% vs 1.1%) and depression (1.4% vs -0.3%).

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 the national reporting system listed in Appendix V.

4.9 Overdose

Symptoms

In Phase I studies, the following symptoms were observed at an increased rate after single oral doses of 2,500 micrograms and one single dose of 5,000 micrograms (ten times the recommended dose): headache, gastrointestinal disorders, dizziness, palpitations, light-headedness, clamminess and arterial hypotension.

Management

In case of overdose, it is recommended that the appropriate supportive medical care is provided. Since roflumilast is highly protein bound, haemodialysis is not likely to be an efficient method of its removal. It is not known whether roflumilast is dialysable by peritoneal dialysis.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Drugs for obstructive airway diseases, other systemic drugs for obstructive airway diseases, ATC code: R03DX07

Mechanism of action

Roflumilast is a PDE4 inhibitor, a non-steroid, anti-inflammatory active substance designed to target both the systemic and pulmonary inflammation associated with COPD. The mechanism of action is the inhibition of PDE4, a major cyclic adenosine monophosphate (cAMP)-metabolizing enzyme found in structural and inflammatory cells important to the pathogenesis of COPD. Roflumilast targets the PDE4A, 4B and 4D splicing variants with similar potency in the nanomolar range. The affinity to the PDE4C splicing variants is 5 to 10-fold lower. This mechanism of action and the selectivity also apply to roflumilast N-oxide, which is the major active metabolite of roflumilast.

Pharmacodynamic effects

Inhibition of PDE4 leads to elevated intracellular cAMP levels and mitigates COPD-related malfunctions of leukocytes, airway and pulmonary vascular smooth muscle cells, endothelial and airway epithelial cells and fibroblasts in experimental models. Upon *in vitro* stimulation of human neutrophils, monocytes, macrophages or lymphocytes, roflumilast and roflumilast N-oxide suppress the release of inflammatory mediators e.g. leukotriene B4, reactive oxygen species, tumour necrosis factor α, interferon γ and granzyme B.

In patients with COPD, roflumilast reduced sputum neutrophils. Furthermore, roflumilast attenuated influx of neutrophils and eosinophils into the airways of endotoxin challenged healthy volunteers.

Clinical efficacy and safety

In two confirmative replicate one-year studies (M2-124 and M2-125) and two supplementary six-month studies (M2-127 and M2-128), a total number of 4,768 patients were randomised and treated of whom 2,374 were treated with roflumilast. The design of the studies was parallel-group, double-blind and placebo-controlled.

The one-year studies included patients with a history of severe to very severe COPD [FEV1 (forced expiratory volume in one second) $\leq 50\%$ of predicted] associated with chronic bronchitis, with at least one documented exacerbation in the previous year and with symptoms at baseline as determined by cough and sputum score. Long-acting beta-agonists (LABAs) were allowed in the studies and were used in approximately 50% of the study population. Short-acting anticholinergics (SAMAs) were allowed for those patients not taking LABAs. Rescue medicinal products (salbutamol or albuterol) were allowed on an as-needed basis. The use of inhaled corticosteroids and theophylline was prohibited during the studies. Patients with no history of exacerbations were excluded.

In a pooled analysis of the one-year studies M2-124 and M2-125, roflumilast 500 micrograms once daily significantly improved lung function compared to placebo, on average by 48 ml (pre-bronchodilator FEV₁, primary endpoint, p<0.0001), and by 55 ml (post-bronchodilator FEV₁, p<0.0001). The improvement in lung function was apparent at the first visit after 4 weeks and was maintained up to one year (end of treatment period). The rate (per patient per year) of moderate exacerbations (requiring intervention with systemic glucocorticosteroids) or severe exacerbations (resulting in hospitalisation and/or leading to death) after 1 year was 1.142 with roflumilast and 1.374 with placebo corresponding to a relative risk reduction of 16.9% (95% CI: 8.2% to 24.8%) (primary endpoint, p=0.0003). Effects were similar, independent of previous treatment with inhaled corticosteroids or underlying treatment with LABAs. In the subgroup of patients with history of frequent exacerbations (at least 2 exacerbations during the last year), the rate of exacerbations was 1.526 with roflumilast and 1.941 with placebo corresponding to a relative risk reduction of 21.3% (95% CI: 7.5% to 33.1%). Roflumilast did not significantly reduce the rate of exacerbations compared with placebo in the subgroup of patients with moderate COPD.

The reduction of moderate or severe exacerbations with roflumilast and LABA compared to placebo and LABA was on average 21% (p=0.0011). The respective reduction in exacerbations seen in patients without concomitant LABAs was on average 15% (p=0.0387). The numbers of patients who died due to any reason were equal for those treated with placebo or roflumilast (42 deaths each group; 2.7% each group; pooled analysis).

A total of 2,690 patients were included and randomised in two supportive 1-year studies (M2-111 and M2-112). In contrast to the two confirmative studies, a history of chronic bronchitis and of COPD exacerbations was not requested for patients' inclusion. Inhaled corticosteroids were used in 809 (61%) of the roflumilast treated patients, whereas the use of LABAs and theophylline was prohibited. Roflumilast 500 micrograms once daily significantly improved lung function compared to placebo, on average by 51 ml (pre-bronchodilator FEV₁, p<0.0001), and by 53 ml (post-bronchodilator FEV₁, p<0.0001). The rate of exacerbations (as defined in the protocols) were not significantly reduced by roflumilast in the individual studies (relative risk reduction: 13.5% in Study M2-111 and 6.6% in Study M2-112; p= not significant). Adverse events rates were independent of concomitant treatment with inhaled corticosteroids.

Two six-month supportive studies (M2-127 and M2-128) included patients with a history of COPD for at least 12 months prior to baseline. Both studies included moderate to severe patients with a non-reversible airway obstruction and a FEV $_1$ of 40% to 70% of predicted. Roflumilast or placebo treatment was added to continuous treatment with a long-acting bronchodilator, in particular salmeterol in Study M2-127 or tiotropium in Study M2-128. In the two six-month studies, pre-bronchodilator FEV $_1$ was significantly improved by 49 ml (primary endpoint, p<0.0001) beyond the bronchodilator effect of concomitant treatment with salmeterol in Study M2-127 and by 80 ml (primary endpoint, p<0.0001) incremental to concomitant treatment with tiotropium in Study M2-128.

Study RO-2455-404-RD was a one-year study in COPD patients with a baseline (pre-bronchodilator) FEV $_1$ <50% of predicted normal and a history of frequent exacerbations. The study assessed the effect of roflumilast on COPD exacerbation rate in patients treated with fixed combinations of LABA and inhaled corticosteroids, compared to placebo. A total of 1935 patients were randomised to double-blind medication and approximately 70% were also using a long-acting muscarinic antagonist (LAMA) through the course of the trial. The primary endpoint was reduction in rate of moderate or severe COPD exacerbations per patient per year. The rate of severe COPD exacerbations and changes in FEV $_1$ were evaluated as key secondary endpoints.

Table 2. Summar	v of	f COPD exaceri	bation endn	oints in Stud	v RO-2455-404-RD

		Roflumilast	Placebo	Ratio F	Roflumilast	/Placebo	2-Sided
Exacerbation Category	Analysis model	(N=969) Rate (n)	(N=966) Rate (n)	Rate Ratio	Change (%)	95% CI	p- value
Moderate or severe	Poisson regression	0.805 (380)	0.927 (432)	0.868	-13.2	0.753, 1.002	0.0529
Moderate	Poisson regression	0.574 (287)	0.627 (333)	0.914	-8.6	0.775, 1.078	0.2875
Severe	Negative binomial regression	0.239 (151)	0.315 (192)	0.757	-24.3	0.601, 0.952	0.0175

There was a trend towards a reduction in moderate or severe exacerbations in subjects treated with roflumilast compared with placebo over 52 weeks, which did not achieve statistical significance (Table 2). A pre-specified sensitivity analysis using the negative binomial regression model treatment showed a statistically significant difference of -14.2% (rate ratio: 0.86; 95% CI: 0.74 to 0.99).

The per-protocol Poisson regression analysis and the non-significant sensitivity to drop-out Poisson regression intention-to-treat analysis rate ratios were 0.81 (95% CI: 0.69 to 0.94) and 0.89 (95% CI: 0.77 to 1.02), respectively.

Reductions were achieved in the subgroup of patients concomitantly treated with LAMA (rate ratio: 0.88; 95% CI: 0.75 to 1.04) and in the subgroup not treated with LAMA (rate ratio: 0.83; 95% CI: 0.62 to 1.12).

The rate of severe exacerbations was reduced in the overall patient group (rate ratio: 0.76; 95% CI: 0.60 to 0.95) with a rate of 0.24 per patient/year compared to a rate of 0.32 per patient/year in patients treated with placebo. A similar reduction was achieved in the subgroup of patients concomitantly treated with LAMA (rate ratio: 0.77; 95% CI: 0.60 to 0.99) and in the subgroup not treated with LAMA (rate ratio: 0.71; 95% CI: 0.42 to 1.20).

Roflumilast improved lung function after 4 weeks (sustained over 52 weeks). Post-bronchodilator FEV_1 increased for the roflumilast group by 52 mL (95% CI: 40, 65 mL) and decreased for the placebo group by 4 mL (95% CI: -16, 9 mL). Post-bronchodilator FEV_1 showed a clinically significant improvement in favour of roflumilast by 56 mL over placebo (95% CI: 38, 73 mL).

Seventeen (1.8%) patients in the roflumilast group and 18 (1.9%) patients in the placebo group died during the double-blind treatment period due to any reason and 7 (0.7%) patients in each group due to a COPD exacerbation. The proportion of patients who experienced at least 1 adverse event during the double-blind treatment period were 648 (66.9%) patients and 572 (59.2%) patients in the roflumilast and placebo groups, respectively. The observed adverse reactions for roflumilast in Study RO-2455-404-RD were in line with those already included in section 4.8.

More patients in the roflumilast group (27.6%) than placebo (19.8%) withdrew study medication due to any reason (risk ratio: 1.40; 95% CI: 1.19 to 1.65). The major reasons for trial discontinuation were withdrawal of consent and reported adverse events.

Starting dose titration trial

The tolerability of roflumilast was evaluated in a 12-week randomised, double-blind, parallel group trial (RO-2455-302-RD) in patients with severe COPD associated with chronic bronchitis. At screening, patients were required to have had at least one exacerbation in the previous year and on standard of care COPD maintenance treatment for at least 12 weeks. A total of 1323 patients were randomised to receive roflumilast 500 micrograms once a day for 12 weeks (n=443), roflumilast 500 micrograms every other day for 4 weeks followed by roflumilast 500 micrograms once a day for 8 weeks (n=439), or roflumilast 250 micrograms once a day for 4 weeks followed by roflumilast 500 micrograms once a day for 8 weeks (n=441).

Over the entire study period of 12 weeks, the percentage of patients discontinuing treatment due to any reason was statistically significantly lower in patients initially receiving roflumilast 250 micrograms once a day for 4 weeks followed by roflumilast 500 micrograms once a day for 8 weeks (18.4%) compared to those receiving roflumilast 500 micrograms once a day for 12 weeks (24.6%; Odds Ratio 0.66, 95% CI [0.47, 0.93], p=0.017). The discontinuation rate for those receiving 500 micrograms every other day for 4 weeks followed by 500 micrograms once a day for 8 weeks was not statistically significantly different to those receiving 500 micrograms once a day for 12 weeks. The percentage of patients experiencing a Treatment Emergent Adverse Event (TEAE) of interest, defined as diarrhoea, nausea, headache, decreased appetite, insomnia, and abdominal pain (secondary endpoint), was nominally statistically significantly lower in patients initially receiving roflumilast 250 micrograms once a day for 4 weeks followed by roflumilast 500 micrograms once a day for 8 weeks (45.4%) compared to those receiving roflumilast 500 micrograms once a day for 12 weeks (54.2%, Odds Ratio 0.63, 95% CI [0.47, 0.83], p=0.001). The rate of experiencing a TEAE of interest for those receiving 500 micrograms every other day for 4 weeks followed by 500 micrograms once a day for 8 weeks was not statistically significantly different to those receiving 500 micrograms once a day for 12 weeks.

Patients receiving a 500 micrograms dose once a day had a median PDE4 inhibitory activity of 1.2 (0.35, 2.03) and those receiving a 250 micrograms dose once a day had a median PDE4 inhibitory activity of 0.6 (0.20, 1.24). Long-term administration at the 250 micrograms dose level may not induce

sufficient PDE4 inhibition to exert clinical efficacy. 250 micrograms once a day is a sub-therapeutic dose, and should be used only as a starting dose for the first 28 days (see sections 4.2 and 5.2).

Paediatric population

The European Medicines Agency has waived the obligation to submit the results of studies with roflumilast in all subsets of the paediatric population in chronic obstructive pulmonary disease (see section 4.2 for information on paediatric use).

5.2 Pharmacokinetic properties

Roflumilast is extensively metabolised in humans, with the formation of a major pharmacodynamically active metabolite, roflumilast N-oxide. Since both roflumilast and roflumilast N-oxide contribute to PDE4 inhibitory activity *in vivo*, pharmacokinetic considerations are based on total PDE4 inhibitory activity (i.e. total exposure to roflumilast and roflumilast N-oxide).

Absorption

The absolute bioavailability of roflumilast following a 500 micrograms oral dose is approximately 80%. Maximum plasma concentrations of roflumilast typically occur approximately one hour after dosing (ranging from 0.5 to 2 hours) in the fasted state. Maximum concentrations of the N-oxide metabolite are reached after about eight hours (ranging from 4 to 13 hours). Food intake does not affect the total PDE4 inhibitory activity, but delays time to maximum concentration (t_{max}) of roflumilast by one hour and reduces C_{max} by approximately 40%. However, C_{max} and t_{max} of roflumilast N-oxide are unaffected.

Distribution

Plasma protein binding of roflumilast and its N-oxide metabolite is approximately 99% and 97%, respectively. Volume of distribution for single dose of 500 micrograms roflumilast is about 2.9 l/kg. Due to the physico-chemical properties, roflumilast is readily distributed to organs and tissues including fatty tissue of mice, hamster and rat. An early distribution phase with marked penetration into tissues is followed by a marked elimination phase out of fatty tissue most probably due to pronounced break-down of parent compound to roflumilast N-oxide. These studies in rats with radiolabelled roflumilast also indicate low penetration across the blood-brain barrier. There is no evidence for a specific accumulation or retention of roflumilast or its metabolites in organs and fatty tissue.

Biotransformation

Roflumilast is extensively metabolised via Phase I (cytochrome P450) and Phase II (conjugation) reactions. The N-oxide metabolite is the major metabolite observed in the plasma of humans. The plasma AUC of the N-oxide metabolite on average is about 10-fold greater than the plasma AUC of roflumilast. Thus, the N-oxide metabolite is considered to be the main contributor to the total PDE4 inhibitory activity *in vivo*.

In vitro studies and clinical interaction studies suggest that the metabolism of roflumilast to its N-oxide metabolite is mediated by CYP1A2 and 3A4. Based on further *in vitro* results in human hepatic microsomes, therapeutic plasma concentrations of roflumilast and roflumilast N-oxide do not inhibit CYP1A2, 2A6, 2B6, 2C8, 2C9, 2C19, 2D6, 2E1, 3A4/5, or 4A9/11. Therefore, there is a low probability of relevant interactions with substances metabolised by these P450 enzymes. In addition, *in vitro* studies demonstrated no induction of the CYP1A2, 2A6, 2C9, 2C19, or 3A4/5 and only a weak induction of CYP2B6 by roflumilast.

Elimination

The plasma clearance after short-term intravenous infusion of roflumilast is about 9.6 l/h. Following an oral dose, the median plasma effective half-life of roflumilast and its N-oxide metabolite are approximately 17 and 30 hours, respectively. Steady state plasma concentrations of roflumilast and its N-oxide metabolite are reached after approximately 4 days for roflumilast and 6 days for roflumilast N-oxide following once-daily dosing. Following intravenous or oral administration of radiolabelled roflumilast, about 20% of the radioactivity was recovered in the faeces and 70% in urine as inactive metabolites.

Linearity/non-linearity

The pharmacokinetics of roflumilast and its N-oxide metabolite are dose-proportional over a range of doses from 250 micrograms to 1,000 micrograms.

Special populations

In older people, females and in non-Caucasians, total PDE4 inhibitory activity was increased. Total PDE4 inhibitory activity was slightly decreased in smokers. None of these changes were considered to be clinically meaningful. No dose adjustment is recommended in these patients. A combination of factors, such as in black, non-smoking females, might lead to an increase of exposure and persistent intolerability. In this case, roflumilast treatment should be reassessed (see section 4.4).

In Study RO-2455-404-RD when compared with the overall population, the total PDE4 inhibitory activity determined from $ex\ vivo$ unbound fractions was found to be 15% higher in patients \geq 75 years of age, and 11% higher in patients with baseline body weight <60 kg (refer to section 4.4).

Renal impairment

Total PDE4 inhibitory activity decreased by 9% in patients with severe renal impairment (creatinine clearance 10-30 ml/min). No dose adjustment is necessary.

Hepatic impairment

The pharmacokinetics of roflumilast 250 micrograms once-daily was tested in 16 patients with mild to moderate hepatic impairment classified as Child-Pugh A and B. In these patients, the total PDE4 inhibitory activity was increased by about 20% in patients with Child-Pugh A and about 90% in patients with Child-Pugh B. Simulations suggest dose proportionality between roflumilast 250 and 500 micrograms in patients with mild and moderate hepatic impairment. Caution is necessary in Child-Pugh A patients (see section 4.2). Patients with moderate or severe hepatic impairment classified as Child-Pugh B or C should not take roflumilast (see section 4.3).

5.3 Preclinical safety data

There is no evidence for an immunotoxic, skin sensitising or phototoxic potential.

A slight reduction in male fertility was seen in conjunction with epididymal toxicity in rats. No epididymal toxicity or changes in semen parameters were present in any other rodent or non-rodent species including monkeys in spite of higher exposures.

In one of two rat embryofetal development studies, a higher incidence of incomplete skull bone ossification was seen at a dose producing maternal toxicity. In one of three rat studies on fertility and embryofetal development, post-implantation losses were observed. Post-implantation losses were not seen in rabbits. Prolongation of gestation was seen in mice.

The relevance of these findings to humans is unknown.

Most relevant findings in safety pharmacology and toxicology studies occurred at higher doses and exposure than that intended for clinical use. These findings consisted mainly of gastrointestinal

findings (i.e. vomiting, increased gastric secretion, gastric erosions, intestine inflammation) and cardiac findings (i.e. focal haemorrhages, haemosiderin deposits and lympho-histiocytic cell infiltration in the right atria in dogs, and decreased blood pressure and increased heart rate in rats, guinea pigs and dogs).

Rodent-specific toxicity in the nasal mucosa was observed in repeat-dose toxicity and carcinogenicity studies. This effect seems to be due to an ADCP (4-Amino-3,5-dichloro-pyridine) N-oxide intermediate specifically formed in rodent olfactory mucosa, with special binding affinity in these species (i.e. mouse, rat and hamster).

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Core
Lactose monohydrate
Maize starch
Povidone
Magnesium stearate

Coating
Hypromellose
Macrogol (4000)
Titanium dioxide (E171)
Iron oxide yellow (E172)

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

3 years.

6.4 Special precautions for storage

This medicinal product does not require any special storage conditions.

6.5 Nature and contents of container

PVC/PVDC aluminium blisters in packs of 10, 14, 28, 30, 84, 90 or 98 film-coated tablets.

Not all pack sizes may be marketed.

6.6 Special precautions for disposal

No special requirements.

7. MARKETING AUTHORISATION HOLDER

AstraZeneca AB SE-151 85 Södertälje Sweden

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/10/636/001	10 film-coated tablets
EU/1/10/636/002	30 film-coated tablets
EU/1/10/636/003	90 film-coated tablets
EU/1/10/636/004	14 film-coated tablets
EU/1/10/636/005	28 film-coated tablets
EU/1/10/636/006	84 film-coated tablets
EU/1/10/636/007	98 film-coated tablets

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 05 July 2010 Date of latest renewal: 20 May 2020

10. DATE OF REVISION OF THE TEXT

Detailed information on this medicinal product is available on the website of the European Medicines Agency http://www.ema.europa.eu

ANNEX II

- A. MANUFACTURER RESPONSIBLE FOR BATCH RELEASE
- B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE
- C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION
- D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT

A. MANUFACTURER RESPONSIBLE FOR BATCH RELEASE

Name and address of the manufacturer responsible for batch release

Corden Pharma GmbH Otto-Hahn-Str. 68723 Plankstadt Germany

B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE

Medicinal product subject to medical prescription.

C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION

• Periodic safety update reports (PSURs)

The marketing authorisation holder shall submit PSURs for this product in accordance with the requirements set out in the list of Union reference dates (EURD list) provided for under Article 107c(7) of Directive 2001/83/EC and published on the European medicines web-portal.

D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT

• Risk management plan (RMP)

The marketing authorisation holder (MAH) shall perform the required pharmacovigilance activities and interventions detailed in the agreed RMP presented in Module 1.8.2 of the marketing authorisation and any agreed subsequent updates of the RMP.

An updated RMP should be submitted

- At the request of the European Medicines Agency;
- Whenever the risk management system is modified, especially as the result of new information being received that may lead to a significant change to the benefit/risk profile or as the result of an important (pharmacovigilance or risk minimisation) milestone being reached.

If the dates for submission of a PSUR and the update of a RMP coincide, they can be submitted at the same time.

ANNEX III LABELLING AND PACKAGE LEAFLET

A. LABELLING

PARTICULARS TO APPEAR ON THE OUTER PACKAGING
OUTER CARTON FOR BLISTER
1. NAME OF THE MEDICINAL PRODUCT
Daxas 250 micrograms tablets roflumilast
2. STATEMENT OF ACTIVE SUBSTANCE(S)
Each tablet contains 250 micrograms roflumilast.
3. LIST OF EXCIPIENTS
Contains lactose. See package leaflet for further information.
4. PHARMACEUTICAL FORM AND CONTENTS
28 tablets – 28-day starter pack
5. METHOD AND ROUTE(S) OF ADMINISTRATION
Read the package leaflet before use. Oral use
6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN
Keep out of the sight and reach of children.
7. OTHER SPECIAL WARNING(S), IF NECESSARY
8. EXPIRY DATE
EXP
9. SPECIAL STORAGE CONDITIONS
10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE

11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
AstraZeneca AB SE-151 85 Södertälje Sweden
12. MARKETING AUTHORISATION NUMBER(S)
EU/1/10/636/008 28 tablets
13. BATCH NUMBER
Lot
14. GENERAL CLASSIFICATION FOR SUPPLY
15. INSTRUCTIONS ON USE
16. INFORMATION IN BRAILLE
daxas 250 mcg
17. UNIQUE IDENTIFIER – 2D BARCODE
2D barcode carrying the unique identifier included.
18. UNIQUE IDENTIFIER - HUMAN READABLE DATA
PC SN NN

MIN	MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS		
BLIS	STERS		
1.	NAME OF THE MEDICINAL PRODUCT		
Daxa roflu	s 250 micrograms tablets milast		
2.	NAME OF THE MARKETING AUTHORISATION HOLDER		
Astra	Zeneca (AstraZeneca logo)		
3.	EXPIRY DATE		
EXP			
4.	BATCH NUMBER		
Lot			
5.	OTHER		

PARTICULARS TO APPEAR ON THE OUTER PACKAGING **OUTER CARTON FOR BLISTER** 1. NAME OF THE MEDICINAL PRODUCT Daxas 500 micrograms film-coated tablets roflumilast 2. STATEMENT OF ACTIVE SUBSTANCE(S) Each tablet contains 500 micrograms roflumilast. 3. LIST OF EXCIPIENTS Contains lactose. See package leaflet for further information. 4. PHARMACEUTICAL FORM AND CONTENTS 10 film-coated tablets 14 film-coated tablets 28 film-coated tablets 30 film-coated tablets 84 film-coated tablets 90 film-coated tablets 98 film-coated tablets 5. METHOD AND ROUTE(S) OF ADMINISTRATION Read the package leaflet before use. Oral use. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN Keep out of the sight and reach of children. 7. OTHER SPECIAL WARNING(S), IF NECESSARY 8. **EXPIRY DATE EXP**

9.

SPECIAL STORAGE CONDITIONS

10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE

11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

AstraZeneca AB SE-151 85 Södertälje Sweden

12. MARKETING AUTHORISATION NUMBER(S)

EU/1/10/636/001	10 film-coated tablets
EU/1/10/636/002	30 film-coated tablets
EU/1/10/636/003	90 film-coated tablets
EU/1/10/636/004	14 film-coated tablets
EU/1/10/636/005	28 film-coated tablets
EU/1/10/636/006	84 film-coated tablets
EU/1/10/636/007	98 film-coated tablets

13. BATCH NUMBER

Lot

14. GENERAL CLASSIFICATION FOR SUPPLY

15. INSTRUCTIONS ON USE

16. INFORMATION IN BRAILLE

daxas 500 mcg

17. UNIQUE IDENTIFIER – 2D BARCODE

2D barcode carrying the unique identifier included.

18. UNIQUE IDENTIFIER - HUMAN READABLE DATA

PC

SN

NN

MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS	
BLISTERS	
1.	NAME OF THE MEDICINAL PRODUCT
Daxas	s 500 micrograms tablets milast
2.	NAME OF THE MARKETING AUTHORISATION HOLDER
Astra	Zeneca (AstraZeneca logo)
3.	EXPIRY DATE
EXP	
4.	BATCH NUMBER
Lot	
5.	OTHER

MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS	
CALENDAR BLISTERS	
1. NAME OF THE MEDICINAL PRODUCT	
Daxas 500 micrograms tablets roflumilast	
2. NAME OF THE MARKETING AUTHORISATION HOLDER	
AstraZeneca (AstraZeneca logo)	
3. EXPIRY DATE	
EXP	
4. BATCH NUMBER	
Lot	
5. OTHER	

B. PACKAGE LEAFLET

Package leaflet: Information for the patient

Daxas 250 micrograms tablets roflumilast

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

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

What is in this leaflet:

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

1. What Daxas is and what it is used for

Daxas contains the active substance roflumilast, which is an anti-inflammatory medicine called phosphodiesterase-4 inhibitor. Roflumilast reduces the activity of phosphodiesterase-4, a protein occurring naturally in body cells. When the activity of this protein is reduced, there is less inflammation in the lungs. This helps to stop narrowing of airways occurring in **chronic obstructive pulmonary disease (COPD).** Thus Daxas eases breathing problems.

Daxas is used for maintenance treatment of severe COPD in adults who in the past had frequent worsening of their COPD symptoms (so-called exacerbations) and who have chronic bronchitis. COPD is a chronic disease of the lungs that results in tightening of the airways (obstruction) and swelling and irritation of the walls of the small air passages (inflammation). This leads to symptoms such as coughing, wheezing, chest tightness or difficulty in breathing. Daxas is to be used in addition to bronchodilators.

2. What you need to know before you take Daxas

Do not take Daxas

- if you are allergic to roflumilast or any of the other ingredients of this medicine (listed in section 6)
- if you have moderate or severe liver problems.

Warnings and precautions

Talk to your doctor or pharmacist before taking Daxas.

Sudden attack of breathlessness

Daxas is not intended for the treatment of a sudden attack of breathlessness (acute bronchospasms). In order to relieve a sudden attack of breathlessness it is very important that your doctor provides you with another medicine to be available to you at all times that can cope with such an attack. Daxas will not help you in this situation.

Body weight

You should check your body weight on a regular basis. Talk to your doctor if, while taking this medicine, you observe an unintentional loss of body weight (not related to a diet or exercise programme).

Other diseases

Daxas is not recommended if you have one or more of the following diseases:

- severe immunological diseases such as HIV infection, multiple sclerosis (MS), lupus erythematosus (LE) or progressive multifocal leukoencephalopathy (PML)
- severe acute infectious diseases such as acute hepatitis
- cancer (except basal-cell carcinoma, a slow-growing type of skin cancer)
- or severe impairment of the heart function

There is a lack of relevant experience with Daxas under these conditions. You should talk to your doctor, if you are diagnosed with any of these diseases.

Experience is also limited in patients with a previous diagnosis of tuberculosis, viral hepatitis, herpes viral infection or herpes zoster. Please talk to your doctor if you have one of these diseases.

Symptoms you should be aware of

You may experience diarrhoea, nausea, abdominal pain or headache during the first weeks of treatment with Daxas. Talk to your doctor if these side effects do not resolve within the first weeks of treatment.

Daxas is not recommended in patients with a history of depression associated with suicidal thinking or behaviour. You may also experience sleeplessness, anxiety, nervousness, or depressive mood. Before starting treatment with Daxas, inform your doctor if you are suffering from any symptoms of this kind and of any additional medicines you may take since some of those could increase the probability of these side effects. You or your caregiver should also immediately inform your doctor of any changes in behaviour or mood and of any suicidal thoughts you may have.

Children and adolescents

Do not give this medicine to children and adolescents under 18 years of age.

Other medicines and Daxas

Tell your doctor or pharmacist if you are taking, have recently taken or might take any other medicines, especially the following:

- a medicine containing theophylline (a medicine to treat respiratory diseases), or
- a medicine used for treatment of immunological diseases, such as methotrexate, azathioprine, infliximab, etanercept, or oral corticosteroids to be taken long-term.
- a medicine containing fluvoxamine (a medicine to treat anxiety disorders and depression), enoxacin (a medicine to treat bacterial infections) or cimetidine (a medicine to treat stomach ulcers or heartburn).

The effect of Daxas may be reduced if taken together with rifampicin (an antibiotic medicine) or with phenobarbital, carbamazepine or phenytoin (medicines usually prescribed for the treatment of epilepsy). Ask your doctor for advice.

Daxas may be taken with other medicines used in the treatment of COPD such as inhaled or oral corticosteroids or bronchodilators. Do not stop taking these medicines or reduce their dose unless advised by your doctor.

Pregnancy and breast-feeding

If you are pregnant or breast-feeding, think you may be pregnant or are planning to have a baby, ask your doctor or pharmacist for advice before taking this medicine.

You should not become pregnant during treatment with this medicine and should use an effective method of contraception during therapy, because Daxas may be harmful for the unborn baby.

Driving and using machines

Daxas has no influence on the ability to drive and use machines.

Daxas contains lactose

If you have been told by your doctor that you have an intolerance to some sugars, contact your doctor before taking this medicinal product.

3. How to take Daxas

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

- For the first 28 days the recommended starting dose is one 250 micrograms tablet once daily.
 - The starting dose is a low dose used to help your body get used to the medicine before you start taking the full dose. At this low dose you will not get the full effect from the medicine therefore it is important that you move on to the full dose (called a 'maintenance dose') after 28 days.
- After 28 days the recommended maintenance dose is one 500 micrograms tablet once daily.

Swallow the tablet with some water. You may take this medicine with or without food. Take the tablet at the same time every day.

You may need to take Daxas for several weeks to achieve its beneficial effect.

If you take more Daxas than you should

If you have taken more tablets than you should, you may experience the following symptoms: headache, nausea, diarrhoea, dizziness, throbbing of your heart, light-headedness, clamminess and low blood pressure. Tell your doctor or pharmacist straight away. If possible take your medicine and this leaflet with you.

If you forget to take Daxas

If you forget to take a tablet at the usual time, take the tablet as soon as you remember on the same day. If on one day you have forgotten to take a tablet of Daxas, just carry on the next day with the next tablet as usual. Continue taking your medicine at the usual times. Do not take a double dose to make up for a forgotten dose.

If you stop taking Daxas

It is important to continue taking Daxas for as long as prescribed by your doctor, even when you have no symptoms, in order to maintain control of your lung function.

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

4. Possible side effects

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

You may experience diarrhoea, nausea, stomach ache or headache during the first weeks of treatment with Daxas. Talk to your doctor if these side effects do not resolve within the first weeks of treatment.

Some side effects could be serious. In clinical studies and post-marketing experience, rare instances of suicidal thinking and behaviour (including suicide) were reported. Please notify your doctor immediately of any suicidal thoughts you may have. You may also experience sleeplessness (common), anxiety (uncommon), nervousness (rare), panic attack (rare) or depressive mood (rare).

In uncommon cases allergic reactions may occur. Allergic reactions may affect the skin and in rare cases cause swelling of the eyelids, face, lips and tongue, possibly leading to difficulties in breathing and/or a drop in blood pressure and accelerated heartbeat. In case of an allergic reaction, stop taking Daxas and contact your doctor immediately, or go immediately to the emergency department in the nearest hospital. Take all your medicines and this leaflet with you and provide full information of your current medications.

Other side effects include the following:

Common side effects (may affect up to 1 in 10 people)

- diarrhoea, nausea, stomach ache
- weight decrease, decreased appetite
- headache

Uncommon side effects (may affect up to 1 in 100 people)

- trembling, sensation of spinning head (vertigo), dizziness
- sensation of rapid or irregular heartbeat (palpitations)
- gastritis, vomiting
- reflux of stomach acid to the gullet (acid regurgitations), indigestion
- rash
- muscle pain, muscle weakness or cramps
- back pain
- feeling of weakness or tiredness, feeling unwell.

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

- male breast enlargement
- decreased sense of taste
- respiratory tract infections (excluding pneumonia)
- bloody stools, constipation
- elevation of liver or muscle enzymes (seen in blood tests)
- wheals (urticaria).

Reporting of side effects

If you get any side effects, talk to your doctor or pharmacist. This includes any possible side effects not listed in this leaflet. You can also report side effects directly via the national reporting system listed in Appendix V. By reporting side effects you can help provide more information on the safety of this medicine.

5. How to store Daxas

Keep this medicine out of the sight and reach of children.

Do not use this medicine after the expiry date which is stated on the carton and blister after EXP. The expiry date refers to the last day of that month.

This medicine does not require any special storage conditions.

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

6. Contents of the pack and other information

What Daxas contains

The active substance is roflumilast.

Each Daxas 250 micrograms tablet contains 250 micrograms roflumilast. The other ingredients are lactose monohydrate (see section 2 under "Daxas contains lactose"), maize starch, povidone, magnesium stearate.

What Daxas looks like and contents of the pack

Daxas 250 micrograms tablets are white to off-white, embossed with 'D' on one side and '250' on the other side.

Each pack contains 28 tablets.

Marketing Authorisation Holder

AstraZeneca AB SE-151 85 Södertälje Sweden

Manufacturer

Corden Pharma GmbH Otto-Hahn-Str. 68723 Plankstadt Germany

For any information about this medicine, please contact the local representative of the Marketing Authorisation Holder:

België/Belgique/Belgien

AstraZeneca S.A./N.V. Tel: +32 2 370 48 11

България

АстраЗенека България ЕООД Тел.: +359 24455000

Česká republika

AstraZeneca Czech Republic s.r.o. Tel: +420 222 807 111

Danmark

AstraZeneca A/S Tlf: +45 43 66 64 62

Deutschland

AstraZeneca GmbH Tel: +49 40 809034100

Eesti

AstraZeneca

Tel: +372 6549 600

Ελλάδα

AstraZeneca A.E. Τηλ: +30 210 6871500

España

AstraZeneca Farmacéutica Spain, S.A. Tel: +34 91 301 91 00

France

Lietuva

UAB AstraZeneca Lietuva Tel: +370 5 2660550

Luxembourg/Luxemburg

AstraZeneca S.A./N.V. Tél/Tel: +32 2 370 48 11

Magyarország

AstraZeneca Kft. Tel.: +36 1 883 6500

Malta

Associated Drug Co. Ltd Tel: +356 2277 8000

Nederland

AstraZeneca BV Tel: +31 85 808 9900

Norge

AstraZeneca AS TIf: +47 21 00 64 00

Österreich

AstraZeneca Österreich GmbH Tel: +43 1 711 31 0

Polska

AstraZeneca Pharma Poland Sp. z o.o.

Tel.: +48 22 245 73 00

Portugal

AstraZeneca

Tél: +33 1 41 29 40 00

Hrvatska

AstraZeneca d.o.o. Tel: +385 1 4628 000

Ireland

AstraZeneca Pharmaceuticals (Ireland) DAC

Tel: +353 1609 7100

Ísland Vistor hf.

Sími: +354 535 7000

Italia

Simesa S.p.A.

Tel: +39 02 00704500

Κύπρος

Αλέκτωρ Φαρμακευτική Λτδ

Τηλ: +357 22490305

Latvija

SIA AstraZeneca Latvija Tel: +371 67377100 AstraZeneca Produtos Farmacêuticos, Lda.

Tel: +351 21 434 61 00

România

AstraZeneca Pharma SRL Tel: +40 21 317 60 41

Slovenija

AstraZeneca UK Limited Tel: +386 1 51 35 600

Slovenská republika

AstraZeneca AB, o.z. Tel: +421 2 5737 7777

Suomi/Finland

AstraZeneca Oy

Puh/Tel: +358 10 23 010

Sverige

AstraZeneca AB Tel: +46 8 553 26 000

United Kingdom (Northern Ireland)

AstraZeneca UK Ltd Tel: +44 1582 836 836

This leaflet was last approved in

Detailed information on this medicine is available on the European Medicines Agency web site: http://www.ema.europa.eu

Package leaflet: Information for the patient

Daxas 500 micrograms film-coated tablets roflumilast

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

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

What is in this leaflet:

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

1. What Daxas is and what it is used for

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Daxas is used for maintenance treatment of severe COPD in adults who in the past had frequent worsening of their COPD symptoms (so-called exacerbations) and who have chronic bronchitis. COPD is a chronic disease of the lungs that results in tightening of the airways (obstruction) and swelling and irritation of the walls of the small air passages (inflammation). This leads to symptoms such as coughing, wheezing, chest tightness or difficulty in breathing. Daxas is to be used in addition to bronchodilators.

2. What you need to know before you take Daxas

Do not take Daxas

- if you are allergic to roflumilast or any of the other ingredients of this medicine (listed in section 6)
- if you have moderate or severe liver problems.

Warnings and precautions

Talk to your doctor or pharmacist before taking Daxas.

Sudden attack of breathlessness

Daxas is not intended for the treatment of a sudden attack of breathlessness (acute bronchospasms). In order to relieve a sudden attack of breathlessness it is very important that your doctor provides you with another medicine to be available to you at all times that can cope with such an attack. Daxas will not help you in this situation.

Body weight

You should check your body weight on a regular basis. Talk to your doctor if, while taking this medicine, you observe an unintentional loss of body weight (not related to a diet or exercise programme).

Other diseases

Daxas is not recommended if you have one or more of the following diseases:

- severe immunological diseases such as HIV infection, multiple sclerosis (MS), lupus erythematosus (LE) or progressive multifocal leukoencephalopathy (PML)
- severe acute infectious diseases such as acute hepatitis
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- or severe impairment of the heart function

There is a lack of relevant experience with Daxas under these conditions. You should talk to your doctor, if you are diagnosed with any of these diseases.

Experience is also limited in patients with a previous diagnosis of tuberculosis, viral hepatitis, herpes viral infection or herpes zoster. Please talk to your doctor if you have one of these diseases.

Symptoms you should be aware of

You may experience diarrhoea, nausea, abdominal pain or headache during the first weeks of treatment with Daxas. Talk to your doctor if these side effects do not resolve within the first weeks of treatment.

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Children and adolescents

Do not give this medicine to children and adolescents under 18 years of age.

Other medicines and Daxas

Tell your doctor or pharmacist if you are taking, have recently taken or might take any other medicines, especially the following:

- a medicine containing theophylline (a medicine to treat respiratory diseases), or
- a medicine used for treatment of immunological diseases, such as methotrexate, azathioprine, infliximab, etanercept, or oral corticosteroids to be taken long-term.
- a medicine containing fluvoxamine (a medicine to treat anxiety disorders and depression), enoxacin (a medicine to treat bacterial infections) or cimetidine (a medicine to treat stomach ulcers or heartburn).

The effect of Daxas may be reduced if taken together with rifampicin (an antibiotic medicine) or with phenobarbital, carbamazepine or phenytoin (medicines usually prescribed for the treatment of epilepsy). Ask your doctor for advice.

Daxas may be taken with other medicines used in the treatment of COPD such as inhaled or oral corticosteroids or bronchodilators. Do not stop taking these medicines or reduce their dose unless advised by your doctor.

Pregnancy and breast-feeding

If you are pregnant or breast-feeding, think you may be pregnant or are planning to have a baby, ask your doctor or pharmacist for advice before taking this medicine.

You should not become pregnant during treatment with this medicine and should use an effective method of contraception during therapy, because Daxas may be harmful for the unborn baby.

Driving and using machines

Daxas has no influence on the ability to drive and use machines.

Daxas contains lactose

If you have been told by your doctor that you have an intolerance to some sugars, contact your doctor before taking this medicinal product.

3. How to take Daxas

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

- For the first 28 days the recommended starting dose is one 250 micrograms tablet once daily.
 - The starting dose is a low dose used to help your body get used to the medicine before you start taking the full dose. At this low dose you will not get the full effect from the medicine therefore it is important that you move on to the full dose (called a 'maintenance dose') after 28 days.
- After 28 days the recommended maintenance dose is one 500 micrograms tablet once daily.

Swallow the tablet with some water. You may take this medicine with or without food. Take the tablet at the same time every day.

You may need to take Daxas for several weeks to achieve its beneficial effect.

If you take more Daxas than you should

If you have taken more tablets than you should, you may experience the following symptoms: headache, nausea, diarrhoea, dizziness, throbbing of your heart, light-headedness, clamminess and low blood pressure. Tell your doctor or pharmacist straight away. If possible take your medicine and this leaflet with you.

If you forget to take Daxas

If you forget to take a tablet at the usual time, take the tablet as soon as you remember on the same day. If on one day you have forgotten to take a tablet of Daxas, just carry on the next day with the next tablet as usual. Continue taking your medicine at the usual times. Do not take a double dose to make up for a forgotten dose.

If you stop taking Daxas

It is important to continue taking Daxas for as long as prescribed by your doctor, even when you have no symptoms, in order to maintain control of your lung function.

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

4. Possible side effects

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

You may experience diarrhoea, nausea, stomach ache or headache during the first weeks of treatment with Daxas. Talk to your doctor if these side effects do not resolve within the first weeks of treatment.

Some side effects could be serious. In clinical studies and post-marketing experience, rare instances of suicidal thinking and behaviour (including suicide) were reported. Please notify your doctor immediately of any suicidal thoughts you may have. You may also experience sleeplessness (common), anxiety (uncommon), nervousness (rare), panic attack (rare) or depressive mood (rare).

In uncommon cases allergic reactions may occur. Allergic reactions may affect the skin and in rare cases cause swelling of the eyelids, face, lips and tongue, possibly leading to difficulties in breathing and/or a drop in blood pressure and accelerated heartbeat. In case of an allergic reaction, stop taking Daxas and contact your doctor immediately, or go immediately to the emergency department in the nearest hospital. Take all your medicines and this leaflet with you and provide full information of your current medications.

Other side effects include the following:

Common side effects (may affect up to 1 in 10 people)

- diarrhoea, nausea, stomach ache
- weight decrease, decreased appetite
- headache

Uncommon side effects (may affect up to 1 in 100 people)

- trembling, sensation of spinning head (vertigo), dizziness
- sensation of rapid or irregular heartbeat (palpitations)
- gastritis, vomiting
- reflux of stomach acid to the gullet (acid regurgitations), indigestion
- rash
- muscle pain, muscle weakness or cramps
- back pain
- feeling of weakness or tiredness, feeling unwell.

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

- male breast enlargement
- decreased sense of taste
- respiratory tract infections (excluding pneumonia)
- bloody stools, constipation
- elevation of liver or muscle enzymes (seen in blood tests)
- wheals (urticaria).

Reporting of side effects

If you get any side effects, talk to your doctor or pharmacist. This includes any possible side effects not listed in this leaflet. You can also report side effects directly via the national reporting system listed in Appendix V. By reporting side effects you can help provide more information on the safety of this medicine.

5. How to store Daxas

Keep this medicine out of the sight and reach of children.

Do not use this medicine after the expiry date which is stated on the carton and blister after EXP. The expiry date refers to the last day of that month.

This medicine does not require any special storage conditions.

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

6. Contents of the pack and other information

What Daxas contains

The active substance is roflumilast.

Each film-coated tablet (tablet) contains 500 micrograms roflumilast.

- The other ingredients are:
 - Core: lactose monohydrate (see section 2 under "Daxas contains lactose"), maize starch, povidone, magnesium stearate,
 - Coating: hypromellose, macrogol (4000), titanium dioxide (E171), and iron oxide yellow (E172).

What Daxas looks like and contents of the pack

Daxas 500 micrograms film-coated tablets are yellow, D-shaped film-coated tablets, embossed with 'D' on one side.

Each pack contains 10, 14, 28, 30, 84, 90 or 98 film-coated tablets. Not all pack sizes may be marketed.

Marketing Authorisation Holder

AstraZeneca AB SE-151 85 Södertälje Sweden

Manufacturer

Corden Pharma GmbH Otto-Hahn-Str. 68723 Plankstadt Germany

For any information about this medicine, please contact the local representative of the Marketing Authorisation Holder:

België/Belgique/Belgien

AstraZeneca S.A./N.V. Tel: +32 2 370 48 11

България

АстраЗенека България ЕООД Тел.: +359 24455000

Česká republika

AstraZeneca Czech Republic s.r.o. Tel: +420 222 807 111

Danmark

AstraZeneca A/S Tlf: +45 43 66 64 62

Deutschland

AstraZeneca GmbH Tel: +49 40 809034100

Eesti

AstraZeneca Tel: +372 6549 600

Ελλάδα

AstraZeneca A.E. Tηλ: +30 210 6871500

España

Lietuva

UAB AstraZeneca Lietuva Tel: +370 5 2660550

Luxembourg/Luxemburg

AstraZeneca S.A./N.V. Tél/Tel: +32 2 370 48 11

Magyarország

AstraZeneca Kft. Tel.: +36 1 883 6500

Malta

Associated Drug Co. Ltd Tel: +356 2277 8000

Nederland

AstraZeneca BV Tel: +31 85 808 9900

Norge

AstraZeneca AS Tlf: +47 21 00 64 00

Österreich

AstraZeneca Österreich GmbH

Tel: +43 1 711 31 0

Polska

AstraZeneca Farmacéutica Spain, S.A.

Tel: +34 91 301 91 00

France

AstraZeneca

Tél: +33 1 41 29 40 00

Hrvatska

AstraZeneca d.o.o. Tel: +385 1 4628 000

Ireland

AstraZeneca Pharmaceuticals (Ireland) DAC

Tel: +353 1609 7100

Ísland

Vistor hf.

Sími: +354 535 7000

Italia

Simesa S.p.A.

Tel: +39 02 00704500

Κύπρος

Αλέκτωρ Φαρμακευτική Λτδ

Τηλ: +357 22490305

Latvija

SIA AstraZeneca Latvija

Tel: +371 67377100

AstraZeneca Pharma Poland Sp. z o.o.

Tel.: +48 22 245 73 00

Portugal

AstraZeneca Produtos Farmacêuticos, Lda.

Tel: +351 21 434 61 00

România

AstraZeneca Pharma SRL

Tel: +40 21 317 60 41

Slovenija

AstraZeneca UK Limited

Tel: +386 1 51 35 600

Slovenská republika

AstraZeneca AB, o.z.

Tel: +421 2 5737 7777

Suomi/Finland

AstraZeneca Oy

Puh/Tel: +358 10 23 010

Sverige

AstraZeneca AB

Tel: +46 8 553 26 000

United Kingdom (Northern Ireland)

AstraZeneca UK Ltd

Tel: +44 1582 836 836

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Detailed information on this medicine is available on the European Medicines Agency web site: http://www.ema.europa.eu