## VI.2 Elements for a public summary

## VI.2.1 Overview of disease epidemiology

## Major depressive disorder (MDD)

Major depressive disorder (MDD) is a mental disorder characterized by a pervasive and persistent low mood that is accompanied by low self-esteem and by a loss of interest or pleasure in normallyenjoyable activities. Chemical changes in the brain are believed responsible for this illness that may be due to a problem with person genes or may be triggered by certain stressful events. More likely, it is a combination of both.

The Western European countries 12-month occurrence rate is around 5%, with higher prevalence in women, the middle-aged, less privileged groups, and those experiencing social adversity.

The US 12-month respective rate in adults is 6.7%. The frequency in 18- to 29-year-old individuals is threefold higher than the frequency in those aged 60 or older. The occurrence rates appear to be unrelated to ethnicity, education, income, or marital status. In childhood, boys and girls are equally affected. However, in adolescence and adulthood, the frequency is 1.5- to 3-fold higher in females compared to males.

## **Diabetic Peripheral Neuropathic Pain (DPNP)**

Peripheral neuropathy, also called distal symmetric neuropathy or sensorimotor neuropathy, is nerve damage in the arms and legs. Feet and legs are likely to be affected before hands and arms. About 60-70% of people with diabetes have some form of neuropathy. People with diabetes can develop nerve problems at any time, but risk rises with age and longer duration of diabetes. The highest rates of neuropathy are among people who have had diabetes for at least 25 years. Diabetic neuropathies also appear to be more common in people who have problems controlling their blood glucose, also called blood sugar, as well as those with high levels of blood fat and blood pressure and those who are overweight.

In a landmark study, over 4400 patient with diabetes were serially evaluated over 25 years. Neuropathy was defined as decreased sensation in the feet and depressed or absent ankle reflexes. The onset of neuropathycorrelated positively with the duration of diabetes and, by 25 years, 50 percent of patients had neuropathy.

#### Generalized anxiety disorder (GAD)

Generalized anxiety disorder (GAD) is a long-term condition that causes patient to feel anxious about a wide range of situations and issues, rather than one specific event.

People with GAD feel anxious most days and often struggle to remember the last time they felt relaxed. GAD can cause both mental and physical symptoms. These vary from person to person, but can include feeling restless or worried and having trouble concentrating or sleeping. In Europe, studies suggest that GAD is a relatively rare disorder in the community with a 12-month frequency of about 2%, with a higher incidence in women and most common among older age groups.

Nevertheless, GAD affects about 3.1% American adults age 18 years and older (about 18%) in a given year. The average age of onset is 31 years old. GAD affects about 6.8 million American adults, including twice as many women as men. The disorder develops gradually and can begin at any point in the life cycle, although the years of highest risk are between childhood and middle age.

### VI.2.2 Summary of treatment benefits

**Depression** is commonly associated with physical or painful symptoms.

Serotonin and norepinephrine appear to be involved in the mechanisms of both depression and pain, and these conditions may be mediated through a common pathway. Antidepressants that act via serotonergic or noradrenergic mechanisms (or both) have analgesic properties independent of their effects on mood and have been used successfully to manage the symptoms of various pain states.

Duloxetine, a dual-acting appear to possess analgesic efficacy similar to that of the tricyclicantidepressants (TCAs), but have a more favorable safety and tolerability profile. In addition it has an efficacy advantage over selective serotonin reuptake inhibitors (SSRIs) in treating the painful physical symptoms of depression and in achieving remission of all symptoms of depression.

Studies suggest that diabetic peripheral neuropathic pain is related to an unbalanced release of norepinephrine and serotonin from neurons. Serotonin-norepinephrine reuptake inhibitors (SNRIs), including venlafaxine and duloxetine, are a category of antidepressants for treatment of diabetic peripheral neuropathic pain. They are better tolerated and have fewer drug interactions than tricyclic antidepressants (TCAs).

### VI.2.3 Unknowns relating to treatment benefits

Duloxetine should not be used in children and adolescents aged < 18 years. There is no evidence for its use in this population and antidepressants increase the risk of suicidal thinking and behaviour in children, adolescents and young adults.

There are no adequate data on the use of duloxetine in pregnant women. The potential risk of reproductive toxicity for humans is unknown.

## VI.2.4 Summary of safety concerns

### Important identified risks

Risk	What is known	Preventability

Risk	What is known	Preventability
Liver disease(Hepatic risks)	Duloxetine undergoes Predominantly hepatic metabolism (via two cytochrome P450 isozymes, CYP2D6 and CYP1A2). Circulating metabolites are pharmacologically inactive. However, moderate liver diseases affect pharmacokinetics of duloxetine. In patients with liver problems plasma clearance of duloxetine is lower than in healthy patients. Thus, use of duloxetine in patients with liver diseases may result in hepatic impairment.	Yes.  Duloxetine must not be used in patients with liver disease.  Physician should be aware if patient has liver problems
Thoughts of harming or killing yourself (Suicidality)	Suicide attempts and thoughtsabout committing suicide areuncommon and may affect up to 1in 100 people.	This risk is not preventable.  Patients should contact their doctor or go to a hospital straightaway if they have thoughts of harming or killing themselves at any time. Patients may find it helpful to tell a relative or close friend that they are depressed or have an anxiety disorder and ask them to read the

Risk	What is known	Preventability
		patient leaflet. Patients mightask family or friends to tell them if they think the depression or anxiety symptoms are getting worse or if they are worried about changes in the patient's behaviour. Patients with a history of suiciderelatedbehaviours should be carefully monitored during treatment.
High blood sugar (Hyperglycaemia)	Hyperglycaemia has been reported uncommonly, especially in diabetic patients (may affect up to 1 in 100 people)	Yes.  Blood sugar level in patient with diabetic neuropathy pain should be monitored.
Serious allergic reaction(Stevens-JohnsonSyndrome)	Stevens-Johnson syndrome is a serious illness with blistering of the skin, mouth, eyes and genitals. Based on the postmarketing surveillance of duloxetine adverse events it may occur rarely (may affect up to 1 in 1000 people)	Yes.  Duloxetine administration should be discontinued immediately if Stevens- Johnson syndrome symptoms occur.  Patient should ask physician advice
Loss of blood in thegastrointestinal tract(from the pharynx	There have been reports of bleeding abnormalities, such as ecchymoses, purpura and	Yes.  Caution is advised in patients taking anticoagulants and/or

Risk	What is known	Preventability
to the	gastrointestinal haemorrhage	medicinal products known to
rectum)(Gastrointestin	with selective serotonin	affect platelet function (e.g.
al Tract Bleeding	reuptake inhibitors (SSRIs)	NSAIDs or acetylsalicylic acid
(GITbleeding))	and serotonin/noradrenaline	(ASA)), and in patients with
	reuptake inhibitors (SNRIs),	known bleeding tendencies.
	including duloxetine. Adverse	
	events such as gastrointestinal	
	haemorrhage, haematochezia	
	(black tarry stools) have been	
	reported on the post marketing	
	surveillance of duloxetine	
	adverse events.	

# Important potential risks

Risk	What is known
Cardiovascular events	Duloxetine has been associated with an increase in blood
including those with	pressure and clinically significant hypertension in some
concomitant use of NSAIDs	patients. This may be due to the noradrenergic effect of
(including myocardial	duloxetine. Cases of hypertensive crisis have been reported
infarction heart failure	with duloxetine, especially in patients with preexisting
andstroke)	hypertension. Therefore, in patients with known
	hypertension and/or other cardiac disease, blood pressure
	monitoring is recommended, especially during the first
	month of treatment. Duloxetine should be used with
	caution in patients whose conditions could be compromised
	by an increased heart rate or by an increase in blood
	pressure. Caution should also be exercised when duloxetine

Risk	What is known
	is used with medicinal products that may impair its metabolism. For patients who experience a sustained increase in blood pressure while receiving duloxetine either dose reduction or gradual discontinuation should be considered. In patients with uncontrolled hypertension duloxetine should not be initiated.
Upper gastrointestinal tract bleeding events with concomitant use of NSAIDs	Duloxetine has been associated with an increase in blood pressure andclinically significant hypertension in some patients. This may be due to the noradrenergic effect of duloxetine. Cases of hypertensive crisishave been reported with duloxetine, especially in patients with preexistinghypertension. Therefore, in patients with known hypertensionand/or other cardiac disease, blood pressure monitoring isrecommended, especially during the first month of treatment.  Duloxetine should be used with caution in patients whose conditionscould be compromised by an increased heart rate or by an increase inblood pressure. Caution should also be exercised when duloxetine issued with medicinal products that may impair its metabolism. Forpatients who experience a sustained increase in blood pressure whilereceiving duloxetine either dose reduction or gradual discontinuationshould be considered. In patients with uncontrolled hypertensionduloxetine should not be initiated.  There have been reports of bleeding abnormalities, such asecchymoses, purpura and gastrointestinal haemorrhage with selective serotonin reuptake inhibitors (SSRIs) and

Risk	What is known
	serotonin/noradrenaline reuptake inhibitors (SNRIs), including duloxetine. Caution is advised in patients taking anticoagulants and/or medicinal products known to affect platelet function (e.g. NSAIDs or acetylsalicylic acid (ASA)), and in patients with known bleeding tendencies.
Renal failure	No dosage adjustment is necessary for patients with mild or moderate renal dysfunction (creatinine clearance 30 to 80 ml/min).  However, increased plasma concentrations of duloxetine occur in patients with severe renal impairment on haemodialysis (creatinineclearance <30 ml/min).  Duloxetine must not be used in patients with severe renal impairment(creatinine clearance <30 ml/min)

# Missing information

Risk	What is known
Characterization of the safety and tolerability of duloxetine in paediatric patients	Duloxetine should not be used in children and adolescents under the age of 18 years for the treatment of major depressive disorder because of safety and efficacy concerns. The safety and efficacy of duloxetine for the treatment of diabetic peripheral neuropathic pain or generalized anxiety disorder have not been studied. No data are available.  Use in children and adolescents under 18 years of age
	Duloxetine should not be used in the treatment of children

Risk	What is known
	and adolescents under the age of 18 years. Suicide-related
	behaviours (suicide attempts and suicidal thoughts), and
	hostility (predominantly aggression, oppositional behaviour
	and anger), were more frequently observed in clinical trials
	among children and adolescents treated with
	antidepressants compared to those treated with placebo. If,
	based on clinical need, a decision to treat is nevertheless
	taken, the patient should be carefully monitored for the
	appearance of suicidal symptoms (please refer to 'clinical
	efficacy and safety' below). In addition, long-term safety
	data in children and adolescents concerning growth,
	maturation and cognitive and behavioural development are
	lacking.
	Adverse events in paediatric population
	A total of 509 paediatric patients aged 7 to 17 years with
	MDD were treated with duloxetine in clinical trials. In
	general, the adversereaction profile of duloxetine in
	children and adolescents was similar to that seen for adults.
	Three hundred and thirty two paediatric patients initially
	randomized to duloxetine in clinical trials experienced a
	0.2 kg mean decrease in weight at 10-weeks. Subsequently,
	over a six-month extension period, most of these patients
	trended toward recovery to their baseline weight percentile
	expected based on population data from age- andgender-
	matched peers.
	Clinical efficacy and safety
	Duloxetine has not been studied in patients under the age

Risk	What is known
	of 7. Two randomized, double-blind, parallel clinical trials were performed in 800 paediatric patients aged 7 to 17 years with major depressive disorder. These two studies included a 10 week placebo and active (fluoxetine) controlled acute phase followed by six months period of active controlled extension treatment. Neither duloxetine (30-120 mg) nor the active control arm (fluoxetine 20-40 mg) statistically separated from placebo on change from baseline to endpoint in the Children's Depression Rating Scale-Revised (CDRS-R) total score. Discontinuation due to adverse events was higher in patients taking duloxetine compared with those treated with fluoxetine, mostly due to nausea. During the 10-week acute treatment period, suicidal behaviours were reported (duloxetine 0/333 [0%], fluoxetine 2/225 [0.9%], placebo 1/220 [0.5%]). Over the entire 36-week course of the study, 6 out of 333 patients initially randomized to duloxetine and 3 out of 225 patients initially randomized to fluoxetine experienced suicidal behaviour (exposure adjusted incidence 0.039 events per patient year for duloxetine, and 0.026 for fluoxetine). In addition, one patient who transitioned from placebo to duloxetine experienced a suicidal behaviour while taking
Description det de la constitue de la constitu	There are no adequate data on the use of dule veting in
Prospective data about potential risks of exposure to duloxetine during pregnancy)	There are no adequate data on the use of duloxetine in pregnant women. Studies in animals have shown reproductive toxicity at systemic exposure levels (AUC) of duloxetine lower than the maximum clinical exposure (please refer to the 'preclinical safety data' below).

Risk	What is known
	The potential risk for humans is unknown.
	Epidemiological data have suggested that the use of SSRIs
	in pregnancy, particularly in late pregnancy, may increase
	the risk of persistent pulmonary hypertension in the
	newborn (PPHN). Although no studies have investigated
	the association of PPHN to SNRI treatment, this potential
	risk cannot be ruled out with duloxetine taking into account
	the related mechanism of action (inhibition of the re-uptake
	of serotonin).
	As with other serotonergic medicinal products,
	discontinuation symptoms may occur in the neonate after
	maternal duloxetine use near term. Discontinuation
	symptoms seen with duloxetine may include hypotonia,
	tremor, jitteriness, feeding difficulty, respiratory distress
	and seizures. The majority of cases have occurred either at
	birth or within a few days of birth.
	Duloxetine should be used in pregnancy only if the
	potential benefit justifies the potential risk to the foetus.
	Women should be advised to notify their physician if they
	become pregnant, or intend to become pregnant, during
	therapy.
	Preclinical safety data
	Duloxetine was not genotoxic in a standard battery of tests
	and was not carcinogenic in rats. Multinucleated cells were
	seen in the liver in the absence of other histopathological
	changes in the rat carcinogenicity study. The underlying
	mechanism and the clinical relevance are unknown. Female

Risk			What is known
			mice receiving duloxetine for 2 years had an increased
			incidence of hepatocellular adenomas and carcinomas at
			the high dose only (144 mg/kg/day), but these were
			considered to be secondary to hepatic microsomal enzyme
			induction. The relevance of this mouse data to humans is
			unknown. Female rats receiving duloxetine (45 mg/kg/day)
			before and during mating and early pregnancy had a
			decrease in maternal food consumption and body weight,
			oestrous cycle disruption, decreased live birth indices and
			progeny survival, and progeny growth retardation at
			systemic exposure levels estimated to be at the most at
			maximum clinical exposure (AUC). In an embryotoxicity
			study in the rabbit, a higher incidence of cardiovascular
			and skeletal malformations was observed at systemic
			exposure levels below the maximum clinical exposure
			(AUC). No malformations were observed in another study
			testing a higher dose of a different salt of duloxetine. In
			prenatal/postnatal toxicity studies in the rat, duloxetine
			induced adverse behaviouraleffects in the offspring at
			exposures below maximum clinical exposure (AUC).
			Studies in juvenile rats reveal transient effects on
			neurobehaviour, as well as significantly decreased body
			weight and food consumption; hepatic enzyme induction;
			and hepatocellular vacuolation at 45 mg/kg/day. The
			general toxicity profile of duloxetine in juvenile rats was
			similar to that in adult rats. The no-adverse effect level was
			determined to be 20 mg/kg/day
Safety of	duloxetine	in	Data on the use of duloxetine 120mg in elderly patients

Risk	What is known	
elderly patients ≥75 years old	with majordepressive disorders are limited. The effect of duloxetine 60 mgonce a day in elderly depressed patients (≥65 years) was specifically examined in a study that showed a statistically significative difference in the reduction of the 17-item Hamilton Depression Rating Scale(HAMD17 score) for duloxetine-treated patients compared to placebo.  Tolerability of duloxetine 60 mg once daily in elderly patients was comparable to that seen in the younger adults. However, data onelderly patients exposed to the maximum dose (120mg per day) are limited and thus, caution is recommended when treating this population.	
Long-term safety data in chronic pain patient	The efficacy of duloxetine used in women for the treatment of moderate to severe Stress Urinary Incontinence (SUI) has not been evaluated for longer than 3 months in placebo-controlled studies.  In addition, the safety and efficacy of duloxetine for the treatment of diabetic peripheral neuropathic pain or generalized anxiety disorder have not been studied. No data are available.	

## VI.2.5 Summary of risk minimisation measures by safety concern

All medicines have a Summary of Product Characteristics (SPC) which provides physicians, pharmacists and other health care professionals with details on how to use the medicine, the risks and recommendations for minimizing them. An abbreviated version of this in lay language is provided in the form of the package leaflet (PL). The measures in these documents are known as routine risk minimisation measures.

This medicine has no additional risk minimisation measures.

## VI.2.6 Planned post authorisation development plan

No studies planned

## VI.2.7 Summary of changes to the risk management plan over time

Version	Date	Safety Concerns	Comment
Version 3.0	10.07.2015	No change in safety concerns	As per Day 120 Draft Assessment Report, RMP has been updated. RMP updated with follow- up targeted questionnaire for collecting prospective data of exposure during pregnancy.

Version 10-04- 2.0 2015	Following safety concerns have been added/corrected  • Important potential risk, Cardiac events expanded with "toinclude those with concomitant use of NSAIDs'  Missing information included following safety concerns:  • Safety of duloxetine in elderly patients ≥75 years old.	As per Day 70 Preliminary Assessment Report of Duloxetine Capsules dated 11 December 2014, safety concerns have been revised	
		<ul> <li>Long-term safety data in chronic pain patient</li> <li>Following safety concerns were removed</li> <li>Elevated plasma concentrations of duloxetine in co-administration with potent CYP1A2 inhibitors</li> </ul>	
		<ul> <li>Serotonin syndrome due to coadministration with nonselective, irreversible monoamine oxidase inhibitors (MAOIs)</li> <li>Hyponatraemia</li> <li>Mydriasis</li> <li>Akathisia/psychomotor restlessness</li> <li>Characterization of drug utilization in unapproved</li> </ul>	