

## **Nodetrip**

Procedural steps taken and scientific information after the authorisation

Application number	Scope	Opinion/ Notification <sup>1</sup> issued cn	Commission Decision Issued <sup>2</sup> / amended on	Product Information affected <sup>3</sup>	Summary
PSUSA/1187/ 202008	Periodic Safety Update EU Single assessment - duloxetine	11/03/2021	n/a		PRAC Recommendation - maintenance
IAIN/0089	A.2.a - Administrative change - Change in the (invented) name of the medicinal product for CAPs	29/09/2020		SmPC, Labelling and	

<sup>&</sup>lt;sup>1</sup> Notifications are issued for type I variations and Article 61(3) notifications (unless part of a group including a type II variation or extension application or a worksharing application). Opinions are issued for all other procedures.



<sup>&</sup>lt;sup>2</sup> A Commission decision (CD) is issued for procedures that affect the terms of the marketing authorisation (e.g. summary of product characteristics, annex II, labelling, package leaflet). The CD is issued within two months of the opinion for variations falling under the scope of Article 23.1a(a) of Regulation (EU) No. 712/2012, or within one year for other procedures.

<sup>&</sup>lt;sup>3</sup> SmPC (Summary of Product Characteristics), Annex II, Labelling, PL (Package Leaflet).

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IB/0088	C.I.z - Changes (Safety/Efficacy) of Human and Veterinary Medicinal Products - Other variation	24/09/2020	SmPC and PL	iseo
WS/1755	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  As agreed in the procedure WS-1527G in order to address the foetal outcomes, submission of the final report from study FIJ-MC-B059 'Observational Study to Assess Fetal Outcomes Following Maternal Exposure to Duloxetine' and the revised final report from study Study F1J-MC-B057 'Observational Studies to Assess Maternal and Fetal Outcomes Following Exposure to Duloxetine'.  Section 4.6 of the SmPc and section 2 of the PL were updated to reflect the available knowledge with regard to the usBe of duloxetine during pregnancy.  The MAH took also the opportunity to include the declaration of sodium in the Product Information following the guideline on 'Excipients in the labelling and package leaflet of medicinal products for human use'.  During the assessment and following a transfer of MAH, Xeristar was removed from the WS procedure.  C.I.13 - Other variations not specifically covered	11/06/2020	SmPC and PL	As agreed in the procedure WS-1527G in order to address the roetal outcomes, submission of the final report from study FIJ-MC-B059 'Observational Study to Assess Fetal Outcomes Following Maternal Exposure to Duloxetine' and the revised final report from study Study F1J-MC-B057 'Observational Studies to Assess Maternal and Fetal Outcomes Following Exposure to Duloxetine'.  Section 4.6 of the SmPc and section 2 of the PL were updated to reflect the available knowledge with regard to the usBe of duloxetine during pregnancy.  The MAH took also the opportunity to include the declaration of sodium in the Product Information following the guideline on 'Excipients in the labelling and package leaflet of medicinal products for human use'.  During the assessment and following a transfer of MAH, Xeristar was removed from the WS procedure.

	elsewhere in this Annex which involve the submission of studies to the competent authority				60
T/0087	Transfer of Marketing Authorisation	30/01/2020	28/02/2020	SmPC, Labelling and PL	volise
IG/1134	B.II.b.2.a - Change to importer, batch release arrangements and quality control testing of the FP - Replacement/addition of a site where batch control/testing takes place	14/08/2019	n/a	नहां अ	thoiised
IG/1126	C.I.z - Changes (Safety/Efficacy) of Human and Veterinary Medicinal Products - Other variation	25/07/2019	23/01/2020	SmPC and PL	
WS/1527/G	This was an application for a group of variations following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  C.I.11.z (Type IB) - to stop enrolment of Study F1J-MC-B034 (study B034), another study included in the current EU-RMP as an additional pharmacovigilance activities to address missing information regarding duloxetine exposure due to pregnancy.  C.I.4 (Type II) - Update of sections 4.4, 4.6 and 4.8 of the SmPC in order to add a warning on the risk of postpartum haemorrhage based on final results from study Study F1J-MC-B057 listed as a category 3 in the RMP; this is an observational study to assess maternal and foetal outcomes following exposure to duloxetine. The Package Leaflet is updated	25/07/2019	23/01/2020	SmPC, Labelling and PL	Observational data have provided evidence of an increased risk (less than 2 -fold) of postpartum haemorrhage following duloxetine exposure within the month prior to birth. This risk is now reflected in the SmPC sections 4.4, 4.6 and 4.8.

	The RMP version 13 has also been submitted. In addition, the Worksharing applicant (WSA) took the opportunity to correct the term "sucraseisomaltase" in section 4.4 of the SmPC in line with the Annex to the EC guideline on 'Excipients in the labelling and package leaflet of medicinal products for human use' (EMA/CHMP/302620/2017 corr. 1*) and to bring the PI in line with the latest QRD template version 10.  The Xeristar 30 mg SmPC & Xeristar 60 mg SmPC and the Yentreve 20 mg SmPC & Yentreve 40 mg SmPC have been combined in a single SmPC, respectively, following the Policy on combined SmPCs (EMA/333423/2015).  C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data C.I.11.z - Introduction of, or change(s) to, the obligations and conditions of a marketing authorisation, including the RMP - Other variation	codinci		OSI OSI	inoiised
WS/1598	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  C.I.3.b - Change(s) in the SPC, Labelling or PL intended to implement the outcome of a procedure concerning PSUR or PASS or the outcome of the	14/06/2019	23/01/2020	SmPC	

	assessment done under A 45/46 - Change(s) with				
	new additional data submitted by the MAH				00
WS/1619	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  B.I.a.1.z - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - Other variation	06/06/2019	n/a		inorised
IG/1055	C.I.z - Changes (Safety/Efficacy) of Human and Veterinary Medicinal Products - Other variation	18/01/2019	23/01/2020	SmPC and PL	
IG/0996	A.7 - Administrative change - Deletion of manufacturing sites	23/10/2018	n/a		
PSUSA/1187/ 201708	Periodic Safety Update EU Single assessment - duloxetine	12/04/2018	n/a		PRAC Recommendation - maintenance
WS/1264	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  Submission of the final report from study F1J-MC-B056 listed as a category 3 study in the RMP. This is a non-interventional non-imposed study aimed to investigate the association between duloxetine exposure and suicide-related behaviours and ideation in women with stress urinary inconsistence (SUI). The RMP version 12.4 has also been updated to	08/02/2018	n/a		The association between suicide attempts and receipt of duloxetine treatment in women with stress urinary inconsistence (SUI) compared to women with SUI without duloxetine treatment has been assessed in study F1J-MC-B056. Study B056 has several limitations but in the light of the results the association between the risk of suicidality and duloxetine treatment cannot be completely ruled out. Currently, the risk of suicidality is an important identified risk for duloxetine-containing products and adequate warnings concerning this risk are already included in the SmPC. No further changes to the product information are

	reflect the study results.  C.I.13 - Other variations not specifically covered elsewhere in this Annex which involve the submission of studies to the competent authority			ā	warranted. The RMP is updated to reflect the study results and limitations and update the pharmacovigilance plan regarding this study.
WS/1331	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  B.II.d.1.h - Change in the specification parameters and/or limits of the finished product - Update of the dossier to comply with the provisions of an updated general monograph of the Ph. Eur. for the finished product	01/02/2018	n/a	ider an	
IG/0759/G	This was an application for a group of variations.  B.III.1.b.2 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability - New certificate for a starting material/reagent/intermediate/or excipient from a new or an already approved manufacturer  B.III.1.b.2 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability - New certificate for a starting material/reagent/intermediate/or excipient from a new or an already approved manufacturer  B.III.1.b.3 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability - Updated certificate from an already approved	11/01/2017	n/a		

	manufacturer B.III.1.b.3 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability - Updated certificate from an already approved manufacturer B.III.1.b.3 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability - Updated certificate from an already approved manufacturer				thoiised
N/0073	Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)	09/12/2016	23/01/2020	Labelling	
WS/1015	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  To update the RMP to add a new Observational Study to Assess Maternal and Fetal Outcomes Following Exposure to Duloxetine (F1J-MC-B057), and to update the plans for the existing pregnancy registry (F1JMC-B034) in section III.4.3 of the RMP.  C.I.11.b - Introduction of, or change(s) to, the obligations and conditions of a marketing authorisation, including the RMP - Implementation of change(s) which require to be further substantiated by new additional data to be submitted by the MAH where significant assessment is required	13/10/2016	n/a O		
IG/0664	B.I.a.1.f - Change in the manufacturer of AS or of a	25/02/2016	n/a		

	starting material/reagent/intermediate for AS - Changes to quality control testing arrangements for the AS -replacement or addition of a site where batch control/testing takes place				thorised
IG/0662	A.1 - Administrative change - Change in the name and/or address of the MAH	23/02/2016	22/07/2016	SmPC, Labelling and PL	IIIO,
WS/0758	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  Update of section 4.8 of the SmPC in order to add microscopic colitis with frequency category 'rare' as a new ADR identified from post marketing experience. The Package Leaflet is updated accordingly. In addition, the Worksharing applicant took the opportunity to make minor editorial changes in the SmPC and PL and to update the local representative for Italy in the Package Leaflet for Xeristar. Moreover, the Worksharing applicant took the opportunity to correct the stated mass of sucrose in capsule in section 2 of the SmPC.  C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data	09/07/2015	22/07/2016	SmPC and PL	
PSUSA/1187/ 201408	Periodic Safety Update EU Single assessment - duloxetine	26/03/2015	27/05/2015	SmPC and PL	Refer to Scientific conclusions and grounds recommending the variation to terms of the Marketing Authorisation(s)' for

				PSUSA/1187/201408.
IG/0472	B.II.d.2.a - Change in test procedure for the finished product - Minor changes to an approved test procedure	18/08/2014	n/a	disec
IG/0457/G	B.III.1.b.2 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability - New certificate for a starting material/reagent/intermediate/or excipient from a new or an already approved manufacturer B.III.1.b.2 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability - New certificate for a starting material/reagent/intermediate/or excipient from a new or an already approved manufacturer B.III.1.b.2 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability - New certificate for a starting material/reagent/intermediate/or excipient from a new or an already approved manufacturer B.III.1.b.2 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability New certificate for a starting material/reagent/intermediate/or excipient from a new or an already approved manufacturer B.III.1.b.2 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability - New certificate for a starting material/reagent/intermediate/or excipient from a deletion of Ph. Eur. TSE Certificate of Suitability - New certificate for a starting material/reagent/intermediate/or excipient from a meterial/reagent/intermediate/or excipient from a new certificate for a starting material/reagent/intermediate/or excipient from a	14/07/2014	n/a	PSUSA/1187/201408.

new or an already approved manufacturer B.III.1.b.2 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability -New certificate for a starting material/reagent/intermediate/or excipient from a new or an already approved manufacturer B.III.1.b.3 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability -Updated certificate from an already approved manufacturer B.III.1.b.3 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability -Updated certificate from an already approved manufacturer B.III.1.b.3 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability -Updated certificate from an already approved manufacturer B.III.1.b.3 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability -Updated certificate from an already approved manufacturer B.III.1.b.3 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability Updated certificate from an already approved manufacturer B.III.1.b.3 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability -Updated certificate from an already approved manufacturer B.III.1.b.3 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability -

	Updated certificate from an already approved manufacturer B.III.1.b.3 - Submission of a new/updated or deletion of Ph. Eur. TSE Certificate of Suitability - Updated certificate from an already approved manufacturer				.hojised
WS/0490	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  Update of sections 4.2, 4.8 and 5.1 of the SmPC following the completion of study HMGI in paediatric patients with generalised anxiety disorder.  C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data	26/06/2014	40/0	SmPC	This variation proposed changes to the duloxetine product information in order to reflect the data obtained from the clinical study conducted in children with a condition called generalised anxiety disorder (GAD). Treatment with duloxetine showed greater improvement in GAD symptoms after 10 weeks of treatment. There was no difference in numbers of patents stopping the treatment due to side effects between duloxetine and placebo (dummy treatment) groups during the 10 week treatment phase.
WS/0513	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  Update of sections 4.2, 4.4 and 5.1 of the SmPC in order to add information from the study HMGF in elderly patients with generalised anxiety disorder (GAD). Minor typographical corrections have been made in the product information.  C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance	20/03/2014	15/10/2014	SmPC	In this variation the Marketing Authorisation Holder updated the Summary of Product Characteristics for Cymbalta and Xeristar to include the additional efficacy and safety information regarding the elderly patients with generalised anxiety disorder following the completion of a clinical study in this population.

	data				8
IG/0383	A.4 - Administrative change - Change in the name and/or address of a manufacturer or an ASMF holder or supplier of the AS, starting material, reagent or intermediate used in the manufacture of the AS or manufacturer of a novel excipient	06/12/2013	n/a		thoiised
WS/0444	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  Update of sections 4.4 and 4.5 of the SmPC to introduce a description of the signs and symptoms of 'serotonin syndrome' and provided an updated list of examples of MAOIs and serotonergic agents. These changes have been proposed by PRAC and endorsed by the CHMP. The Package Leaflet was updated accordingly.  C.I.z - Changes (Safety/Efficacy) of Human and Veterinary Medicinal Products - Other variation	24/10/2013	15/10/2014	SmPC and PL	This variation updated the product information with details of a condition called serotonin syndrome which may occur when medicines acting on the serotonin system in the brain are given concomitantly. Examples of such medicines have been also provided and included in the product information. These changes have been proposed by Pharmacovigilance Risk Assessment Committee (PRAC) and endorsed by the Committee for Medicinal Products for Human Use (CHMP).
IG/0321	C.I.z - Changes (Safety/Efficacy) of Human and Veterinary Medicinal Products - Other variation	17/07/2013	n/a		
WS/0334/G	This was an application for a group of variations following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.	30/05/2013	04/07/2013	SmPC, Annex II and PL	The MAH conducted an updated review of ADR. As a result of this analysis a number of changes to the PI were introduced. The following new ADRs were added to section 4.8 of the SmPC: 'dysphagia', 'testicular pain' (not relevant for YENTREVE) and 'pollakiuria'. The frequency for the ADR

Scope (Ariclaim)

The variations introduced several updates to different sections of the SmPCs of Ariclaim, Cymbalta Xeristar and Yentreve, following the assessment of the data acquired form the performed database analysis of observed ADRs, and the data from the performed two studies in the paediatric population. The changes were specific for the different products, according to the indications they were licensed for. For Ariclaim the updates included:

Update to sections 4.4, 4.8 of the SmPC in order to remove the statement that no clinical trials have been conducted in paediatric population, to add the adverse events (AEs) 'dysphagia' and 'pollakiuria', and to update the description of sensory disturbances as requested by the CHMP further to the assessment of PSUR 11.

Update of section 4.8 of the SmPC to add the AE 'testicular pain' and to reflect the increased reporting frequency of the adverse event 'falls'.

Update to the list of the most commonly reported discontinuation symptoms in Section

4.8c:'Description of selected adverse reactions' to include the term 'myalgia' and update the term 'sensory disturbances (including paraesthesia or electric shock-like sensations, particularly in the head)' as requested by the CHMP further to the assessment of the cumulative review of the cases with "electric shock-like sensations".

Furthermore, the MAH used the opportunity to bring the PI in line with the QRD template version 8.3.

The Package Leaflet was updated accordingly.

'falls' was also updated. Additionally, the MAH updated the list of most commonly reported discontinuation symptoms to include "mya(gia". The term "sensory disturbances (including paresthesia)" was changed to "sensory disturbances (including paresthesia or electric shock-like sensations, particularly in the head)".

Additionally, the data from two completed studies of dufoxetine in paediatric patients with major depressive disorder were available for analysis and resulted in a number of other changes to the PI. The wording of Section 4.2 – Posology and Method of Administration, describing the fact that duloxetine should not be used in children and adolescents was updated to read: "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".

## Scope (Cymbalta)

The variations introduced several updates to

different sections of the SmPCs of Ariclaim, Cymbalta Xeristar and Yentreve, following the assessment of the data acquired form the performed database analysis of observed ADRs, and the data from the performed two studies in the paediatric population. The changes were specific for the different products, according to the indications they were licensed for. For Cymbalta the updates included: Update to sections 4.4, 4.8 of the SmPC in order to remove the statement that no clinical trials have been conducted in paediatric population, to add the adverse events (AEs) 'dysphagia' and 'pollakiuria', and to update the description of sensory disturbances as requested by the CHMP further to the assessment of PSUR 11. Update of section 4.8 of the SmPCs, to add the AE 'testicular pain' and to reflect the increased reporting frequency of the adverse event 'falls'. Update to the list of the most commonly reported discontinuation symptoms in Section 4.8c: Description of selected adverse reactions to include the term 'myalgia' and update the term 'sensory disturbances (including paraesthesia or electric shock-like sensations, particularly in the head)' as requested by the CHMP further to the assessment of the cumulative review of the cases with "electric shock-like sensations". Update of sections 4.2, 4.8 and 5.1 with text reflecting the results of two placebo-controlled trials

with 800 paediatric major depressive disorder (MDD) patients. Furthermore, the MAH used the opportunity to bring the PI in line with the QRD template version 8.3. The Package Leaflet was updated accordingly. Scope (Xeristar) The variations introduced several updates to different sections of the SmPCs of Ariclaim, Cymbalta Xeristar and Yentreve, following the assessment of the data acquired form the performed database analysis of observed ADRs, and the data from the performed two studies in the paediatric population. The changes were specific for the different products, according to the indications they were licensed for. For Xeristar the updates included: Update to sections 4.4, 4.8 of the SmPC in order to remove the statement that no clinical trials have been conducted in paediatric population, to add the adverse events (AEs) 'dysphagia' and 'pollakiuria', and to update the description of sensory disturbances as requested by the CHMP further to the assessment of PSUR 11. Update of section 4.8 of the SmPC, to add the AE 'testicular pain' and to reflect the increased reporting frequency of the adverse event 'falls' Update to the list of the most commonly reported discontinuation symptoms in Section 4.8c:'Description of selected adverse reactions' to include the term 'myalgia' and update the term

'sensory disturbances (including paraesthesia or electric shock-like sensations, particularly in the head)' as requested by the CHMP further to the assessment of the cumulative review of the cases with "electric shock-like sensations".

Update of sections 4.2, 4.8 and 5.1 with the results of two placebo-controlled trials with 800 paediatric major depressive disorder (MDD) patients.

Furthermore, the MAH used the opportunity to bring the PI in line with the QRD template version 8.3.

The Package Leaflet was updated accordingly.

## Scope (Yentreve)

The variations introduced several updates to different sections of the SmPCs of Ariclaim, Cymbalta Xeristar and Yentreve, following the assessment of the data acquired form the performed database analysis of observed ADRs, and the data from the performed two studies in the paediatric population. The changes were specific for the different products, according to the indications they were licensed for. For Yentreve the updates included: Update to sections 4.4, 4.8 of the SmPC in order to remove the statement that no clinical trials have been conducted in paediatric population, to add the adverse events (AEs) 'dysphagia' and 'pollakiuria' and to update the description of sensory disturbances as requested by the CHMP further to the assessment of PSUR 11. Update to the list of the most commonly reported discontinuation symptoms in Section 4.8c: 'Description of selected adverse reactions' to include the term 'myalgia' and update the term 'sensory disturbances (including paraesthesia or

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IG/0239	electric shock-like sensations, particularly in the head)' as requested by the CHMP further to the assessment of the cumulative review of the cases with "electric shock-like sensations".  Furthermore, the MAH used the opportunity to bring the PI in line with the QRD template version 8.3.  The Package Leaflet was updated accordingly.  C.I.3.b - Implementation of change(s) requested following the assessment of an USR, class labelling, a PSUR, RMP, FUM/SO, data submitted under Article 45/46, or amendments to reflect a Core SPC - Change(s) with new additional data submitted by the MAH  C.I.4 - Variations related to significant modifications of the SPC due in particular to new quality, preclinical, clinical or pharmacovigilance data  B.III.2.a.1 - Change of specification('s) of a former	28/11/2012	n/a	oer al	Moiised
16/0239	non Pharmacopoeial substance to comply with the Ph. Eur. or with a national pharmacopoeia of a Member State - AS	28/11/2012	nya		
IG/0178	A.7 - Administrative change - Deletion of manufacturing sites	11/05/2012	n/a		
N/0057	Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)	13/03/2012	04/07/2013	PL	
WS/0181/G	This was an application for a group of variations following a worksharing procedure according to	17/11/2011	n/a		

	Article 20 of Commission Regulation (EC) No 1234/2008.  Changes in testing of the active substance  B.I.b.2.c - Change in test procedure for AS or starting material/reagent/intermediate - Other changes to a test procedure for a reagent, which does not have a significant effect on the overall quality of the AS  B.I.b.2.e - Change in test procedure for AS or starting material/reagent/intermediate - Other changes to a test procedure (including replacement or addition) for the AS or a starting material/intermediate			logi al	Moiised
N/0050	Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)	05/08/2011	n/a	PL	
WS/0135	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  This variation application followed a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008 and concerned:  - Update of the SmPC section 4.4 to include NSAIDS and ASA as examples of antiplatelet agents.  - Update of the SmPC section 4.8 to include terms 'menstrual disorder', 'blood potassium increased', 'dry eye' and 'falls' to the tabulated summary of	23/06/2011	26/07/2011	SmPC and PL	In this variation sections of the product information which provide information on precautions one should take before taking duloxetines were updated with examples of medicines that prevent the blood from clotting, e.g. non steroidal anti-inflammatory drugs and acetylsalicylic acid. New information was also added to the sections of the product information describing possible side effects, for example: menstrual disorder, increase in blood potassium levels, dry eye and falls. Somnolence was added to the list of most commonly reported withdrawal symptoms. It was also mentioned that the patients older than 65 years might experience falls more often. In addition, frequencies of

IB/0049	adverse reactions and to add 'somnolence' to the list of most commonly reported withdrawal symptoms. A footnote "falls were more common in the elderly (more than 65 years old)" was added to 'falls'.  Additionally frequencies of some currently listed ADRs were changed. These updates were based on the most recent CCDS from February 2011.  The Package Leaflet has been updated accordingly. Furthermore, minor editorial changes were also introduced to the product information.  C.I.4 - Variations related to significant modifications of the SPC due in particular to new quality, preclinical, clinical or pharmacovigilance data  The application proposes the introduction of a new	11/03/2011	n/a	loer al	several side effects were updated.
	Pharmacovigilance system, which has been assessed by the EMA for another Lilly product.  C.I.8.b - Introduction of a new Pharmacovigilance system - which has been assessed by the relevant NCA/EMA for another product of the same MAH	roduci			
WS/0071	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  C.I.4 - Variations related to significant modifications of the SPC due in particular to new quality, preclinical, clinical or pharmacovigilance data	16/12/2010	27/01/2011	SmPC, Annex II, Labelling and PL	This variation updates the SmPC section 4.4 with the laboratory measure of the seriousness of low sodium levels in blood and underlines the fact that the elderly are at risk of low sodium levels. The Package Leaflet has been updated accordingly.  Additionally, the contact details for the local representatives in Estonia for the Ariclaim, Cymbalta, Xeristar and Yentreve Package Leaflets have been updated.

IG/0031	A.4 - Administrative change - Change in the name and/or address of a manufacturer or supplier of the AS, starting material, reagent or intermediate used in the manufacture of the AS	17/12/2010	n/a		ised
WS/0011/G	This was an application for a group of variations following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  C.I.z - Changes (Safety/Efficacy) of Human and Veterinary Medicinal Products - Other variation C.I.3.a - Implementation of change(s) requested following the assessment of an USR, class labelling, a PSUR, RMP, FUM/SO, data submitted under A 45/46, or amendments to reflect a Core SPC - Changes with NO new additional data are submitted by the MAH	22/07/2010	06/09/2010	SmPC and PL	This application was submitted for a group of variations consisting of two type 1B variations.  In the variation C.I.z the MAH updated the section 4.6 'Pregnancy and lactation' of the SmPC with symptoms and time to onset of neonatal drug withdrawal syndrome and added galactorrhoea and hyperprolactinaemia to section 4.8 'Undesirable effects' of the SmPC as the result of the assessment of PSUR-9. The Package Leaflet has been updated accordingly.  In the variation C.I.3.a the MAH updated the section 'Pregnancy and lactation' of the Product Information following the class review for SSRIs/SNRIs to inform that when taken during pregnancy SSRI/SNRIs may increase the risk of persistent pulmonary hypertension in neonates. In addition the MAH introduced minor administrative, editorial and linguistic changes to the Product Information.
T/0048	Transfer of Marketing Authorisation	16/04/2010	10/05/2010	SmPC, Labelling and PL	
IB/0046	IB_35_b_Change in weight of coating/capsule shells - gastro-res., modif., prol. release ph. forms	24/11/2009	n/a		
IA/0047	To submit new, updated and unchanged TSE Ph. Eur. certificates of suitability for the gelatine used by the current authorised manufacturer of the capsules and	24/11/2009	n/a		

	for an alternative new suppplier.  IA_22_a_Submission of TSE Ph. Eur. certificate for exc Approved/new manufacturer				ised
II/0036	Extension of indication to include treatment of major depressive disorder.  Extension of Indication	22/10/2009	20/11/2009	SmPC	The CHMP assessment report will be published after deletion of confidential information.
IB/0045	IB_34_b_01_Change in colour/flavour - Increase or addition: colouring system	13/11/2009	n/a	10e/	
II/0044	Update of the Detailed Description of Pharmacovigilance System (DDPS) in the Annex II, including formal notification of a change in the administrative data of the marketing authorisation application form.  Changes to QPPV Update of DDPS (Pharmacovigilance)	23/07/2009	27/08/2009	Annex II	The Detailed Description of the Pharmacovigilance System has been updated (Version 5.2 dated April 2009) to notify changes performed since the last approved version. Consequently, Annex II has been updated with the new version number and date of the agreed DDPS.
II/0041	Update of section 4.5 of the Summary of Product Characteristics to reflect the results of a recent duloxetine/warfarin interaction study (study F1) MC- HMFP).  Furthermore, the term "adolescents" was included in section 4.2 of the SPC in order to align this section with the current QRD template.	29/05/2009	01/07/2009	SmPC	Study F1J-MC-HMFP was an open-label study with the primary objective to evaluate the anticoagulant effects of multiple doses of warfarin when taken at the same time with multiple doses of duloxetine as measured by changes in the international normalized ratio (INR).  Increases in INR values were reported when duloxetine was co-administered with warfarin. However, concomitant administration of duloxetine with warfarin under steady

	Update of Summary of Product Characteristics				state conditions, in healthy volunteers, as part of a clinical pharmacology study, did not result in a clinically significant change in INR from baseline or in the pharmacokinetics of
R/0038	Renewal of the marketing authorisation.	23/04/2009	24/06/2009	SmPC, Annex	warfarin.  Based on the review of the available information the CHMP
				II, Labelling and PL	is of the opinion that the quality, the safety and the efficacy of this medicinal product continues to be adequately and sufficiently demonstrated and therefore considers that the benefit/risk profile of Xeristar continues to be favorable.
				CO,	The MAH will continue to submit a yearly PSUR.
II/0040	to change the finished product specification.	23/04/2009	28/04/2009	(6)	
	Quality changes		.0		
IB/0042	IB_07_c_Replacement/add. of manufacturing site: All other manufacturing operations ex. batch release	27/04/2009	n/a		
IA/0043	IA_32_b_Change in batch size of the finished product - downscaling down to 10-fold	02/04/2009	n/a		
II/0037	Update of Sections 4.8 "Undesirable effects" and 4.9 "Overdose" of the Summary of Product Characteristics (SPC) to reflect the most recent clinical trial data findings of the 7th PSUR. The Package Leaftlet (PL) was updated accordingly.  In addition, this variation implements the outcome of	19/02/2009	25/03/2009	SmPC and PL	A new data lock point for all placebo-controlled clinical studies resulted in a significant increase in the size of the overall database and thus a more robust basis for the determination of Adverse Drug Reactions (ADRs). As a consequence, the frequency of some ADRs was updated in the SPC.
	a recent user testing of the PL of duloxetine- containing products.				Regarding spontaneous data, the MAH identified one new ADR ("restless legs syndrome") as well as new information

	Update of Summary of Product Characteristics and Package Leaflet				on overdose in the most recent PSUR (PSUR 7) submitted in September 2008, and updated the SPC accordingly.  Finally, the MAH has undertaken a user testing of the PL of duloxetine-containing products in 2008 and the results of these were included within this variation.
II/0035	Update of sections 4.2 and 5.1 of the Summary of Product Characteristics (SPC) to reflect new data from a clinical study that investigated the maintenance of effect of duloxetine over 6 months of treatment. The opportunity is also taken to correct some minor typos in the SPC, Labelling and Package Leaflet.  Update of Summary of Product Characteristics, Labelling and Package Leaflet	22/01/2009	26/02/2009	SmPC, Labelling and PL	Study 'HMEM' was designed to investigate the maintenance of effect of duloxetine 60 mg once daily in patients with Diabetic Peripheral Neuropathic Pain (DPNP) who responded to an initial 8 weeks of therapy.  This variation application was submitted in order to update the SPC to reflect the results of study HMEM. The study demonstrated that, for patients who showed an initial response to DPNP therapy with duloxetine, the pain relief observed with duloxetine 60 mg is maintained over a 6-month period.  The variation resulted in the following SPC wording:  Section 4.2: [Diabetic Peripheral Neuropathic Pain Response to treatment should be evaluated after 2 months. In patients with inadequate initial response, additional response after this time is unlikely]  Section 5.1 [In an open label long-term uncontrolled study, the pain reduction in patients responding to 8-weeks of acute treatment of XERISTAR 60 mg once daily was maintained for a further 6-months as measured by change on the Brief

					Pain Inventory (BPI) 24-hour average pain item]
IA/0039	IA_38_a_Change in test procedure of finished product - minor change to approved test procedure	24/02/2009	n/a		ilse0.
IB/0033	IB_13_b_Change in test proc. for active substance - other changes (replacement/addition)	31/07/2008	n/a		"Holl"
II/0027	Extension of indication for Xeristar to include the treatment of generalised anxiety disorder.  Extension of Indication	26/06/2008	28/07/2008	SmPC, Annex II and PL	Please refer to the Scientific Discussion: Xeristar EMEA/H/C/573/II/27.
IB/0034	IB_33_Minor change in the manufacture of the finished product	24/07/2008	n/a		
IB/0032	IB_18_Replacement of an excipient with a comparable excipient	07/07/2008	n/a		
II/0029	Update of Summary of Product Characteristics sections 4.8 and 4.9. The Package Leaflet has been updated.  Update of Summary of Product Characteristics, Labelling and Package Leaflet	19/03/2008	21/04/2008	SmPC, Labelling and PL	The MAH following a search in their in-house clinical trial database as well as the post-marketing data from the spontaneous reporting has applied for changes in the section 4.8 "Undesirable effects" with the inclusion of new adverse drug reactions ("tinnitus", "gait disturbance", "poor quality sleep", "polyuria", "urine flow decreased", "sexual dysfunction" and "dermatitis contact") as well with the modification of the frequency in already known ones. In addition the cases for overdose have been reviewed and the wording has been modified in section 4.9 "Overdose" to include "coma" and "tachycardia" as symptoms of overdosing.

					Changes were implemented to the Product Information
					according to the latest QRD template. The contact details of the representatives of Latvia have been updated.
IA/0030	IA_22_a_Submission of TSE Ph. Eur. certificate for exc Approved/new manufacturer	31/01/2008	n/a		Molis
IB/0028	IB_10_Minor change in the manufacturing process of the active substance	05/12/2007	n/a	0	ithorite
IB/0025	IB_30_b_Change in supplier of packaging components - replacement/addition	30/08/2007	n/a	del	
II/0023	To update section 4.8 of the SPC regarding gastrointestinal bleedings and withdrawal symptoms, section 4.9 of the SPC regarding dosing and as requested by the CHMP and following discussions at the PhVWP to also update the wording on suicidality in section 4.4 of the SPC. The relevant sections of the Package Leaflet are amended accordingly. In addition the contact details of the local representatives of Spain and Denmark have been updated.  Update of Summary of Product Characteristics and Package Leaflet	19/07/2007	28/08/2007	SmPC and PL	Following the PSUR 4 (covering period 3 February 2006 to 2 August 2006) the MAH was requested to update the Product Information with the latest undesirable effects as well as the dosing of the product. In addition and following a meta-analysis published by the FDA regarding the suicidality of the patients administered duloxetine and further to scientific discussions at the PhVWP in June 2007 re-wording of that information has been performed in the section 4.4 of the SPC.
IB/0021	IB_12_b_02_Change in spec. of active subst./agent in manuf. of active subst test parameter	08/06/2007	n/a		
IB/0024	IB_14_b_Change in manuf. of active substance without Ph. Eur. certificate - new manufacturer	24/05/2007	n/a		

IA/0022	IA_11_b_Change in batch size of active substance or intermediate - downscaling	29/03/2007	n/a		volised
N/0019	Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)	08/03/2007	n/a	PL	NOTIS
II/0018	The Marketing Authorisation Holder applied for an update of the Summary of Product Charasteristics (SPC) and the Package Leaflet (PL) following the review of the 3rd PSUR and review of duloxetine placebo-controlled clinical trial database. Sections 4.3, 4.4, 4.5, 4.6, 4.8, and 4.9 of the SPC and sections 2 and 4 of the PL have been amended.  Update of Summary of Product Characteristics and Package Leaflet	18/10/2006	24/11/2006	SmPC and PL	Following the assessment of the third Periodic Safety Update Report (PSUR) the CHMP requested to the MAH to submit a variation to reflect the new safety information. In addition, the MAH also proposed some changes to the SPC following the review of the placebo-controlled clinical trial database. In this variation, the following sections have been updated:  Section 4.3 - Contraindications Addition of contraindication with regards to the initiation of treatment in patients with uncontrolled hypertension.  Section 4.4. Special warning and precautions of use In this variation warnings have been included in this section: -to update information on extrapyramidal disorders -to update information on blood pressure and heart rate.  Section 4.5 Interaction with other medicinal products and other forms of interactions This section was updated to state that "Caution should be exercised when duloxetine is combined with oral anticoagulants or antiplatelets drugs due to an increased risk of bleeding. Furthermore, increases in INR values have

		oduc		Joek al	been reported when duloxetine was co-administered with warfarin". In addition to this, the following information has been included: Duloxetine is an inhibitor of CYP2D6 and therefore caution is advised when duloxetine is co-administered with medicinal products predominantly metabolised by this route (i.e. risperidone, tricyclic antidepressants such as nortriptyline, amitriptyline, and impramine).  Section 4.6 Pregnancy and Lactation Section 4.6 was updated to reflect that duloxetine is very weakly excreted into human milk based on a study of 6 lactating patients.  Section 4.8 Undesirable effects The following Adverse Drug Reactions (ADRs) have been included in the section 4.8: Hypertensive crisis, supraventricular arrhythmia mainly, atrial fibrillation, paresthesia, hepatic failure, trismus, mania. In addition to this, the MAH updated the frequency of ADRs in section 4.8 to reflect the most recent clinical trials data.
II/0017	Change(s) to the manufacturing process for the active substance	27/07/2006	18/08/2006		
II/0014	The Marketing Authorisation Holder (MAH) applied for an update of the Summary of Product Characteristics (SPC) and Package Leaflet (PL) to include new safety information following the review of the 2nd PSUR and results from a pharmacokinetic study in lactating women. Additional changes in the SPC and PL have been made.	27/04/2006	31/05/2006	SmPC, Annex II, Labelling and PL	Following the assessment of the second Periodic Safety Update Report (PSUR) the CHMP identified a number of adverse reactions clinical relevant and which were not yet reflected in the Product Information. Therefore the MAH was requested to submit a variation to reflect the new safety information. In this variation warnings have been included in section 4.4 of the SPC:

	Update of Summary of Product Characteristics, Labelling and Package Leaflet	coduc			<ul> <li>to recommend caution in patients whose conditions could be compromised by an increased heart rate or by an increase in blood pressure         <ul> <li>to update information on withdrawal syndrome seen on discontinuation of treatment</li> <li>to update information on extrapiramidal disorders</li> </ul> </li> <li>Section 4.8 (Undesirable Effects) of the SPC was also updated with regards to withdrawal symptoms and on the effects of duloxetine in Hb1Ac. In addition, the following ADRs have been added to section 4.8: chest pain, seizures, hypertension, hallucinations, akathisia, psychomotor restlessness.</li> <li>Section 4.5 of the SPC (Interaction with other medicinal products) was updated to state that "Increases in INR have been reported when duloxetine was co-administered with warfarin" and to include some examples of drugs metabolised by CYP2D6 with a narrow therapeutic range (such as flecainide, propafenone and metoprolol) in which case caution is advised if Xeristar is co-administered. In this variation the MAH also update sections 4.6 and 5.2 of the SPC to reflect the results of a pharmacokinetic study in lactating women.</li> <li>The Package Leaflet was updated to reflect the above changes.</li> </ul>
IA/0016	IA_41_a_01_Change in pack size change in no. of units within range of appr. pack size	25/04/2006	25/04/2006	SmPC, Labelling and PL	

This variation relates to an update of sections 4.4 23/02/2006 29/03/2006 SmPC and PL The MAH has updated the SPC and PL with safety and 4.8 of the SPC with safety information following assessment of PSUR 1 and a minor addition to section 4.7 concerning the potential for dizziness,  23/02/2006 29/03/2006 SmPC and PL The MAH has updated the SPC and PL with safety information following assessment of the first Pe Update Report (PSUR).  During the assessment of PSUR 1, 3 cases of S	
with consequential changes to the relevant sections of the PL.  Update of Summary of Product Characteristics and Package Leaflet  Package Leaflet  Package Leaflet  Update of Summary of Product Characteristics and Package Leaflet  Package Leafle	IADH ecretion), 7 dium elderly s risk factors sodium, or diuretics). ne SPC on risk of nydrated onatraemia uretic  umber of there are a c drugs that ing, has been of

				Opt al	duloxetine, information is added to section 4.4 of the SPC on that duloxetine is associated with an increase in blood pressure in some patients. In patients with known hypertension and/or other cardiac disease, blood pressure monitoring is recommended as appropriate.  Cases of liver injury, including severe elevations of liver enzymes (>10 times upper limit of normal), hepatitis and iaundice have been reported with duloxetine (see section 4.8). The pattern of liver damage was predominantly hepatocellular. This information was included in the SPC and it was recommended that duloxetine should be used with caution in pa
IB/0013	IB_41_a_02_Change in pack size - change in no. of units outside range of appr. pack size	01/12/2005	01/12/2005	SmPC, Labelling and PL	
A18/0012	Procedure under Article 18 of Council Regulation (EEC) No. 2309/93, as amended, to review suiciderelated behaviours in children and adolescents.  Article 18 Review	23/06/2005	15/09/2005	SmPC and PL	Please refer to Scientific Conclusion Xeristar-EMEA/H/A-18/652.
IB/0010	IB_38_c_Change in test procedure of finished product - other changes	21/07/2005	n/a		
IB/0009	IB_38_c_Change in test procedure of finished product - other changes	21/07/2005	n/a		
II/0005	This variation relates to an update of sections 4.2, 4.4, 5.1 and 5.2 of the SPC with data from a clinical study on the efficacy and safety of duloxetine in the	26/05/2005	04/07/2005	SmPC	The SPC was updated with data on that no dosage adjustment is recommended for elderly patients solely on basis of age, but that caution should be exercised when

	elderly and the very elderly.  Update of Summary of Product Characteristics				treating the elderly, especially with the maximum dose (120 mg/day) for which data are limited. A summary of the results of this study in elderly depressed patients was also added to the SPC.
II/0004	The variation relates to an update of the section 4.1 of the SPC to include the indication "Diabetic Peripheral Neuropathic Pain (DPNP) in adults".  Consequential changes were introduced in sections 4.2, 4.4, 4.8 and 5.1 of the SPC and corresponding section of the PL.  Extension of Indication	26/05/2005	04/07/2005	SmPC and PL	Please refer to Scientific Discussion: Xeristar EMEA/H/C/573/II/04.
IB/0008	IB_42_a_01_Change in shelf-life of finished product - as packaged for sale	22/06/2005	n/a O	SmPC	
IB/0007	IB_41_a_02_Change in pack size - change in no. of units outside range of appr. pack size	20/05/2005	20/05/2005	SmPC, Labelling and PL	
IA/0006	IA_22_a_Submission of TSE Ph. Eur. certificate for exc Approved/new manufacturer	29/03/2005	n/a		
IA/0003	IA_43_a_01_ Add./replacement/del. of measuring or administration device - addition or replacement	28/02/2005	28/02/2005	SmPC, Labelling and PL	
IB/0002	IA_08_a_Change in BR/QC testing - repl./add. of batch control/testing site IB_07_c_Replacement/add. of manufacturing site: All other manufacturing operations ex. batch release	03/02/2005	n/a		

IA/0001	IA_08_b_01_Change in BR/QC testing - repl./add. manuf. responsible for BR - not incl. BC/testing	27/01/2005	n/a	Annex II and PL	60	
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