

## **Icandra**

Procedural steps taken and scientific information after the authorisation

Application number	Scope	Opinion/ Notification  1 issued on	Commission Decision Issued <sup>2</sup> / amended on	Product Information affected <sup>3</sup>	Summary
PSUSA/3113/ 202402	Periodic Safety Update EU Single assessment - vildagliptin, metformin / vildagliptin	03/10/2024	n/a		PRAC Recommendation - maintenance
IG/1771	A.7 - Administrative change - Deletion of manufacturing sites	25/07/2024	n/a		

<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).

WS/2687	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  B.I.b.z - Change in control of the AS - Other variation	16/05/2024	n/a		
IG/1733	B.II.b.1.a - Replacement or addition of a manufacturing site for the FP - Secondary packaging site	27/03/2024	n/a		
IG/1728	B.III.1.a.2 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer	15/03/2024	n/a		
WS/2590	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  B.I.z - Quality change - Active substance - Other variation	18/01/2024	n/a		
IG/1689/G	This was an application for a group of variations.  B.II.b.1.b - Replacement or addition of a manufacturing site for the FP - Primary packaging site  B.II.b.1.a - Replacement or addition of a manufacturing site for the FP - Secondary packaging site  B.II.b.2.c.1 - Change to importer, batch release	04/01/2024		Annex II and PL	

	arrangements and quality control testing of the FP - Replacement or addition of a manufacturer responsible for importation and/or batch release - Not including batch control/testing B.II.b.2.c.1 - Change to importer, batch release arrangements and quality control testing of the FP - Replacement or addition of a manufacturer responsible for importation and/or batch release - Not including batch control/testing B.II.e.2.b - Change in the specification parameters and/or limits of the immediate packaging of the finished product - Addition of a new specification parameter to the specification with its corresponding test method			
WS/2602/0	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.	30/11/2023	n/a	
	B.II.d.2.a - Change in test procedure for the finished product - Minor changes to an approved test procedure  B.II.b.3.a - Change in the manufacturing process of the finished or intermediate product - Minor change in the manufacturing process			
WS/2528/0	This was an application for a group of variations following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.	16/11/2023	n/a	

	C.I.z - Changes (Safety/Efficacy) of Human and Veterinary Medicinal Products - Other variation C.I.z - Changes (Safety/Efficacy) of Human and Veterinary Medicinal Products - Other variation				
IG/1662/G	This was an application for a group of variations.  A.5.b - Administrative change - Change in the name and/or address of a manufacturer/importer of the finished product, including quality control sites (excluding manufacturer for batch release)  A.5.b - Administrative change - Change in the name and/or address of a manufacturer/importer of the finished product, including quality control sites (excluding manufacturer for batch release)	25/08/2023	n/a		
IG/1585	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	01/02/2023	n/a		
IG/1583	B.I.a.1.a - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - The proposed manufacturer is part of the same pharmaceutical group as the currently approved manufacturer	19/12/2022	n/a		
WS/2283/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.	06/10/2022	n/a		

	B.I.a.1.b - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - Introduction of a manufacturer of the AS supported by an ASMF B.I.a.1.f - Change in the manufacturer of AS or of 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			
WS/2251/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.  B.II.b.3.z - Change in the manufacturing process of the finished or intermediate product - Other variation B.II.b.1.e - Replacement or addition of a manufacturing site for the FP - Site where any manufacturing operation(s) take place, except batch-release, batch control, primary and secondary packaging, for non-sterile medicinal products 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	07/07/2022	n/a	
WS/2253	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	07/07/2022	14/10/2022	SmPC and PL

	the new ADRs 'cutaneous vasculitis' with the frequency "not known". Package Leaflet has been updated accordingly.  C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data				
WS/2224	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 update the list of ADRs and update the ADR table in line with the SmPC guideline (a recommendation of EMEA/H/C/WS1970). The Package Leaflet is updated accordingly.  C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data	23/06/2022	14/10/2022	SmPC and PL	For more information, please refer to the Summary of Product Characteristics
N/0102	Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)	04/05/2022	14/10/2022	PL	
WS/2228	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  B.I.b.1.z - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Other variation	24/03/2022	n/a		

WS/2221/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.  B.I.a.1.z - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - Other variation  B.I.a.1.z - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - Other variation  B.I.a.1.z - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - Other variation	17/03/2022	n/a	
N/0099	Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)	01/03/2022	14/10/2022	PL
WS/2111/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.  B.II.b.1.e - Replacement or addition of a manufacturing site for the FP - Site where any manufacturing operation(s) take place, except batch-release, batch control, primary and secondary packaging, for non-sterile medicinal products  B.II.b.3.a - Change in the manufacturing process of the finished or intermediate product - Minor change in the manufacturing process	13/01/2022	n/a	

B.II.b.4.a - Change in the batch size ranges) of the finished prod compared to the originally appro B.II.b.2.a - Change to importer, arrangements and quality control Replacement/addition of a site w control/testing takes place	uct - Up to 10-fold oved batch size batch release ol testing of the FP -		
WS/2128/G  This was an application for a gro following a worksharing procedul Article 20 of Commission Regular 1234/2008.  B.II.d.1.g - Change in the specificand/or limits of the finished proceducty of a specification parameter (excluding biological product) of a specification parameter corresponding test method as a quality issue  B.II.b.2.a - Change to importer, arrangements and quality control Replacement/addition of a site we control/testing takes place	re according to tion (EC) No  ication parameters duct - Addition or al or immunological meter wit its result of a safety or  batch release of testing of the FP -	n/a	
IG/1442/G  This was an application for a ground B.I.b.2.b - Change in test proceds starting material/reagent/interm a test procedure for the AS or a material/reagent/intermediate, it procedure is already authorised B.I.b.2.b - Change in test procedure.	dure for AS or nediate - Deletion of starting f an alternative test	n/a	

IG/1445	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	20/10/2021	n/a	
N/0091	Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)	12/10/2021	14/10/2022	PL
WS/1970	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  Submission of an updated RMP (version 15.2) in order to bring it in line with revision 2 of GVP module V on 'Risk management systems' and aligned with the conclusions of the PSUR single assessment (PSUSA) procedure (PSUSA/00003113/201802) adopted in October 2018. Annex II.D of the product information is updated to remove the statement around submission of an RMP update every 3 years.  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	30/09/2021	14/10/2022	Annex II
IG/1435	A.4 - Administrative change - Change in the name and/or address of a manufacturer or an ASMF holder	26/08/2021	n/a	

	or supplier of the AS, starting material, reagent or intermediate used in the manufacture of the AS or manufacturer of a novel excipient				
WS/1937/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.	20/05/2021	05/07/2021	SmPC, Annex II and PL	Please refer to Scientific Discussion  `Eucreas_Icandra_Zomarist-H-C-WS-1937-G'
	Update of sections 4.1, 4.4, 4.5, 5.1 and 6.6 of the SmPC to change the existing indication with regards to the use in combination with other diabetes medicines, to reflect the VERIFY study data (on initial combination of vildagliptin with metformin) and expand existing warning on drugs that may affect renal function or metformin disposition by including drugs that inhibit renal transporter (OCT2/MATE inhibitors) and corresponding update in drug interactions. PI update to QRD v10.1. The Package Leaflet and Annex II are updated in accordance.				
	C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data C.I.6.a - Change(s) to therapeutic indication(s) - Addition of a new therapeutic indication or modification of an approved one				

IG/1398	B.III.1.a.2 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer	13/05/2021	n/a		
IG/1377	B.III.1.a.2 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer	26/03/2021	n/a		
IG/1354	B.II.e.6.b - Change in any part of the (primary) packaging material not in contact with the finished product formulation - Change that does not affect the product information	12/02/2021	n/a		
IG/1316/G	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  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	16/12/2020	n/a		
IG/1222/G	This was an application for a group of variations.  B.II.e.1.a.1 - Change in immediate packaging of the finished product - Qualitative and quantitative	22/05/2020	30/09/2020	SmPC, Labelling and PL	

composition - Solid pharmaceutical forms B.II.e.2.b - Change in the specification parameters and/or limits of the immediate packaging of the finished product - Addition of a new specification parameter to the specification with its corresponding test method B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g.

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	tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes			
IG/1201	B.III.1.a.2 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer	31/01/2020	n/a	
IG/1173	B.III.1.a.2 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer	04/12/2019	n/a	
IG/1129/G	This was an application for a group of variations.  A.7 - Administrative change - Deletion of manufacturing sites  B.II.b.1.a - Replacement or addition of a manufacturing site for the FP - Secondary packaging site  B.II.b.1.a - Replacement or addition of a manufacturing site for the FP - Secondary packaging site  B.II.b.2.c.1 - Change to importer, batch release	20/09/2019	30/09/2020	Annex II and PL

	arrangements and quality control testing of the FP - Replacement or addition of a manufacturer responsible for importation and/or batch release - Not including batch control/testing			
WS/1658/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.	12/09/2019	n/a	
	B.I.a.1.f - Change in the manufacturer of AS or of 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 B.I.a.1.f - Change in the manufacturer of AS or of 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 B.I.a.1.f - Change in the manufacturer of AS or of 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 B.I.a.1.z - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - Other			
	variation  B.I.a.1.z - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - Other variation  B.I.a.3.a - Change in batch size (including batch size			

	ranges) of AS or intermediate - Up to 10-fold increase compared to the originally approved batch size  B.I.a.3.a - Change in batch size (including batch size ranges) of AS or intermediate - Up to 10-fold increase compared to the originally approved batch size  B.I.b.2.z - Change in test procedure for AS or starting material/reagent/intermediate - Other variation  B.I.b.2.z - Change in test procedure for AS or starting material/reagent/intermediate - Other variation  B.I.b.2.z - Change in test procedure for AS or starting material/reagent/intermediate - Other variation  B.I.b.2.z - Change in test procedure for AS or starting material/reagent/intermediate - Other variation  B.I.c.1.a - Change in immediate packaging of the AS - Qualitative and/or quantitative composition			
IG/1102	B.III.1.a.2 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer	17/05/2019	n/a	
N/0076	Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)	22/02/2019	30/09/2020	PL
WS/1513	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.	17/01/2019	n/a	
	B.I.a.1.z - Change in the manufacturer of AS or of a			

	starting material/reagent/intermediate for AS - Other variation				
IG/1000/G	B.I.a.1.f - Change in the manufacturer of AS or of 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 B.II.c.1.a - Change in the specification parameters and/or limits of an excipient - Tightening of specification limits B.III.1.a.2 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer	22/11/2018	n/a		
N/0073	Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)	07/11/2018	30/09/2020	PL	
PSUSA/3113/ 201802	Periodic Safety Update EU Single assessment - vildagliptin, metformin / vildagliptin	04/10/2018	n/a		PRAC Recommendation - maintenance
IG/0942	B.I.b.1.c - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Addition of a new specification parameter to the specification with its corresponding test method	26/06/2018	n/a		
IG/0943	B.I.a.1.f - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS -	01/06/2018	n/a		

	Changes to quality control testing arrangements for the AS -replacement or addition of a site where batch control/testing takes place			
IG/0928/G	This was an application for a group of variations.  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  B.I.a.2.a - Changes in the manufacturing process of the AS - Minor change in the manufacturing process of the AS	03/05/2018	n/a	
IG/0920/G	This was an application for a group of variations.  B.I.a.1.f - Change in the manufacturer of AS or of 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  B.I.a.1.f - Change in the manufacturer of AS or of 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  B.I.a.1.f - Change in the manufacturer of AS or of 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	26/04/2018	n/a	

	B.I.a.1.f - Change in the manufacturer of AS or of 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			
T/0067	Transfer of Marketing Authorisation	26/03/2018	12/04/2018	SmPC, Labelling and PL
IG/0895	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	21/02/2018	n/a	
IG/0843/G	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 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 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	04/10/2017	n/a	

	manufacturer of a novel excipient  A.7 - Administrative change - Deletion of manufacturing sites			
WS/1184	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  B.II.b.3.z - Change in the manufacturing process of	14/09/2017	n/a	
	the finished or intermediate product - Other variation			
IG/0797/G	This was an application for a group of variations.  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  A.5.b - Administrative change - Change in the name and/or address of a manufacturer/importer of the finished product, including quality control sites (excluding manufacturer for batch release)  A.5.b - Administrative change - Change in the name and/or address of a manufacturer/importer of the finished product, including quality control sites (excluding manufacturer for batch release)  B.I.a.1.f - Change in the manufacturer of AS or of 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	12/05/2017	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 B.III.1.a.2 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer				
WS/1072	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 5.1 of the SmPC, subsection 'cardiovascular risk', with results from a new meta-analysis evaluating the cardiovascular safety of vildagliptin. In addition, the Worksharing applicant (WSA) took the opportunity to bring the annexes in line with the latest QRD template version 10, and to merge the two SmPCs into one single SmPC for Eucreas, Icandra and Zomarist. Moreover, the section on pregnancy and breast-feeding in the PL for Eucreas/Icandra/Zomarist has been aligned with the wording used for Galvus/Jalra/Xiliarx.  C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data	21/04/2017	19/02/2018	SmPC, Annex II, Labelling and PL	A meta-analysis of independently and prospectively adjudicated cardiovascular events from 37 phase III and IV monotherapy and combination therapy clinical studies of up to more than 2 years duration (mean exposure 50 weeks for vildagliptin and 49 weeks for comparators) was performed and showed that vildagliptin treatment was not associated with an increase in cardiovascular risk versus comparators. The composite endpoint of adjudicated major adverse cardiovascular events (MACE) including acute myocardial infarction, stroke or cardiovascular death was similar for vildagliptin versus combined active and placebo comparators [Mantel–Haenszel risk ratio (M-H RR) 0.82 (95% CI 0.61 1.11)]. A MACE occurred in 83 out of 9,599 (0.86%) vildagliptin-treated patients and in 85 out of 7,102 (1.20%) comparator-treated patients. Assessment of each individual MACE component showed no increased risk (similar M-H RR). Confirmed heart failure (HF) events defined as HF requiring hospitalisation or new onset of HF were reported in 41 (0.43%) vildagliptin-treated patients and 32 (0.45%) comparator-treated patients with M-H RR 1.08 (95% CI 0.68 1.70).
WS/1088	This was an application for a variation following a	21/04/2017	n/a		

	worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  C.I.11.z - Introduction of, or change(s) to, the obligations and conditions of a marketing authorisation, including the RMP - Other variation				
IG/0786	C.I.z - Changes (Safety/Efficacy) of Human and Veterinary Medicinal Products - Other variation	09/03/2017	19/02/2018	SmPC	
WS/1094/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.  B.I.a.2.a - Changes in the manufacturing process of the AS - Minor change in the manufacturing process of the AS B.I.a.2.a - Changes in the manufacturing process of the AS - Minor change in the manufacturing process of the AS - Minor change in the manufacturing process of the AS B.I.a.4.c - Change to in-process tests or limits applied during the manufacture of the AS - Deletion of a non-significant in-process test	16/02/2017	n/a		
A31/0056	Pursuant to Article 31 of Regulation (EC) No 726/2004, the European Commission requested on 25 January 2016 the opinion of the European Medicines Agency on the adequacy of the current recommendations for metformin containing products with respect to the use in patients with moderate renal failure, taking into account the available	13/10/2016	08/12/2016	SmPC and PL	Please refer to the assessment report:  Metformin containing medicinal products - EMEA/H/A- 31/1432

	information on the risk of lactic acidosis. The CHMP was requested to assess the impact thereof on the benefit-risk balance of metformin containing products and to give its recommendation whether the marketing authorisation of this product should be maintained, varied, suspended or revoked. The notification for the procedure is appended to this opinion.			
N/0057	Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)	21/03/2016		PL
WS/0905/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.  B.I.b.1.b - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Tightening of specification limits  B.I.b.1.c - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Addition of a new specification parameter to the specification with its corresponding test method  B.I.b.2.a - Change in test procedure for AS or starting material/reagent/intermediate - Minor changes to an approved test procedure  B.I.b.2.e - Change in test procedure for AS or	10/03/2016	n/a	

	starting material/reagent/intermediate - Other changes to a test procedure (including replacement or addition) for the AS or a starting material/intermediate			
WS/0887/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.  B.III.1.a.2 - Submission of a new/updated or	11/02/2016	n/a	
	deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer  B.III.1.a.2 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer  B.I.d.1.a.4 - Stability of AS - Change in the re-test period/storage period - Extension or introduction of a re-test period/storage period supported by real time data			
WS/0883/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.	04/02/2016	n/a	
	<ul><li>A.7 - Administrative change - Deletion of manufacturing sites</li><li>B.II.b.1.a - Replacement or addition of a manufacturing site for the FP - Secondary packaging</li></ul>			

	site  B.II.b.1.b - Replacement or addition of a manufacturing site for the FP - Primary packaging site  B.II.b.3.z - Change in the manufacturing process of the finished or intermediate product - Other variation				
WS/0791	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  C.I.13 - Other variations not specifically covered elsewhere in this Annex which involve the submission of studies to the competent authority	28/01/2016	n/a		
PSUSA/3113/ 201502	Periodic Safety Update EU Single assessment - vildagliptin, metformin / vildagliptin	22/10/2015	16/12/2015	SmPC	Refer to Scientific conclusions and grounds recommending the variation to terms of the Marketing Authorisation(s)' for PSUSA/3113/201502.
IG/0637	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	02/12/2015	n/a		
WS/0835/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.  B.II.b.3.a - Change in the manufacturing process of	12/11/2015	n/a		

	in the manufacturing process B.II.b.5.b - Change to in-process tests or limits applied during the manufacture of the finished product - Addition of a new test(s) and limits				
IG/0566/G	This was an application for a group of variations.  B.I.b.2.a - Change in test procedure for AS or starting material/reagent/intermediate - Minor changes to an approved test procedure  B.I.b.2.a - Change in test procedure for AS or starting material/reagent/intermediate - Minor changes to an approved test procedure  B.I.d.1.a.1 - Stability of AS - Change in the re-test period/storage period - Reduction	02/06/2015	n/a		
WS/0722	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 line with a PRAC recommendation dated 4 Dec 2014, in order to add the ADR 'myalgia' with frequency "not known". The Package Leaflet has been updated accordingly. In addition, the MAH took the opportunity to update the contact details of the local representative in Spain in the Package Leaflet for Xiliarx and Icandra.  C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data	21/05/2015	16/12/2015	SmPC and PL	N/A

WS/0724/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.  A.1 - Administrative change - Change in the name and/or address of the MAH B.II.f.1.b.1 - Stability of FP - Extension of the shelf life of the finished product - As packaged for sale (supported by real time data)	23/04/2015	16/12/2015	SmPC, Labelling and PL
WS/0697/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.  B.II.d.3 - Variations related to the introduction of real-time release or parametric release in the manufacture of the finished product B.II.d.3 - Variations related to the introduction of real-time release or parametric release in the manufacture of the finished product B.II.d.3 - Variations related to the introduction of real-time release or parametric release in the manufacture of the finished product B.II.d.3 - Variations related to the introduction of real-time release or parametric release in the manufacture of the finished product B.II.d.2.d - Change in test procedure for the finished product - Other changes to a test procedure (including replacement or addition)	26/02/2015	n/a	
WS/0695	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.	26/02/2015	n/a	

	To updated the RMP to version 12.1 following the change to the due date of the final CSR report for Study CLAF237A2401 from 'Q4 2014' to 'Q2 2015'.  C.I.11.z - Introduction of, or change(s) to, the obligations and conditions of a marketing authorisation, including the RMP - Other variation			
IG/0506/G	B.III.1.a.2 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer B.III.1.a.2 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer B.III.1.a.2 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer A.7 - Administrative change - Deletion of manufacturing sites 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	28/11/2014	n/a	

IG/0504	B.II.b.1.a - Replacement or addition of a manufacturing site for the FP - Secondary packaging site	12/11/2014	n/a	
WS/0584/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.	24/07/2014	n/a	
	This was an application for a group of variations following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.			
	B.I.b.1.c - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Addition of a new specification parameter to the specification with its corresponding test method B.I.b.1.c - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Addition of a new specification parameter to the specification with its			
	corresponding test method B.I.b.1.c - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Addition of a new specification parameter to the specification with its corresponding test method B.I.b.1.c - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Addition of a new			

specification parameter to the specification with its
corresponding test method
B.I.b.1.c - Change in the specification parameters
and/or limits of an AS, starting
material/intermediate/reagent - Addition of a new
specification parameter to the specification with its
corresponding test method
B.I.b.1.c - Change in the specification parameters
and/or limits of an AS, starting
material/intermediate/reagent - Addition of a new
specification parameter to the specification with its
corresponding test method
B.I.b.1.c - Change in the specification parameters
and/or limits of an AS, starting
material/intermediate/reagent - Addition of a new
specification parameter to the specification with its
corresponding test method
B.I.b.1.c - Change in the specification parameters
and/or limits of an AS, starting
material/intermediate/reagent - Addition of a new
specification parameter to the specification with its
corresponding test method
B.I.b.1.c - Change in the specification parameters
and/or limits of an AS, starting
material/intermediate/reagent - Addition of a new
specification parameter to the specification with its
corresponding test method
B.I.b.1.c - Change in the specification parameters
and/or limits of an AS, starting
material/intermediate/reagent - Addition of a new
specification parameter to the specification with its
corresponding test method
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B.I.b.1.c - Change in the specification parameters
and/or limits of an AS, starting
material/intermediate/reagent - Addition of a new
specification parameter to the specification with its
corresponding test method
B.I.b.1.b - Change in the specification parameters
and/or limits of an AS, starting
material/intermediate/reagent - Tightening of
specification limits
B.I.b.1.b - Change in the specification parameters
and/or limits of an AS, starting
material/intermediate/reagent - Tightening of
specification limits
B.I.b.1.b - Change in the specification parameters
and/or limits of an AS, starting
material/intermediate/reagent - Tightening of
specification limits
B.I.b.1.b - Change in the specification parameters
and/or limits of an AS, starting
material/intermediate/reagent - Tightening of
specification limits
B.I.b.2.a - Change in test procedure for AS or
starting material/reagent/intermediate - Minor
changes to an approved test procedure
B.I.b.1.c - Change in the specification parameters
and/or limits of an AS, starting
material/intermediate/reagent - Addition of a new
specification parameter to the specification with its
corresponding test method
B.I.b.1.c - Change in the specification parameters
and/or limits of an AS, starting
material/intermediate/reagent - Addition of a new

IG/0460	specification parameter to the specification with its corresponding test method B.I.b.1.c - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Addition of a new specification parameter to the specification with its corresponding test method  B.II.b.1.a - Replacement or addition of a manufacturing site for the FP - Secondary packaging site	14/07/2014	n/a		
WS/0518	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 4.4 of the SmPC to update the safety information on acute pancreatitis. The Package Leaflet was updated accordingly.  C.I.z - Changes (Safety/Efficacy) of Human and Veterinary Medicinal Products - Other variation	25/04/2014	13/04/2015	SmPC, Labelling and PL	The MAH has provided information on clinical studies that may provide further safety data with regards to pancreatic safety in patients treated with vildagliptin. In the absence of a CV outcome study, the VERIFY study is discussed. This study is relatively small (2000 patients) when it comes to investigating uncommon events but instead of long duration (5 years). Adverse events will be captured with routine methods in the study with specific analyses of pancreatic events which is considered sufficient. Timelines have been provided and are acceptable.  The MAH has also provided information on the only observational study currently ongoing and planned. The study protocol already includes pancreatitis as an outcome of interest and the protocol has been amended with pancreatic cancer. The updated protocol is included in the updated RMP.  The benefit/risk balance for Galvus/Eucreas remains positive.
IG/0398/G	This was an application for a group of variations.	24/01/2014	n/a		

	B.I.b.2.a - Change in test procedure for AS or starting material/reagent/intermediate - Minor changes to an approved test procedure B.I.b.1.c - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Addition of a new specification parameter to the specification with its corresponding test method			
IAIN/0037/G	This was an application for a group of variations.  B.I.b.1.c - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Addition of a new specification parameter to the specification with its corresponding test method  B.I.b.1.c - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Addition of a new specification parameter to the specification with its corresponding test method  B.III.1.a.2 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer  B.III.1.a.3 - Submission of a new/updated or deletion of Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - New certificate from a new manufacturer (replacement or addition)	03/12/2013	n/a	

14/0/02/50		27/26/2015	05/00/00/5	0.00.4	
WS/0358	This was an application for a variation following a	27/06/2013	05/08/2013	SmPC, Annex	With the current variation new clinical data in patients with
	worksharing procedure according to Article 20 of			II and PL	T2DM and CHF NYHA class I-III has been provided. Based
	Commission Regulation (EC) No 1234/2008.				on previously available data, the use of vildagliptin in
					patients with CHF in NYHA class I-II has been accepted,
	Update of sections 4.4 and 5.1 of the SmPC in order				whereas the use of vildagliptin in patients with CHF in NYHA
	to include data in patients with type 2 diabetes				class III-IV has not been recommended due to lack of data.
	mellitus and congestive heart failure NYHA class I-				The number of patients in NYHA class III treated with
	III. The Package Leaflet is updated accordingly.				vildagliptin is still limited (47) and the data indicate that
	In addition, the MAH took the opportunity to update				there were imbalances with regards to background
	the list of local representatives in the Package				morbidity, thus the data has to be interpreted with caution
	Leaflet.				and the data are insufficient to make claims on the efficacy
					and safety of vildagliptin in this patient group.
	Furthermore, the PI is being brought in line with the				In the study submitted, the primary objective was to
	latest version of the QRD template.				investigate the effect of vildagliptin on cardiac safety with
					change in LVEF being the primary outcome. The data on
	C.I.4 - Variations related to significant modifications				LVEF, as well as on worsening of CHF does not indicate any
	of the SPC due in particular to new quality, pre-				adverse effect of vildagliptin on cardiac function and there
	clinical, clinical or pharmacovigilance data				is no mechanistic/biological rationale for a negative effect.
					There were, however, imbalances in AE reporting with more
					SAEs and deaths observed in the vildagliptin treated NYHA
					class III population. This was most likely related to the
					background morbidity. The safety data from the study is in
					line with the known safety profile of vildagliptin. However,
					data in patients with CHF NYHA III is still limited and this
					issue will be continuously monitored in future PSURs which
					is endorsed.
					The outcome with regards to HbA1c, responder analysis
					and FPG in the overall study population (CHF, NYHA class I-
					III) was in line with that observed in previous studies with
					vildagliptin although the effect was less prominent at week
					52. In patient with CHF, NYHA class III, the HbA1c was
					modest (-0.3 %), however, this subgroup of patients was

R/0035	Renewal of the marketing authorisation.	30/05/2013	31/07/2013		small and there are indications of heterogeneity in the data which may explain the modest reduction in HbA1c. This was reflected in the study description in section 5.1 of the SmPC.  Due to the limited number of patients, the data provided is still deemed insufficient to decide on the overall benefit risk balance of vildagliptin in patients with CHF NYHA class III. The submitted study, which focused on changes in LVEF in patients treated with vildagliptin, however, provides sufficient data to allow a cautious use in these patients support that vildagliptin has no negative effect on cardiac function and there is no mechanistic/biological rationale for such an effect.  The benefit-risk balance for vildagliptin containing products remains positive.  Based upon the data that have become available since the granting of the initial Marketing Authorisation, the CHMP considers that the benefit-risk balance of Icandra remains positive. The CHMP recommends that the renewal be granted with unlimited validity.
IA/0036/G	This was an application for a group of variations.  B.II.b.4.a - Change in the batch size (including batch size ranges) of the finished product - Up to 10-fold compared to the currently approved batch size  B.II.b.4.a - Change in the batch size (including batch size ranges) of the finished product - Up to 10-fold compared to the currently approved batch size	25/06/2013	n/a		
IB/0031/G	This was an application for a group of variations.	08/02/2013	31/07/2013	SmPC and	

B.II.e.1.z - Change in immediate packaging of the
finished product - Other variation
B.II.e.5.a.1 - Change in pack size of the finished
product - Change in the number of units (e.g.
tablets, ampoules, etc.) in a pack - Change within
the range of the currently approved pack sizes
B.II.e.5.a.1 - Change in pack size of the finished
product - Change in the number of units (e.g.
tablets, ampoules, etc.) in a pack - Change within
the range of the currently approved pack sizes
B.II.e.5.a.1 - Change in pack size of the finished
product - Change in the number of units (e.g.
tablets, ampoules, etc.) in a pack - Change within
the range of the currently approved pack sizes
B.II.e.5.a.1 - Change in pack size of the finished
product - Change in the number of units (e.g.
tablets, ampoules, etc.) in a pack - Change within
the range of the currently approved pack sizes
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product - Change in the number of units (e.g.
tablets, ampoules, etc.) in a pack - Change within
the range of the currently approved pack sizes
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product - Change in the number of units (e.g.
tablets, ampoules, etc.) in a pack - Change within
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product - Change in the number of units (e.g.
tablets, ampoules, etc.) in a pack - Change within
the range of the currently approved pack sizes
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product - Change in the number of units (e.g.

tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g.

	tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes B.II.e.5.a.1 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change within the range of the currently approved pack sizes				
WS/0345/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.  - To add a manufacturer of a starting material To delete two manufacturing sites for a starting material.  B.I.a.1.z - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - Other variation A.7 - Administrative change - Deletion of manufacturing sites A.7 - Administrative change - Deletion of manufacturing sites	17/01/2013	17/01/2013		
IG/0248	C.I.z - Changes (Safety/Efficacy) of Human and Veterinary Medicinal Products - Other variation	17/12/2012	n/a		
IB/0033/G	This was an application for a group of variations.  B.I.b.1.z - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Other variation	14/12/2012	n/a		

	B.I.b.2.a - Change in test procedure for AS or starting material/reagent/intermediate - Minor changes to an approved test procedure				
WS/0330	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.	15/11/2012	n/a		
	To add an additional manufacturing site of the finished product.				
	B.II.b.1.e - Replacement or addition of a manufacturing site for the FP - Site where any manufacturing operation(s) take place, except batch-release, batch control, primary and secondary packaging, for non-sterile medicinal products				
WS/0272	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.	20/09/2012	29/10/2012	SmPC and PL	For further information please refer to the scientific conclusion: H-XXX-WS-0272-AR
	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  Extension of indication for the use of vildagliptin and				
	vildagliptin/metformin in triple therapy with a sulphonylurea and metformin affecting sections 4.1, 4.2, 4.4, 4.8 and 5.1 of the SmPC. The Package				
	Leaflet was proposed to be updated in accordance. In addition, the MAH took the opportunity to update the list of local representatives in the Package Leaflet.				
	C.I.6.a - Change(s) to therapeutic indication(s) -				

	Addition of a new therapeutic indication or modification of an approved one.  C.I.6.a - Change(s) to therapeutic indication(s) - Addition of a new therapeutic indication or modification of an approved one				
WS/0257	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  Extension of indication for use of vildagliptin and vildagliptin/metformin in combination with insulin affecting sections 4.1, 4.2, 4.8 and 5.1 of the SmPC. The Package Leaflet is updated accordingly.  C.I.6.a - Change(s) to therapeutic indication(s) - Addition of a new therapeutic indication or modification of an approved one.  C.I.6.a - Change(s) to therapeutic indication(s) - Addition of a new therapeutic indication or modification of an approved one	20/09/2012	29/10/2012	SmPC and PL	For further information please refer to the scientific conclusion: H-XXX-WS-0257-AR
IG/0209/G	This was an application for a group of variations.  C.I.9.b - Changes to an existing pharmacovigilance system as described in the DDPS - Change in the contact details of the QPPV  C.I.9.h - Changes to an existing pharmacovigilance	17/08/2012	n/a		

	system as described in the DDPS - Other change(s) to the DDPS that does not impact on the operation of the pharmacovigilance system				
WS/0256	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  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 4.8 of the SmPC in order to add a warning of bullous or exfoliative skin lesions following the review of the Galvus PSUR 7. The Package Leaflet is updated accordingly.  C.I.3.z - 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 - Other variation.  C.I.3.z - 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 - Other variation.	24/05/2012	28/06/2012	SmPC and PL	At the time of granting marketing license, skin lesions were considered a potential risk based on pre-clinical findings in monkeys. In the assessment of the PSUR No 7 of Galvus, 19 post marketing reports of blistering dermatitis, bullous rash pemphigus were identified. Of these, a causal association to vildagliptin was suspected in 9 cases. To further evaluate the possible risk of skin adverse reactions associated with vildagliptin, the MAH was requested to submit a thorough evaluation of all relevant cases. This evaluation concluded that "bullous or exfoliative lesions" should be added to the description of the post-marketing experience of sections 4.4 and 4.8 of the SmPC.
IG/0148/G	This was an application for a group of variations.  C.I.9.e - Changes to an existing pharmacovigilance system as described in the DDPS - Changes in the	22/02/2012	n/a		

	major contractual arrangements with other persons or organisations involved in the fulfilment of pharmacovigilance obligations and described in the DD C.I.9.h - Changes to an existing pharmacovigilance system as described in the DDPS - Other change(s) to the DDPS that does not impact on the operation of the pharmacovigilance system			
IG/0143/G	A.7 - Administrative change - Deletion of manufacturing sites  B.I.a.1.f - Change in the manufacturer of AS or of 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  B.I.a.2.a - Changes in the manufacturing process of the AS - Minor change in the manufacturing process of the AS  B.I.b.1.b - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Tightening of specification limits  B.II.b.2.a - Change to batch release arrangements and quality control testing of the FP - Replacement or addition of a site where batch control/testing takes place  B.III.1.a.2 - Submission of a new or updated Ph. Eur. Certificate of Suitability to the relevant Ph. Eur.	20/12/2011	n/a	

	approved manufacturer				
IB/0022	B.I.b.2.z - Change in test procedure for AS or starting material/reagent/intermediate - Other variation	13/12/2011	n/a		
WS/0125	This was an application for a variation following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  Update of product information with data on liver dysfunction received by the MAH from marketed use of vildagliptin. In addition, the MAH took the opportunity to update the list of local representatives in the Package Leaflet.  C.I.4 - Variations related to significant modifications of the SPC due in particular to new quality, preclinical, clinical or pharmacovigilance data	20/10/2011	24/11/2011	SmPC and PL	The clinical safety database for vildagliptin consists of pooled analysis of data from 38 clinical trials including more than 11,500 patients treated with vildagliptin. Furthermore, post-marketing data are available from more than 1.24 million patient years.  Data from clinical trials, although not statistically significant, indicate a slightly increased risk of persistent transaminase elevation in patients treated with vildagliptin relative to the comparators. This risk is reflected in the current labeling. Following a request by the CHMP (EMA/549257/2010), the MAH has conducted an evaluation of all reports related to liver dysfunction received from marketed use of vildagliptin (including Galvus and Eucreas) since the original placing on the market.  The cumulative post-marketing experience has identified seven cases consistent with a drug-related liver event and a further 15 cases where a causal association cannot be excluded. Based on this, the CHMP considered that the safety information referring to the post-marketing experience in section 4.8 should be updated to reflect this safety concern.  Additionally, the MAH agreed to the CHMP request that adverse liver events should be reviewed again in future

					PSURs.
IG/0118/G	This was an application for a group of variations.  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  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  A.7 - Administrative change - Deletion of manufacturing sites  B.I.a.3.a - Change in batch size (including batch size ranges) of AS or intermediate - Up to 10-fold increase compared to the currently approved batch size  B.I.b.1.c - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Addition of a new specification parameter to the specification with its corresponding test method  B.I.b.2.a - Change in test procedure for AS or starting material/reagent/intermediate - Minor changes to an approved test procedure	27/10/2011	n/a		
WS/0164	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 4.4 of the SPC and relevant	22/09/2011	24/10/2011	SmPC and PL	Following the review of the latest vildagliptin PSUR, the CHMP requested the addition to the SmPC of a warning statement on acute pancreatitis including information that patients should be informed on the characteristic symptoms of acute pancreatitis, that resolution has been observed

IR/0012	section of the PL to include a warning on pancreatitis, following assessment of the latest PSUR.  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  B.I. a. 1. 7 Change in the manufacturer of AS or of a	20/07/2011	n/a	after discontinuation and that treatment should not be resumed after pancreatitis has been diagnosed.  Few cases of pancreatitis were reported in the clinical trials safety data base and there was no apparent imbalance between vildagliptin and comparators/placebo. In five of the cases the pancreatitis occurred in the context of cholelithiasis or cholecystitis and all the four cases in which vildagliptin was re-initiated belonged to this group. That is, all the rechallenged cases had an easily identifiable risk factor for pancreatitis present and all four rechallenges were negative.  The post-marketing data presented with the current variation covers the period up to 30 April 2011. The cut-off date for the latest PSUR was 30 Nov 2010, thus the current report covers an additional time period of five months.  Overall, the analysis of post-marketing events consistent with pancreatitis in vildagliptin agents revealed 42 cases of pancreatitis/acute pancreatitis and 15 cases of reported elevated lipase/amylase. The majority of reported cases had additional risk factors for pancreatitis and that patients should be informed on the characteristic symptoms of acute pancreatitis has been introduced in the product information, in line with the current warnings given for other DPP4-inhibitors.  The data provided with this variation is not considered to alter the benefit risk balance for vildagliptin, which remains positive.
IB/0012	B.I.a.1.z - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - Other variation	20/07/2011	n/a	

IG/0088/G	This was an application for a group of variations.  C.I.9.e - Changes to an existing pharmacovigilance system as described in the DDPS - Changes in the major contractual arrangements with other persons or organisations involved in the fulfilment of pharmacovigilance obligations and described in the DD  C.I.9.h - Changes to an existing pharmacovigilance system as described in the DDPS - Other change(s) to the DDPS that does not impact on the operation of the pharmacovigilance system	11/07/2011	n/a		
IG/0057/G	This was an application for a group of variations.  B.II.b.1.a - Replacement or addition of a manufacturing site for the FP - Secondary packaging site  B.II.b.1.b - Replacement or addition of a manufacturing site for the FP - Primary packaging site	28/03/2011	n/a		
WS/0070	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 4.2 of the SPC following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008, to reflect the safety and efficacy data that the MAH has accumulated in patients > 75 years. The MAH also	16/12/2010	27/01/2011	SmPC, Annex II and PL	During the review of the original marketing authorization application for vildagliptin, there was only a limited number of patients > 75 years who had been treated with vildagliptin (N=113) in clinical studies. Since then, the number of elderly patients ? 75 years treated with vildagliptin has increased. In the enlarged database, 334 patients > 75 years were exposed to vildagliptin, 295 of which are included in the main safety population and up to 187 in the efficacy populations. The majority of the patients

	takes this opportunity to implement the latest QRD template, to update Annex IIB with the new RMP version and to update some local representatives' details in the PL.  C.I.4 - Variations related to significant modifications of the SPC due in particular to new quality, preclinical, clinical or pharmacovigilance data			were treated with the approved dose 50 mg bid.  The submitted analyses support a similar efficacy in elderly patients compared to the younger population with respect to reduction of HbA1c and FPG. The treatment was largely weight neutral and the incidence of hypoglycaemia was low. No other safety issues compared to comparators or the younger population were identified.  There are limited safety data on the use of vildagliptin in patients with moderately and severely impaired renal function at risk for higher drug exposure. Until data from ongoing studies in patients with moderate and severe renal impairment become available, the use of vildagliptin is not recommended in this population. This would also apply for the elderly patient population > 75 years.
IG/0032/G	This was an application for a group of variations.  To update the Detailed Description of the Pharmacovigilance System (DDPS) to version 9.0, to include:  - a change in the deputy of the Qualified Person for Pharmacovigilance (QPPV);  - a change in the major contractual arrangements.  - administrative changes not impacting the operation of the pharmacovigilance system.  Annex II.B has also been updated with the latest wording as per October 2010 CHMP procedural announcement.  C.I.9.c - Changes to an existing pharmacovigilance system as described in the DDPS - Change of the back-up procedure of the QPPV	21/12/2010	n/a	

	C.I.9.e - Changes to an existing pharmacovigilance system as described in the DDPS - Changes in the major contractual arrangements with other persons or organisations involved in the fulfilment of pharmacovigilance obligations and described in the DD C.I.9.h - Changes to an existing pharmacovigilance system as described in the DDPS - Other change(s) to the DDPS that does not impact on the operation of the pharmacovigilance system				
WS/0005/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.  This was an application for a group of variations following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  Section 5.1 of the Summary of Products Characteristics (SPC) has been updated to include	20/05/2010	05/07/2010	SmPC, Annex II and PL	The MAH has provided results from two clinical studies (LAF237A2338 and LMF237A2302) and proposed to include information from these studies in section 5.1 of the Summary of Products Characteristics (SPC).  The first one was a multicenter, randomized, double-blind, active-controlled study to compare the efficacy and safety of long-term treatment (52 weeks) with vildagliptin to gliclazide in patients with T2DM inadequately controlled with metformin monotherapy. The presented results from this study are in line with what has been seen in previous studies with vildagliptin in combination with sulfonylurea.
	information on 2 new vildagliptin studies.  Additionally, the adverse event pancreatitis has been added to section 4.8 of the SPC.  C.I.4 - Variations related to significant modifications of the SPC due in particular to new quality, preclinical, clinical or pharmacovigilance data  C.I.3.b - Implementation of change(s) requested following the assessment of an USR, class labelling, a				The second study was a 24-week multicenter, randomized, double-blind, active-controlled initial combination therapy study with the fixed dose combination of vildagliptin plus metformin in drug naïve patients with T2DM. The results of this study demonstrated that initial combination therapy with the fixed dose combination of vildagliptin plus metformin was statistically superior to that of both individual monotherapy components. Initial combination

	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				therapy is not recommended as first line treatment in treatment guidelines and is not in line with the approved indication for vildagliptin. However, the CHMP accepted that information on this study was added to section 5.1 of the SPC.  A new pooled safety analysis has been performed which integrates data from these new studies and provides an updated assessment of the safety and tolerability of vildagliptin based on data from more than 11,500 patients. The increased database was also utilized to conduct an extensive analysis of the cardiovascular safety of vildagliptin. The CHMP agreed that this information is consistent with the previously submitted pooled safety data and does not signal any new, unidentified concerns.  Additionally, section 4.8 of the SPC and relevant section of the Package Leaflet have been updated to include pancreatitis as a post-marketing adverse event.
IB/0011	B.II.b.1.e - Replacement or addition of a manufacturing site for the FP - Site where any manufacturing operation(s) take place, except batch-release, batch control, primary and secondary packaging, for non-sterile medicinal products	29/06/2010	n/a		
IA/0010	B.III.1.a.2 - Submission of a new or updated Ph. Eur. Certificate of Suitability to the relevant Ph. Eur. Monograph - Updated certificate from an already approved manufacturer	08/06/2010	n/a		
II/0009	Update of the Detailed Description of the	18/02/2010	15/03/2010	Annex II and	With this variation the MAH submitted a new version of the

	Pharmacovigilance system (DDPS).  Changes to QPPV Update of DDPS (Pharmacovigilance)			PL	DDPS (core version 8.0) in accordance with the current Pharmacovigilance guideline. After assessing the documentation the CHMP concluded that the submitted DDPS contained all required elements. Consequently, Annex II has been updated with the new version number of the agreed DDPS.  In addition, the Marketing Authorisation Holder took the opportunity to update the local representatives contact details for Finland, Latvia and Luxembourg in the Package Leaflet and to introduce some linguistic changes in the German and Romanian Annexes.
IA/0007	IA_08_a_Change in BR/QC testing - repl./add. of batch control/testing site	15/12/2009	n/a		
IA/0006	IA_07_a_Replacement/add. of manufacturing site: Secondary packaging site IA_07_b_01_Replacement/add. of manufacturing site: Primary packaging site - Solid forms	15/12/2009	n/a		
IA/0008	IA_15_b_02_Submission of Ph. Eur. certificate for active substance - new manuf./other substances	10/12/2009	n/a		
II/0003	Update of SPC sections 4.8 and 5.1 and PL to include information on the results of 3 active-controlled long-term clinical studies.  Update of Summary of Product Characteristics and Package Leaflet	23/07/2009	31/08/2009	SmPC and PL	During the review of the initial Marketing Authorisation application for vildagliptin, the Marketing Authorisation Holder committed to submit the results of ongoing, active-controlled, long-term, comparative studies. This included one 2-year monotherapy study, one ? 2-year add-on combination therapy study with metformin, and one 1-year (24 week + 28 week extension) add-on combination therapy study with metformin.

					The results of these studies have now become available and are discussed in the current variation application.  Furthermore, a review of the safety of vildagliptin has been performed, integrating data from the above studies of up to more than 2 years in duration as well as from additional studies.  Concerning efficacy, non-inferiority to the comparators (glimepiride and pioglitazone), according to the prespecified non-inferiority margins, was achieved in the addon to metformin studies, but not in the monotherapy study. In all studies the absolute reduction of HbA1c with vildagliptin was lower compared to the comparators.  Concerning safety, vildagliptin has a hypoglycemia profile superior to that of glimepiride and similar to that of pioglitazone and metformin. Previously, signals concerning hepatic and skin safety have been identified. In the current updated safety data set, slightly higher odds of having a persistent transaminase elevation with vildagliptin were found. Concerning skin safety, the incidences of skin/vascular AEs were low and not statistically significant.  No additional safety signals were seen.
N/0004	Update of the list of local representatives in section 6 of the Package Leaflet.  Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)	04/08/2009	n/a	PL	
II/0002	To change the testing monographs of the finished	29/05/2009	19/06/2009		

	product.				
	Change(s) to the test method(s) and/or specifications for the finished product				
IB/0001	IB_02_Change in the name of the medicinal product	06/02/2009	n/a	SmPC, Annex II, Labelling and PL	