



Adempas

Procedural steps taken and scientific information after the authorisation

| Application number | Scope | Opinion/ Notification ¹ issued on | Commission Decision Issued ² / amended on | Product Information affected ³ | Summary |
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| PSUSA/10174 /201809 | Periodic Safety Update EU Single assessment - riociguat | 11/04/2019 | n/a | | PRAC Recommendation - maintenance |
| II/0028 | Update of sections 4.2, 4.4 and 4.5 of the SmPC to include information for recommended starting dose for riociguat for patients who are on stable doses of strong multi pathway cytochrome P450 proteins (CYP) and P-gp/BCRP inhibitors based on data from Study 17957 which investigated the potential pharmacokinetic (PK) | 28/02/2019 | | SmPC and PL | Study 17957 investigated the effect of fixed-dose antiretroviral therapies, Atripla, Complera, Stribild, Triumeq, or any approved antiretroviral protease inhibitor in combination with (preferably) Triumeq, on the exposure to riociguat in HIV patients on a stable dose of one of these therapies. Comparison with limited historical data indicated |

¹ 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.

² 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.

³ SmPC (Summary of Product Characteristics), Annex II, Labelling, PL (Package Leaflet).



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| | <p>interaction of human immunodeficiency virus (HIV) antiretroviral agents as fixed-dose combination and riociguat in HIV patients, data from a statistical drug-drug interaction (DDI) which was evaluated in study 18634, in which PK data from study 17957 was compared to the historical PK data and data from a nonclinical study to elucidate the DDI potential of the different components included in the HIV combination products in vitro.</p> <p>C.1.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data</p> | | | | <p>no effect of Atripla on the AUC of riociguat, while HIV combined therapy including ritonavir boosted protease inhibitors, Complera, Stribild and Triumeq increased the exposure by 18, 88, 88 and 159%, respectively. In vitro data showed that the increase is mainly due to inhibition of CYP1A1, and partial due to inhibition of CYP3A4. The plasma concentrations in all HIV non-PH patients were in the range of the observed plasma riociguat concentrations in healthy subjects and healthy subjects receiving riociguat on top of ketoconazole. Taking into account the observed safety (albeit after a single dose) and vital signs, it is considered that, with application of the reduced starting dose of 0.5 mg here times a day, prescribers can manage this risk of hypotension caused by riociguat. Sections 4.2, 4.4 and 4.5 of the SmPC have been updated as a consequence. The Package leaflet Labelling has been updated accordingly.</p> |
| R/0026 | Renewal of the marketing authorisation. | 15/11/2018 | 18/01/2019 | SmPC, Labelling and PL | Based on the review of data on quality, safety and efficacy, the CHMP considered that the benefit-risk balance of Adempas in the approved indication remains favourable and therefore recommended the renewal of the marketing authorisation with unlimited validity. |
| PSUSA/10174 /201709 | Periodic Safety Update EU Single assessment - riociguat | 12/04/2018 | n/a | | PRAC Recommendation - maintenance |
| II/0023 | Update of section 4.2 of the SmPC in order to add new information regarding posology for transitioning to and from riociguat based on results from study 16719: An open-label, international, multicentre, single-arm, uncontrolled, phase IIIb study of riociguat in patients with pulmonary arterial hypertension (PAH) who demonstrate an insufficient response to treatment | 25/01/2018 | 07/03/2018 | SmPC and PL | The following recommendations with regards to transitioning between sildenafil or tadalafil and Adempas were made: Patients stopping sildenafil must wait at least 24 hours before taking Adempas. Patients stopping tadalafil must wait at least 48 hours before taking Adempas. Patients stopping Adempas to change to another medicine called a PDE5 inhibitor (e.g. sildenafil or tadalafil) must wait at least 24 |

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| | <p>with phosphodiesterase-5 inhibitors (PDE-5i). Section 4.5 of the SmPC was updated in parallel to reflect on the main study results concerning pharmacodynamic interactions. Minor editorial changes were also implemented throughout the SmPC.</p> <p>The Package Leaflet was updated accordingly.</p> <p>C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data</p> | | | | <p>hours from their last dose of Adempas before taking the PDE5 inhibitor. Signs and symptoms of hypotension should be monitored after any transition.</p> |
| II/0024/G | <p>This was an application for a group of variations.</p> <p>C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data</p> <p>C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data</p> <p>C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data</p> | 12/10/2017 | 07/03/2018 | SmPC | |
| T/0022 | Transfer of Marketing Authorisation | 07/04/2017 | 02/05/2017 | SmPC, Labelling and PL | |
| PSUSA/10174 /201609 | Periodic Safety Update EU Single assessment - riociguat | 06/04/2017 | n/a | | PRAC Recommendation - maintenance |
| II/0018/G | This was an application for a group of variations. | 30/03/2017 | n/a | | |

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| | C.I.13 - Other variations not specifically covered elsewhere in this Annex which involve the submission of studies to the competent authority C.I.13 - Other variations not specifically covered elsewhere in this Annex which involve the submission of studies to the competent authority | | | | |
| IAIN/0021/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.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.a - Administrative change - Change in the name and/or address of a manufacturer/importer responsible for batch release | 14/03/2017 | 02/05/2017 | Annex II and PL | |
| II/0014 | 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 | 23/02/2017 | n/a | | |
| II/0019 | Update of section 4.5 of the SmPC in order to add information about interactions of riociguat when | 26/01/2017 | 02/05/2017 | SmPC, Annex II, Labelling | The CHMP considered that results of an interaction study indicated that ethinyl estradiol and levonorgestrel exposure |

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| | <p>administered concomitantly with combined oral contraceptives containing levonorgestrel and ethinyl estradiol to healthy female subjects. Section 4.4 of the SmPC was updated to reinforce the existing messages in sections 4.3 and 4.6 with regards to pregnancy. Furthermore, section 4.5 of the SmPC was updated to correct the list of CYP isoforms involved in the metabolism of riociguat based on in vitro data. In addition, the Marketing authorisation holder (MAH) took the opportunity to bring the Product Information in line with the latest QRD template version 10.0 and to update the contact details of the German local representative.</p> <p>C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data</p> | | | and PL | <p>was not affected when administered on top of a treatment with riociguat. The following text has been added to section 4.5 of the SmPC:</p> <p>“Patients must not get pregnant during Adempas therapy (see section 4.3). Riociguat (2.5 mg three times per day) did not have a clinically meaningful effect on the plasma levels of combined oral contraceptives containing levonorgestrel and ethinyl estradiol when concomitantly administered to healthy female subjects. Based on this study and as riociguat is not an inducer of any of the relevant metabolic enzymes, also no pharmacokinetic interaction is expected with other hormonal contraceptives.”</p> <p>Furthermore, the existing messages conveyed in sections 4.3 and 4.6 with regards to pregnancy have been reinforced in section 4.4 of the SmPC as follows:</p> <p>“Pregnancy/contraception Adempas is contraindicated during pregnancy (see section 4.3). Therefore, female patients at potential risk of pregnancy must use an effective method of contraception. Monthly pregnancy tests are recommended.”</p> |
| IB/0017 | B.I.a.3.z - Change in batch size (including batch size ranges) of AS or intermediate - Other variation | 11/11/2016 | n/a | | |
| IA/0015/G | <p>This was an application for a group of variations.</p> <p>B.I.b.1.b - Change in the specification parameters and/or limits of an AS, starting material/intermediate/reagent - Tightening of specification limits</p> <p>B.I.b.2.c - Change in test procedure for AS or starting material/reagent/intermediate - Other changes to a</p> | 27/10/2016 | n/a | | |

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| | test procedure for a reagent, which does not have a significant effect on the overall quality of the AS | | | | |
| IA/0016/G | <p>This was an application for a group of variations.</p> <p>B.II.c.1.a - Change in the specification parameters and/or limits of an excipient - Tightening of specification limits</p> <p>B.II.c.1.a - Change in the specification parameters and/or limits of an excipient - Tightening of specification limits</p> <p>B.II.c.1.a - Change in the specification parameters and/or limits of an excipient - Tightening of specification limits</p> <p>B.II.c.1.b - Change in the specification parameters and/or limits of an excipient - Addition of a new specification parameter to the specification with its corresponding test method</p> <p>B.II.c.1.c - Change in the specification parameters and/or limits of an excipient - Deletion of a non-significant specification parameter (e.g. deletion of an obsolete parameter)</p> <p>B.II.c.1.c - Change in the specification parameters and/or limits of an excipient - Deletion of a non-significant specification parameter (e.g. deletion of an obsolete parameter)</p> <p>B.II.c.1.c - Change in the specification parameters and/or limits of an excipient - Deletion of a non-significant specification parameter (e.g. deletion of an obsolete parameter)</p> <p>B.II.c.1.c - Change in the specification parameters and/or limits of an excipient - Deletion of a</p> | 26/10/2016 | n/a | | |

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| | non-significant specification parameter (e.g. deletion of an obsolete parameter) | | | | |
| PSUSA/10174 /201603 | Periodic Safety Update EU Single assessment - riociguat | 29/09/2016 | n/a | | PRAC Recommendation - maintenance |
| IAIN/0013 | C.I.z - Changes (Safety/Efficacy) of Human and Veterinary Medicinal Products - Other variation | 03/08/2016 | 22/08/2016 | SmPC and PL | |
| PSUSA/10174 /201509 | Periodic Safety Update EU Single assessment - riociguat | 14/04/2016 | n/a | | PRAC Recommendation - maintenance |
| N/0009 | Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification) | 23/11/2015 | 22/08/2016 | PL | |
| PSUSA/10174 /201503 | Periodic Safety Update EU Single assessment - riociguat | 08/10/2015 | n/a | | PRAC Recommendation - maintenance |
| II/0007/G | This was an application for a group of variations. C.I.z - Changes (Safety/Efficacy) of Human and Veterinary Medicinal Products - Other variation C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data | 23/07/2015 | 15/10/2015 | SmPC, Labelling and PL | |
| PSUSA/10174 /201409 | Periodic Safety Update EU Single assessment - riociguat | 10/04/2015 | n/a | | PRAC Recommendation - maintenance |
| II/0006 | C.I.13 - Other variations not specifically covered elsewhere in this Annex which involve the submission of studies to the competent authority | 26/02/2015 | n/a | | |

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| IB/0004 | B.I.a.1.z - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - Other variation | 11/12/2014 | n/a | | |
| II/0001 | <p>Submission of non-clinical study report R-9318, an in vitro study undertaken to determine the M-1 potential to inhibit renal efflux transporters MATE1 and MATE2-K. The study was included as a category 3 study in the RMP, and a revised RMP version 3.0 was provided accordingly.</p> <p>C.I.13 - Other variations not specifically covered elsewhere in this Annex which involve the submission of studies to the competent authority</p> | 23/10/2014 | n/a | | <p>The applicant provided the requested information regarding the inhibition potential of M-1 towards MATE1 and MATE2-K uptake transporters. Based on the data from these in vitro studies, at clinical relevant concentrations of M-1, no drug-drug interactions due to inhibition of MATEs by M-1 are expected. Therefore, no changes to the product information are necessary.</p> <p>The study was included as a category 3 study in the RMP, and a revised RMP version 3.0 was provided accordingly. The revised RMP version 3.0 is agreed.</p> <p>These study results do not influence the benefit / risk balance of riociguat, which remains unchanged in the authorised indication(s).</p> |
| IB/0003/G | <p>This was an application for a group of variations.</p> <p>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</p> <p>B.II.e.5.a.2 - Change in pack size of the finished product - Change in the number of units (e.g. tablets, ampoules, etc.) in a pack - Change outside the range of the currently approved pack sizes</p> <p>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</p> | 14/10/2014 | 15/10/2015 | SmPC, Labelling and PL | |

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| | <p>the currently approved pack sizes</p> <p>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</p> <p>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</p> | | | | |
| PSUV/0002 | Periodic Safety Update | 09/10/2014 | n/a | | PRAC Recommendation - maintenance |