

Rozlytrek

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
R/0015	Renewal of the marketing authorisation.	26/04/2023	07/07/2023	SmPC	The CHMP, having reviewed the available information on the status of the fulfilment of Specific Obligations and having confirmed the positive benefit risk balance, is of the opinion that the quality, safety and efficacy of this medicinal product continue to be adequately and sufficiently demonstrated and therefore recommends the

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.

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



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.

PSUSA/10874 /202212	Periodic Safety Update EU Single assessment - entrectinib	06/07/2023	n/a		renewal of the conditional MA for Rozlytrek, subject to the Specific Obligations and Conditions as laid down in Annex II to the opinion. PRAC Recommendation - maintenance
IA/0018	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)	29/06/2023	n/a		
II/0014	Update of sections 4.2 and 5.2 of the SmPC in order to update the pharmacokinetic information based on final results from study GP411174 listed as an additional pharmacovigilance activity in the RMP; this is a Phase I, non-randomized, single-dose, openlabel study to investigate the effect of impaired hepatic function on the pharmacokinetics of entrectinib in volunteers with different levels of hepatic function. The RMP version 4.1 has also been agreed. In addition, the MAH took the opportunity to update in Annex II section C and to update the list of local representatives in the Package Leaflet. C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data	26/01/2023	09/03/2023	SmPC, Annex II and PL	The pharmacokinetics of entrectinib were studied in subjects with mild (Child-Pugh A), moderate (Child-Pugh B) and severe (Child-Pugh C) hepatic impairment, relative to subjects with normal hepatic function. Following administration of a single oral dose of 100 mg entrectinib, the combined AUClast of entrectinib and M5 showed no relevant change in the hepatic impaired groups compared to the normal function group. The AUClast geometric mean ratio (90% CI) was 1.30 (0.889, 1.89) for the mild, 1.24 (0.886, 1.73) for the moderate, and 1.39 (0.988, 1.95) for the severe hepatic impaired groups compared to the normal hepatic function group. For the unbound entrectinib and M5, the AUClast (fu) geometric mean ratio (90% CI) was 1.91 (1.21, 3.02) for the mild, 1.57 (1.06, 2.31) for the moderate, and 2.34 (1.57, 3.48) for the severe hepatic impaired groups compared to the normal hepatic function group. Although the effect of hepatic impairment on unbound PK parameters generally followed a similar direction as total PK parameters, due to the high non-specific binding in buffer and high variability, results should

					be interpreted with caution. No dose adjustment is recommended for patients with mild, moderate or severe hepatic impairment. Patients with severe hepatic impairment should be carefully monitored for hepatic function and adverse reactions. For more information, please refer to the Summary of Product Characteristics.
PSUSA/10874 /202206	Periodic Safety Update EU Single assessment - entrectinib	12/01/2023	n/a		PRAC Recommendation - maintenance
II/0012	Submission of the final integrated analysis report for cardiac risks, listed as a category 3 study in the RMP, in order to fulfil MEA/003. This is an integrated safety analysis report to assess cardiac risks based on GO40782 [STARTRK-2], CO40778 [STARTRK-NG], and BO41932 [TAPISTRY] studies (PAESs). Section 4.4 of the SmPC has been updated to better reflect the cardiac safety findings. C.I.13 - Other variations not specifically covered elsewhere in this Annex which involve the submission of studies to the competent authority	08/09/2022	09/03/2023	SmPC	Congestive heart failure (CHF) has been reported in less than 5% of patients across clinical trials with Rozlytrek. These reactions were observed in patients with or without a history of cardiac disease and resolved in 70% of those patients upon institution of appropriate clinical management and/or Rozlytrek dose reduction/interruption. For more information, please refer to the Summary of Product Characteristics.
PSUSA/10874 /202112	Periodic Safety Update EU Single assessment - entrectinib	07/07/2022	n/a		PRAC Recommendation - maintenance
IB/0011/G	This was an application for a group of variations. B.II.f.1.d - Stability of FP - Change in storage conditions of the finished product or the diluted/reconstituted product	27/06/2022	09/03/2023	SmPC, Labelling and PL	

	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)				
II/0010	Submission of the final report from study (RO7102122) to address the non-clinical recommendation issued within the initial MAA. This is an in-vitro study for the evaluation of entrectinib against novel clinically-relevant NTRK fusions using the Ba/F3 cell line. C.I.13 - Other variations not specifically covered elsewhere in this Annex which involve the submission of studies to the competent authority	02/06/2022	n/a		
R/0007	Renewal of the marketing authorisation.	24/03/2022	30/05/2022	SmPC and PL	The CHMP, having reviewed the available information on the status of the fulfilment of Specific Obligations and having confirmed the positive benefit risk balance, is of the opinion that the quality, safety and efficacy of this medicinal product continue to be adequately and sufficiently demonstrated and therefore recommends the renewal of the conditional MA for Rozlytrek, subject to the Specific Obligations and Conditions as laid down in Annex II to the Opinion.
IB/0009/G	This was an application for a group of variations. 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 B.II.b.2.a - Change to importer, batch release	05/04/2022	n/a		

	arrangements and quality control testing of the FP - Replacement/addition of a site where batch control/testing takes place				
PSUSA/10874 /202106	Periodic Safety Update EU Single assessment - entrectinib	13/01/2022	n/a		PRAC Recommendation - maintenance
IB/0005/G	This was an application for a group of variations. A.6 - Administrative change - Change in ATC Code/ATC Vet Code 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)	10/08/2021	30/05/2022	SmPC and PL	
PSUSA/10874 /202012	Periodic Safety Update EU Single assessment - entrectinib	08/07/2021	n/a		PRAC Recommendation - maintenance
R/0002	Renewal of the marketing authorisation.	22/04/2021	21/06/2021		
IB/0004/G	This was an application for a group of variations. B.II.d.1.z - Change in the specification parameters and/or limits of the finished product - Other variation 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 B.I.b.2.c - Change in test procedure for AS or starting material/reagent/intermediate - Other	31/05/2021	n/a		

	st procedure for a reagent, which
does not have a	a significant effect on the overall
quality of the A	S
B.I.b.1.z - Char	nge in the specification parameters
and/or limits of	an AS, starting
material/interm	nediate/reagent - Other variation
B.I.b.1.c - Char	nge in the specification parameters
and/or limits of	an AS, starting
material/interm	nediate/reagent - Addition of a new
specification pa	rameter to the specification with its
corresponding t	test method
B.I.b.1.b - Cha	nge in the specification parameters
and/or limits of	an AS, starting
material/interm	nediate/reagent - Tightening of
specification lin	nits
B.I.a.4.z - Char	nge to in-process tests or limits
applied during	the manufacture of the AS - Other
variation	
B.I.a.4.c - Char	nge to in-process tests or limits
applied during	the manufacture of the AS - Deletion
of a non-signifi	cant in-process test
B.I.a.4.a - Chai	nge to in-process tests or limits
applied during	the manufacture of the AS -
Tightening of ir	n-process limits
B.I.a.3.a - Chai	nge in batch size (including batch size
ranges) of AS of	or intermediate - Up to 10-fold
increase compa	red to the originally approved batch
size	
B.I.a.2.a - Chai	nges in the manufacturing process of
the AS - Minor	change in the manufacturing process
of the AS	
B.I.a.1.a - Chai	nge 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 B.I.a.1.z - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - Other variation				
IB/0001	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)	15/10/2020	21/06/2021	SmPC, Labelling and PL	