

22 May 2015
EMA/COMP/435101/2014 Rev.2
Committee for Orphan Medicinal Products

Public summary of opinion on orphan designation

Recombinant fusion protein consisting of a modified form of the extracellular domain of human activin receptor IIB linked to the human IgG1 Fc domain for the treatment of myelodysplastic syndromes

First publication	23 September 2014
Rev.1: sponsor's change of address	10 March 2015
Rev.2: sponsor's change of address	22 May 2015
Disclaimer Please note that revisions to the Public Summary of Opinion are purely administrative updates. Therefore, the scientific content of the document reflects the outcome of the Committee for Orphan Medicinal Products (COMP) at the time of designation and is not updated after first publication.	

On 22 August 2014, orphan designation (EU/3/14/1331) was granted by the European Commission to IDEA Innovative Drug European Associates Limited, the United Kingdom, for recombinant fusion protein consisting of a modified form of the extracellular domain of human activin receptor IIB linked to the human IgG1 Fc domain for the treatment of myelodysplastic syndromes.

What are myelodysplastic syndromes?

Myelodysplastic syndromes are a group of disorders in which the red blood cells, white blood cells and platelets produced by the bone marrow (the spongy tissue inside the large bones) do not grow and mature normally. Patients with myelodysplastic syndromes can develop several symptoms including tiredness or weakness due to anaemia (low red blood cell counts), infections due to low white blood cells, and bruising or abnormal bleeding due to low platelet counts.

Myelodysplastic syndromes are long-term debilitating and life-threatening diseases because they can lead to severe anaemia, infections or bleeding, and can result in leukaemia (cancer of the white blood cells).

What is the estimated number of patients affected by the condition?

At the time of designation, myelodysplastic syndromes affected approximately 2 in 10,000 people in the European Union (EU). This was equivalent to a total of around 103,000 people*, and is below the ceiling for orphan designation, which is 5 people in 10,000. This is based on the information provided by the sponsor and the knowledge of the Committee for Orphan Medicinal Products (COMP).

What treatments are available?

At the time of designation, some medicines were authorised in the EU for the treatment of MDSs. The choice of treatment for MDSs depended on a number of factors, including the type and the extent of the disease, whether it had been treated before, and the patient's age, symptoms and general state of health. The main treatments for MDSs included chemotherapy (medicines to treat cancer) and bone marrow transplantation. This is a complex procedure where the bone marrow of the patient is cleared of cells and replaced with healthy bone marrow cells from a matched donor.

The sponsor has provided sufficient information to show that this medicine might be of significant benefit for patients with myelodysplastic syndromes because it works in a different way to existing treatments and early studies show that it may improve anaemia, an important aspect of these conditions. This assumption will need to be confirmed at the time of marketing authorisation, in order to maintain the orphan status.

How is this medicine expected to work?

This medicine is an engineered protein that has been designed to attach to certain proteins in the body which slow down (or inhibit) the development of red blood cells. By attaching to these 'inhibitory' proteins, it is expected to trap them so they don't have their normal effect on the red blood cells. As a result, production of red blood cells is increased. This is expected to improve the anaemia of patients with myelodysplastic syndromes.

What is the stage of development of this medicine?

The effects of this medicine have been evaluated in experimental models.

At the time of submission of the application for orphan designation, clinical trials with the medicine in patients with myelodysplastic syndromes were ongoing.

At the time of submission, the medicine was not authorised anywhere in the EU for myelodysplastic syndromes. Orphan designation of the medicine had been granted in the United States for these conditions.

In accordance with Regulation (EC) No 141/2000 of 16 December 1999, the COMP adopted a positive opinion on 10 July 2014 recommending the granting of this designation.

*Disclaimer: For the purpose of the designation, the number of patients affected by the condition is estimated and assessed on the basis of data from the European Union (EU 28), Norway, Iceland and Liechtenstein. At the time of designation, this represented a population of 512,900,000 (Eurostat 2014).

Opinions on orphan medicinal product designations are based on the following three criteria:

- the seriousness of the condition;
- the existence of alternative methods of diagnosis, prevention or treatment;
- either the rarity of the condition (affecting not more than 5 in 10,000 people in the EU) or insufficient returns on investment.

Designated orphan medicinal products are products that are still under investigation and are considered for orphan designation on the basis of potential activity. An orphan designation is not a marketing authorisation. As a consequence, demonstration of quality, safety and efficacy is necessary before a product can be granted a marketing authorisation.

For more information

Sponsor's contact details:

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For contact details of patients' organisations whose activities are targeted at rare diseases see:

- [Orphanet](#), a database containing information on rare diseases, which includes a directory of patients' organisations registered in Europe;
- [European Organisation for Rare Diseases \(EURORDIS\)](#), a non-governmental alliance of patient organisations and individuals active in the field of rare diseases.

Translations of the active ingredient and indication in all official EU languages¹, Norwegian and Icelandic

Language	Active ingredient	Indication
English	Recombinant fusion protein consisting of a modified form of the extracellular domain of human activin receptor IIB linked to the human IgG1 Fc domain	Treatment of myelodysplastic syndromes
Bulgarian	Рекомбинантен фузионен протеин, състоящ се от модифицирана форма на екстрацелуларния домейн на човешки активин рецептор IIB, свързан към Fc домейн на човешки IgG1	Лечение на миелодиспластичен синдром
Croatian	Rekombinantni fuzijski protein sačinjen od modificiranog oblika ljudskog aktivinskog receptora IIB povezanog na ljudsku IgG1 Fc domenu	Liječenje mijelodisplastičnih sindroma
Czech	Rekombinantní fúzní protein sestávající z modifikované formy extracelulární domény lidského receptoru typu II B pro aktivin navázané na Fc doménu lidského IgG1	Léčba myelodysplastického syndromu
Danish	Rekombinant fusionsprotein bestående af en modificeret form af ekstracellulære dele af human activin receptor IIB knyttet til det humane IgG1 Fc domæne	Behandling af myelodysplastiske syndromer
Dutch	Recombinant fusie-eiwit dat bestaat uit een gemodificeerde vorm van het extracellulaire domein van de humane activinereceptor IIB gekoppeld aan het humane IgG1 Fc-domein	Behandeling van myelodysplastische syndromen
Estonian	Rekombinantne liitvalk, mis koosneb inimese IgG1 Fc domeeniga seotud inimese aktiiviini retseptori IIB rakuvälise domeeni modifitseeritud vormist	Müelodüsplastiliste sündroomide ravi
Finnish	Ihmisen IgG1 Fc -domeeniin yhdistetystä muunnellusta solun ulkopuolisesta aktiiviini IIB -reseptorin domeenimuodosta koostuva rekombinantti fuusioproteiini	Myelodysplastisten syndroomien hoito
French	Protéine de fusion recombinante consistant en une forme modifiée du domaine extracellulaire du récepteur IIB de l'activine humaine qui se lie au domaine Fc de l'IgG1 humaine	Traitement des syndromes myélodysplasiques
German	Rekombinantes Fusionsprotein, das aus einer modifizierten Form der extrazellulären Domäne des humanen Activin-Rezeptors IIB besteht, der an die Fc-Domäne von humanem IgG1 angekoppelt wurde	Behandlung der myelodysplastischen Syndrome

¹ At the time of designation

Language	Active ingredient	Indication
Greek	Ανθρώπινη ανασυνδυασμένη πρωτεΐνη σύντηξης αποτελούμενη από τροποποιημένη μορφή του εξωκυττάριου τμήματος του υποδοχέα της ακτιβίνης IIB συνδεδεμένο με την περιοχή Fc της ανθρώπινης IgG1	Θεραπεία των μυελοδυσπλαστικών συνδρόμων
Hungarian	A humán IgG1 Fc doménhez kötődő humán aktivin receptor módosított IIB extracelluláris doménjéből álló rekombináns fúziós fehérje	Myelodysplasias syndroma kezelése
Italian	Proteina ricombinante di fusione costituita da una forma modificata del dominio extracellulare del recettore dell'activina umana di tipo IIB legata al dominio Fc di IgG1 umana	Trattamento delle sindromi mielodisplastiche
Latvian	Rekombinants fūzijas proteīns, ko veido cilvēka aktīvina IIB receptora ekstracelulārā domēna modificēta forma, kas savienota ar cilvēka IgG1 Fc domēnu	Mielodisplastisko sindromu ārstēšana
Lithuanian	Rekombinantinis sulietas baltymas, sudarytas iš modifikuotos formos ekstraląstelinio žmogaus aktivino IIB receptoriaus domeno, sujungto su žmogaus IgG1 Fc domenu	Mielodisplastinių sindromų gydymas
Maltese	Proteina ta' fużjoni rikombinanti li tikkonsisti minn għamla modifikata tal-qasam ekstraċellulari tar-riċettur uman għal attivin IIB magħqud mal-qasam IgG1 Fc uman	Kura tas-sindromi mjelodisplastici
Polish	Rekombinowane białko fuzyjne składające się ze zmodyfikowanej postaci zewnątrzkomórkowej domeny receptora ludzkiej aktywiny typu IIB połączonego z domeną Fc ludzkiej IgG1	Leczenie zespołów mielodysplastycznych
Portuguese	Proteína de fusão recombinante composta por uma forma modificada do domínio extracelular do receptor IIB da activina humana ligada ao domínio Fc da IgG1 humana	Tratamento dos síndromes mielodisplásicas
Romanian	Proteină de fuziune recombinantă constând dintr-o formă modificată a domeniului extracelular al receptorului activinei umane de tip IIB legată de domeniul Fc al IgG1 umane	Tratamentul sindromului mielodisplazic
Slovak	Rekombinantný fúzny proteín zložený z modifikovanej formy extracelulárnej časti ľudského receptora pre aktívín IIB viazaného na Fc doménu ľudského IgG1	Liečba myelodysplastického syndrómu
Slovenian	Rekombinantna fuzijska beljakovina, sestavljena iz prilagojene oblike zunajcelične domene humanega receptorja IIB za aktivin, vezanega na domeno Fc humanega IgG1	Zdravljenje mielodisplastičnega sindroma

Language	Active ingredient	Indication
Spanish	Proteína recombinante de fusión constituida por una forma modificada del dominio extracelular del receptor humano de activina tipo IIB unida al dominio Fc de la IgG1 humana	Tratamiento de los síndromes mielodisplásicos
Swedish	Rekombinant fusionsprotein bestående av en modifierad form av den extracellulära domänen av human aktivinreceptor IIB länkad till den humana IgG1 Fc-domänen	Behandling av myelodysplastiska syndrom
Norwegian	Rekombinant fusjonsprotein som består av en modifisert form av det ekstracellulære domenet for human aktivinreseptor IIB, bundet til det humane IgG1 Fc-domenet	Behandling av myelodysplastisk syndrom
Icelandic	Raðbrigða samrunaprótein sem er samsett úr breytttri mynd af utanfrumu léni aktivín viðtaka IIB manna sem er tengt við IgG1 Fc lén manna	Til meðferðar við mergmisþroskaheilkenni