



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

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Committee for Orphan Medicinal Products

Public summary of opinion on orphan designation

Heterologous human adult liver-derived progenitor cells for the treatment of ornithine translocase deficiency (hyperornithinaemia-hyperammonaemia homocitrullinuria (HHH) syndrome)

On 17 July 2013, orphan designation (EU/3/13/1167) was granted by the European Commission to Promethera Biosciences, Belgium, for heterologous human adult liver-derived progenitor cells for the treatment of ornithine translocase deficiency (hyperornithinaemia-hyperammonaemia homocitrullinuria (HHH) syndrome).

What is ornithine translocase deficiency (hyperornithinaemia-hyperammonaemia homocitrullinuria (HHH) syndrome)?

Ornithine translocase deficiency (also known as hyperornithinaemia-hyperammonaemia homocitrullinuria (HHH) syndrome) is one of the inherited disorders known as 'urea cycle disorders', which cause ammonia to accumulate in the blood. Patients with this disorder lack 'ornithine translocase', one of the liver enzymes that are needed to get rid of excess nitrogen. In the absence of this liver enzyme, excess nitrogen accumulates in the body in the form of ammonia, which can be toxic at high levels, especially to the brain. Symptoms of the disease may appear during adulthood and include lethargy (lack of energy), vomiting, loss of appetite, seizures (fits) and coma.

Ornithine translocase deficiency is a long-term debilitating and life-threatening disease that leads to mental retardation and is associated with poor overall survival.

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

At the time of designation, ornithine translocase deficiency affected less than 0.01 in 10,000 people in the European Union (EU). This was equivalent to a total of fewer than 500 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).

*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 27), Norway, Iceland and Liechtenstein. This represents a population of 509,000,000 (Eurostat 2013).



What treatments are available?

At the time of designation, no satisfactory methods were authorised in the EU for the treatment of ornithine translocase deficiency. Patients were advised to control their dietary intake of proteins, which are rich in nitrogen, to reduce the amount of ammonia formed in the body.

How is this medicine expected to work?

This medicine is an advanced therapy medicine that belongs to the group called 'somatic cell therapy products'. These are medicines that contain cells or tissues that have been manipulated to change their biological characteristics so that they can be used to cure, diagnose or prevent a disease. The medicine is made up of progenitor (immature) cells derived from the liver of an adult donor ('heterologous'). When implanted into the liver of a patient, it is believed that these heterologous liver-derived progenitor cells will develop into mature, healthy liver cells that can produce the ornithine translocase enzyme. The new cells are thereby expected to reduce the accumulation of ammonia and to relieve the symptoms of the disease.

What is the stage of development of this medicine?

The effects of heterologous human adult liver-derived progenitor cells 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 urea cycle disorders were ongoing.

At the time of submission, the medicine was not authorised anywhere in the EU for ornithine translocase deficiency. Orphan designation of the medicine had been granted in the United States for urea cycle disorders.

In accordance with Regulation (EC) No 141/2000 of 16 December 1999, the COMP adopted a positive opinion on 13 June 2013 recommending the granting of this designation.

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	Heterologous human adult liver-derived progenitor cells	Treatment of ornithine translocase deficiency (hyperornithinaemia-hyperammonaemia homocitrullinuria (HHH) syndrome)
Bulgarian	Хетероложни човешки прогениторни клетки, получени от черен дроб на възрастни	Лечение на орнитин транслоказна недостатъчност(синдром на хиперорнитинемия-хиперамонемия-хомоцитрулинурия)
Czech	Heterologní progenitorové buňky získané z jater dospělého člověka	Léčba deficitu translokázy ornitinu (hyperornitinémie, hyperamonémie, homocitrullinurie (HHH) syndromu)
Croatian	Heterologne progenitorske stanice izolirane iz jetre odraslog čovjeka	Liječenje nedostatka ornitinske translokaze (sindrom hiperornitinemije, hiperamoniemije, homocitrulinurije (HHH))
Danish	Heterologe humane leverderiverede progenitorceller fra voksne	Behandling af ornitintranslocase mangel (hyperornitinæmi hyperammonæmi homocitrullinuri (HHH) syndrom)
Dutch	Uit de adulte lever afgeleide heterologe humane progenitorcellen	Behandeling van ornithinetranslocase deficiëntie, (hyperornithinemia-hyperammonemia homocitrullinurie (HHH) syndroom)
Estonian	Heteroloogilised täiskasvanud inimese maksast pärinevad eellasrakud	Ornitiini translokaasi puudulikkuse ravi (hüperornitineemia-hüperammonemia-homotsitrullinuuria (HHH) sündroom)
Finnish	Heterologiset aikuisen ihmisen maksaperäiset progenitorisolut	Ornitiinitranslokaasin puutoksen (hyperornitinemia-hyperammonemia-homositrullinurian eli HHH-oireyhtymän) hoito
French	Cellules progénitrices hétérologues dérivées du foie adulte humain	Traitement du déficit en ornithine translocase; (syndrome d'hyperammoniémie, hyperornithinémie, homocitrullinurie (HHH))
German	Aus der adulten Leber abgeleitete heterologe Vorläuferzellen	Behandlung eines Ornithin-Translokase Mangels (Hyperornithinämie, Hyperammonämie, Homocitrullinurie oder HHHSyndrom)
Greek	Ανθρώπινα ετερόλογα προγονικά κύτταρα προερχόμενα από ήπαρ ενηλίκου	Θεραπεία της ανεπάρκειας της τρανσλοκάσης της ορνιθίνης, (σύνδρομο υπερορνιθιναιμίας-υπεραμμωναιμίας-ομοκιτρουλινουρίας (HHH))
Hungarian	Heterológ humán felnőttek májából származó progenitor sejtek	Ornithinhordozó elégtelenség (hiperornithinaemia-hiperammoniaemia-homocitrullinuria (HHH szindróma)) kezelésére
Italian	Cellule progenitrici eterologhe di fegato umano adulto	Trattamento della sindrome da carenza di ornitina-translocasi (iperornitinemia-iperammoniemia-omocitrullinemia)

¹ At the time of designation

Language	Active ingredient	Indication
Latvian	Heterologas pieauguša cilvēka aknu priekšgājēju šūnas	Ornitīna translokāzes deficīta (hiperornitinēmijas-hiperamoniēmijas, homocitrulinūrijas sindroma) ārstēšana
Lithuanian	Heterologinės pirminės ląstelės, išskirtos iš suaugusio žmogaus kepenų	Ornitino translokazės stoka (hiperornitinemijos-hiperamoniemijos-homocitrulinurijos (HHH) sindromo) gydymas
Maltese	Ċelloli proġenituri eteroloġi minn fwiċċ adult uman	Kura ta' nuqqas ta' ornithine translocase (sindrome ta' iperornitinemija-iperamoniemija-homocitrullinurja) (HHH syndrome))
Polish	Ludzkie heterologiczne komórki progenitorowe izolowane z wątroby osoby dorosłej	Leczenie niedoboru translokazy ornitynowej (zespół (HHH) hiperornitynemii-hiperamoniemii-homocytulinurii)
Portuguese	Células progenitoras heterólogas derivadas do fígado de adultos humanos	Tratamento da deficiência de ornitina translocase (hiperornitinémia – hiperamoniémia – homocitrulinúria (HHH))
Romanian	Celule progenitoare heterologe umane derivate din celule hepatice adulte	Tratamentul deficienței de ornitin translocază (sindromul hiperornitinemie-hiperamoniemie-homocitrulinurie (HHH))
Slovak	Heterológne progenitorové bunky derivované z pečene dospelého človeka	Liečba nedostatku ornitíntranslokázy (hyperornitinémia-hyperamoniémia-homocitrulínúria (HHH) syndróm)
Slovenian	Heterologne progenitorne celice, pridobljene iz jeter odraslega človeka	Zdravljenje pomanjkanja ornitinske translokaze (hiperornitinemija-hiperamoniemija-homocitrulinurija (HHH) sindrom)
Spanish	Células progenitoras heterólogas extraídas de hígado humano adulto	Tratamiento de la deficiencia de ornitina translocasa (hiperornitinemia-hiperamoniemia-homocitrulinuria; síndrome HHH)
Swedish	Humana heterologa leverderiverade progenitorceller från vuxna	Behandling av ornitintranslokasbrist (hyperornitinemi -hyperamoniemi homocitrullinuri (HHH) syndrom)
Norwegian	Heterologe leverderiverte progenitorceller fra voksne	Behandling av ornitin translokasemangel (hyperornitinemi-hyperamoniemi homocitrullinuri (HHH)syndrom)
Icelandic	Ósamgena lifrarforstigsfrumur úr fullorðnum	Meðferð á ornithín translókasa skorti (hyper ornithíníndreyri-hýperammóníumdreyri hómósítrúllinmigu (HHH) heilkenni)