

Damoctocog alfa pegol

Resolution of: 20 June 2019 / 27 August 2018
Entry into force on: 20 June 2019 / 27 August 2018
BANz AT 22 07 2019 B1 / BANz AT14 11 2019 B2

Valid until: unlimited

Therapeutic indication (according to the marketing authorisation of 22 November 2018):

Treatment and prophylaxis of bleeding in previously treated patients 12 years of age and older with haemophilia A (congenital factor VIII deficiency).

1. Additional benefit of the medicinal product in relation to the appropriate comparator therapy

Patients aged 12 and older with haemophilia A (congenital factor VIII deficiency)

Appropriate comparator therapy:

- Recombinant or human plasma-derived blood coagulation factor VIII preparations

Extent and probability of additional benefit of Damoctocog alfa pegol compared to the appropriate comparator therapy:

An additional benefit is not proven.

2. Number of patients or demarcation of patient groups eligible for treatment

Patients aged 12 and older with haemophilia A (congenital factor VIII deficiency)

Approx. 2,840–3,190 patients

3. Requirements for a quality-assured application

The requirements of the product information are to be taken into account. The European Medicines Agency (EMA) provides the contents of the product information (summary of product characteristics, SmPC) for Jivi® (active ingredient: Damoctocog alfa pegol) at the following publicly accessible link (last access: 29 April 2019):

https://www.ema.europa.eu/documents/product-information/jivi-epar-product-information_de.pdf

Treatment with Damoctocog alfa pegol should be initiated and monitored by specialists experienced in the treatment of haemophilia.

4. Treatment costs

Patients aged 12 and older with haemophilia A (congenital factor VIII deficiency)

Annual treatment costs¹:

Designation of the therapy	Annual treatment costs/patient	
Medicinal product to be assessed:		
Damoctocog alfa pegol	Adults	€ 484.056,30 – € 679,540.58
	12 – <18 years	€ 345.754,50 – € 485,386.13
Appropriate comparator therapy:		
Recombinant blood coagulation factor VIII		
Rurioctocog alfa pegol	Adults	€ 506,797.20 – € 615,396.60
	12 – <18 years	€ 361,998.00 – € 470,597.40
Efmoctocog alfa	Adults	€ 184,120.97 – € 786,367.47
	12 – <18 years	€ 143,205.20 – € 581,228.13
Lonoctocog alfa	Adults	€ 200,986.24 – € 732,164.16
	12 – <18 years	€ 143,561.60 – € 559,890.24
Moroctocog alfa	Adults	€ 246,443.05 – € 739,329.15
	12 – <18 years	€ 176,030.75 – € 528,092.25
Octocog alfa ²	Adults	€ 237,718.21 – € 713,154.62
	12 – <18 years	€ 169,798.72 – € 509,396.16
Simoctocog alfa ³	Adults	€ 222,306.88 – € 666,920.63
	12 – <18 years	€ 158,790.63 – € 476,371.88
Turoctocog alfa	Adults	€ 269,642.10 – € 654,845.10
	12 – <18 years	€ 192,601.50 – € 500,763.90
Blood coagulation factor VIII derived from human plasma		
Human plasma preparations ⁴	Adults	€ 210,873.95 – € 632,621.85
	12 – <18 years	€ 150,624.25 – € 451,872.75

Costs after deduction of statutory rebates (LAUER-TAXE®) as last revised: 01 June 2019)

Costs for additionally required SHI services: not applicable

¹ The prices are not subject to the Pharmaceutical Price Ordinance (AMPreisV).

² Cost representation based on the information provided in the summary of product characteristics for Kovaltry®. Further proprietary medicinal products are available.

³ Cost representation based on the information provided in the summary of product characteristics for Nuwiq®. Further proprietary medicinal products are available.

⁴ Cost representation based on the information provided in the summary of product characteristics for Fanhdi®. Further proprietary medicinal products are available.