



Effective: July 1, 2025

	⊠ Prior Authorization
	□ Non-Formulary
Guideline Type	□ Step-Therapy

Applies to:

Commercial Products

- □ Harvard Pilgrim Health Care Commercial products; Fax 617-673-0988
- □ Tufts Health Plan Commercial products; Fax 617-673-0988 CareLinkSM – Refer to CareLink Procedures, Services and Items Requiring Prior Authorization

Public Plans Products

- □ Tufts Health Direct A Massachusetts Qualified Health Plan (QHP) (a commercial product); Fax 617-673-0988
- □ Tufts Health Together MassHealth MCO Plan and Accountable Care Partnership Plans; Fax 617-673-0939
- □ Tufts Health RITogether A Rhode Island Medicaid Plan; Fax 617-673-0939
- ☑ Tufts Health One Care* A Medicare-Medicaid Plan (a dual eligible product); Fax 617-673-0956
 *The MNG applies to Tufts Health One Care members unless a less restrictive LCD or NCD exists.

Senior Products

- In Tufts Health Plan Senior Care Options (SCO), (a dual-eligible product); Fax 617-673-0956
- In Tufts Medicare Preferred HMO, (a Medicare Advantage product); Fax 617-673-0956
- ☑ Tufts Medicare Preferred PPO, (a Medicare Advantage product); Fax 617-673-0956

Note: While you may not be the provider responsible for obtaining prior authorization, as a condition of payment you will need to ensure that prior authorization has been obtained.

Overview

Hemophilia is an inherited, lifelong bleeding disorder caused by a deficiency of coagulation factors. The two most common types of hemophilia are hemophilia A (Factor VIII deficiency) and B (Factor IX deficiency), and either type can lead to spontaneous bleeding and prolonged bleeding following an injury or surgical procedure. There are varying severities of hemophilia A and B depending on the level of factor produced by the patient. Severe hemophilia frequently results in bleeding even in the absence of trauma; moderate hemophilia is associated with less bleeding, and mild hemophilia usually results in bleeding only after obvious trauma. Historically hemophilia treatment involves replacing the deficient coagulation factor through episodic (on-demand) treatment or prophylaxis. Newer, easier-to-administer products have provided options for the management of patients with hemophilia A and B and include Alhemo. Alhemo is a tissue factor pathway inhibitor (TFPI) antagonist. Alhemo works by reducing the amount, and therefore, the activity of, the naturally occurring anticoagulation protein TFPI. This results in increased amounts of thrombin, an enzyme that is critical in blood clotting.

Approval of Alhemo was based on the open-label explorer 7 trial which included 133 patients (male patients aged 12 years and older and weighing ≥25 kg) with hemophilia A or B with inhibitors. The primary objective compared the number of treated spontaneous and traumatic bleeding episodes (measured by annualized bleeding rate [ABR]), showed an 86% reduction of ABR in patients randomized to receive Alhemo prophylaxis compared to no prophylaxis.

Food and Drug Administration (FDA) - Approved Indications:

Alhemo (concizumab-mtci) is a TFPI antagonist indicated for routine prophylaxis to prevent or reduce the frequency of bleeding in episodes in adult and pediatric patients 12 years of age and older with hemophilia A (congenital factor VIII deficiency) with factor VIII inhibitors, or hemophilia B (congenital factor IX deficiency) with factor IX inhibitors.

Clinical Guideline Coverage Criteria			
The plan may authorize coverage of Alhemo for Members when ALL of the following criteria are met:			
Initial Authorization Criteria			
1.	1. Documented diagnosis of hemophilia A or B		
	AND		
2.	2. Documentation for use as prophylaxis to prevent or reduce the frequency of bleeding episodes		
	AND		
3.			
	AND		
4. Documentation the patient does not have inhibitors (hemophilia A: FVIII, hemophilia B: FIX)			
	AND		
5.	5. Prescribed by or in consultation with a hematologist		
Reauthorization Criteria			
1.	1. Documented diagnosis of hemophilia A or B		
	AND		
2.	The patient is at least 12 years of age or older		
	AND		
3.	Prescribed by or in consultation with a hematologist		
	AND		
4.	•		
	AND		
5.	 Documentation the Member has experienced a therapeutic res frequency of bleeds 	oonse from Alhemo as defined by a reduction in the	

Limitations

- Coverage of Alhemo will be authorized for 12 months
- Members new to the plan stable on Alhemo should be reviewed against Reauthorization Criteria.

Codes

None

References

- 1. Alhemo (concizumab-mtci) [package insert]. Plainsboro, NJ; Novo Nordisk Inc: December 2024.
- 2. Astermark J, et al. Efficacy and safety of concizumab prophylaxis in patients with hemophilia a or b without inhibitors: 56week cut-off results of the Phase 3 explorer8 study. Blood. 2023;142(suppl 1):2609.
- Chowdary P, et al. Concizumab prophylaxis in people with haemophilia A or haemophilia B without inhibitors (explorer8): a prospective, multicentre, open-label, randomised, phase 3a trial [published correction appears in Lancet Haematol. 2024;11(12):e886. doi:10.1016/S2352-3026(24)00353-3]. Lancet Haematol. 2024;11(12):e891–e904.
- 4. Matsushita T, et al. Phase 3 Trial of Concizumab in hemophilia with inhibitors. N Engl J Med. 2023;389(9):783–794.
- Alok Srivastava et al on behalf of the WFH Guidelines for the Management of Hemophilia panelists and co-authors. World Federation of Hemophilia guidelines for the management of hemophilia. 3rd edition. 2020. Accessed February 2025. Available at: <u>https://elearning.wfh.org/resource/treatment-guidelines/</u>.

Approval And Revision History

May 13, 2025: Reviewed by Pharmacy and Therapeutics Committee (P&T). June 2025: Joint Medical Policy and Health Care Services UM Committee review (eff 7/1/25)

Background, Product and Disclaimer Information

Point32Health prior authorization criteria to be applied to Medicare Advantage plan members is based on guidance from Medicare laws, National Coverage Determinations (NCDs) or Local Coverage Determinations (LCDs). When no guidance is provided, Point32Health uses clinical practice guidance published by relevant medical societies, relevant medical literature, Food and Drug Administration (FDA)-approved package labeling, and drug compendia to develop prior authorization criteria to apply to Medicare Advantage plan members. Medications that require prior authorization generally meet one or more of the following criteria: Drug product has the potential to be used for cosmetic purposes; drug product is not considered as first-line

Point32Health companies

treatment by medically accepted practice guidelines, evidence to support the safety and efficacy of a drug product is poor, or drug product has the potential to be used for indications outside of the indications approved by the FDA. Prior authorization and use of the coverage criteria within this Medical Necessity Guideline will ensure drug therapy is medically necessary, clinically appropriate, and aligns with evidence-based guidelines. We revise and update Medical Necessity Guidelines annually, or more frequently if new evidence becomes available that suggests revisions.

Treating providers are solely responsible for the medical advice and treatment of Members. The use of this guidelines not a guarantee of payment or a final prediction of how specific claim(s) will be adjudicated. Claims payment is subject to eligibility and benefits on the date of service, coordination of benefits, referral/authorization, utilization management guidelines when applicable, and adherence to plan policies, plan procedures, and claims editing logic.