Recombinant FVIII Products (Turoctocog Alfa and Turoctocog Alfa Pegol) Stable Up to 40°C

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Purpose: The stability under high-temperature conditions of factor VIII (FVIII) concentrates for replacement therapy is of critical importance to patients, particularly those who reside in, or travel to, regions with high ambient temperatures. Concerns about product stability may limit or prevent access to treatment for patients and may limit their ability to live a close-to-normal life. This study evaluated the effect of hot and humid storage conditions on the long-term stability of the recombinant FVIII products, turoctocog alfa and turoctocog alfa pegol.

Methods: Turoctocog alfa samples were assessed for stability at 30°C for 9 months or 40°C for 3 months following storage at 5°C for 21 or 27 months, respectively, while turoctocog alfa pegol samples were assessed at 30°C for 12 months or 40°C for 3 months following storage at 5°C for 18 or 27 months, respectively. In addition, turoctocog alfa and turoctocog alfa pegol dry powders were evaluated for stability at 5°C/ambient humidity (AH) for 30 months, 30°C/75% relative humidity (RH) for 12 months and 40°C/75% RH for 6 months. Both studies utilized a range of product strengths. Key stability assessments included oxidized forms, potency, water content and high molecular weight protein (HMWP).

Results: Both turoctocog alfa and turoctocog alfa pegol remained stable following storage at 40°C/75% RH for 3 months, and at single temperatures (5°C/AH, 30 and 40°C/75% RH), without any major increase in HMWP or any impairment of potency or water content.

Conclusion: Turoctocog alfa and turoctocog alfa pegol offer stability at 40°C for up to 3 months without jeopardizing the quality of each product. These stability characteristics may offer patients flexibility with product storage and daily use.

Keywords: factor VIII, hemophilia A, storage flexibility, temperature stability, turoctocog alfa, turoctocog alfa pegol

Plain Language Summary

For patients with hemophilia A and their caregivers, it is crucial that factor VIII (FVIII) products used for replacement therapy can be stored under intense heat for long periods. Concerns about whether their FVIII products are affected by intense heat may prevent patients from accessing treatment and living a normal life when they are in hot and humid regions. This study therefore evaluated the effect of temperatures up to 40°C and humid storage conditions on the long-term stability of the FVIII products: turoctocog alfa and turoctocog alfa pegol.

Turoctocog alfa samples were assessed for stability at 30°C for 9 months or 40°C for 3 months following storage at 5°C for 21 or 27 months, respectively, while turoctocog alfa pegol samples were assessed at 30°C for 12 months or 40°C for 3 months following storage at 5°C for 18 or 27 months, respectively. In addition, turoctocog alfa and turoctocog alfa pegol were evaluated for stability at 5°C/ambient humidity (AH) for 30 months, 30°C/75% relative humidity (RH) for 12 months and 40°C/75% RH for 6 months.
We found both FVIII products remained stable following storage at 40°C/75% RH for 3 months, and at single temperatures (5°C/AH, 30 and 40°C/75% RH), without any major increase in protein products or impairment of potency or water content.

Turoctocog alfa and turoctocog alfa pegol offer stability at 40°C for up to 3 months without jeopardizing the quality of each product. These stability characteristics offer patients flexibility with product storage and daily use.

**Introduction**

In patients with hemophilia A, it is recommended to treat bleeds with factor VIII (FVIII) within 2 hours, which requires access to replacement factor at all times. Furthermore, FVIII prophylaxis—the gold standard of care for severe hemophilia A patients—requires patients or their caregivers to administer regular intravenous injections of FVIII (usually every other day or 3 times a week with standard half-life FVIII concentrates and once or twice weekly with extended half-life [EHL] FVIII products).

Manufacturers recommend storing both standard and EHL FVIII products in a refrigerator (or below 25°C) and reconstituting them at room temperature. However, some patients are unaware that most FVIII products remain stable for a limited time at higher temperatures, therefore, their adherence to (and satisfaction with) therapy may be adversely affected by the perceived limitations of FVIII product storage conditions, particularly when patients are traveling. With reference to product-related factors for hemophilia treatment, patients have reported a preference for storage-flexible products that allow reduced injection times, greater patient adherence, less waste, greater ease of handling, and fewer restrictions in daily activities such as travel and sports.

The structural stability of recombinant therapeutic products can affect biological activity, clearance, and immunogenicity, with increasing instability occurring over long-term storage. While the World Health Organization recommendations are that the stability of therapeutic products should be assessed based on climatic zones with a maximum temperature of 30°C and 75% relative humidity (RH; Figure 1), they also recognize that storage conditions often exceed these temperatures. It is thus crucial to also assess the stability of FVIII products at temperatures above 30°C.

Turoctocog alfa (NovoEight®, Novo Nordisk A/S, Bagsværd, Denmark) and turoctocog alfa pegol (an extended half-life glycoPEGylation of turoctocog alfa, Esperoct®, Novo Nordisk A/S, Bagsværd, Denmark) are recombinant FVIII (rFVIII) products developed to prevent and treat bleeding

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**Figure 1** ICH quality guidelines for pharmaceutical stability storage. ([https://q1scientific.com/ich-quality-guidelines/](https://q1scientific.com/ich-quality-guidelines/))

**Notes:** Long-term testing conditions for: Zone I, 21°C/45% RH (temperate); Zone II, 25°C/60% RH (subtropical and Mediterranean); Zone III, 30°C/35% RH (hot/dry); Zone IV, 30°C/75% RH (hot/very humid). Data from WHO.19

**Abbreviations:** ICH, International Council on Harmonisation; RH, relative humidity; WHO, World Health Organization.
episodes in patients with hemophilia A.20–26 The activated forms of both products have the same primary structure as native activated FVIII.27 The production steps for turoctocog alfa are the same as for turoctocog alfa pegol prior to PEGylation;27–29 importantly, PEGylation of turoctocog alfa does not affect product stability.27 Both products are supplied as a dry powder in single-dose vials of 250, 500, 1000, 1500, 2000, and 3000 IU (turoctocog alfa)7,8 and 500, 1000, 1500, 2000, and 3000 IU (turoctocog alfa pegol).13,14 The current approved storage and shelf life of the dry powder for both products is storage at 5°C for up to 30 months, during which time the products may be stored at room temperature (≤30°C) for ≤12 months or up to 40°C for ≤3 months.7,8,13,14

Patients may live in, or travel to, countries with a hot and humid climate. To allow these patients to use turoctocog alfa and turoctocog alfa pegol safely, we evaluated the effect of high-temperature, high-humidity storage conditions on the long-term stability of both products. In the first study, turoctocog alfa samples were assessed at 30°C for 9 months or 40°C for 3 months following storage at 5°C for 21 or 27 months, respectively, while turoctocog alfa pegol samples were assessed at 30°C for 12 months or 40°C for 3 months following storage at 5°C for 18 or 27 months, respectively. The second study assessed the dry powders of turoctocog alfa and turoctocog alfa pegol when stored at 5°C/ambient humidity (AH) for 30 months, 30°C/75% RH for 12 months, and 40°C/75% RH for 6 months (Figure 2).

Patients and Methods
Product and Sample Preparation
Commercially available turoctocog alfa and turoctocog alfa pegol in dry-powder batches were each evaluated at their lowest and two highest strengths: 250, 2000, and 3000 IU for turoctocog alfa, and 500, 2000, and 3000 IU for turoctocog alfa pegol (Figure 2).

Long-Term Stability Study
Turoctocog Alfa
The long-term stability of turoctocog alfa dry-powder batches was assessed during single temperature storage at 40°C (±2°C) and 75% RH (±5% RH) for 3 months, and at 30°C (±2°C) and 75% RH (±5% RH) for 9 months. A reference sample for each dose was maintained at 5°C (±3°C) in darkness and AH for 3 months. Prior to the start of this study, the turoctocog alfa samples had been stored at 5°C (±3°C) in darkness and AH for 21 and 27 months (Figure 2).

Turoctocog Alfa Pegol
Dry-powder batches of turoctocog alfa pegol were assessed for long-term stability at 40°C (±2°C) in darkness and 75% RH (±5% RH) for 3 months, and at 30°C (±2°C) and 75% RH (±5% RH) for 12 months. Prior to the start of this study, the dry products had been stored at 5°C (±2°C) in darkness and AH for 18 and 27 months for consecutive temperature storage (Figure 2).

Stability Assessments and Assay Methods
All studies were performed according to current International Conference on Harmonisation (ICH) bracketing design guidelines.30 The key stability parameters assessed for both products were those from the drug product specifications that are susceptible to change during storage and/or those that could potentially influence product quality: namely, oxidized forms, potency, water content, and high-molecular-weight protein (HMWP). For

![Figure 2](image-url)
oxidized forms, potency, water content, and HMWP, the results from all assays were evaluated against predetermined acceptance limits and calculated using appropriate statistical methods. Potency was expressed as % target. The specific ranges for each assay are noted within the appropriate sections below.

**Oxidized Forms**

Oxidized forms within the reconstituted turoctocog alfa and turoctocog alfa pegol products were assessed using reverse-phase-high performance liquid chromatography (RP-HPLC). Analysis was performed on an HPLC system equipped with processing software and a 4.0 x 250-mm, C₄ 5 μm, 300 Å column (Novo Nordisk Pharmatech A/S, Køge, Denmark). The column temperature was set at 40°C for turoctocog alfa and 40°C ± 2°C for turoctocog alfa pegol, with a detection wavelength of 215 nm. A gradient of 35–100% eluent B (0.09% [turoctocog alfa] or 0.14% [turoctocog alfa pegol] trifluoroacetic acid [TFA] in 80% acetonitrile in purified water) and 65–0% eluent A (0.1% [turoctocog alfa] or 0.15% [turoctocog alfa pegol] TFA in purified water) was applied over a 40-min period with a flow rate of 1 mL/min. The composition of 100% eluent B was then maintained for 5 min, before being changed back to the initial conditions over 1 min, followed by column equilibration for 14 min, resulting in a total run time of 60 min. Oxidized forms were calculated as the percentage area on the resulting chromatograms. RP-HPLC chromatograms showed all components were present in the drug substances and therefore in the drug products (chromatograms not shown).

**Potency**

The potency of turoctocog alfa and turoctocog alfa pegol was investigated using the chromogenic assay (Chromogenix Coamatic® Factor VIII) and the SynthASil (HemosIL, Instrumentation Laboratory) activated partial thromboplastin time reagent.

Turoctocog alfa and turoctocog alfa pegol product samples, calibrator, and blank samples were analyzed in triplicate on the ACL® Elite Pro analyzer. Absorbance readings for product, reference samples, and calibrator were used to calculate potency using slope–ratio analysis or parallel line regression (chromogenic assays). Potency was expressed as % target (European Pharmacopoeia acceptance limits are 80–125% of target).

**Water Content**

The water content of turoctocog alfa and turoctocog alfa pegol samples was evaluated using near-infrared (NIR) spectroscopy for the high-temperature stability samples only. NIR spectroscopy was conducted using a Fourier transform-NIR Spectrometer (MPA, Bruker, Billerica, MA, USA) equipped with an integrating sphere (or equivalent). Both products were scanned in the frequency ranges of 7502.1–6098.1 and 5450.1–4597.7 nm at a resolution of 8 cm⁻¹, with an average of 32 scans being recorded for each spectrum. The spectrometer was equipped with OPUS software (Bruker) (or equivalent) and data were analyzed using a partial least squares fit method. Turoctocog alfa and turoctocog alfa pegol samples were analyzed without any pretreatment; however, NIR spectroscopy data were pretreated using vector normalization and a first derivative to enhance spectral information and correct interferences from the analyzed material that might otherwise induce baseline drift and changes in maximum absorbance. Karl–Fischer coulometry was used as the reference method for determining the calibration function and to analyze samples in cases where NIR results were not accepted (for example, due to persistent outliers).

**High Molecular Weight Protein**

Size exclusion-HPLC (SE-HPLC) was used to determine the presence of protein aggregates in the turoctocog alfa and turoctocog alfa pegol samples. Prior to SE-HPLC analysis, the samples were reconstituted in either 1.0 mL (250-IU samples [turoctocog alfa] and 500-IU samples [turoctocog alfa pegol]) or 4.3 mL (2000- and 3000-IU samples, both products) of 0.9% sodium chloride solution. SE-HPLC measurements for turoctocog alfa were performed using an HPLC system equipped with a Shodex PROTEIN KW-803, 8 × 300-mm column (Shodex), or equivalent. For turoctocog alfa pegol, SE-HPLC measurements were performed using a Sepax Technologies SRT SEC-500, 7.8 × 300-mm, 5-μm, 500 Å column (Sepax Technologies, Newark, DE, USA). An elution flow rate of 0.4 mL/min was employed for turoctocog alfa, using a column temperature of 30°C and excitation and emission detection wavelengths of 285 and 335 nm, respectively. For turoctocog alfa pegol, an elution flow rate of 0.3 mL/min was used, with a column temperature of 23°C, an excitation detection wavelength of 285 nm, and an emission detection wavelength of 345 nm. The eluent buffer consisted of 10-mM TRIS, 10-mM CaCl₂, 300-mM NaCl,
and 5% 2-propanol at pH 7.0 (turoctocog alfa) or 10-M BIS-TRIS propanol, 10-mM calcium acetate hydrate, 0.5-M sodium chloride, and 10% isopropanol at a pH adjusted to 6.8 with acetic acid (turoctocog alfa pegol). The injection load volume was within the validated range of the method (0.95–7.8 μg). Run times for products reconstituted in 4.3- and 1.0-mL 0.9% sodium chloride solutions were ≥70 and ≥80 min, respectively, for turoctocog alfa, and 60 min for turoctocog alfa pegol. HMWP content was determined by calculating the area percentage of the HMWP peak on the resulting chromatogram.

**Results**

**Oxidized Forms**

Oxidized forms, which are inactive forms of rFVIII that can reduce potency, are the stability-limiting factor. Oxidized forms increased as a function of time and temperature following an Arrhenius correlation for both products (Figure 3A). Results for turoctocog alfa samples stored at consecutive temperatures of 30°C/75% RH for 9 months (Figure 3B) or 40°C/75% RH for 3 months (Figure 3C), following storage at 5°C for 21 or 27 months, respectively, showed that levels of oxidized forms were within the predetermined acceptance limits for all batches. Similar results were seen for turoctocog alfa pegol samples stored at consecutive temperatures of 30°C/75% RH for 12 months (Figure 3G) or 40°C/75% RH for 3 months (Figure 3H), following storage at 5°C for 18 or 27 months, respectively. There was a small increase in the occurrence of oxidized forms of both turoctocog alfa and turoctocog alfa pegol following storage at single temperatures of 5°C, 30°C/75% RH, and 40°C/75% RH for 30, 12, and 6 months, respectively (Figure 3D–F and Figure 3I–K, respectively). For both products, oxidized forms were close to 5% after storage at 30°C/75% RH for 12 months (Figures 3E and J) and approaching 7% after storage at 40°C/75% RH for 6 months (Figures 3F and K).

**Potency**

As potency is used as the basis for dosage calculations and treatment strategies, it is important to determine whether potency is affected by hot and humid storage conditions. Changes in potency were within the limits for all turoctocog alfa batches when stored at consecutive temperature conditions for 9 months (Figure 4A) or 3 months (Figure 4B), respectively. Potency was also within the limits for all turoctocog alfa pegol batches when stored at consecutive temperature conditions for 12 months (Figure 4F) or 3 months (Figure 4G), respectively. These limits are also in line with the European Pharmacopoeia acceptance criteria of −20% to +125% from target. The results of the potency test of dry turoctocog alfa and turoctocog alfa pegol samples stored at single temperatures for 30, 12, and 6 months, respectively, are shown in Figure 4C–E and Figure 4H–J, respectively. Potency was maintained within the limits for all turoctocog alfa and turoctocog alfa pegol sample strengths tested.

**Water Content**

Excessive water content can increase degradation and reduce the stability of a therapeutic product. When turoctocog alfa batches were stored at consecutive temperature conditions for 9 months (Figure 5A) or 3 months (Figure 5B), respectively, water content for all batches was within the range of 1–2%, corresponding to the observed ranges in stability studies. Likewise, water content was also within this upper range for all turoctocog alfa pegol batches when stored at consecutive temperature conditions for 12 months (Figure 5F) or 3 months (Figure 5G), respectively. There was a small increase in the water content with increasing temperature for both turoctocog alfa and turoctocog alfa pegol samples following storage at single temperatures for 30, 12, and 6 months, respectively (Figure 5C–E and Figure 5H–J, respectively). Water content was approximately 1% after storage at 30°C/75% RH for 12 months (Figure 5D and I) and <2% after storage at 40°C/75% RH for 6 months (Figure 5E and J).

**High Molecular Weight Protein**

Protein product aggregates, including HMWP, are described as potent inducers of immune responses to therapeutic protein products. When turoctocog alfa batches were stored at consecutive temperature conditions for 9 months (Figure 6A) or 3 months (Figure 6B), respectively, HMWP for all batches was within the range of 2–4%. Similarly, HMWP was also within this upper range for all turoctocog alfa pegol batches when stored at consecutive temperature conditions for 12 months (Figure 6F) or 3 months (Figure 6G), respectively. Small increases in HMWP were observed for both turoctocog alfa and turoctocog alfa pegol samples following storage at single temperatures for 30, 12, and 6 months, respectively (Figure 6C–E and Figure 6H–J, respectively); for both products, HMWP levels were ≤3% after storage at 30°C/
Figure 3 Oxidized forms of turoctocog alfa dry powder samples as the rate of oxidation (Arrhenius correlation) (A), when stored at 5°C for 21 months followed by storage at 30°C for 9 months (B), 5°C for 27 months followed by storage at 40°C for 3 months (C), and when stored at 5°C for 30 months (D), 30°C for 12 months (E), and 40°C for 6 months (F). Oxidized forms of turoctocog alfa pegol dry powder samples when stored at 5°C for 18 months followed by storage at 30°C for 12 months (G), 5°C for 27 months followed by storage at 40°C for 3 months (H), and when stored at 5°C for 30 months (I), 30°C for 12 months (J), and 40°C for 6 months (K). Shelf-life limits for both products are shown by the horizontal dotted lines.

Abbreviations: IU, international units; T, temperature (in degrees Kelvin).
Figure 4 Potency of turoctocog alfa dry powder samples when stored at 5°C for 21 months followed by storage at 30°C for 9 months (A), 5°C for 27 months followed by storage at 40°C for 3 months (B), 5°C for 30 months (C), 30°C for 12 months (D), and 40°C for 6 months (E). Potency of turoctocog alfa pegol dry powder samples when stored at 5°C for 18 months followed by storage at 30°C for 12 months (F), 5°C for 27 months followed by storage at 40°C for 3 months (G), 5°C for 30 months (H), 30°C for 12 months (I), and 40°C for 6 months (J). Upper and lower shelf-life limits for both products are shown by the horizontal dotted lines.

Abbreviation: IU, international units.
Figure 5 Water content of turoctocog alfa dry powder samples when stored at 5°C for 21 months followed by storage at 30°C for 9 months (A), 5°C for 27 months followed by storage at 40°C for 3 months (B), 5°C for 30 months (C), 30°C for 12 months (D), and 40°C for 6 months (E). Water content of turoctocog alfa pegol dry powder samples when stored at 5°C for 18 months followed by storage at 30°C for 12 months (F), 5°C for 27 months followed by storage at 40°C for 3 months (G), 5°C for 30 months (H), 30°C for 12 months (I), and 40°C for 6 months (J). Shelf-life limits for both products are shown by the horizontal dotted lines.

Abbreviation: IU, international units.
Figure 6 HMWP of turoctocog alfa dry powder samples when stored at 5°C for 21 months followed by storage at 30°C for 9 months (A), 5°C for 27 months followed by storage at 40°C for 3 months (B), 5°C for 30 months (C), 30°C for 12 months (D), and 40°C for 6 months (E). HMWP of turoctocog alfa pegol dry powder samples when stored at 5°C for 18 months followed by storage at 30°C for 12 months (F), 5°C for 27 months followed by storage at 40°C for 3 months (G), 5°C for 30 months (H), 30°C for 12 months (I), and 40°C for 6 months (J). Shelf-life limits for both products are shown by the horizontal dotted lines.

Abbreviations: HMWP, high molecular weight protein; IU, international units.
75% RH for 12 months (Figure 6D and I) and, for all but one batch (turoctocog alfa pegol at 500 IU), <4% after storage at 40°C/75% RH for 6 months (Figure 6E and J).

It was noted that there was some variability in HMWP for both turoctocog alfa and turoctocog alfa pegol due to analytical issues encountered at low concentrations. HMWP is oligomeric forms of the monomer of turoctocog alfa and turoctocog alfa pegol, eluting ahead of the monomer in SE-HPLC.

Discussion

The ICH guidelines state that well-defined stability studies should be developed to confirm product stability during the intended storage period.30 We assessed key stability parameters for both turoctocog alfa and turoctocog alfa pegol; these parameters are susceptible to change during storage and/or could potentially influence product quality, namely oxidized forms, potency, water content, and HMWP. There were no major increases in oxidized forms or HMWP and no impairment in potency or water content for either product; most results were within each product’s range. Although FVIII activity (potency) and protein aggregates (HMWP) were largely unaffected, a small increase in oxidized forms was observed after storage at 30°C/75% RH for 12 months and after storage at 40°C/75% RH for 6 months. However, the changes in oxidized forms for both turoctocog alfa and turoctocog alfa pegol at high temperatures up to 40°C did not affect the quality of either product. Oxidized forms are inactive forms of rFVIII and a correlation between potency and oxidized forms can be expected.33 For example, oxidation of FVIII has been reported to correlate with the oxidation of free Cys and disulfide bonds, leading to structural change and loss of potency.33

Interestingly, turoctocog alfa can be subjected to variable storage conditions, including cycling between 5°C and ≤40°C, and subsequent storage for 3 months up to 40°C, without loss of stability.37 As of yet, no temperature cycling data are available for turoctocog alfa pegol. However, among commercially available EHL-rFVIII products, turoctocog alfa pegol is the only one that has shown stability at 40°C.13 Thus, the current stability data should reassure patients who are receiving turoctocog alfa and considering switching to turoctocog alfa pegol. Furthermore, as the current study shows that both products can be safely stored at temperatures up to 40°C/75% RH for 3 months without compromising product potency, patients can also be reassured that the dose on the vial label is unaffected—and therefore dose adjustments are not required—when either product is stored outside the refrigerator for several months. Patients should note the date when turoctocog alfa and turoctocog alfa pegol are taken out of the fridge so they can ensure that the duration of room-temperature storage does not exceed 3 months.

Home treatment of hemophilia is necessary and recommended: the World Federation of Hemophilia recommends acute bleeds should be treated as quickly as possible, preferably within 2 hrs.2 Patients/caregivers of those with hemophilia A have reported preferences for FVIII products that can be stored at higher temperatures for long periods of time, both overall and when travelling.15,16 However, in one small study, only 13% of patients and caregivers were aware of their product’s storage conditions,38 while another study showed that 80% of patients/caregivers who stored FVIII in the refrigerator did not know that their product may be safely stored at room temperature for a limited number of months.15 Therefore, it is important for patients to know if there is a meaningful, clinical difference between an FVIII product kept outside of the fridge for 48 hrs (for example) compared with one kept in the fridge for 3 months. Our study shows that both turoctocog alfa and turoctocog alfa pegol remained stable when stored at a relatively high ambient room temperature and humidity (40°C/75% RH) for up to 3 months following an extended storage period at 5°C. Two other rFVIII products (Advate® and Recombinate® [Takeda, Lexington, MA]) are thermostable at 30°C–40°C;39,40 however, at the time of writing and to the best of our knowledge, turoctocog alfa and turoctocog alfa pegol are the only FVIII and EHL FVIII products, respectively, licensed for storage at temperatures up to 40°C.

These data are of clinical importance, as they demonstrate that the stability of both products may offer flexibility and greater convenience in product storage and daily use. Benefits of increased storage flexibility and portability include the ability to keep more FVIII product at home, the ability to travel more easily, reduced daily disruption15,16 and, potentially, lead a more active lifestyle. The resulting improvement in convenience may also contribute to better treatment outcomes and quality of life by facilitating early treatment of bleeds15,17 and improving adherence to the recommended prophylaxis schedules. These benefits would be particularly relevant for patients who live in, or who travel to, countries where daytime temperatures are consistently or transiently above 30°C, and even up to 40°C. It is therefore crucial for hemophilia care providers to clearly explain to patients and/or
caregivers that FVIII products such as turoctocog alfa and turoctocog alfa pegol can be kept at room temperature for extended periods (assuming no higher than 40°C) while remaining clinically effective to use.

**Conclusion**

Turoctocog alfa and turoctocog alfa pegol offer stability at 40°C/75% RH for up to 3 months without jeopardizing the quality of either product. These stability characteristics may be beneficial to patients by allowing flexibility in product storage and daily use.

**Data Sharing Statement**

Novo Nordisk’s policy on data sharing may be found at [https://www.novonordisk-trials.com/how-access-clinical-trial-datasets](https://www.novonordisk-trials.com/how-access-clinical-trial-datasets).

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**Author Contributions**

All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis and interpretation, or in all these areas; took part in drafting, revising or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work.

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**Disclosure**

MN has acted as a consultant for Amgen, Bayer, BIOFVIIIx and Novo Nordisk, and received speaker fees from Baxalta, Bayer, CSL Behring, Kedrion, Novo Nordisk, Sobi, Takeda, and Octapharma; AAO and AMN are employees of Novo Nordisk A/S. The authors report no other conflicts of interest in this work.

**References**

1. Pipe S. Antihemophilic factor (recombinant) plasma/albunin-free method for the management and prevention of bleeding episodes in patients with hemophilia A. *Biologics*. 2009;3:117–125. doi:10.2147/bt.2009.2872
2. Srivastava A, Brewer A, Mauser-Bunschoten E, et al. Guidelines for the management of hemophilia. *Haemophilia*. 2013;19(1):e1–47.
3. Mancuso ME, Santagostino E. Outcome of clinical trials with new extended half-life FVIII/IX concentrates. *J Clin Med*. 2017;6(4):39. doi:10.3390/jcm604039
4. Advate [prescribing information]. Baxter Healthcare Corporation; 2011. Available from: [https://www.fda.gov/downloads/BloodB products/ucm059095.pdf](https://www.fda.gov/downloads/BloodB products/ucm059095.pdf). Accessed September 18, 2018.
5. Kogenate [summary of product characteristics]. Bayer HealthCare Manufacturing SRL; 2018. Available from: [http://www.emea.europa.eu/ema/index.jsp?curl=pages/medicines/human/medicines/000275/human_med_000879.jsp&mid=WCO601ac058001d124](http://www.emea.europa.eu/ema/index.jsp?curl=pages/medicines/human/medicines/000275/human_med_000879.jsp&mid=WCO601ac058001d124). Accessed September 28, 2018.
6. NUWIQ [prescribing information]. Octapharma AB; 2015. Available from: [https://www.fda.gov/downloads/BiologyBlood Vessels/BloodB products/ApprovedProducts/LicensedProductsBLAs/FractionatedPlasmaProducts/UCM461328.pdf](https://www.fda.gov/downloads/BiologyBlood Vessels/BloodB products/ApprovedProducts/LicensedProductsBLAs/FractionatedPlasmaProducts/UCM461328.pdf). Accessed September 28, 2018.
7. NovoEight® [summary of product characteristics]. Novo Nordik A/S; 2018. Available from: [http://www.emea.europa.eu/docs/en/GB/document_library/EPAR_-_Product_Information/human/002719/WC5001575.pdf](http://www.emea.europa.eu/docs/en/GB/document_library/EPAR_-_Product_Information/human/002719/WC5001575.pdf). Accessed August 23, 2018.
8. NovoEight®, [Highlights of prescribing information]. Available from: [https://www.novo-pi.com/novoeight.pdf](https://www.novo-pi.com/novoeight.pdf). Accessed August 10, 2018.
9. Jivi [summary of product characteristics]. Bayer AG; 2019. Available from: [https://www.ema.europa.eu/en/documents/product-information/jivi-epar-product-information_en.pdf](https://www.ema.europa.eu/en/documents/product-information/jivi-epar-product-information_en.pdf). Accessed November 14, 2019.
10. ELOCTATE [summary of product characteristics]. Swedish Orphan Biovitrum AB; 2019. Available from: [https://www.ema.europa.eu/en/documents/product-information/elocta-epar-product-information_en.pdf](https://www.ema.europa.eu/en/documents/product-information/elocta-epar-product-information_en.pdf). Accessed November 14, 2019.
11. ADYNOVATE [package insert]. Baxalta US Inc; 2019. Available from: [https://www.fda.gov/files/vaccines%20blood%20%20biolog ics/published/Package-Insert—ADYNOVATE.pdf](https://www.fda.gov/files/vaccines%20blood%20%20biolog ics/published/Package-Insert—ADYNOVATE.pdf). Accessed November 14, 2019.
12. AFSTYLA [summary of product characteristics]. CSL Behring GmbH; 2019. Available from: [https://www.ema.europa.eu/en/docs/ment/product-information/afstyla-epar-product-information_en.pdf](https://www.ema.europa.eu/en/docs/ment/product-information/afstyla-epar-product-information_en.pdf). Accessed November 14, 2019.
13. ESPEROCt. [Summary of product characteristics]. Available from: [https://www.ema.europa.eu/en/documents/product-information/esper oct-epar-product-information_en.pdf](https://www.ema.europa.eu/en/documents/product-information/esper oct-epar-product-information_en.pdf). Accessed 14 November 2019.
14. ESPEROCt [package insert]. Novo Nordik A/S; 2019. Available from: [https://www.fda.gov/media/120351/download](https://www.fda.gov/media/120351/download). Accessed November 14, 2019.
15. Tischer B, Marino R, Napolitano M. Patient preferences in the treatment of hemophilia A: impact of storage conditions on product choice. *Patient Prefer Adherence*. 2018;12:431–441. doi:10.2147/PPA.S151812
16. DiBenedetti D, Coles T, Sharma T, Pericleous L, Kulkarni R. Assessing patients’ and caregivers’ perspectives on stability of factor VIII products for haemophilia A: a web-based study in the United States and Canada. *Haemophilia*. 2014;20(4):c296–c303. doi:10.1111/hae.12459
17. Saxena K. Barriers and perceived limitations to early treatment of hemophilia. *J Blood Med*. 2013;4:49–56. doi:10.2147/JBM.S43734
18. Shannon E, Duffy J, Jones H, Paulson A, Vicik SM. Etanercept (Enbrel) alternative storage at ambient temperature. *Clin Pharmacol*. 2017;9:87–99. doi:10.2147/CPAA.S131832
19. World Health Organization. WHO Expert Committee on Specifications for Pharmaceutical Preparations. WHO Technical Report Series 953; forty-third report, 2009. Available from: http://apps.who.int/iris/bitstream/handle/10665/44553/WHO_TRS_953_eng.pdf?sequence=1. Accessed November 1, 2018.

20. Kulkarni R, Karim F, Glomocanin S, et al. Results from a large multinational clinical trial (guardian3) using prophylactic treatment with turoctocog alfa in paediatric patients with severe haemophilia A: safety, efficacy and pharmacokinetics. Haemophilia. 2015;19(5):698–705. doi:10.1111/hae.12165

21. Santagostino E, Lentz SR, Misgav M, et al. Safety and efficacy of turoctocog alfa (NovoEight®) during surgery in patients with haemophilia A: results from the multinational guardian clinical trials. Haemophilia. 2015;21(1):34–40. doi:10.1111/hae.12518

22. Lentz S, Misgav M, Ozelo M, et al. Results from a large multinational clinical trial (guardian1) using prophylactic treatment with turoctocog alfa in adolescent and adult patients with severe haemophilia A: safety and efficacy. Haemophilia. 2013;19(5):691–697. doi:10.1111/hae.12159

23. Curry N, Albayрак C, Escobar M, et al. Once-weekly prophylaxis with glycoPEGylated recombinant factor VIII (N8-GP) in severe haemophilia A: safety and efficacy results from pathfinder 2 (randomized Phase III trial). Haemophilia. 2019;25(3):373–381. doi:10.1111/hae.13712

24. Giaigrande P, Andreva T, Chowdary P, et al. Clinical evaluation of glycoPEGylated recombinant FVIII: efficacy and safety in severe haemophilia A. Thromb Haemost. 2017;117(2):252–261. doi:10.1160/TH16-06-0444

25. Hampton K, Chowdary P, Dunkley S, et al. First report on the safety and efficacy of an extended half-life glycoPEGylated recombinant FVIII for major surgery in severe haemophilia A. Haemophilia. 2017;23(5):689–696. doi:10.1111/hae.13246

26. Meunier S, Alamelu J, Ehrenfors S, et al. Safety and efficacy of a glycoPEGylated rFVIII (turoctocog alpha pegol, N8-GP) in paediatric patients with severe haemophilia A. Thromb Haemost. 2017;117(9):1705–1713. doi:10.1160/TH17-03-0166

27. Stennicke HR, Kjkalke M, Karpf DM, et al. A novel B-domain O-glycoPEGylated FVIII (N8-GP) demonstrates full efficacy and prolonged effect in hemophilic mouse models. Blood. 2013;121(11):2108–2116. doi:10.1182/blood-2012-01-407494

28. Elgaard TW, Bindshelf L, Kamstrup S. Evaluation of the virus clearance and robustness of the manufacturing process for the recombinant factor VIII protein, turoctocog alfa. Protein Expr Purif. 2017;129:94–100. doi:10.1016/j.pep.2016.09.002

29. Agero H, Stennicke HR, Pelzer H, et al. Pharmacokinetics and pharmacodynamics of turoctocog alfa and N8-GP in haemophilia A dogs. Haemophilia. 2012;18(6):941–947. doi:10.1111/j.1365-2516.2012.02896.x

30. International Conference on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use. ICH Harmonised Tripartite Guideline: quality of Biotechnological Products: stability Testing of Biotechnological/Biological Products Q5C. Current Step 5 version. Available from: https://www.ema.europa.eu/en/documents/scientific-guideline/ich-topic-q-s-c-quality-biotechnological-products-stability-testing-biotechnological/biotechnological-products_en.pdf. Accessed May 19, 2020.

31. European Pharmacopoeia. 10th Edition. Available from: https://www.ephap.eu/en/european-pharmacopoeia-ph-eur-10th-edition. Accessed May 19, 2020.

32. Eldin A, Shalaby A. Comparison of FT-NIR transmission and HPLC to assay montelukast in its pharmaceutical tablets. Am J Analytical Chem. 2011;2(8):885–891. doi:10.4236/ajac.2011.28102

33. Wang W, Wang YJ, Kelner DN. Coagulation factor VIII: structure and stability. Int J Pharm. 2003;259(1–2):1–15. doi:10.1016/S0378-5173(03)00227-8

34. Butenas S, Parhami-Seren B, Gissel MT, Gomperts ED, Fass DN, Mann KG. Potency and mass of factor VIII in FVII products. Haemophilia. 2009;15(1):63–72. doi:10.1111/j.1365-2516.2008.01826.x

35. Yoshioka S, Aso Y. Correlations between molecular mobility and chemical stability during storage of amorphous pharmaceuticals. J Pharm Sci. 2007;96(5):960–961. doi:10.1002/jps.20926

36. Rosenberg AS. Effects of protein aggregates: an immunologic perspective. AAPS J. 2006;8(3):E501–507. doi:10.1208/aaps080359

37. Napolitano M, Nohr AM. The effect of fluctuating temperature on the stability of turoctocog alfa for hemophilia A. Drugs R D. 2019;19(4):381–390. doi:10.1007/s40268-019-00290-3

38. Novais T, Duclos A, Varin R, Lopez I, Chamouard V. Treatment-related knowledge and skills of patients with haemophilia and their informal caregivers. Int J Clin Pharm. 2016;38(1):61–69. doi:10.1007/s11096-015-0207-z

39. Parti R, Ardosa J, Yang L, Mankarious M. In vitro stability of recombinant human factor VIII (Recombinate). Haemophilia. 2000;6(5):513–522. doi:10.1046/j.1365-2516.2000.00410.x

40. Parti R, Schoppmann A, Lee H, Yang L. Stability of lyophilized and reconstituted plasma/albumin-free recombinant human factor VIII (ADVATE rAHF-PFM). Haemophilia. 2005;11(5):492–496. doi:10.1111/j.1365-2516.2005.01128.x

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