## Move beyond the threshold<sup>a</sup>

Esperoct<sup>®</sup> can give you high factor levels for longer.<sup>b</sup>

<sup>a</sup>Of 1% trough factor levels for standard half-life (SHL) products in adults and adolescents. <sup>b</sup>Compared with SHL products.

#### WHAT IS ESPEROCT®?

Esperoct<sup>®</sup> [antihemophilic factor (recombinant), glycopegylated-exei] is an injectable medicine to treat and prevent or reduce the number of bleeding episodes in people with hemophilia A. Your healthcare provider may give you Esperoct<sup>®</sup> when you have surgery

• Esperoct<sup>®</sup> is not used to treat von Willebrand Disease



Please see Important Safety Information throughout. Please see accompanying Prescribing Information.

#### **esperoct**®

### Extend half-life **beyond the standard**

### **22-hour** average half-life in adults<sup>a</sup>



#### Esperoct<sup>®</sup> 50 IU/kg Standard rFVIII

rFVIII=recombinant factor VIII.

<sup>a</sup>Data shown are from 42 adults who received a pharmacokinetic (PK) assessment around the first Esperoct<sup>®</sup> 50 IU/kg dose. <sup>b</sup>Data shown are from a comparison study of 26 previously treated patients (PTPs) 18 years or older who received a 25, 50, or 75 IU/kg dose of their previous SHL product followed by the same dose of Esperoct<sup>®</sup>. To allow for comparison, all results were adjusted to a 50 IU/kg dose of each product.

Esperoct<sup>®</sup> is made by taking the existing Novoeight<sup>®</sup> (rFVIII) molecule and adding **PEGylation technology to extend the half-life** 

#### **IMPORTANT SAFETY INFORMATION**

#### Who should not use Esperoct<sup>®</sup>?

• You should not use Esperoct<sup>®</sup> if you are allergic to factor VIII or any of the other ingredients of Esperoct<sup>®</sup> or if you are allergic to hamster proteins

Please see additional Important Safety Information throughout.

Please see accompanying Prescribing Information.

### Level up prophylaxis with a simple dose

High factor levels from one dose to the next



<sup>c</sup>Trough level goal is 1% for prophylaxis.

<sup>d</sup>Data shown are from a study where 175 previously treated adolescents and adults received routine prophylaxis with Esperoct® 50 IU/kg every 4 days for 76 weeks. Pre-dose factor activity (trough) levels were evaluated at follow-up visits. Mean trough levels for adolescents (12-<18 years) were 2.7 IU/dL.

eSteady-state factor VIII (FVIII) activity levels were estimated in 143 adults and adolescents using PK modeling.

### esperoct<sup>®</sup>

### Stay protected from bleeds

Dose less often<sup>a</sup> without sacrificing protection



<sup>a</sup>Compared to SHL products, 50% fewer infusions when administered every other day and 40% fewer when administered 3x weekly. <sup>b</sup>175 PTPs with severe hemophilia A received Esperoct<sup>®</sup> 50 IU/kg every 4 days for 76 weeks based on median annualized bleed rates shown.

#### **IMPORTANT SAFETY INFORMATION**

#### What is the most important information I need to know about Esperoct<sup>®</sup>?

- Do not attempt to do an infusion yourself unless you have been taught how by your healthcare provider or hemophilia treatment center
- Call your healthcare provider right away or get emergency treatment right away if you get any signs of an

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### **Prepare** for the unexpected

Control for the treatment of bleeding episodes



# Dosing for the treatment of bleeding episodes

40 IU/kg for minor to moderate bleeds 50 IU/kg for major bleeds

For moderate to major bleeds, additional dose(s) may be administered every 24 hours

<sup>c</sup>Data shown are from a study where 12 adult and adolescent PTPs with severe hemophilia chose to be treated on demand and received Esperoct<sup>®</sup> for 532 bleeding episodes.

**esperoct**<sup>®</sup> antihemophilic factor (recombinant), glycopegylated-exei

### **Switching** made easy

One standard dose makes it easy to switch at any age

# 65 IU/kg twice weekly

No dose adjustment needed<sup>a</sup>

Because FVIII products may be cleared from the body faster in children <12 years, higher and more frequent dosing may be needed.

<sup>a</sup>Interval can be adjusted based on individual response to treatment

#### **IMPORTANT SAFETY INFORMATION**

#### What should I tell my healthcare provider before using Esperoct<sup>®</sup>?

• Before taking Esperoct<sup>®</sup>, you should tell your healthcare provider if you have or have had any medical conditions, take any medicines (including non-prescription medicines and dietary supplements), are nursing, pregnant or planning to become pregnant, or have been told that you have inhibitors to factor VIII

Please see additional Important Safety Information throughout.

Please see accompanying Prescribing Information.

Fewer infusions per year compared with SHL dosing regimens





### **Protection** that keeps up with them

Esperoct<sup>®</sup> achieved a

14.3-hour average half-life in children<sup>b</sup> **85% longer** half-life compared to SHL<sup>c</sup>

Low number of bleeds per year for children aged 0-<12 years<sup>d</sup>

**2** Overall bleeds per vear<sup>d</sup>

Joint bleeds Spontaneous bleeds

<sup>b</sup>Comparison to prior FVIII product was performed at the beginning of the study in previously treated children. The geometric mean terminal half-life in 23 children aged 0-<12 years was 14.3 hours. Esperoct<sup>®</sup> geometric mean terminal half-life was 14.7 hours in 12 children ages 0-5 and 13.8 hours in 10 children ages 6-11. <sup>c</sup>Comparison to prior FVIII product was performed at the beginning of the study in previously treated children. Esperoct® half-life was 14.7 hours in 12 children ages 0-6 and 13.8 hours in 10 children ages 6-11.

<sup>d</sup>Data shown are from a study of 68 previously treated children (34 aged 0-5 and 34 aged 6-11) who received an average dose of approximately 65 IU/kg twice weekly for 26 weeks. Median annualized bleeding rates are shown.

### esperoct<sup>®</sup>

### Count on a **proven safety profile**

previously treated



### patients (PTPs)

#### Safety proven across 5 studies, the largest and longest EHL clinical trial program

- 0 blood clots
- No PEG-related safety concerns
- One PTP with a high-risk gene mutation developed an inhibitor to FVIII<sup>a</sup>
- Similar to the reported rate in patients with severe hemophilia A

EHL=extended half-life; PEG=polyethylene glycol.

<sup>a</sup>An 18-year-old African-American male developed an inhibitor after 93 infusion days of Esperoct<sup>®</sup>. The inhibitor rose to 13.5 Bethesda units and the patient stopped participation in the study. There was no change in efficacy, and the inhibitor eventually went away on its own (without use of immune tolerance induction therapy).

#### **IMPORTANT SAFETY INFORMATION**

#### What should I tell my healthcare provider before using Esperoct<sup>®</sup>? (cont'd)

- Your body can make antibodies called "inhibitors" against Esperoct<sup>®</sup>, which may stop Esperoct<sup>®</sup> from working properly. Call your healthcare provider right away if your
- bleeding does not stop after taking Esperoct<sup>®</sup>

### Designed to **fit into your life**



Please see additional Important Safety Information throughout. Please see accompanying Prescribing Information.

#### esperoct<sup>®</sup>

### Flexible on the go

The EHL product with the highest storage temperature for the longest time



#### **IMPORTANT SAFETY INFORMATION**

What are the possible side effects of Esperoct<sup>®</sup>?

• Common side effects of Esperoct<sup>®</sup> include rash or itching, and swelling, pain, rash or redness at the location of infusion 10

### Ready in **3 simple steps**

A prefilled syringe provides convenient administration in 2 minutes

### Attach

Prefilled diluent syringe contains 4 mL of diluent—works with any dose strength

### Twist

Adapter connects the syringe and vial with a 25 µm inline particle filter

### Mix

After mixing, the reconstituted solution can be administered

**5 vial sizes** for personalized treatment



Please see additional Important Safety Information throughout. Please see accompanying Prescribing Information.

#### esperoct<sup>®</sup>

### Considerations when switching to Esperoct®

You may be eligible for:

### Trial Program

Talk to a NovoSecure<sup>™</sup> specialist to find out if you're eligible<sup>a</sup>

### Product Assistance Program

Apply for the Product Assistance Program by calling 1-844-NOVOSEC (1-844-668-6732) for more information<sup>b</sup>

### **Co-pay Assistance Program**

Get help with co-pay costs for Esperoct<sup>®c</sup>

<sup>a</sup>Patients who have been prescribed a Novo Nordisk hemophilia and rare bleeding disorder product for an FDA-approved indication, and who have commercial insurance, may be eligible to receive a limited supply of free product. Patients who participate in any government, state, or federally funded medical or prescription benefit programs, including Medicare, Medicaid, Medigap, VA, DOD, and TRICARE, including patients who participate in a managed Medicaid program or have Medicaid as secondary insurance, are not eligible to receive product support. Product is provided at no cost to the patient or the HCP, is not contingent on any product purchase, and the patient and HCP must not: (1) bill any third party for the free product, or (2) resell the free product.

<sup>b</sup>The Novo Nordisk Patient Assistance Program (PAP) is administered by NovoSecure<sup>TM</sup>. To qualify for the PAP, patients must demonstrate financial need and must have attempted to find alternative reimbursement. Several factors are considered in evaluating financial need, including cost of living, size of household, and burden of total medical expenses. If the applicant qualifies under the PAP guidelines, a limited supply of the requested medication(s) will be shipped to the patient. Patients who qualify for PAP may be eligible to receive the prescribed Novo Nordisk product, for up to 1 year from the approval date. Product limits vary.

<sup>c</sup>Novo Nordisk Hemophilia and Rare Bleeding Disorders Copay/Coinsurance Terms and Conditions: Enrolled patients are eligible for up to \$12,000 in co-pay/coinsurance assistance per calendar year for each NNI hemophilia or rare bleeding disorder product. Assistance is retroactive to 60 days. Patients must be commercially insured and may not be participating in any government, state, or federally funded medical or prescription benefit programs, including Medicare, Medicaid, Medigap, VA, DOD, and TRICARE, including patients who participate in a managed Medicaid program or have Medicaid as secondary insurance. Uninsured, cash-paying patients are not eligible to participate. Patients are eligible to receive co-pay/coinsurance assistance on an annual basis (12 months). Offer good only in the USA, Puerto Rico, Guam, Saipan, and Virgin Islands with participating pharmacies and cannot be redeemed at government-subsidized clinics. Void where taxed, restricted, or prohibited by law. Absent of a change in Massachusetts law, effective July 1, 2019, the Savings Card will no longer be valid for residents of Massachusetts. Patient is responsible for complying with any insurance carrier co-payment disclosure requirements, including disclosing any savings received from this program. Re-confirmation of information may be requested periodically to ensure accuracy of data and compliance with terms. This is not an insurance program. Novo Nordisk reserves the right to rescind, revoke, or amend this offer without notice at any time. Non-medication expenses, such as ancillary supplies or administration-related costs, are not eligible. Must have a current prescription for an FDA-approved indication.

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- 12 if you are allergic to hamster proteins



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### esperoct<sup>®</sup>

# Move beyond the threshold<sup>a</sup>

# Fewer infusions per year compared with SHL for adults and adolescents

- 50% fewer infusions if previously infused every other day
- 40% fewer infusions if previously infused 3x/week

#### High factor levels in adults and adolescents<sup>b</sup>

- At or above 3% for 100% of the time<sup>c</sup>
- At or above 5% for 90% of the time<sup>d</sup>

#### Flexible on the go<sup>a</sup>

 $\bullet$  The only EHL product with stability up to  $104^{\circ}F^{\rm e}$ 

#### The largest and longest EHL clinical trial program

°Of 1% trough factor levels for SHL products in adults and adolescents.

<sup>b</sup>Data shown are from 42 adults who received a PK assessment around the first Esperoct<sup>®</sup> 50 IU/kg dose. <sup>c</sup>Data shown are from a study where 175 previously treated adolescents and adults received routine prophylaxis with Esperoct<sup>®</sup> 50 IU/kg every 4 days for 76 weeks. Pre-dose factor activity (trough) levels were evaluated at follow-up visits. Mean trough levels for adolescents (12-<18 years) were 2.7 IU/dL. <sup>d</sup>Steady-state FVIII activity levels were estimated in 143 adults and adolescents using PK modeling. <sup>e</sup>For up to 3 months.

#### **IMPORTANT SAFETY INFORMATION**

#### What is the most important information I need to know about Esperoct<sup>®</sup>?

• Do not attempt to do an infusion yourself unless you have been taught how by your healthcare provider or hemophilia treatment center

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#### Please see accompanying Prescribing Information.



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### esperoct<sup>®</sup>