

IXINITY® [coagulation factor IX (recombinant)] Free Trial Request Form

Fax both sides of completed form to 1-888-747-9329

New patients are eligible for 1 free month of IXINITY up to 20,000 IU. The IXINITY Trial Program is available only to those patients who have not previously enrolled in this program and are not currently using IXINITY.

Instructions:

1. Complete this side of the form.
2. If requesting Custom Ancillaries to be shipped with the Free Trial, please indicate selections on the reverse side of this form.
3. Fax both sides of the completed form to 1-888-747-9329 or email to IXINITY@thealliancepharmacy.org.

Important: This form must be filled out completely and signed by your healthcare professional or it will not be processed. Your Free Trial product and Custom Ancillaries will be shipped via overnight courier directly to the patient's or physician's address of choice as indicated below:

Please ship to (select one): Patient's address Physician's address

Patient Information:

First Name: _____ Last Name: _____ DOB: _____

Contact Phone #: _____ Email Address: _____

Please include your phone number and email address so shipment arrangements can be confirmed.

Shipping Address (No PO Boxes): _____

City: _____ State: _____ Zip Code: _____

Primary Health Insurance Provider: _____

I would like to receive Custom Ancillaries with my Free Trial Shipment (select one):

Yes Please indicate your selections on the back of this form. No

Important: Your answers to the following questions do not disqualify you from participation in the IXINITY® Free Trial or Custom Ancillary Programs.

1. I authorize an independent, third party to contact me for a follow-up survey about my experience with this program (select one): Yes No
2. I authorize the administrator of this program to share my email address with Aptevio BioTherapeutics LLC so I may receive information on product updates and new developments (select one): Yes No

Prescriber Information:

Physician's Name: _____ Facility Name: _____

State License #: _____

Physician's Address: _____

City: _____ State: _____ Zip Code: _____

Contact name for this product request: _____

Phone #: _____ Email Address: _____

Prescription Information:

Patient weight: _____ lb _____ kg Baseline FIX: _____ % Target FIX activity desired: _____ %

Total IXINITY IU required for 1 dose: _____ Number of doses requested: _____ (Max 1 month up to 20,000 IU)

Special Instructions: _____

Authorized refills - 0. The free trial prescription is valid for one time only with no refills.

Prescriber Authorization: I hereby verify that, to my knowledge, the above patient has no treatment history with the brand-named product requested. This trial product will not be exported or transferred in exchange for money, other property, or services. No portion of this trial product will be used for reimbursement purposes from Medicaid/Medicare or any other third-party program, which provides cost or charge-based reimbursement to the participating institution, either directly or indirectly.

Physician/Prescriber Signature: _____ Date: _____

NPI #: _____

Patient Last Name: _____

IXINITY® [coagulation factor IX (recombinant)] Free Trial Request Form

Fax both sides of completed form to 1-888-747-9329

Custom Ancillary supplies are available with IXINITY at no additional cost to the patient or physician. Following the Free Trial, patients will be contacted to confirm their supplies selections. Patients are eligible to receive Custom Ancillaries as long as they remain on IXINITY.

Instructions:

1. Check 1 selection for each category of ancillary supplies below.
2. If your preferred item is not listed, please check "Other" and describe it in detail. We will do our best to accommodate your request. However, availability of specially requested items is not guaranteed.
3. To make changes to your ancillary supply selections at any time, please call **1-855-IXINITY**.

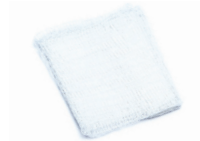
Winged Infusion Sets



- Wing Infusion Set Long 12" 23 gauge
- Wing Infusion Set Long 12" 25 gauge
- Wing Infusion Set Short 12" 23 gauge
- Wing Infusion Set Short 12" 25 gauge

Other _____

Gauze



- Sponge Gauze 8 Ply Sterile 2" x 2"
- Sponge Gauze 8 Ply Sterile 4" x 4"

Other _____

Bandages



- Adhesive Strip Sheer Plastic $\frac{1}{2}$ " x 3"
- Adhesive Bandage Woven $\frac{3}{4}$ " x 3"
- Bandage Adhesive Spot Oval Coverlet 1 $\frac{1}{4}$ "

Other _____

Extra Syringes



- 5 mL
- 10 mL
- 20 mL

Other _____

Flex Wrap



- Bandage Cohesive Flex Wrap 2" Wide
- Bandage Cohesive 3" Wide

Other _____

Tourniquet



- Seraket® Automatic Tourniquet by Proper
- Traditional velcro tourniquet
- Traditional elastic tourniquet

Please Include

Sterile Alcohol Prep Pads

Disposable Infusion Mats

Seraket® is a registered trademark of Proper Manufacturing Co., Inc.

Aptevo BioTherapeutics LLC, Berwyn, PA 19312

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CM-FIX-0035



12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Hemophilia B is a sex-linked hereditary disorder of blood coagulation caused by a deficiency in factor IX and results in bleeding into joints, muscles or internal organs, either spontaneously or as a result of accidental or surgical trauma. Treatment with IXINITY replaces factor IX, thereby enabling a temporary correction of the factor deficiency and correction of the bleeding tendencies.

12.2 Pharmacodynamics

The administration of IXINITY increases plasma levels of factor IX and can temporarily correct the coagulation defect in these patients, as reflected by decrease in the aPTT.

12.3 Pharmacokinetics

Pharmacokinetic studies with IXINITY were conducted in 32 previously treated patients (PTPs) \geq 12 years of age with severe to moderately severe hemophilia B (factor IX \leq 2 IU/dL). Intravenous administration of 75 \pm 5 IU/kg of IXINITY to 32 PTPs showed an initial recovery ranging from 51 to 113% (median 70%). The results of pharmacokinetic studies are summarized below in Table 4.

Table 4 Pharmacokinetic Parameters for IXINITY (n = 32)

Parameters	Mean (\pm SD) (Range)
AUC_{0-∞} (IU/dL/hr)	1573 (\pm 451) (862-2643)
Incremental Recovery (IU/dL per IU/kg)	0.98 (\pm 0.21) (0.67-1.50)
Terminal Half-life (hours)	24 (\pm 7) (13-43)
C_{max} (IU/dL)	73 (\pm 17) (51-113)
Mean Residence Time (hours)	32 (\pm 6) (19-47)
VD_s (mL/kg)	175 (\pm 57) (102-314)
Clearance [mL/(kg-hr)]	5.1 (\pm 1.3) (2.8-7.7)

Pharmacokinetic parameters were re-assessed in a subset of 14 subjects after routine treatment with IXINITY for a median of 5.8 months (range 3.1 to 18.6 months) as summarized in Table 5 below.

Table 5 Pharmacokinetic Parameters for IXINITY Following Repeat-Dosing (n = 14)

Parameters	Initial Mean (\pm SD)	Repeat-Dosing PK Mean (\pm SD)
AUC_{0-∞} (IU/dL/hr)	1438 (\pm 409)	1530 (\pm 435)
Incremental Recovery (IU/dL per IU/kg)	0.96 (\pm 0.22)	0.95 (\pm 0.18)
Terminal Half-life (hours)	24 (\pm 7)	24 (\pm 6)
C_{max} (IU/dL)	73 (\pm 16)	73 (\pm 15)
Mean Residence Time (hours)	30 (\pm 6)	31 (\pm 5)
VD_s (mL/kg)	193 (\pm 62)	185 (\pm 70)
Clearance [mL/(kg-hr)]	5.6 (\pm 1.3)	5.3 (\pm 1.5)

Repeat dosing did not impact the pharmacokinetics of IXINITY.

The PK data were divided into two subgroups of subjects with a BMI \leq 30 (n = 26) or BMI > 30 (n = 6). The AUC_{0-∞} and C_{max} values of IXINITY were 40% and 34% higher, respectively, in subjects with BMI > 30.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

No macroscopic or microscopic pathologies in reproductive organs were observed in repeated dose toxicity studies of IXINITY in animals. Animal studies regarding impairment of fertility were not conducted.

No nonclinical investigations of genotoxicity, carcinogenicity, or toxicity to reproduction and development have been conducted with IXINITY.

14 CLINICAL STUDIES

The efficacy of IXINITY was evaluated in a prospective, open-label, uncontrolled multicenter study in which a total of 77 subjects (76 male, 1 female carrier in surgery study) were exposed to IXINITY for treatment of hemophilia B or for perioperative management. All male subjects either had severe or moderately severe (factor IX level \leq 2 IU/dL) hemophilia B, or had factor IX levels between 2-8 IU/dL and clinically severe hemophilia B with recurrent hemarthroses and required surgery (n = 3 in surgery study, 1 continued to treatment phase). Previously treated patients (PTPs) were defined as patients with a minimum of 150 exposures to another factor IX preparation. Of the 77 subjects, 68 PTPs between 7 and 64 years of age received IXINITY either as routine or on-demand treatment. Routine treatment was defined as PTPs who received a starting dose of 40-70 international units (IU) per kg twice weekly. Excluded from the study were patients with a history of a detectable factor IX inhibitor (\geq 0.6 BU), a history of hypersensitivity reactions following exposure to factor IX-containing products, a known allergic reaction to hamster proteins, evidence of severe liver impairment, evidence of impaired renal function, CD4 count < 400 cells/mm³, or any coagulation defect other than hemophilia B. In addition, there was a prospective, open-label, uncontrolled, multicenter substudy where 17 subjects (16 male, 1 female carrier) underwent surgeries (19 major procedures in males) receiving IXINITY for perioperative management; some of the surgery subjects also participated in the treatment trial.

Of the 68 PTPs in the treatment group, subjects were primarily prescribed a routine (n = 58) or an on-demand regimen (n = 9); one subject was not assigned a regimen. Subjects were allowed to switch regimens during the course of the study. As a result, 61 subjects were treated at some point with routine treatment and 12 were treated at some point with an on-demand regimen. Subjects in the routine therapy group received mean intravenous doses of 55 \pm 12.8 IU/kg of IXINITY twice weekly. Subjects in the on-demand therapy group received mean doses of 60 \pm 18.2 IU/kg (median 59.3, interquartile range 49.9, 71.8) for bleeding episodes. The mean number of exposure days (ED) was 138.2 (median 127.5), including 45 subjects with \geq 100 ED and 55 subjects with \geq 50 ED. Median duration on study for the on-demand group was 14.1 months (range 2.3-36.9).

Control and Prevention of Bleeding Episodes

A total of 508 bleeding episodes were treated with IXINITY, of which 286 bleeds were recorded for subjects treated with the routine treatment regimen and 222 with the on-demand regimen. Bleeding resolved in 360 episodes (70.9%) after a single infusion of IXINITY and in 66 (13.0%) episodes after two infusions. For 24 bleeding episodes (4.7%), five or more infusions were required; these 24 bleeding episodes were predominantly related to trauma, target joints, or muscle bleeds.

Hemostatic efficacy to resolve a bleed was rated by subjects as excellent or good in 84% of treated bleeding episodes. Excellent was defined as a dramatic response with abrupt pain relief and clear reduction in joint or hemorrhage site size, and good was defined as pain relief or reduction in hemorrhage size that may have required an additional infusion for resolution.

Perioperative Management

The efficacy analysis of IXINITY in perioperative management included 19 major surgeries performed in 16 male PTPs between 12 and 56 years of age (female carrier not included in efficacy analysis). Efficacy of IXINITY for support of major surgery was based on the surgeon's assessment of efficacy including: a) at the time of surgery as estimation of blood loss as 'less than expected', 'expected', or 'more than expected'; and b) at 12 and 24 hours post-surgical assessments of hemostasis as 'adequate', 'better than adequate', or 'poorly controlled'. Transfusion requirements to support surgery were also monitored. There were no transfusions required during the procedures.

IXINITY was administered during major surgical procedures as bolus (n = 13) or continuous infusion (n = 6). IXINITY was rated as adequate or better in controlling hemostasis post-surgery as assessed by the surgeon when used in various procedures, including, knee arthroplasty (n = 8), elbow arthroplasty (n = 2), knee amputation (n = 1), percutaneous Achilles tendon lengthening (n = 1), open inguinal hernia repair (n = 1), tibiotalar fusion (n = 1), arthroscopic synovectomy (n = 2), and debridement (ankle, knee) (n = 3). In all instances, blood loss at surgery was 'expected' or 'less than expected' as assessed by the surgeon.

15 REFERENCES

1. Srivastava A, et al. World Federation of Hemophilia, Guidelines for the management of hemophilia. Haemophilia. 2013; 19:e1-e47.
2. Ingerslev J, Christiansen K, Ravn HB, et al. Antibodies to heterologous proteins in hemophilia A patients receiving recombinant factor VIII (Recombinate™). Thromb Haemost. 2002; 87:626-634.

16 HOW SUPPLIED/STORAGE AND HANDLING

IXINITY is supplied as a lyophilized powder in single-use glass vials containing the labeled amount of factor IX activity, expressed in international units (IU). The actual factor IX activity in IU is stated on the label of each vial.

Kits include one single-use vial (containing nominally 250, 500, 1000, 1500, 2000, or 3000 IU per vial), a 10 mL syringe pre-filled with 5 mL of Sterile Water for Injection with plunger rod attached, and a vial adapter with filter. None of the kit components are made with natural rubber latex.

Color Code	Nominal Strength	Kit NDC Number
Yellow	250 IU	70504-0287-5
Blue	500 IU	70504-0282-5
Green	1000 IU	70504-0283-5
Orange	1500 IU	70504-0284-5
Red	2000 IU	70504-0288-5
Brown	3000 IU	70504-0289-5

250 IU strength only; store at 2 to 8°C (36 to 46°F).

500, 1000, 1500, 2000, and 3000 IU strengths; store at 2 to 25°C (36 to 77°F).

Do not freeze.

Keep the vial in the carton and protect from light.

Infuse reconstituted solution immediately or within 3 hours of storage at room temperature after reconstitution. Do not refrigerate after reconstitution.

17 PATIENT COUNSELING INFORMATION

- Advise patients to read the FDA-approved patient labeling (Patient Information and Instructions for Use).
- Inform patients of the early signs of hypersensitivity reactions (including hives, generalized urticaria, chest tightness, wheezing, and hypotension) and anaphylaxis. Instruct patients to discontinue use of the product and contact their physician if these symptoms occur.
- Advise patients to contact their physician or treatment facility for further treatment and/or assessment if they experience a lack of clinical response to factor IX replacement therapy, as in some cases this may be a manifestation of an inhibitor.

Manufactured by:

Aptevo BioTherapeutics LLC

Berwyn PA, 19312

U.S. License No. 2054

Patient Information

IXINITY® [coagulation factor IX (recombinant)]

This leaflet summarizes important information about IXINITY. Please read it carefully before using this medicine. This information does not take the place of talking with your healthcare provider, and it does not include all of the important information about IXINITY. If you have any questions after reading this, ask your healthcare provider.

What is IXINITY?

IXINITY is a medicine used to replace clotting factor (factor IX) that is missing in people with hemophilia B. Hemophilia B is also called congenital factor IX deficiency or Christmas disease. Hemophilia B is an inherited bleeding disorder that prevents clotting.

Your healthcare provider may give you IXINITY when you have surgery.

Who should not use IXINITY?

You should not use IXINITY if you:

- Are allergic to hamsters
- Are allergic to any ingredients in IXINITY

Tell your healthcare provider if you are pregnant or breastfeeding because IXINITY may not be right for you.

What should I tell my healthcare provider before using IXINITY?

You should tell your healthcare provider if you:

- Have or have had any medical problems
- Take any medicines, including prescription and non-prescription medicines, such as over-the-counter medicines, supplements, or herbal remedies
- Have any allergies, including allergies to hamsters
- Are breastfeeding. It is not known if IXINITY passes into your milk and if it can harm your baby
- Are pregnant or planning to become pregnant. It is not known if IXINITY may harm your baby
- Have been told that you have inhibitors to factor IX (because IXINITY may not work for you)

How should I infuse IXINITY?

IXINITY is given directly into the bloodstream. IXINITY should be administered as ordered by your healthcare provider. You should be trained on how to do infusions by your healthcare provider or hemophilia treatment center. Many people with hemophilia B learn to infuse their IXINITY by themselves or with the help of a family member.

See the step-by-step guide (Instructions for Use) provided at the end of this leaflet.

Your healthcare provider will tell you how much IXINITY to use based on your weight, the severity of your hemophilia B, and where you are bleeding. You may have to have blood tests done after getting IXINITY to be sure that your blood level of factor IX is high enough to stop the bleeding. Call your healthcare provider right away if your bleeding does not stop after taking IXINITY.

What are the possible side effects of IXINITY?

Allergic reactions may occur with IXINITY. Call your healthcare provider or get emergency treatment right away if you get any of the following symptoms: rash, hives, itching, tightness of the throat, chest pain or tightness, difficulty breathing, lightheadedness, dizziness, nausea, or fainting.

Tell your healthcare provider about any side effect that bothers you or does not go away.

The most common side effect of IXINITY in clinical trials was headache.

These are not all the side effects possible with IXINITY. You can ask your healthcare provider for information that is written for healthcare professionals.

Call your healthcare provider for medical advice about side effects. You may report side effects to the FDA at 1-800-FDA-1088.

What are the IXINITY dosage strengths?

IXINITY comes in vials containing six different dosage strengths: 250, 500, 1000, 1500, 2000 and 3000 international units (IU). The actual strength will be printed on the label of the vial and on the box. The six different strengths in the vials are color coded as follows:

Color Code	Nominal Strength
Yellow	250 IU
Blue	500 IU
Green	1000 IU
Orange	1500 IU
Red	2000 IU
Brown	3000 IU

Always check the actual dosage strength printed on the label to make sure you are using the strength prescribed by your healthcare provider.

How should I store IXINITY?

250 IU strength only; store at 2 to 8°C (36 to 46°F). Do not freeze.

500, 1000, 1500, 2000, and 3000 IU strengths; store at 2 to 25°C (36 to 77°F). Do not freeze.

Do not use IXINITY after the expiration date printed on the label. Throw away any unused IXINITY and diluents after it reaches this date.

Reconstituted product (after mixing dry product with Sterile Water for Injection) must be used within 3 hours and cannot be stored or refrigerated. Discard any IXINITY left in the vial at the end of your infusion.

What else should I know about IXINITY?

Your body may form inhibitors to factor IX. An inhibitor is part of the body's immune system. If you form inhibitors, it may stop IXINITY from working properly. Consult with your healthcare provider to make sure you are carefully monitored with blood tests to check for the development of inhibitors to factor IX. Consult your doctor promptly if bleeding is not controlled with IXINITY as expected.

Medicines are sometimes prescribed for purposes other than those listed here. Do not use IXINITY for a condition for which it is not prescribed. Do not share IXINITY with other people, even if they have the same symptoms as you.

Resources available to patients

For information on patient assistance programs that may be available to you, please call our IXINITY Patient Care Center at 1-855-IXINITY (1-855-494-6489).

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Instructions for Use

IXINITY [coagulation factor IX (recombinant)]

For intravenous use after reconstitution only

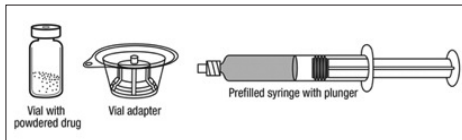
Do not attempt to do an infusion to yourself unless you have been taught how by your healthcare provider or hemophilia center.

Always follow the specific instructions given by your healthcare provider. The steps listed below are general guidelines for using IXINITY. If you are unsure of the procedures, please call your healthcare provider before using IXINITY. Your healthcare provider will prescribe the dose that you should take.

Before starting reconstitution and administration you will need the following items:

- One (or more) vial(s) of IXINITY 250, 500, 1000, 1500, 2000, or 3000 IU powder, as prescribed by your healthcare provider
- One (or more) 10 mL syringe(s), pre-filled with 5 mL of Sterile Water for Injection (Pre-filled Syringe) with plunger rod attached
- Sterile vial adapter with filter
- One sterile LUER-LOK syringe (Administration Syringe); additional or larger syringes may be required if pooling multiple vials
- Sterile alcohol swabs
- Sterile infusion set
- Sterile gauze pad
- Sterile bandage

IXINITY is supplied in kits that include single-use vials which contain vials of IXINITY (250, 500, 1000, 1500, 2000, or 3000 IU of powder), a 10 mL syringe pre-filled with 5 mL of Sterile Water for Injection with plunger rod attached (to be used for reconstitution only), and a sterile vial adapter with filter.



RECONSTITUTION INSTRUCTIONS

Wash your hands and then clean a flat area before starting the steps for reconstituting IXINITY. Use an aseptic technique during reconstitution.

1. Remove the Pre-filled Syringe and IXINITY vial from storage and allow them to reach room temperature before use. Check the expiration date on the IXINITY vial.
2. Remove the plastic cap from the IXINITY vial and place the vial top up on the clean surface. You will see a rubber circle on the top of the vial.
3. Wipe the top of the IXINITY vial with a sterile alcohol swab and allow it to dry. After cleaning, do not touch the rubber circle with your hands or allow it to touch another object.
4. Peel back the paper cover of the vial adapter package. Be careful not to touch the LUER-LOK (tip) in the center of the vial adapter. **Do not remove the adapter from the package.**
5. Leave the vial adapter in the package and place it open end up on the clean surface with the LUER-LOK pointing up.
6. Twist off the tip cap counter clockwise from the Pre-filled Syringe. **Do not touch the inside of the cap or the syringe tip.**
7. While firmly holding the package containing the adapter with one hand and the barrel of the Pre-filled Syringe with the other, connect the Pre-filled Syringe to the vial adapter by pushing the syringe tip down onto the LUER-LOK in the center of the vial adapter, turning clockwise until the syringe is secured.
8. Carefully lift up the combined syringe-and-vial-adapter and remove it from the plastic package and discard packaging.
9. With one hand, continue to hold the combined syringe-and-vial-adapter. With the other hand, hold the IXINITY vial tightly on the clean, flat surface. **Do not touch the top of the IXINITY vial or the filter spike of the combined syringe and vial adapter.**
10. Place the vial adapter over the IXINITY vial on the table; firmly push the filter spike of the vial adapter through the center of the IXINITY vial rubber circle until the clear plastic cap snaps onto the IXINITY vial.
11. Slowly push the plunger rod down to transfer all of the liquid from the syringe into the IXINITY vial.
- With the syringe and the vial still attached, gently swirl, in a circular motion, the IXINITY vial until the product is fully dissolved. IXINITY is a clear, colorless solution without visible particles. Inspect the final solution for specks before administration. Do not use contents of vial if specks or particles persist after proper reconstitution.
12. **NOTE:** If you use more than one vial of IXINITY per infusion, reconstitute each vial as per the previous instructions.
13. Remove the diluent syringe from the vial adapter by turning syringe counterclockwise until it is completely detached. **Do not touch the luer tip of the vial adapter.**
14. Remove the LUER-LOK syringe (Administration Syringe) from its sterile packaging, taking care to not touch the luer tip of the syringe. Attach to the reconstituted vial and vial adapter by turning syringe clockwise until it is securely attached.
15. Press and keep the plunger down and turn the IXINITY vial/vial adapter/Administration Syringe upside down to transfer the solution. Draw up the solution into the Administration Syringe slowly until all solution is transferred into the syringe. Inspect the vial to confirm as much liquid as possible has been extracted into the Administration Syringe. **NOTE:** If you use more than one vial of IXINITY per infusion, extract reconstituted liquid from each vial, as per the previous instructions.
16. Hold onto the vial adapter with one hand and firmly grasp the Administration Syringe with the other and unscrew the Administration Syringe from the vial adapter turning either counter clockwise. Do not touch the tip of the syringe to any object or surface. **NOTE:** If multiple reconstituted vials are required for infusion, do not detach the large *LUER-LOK Administration Syringe from the first vial until you are ready to attach the next vial (with vial adapter attached).*
17. Prior to administering the solution, invert the Administration Syringe so that the tip is pointed toward the ceiling and express any air in the syringe. Place the Administration Syringe containing the IXINITY solution on the clean surface, making sure that the tip does not touch anything.

The reconstituted solution should be infused immediately or within 3 hours of storage at room temperature after reconstitution. **NOTE:** The luer tip of the syringe must not be touched by any objects or surfaces, when disconnecting the syringe from the vial adapter, and when transferring the Administration Syringe to the infusion set.

If you are using more than one vial, stop here and proceed to the Pooling Instructions.

POOLING INSTRUCTIONS

POOLING is the process of combining two or more reconstituted vials into a larger Administration Syringe prior to intravenous administration.

Do not detach the large LUER-LOK Administration Syringe until you are ready to attach the next vial (with vial adapter attached).

Follow the instructions above for reconstitution of the second vial.

1. Remove the Administration Syringe from the first vial adapter by turning it counter clockwise until it is completely detached.
2. Attach the Administration Syringe to the second reconstituted vial by turning clockwise until it is securely attached.
3. Turn the IXINITY vial/vial adapter/Administration Syringe upside down, slowly pull on the plunger rod to draw the solution into the Administration Syringe (see Step 15 above).

Repeat this POOLING procedure with each vial you will be using.

Once you have pooled the required dose, proceed to administration using the Administration Syringe.

ADMINISTRATION INSTRUCTIONS

For intravenous use after reconstitution only.

IXINITY is administered by intravenous (IV) infusion after reconstitution with diluent (Sterile Water for Injection) supplied in the Pre-filled Syringe.

IXINITY must not be mixed with other medicinal products for infusion.

Reconstituted IXINITY must be pulled into an Administration Syringe prior to infusion.

IXINITY is normally administered intravenously over about 5 minutes at a maximum infusion rate of 10 mL per minute. The infusion rate should be adapted to the comfort level of each patient.

1. Attach the Administration Syringe containing the reconstituted IXINITY solution to the luer end of the sterile infusion set. Inspect for and remove any air bubbles in the infusion set and Administration Syringe. **NOTE:** The luer tip of the Administration Syringe and the luer connection of the infusion set must not be touched by any object or surface, prior to connection of the Administration Syringe.
2. Transfer IXINITY solution into the tube by pressing the syringe plunger until the tubing is completely filled. Once again, inspect for and remove any air bubbles in the infusion set and Administration Syringe.
3. Perform venipuncture as directed by your healthcare provider.
4. Limit the amount of blood entering the tubing. Blood must never enter the syringe. If blood is observed in the tubing or syringe, discard all material and resume administration with a new package.
5. Following completion of the infusion, remove the infusion set, press the sterile gauze on the infusion site until bleeding has stopped, then apply a sterile bandage. The amount of drug product remaining in the infusion set should be minimal. Log the batch number of the IXINITY used; it is located on the container.

Dispose of all unused solution, empty vials, and used needles and syringes in an appropriate container for throwing away medical waste as it may hurt others if not handled properly.

Contact your healthcare provider or local hemophilia treatment center if you experience any problems.

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Aptevo BioTherapeutics LLC

Berwyn PA, 19312

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Part Number: 1003240

CM-FIX-0239