THE HEMOPHILIA FOUNDATION OF SOUTHERN CALIFORNIA'S

11TH ANNUAL

UNITE FOR BLEEDING DISORDERS WALK

Saturday, November 2nd, 2019 | Griffith Park
LETTER FROM
THE EXECUTIVE DIRECTOR

Dear Walk Supporters,

On behalf of the bleeding disorders community in Southern California, please accept my utmost gratitude and humble thanks for providing your generous support and funding for our 11th Annual Unite for Bleeding Disorders Walk. There have been many changes in the past few years with new treatments and various battles that have been fought with proposed changes to insurance. One thing that hasn’t changed is the bleeding disorders’ community passion, commitment, and love for each other. Watching a child with a bleeding disorder grow up at camp and later transform into a Leader In Training, Volunteer and then Counselor in order to give back and be a mentor exemplifies this culture. It is an honor and privilege to serve as the Executive Director in Southern California. Thank you again for all your support!

LETTER FROM
THE BOARD PRESIDENT

It’s time to Walk! The Unite for Bleeding Disorders Walk is our primary fundraiser for Camp Blood Brothers and Sisters. Camp is a remarkable place. My eldest son has attended camp since he was in second grade. Every year he looks forward to camp and continues to do so. When I asked him to sum up his experience at camp, he said, “Camp was a life-changing experience. As a camper, it provided me with friends and experiences that I will remember forever and gave me a sense of community so that I don’t feel alone with my disorder. As a counselor, it gave me the best leadership experience of my life and invaluable skills for the future.”

As a parent, camp is also wonderful because you know that your child is being well taken care of by people who truly understand bleeding disorders. There really weren’t that many times that I was away from my son when he was younger, except for the times he was at camp.

In addition to being a fundraiser, the Walk is another opportunity to spend time with members of the Bleeding Disorders Community. Many times, it can be difficult to find people who truly understand about bleeding disorders, except for other people in the bleeding disorders community. We look forward to seeing you at the Walk. Even if you aren’t able to attend in person, who can still participate and show your support by donating. Thank you to those who already donated!

Sincerely,
Sandra Knight

2019 - 2020 CALENDAR

<table>
<thead>
<tr>
<th>Month</th>
<th>Event</th>
<th>Location</th>
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| November 11 | **Men’s Bleeding Disorder Dinner**  
DISTRICT • Los Angeles, CA | |
| December 2  | **Mujeres Enlazadas por Sangre Dinner**  
Maggiano’s Little Italy • Cosa Mesa, CA | |
| December 7  | **Santa Barbara Snowflake Festival**  
Santa Barbara Zoo • Santa Barbara, CA | |
| December 14 | **Santa Ana Snowflake Festival**  
Santa Ana Zoo • Santa Ana, CA | |
| December 18 | **Riverside Snowflake Dinner**  
Mission Inn Hotel • Riverside, CA | |
| December 29 | **Blood Brothers New Years Celebration**  
LA Memorial Coliseum • Los Angeles, CA | |
| January 25 | **Industry Forum**  
Mount Saint Mary’s University • Los Angeles, CA | |
PAINTED TURTLE 2019

START LINE

REGISTRATION

MAIN STAGE

VIP TENTS

BOOTH & ACTIVITIES

WAY IN

December 14     Santa Ana Snowflake Festival
Santa Ana Zoo • Santa Ana, CA

December 18     Riverside Snowflake Dinner
Mission Inn Hotel • Riverside, CA

December 29     Blood Brothers New Year’s Celebration
LA Memorial Coliseum • Los Angeles, CA

January 25     Industry Forum
Mount Saint Mary’s University • Los Angeles, CA
We join our Hemophilia community to walk once again in support of our son who lives with Hemophilia B. Hemophilia is a big part of our lives, therefore, we join forces with the community who understands, shares in our struggles, educates us, and has held our hand for the last 8 years. Simply, we walk to support a cause we love. We are hopeful that by raising awareness through this walk, others will be encouraged to do the same. Our children and family units alike clearly benefit from the funds generated. Hemophilia is difficult on many levels, and the funds raised during this annual event are able to fund recreational activities that not only benefit our children’s mental state of being, but also the parents and caretakers. This is our effort to give back to a community that serves its people and from which we positively benefit. Please join us in this cause.

Rose Cardenas, Walk Co-Chair

I am so honored to be a Walk Co-Chair. As a social worker for Brothers Healthcare, I lean on the Hemophilia Foundation of Southern California (HFSC) often as a reliable resource for the families I work with. When I conduct an intake of a newly diagnosed patient, I tell them about resources available to them, HFSC being one of them. It puts the families at ease to know there is a foundation that they can be a part of. I have the desire to help raise money for the UNITE Walk because I see how the foundation benefits people who are affected by a bleeding disorder. Because of HFSC, children are able to experience summer camp and interact with others who have the similar medical conditions they do. HFSC hosts events that bring families together, thus preventing them from feeling isolated. Through my fundraising efforts, it is my hope that more children will be able to attend summer camp, more families are able to receive assistance through the emergency fund program, and more people become educated on topics related to their bleeding disorder though educational events. My “WHY” is simply the experiences families and children will be able to partake in because of the money I helped raise.

Veronica Diaz, Walk Co-Chair
<table>
<thead>
<tr>
<th>Time</th>
<th>Event</th>
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<tbody>
<tr>
<td>9:00 AM</td>
<td>Registration, Sponsor and Activity Booths Open</td>
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<tr>
<td>10:30 AM</td>
<td>National Anthem</td>
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<tr>
<td></td>
<td>Welcome</td>
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<tr>
<td></td>
<td>Michelle Kim, Executive Director</td>
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<td>Vickere Murphy, Representative, Senator Anthony Portantino</td>
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<td>Jordana Zeger, Chief Operating Officer, National Hemophilia Foundation</td>
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<tr>
<td></td>
<td>Thank You</td>
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<td></td>
<td>Rose Cardenas and Veronica Diaz, Walk Co-Chairs</td>
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<tr>
<td></td>
<td>Top Team Award Presented by Takeda</td>
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<td>Lindsey Adey, Takeda</td>
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<td>Top Individual Award</td>
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<td>HFSC Board Members</td>
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<td>Largest Team Award</td>
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<td>HFSC Board Members</td>
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<td></td>
<td>Best T-Shirt Award</td>
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<td>HFSC Board Members</td>
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<td>Best Costume Award</td>
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<td>HFSC Board Members</td>
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<td>Factor Club Winners</td>
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<td>HFSC Board Members</td>
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<td></td>
<td>Pinwheel Ceremony</td>
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<tr>
<td>11:00 AM</td>
<td>Walk Begins</td>
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<td></td>
<td>Water Station Halfway, No Restrooms on Walk Route</td>
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<tr>
<td>11:45 AM</td>
<td>Walk Ends, Booths Re-Open</td>
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<td></td>
<td>Pizza and Ice Cream, Thank You to Ice Cream Sponsor Takeda</td>
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<tr>
<td>12:00 PM</td>
<td>Breakdancing Performance and Lesson from</td>
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<tr>
<td></td>
<td>Dance Stylez Entertainment B-Boyz</td>
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<tr>
<td>1:00 PM</td>
<td>Departure</td>
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Takeda is here to support you throughout your journey and help you embrace life’s possibilities. Our focus on factor treatments and educational programs, and our dedication to the bleeding disorders community, remain unchanged. And our commitment to patients, inspired by our vision for a bleed-free world, is stronger than ever. Let’s make today brilliant.

bleedingdisorders.com
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Let's make today brilliant.
We walk for my son Jason and to help raise money to send children with bleeding disorders to camp. Jason has had the privilege to attend camp the last three years with the money raised by the Unite for Bleeding Disorders Walk. Jason says, “Camp Blood Brothers and Sisters is magical!” This year, his favorite part of camp was the pine wood derby race and no-hands spaghetti. I have noticed he is taking on more responsibilities since attending camp; he helps me set up and prepare his factor, and he is also learning to self-infuse and better advocate for himself and his bleeding disorder. Camp gives these children hope, strength and inspiration! Although they may have a bleeding disorder, the bleeding disorder doesn’t have them, and they can do anything they set their minds to.

RIGO MANZO
HFSC COMMUNITY LIAISON

Ever since I was diagnosed with Hemophilia, I have been surrounded by incredible people in the medical field, as well as great friends that also know what it is like to have a bleeding disorder. I am very fortunate to be a part of a community where there is so much compassion for our fellow blood brothers and sisters. I’m also proud of how empowered the bleeding disorder community becomes when forward-thinking leaders continue to fight for access to valuable information and services. I walk so that I can provide that same love and support I was given throughout my life to the youth of the bleeding disorder community. I hope to do that with my work as one of the Community Liaisons of Hemo SoCal. I walk so that future generations can run.
From July 23-28, 2019, HFSC sponsored over 100 campers who attended the 40th Annual Camp Blood Brothers and Sisters held at The Painted Turtle in Lake Hughes, CA. Campers had a blast and participated in many activities including archery, ropes course with zip line, talent show and of course the famous Silly Olympics! HFSC is so proud of the Big Stick Award Winners: Benjamin Cardenas, Jesus Fabian, Michael Lowe, Bentley Martinez, Andrew Turrey. Congratulations to our All Access Award winner, Lucas Brolezi.

HFSC is particularly thankful for our major camp donors Takeda, Novo Nordisk, Pfizer, CVS, Bayer, CSL Behring, and Sanofi Genzyme. We are also thankful to our many walkers and individual donors who stepped up to generously support camp! Thank you also to our many volunteers; camp could not run without you!
What is HEMLIBRA?

HEMLIBRA is a prescription medicine used for routine prophylaxis to prevent or reduce the frequency of bleeding episodes in adults and children, ages newborn and older, with hemophilia A with or without factor VIII inhibitors.

What is the most important information I should know about HEMLIBRA?

HEMLIBRA increases the potential for your blood to clot. Carefully follow your healthcare provider’s instructions regarding when to use an on-demand bypassing agent or factor VIII, and the dose and schedule to use for breakthrough bleed treatment. HEMLIBRA may cause serious side effects when used with activated prothrombin complex concentrate (aPCC; FEIBA®), including thrombotic microangiopathy (TMA), and blood clots (thrombotic events). If aPCC (FEIBA®) is needed, talk to your healthcare provider in case you feel you need more than 100 U/kg of aPCC (FEIBA®) total.

Please see Brief Summary of Medication Guide on following page for Important Safety Information, including Serious Side Effects.
What is the most important information I should know about HEMLIBRA?

HEMLIBRA increases the potential for your blood to clot. Carefully follow your healthcare provider’s instructions regarding when to use an on-demand bypassing agent or factor VIII (FVIII) and the recommended dose and schedule to use for breakthrough bleed treatment.

HEMLIBRA may cause the following serious side effects when used with activated prothrombin complex concentrate (aPCC; FEIBA®), including:

- Thrombotic microangiopathy (TMA). This is a condition involving blood clots and injury to small blood vessels that may cause harm to your kidneys, brain, and other organs. Get medical help right away if you have any of the following signs or symptoms during or after treatment with HEMLIBRA:
  - confusion – stomach (abdomen)
  - weakness or back pain
  - swelling of arms and legs – nausea or vomiting
  - yellowing of skin and eyes – feeling sick
  - decreased urination

- Blood clots (thrombotic events). Blood clots may form in blood vessels in your arm, leg, lung, or head. Get medical help right away if you have any of these signs or symptoms of blood clots during or after treatment with HEMLIBRA:
  - swelling in arms or legs – cough up blood
  - pain or redness in your arms or legs – feel faint
  - shortness of breath – headache
  - chest pain or tightness – numbness in your face
  - joint pain – eye pain or swelling
  - fast heart rate – trouble seeing

If aPCC (FEIBA®) is needed, talk to your healthcare provider in case you feel you need more than 100 U/kg of aPCC (FEIBA®) total.

See “What are the possible side effects of HEMLIBRA?” for more information about side effects.

What is HEMLIBRA?

HEMLIBRA is a prescription medicine used for routine prophylaxis to prevent or reduce the frequency of bleeding episodes in adults and children, ages newborn and older, with hemophilia A with or without factor VIII inhibitors.

Hemophilia A is a bleeding condition people can be born with where a missing or faulty Blood clotting factor (factor VIII) prevents blood from clotting normally. HEMLIBRA is a therapeutic antibody that bridges clotting factors to help your blood clot.

Before using HEMLIBRA, tell your healthcare provider about all of your medical conditions, including if you:

- are pregnant or plan to become pregnant. It is not known if HEMLIBRA may harm your unborn baby. Women who are able to become pregnant should use birth control (contraception) during treatment with HEMLIBRA.
- are breastfeeding or plan to breastfeed. It is not known if HEMLIBRA passes into your breast milk.

Tell your healthcare provider about all the medicines you take, including prescription medicines, over-the-counter medicines, vitamins, or herbal supplements. Keep a list of them to show your healthcare provider and pharmacist when you get a new medicine.

How should I use HEMLIBRA?

See the detailed “Instructions for Use” that comes with your HEMLIBRA for information on how to prepare and inject a dose of HEMLIBRA, and how to properly throw away (dispose of) used needles and syringes.

- Use HEMLIBRA exactly as prescribed by your healthcare provider.
- Stop (discontinue) prophylactic use of bypassing agents the day before starting HEMLIBRA prophylaxis.
- You may continue prophylactic use of FVIII for the first week of HEMLIBRA prophylaxis.
- HEMLIBRA is given as an injection under your skin (subcutaneous injection) by you or a caregiver.

Your healthcare provider should show you or your caregiver how to prepare, measure, and inject your dose of HEMLIBRA before you inject yourself for the first time.

Do not attempt to inject yourself or another person unless you have been taught how to do so by a healthcare provider.

Your healthcare provider will prescribe your dose based on your weight. If your weight changes, tell your healthcare provider.

You will receive HEMLIBRA 1 time a week for the first 4 weeks. Then you will receive a maintenance dose as prescribed by your healthcare provider.

If you miss a dose of HEMLIBRA on your scheduled day, you should give the dose as soon as you remember. You must give the missed dose as soon as possible before the next scheduled dose, and then continue with your normal dosing schedule.

Do not give two doses on the same day to make up for a missed dose.

HEMLIBRA may interfere with laboratory tests that measure how well your blood is clotting and may cause a false reading. Talk to your healthcare provider about how this may affect your care.

What are the possible side effects of HEMLIBRA?

See “What is the most important information I should know about HEMLIBRA?” for more information about side effects.

The most common side effects of HEMLIBRA include:

- redness, tenderness, warmth, or itching at the site of injection
- headache
- joint pain

These are not all of the possible side effects of HEMLIBRA. Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

How should I store HEMLIBRA?

- Store HEMLIBRA in the refrigerator at 36°F to 46°F (2°C to 8°C). Do not freeze.
- Store HEMLIBRA in the original carton to protect the vials from light.
- Do not shake HEMLIBRA.

If needed, unopened vials of HEMLIBRA can be stored out of the refrigerator and then returned to the refrigerator. HEMLIBRA should not be stored out of the refrigerator for more than a total of 7 days or at a temperature greater than 86°F (30°C).

- After HEMLIBRA is transferred from the vial to the syringe, HEMLIBRA should be used right away.
- Throw away (dispose of) any unused HEMLIBRA left in the vial.

Keep HEMLIBRA and all medicines out of the reach of children.

General information about the safe and effective use of HEMLIBRA.

Medicines are sometimes prescribed for purposes other than those listed in a Medication Guide. Do not use HEMLIBRA for a condition for which it was not prescribed. Do not give HEMLIBRA to other people, even if they have the same symptoms that you have. It may harm them. You can ask your pharmacist or healthcare provider for information about HEMLIBRA that is written for health professionals.

What are the ingredients in HEMLIBRA?

Active ingredient: emicizumab-kxwh

Inactive ingredients: L-arginine, L-histidine, poloxamer 188, and L-aspartic acid.

Manufactured by: Genentech, Inc., A Member of the Roche Group, 1 DNA Way, South San Francisco, CA 94080-4990
U.S License No. 1048

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For more information, go to www.HEMLIBRA.com or call 1-866-HEMLIBRA.

This Medication Guide has been approved by the U.S. Food and Drug Administration. Revised: 10/2018

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Being a teen leader has opened my eyes to a world of new possibilities. In Sacramento we were able to go speak with our district representatives; making a huge difference for our state, one of the largest and most influential in the country. Washington D.C. was even more eye opening, there were tons of people waiting to speak to the ones in charge, not only to the district representatives, but people like the Speaker of the House and our Senator. The feeling of actually being able to make an impact and speak to these important people about what I believed in was liberating. I was finally able to speak up for myself and others who could not, especially where it matters the most. I thank HFSC for all it’s advocacy efforts to provide a better life for teens with bleeding disorders like myself. Please donate to this impactful organization.

MIA CASTANEDA
HFSC TEEN LEADER

At Pfizer Hemophilia, we have always been deeply committed to you and to listening to what you have to say. Over the years, what you’ve shared with us has proven invaluable. The events we sponsor, the technology we develop, and the educational materials we create are all designed in response to the requests, needs, and desires of the hemophilia community.

We are grateful for having the chance to partner with you.

—Your Pfizer Hemophilia Team
MARIELA RODRIGUEZ  
TEAM JONATHAN LOPEZ

Hola nosotros somos la Familia López y tenemos un hijo con la condición llamada Glanzmann’s Thrombasthenia. Es una condición sanguínea y es por eso que quisimos hacer una rifa para poder juntar un poco de dinero. Sabemos que tal vez no será mucho lo que podamos aportar pero también sabemos que de poco en poco llegaremos a cumplir la meta. ¡Nosotros estamos muy agradecidos con la Fundación de HFSC por que nos has apoyado mucho! Gracias a ellos hemos aprendido cómo cuidar a nuestro hijo y a que el tenga un mejor futuro a cómo reconocer cada síntoma de su condición y a como abogar por el y su salud. Para el a tenido un fuerte impacto para bien poder ir a los campamentos. El se ha sentido más seguro de sí mismo. ¡Se ha sentido más aceptado por la sociedad, ha tenido más confianza en sus habilidades, a aprendido a como abogar por el mismo y por su salud pero sobre todo a aprendido a ser solidario con los demás al saber que hay personas que sufren y viven igual que el pero que también disfrutan y se divierten igual que el! Mi hijo a hecho grandes amigos que esperamos en un futuro sean los grandes abogados de los ahora más pequeños y de su propio futuro. ¡Gracias a toda la familia de sangre por siempre ser un gran apoyo para todos y a la fundación de HFSC por ser nuestro apoyo incondicional y por ayudarnos a cuidar a nuestros hijos que son el tesoro más grande que temenos!

CONGRATULATIONS 2019  
FACTOR CLUB WINNERS

Jesus Avila, Paul Avila, Karlyn Johnson Brown, Ana Cardenas, Rose Cardenas, The Carthey Family, Shin Chen, Veronica Diaz, Kathy Dunham, Wayne Guzman, Oscar Horta, Amelie Iavicoli, Elena Iavicoli, Greg Jordan, Joshua Kim, Michelle Kim, Sandra Knight, Judy Mangione, Christina Olivarria, Perry Parker, Pedro Sanchez, Michele Warner, Doris Quon*

*As of October 23, 2019

THANK YOU TO OUR WALKERS!

There’s Still Time to Donate! Donations are accepted until 1/1/2020. Visit uniteforbleedingdisorders.org/event/socal or send donations to Hemophilia Foundation of Southern California 959 E. Walnut Street, Suite 114, Pasadena, CA 91106.
ADYNOVATE® is a treatment that can be personalized to fit your lifestyle so you have more time to spend doing the other things that also make you, you. It has a simple, twice-weekly dosing schedule on the same 2 days every week.1,2

*In clinical trials, ADYNOVATE demonstrated the ability to help prevent bleeding episodes using a prophylaxis regimen.

HEMOPHILIA A IS A PIECE OF YOU. NOT ALL OF YOU.

ADYNOVATE® is a treatment that can be personalized to fit your lifestyle so you have more time to spend doing the other things that also make you, you.
ADYNOVATE® [Antihemophilic Factor (Recombinant), PEGylated]

**Patient Important facts about ADYNOVATE**

This leaflet summarizes important information about ADYNOVATE. 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 ADYNOVATE. If you have any questions after reading this, ask your healthcare provider.

**What is the most important information I need to know about ADYNOVATE?**

Do not attempt to do an infusion to yourself unless you have been taught how by your healthcare provider or hemophilia center. You must carefully follow your healthcare provider's instructions regarding the dose and schedule for infusing ADYNOVATE so that your treatment will work best for you.

**What is ADYNOVATE?**

ADYNOVATE is an injectable medicine that is used to help treat and control bleeding in children and adults with hemophilia A (congenital Factor VIII deficiency). Your healthcare provider may give you ADYNOVATE when you have surgery. ADYNOVATE can reduce the number of bleeding episodes when used regularly (prophylaxis).

ADYNOVATE is not used to treat von Willebrand disease.

**Who should not use ADYNOVATE?**

You should not use ADYNOVATE if you:

- Are allergic to mouse or hamster protein
- Are allergic to any ingredients in ADYNOVATE or ADVATE® [Antihemophilic Factor (Recombinant)]

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

**How should I use ADYNOVATE?**

ADYNOVATE is given directly into the bloodstream. You may infuse ADYNOVATE at a hemophilia treatment center, at your healthcare provider’s office or in your home. You should be trained on how to do infusions by your healthcare provider or hemophilia treatment center. Many people with hemophilia A learn to infuse their ADYNOVATE by themselves or with the help of a family member.

Your healthcare provider will tell you how much ADYNOVATE to use based on your individual weight, level of physical activity, the severity of your hemophilia A, and where you are bleeding.

Reconstituted product (after mixing dry product with wet diluent) must be used within 3 hours and cannot be stored or refrigerated. Discard any ADYNOVATE left in the vial at the end of your infusion as directed by your healthcare professional.

You may have to have blood tests done after getting ADYNOVATE to be sure that your blood level of factor VIII is high enough to clot your blood.

**What should I tell my healthcare provider before I use ADYNOVATE?**

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 mice or hamsters.
- Are breastfeeding. It is not known if ADYNOVATE passes into your milk and if it can harm your baby.
- Are pregnant or planning to become pregnant. It is not known if ADYNOVATE may harm your unborn baby.
- Have been told that you have inhibitors to factor VIII (because ADYNOVATE may not work for you).

**What is the most important information I need to know about ADYNOVATE?**

Do not attempt to do an infusion to yourself unless you have been taught how by your healthcare provider or hemophilia center. You must carefully follow your healthcare provider's instructions regarding the dose and schedule for infusing ADYNOVATE so that your treatment will work best for you.

You should tell your healthcare provider right away if your bleeding does not stop after taking ADYNOVATE.

**What are the possible side effects of ADYNOVATE?**

You can have an allergic reaction to ADYNOVATE. Call your healthcare provider right away and stop treatment if you get a rash or hives, itching, tightness of the throat, chest pain or tightness, difficulty breathing, lightheadedness, dizziness, nausea or fainting.

The common side effects of ADYNOVATE are headache and nausea. Tell your healthcare provider about any side effects that bother you or do not go away. These are not all the possible side effects with ADYNOVATE. You can ask your healthcare provider for information that is written for healthcare professionals.

**What else should I know about ADYNOVATE and Hemophilia A?**

Your body may form inhibitors to Factor VIII. An inhibitor is part of the body’s normal defense system. If you form inhibitors, it may stop ADYNOVATE from working properly. Consult with your healthcare provider to make sure you are carefully monitored with blood tests for the development of inhibitors to Factor VIII.

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

The risk information provided here is not comprehensive. To learn more, talk with your healthcare provider or pharmacist about ADYNOVATE. The FDA-approved product labeling can be found at www.ADYNOVATE.com or 1-877-825-3327.

You are encouraged to report negative side effects of prescription drugs to the FDA. Visit www.fda.gov/medwatch, or call 1-800-FDA-1088.

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Issued 05/2018

S47033

Snowflake Festival

December 7th: Santa Barbara Zoo | 10am-1pm
https://hfscsnowflake2019.eventbrite.com

December 14th: Santa Ana Zoo | 10am-1pm
https://hfscsantaana2019.eventbrite.com

December 18th: Mission Inn, Riverside | 6pm-8:30pm
https://riversideholiday2019.eventbrite.com

Community Members Only

ANIMAL SHOW* CRAFTS * GIFTS FOR YOUTH * ENJOY THE ZOO