HFSC OLYMPICS
LET THE GAMES BEGIN
HFSC’S UNITE FOR BLEEDING DISORDERS WALK
NOVEMBER 13, 2021 • LA MEMORIAL COLISEUM
HTTPS://WWW.UNITEFORBLEEDINGDISORDERS.ORG/EVENT/SOCAL
Dear Supporters and Friends,

On behalf of the Hemophilia Foundation of Southern California Board of Directors and Staff, we sincerely thank you for your tireless efforts and unwavering support of our 13th Annual Unite Walk. We are so thrilled we can be together live, in person, after such a long and stressful pandemic. Our community continues to be a constant source of courage and support. When I reflect upon our walk’s theme, the Olympics, which is symbolized by five rings representing five continents, I think about the great gift of our particular community in Southern California: our beautiful diversity which is our strength.

Thank you all for coming together for an extraordinary day of rallying for our future and to support the rare bleeding disorders community in Southern California. Thank you to each and every one of you for making a direct impact and difference in the lives of our community members.

With deepest love and warmest wishes,
Michelle Kim
HFSC Executive Director

I am unbelievably excited this year’s SoCal Unite for Bleeding Disorders Walk will be in person at the LA Coliseum, and I am so looking forward to seeing everyone, many of you for the first time in almost 2 years! Thank you so much to all of you for joining me in registering and fundraising in support of HFSC’s local initiatives such as advocacy, education and support programs, Camp Blood Brothers & Sisters, scholarships, and research. We are so very grateful!

Due to your extraordinary efforts last year, the Board was recently able to invest in local research to make a greater impact on the future of our bleeding disorders community. Two grants were awarded: one to support research in “Heavy Menstrual Bleeding Management and Quality of Life of Adolescent and Young Adult Females” and one to support research in “Real-Time Feedback on Joint Range of Motion.”

A heartfelt thank you to all of you for such amazing and impactful support in helping HFSC continue to fulfill its mission.

Sincerely,
Rick Kelly
HFSC Board President

Factor Club members have raised $500 or more for the Unite Walk in 2021.*

Jesus Avila
Paul Avila
Michael Bennet
Anita Brenner
Jack Burns
William Campbell
Mathew Casas
Cynthia Chavez
Shin Chen
Yuan Chen
Erika Clement
Linda & Paul Clement
Eli Economou
Kelly Guzman
Larry Hammerness
Amelie Iavicoli
Karlyn Johnson Brown
John Kim
Joshua Kim
Michelle Kim
Patrick Lin
Ricky Martellaro
Richard Metz
Kelly & Joseph Mullen
Bob Numerof
Christina Olivaria
Perry Parker
Doris Quon
Linda Reyes
Deborah Sazdoff
Michele Warner

*as of 10/15/2021
Friday 6:26 pm
Sharing stories by the campfire with friends
Isaac, living with hemophilia B

WE’RE IN THIS TOGETHER.

Let’s make today brilliant.

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.

bleedingdisorders.com
INTRODUCING WALK CHAIR KARLYN BROWN  
WHY WE WALK

I am honored to be the Chair for this year’s Unite Walk. Our first HFSC walk was at the LA Coliseum when our son, Langston, was just a baby. Yes, Langston has hemophilia, but he’s leading a beautiful life. This is in part because we had him diagnosed early, but also largely because of all of the advocacy, education, and support provided by HFSC and the local bleeding disorders community. He plays baseball and basketball, climbs everything, is interested in parkour, and is a super-fast runner - he had just made the track team as a 6th grader when Covid hit.

Covid brought with it many challenges for everyone, I know. But it also brought some opportunities. One of those opportunities for our family was the ability to participate in this year’s Legislative Day and to meet virtually with lawmakers and share the challenges and triumphs of being a family affected by a bleeding disorder. While we were prepping for the day, my son was nervous and unsure of why anyone would want to hear from him. My husband and I encouraged him to think of all of the folks who championed this cause before now. Those who walked before him, spoke out before him and advocated for those in the bleeding disorders community - and those who support and speak on his and our behalf now - the HFSC Board and Staff, the many scientists, doctors, patients, advocates, and families - ALL of YOU!

Langston spoke tentatively at first but when he found his voice, he spoke with strength and pride about all of the things he is able to do and the many ways he has been supported by HFSC and the bleeding disorders community in Southern California. I heard him speak about the challenges facing those with hemophilia and other bleeding disorders and how other individuals can be allies.

Langston told me what really struck him was thinking about just how many people had to pave the way and trailblaze before him so that he could live this beautiful life - so that we could have opportunities and care and support as a family. Throughout the years we have not only benefited from education, programs, support, and local advocacy, but we have gained valued friendships, thanks to HFSC’s various events. This is why we walk. To honor those who have gone before and to make the road a little less challenging for those who will come after.

THANK YOU TO OUR SPONSORS!
The most comprehensive hemophilia program for children and adults in Southern California.

- Coordinated team of specialists, including physicians, orthopaedic surgeons, pharmacists, physical therapists, a nurse practitioner and a social worker under one roof
- Nearly 60 years of experience, with more orthopaedic procedures done on patients with hemophilia than any other center in the US, according to the National Hemophilia Foundation
- Telehealth and in-person appointments
- In-house pharmacy that delivers directly to your door
- 24/7 on-call patient care provider and pharmacist
- Medication management and education
- On-site assistance with navigating insurance barriers and resolving billing issues
- As a 340B pharmacy, all proceeds are put back into the HTC to sustain and create services for our patients and the hemophilia community

Conveniently located directly off the 110 freeway at 403 W Adams Blvd in downtown Los Angeles.

HTC: 213-742-1402    Pharmacy: 213-742-1128

ortho-institute.org/hemophilia

YOUR HEMO HOME IS HERE.

WALK DAY INFORMATION

Agenda

9:00 am  Sponsor and Activity Booth Time
10:30 am Opening Ceremony

Welcome
National Anthem
Sponsor Welcome
Factor Club Winners
Awards Ceremony
  - Top Fundraising Team
  - Top Fundraising Individual
Pinwheel Ceremony
Warm-Up
11:00 am Walk

Route

Participants will enter LAMC through Gate 31 (The Olympic Gateway). The program will be on the Peristyle, and the start/finish line will be here as well. Please proceed around the LA Memorial Coliseum twice (or more!) in a clockwise direction. Each lap is about 0.6 miles. Restrooms will be available near the start and midpoint of the route at Gates 4 and 19. Water will be available in your bag as well as at the Peristyle after the walk begins.
The most comprehensive hemophilia program for children and adults in Southern California.

- Coordinated team of specialists, including physicians, orthopaedic surgeons, pharmacists, physical therapists, a nurse practitioner and a social worker under one roof
- Nearly 60 years of experience, with more orthopaedic procedures done on patients with hemophilia than any other center in the US, according to the National Hemophilia Foundation
- Telehealth and in-person appointments
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- Medication management and education
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- As a 340B pharmacy, all proceeds are put back into the HTC to sustain and create services for our patients and the hemophilia community

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ortho-institute.org/hemophilia
COMMUNITY MEMBER
CHARLIE FRANZEN COMPETES IN JR. OLYMPICS

This past July, I had the opportunity to compete in the 16u division of the Jr. Olympics for Water Polo. My team and I qualified for the tournament by winning different games throughout the year. Over 200 teams qualify for the tournament every year, but only around 50 teams qualify for the top division. One of the coolest things about the tournament is how teams from all over the country come to California to compete. So, it makes it super special when we win because then we know that we are one of the best teams in the country. I have gone to the Jr. Olympics before in the past, but I have never gotten a medal or even qualified for the top division, so this year was very special. The tournament took place over 4 days, and we played a total of 10 games; by the end, our entire team was exhausted. We ended up playing very well, and we were able to place third. This was a massive accomplishment for our team, especially because we had been training all year long for this specific tournament. Next year I will be moving up to the 18u division, and we have goals of doing better than this year.

Regular exercise is important. But moods vary. No two days are exactly the same. So we asked celebrity trainer Dolvett Quince to create a series of exercise videos called H-Fit. Each video features a different hemophilia-friendly workout. And each workout is built around a different mood. How are you feeling today? H-Fit has got you covered.

Sign up with Novo Nordisk and register at H-Fit.com to get free access to the H-Fit video library.

Always talk to your health care provider before starting any new exercise routine.
HFSC and 42 years of Camp! Though it was virtual, we still were able to connect and experience the magic of Camp with The Painted Turtle from July 23-25!

Families cut loose and enjoyed their favorite camp activities including making their own campfire, marshmallow toasting, Stage Day, the Pinewood Derby, and of course, no-hands spaghetti!

We are so astronomically thankful to our sponsors who joined our camp visitor’s day!

“YES YES YES! YES YES YES!”
INDICATION & IMPORTANT SAFETY INFORMATION

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. People who use activated prothrombin complex concentrate (aPCC; Feiba®) to treat breakthrough bleeds while taking HEMLIBRA may be at risk of serious side effects related to blood clots.

These serious side effects include:

- Thrombotic microangiopathy (TMA), a condition involving blood clots and injury to small blood vessels that may cause harm to your kidneys, brain, and other organs
- Blood clots (thrombotic events), which may form in blood vessels in your arm, leg, lung, or head

Please see Brief Summary of Medication Guide on following page for Important Safety Information, including Serious Side Effects.
Medication Guide
HEMLIBRA® (hem-leeb-roh)
(emicizumab-kxwh)
 injection, for subcutaneous use

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 bleeding 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
  – weakness
  – swelling of arms and legs
  – yellowing of skin and eyes
  – stomach (abdomen) or back pain
  – nausea or vomiting
  – 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
  – pain or redness in your arms or legs
  – shortness of breath
  – chest pain or tightness
  – fast heart rate
  – cough up blood
  – feel faint
  – numbness in your face
  – eye pain or swelling
  – 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. Females 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 four 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?”

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. 1046

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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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The Genentech logo is a registered trademark of Genentech, Inc.

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Due to your extraordinary efforts in 2020, the HFSC Board of Directors created the first ever HFSC Research Grant to directly impact patients in Southern California. The Board awarded two grants in the amount of $30,000 each.

The first grant was awarded to The Center for Inherited Blood Disorders (CIBD) to support research in “Heavy Menstrual Bleeding Management and Quality of Life of Adolescent and Young Adult Females.” The second grant was awarded to the Orthopaedic Hemophilia Treatment Center to support research in “Real-Time Feedback on Joint Range of Motion.”

Supporting research is a crucial way to improve the lives of people living with rare blood disorders and we are so grateful to all of you!

La fundación del Sur de California tendrá una caminata en el mes de noviembre en la cual se recaudarán fondos. Muchos de estos fondos serán para servicios en varios aspectos para todas las personas que padecen de un desorden de sangre y sus familias.

¡Es importante recordar que los fondos recaudados ayudan a las familias de nuestra comunidad en emergencias menos esperadas, ayudan a nuestros jóvenes a permanecer en la escuela y envían a nuestros hijos más pequeños al campamento de verano donde aprenden a ser independientes y a aceptar su desorden sanguíneo! Por eso es importante para mí y mi familia participar y apoyar esta caminata y recaudar fondos para esta buena causa.
CAMP ON THE MOVE

Thank you to The Painted Turtle for hosting “Camp on the Move Parking Lot Party” in August for our campers! We had so much fun singing our favorite Camp songs, painting (cars too!), and dancing! Thank you for gifting us with wood shop items, a giant bag of Paul Newman goodies, arts and crafts, a game of choice, and most importantly, our own turtle pillow!
Prophylaxis with ADVATE prevented bleeds

The ability of ADVATE to treat or prevent bleeds was evaluated in a clinical study using a standard prophylaxis, pharmacokinetic driven prophylaxis, and on-demand treatment. 53 previously treated patients (PTPs) with severe to moderately severe hemophilia A were analyzed. For the first 6 months of the study, patients received on-demand treatment. For the following 12 months of the study, patients received either standard prophylaxis every 48 hours or a pharmacokinetic-driven prophylaxis every 72 hours. The primary goal of the study was to compare annual bleeding rates between those receiving prophylaxis treatment and those receiving treatment on-demand. The number of bleeds per year for the 2 prophylaxis regimens were comparable:

- Those patients experienced a median of 1 overall bleed per year on either prophylaxis treatment vs 44 overall bleeds per year with on-demand treatment. This represented a 98% reduction in overall bleeds per year.
- Zero bleeds were reported in 42% of patients (22 out of 53 patients) during 12 months on prophylaxis

Median is the middle number in a group of numbers arranged from lowest to highest.

ADVATE Important Information

What is ADVATE?

- ADVATE is a medicine used to replace clotting factor (factor VIII or antihemophilic factor) that is missing in people with hemophilia A (also called “classic” hemophilia).
- ADVATE is used to prevent and control bleeding in adults and children (0-16 years) with hemophilia A. Your healthcare provider (HCP) may give you ADVATE when you have surgery.
- ADVATE can reduce the number of bleeding episodes in adults and children (0-16 years) when used regularly (prophylaxis).
- ADVATE is not used to treat von Willebrand disease.

DETAILED IMPORTANT RISK INFORMATION

Who should not use ADVATE?

Do not use ADVATE if you:
- Are allergic to mice or hamsters.
- Are allergic to any ingredients in ADVATE.

Tell your HCP if you are pregnant or breastfeeding because ADVATE may not be right for you.

What should I tell my HCP before using ADVATE?

Tell your HCP 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 ADVATE passes into your milk and if it can harm your baby.

What should I tell my HCP before using ADVATE? (continued)

- Are or become pregnant. It is not known if ADVATE may harm your unborn baby.
- Have been told that you have inhibitors to factor VIII (because ADVATE may not work for you).

What important information do I need to know about ADVATE?

- You can have an allergic reaction to ADVATE. Call your HCP 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.
- Do not attempt to infuse yourself with ADVATE unless you have been taught by your HCP or hemophilia center.

What else should I know about ADVATE 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 ADVATE from working properly. Talk with your HCP to make sure you are carefully monitored with blood tests for the development of inhibitors to factor VIII.

What are possible side effects of ADVATE?

- Side effects that have been reported with ADVATE include: cough, headache, joint swelling/aching, sore throat, fever, itching, unusual taste, diziness, hematoma, abdominal pain, hot flashes, swelling of legs, diarrhea, chills, runny nose/congestion, nausea/vomiting, sweating, and rash. Tell your HCP about any side effects that bother you or do not go away or if your bleeding does not stop after taking ADVATE.

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.

Please see Important Facts about ADVATE on the following page and discuss with your HCP.

For Full Prescribing Information, visit www.ADVA TE.com.
**ADVATE [Antihemophilic Factor (Recombinant)]**

**Important facts about ADVATE [Antihemophilic Factor (Recombinant)]**

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

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

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 ADVATE so that your treatment will work best for you.

**What is ADVATE?**

ADVATE is a medicine used to replace clotting factor (factor VIII or antihemophilic factor) that is missing in people with hemophilia A (also called “classic” hemophilia). The product does not contain plasma or albumin. Hemophilia A is an inherited bleeding disorder that prevents blood from clotting normally. ADVATE is used to prevent and control bleeding in adults and children (0-16 years) with hemophilia A.

Your healthcare provider may give you ADVATE when you have surgery. ADVATE can reduce the number of bleeding episodes in adults and children (0-16 years) when used regularly (prophylaxis). ADVATE is not used to treat von Willebrand disease.

**Who should not use ADVATE?**

You should not use ADVATE if you:
- Are allergic to mice or hamsters.
- Are allergic to any ingredients in ADVATE.

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

**How should I use ADVATE?**

ADVATE is given directly into the bloodstream.

You may infuse ADVATE 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 ADVATE by themselves or with the help of a family member.

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

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

Call your healthcare provider right away if your bleeding does not stop after taking ADVATE.

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

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

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

You can have an allergic reaction to ADVATE.

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.

Side effects that have been reported with ADVATE include:
- Cough
- Headache
- Joint swelling/aching
- Sore throat
- Fever
- Itching
- Unusual taste
- Dizziness
- Hematoma
- Abdominal pain
- Hot flashes
- Swelling of legs
- Diarrhea
- Chills
- Runny nose/congestion
- Nausea/vomiting
- Sweating
- Rash

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 ADVATE. You can ask your healthcare provider for information that is written for healthcare professionals.

**What else should I know about ADVATE 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 ADVATE 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 ADVATE for a condition for which it is not prescribed. Do not share ADVATE 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 health care provider or pharmacist about ADVATE. The FDA-approved product labeling can be found at www.ADVATE.com or 1-877-828-3327.

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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Patented: see https://www.takeda.com/en-us/patents/

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Issued: 12/2018

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**Hemophilia Foundation of Southern California**

959 East Walnut Street, Suite 114
Pasadena, CA 91106

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**INDUSTRY FORUM 2022**

**JANUARY 19:**
HTC TOWNHALL: PIPELINE PRODUCTS WEBINAR
1:00-8:00PM
HTTPS://BIT.LY/3G0D2PO

**JANUARY 22:**
INDUSTRY FORUM
PASADENA CITY COLLEGE
9:00AM - 1:00PM

GAIN KNOWLEDGE OF NEW PRODUCTS AND TREATMENTS!
INCLUDES SIX FLAGS TICKET TO EACH ATTENDEE AGES 3+
HTTPS://HFSINDUSTRYFORUM2022.EVENTBRITE.COM

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**SAVE THE DATE 2022 WOMEN'S RETREAT**

Join us in person for an overnight retreat of support, self-empowerment, education and fun!

**Saturday, March 19 - Sunday, March 20, 2022**

Marriott Irvine Spectrum
7905 Irvine Center Drive
Irvine, CA

Begins Saturday 9:00am | Ends Sunday 11:00am