Hemophilia

GUIDELINES FOR PARENTS

Indian Academy of Pediatrics (IAP)

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34 FAQs on HEMOPHILIA

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Hemophilia

Hemophilia is a condition where our blood does not clot easily. For blood to clot, we need to have normal platelets and clotting factors in our blood. In hemophilia, the clotting factors VIII or IX are deficient from birth. Children born with hemophilia have increased tendency to bleed either spontaneously or after trauma.

Q1
Why is my child getting bruising and bleeding?

Hemophilia is a condition where our blood does not clot easily. For blood to clot, we need to have normal platelets and clotting factors in our blood. In hemophilia, the clotting factors VIII or IX are deficient from birth. Children born with hemophilia have increased tendency to bleed either spontaneously or after trauma.

Q2
Our son has been diagnosed with hemophilia. Nobody on both sides of the family has this disease. Why did our child still get it?

Hemophilia is a genetic disease which is inherited in two-thirds of the cases. In the rest, it is a new mutation, and hence family history may not be there.
We have not told anyone in the family or friends regarding this illness yet. Should we tell them?

Yes. Since it is a chronic condition which will need quite a few lifestyle adjustments, it is better to inform close family members and friends. Also at school entry, school teachers and others who take care of your child such as bus drivers also should be aware that your child has increased tendency to bleed, especially following trauma. Your child may need to be taken care by others when you are not available.

What are the common sites of bleeding?

Children with hemophilia usually present with bleeding into joints and muscles. At a young age, when children are crawling, they can present with bruises on their shins and chin due to trauma. When they start walking, they usually present with bruises and joint bleeding into ankles and knees (Fig. 1).

Fig. 1: Joint bleeding in hemophilia.
Bleeding could be spontaneous or following any trauma. Usually, they do not have bleeding from nose and mouth, unless there is an injury on the tongue or following tooth fall (Fig. 2). Joint and muscle bleeds account for >80% of bleeding observed. Other sites of bleeding include blood in urine and stools, blood in vomitus, bleeding into the brain (Fig. 3), airway, etc. which are more serious. Child should be brought to the Hemophilia Treatment Center or nearest pediatrician at the earliest suspicion of severe bleeding. Repeated bleeding into joints can result in “target joints” and permanent damage to the joints leading to disability in future. With the present treatment options available in our country, children with hemophilia are able to lead near normal lives with less risk for disability.

Fig. 2: Trauma-related oral bleeding.

Fig. 3: Intracranial bleeding in hemophilia.
Hemophilia

How severe is the disease in my child? Will my child be able to lead a normal life?

Normally the level of factor VIII and IX in human body ranges from 50 to 150%. Hemophilia is graded as severe (factor level <1%), moderate (factor level 1–5%), and mild (factor level 5–40%) disease. In those with mild deficiency, bleeding usually occurs following any major trauma or surgery. Those with moderate deficiency develop bleeding following minor trauma and those with severe deficiency can have spontaneous bleeding. With the current treatment availability, most children are able to lead a near normal life.

Will this disease last his whole life? Will there be a cure for this illness?

Yes. At present, the treatment available for most patients will not cure the disease. Instead, risk of bleeding can be kept under check either preventing or treating bleeding episodes. Gene therapy is the only curative treatment, which is currently under research trials. Once available freely, it will offer the hope of cure.
Hemophilia is treated with clotting factor concentrates. These clotting factor concentrates replenish the deficient factor in the body and stop bleeding. These factors can be plasma derived (collected from people with normal factor levels) or recombinant (prepared in research laboratories). They can be given either on-demand, viz. after the bleeding occurs or as replacement (or prophylaxis) in order to prevent bleeding. On-demand treatment is given based on the site of bleeding and weight of the child. In most joint and muscle bleeds, 20–40% factor level is targeted. In more difficult bleeds such as psoas muscle bleed, hip joint bleed or severe bleeding, higher factor level is targeted for a longer period. While receiving on-demand treatment, children would be brought to the hospital as early as possible after bleeding. Prophylaxis, on the other hand, is to prevent bleeding. Small doses of factor concentrates are given two to three times a week to reduce the risk of severe bleeding and muscle/joint bleeds. As the bleed frequency reduces following prophylaxis, children have lesser chance of developing target joints and disability in future.

We usually advise parents to do all that is possible to reduce accidental bleeding. When the baby is very young, get childproofing of your home done by padding sharp corners of tables, chairs, cabinets, and doors. Refrain child from using walkers and staircases till his balance is good. Keep sharp objects such as knives, scissors, and similar objects away from the baby. While going to play, child should be encouraged to stay away from sports, where there is risk of injury to the head. Cycling can be taught with adequate precautions such as helmet and knee and ankle pads. Swimming as well as most indoor games are safe. Caretaker has to be cautious to prevent slipping and falling at home. Babies should not be lifted by their hands or forearms only as this might trigger a pulled elbow and bleeding.
Will he be able to play with his siblings and friends?

Yes. In the initial few years of life, we encourage parents to supervise and keep children away from sports with risk of trauma to head such as skating, diving, cricket, etc. Children on prophylaxis/replacement factor treatment can engage in sports and physical activities more freely.

Can my child receive all vaccinations? Should I give factor prior to vaccination?

Yes. All routine vaccinations are to be given to children with hemophilia. Factor administration for vaccination is not recommended. Vaccines can be preferably given subcutaneously with 26G needle. Application of ice for 2–3 minutes before and after the injection can reduce the risk of bleeding and pain significantly.

How will I cut his nails and hair without the risk of bleeding?

In the initial months, care must be taken to prevent trauma. Once the baby grows up, he will understand the risk of bleeding and will cooperate.

My son has just started walking. Should we use kneecaps and helmets to prevent bleeding?

No. There is no need for constant use of kneecaps, braces, or helmets. As discussed earlier, helmets may be used while cycling or playing in the park.
Yes. Nonsteroidal anti-inflammatory drugs (NSAIDs) commonly used for fever and pain are to be avoided in children with hemophilia. Inform your pediatrician or doctor prescribing medicines that they should not be used. Paracetamol is safe for fever and pain.

I have heard about prophylaxis in hemophilia. When should we start it? Where is it available?

Prophylaxis is the process of giving factor concentrate 1–3 times in a week in order to reduce the risk of bleeding especially in the joints. When factor is administered at regular intervals, the trough level of factor in the body usually remains above 1%, which makes the child with severe hemophilia behave-like moderate hemophilia, thus reducing the risk of spontaneous bleeding. Prophylaxis does not prevent all bleeds. It is basically given to reduce the risk of spontaneous joint bleeds, which in-turn reduces target joint formation and long-term disability associated with it. In western and resource plentiful countries, the doses recommended are high which encourages a very active lifestyle for the child-like sports. In India, we follow intermediate-to-low dose regimens aiming at reducing joint morbidity in the long term.

He fell while running and hit his head. How do I know if there is internal bleeding? Should I give factor immediately?

The child should be watched for any danger signs for head injury such as vomiting, altered behavior such as excessive cry or reduced activity, inability to move a limb and seizures. As the risk of bleeding can be late, any such symptom in the next 10–14 days should be reported to your treating doctor. In some cases, a dose of factor is usually given immediately. In children who have any of the danger signs, immediate factor administration followed by investigations such as CT scan and further management in an experienced center is advisable.
Joint bleeds usually present with pain and difficulty in movement of the joints. Unlike superficial bleeds, blue/black discoloration of a bruise is not seen in joint bleeds. Children usually develop painful swelling, which makes joint movement difficult. Young babies may cry while joint is handled and prefer not to move it. At home, if a joint bleed is expected, the joint should be kept immobile and elevated. Application of ice/cold compression can help in reducing pain. For upper limb joints, sling can be applied around the neck, and for lower limb joints bed rest is encouraged to prevent movement of the joint. In most bleeds, factor concentrate will be required to stop the bleeding. Early access to factor concentrates can reduce the bleeding and restore joint health at the earliest.

No. For serious (joint and muscle bleeds, mucosal bleeds) and life-threatening bleeds (neck/airway bleeds, bleeding inside the head, and blood in urine) treatment with factor concentrates is vital, and it is best to reach the treatment center at the earliest. If you have access to home therapy, factor treatment should be initiated at the earliest. In small bruises, small oral bleeds and other superficial bleeds, one can wait and watch and go to a hospital only if bleeding is persisting.

Yes. Most definitely. It is prudent to store few vials of factor at home, in the refrigerator at 2–8°C. As the child grows, self-infusion can be taught so that factor can be administered during an emergency by the child himself. Till then, one of the parents can be taught how to administer factors or the help of a doctor or nurse nearby can be sought.

Tongue injury can result in troublesome bleeding. In case of bleeding, application of crushed/powdered tablet of tranexamic acid can help reduce bleeding. Child should be encouraged to sleep on the side to reduce the risk of aspiration of blood. Warm food is discouraged to reduce the risk of bleeding. Factor support is usually needed for few days.
The first aid that can be done at home, before factor is given which reduces the chance of bleeding is called RICE or PRICE. RICE stands for Rest, Ice, Compression, and Elevation of the joint whereas P-RICE adds Protection. POLICE is the new terminology introduced in World Federation of Hemophilia 2020 guideline where Protection, Optimal Loading, Ice, Compression, and Elevation is the acronym used. Optimal loading refers to early initiation of physiotherapy to reduce the chance of muscle wasting. These techniques should be learnt under the guidance of experienced doctors from your “Hemophilia Treatment Center”.

Prophylaxis helps in reducing long-term joint damage. Regular physiotherapy keeps joints and muscles healthy and reduces the risk of bleeding. If prophylaxis is not feasible, early access to episodic treatment should be encouraged to promote early recovery of joint bleeds.

Children with hemophilia can develop limping subsequent to repeated joint bleeds and joint damage. This can be prevented by adhering to a regular treatment program and daily physiotherapy at home.

Q20 How should I take care of my child in order to prevent long-term joint damage?

Prophylaxis helps in reducing long-term joint damage. Regular physiotherapy keeps joints and muscles healthy and reduces the risk of bleeding. If prophylaxis is not feasible, early access to episodic treatment should be encouraged to promote early recovery of joint bleeds.

Q21 He wants to go out and play cricket. I am scared of sending him for risk of bleeding. What should I do?

Children should be encouraged to play in order to promote an active lifestyle. However, the risk of bleeding in some games such as cricket, hockey, and football is significant and children should be advised to play sports that are safer. If on prophylaxis, they can participate in most sports activities according to the factor level achieved.

Q22 What is RICE/PRICE or POLICE therapy?

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Q23 My 8-year-old son with hemophilia walks with a limp. What should I do to prevent disability in future?

Children with hemophilia can develop limping subsequent to repeated joint bleeds and joint damage. This can be prevented by adhering to a regular treatment program and daily physiotherapy at home.
Q24
Are there any dietary restrictions for my child? Will any supplements reduce his chances of bleeding?
No. There are no dietary restrictions for children with hemophilia. There are no supplements that reduce bleeding either. However, regular intake of calcium and vitamin supplements can help to improve bone health.

Q25
Can my son with hemophilia go on a school trip? Should I send factor with him?
Yes, once older your child can go on school trips. The school teachers who accompany him should be aware of when to inform you and what to do in case of a bleed. Factor concentrates may be sent with him, which can be used in case of an emergency.

Q26
My wife is a hemophilia carrier and is pregnant. Is it safe for us to have a normal delivery?
Yes. Delivery can be normal unless your gynecologist feels it is not safe for the mother or child due to other reasons.

Q27
My wife is a hemophilia carrier and we have just had a baby. How early can we test the baby for hemophilia? What is the possibility of the baby having hemophilia?
The baby can be tested in the newborn period for hemophilia A. As factor IX is normally also low in newborn babies, it is recommended to wait for few months before testing it to prevent false reports, if the baby is otherwise well. In case of carrier moms, there is 50% risk for boys to develop this disease.

Q28
Can we prevent this disease again in our family? Should I test my daughters for carrier status?
Yes, carrier status is recommended in sisters of the patient with hemophilia. Prenatal testing is available freely which should be encouraged in all carriers to prevent this disease again in families who already have one affected child.
Some patients with hemophilia can develop what are called inhibitors. Inhibitors are antibodies that can destroy the factor, and thus reduce its efficiency. The development of ET inhibitors is a serious complication of hemophilia. In patients who develop inhibitors, further treatment with factor VIII or IX concentrates is not recommended. Instead bypassing agents such as factor eight inhibitor bypass activity (FEIBA) or recombinant factor VIIa is used for these patients.

Inhibitors develop as a result of immune response to the factor injections received. They mostly develop in children with severe hemophilia who have severe mutations, and they run in families. It is not related to the dose or schedule of factor received.

In our country, private medical insurance does not cover most genetic diseases including hemophilia. Employer insurance, however, does cover this disease. Most government sector jobs provide insurance cover for hemophilia.

FEIBA is an acronym for “Factor Eight Inhibitor Bypassing Agent”. This is a drug used in patients of hemophilia A who develop inhibitors.

Yes. Compared to few years back, the availability of factor concentrates has improved dramatically in our country. Most states provide free factor concentrates to patients with hemophilia. Treatment is available in medical colleges, institutes, and many district hospitals under the government. In addition, hemophilia patient welfare organizations also help patients with provision of factor concentrates either free or at a subsidised rate.
Yes. There are many new factor concentrates that have extended half life which reduces the need for frequent doses. Also, currently newer nonfactor treatments are also available that can be administered once in 2–4 weeks, thus reducing the need for frequent injections.

Gene therapy is the only curative treatment available in hemophilia, where the correction of the genetic defect is achieved through insertion of the gene through a viral vector. It is successful in research trials and will be available for use to patients in our country also in a few years’ time. We encourage you to have regular conversations with your treatment center and with other parents of hemophilic children. Their expertise will go a long way in helping you to raise our child. Refrain from unverified knowledge circulated in social media.