Hereditary angioedema (HAE) caused by C1-esterase inhibitor deficiency is an autosomal-dominant disease caused by a mutation in the C1-inhibitor gene. It is a rare disease that is often worsened during pregnancy and childbirth. HAE, though uncommon but if untreated it may lead to maternal death. The case report presents the successful management of a 24 years old, G2P1, with hereditary angioedema caused by C1-esterase inhibitor deficiency. This patient was managed with a multidisciplinary approach by an obstetrician, an immunologist, an anaesthesiologist and a pediatrician. She had an uneventful antenatal period, labor was induced. She had precipitate delivery and soon after delivery had a flare up of the disease. It was successfully managed with fresh frozen plasma and close observation.

Keywords: C1-esterase inhibitor deficiency (HAE-C1-INH), CI inhibitor concentrate, Fresh frozen plasma (FFP), Hereditary angioedema (HAE), Plasma derived human C1 inhibitor concentrate (pdhC1INH), Pregnancy

INTRODUCTION

Hereditary angioedema (HAE) caused by C1-esterase inhibitor deficiency has an autosomal-dominant pattern of inheritance. It is characterized by frequent attacks of extensive, localized subcutaneous edema of body tissues. It may involve the extremities, genitalia, face, trunk or submucosal edema of upper airway or bowels. These symptoms may be debilitating. On occasions the flare-ups may be life-threatening when affecting the upper airways.\(^1\) Women with hereditary angioedema (HAE) are more likely to be symptomatic that men. Hormonal factors as puberty, contraception, pregnancy etc. Play a significant role in the precipitation of worsening of the condition in women.\(^2-4\) During pregnancy, attack rate may increase.\(^3-5\) Some studies have demonstrated increase in the attack rates with the subsequent pregnancies.\(^6,7\) Different effects of HAE on pregnancy and postpartum period have been noted. Some patients have flare-ups while some experience improvement of symptoms.\(^2\) HAE-C1-INH during pregnancy should be treated according to symptoms in pregnancy and previous history of flare-ups in previous child birth. Before labor and delivery prophylaxis may/may not be indicated; use of plasma-derived human C1-inhibitor concentrate (pdC1-INH) for acute attacks or as a short-term preventive treatment for pregnant women is recommended.\(^5,8-11\)

Due to its rare prevalence, there is lack of clinical knowledge in the management of HAE. In countries with very low prevalence of these cases, management is even more challenging. In addition, due to unavailability of the newly recommended drugs for HAE and their safety in pregnancy and lactation, treatment in pregnancy and in post-natal period remains a challenge for the clinicians.\(^10,11\) In places where the prevalence of HAE is very low and the drugs for treatment of HAE are not available, management of these patients with fresh frozen
plasma is promising and effective. The case below, presents a pregnant patient with hereditary angioedema, illustrates the symptomatic presentation in her post-natal period and successful treatment with fresh frozen plasma.

CASE REPORT

A 24 years old female, gravida 2, para 1 had regular antenatal visits to our hospital. She was a known case of hereditary angioedema (C1-Esterase Inhibitor Deficiency). This was diagnosed after she had an attack of the disease in the form of respiratory distress and facial edema after receiving analgesia for a dental procedure. Patient was intubated and hospitalized for a couple of days. She also had a brother and a cousin with the HAE-C1-INH.

Her previous pregnancy and post-natal phase were uneventful. In her current pregnancy she was asymptomatic, and her C1-Esterase Inhibitor protein level was low, 8.2.5mg/dl (normal range:195-345mg/dl). Levels of C2 and C4 were low while C1a was normal. In view of HAE-C1-INH, she had planned induction of labor at 40 weeks to avoid complications in labor. She had consultation with pulmonologist, neonatologist and anesthetist during her antenatal period and prior to decision for timing of delivery. The plan was to transfuse fresh frozen plasma before onset of labor to prevent an attack of HAE. Epidual analgesia was also planned. However, after induction of labor with prostaglandin E2 first dose, she had precipitate labor and delivered soon. Her delivery was spontaneous and uneventful without any complications. Within thirty minutes after her delivery, she developed facial itching followed by rashes all over her face and facial edema commenced. Rashes were non-urticarial and serpiginous. Her vitals were stable with no respiratory discomfort. She was assessed by the anesthesiologist immediately for the need of airway support. She received 4 units of fresh frozen plasma. Gradually her symptoms subsided, and she was monitored for next 6 hours in labor room. She had uneventful post-natal period later in the post-natal ward and was discharged home in stable condition after medical and anesthetic consultation. Follow up with immunologist was given. She had an uneventful postnatal period.

DISCUSSION

The above case presents successful management of patient with C1-INH-HAE with a multidisciplinary approach and hence preventing life threatening complications. HAE appears to have a benign course especially in third trimester and those with anticipated instrumental delivery, prophylaxis should be considered.

Vaginal delivery is the preferred mode of delivery to avoid anesthetic risks during cesarean section. Routine prophylaxis in uncomplicated labor is not recommended. In patients with recurrent attacks during pregnancy especially in third trimester and those with anticipated instrumental delivery, prophylaxis should be considered.

Prophylaxis with pdhC1INH is advised before forceps or vacuum extraction or cesarean section. Regional Anesthesia is preferred to endotracheal intubation. In patients with planned and emergency Cesarean pdC1-INH prophylaxis is recommended. Epidural analgesia is considered as the analgesia of choice.

HAE, though uncommon but if untreated it may lead to maternal death. It may also cause fetal and neonatal death. According to International consensus and practical guidelines on the gynecologic and obstetric management of female patients with hereditary angioedema caused by C1 inhibitor deficiency, “Attenuated androgens are contraindicated and should be discontinued before attempting conception. Plasma derived human C1 inhibitor concentrate (pdhC1INH) is preferred for acute treatment, short-term prophylaxis, or long-term prophylaxis. Tranexamic acid or virally inactivated fresh frozen plasma can be used for long-term prophylaxis if human plasma-derived C1-INH is not available. No safety data are available on icatibant, ecallantide, or recombinant human C1-INH (rhC1INH) as of now.” FFP may be used as a life-saving technique in emergency rooms to immediately treat cases of life-threatening angioedema. Though it is not the drug of choice, but it is fast-acting, effective, relatively safe and easily available in most emergency rooms. At places where pdC1-INH is not available, it can be used for acute flare-ups and short-term prophylaxis as well.

Some cases have also been reported to aggravate the symptoms, hence when FFP is used, patients should be closely monitored. WAO recommends at least 72 hours follow up in uncomplicated delivery. Lactation may also cause aggravation of symptoms and flare up but is still recommended as benefits outweigh the risks.
In patients with recurrent attacks during lactation and in cases where androgens are introduced, breast feeding should be discontinued. These patients should be given advice on contraception on discharge to prevent unwanted pregnancy, till the diagnosis if confirmed and patient is stabilized. Estrogen has been proved to precipitate attacks, therefore combined contraceptive pills are not indicated and progesterone pill must be preferred for contraception. Evaluation of C1-INH level and function should be done in post-natal period for newly diagnosed cases in pregnancy/lactation for confirmation of diagnosis. Family members should be counseled for support at home and to report to Emergency in case any flare-up is noted.

**CONCLUSION**

Essentially, diagnosis and management of HAE patients in pregnancy and post-natal period is challenging, notably in places with low prevalence. Early diagnosis and prompt treatment with FFP can be lifesaving. Physicians should be aware of the presentation of the disease and treatment of flare-ups to prevent maternal and fetal morbidity and mortality.

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**REFERENCES**


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