**ORIGINAL ARTICLE**

**Management of Pregnancy and Delivery in Patients With Hereditary Angioedema Due to C1 Inhibitor Deficiency**

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**Abstract**

Background and Objective: There is little information on pregnancy and delivery in patients with hereditary angioedema due to C1 inhibitor deficiency (C1INH-HAE). The aim of this study was to describe the effect of pregnancy and deliveries on symptoms of C1INH-HAE and review the need for and safety of treatments available during the study period.

Methods: Retrospective review using a purpose-designed questionnaire of 61 C1INH-HAE patients from 5 hospitals specialized in the management of HAE in Spain. The outcomes measured were number of pregnancies, changes in symptoms during pregnancy and delivery, mode of delivery, type of anesthesia during delivery, treatments received, and tolerance of treatments.

Results: We reviewed 125 full-term pregnancies (89 without a prior diagnosis of C1INH-HAE), 14 miscarriages, and 4 induced abortions. Patients reported an increased frequency of C1INH-HAE symptoms in 59.2% of pregnancies (74/125) and the presence of symptoms throughout pregnancy in 40% (50/125). Prophylactic C1INH-HAE therapy was used during 9 (7.2%) of the 125 pregnancies. Nine patients—in 11 pregnancies (8.8 %)—received treatment for acute attacks. Most deliveries (n=110, 88%) were vaginal. A cesarean section was necessary in 15 cases (12%). Short-term prophylaxis with pdhC1INH was administered before 14 deliveries (11.2 %); 111 deliveries (88.8 %) were performed without premedication and were well tolerated. Anesthesia was used in 51 deliveries (40.8%).

Conclusions: Pregnancy has a variable influence on the clinical expression of C1INH-HAE. Attacks tend to occur more frequently but not to increase in severity. Vaginal delivery was mostly well tolerated. pdhC1INH prophylaxis should be administered prior to cesarean delivery and is also recommended before vaginal delivery if there are additional risk factors. pdhC1INH should always be available in the delivery room.


**Resumen**

Antecedentes y Objetivo: Existe escasa información sobre la evolución del embarazo y el parto en pacientes con angioedema hereditario con déficit de C1 Inhibidor (AEH-C1INH). El objetivo del estudio fue describir el efecto de embarazo y parto en los síntomas de AEH-C1INH y la necesidad y seguridad de las terapias disponibles durante dicho período.
Introduction

Hereditary angioedema (HAE) due to C1 inhibitor deficiency (C1INH-HAE) is a rare disease [1]. In Spain, a minimal prevalence of 1.09 cases per 100 000 inhabitants has been reported [2]. Two subtypes of C1INH-HAE have been described: type I C1INH-HAE with reduced yet functional C1 inhibitor levels and type II C1INH-HAE, with normal or high C1INH protein levels but reduced C1INH function [3]. Another type of HAE with normal C1INH levels has also been described [4,5], but in this paper we focus just on types I and II.

C1INH-HAE is characterized by nonpruritic, nonpitting edema that typically affects different locations. Abdominal pain, distension, nausea, or vomiting may also be present secondary to edema of submucosal tissues of the gastrointestinal tract [6]. Upper airway involvement can be fatal, and mortality due to suffocation can be as high as 33% in inappropriately treated patients [7]. Estrogens, trauma, and stressful situations can worsen the course of disease and consequently, pregnancy and delivery may be special periods for female patients [8-10].

Three treatment options are available for C1INH-HAE: long-term prophylaxis (LTP), short-term prophylaxis (STP), and acute treatment. LTP consists mainly of attenuated androgens (AAs), but these can cross the placental barrier, possibly producing virilization, and should therefore be strictly avoided during pregnancy and lactation; antifibrinolytic agents are also used for LTP, though they are less effective [11,12]. Plasma-derived human C1 inhibitor (pdhC1INH) can be used for LTP when other treatments are contraindicated, ineffective, or poorly tolerated. STP with pdhC1INH is the most effective preventive therapy for patients undergoing surgery. pdhC1INH has traditionally been the treatment of choice for acute attacks during pregnancies; an alternative, though less safe, option is fresh frozen plasma [13,14]. Icatibant acetate, a B2 receptor blocker, as well as a recombinant version of the human C1 inhibitor protein, have been approved by the European Medicines Agency and the US Food and Drug Administration, and together with ecallantide, are licensed for the treatment of acute edema attacks in adult patients; they have not yet, however, been approved for use in pregnancy [15-17]. In this study, we describe the experience of 5 major HAE centers in Spain in managing C1INH-HAE during pregnancy and delivery with the aim of adding to the body of knowledge regarding the management of obstetric events in this setting.

Patients and Methods

Patients from 5 HAE reference hospitals in Spain were recruited for the study. The participating centers were Hospital Universitario Virgen del Rocío in Sevilla, Andalusia, Hospital Universitario La Paz in Madrid, Complejo Hospitalario Universitario de Vigo in Vigo, Galicia, Hospital Universitario Vall d’Hebron in Barcelona, Catalonia, and Hospital General Universitario Gregorio Marañón in Madrid. We performed a retrospective study of C1INH-HAE patients who had become pregnant before or after C1INH-HAE diagnosis. A specific questionnaire was designed and filled out with information from clinical charts or from telephone interviews when data were missing. Information about full-term pregnancies, miscarriages, and induced abortions was also included. The research ethics committees of Hospital Universitario Virgen del Rocío and Hospital Universitario La Paz approved the study.

The criteria to define worsening of symptoms compared with the baseline condition included an increase in attack frequency as well as the duration of single attacks. Increased severity was also considered a criterion of worsening but there were no cases (ie, there were no changes in the distribution of sites of involvement). The criteria to define an improvement in HAE symptoms included no symptoms, fewer symptoms, or milder attacks.

Descriptive statistics were used to summarize and analyze the collected data. Analysis was performed using the IBM SPSS v19.0 statistical package. Quantitative variables were described as means and ranges; qualitative variables were recorded as absolute values and percentages. Categorical variables were compared using the χ² Pearson test. P values of less than .05 was considered to be statistically significant.
Results

We reviewed 125 pregnancies in 61 patients as well as 14 miscarriages and 4 induced abortions. The percent distribution of patients and full-term pregnancies among the 5 study hospitals are shown in the Figure.

The mean age of patients at the time of C1INH-HAE diagnosis was 30.6 years (range, 6-60 years), and the mean age at the start of pregnancy was 27.1 years (range, 16-42 years). There was a prior diagnosis of C1INH-HAE in just 44 (30.7%) of the 143 initial gestations and in 36 (28.8%) of the 125 full-term pregnancies. Thirty-five women (57.4%) were not aware that they had HAE during 89 pregnancies (71.2%). Twenty of the 61 patients had only 1 pregnancy. Data were collected between the years 2006 and 2010 and corresponded to pregnancies that had occurred more than 20 years previously in nearly half of the patients (49.2%), 6 of whom had only been pregnant once.

Only 13 patients (21.3%) were receiving LTP before they became pregnant. Nine patients had been receiving AAs. Seven of these stopped AA therapy before they conceived and the other 2 stopped 1 month after conception, on confirmation of their pregnancy.

A greater frequency or duration of acute attacks was reported for 59.2% of the pregnancies, no changes with respect to baseline symptoms were reported in 26.4% of cases, and symptoms improved in 14% of cases. There were no differences in attack severity from one trimester to the next, and in 40.0% (50/125) of pregnancies HAE symptoms were present throughout the pregnancy. Patients with more than 1 gestation (67.2%) generally described a similar course for each of their pregnancies (similar C1INH-HAE symptoms in 85.4% of cases). There were no changes in sites of involvement but there was an increase in the frequency of mild abdominal crises. Non-life-threatening symptoms were reported in the 125 full-term pregnancies or abortions.

We found no statistically significant differences for disease course on comparing percentages between the group of women with a known C1INH-HAE diagnosis before pregnancy (n=36) and the group of women with an unknown C1INH-HAE diagnosis at the time of pregnancy (n=89) (Table 1).

LTP was only used during 9 of the 125 pregnancies: epsilon-amino-caproic acid was used in 1 case, tranexamic acid in 2 cases, AAs in 2 cases, and pdhC1INH (Berinert, CSL-Behring) in 4 cases. In the 2 cases in which AAs were temporarily used (for 8 and 12 weeks), the drug was administered after confirmation that the fetus was male. (As mentioned previously, 2 other patients interrupted LTP with AAs when they became aware they were pregnant.)

Our patients reported 14 miscarriages, most of which occurred during the first trimester. Additionally, there were 4 registered abortions. There were also 3 fetal deaths and 1 premature delivery with complications (deafness and visual problems). None of the miscarriages or cases of fetal damage occurred in patients temporarily exposed to AAs.

Nine patients received treatment for an acute attack during 11 pregnancies. pdhC1INH (Berinert) was administered in all 9 cases and 1 of the patients additionally received tranexamic acid and other corticosteroids. This last patient was not diagnosed with C1INH-HAE until 5 years later.

Seven patients were administered a total of 618 vials of pdhC1INH 500 U (Berinert) to treat acute attacks and as LTP (4 cases) in 9 pregnancies (average of 4.16 vials/mo); 1 patient received, as LTP, a total of 356 vials in 2 consecutive pregnancies (average 4.94 vials/wk); no adverse effects were reported.

None of the newborns developed health problems or experienced adverse effects attributable to any of the drugs used.

The vast majority of deliveries (n=110, 88%) were vaginal; forceps and vacuum extraction were used in 9 and 5 deliveries, respectively (Table 2). Cesarean sections were necessary in 15

<table>
<thead>
<tr>
<th>Previous diagnosis of C1INH-HAE</th>
<th>Yes (n=36) (28.8%)</th>
<th>No (n=89) (71.2%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Worsening of C1INH-HAE symptoms during pregnancy (%)</td>
<td>No 51 (40.8) 12 (33.3) 39 (43.8)</td>
<td>Yes 74 (59.2) 24 (66.7) 50 (56.2)</td>
</tr>
</tbody>
</table>

Abbreviation: C1INH-HAE, hereditary angioedema due to C1 inhibitor deficiency.

Table 2. Mode of Delivery and Type of Anesthesia

<table>
<thead>
<tr>
<th>Mode of Delivery</th>
<th>Type of Anesthesia</th>
<th>Total (n=125)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cesarean section (n=15)</td>
<td>Epidural General</td>
<td>31 17 3 74</td>
</tr>
<tr>
<td>Vaginal delivery</td>
<td>Pudendal Block</td>
<td>21 4 0 71</td>
</tr>
<tr>
<td>Noninstrumental (n=96)</td>
<td>No Anesthesia</td>
<td>21 4 0 71</td>
</tr>
<tr>
<td>Vacuum (n=5)</td>
<td>Forceps (n=9)</td>
<td>4 2 3 0 3</td>
</tr>
</tbody>
</table>

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HAE and pregnancy report on few pregnancies and deliveries (between 1 and 3). Our study is the largest to date and can be considered representative of the approach to and management of pregnancies and deliveries in C1INH-HAE patients in Spain in recent decades.

Mild aggravation of C1INH-HAE symptoms was experienced in 59.2% of the pregnancies in our series; this figure is similar to previous reports [18,19]. The mild worsening detected is more likely to be attributable to changes associated with pregnancy rather than to discontinuation of HAE treatment, as most patients were not receiving regular treatment before they became pregnant.

No statistically significant differences were observed for course of disease on comparing patients with a previous diagnosis of C1INH-HAE and those without one (Table 1). Of the 36 pregnancies in which there was a confirmed diagnosis of HAE before pregnancy, there was an improvement in clinical signs in 6 cases (16.7%) and no changes in another 6.

The majority of the 41 patients with more than 1 pregnancy (85.4%) noted that the disease manifested itself in a similar way during each pregnancy and only 4 patients described a change in symptoms from one pregnancy to the next. The course of symptoms has been reported to vary greatly between pregnancy trimesters [18,19], but in more than half of the pregnancies in our series, there were reports of worsening of symptoms throughout the pregnancies, with no clear differences observed between trimesters.

Chinniah and Katelaris [20] described a group of 7 patients with 16 pregnancies who experienced fewer attacks as the pregnancy progressed and more attacks during the first trimester. In all cases, the patients had edema attacks, but the authors did not report whether symptoms worsened during pregnancy in comparison with previous or posterior periods.

### Discussion

We have reported on the course and management of 125 full-term pregnancies and deliveries, 14 miscarriages, and 4 induced abortions in 61 C1INH-HAE patients through a nationwide cooperation study. A clear set of international C1INH-HAE guidelines containing recommendations for delivery and pregnancy follow-up and information on the benefits and risks associated with each of the available treatments was published only recently [12,13]. Three case series have been published [18-20], but most publications on

<table>
<thead>
<tr>
<th>Patient</th>
<th>Year</th>
<th>Pregnancy Number</th>
<th>Age, y</th>
<th>Type</th>
<th>Reasons</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1975</td>
<td>Second</td>
<td>28</td>
<td>Acute</td>
<td>Dystocia</td>
</tr>
<tr>
<td>2</td>
<td>1983</td>
<td>First</td>
<td>19</td>
<td>Elective</td>
<td>Narrow pelvic opening</td>
</tr>
<tr>
<td>3</td>
<td>1983</td>
<td>First</td>
<td>23</td>
<td>Acute</td>
<td>Anomalies of umbilical cord</td>
</tr>
<tr>
<td>4</td>
<td>1984</td>
<td>Third</td>
<td>39</td>
<td>Elective</td>
<td>Other complications</td>
</tr>
<tr>
<td>5</td>
<td>1985</td>
<td>Third</td>
<td>25</td>
<td>Acute</td>
<td>Dystocia</td>
</tr>
<tr>
<td>6</td>
<td>1993</td>
<td>Second</td>
<td>40</td>
<td>Elective</td>
<td>Other complications</td>
</tr>
<tr>
<td>7</td>
<td>1994</td>
<td>First</td>
<td>29</td>
<td>Acute</td>
<td>Dystocia</td>
</tr>
<tr>
<td>8</td>
<td>1994</td>
<td>Second</td>
<td>27</td>
<td>Elective</td>
<td>HAE diagnosed during pregnancy</td>
</tr>
<tr>
<td>9</td>
<td>1996</td>
<td>First</td>
<td>27</td>
<td>Elective</td>
<td>Narrow pelvic opening</td>
</tr>
<tr>
<td>10</td>
<td>1998</td>
<td>First</td>
<td>28</td>
<td>Elective</td>
<td>Unknown. Known diagnosis of HAE</td>
</tr>
<tr>
<td>11</td>
<td>2001</td>
<td>Second</td>
<td>37</td>
<td>Elective</td>
<td>Narrow pelvic opening</td>
</tr>
<tr>
<td>12</td>
<td>2002</td>
<td>First</td>
<td>27</td>
<td>Acute</td>
<td>Fetal bradycardia. Known diagnosis of HAE</td>
</tr>
<tr>
<td>13</td>
<td>2005</td>
<td>First</td>
<td>36</td>
<td>Elective</td>
<td>Unknown. Known diagnosis of HAE</td>
</tr>
<tr>
<td>14</td>
<td>2006</td>
<td>Second</td>
<td>35</td>
<td>Elective</td>
<td>Fetus in breech position. Known diagnosis of HAE</td>
</tr>
<tr>
<td>15</td>
<td>2007</td>
<td>Third</td>
<td>28</td>
<td>Acute</td>
<td>Dystocia</td>
</tr>
</tbody>
</table>

Abbreviation: HAE, hereditary angioedema.
In our case series, none of the 14 miscarriages were directly related to angioedema attacks and this rate of 10.1% can be considered low compared with average rates reported for the general European population (10%-20%) [10]. The decision to terminate the pregnancy was at least partially related to the diagnosis of HAE in 2 of the 4 voluntary interruptions. One of the 3 fetal deaths was due to a congenital cardiopathy and 2 (in the same woman) were due to Rh incompatibility. There was just 1 newborn who developed complications because of premature delivery (at 6 months). None of these cases were related to the temporary use of androgens.

Almost all studies of patients with C1INH-HAE have reported good progression of HAE symptoms during pregnancy and delivery. Although HAE symptoms may be present, they are usually mild and rarely life-threatening. Mc Glinchey et al [21] reported a life-threatening laryngeal angioedema attack in the 25th week of gestation that was treated with fresh frozen plasma and 2000 U of pdhC1INH and resolved completely. Postnikoff and Pritzker [22] wrote about the only death reported of a pregnant HAE woman 110 hours after delivery. This patient developed perineal swelling and a purulent discharge from the episiotomy 48 hours postpartum. Autopsy revealed edema of the subcutaneous tissues, which was most prominent in the perineal region, with severe effusions present in all body cavities and septic shock.

LTP during pregnancy was seldom used in the pregnancies reported in this case series; in some cases the patients had not yet been diagnosed with C1INH-HAE, while in others presumably little was known at the time about the safety of C1INH-HAE treatments. We administered C1INH concentrate as LTP in 4 pregnancies in 2009 and 2010; the drug was administered off-label, with the patients’ informed consent, due to a clear exacerbation of symptoms following discontinuation of effective AA therapy.

In our series vaginal delivery without prophylactic treatment was well tolerated, and although trauma and stress are known to trigger attacks, very few patients developed mild local edema. There have been many isolated reports of pregnant C1INH-HAE patients who experienced no edema attacks after a vaginal or cesarean delivery with prophylactic pdhC1INH treatment [23-28]. There have also been reports of 2 cesarean deliveries [29] and 2 vaginal deliveries [30] that were well tolerated without prophylactic therapy. Anecdotal reports of STP with fresh frozen plasma and danazol prior to delivery, with good outcomes, have also been published [31,32]. In the series reported by Martinez-Saguer et al [19], all the patients received prophylactic pdhC1INH treatment prior to delivery. Chinniah and Katelaris [20] did not report any symptomatic deliveries in their case series, although prophylactic pdhC1INH was used in only 5 of the 16 deliveries. In the series by Czaller et al [18], prophylactic treatment prior to delivery was used in 9 of 82 deliveries, with good tolerance in all cases; the same authors described asymptomatic deliveries in the 73 deliveries performed without prophylaxis. The proportion of deliveries without pretreatment and the good outcomes in cases in which prophylaxis was not used matched the findings of our case series.

Only 5 of 15 women who underwent a cesarean section in our series knew about their condition and were treated with pdhC1INH prior to delivery. Of the 10 cesarean deliveries that did not include prophylactic treatment, a single patient experienced mild HAE symptoms during delivery and 48 hours postpartum and subsequently required treatment with pdhC1INH. Czaller et al [18] described the use of prophylactic treatment prior to cesarean delivery in 1 of 8 cases and reported no symptoms in any of these deliveries. Martinez-Saguer et al [19], in turn, reported 8 cesarean sections out of 35 deliveries; in all cases, the patients underwent prophylactic treatment with pdhC1INH immediately before delivery and no attacks were reported.

Our study has certain limitations. Information was obtained retrospectively through a questionnaire completed using clinical records and telephone interviews in the case of missing data. Due to the retrospective nature of the study, the information corresponded to pregnancies from several years earlier (up to 20 years in nearly half of the patients). It is also important to note that C1INH-HAE had not been previously diagnosed in 89 pregnancies, and in the majority of cases, a diagnosis was not made until several years later. Therefore, the patients may have underestimated the severity of symptoms or failed to associate them with C1INH-HAE. Nevertheless, the fact that most patients had not been diagnosed with C1INH-HAE before conception and did not receive any treatment during pregnancy provides interesting insights into the natural course of C1INH-HAE during pregnancy and labor.

In conclusion, after reviewing data from our series we can conclude that pregnancy appears to have a variable influence on the clinical expression of C1INH-HAE, and may differ from one patient to another and to a lesser extent, from one pregnancy to the next. Attacks tend to occur more frequently but they do not appear to increase in severity. pdhC1INH prophylaxis should be administered prior to cesarean delivery and is also highly recommended for vaginal delivery in patients with additional risks factors or severe C1INH-HAE symptoms during pregnancy or previous deliveries. pdhC1INH (2000 U) should always be available in the delivery room and during hospitalization. To improve outcomes in pregnant women with C1INH-HAE, it would be wise to maintain observation for 48 hours postpartum.

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Conflicts of Interest

Dr Caballero has received sponsorship for educational purposes, has been paid for providing consultancy services, and has taken part in clinical trials sponsored by Jerini AG/Shire, CSL-Behring, Dyax Corp, Pharming NV, and Viropharma Pharmaceutical.

Dr Cimbollek has received sponsorship for educational purposes and has taken part in clinical trials sponsored by CSL-Behring, and Pharming NV.
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Dr. Ignacio Larco has taken part in a clinical trial sponsored by CSL-Behring.

Dr López-Serrano has received sponsorship for educational purposes, been paid for providing consultancy services, and taken part in clinical trials sponsored by Jerini AG/Shire, CSL-Behring, Dyax Corp, and Pharming NV.

Dr. Marcos has been paid for providing consultancy services and received speaker's fees and sponsorship for educational purposes from Jerini AG/Shire.

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