Introduction

Hereditary Angioedema (HAE) is a rare disease with an incidence of about 1:50,000 [1]. According to current estimates prevalence of this autosomal dominant congenital disorder in Germany is roughly 1,600. As HAE is still relatively unknown and the symptoms are quite unspecific, resembling those of other diseases, a considerable number of unidentified cases is to be expected.

Due to a genetic variant HAE patients develop lifelong recurrent attacks of swellings, which may last several days if left untreated [2]. The edema may form in almost any part of the human body, predominantly the extremities and the digestive tract is affected [3]. A large part of the patients experiences prodromal symptoms like thirst or fatigue. There are also some well-known trigger factors like mechanical irritation and stress [4]. Interindividual variability in disease manifestation is huge, this including also attack frequency. Edema in the upper aero-digestive tract can be life threatening [5].

In most patients HAE is caused by a mutation of the “SERPING1” gene on the chromosome 11, which codes the C1 esterase inhibitor (C INH) [6]. C1 INH physiologically can implicate uncontrolled formation of the tissue hormone bradykinin. Accumulation of bradykinin can lead to edema development. The underlying gene defect can cause either a reduced C1 INH synthesis (HAE type 1, about 85% of patients) or the formation of a non-functional C1 INH (HAE type II, about 15% of patients) [7]. Rarely a third type is encountered, which was described simultaneously by Bork et al. and Binkely and Davis only in 2000. It is characterized by normal levels of functional C1 INH (in former times also called HAE type III) [8,9]. In some of these patients a mutation in the factor XII or the plasminogen coding gene can be detected [10,11].

The patient’s history including predominantly also the family history is mostly paving the way for suspecting HAE. However, there are de novo mutations in as many as a quarter of the patients, i.e. a negative family history does not rule out HAE [12].

The diagnosis is then based on the qualitative (=activity) and quantitative (=concentration) C1 INH values in blood plasma. In the typical HAE patient also the values of the C4 complement are too low [13]. The diagnosis of a HAE in case of normal C1 INH values is based on a positive family history in combination with the typical clinical signs. In some patients it can be further supported by the confirmation of a gene mutation.

In Germany there are two approved treatment options for HAE: Either substituting the missing or non-functional enzyme as intravenous C1 INH concentrate or controlling the bradykinin by subcutaneous injections of icatibant (investigational name...
HOE 140), which is a bradykinin-2-receptor antagonist [1,14]. Most of the therapeutic options are approved for home therapy by the patients or their relatives. Based on the attack frequency, the perceived burden of suffering and the potential trigger factors a short-term or long-term prophylaxis can be indicated in addition to acute treatment [15,16]. Unlike the more frequent mast cell mediated urticarial or allergic swellings, edema in HAE do not respond at all or only very little to antihistamines or corticosteroids [5]. Other drugs like kallikrein inhibitors were approved recently in the US, their market authorization for European countries is still pending.

The variability of the clinical course in combination with the unspecific symptoms and the scarce knowledge of the disease often results in a situation where the patients see many different specialty doctors until their suffering is finally diagnosed. This comprises general practitioners, dermatologists, ENT specialists and pediatricians. The time between the first symptoms and the correct diagnosis can be several decades and can include inadequate treatments based on false diagnoses [17]. Misdiagnoses associated the disease with the psychiatric field and led to the name “angioneurotic edema”, which was replaced by HAE after the identification of the underlying congenital defect.

Relatively little is known of the risk factors that may influence the course of the disease. The objectives of our survey study were to evaluate important parameters of HAE in Germany and to assess, if the BMI might have a role in disease severity.

Patients and Methods

This was a questionnaire-based study in adult patients with HAE (>18 years of age). The following parameters were addressed—also the effect of the introduction of adequate therapy was tried to assess retrospectively wherever possible:

- Demographic characteristics (age, sex).
- Height, weight, BMI.
- HAE type.
- Family history.
- Specialty responsible for making correct diagnosis.
- Prodromes.
- Attack sites.
- Frequency and duration of attacks.
- Days of sick leave.
- Medication.

After consultation with the statisticians and regarding the incidence of HAE, a minimum of 50 participants was targeted to obtain a realistic number and representative sample of patients with this rare disease.

We asked all HAE patients treated in our ENT department to participate. All patients were diagnosed by laboratory tests (levels of C1 INH protein <0.17g/l and/or dysfunctional C1 INH protein <70%). HAE with normal C1 INH was diagnosed when patients presented the above-mentioned clinical symptoms, had family members with history of angioedema and chronic urticaria was excluded. Patient questionnaires were collected between July 2015 and July 2016.

The evaluation of the questionnaires was done with descriptive statistics for all parameters. A potential correlation of BMI to attack frequency, location, duration of prodromes was statistically tested. Patients were assigned to two groups: Group 1 = BMI <30 kg/m²; group 2 = BMI ≥ 30 kg/m². T-tests were carried out for these intergroup comparisons. A p-value of ≤ 0.05 was considered statistically significant, a p-value of ≤ 0.01 was considered highly significant. For all statistical evaluation the IBM SPSS Statistics 21 software was used. The study was approved by the review board of Ulm University (Application number 104/15).

Result

Demographic data

Filled in questionnaires from 61 patients were collected during the recruiting period. The mean patient age at inclusion was 43.2 years (range 20-74 years). 72.1% of the participants were female.

The average BMI of female participants was 24.9 ± 5.8 kg/m², male participants had an average BMI of 26.9 ± 4.4 kg/m². This means that both values are at the upper limit of normal weight, defined by the World Health Organization (WHO).

All patients were German nationals and living in Germany upon inclusion. Most of the patients came from southern Germany, which is due to the catchment area of the ENT clinic in Ulm.

Diagnostics: HAE type and family history

29 of the 61 participants (47.5%) were diagnosed as HAE type I, 21.3% as HAE type II, and 8.2% of the patients reported a diagnosis of HAE with normal C INH values. 14 patients (23.0%) did not know their diagnosed type of HAE.

44 patients (72.1%) stated that further family members were affected by HAE. In 41% of the patients only the ancestors were affected, in 18% of cases only the offspring had been diagnosed also with HAE, and in 13.1% of the participants the ancestors as well as the offspring was affected. 24.6% of all patients had no relatives with HAE.

26.2% of the participants reported that their diagnosis was made by a dermatologist. 14.8% had been diagnosed by an ENT specialist or a general practitioner, respectively. 34.4% of the patients were unable to define what specialty doctor made their diagnosis but could state only more general aspects – e.g. that it was done in a hospital or by an allergist (Figure 1).

Symptoms

Fatigue was the most frequently reported prodrome before angioedema attacks in 37.7% of the patients. Psychiatric symptoms comprising mood swings, agitation, and irritability were stated by 18% of the HAE patients as prodromes. Among the 5 most frequently reported prodromes were also thirst, skin conditions like the Erythema marginatum (Figure 2) as well as...
paresthesia’s. Multiple mentions were possible for this question (Figure 3).

Also, regarding the affected locations in the body all answers could be checked that apply. The digestive tract and the extremities were reported most frequently by 70.5% of the patients. 23% of the patients stated that the head/neck region was among the predominantly affected parts of the body (Figure 4).

There was a high variability in the attack frequencies. 34.4% of the HAE patients reported to have acute swellings more than once a month but less than once a week. 26.2% suffered from swellings at least once a week. Only 2 patients (3.3%) reported swelling attacks less than once in a year (Figure 5). If left untreated, the edema lasted on average 3.8 (± 2.7) days. The duration ranged from a couple of hours to more than 10 days.

**Therapy**

The yearly mean number of days of sick leave from work or school was reported as reduced by 81.3% from 24 days without therapy to 4.5 days under adequate treatment.

47.5% of the HAE patients were prescribed a human or recombinant C1 INH concentrate for their acute therapy, 8% had C1 INH concentrate and icatibant for acute therapy. 23% were treated only with the B2 antagonist icatibant for their acute swellings.

34.4% of the participants were on a prophylactic regimen. 32.8% used C1 INH concentrate, which is approved for prophylaxis. 1 patient used icatibant, which means an off-label use in Germany.

**Correlation between BMI and disease severity**

There were no significant correlations between the BMI and the duration or frequency of the attacks, the location of the swellings or the prodromes (p ≥ 0.05). There was a tendency towards a higher frequency of attacks in the HAE patients in the high BMI group (>30 kg/m²): All of them stated to have at least 3 attacks per year while 10.2% of the patients in the low BMI group reported to have less than 3 attacks in a year (Figure 6).
Discussion

Our prospective questionnaire based study evaluated clinical characteristics of HAE and the BMI as a potential risk factor for a more severe clinical course of this rare disease in a group of 61 adult German patients.

It was demonstrated that most of the patients were diagnosed by a dermatologist. In Germany there are currently seven large HAE centers, which are allocated to different medical specialties. They can be led by dermatologists, ENT specialists, internists, or pediatricians. The objective is to allocate each HAE patient to one center that takes care of their individual treatment concept.

Further analyses of our questionnaire study targeted the more general course of the disease. The attack frequency has a major impact on the degree of suffering of the patients. Therefore, frequent attacks are a clear indication for starting a prophylactic treatment. In a Swiss study with 104 HAE patients a comparably broad range of attack frequencies was found. 39% of the HAE patients from Switzerland stated that they had an attack more often than once per month. Another German study by Bork et al. the affected parts of the body were divided into “affecting the skin” — including the extremities and the face, and other sites of manifestation like mucosa, or the internal organs. Out of the 209 HAE patients from this cohort, 100% reported skin manifestations, 97% gastrointestinal attacks, and 54% stated that they had swellings of the larynx [18]. A study comprising Hungarian and Israeli children with HAE showed an even larger variability of the attack frequency than our study. 26.5% of the participants had 1-5 attacks per year, 23.5% reported 25-50 yearly attacks, and almost one third of the patients had been symptom free during the previous year. Also, the negative impact of a high attack frequency on the quality of life was shown in this study [19].

A large German study carried out by Magerl et al. studied the prodromes of HAE. It was shown, that about 79% of the 365 participating patients regularly suffered from prodromal symptoms. Fatigue, nausea, irritability, and skin conditions were the most frequently encountered signs, which is in line with our own results. Significantly more women than men reported to have prodromes in this German trial (83% vs. 73%; p<0.05) [20].

In our patient cohort HAE attacks most frequently affected the gastrointestinal tract and the extremities. This in alignment with the results from the Swiss study by Steiner et al., who found that the angioedema was localized predominantly in the abdomen (43% of cases), followed by swellings in the extremities [18]. Another German study by Bork et al. the affected parts of the body were divided into “affecting the skin” — including the extremities and the face, and other sites of manifestation like mucosa, or the internal organs. Out of the 209 HAE patients from this cohort, 100% reported skin manifestations, 97% gastrointestinal attacks, and 54% stated that they had swellings of the larynx [21]. Already in 2003 Bork et al. found a ratio of 1:70:54 for laryngeal to skin to abdominal attacks [22]. The “Icatibant Outcome Survey” by Caballeró et al. demonstrated 2017 that abdominal attacks predominated (58%) and they also showed, that the majority of attacks (90%) affect only a single anatomical site [23].

The therapy in our cohort proved to be in line with the treatment regimens recommended by the guidelines, which is based on C1 INH concentrates and bradykinin 2 receptor antagonists [24]. According to the individual clinical picture, acute medication schemes with or without additional prophylactic regimen were administered. The efficacy of a long-term treatment with C1 INH concentrates and the associated significant improvements in quality of life in HAE patients with frequent attacks was shown in several studies [15,16,25].

Consistent results from several studies showed that HAE attacks can be triggered by hormones like estrogen and that extraovarian estrogen is predominantly synthesized in visceral fat tissue by the aromatase cytochrome p450 [26,27]. Therefore, we studied a potential correlation between the BMI of the HAE patients with their clinical course. While we found a tendency towards a higher attack frequency in the obese patients, there were no statistically significant differences found in any parameter for the sub-group with a BMI ≥ 30 kg/m² when compared to those with a BMI <30 kg/m². There is a paucity of data regarding obesity as a risk factor in HAE. Bernstein et al. studied if a standardized, weight adjusted dosing of 20 IU/kg of C1 INH concentrate is effective in obese patients. The medication was found equally effective and safe in all patients, irrespective of their BMI [28]. As of now, no biomarker has been identified, that correlates with the individual clinical course of the disease — e.g. with the attack frequency. Not even fundamental factors like the type of HAE, the C1 INH activity, or the underlying mutation are
no reliable predictors for the severity of the disorder. Current basic research studies target to find biomarkers, predictors, and predisposing factors for swelling attacks, in order to better understand this individually so variable disease and to then be able to treat more specifically \[29,30\].

One general limitation of our study is the questionnaire-based set-up. However, this scenario allowed for reaching a relatively large patient group in a short period of time and to provide the patient perspective directly. Another draw-back is the retrospective assessment of some parameters like the days of sick leave before the introduction of adequate therapy. While no accurate numbers can be expected, we believe that the gross picture of the situation is reflected to a sufficient extent.

**Conclusion**

The therapeutic options based on a better understanding of the disease HAE made considerable progress over the past decades. But especially concerning the individual differences of HAE patients, many issues are still unresolved. With future research on the clinical course, fundamental scientific aspects and carefully adjusted therapy, we hope that patients with the incurable disease HAE may be able to lead an almost normal life. Further studies assessing individual factors reflecting the disease severity like trigger factors or biomarkers may be considered.

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JH, TKH and JG designed the study and the questionnaire and wrote the manuscript. JH, PJS and AH edited data and figures, AH and BM developed and performed the statistical analysis. All authors read and approved the final manuscript.

**Conflict of Interests**

JH, TKH and JG report non-financial support from Shire Deutschland GmbH and CSL Behring outside the submitted work. BM, AH and PJS report no conflicts of interest.

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