Increased Frequency of Guillain-Barré Syndrome in HIV Infection: A Prospective Cohort Study

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Abstract

Background: Guillain-Barré syndrome (GBS) has been reported in HIV-positive individuals, but the incidence and characteristics in this group of patients has not systematically been investigated beyond case reports and retrospective series. The aim of this study was to compare the incidence and characteristics of GBS in HIV positive and -negative individuals.

Methods: We performed a prospective, comparative study over a 3 ½ year period in the Western Cape province of South Africa. All adult patients with GBS were included and classified into 2 groups based on HIV status. The two groups were compared with regards to clinical, electrophysiological and laboratory features. Patients were followed until stable or recovered, for a maximum of 12 months.

Results: 28 patients were included in the study, of which 15 were HIV-positive. Using estimated HIV prevalence data for the same geographical area during the study period, the incidence of GBS in HIV-positive patients was calculated to be 18.74, 95% CI [7.69, 40.60] times higher than in HIV-negative patients. Except for the frequency of hyponatraemia, there were no statistically significant differences between the 2 groups with regards to presenting features, severity of illness, GBS subtypes, and treatment response. GBS occurred in all stages of HIV infection, and was the presenting feature of HIV infection in 13 patients.

Interpretation: The incidence of GBS is strikingly increased in HIV infection. The reason for this is still uncertain, but can probably be explained by immune dysregulation. HIV infection does not appear to influence the short term outcome of GBS.

Keywords: Guillain-Barrésyndrome; Acuteinflammatory demyelinating polyneuropathy; Acute inflammatory polyneuropathy; HIV; AIDS

Introduction

Guillain-Barré syndrome (GBS) is an acute inflammatory condition affecting peripheral nerves and nerve roots, often preceded by a flu-like illness or gastroenteritis [1,2]. Rapidly progressive generalized weakness is the hallmark, and 17-30% of cases develop severe respiratory weakness requiring ventilator support [3]. Treatment consists of intravenous immunoglobulin (IVIg) or plasma exchange (PE), and the majority of patients recover well [4]. However, despite advances in intensive care, the mortality rate is still 4-15%, and up to 20% of survivors are permanently disabled [5]. The incidence of GBS is between 1.2 and 1.9 per 100 000 [5–8]. However, because this information is derived from populations with a low prevalence of HIV infection, it does not provide information about the incidence of GBS in HIV-infected individuals. Although GBS has been reported in HIV-positive individuals [9–12] and is regarded as more frequent in this group [13], data beyond case reports and small case series is scarce. Furthermore, although the presentation and course of GBS is thought to be similar in HIV-positive and -negative an individual, no systematic investigation has been performed to explore this.

We performed a prospective study to compare the epidemiology, presentation and course of GBS between HIV-positive and –negative patients. Additionally, we aimed to characterize the profile of HIV-positive patients who develop GBS with regards to CD4+ count, concomitant opportunistic infections, and the use of anti-retroviral treatment.

Patients and Methods

Patients

We performed a prospective, observational study to detect all cases of Guillain-Barré syndrome in HIV-positive and –negative individuals in Tygerberg Hospital, a tertiary referral centre in the Western Cape province of South Africa, for a 3½ year period (July 2008 to December 2011). Patients were included if they were 18 years old, met accepted criteria for GBS [14], and presented within 4 weeks of symptom onset. Patients with pre-existing serious systemic or neurological disease that could complicate the diagnosis or course of GBS were excluded, as well as patients who were unable or unwilling to attend follow-up at Tygerberg Hospital. In the public healthcare system in the Western Cape, specialist neurology and electrodiagnostic services are only available in tertiary hospitals. Consequently, all potential cases of GBS requiring diagnostic workup are referred to tertiary hospitals. Referrals...
Assessments

A thorough disease history was obtained and all patients underwent a detailed clinical evaluation. Disease severity was assessed by means of the GBS disability and arm scales, [16] and the MRC sum score. The MRC sum score is a summation of the MRC grades (0–5) of the following muscle pairs: shoulder abductors, elbow flexors, wrist extensors, hip flexors, knee extensors, and ankle dorsiflexors. The sum score ranges from 0 (total paralysis) to 60 (normal strength).

Bilateral nerve conduction studies, including motor-, sensory- and F-responses were performed on all patients. Motor- and F-responses were investigated in the median, ulnar, common peroneal and posterior tibial nerves. Sensory studies included median palmar, ulnar palmar and sural responses. For each patient, results obtained were classified based on accepted electrophysiological classification criteria for GBS [17]. Diagnostic categories include normal, primary demyelinating, primary axonal, unexcitable and equivocal. Additionally, abnormal studies were further classified according to modalities involved (motor, sensory or both). These categories, in combination with clinical information (absence or presence of sensory abnormalities), were then used to classify individual patients into one of three subtypes of Guillain-Barré syndrome, namely acute inflammatory demyelinating polyneuropathy (AIDP), acute motor and sensory axonal neuropathy (AMSAN) or acute motor axonal neuropathy (AMAN), or undetermined if criteria for none of these were met. HIV testing was performed and patients were divided into the two groups according to serostatus. Results of CSF analysis were available for 13 HIV-positive and 11 HIV-negative patients. In addition, the stage of HIV-infection, CD4+ count and the presence of opportunistic infections were also assessed in HIV-positive patients.

Statistical analysis

The two patient groups were compared with regards to clinical, laboratory and electrophysiological variables. Data was analysed using Microsoft Excel 2010. Differences in metric variables between the two groups were tested using the independent t-test (two-tailed), while differences in categorical variables were tested using Fisher’s exact test. P-values less than 0.05 were regarded as significant.

Follow up

Patients were followed up every threemonths until (1) complete recovery, or (2) no further improvement at two consecutive visits or, (3) one year from admission. At follow-up, patients were reassessed by means of the GBS disability scale, GBS arm scale, and the MRC sum score.

Results

Comparison between the HIV-positive and –negative groups at presentation

A comparison of the clinical and laboratory features between the two groups at the time of presentation to hospitalis provided in Table 1. There were no statistically significant differences between the two groups except for the number of patients with serum hyponatraemia, which was more commonly noted in the HIV-positive group. Of the ten HIV-positive patients who reported a preceding infection, 7 had a respiratory tract infection and 3 had a diarrhoeal illness, whereas 3 of the 8 HIV-negative patients reported a respiratory tract infection and 5 a diarrhoeal illness. This difference was not statistically significant. Eleven (73%) of the HIV-positive patients were diagnosed with AIDP and1 (7%) with AMAN. In the three remaining patients in the HIV-positive group, the subtype could not be determined because the conduction studies were either normal (2 patients) or equivocal (1 patient). Of the HIV-negative patients, 10 patients (77%) were diagnosed with AIDP, 1 (8%) with AMAN and 2 (15%) with AMSAN.

<table>
<thead>
<tr>
<th>Clinical features</th>
<th>HIV-positive (n=15)</th>
<th>HIV-negative (N=13)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median age [years] (range)</td>
<td>42 (28-65)</td>
<td>46 (19-78)</td>
</tr>
<tr>
<td>Male</td>
<td>11 (73%)</td>
<td>10 (77%)</td>
</tr>
<tr>
<td>Symptomatic preceding infection</td>
<td>10 (67%)</td>
<td>8 (62%)</td>
</tr>
<tr>
<td>Median MRC sum score at nadir (range)</td>
<td>32 (12-54)</td>
<td>38 (2-54)</td>
</tr>
<tr>
<td>Median GBS disability grade at nadir (range)</td>
<td>4 (3-5)</td>
<td>4 (2-5)</td>
</tr>
<tr>
<td>Median arm grade at nadir (range)</td>
<td>2 (1-3)</td>
<td>2 (1-4)</td>
</tr>
<tr>
<td>Mean time to nadir in days (range)</td>
<td>10 (2-19)</td>
<td>9 (2-27)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Laboratory features</th>
<th>HIV-positive (n=15)</th>
<th>HIV-negative (N=13)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demyelinating neuropathy (AIDP)</td>
<td>11 (73%)</td>
<td>10 (77%)</td>
</tr>
<tr>
<td>Serum hyponatraemia (&lt;135mmol/l)</td>
<td>9 (60%)</td>
<td>2 (18%)</td>
</tr>
<tr>
<td>Mean CSF protein [g/l] (range)</td>
<td>1.49(0.42-2.69)</td>
<td>1.26 (0.39-5.84)</td>
</tr>
<tr>
<td>CSF lymphocytosis (&gt;5 cells / mm3)</td>
<td>5 (38%) [n=13]</td>
<td>2 (18%) [n=11]</td>
</tr>
</tbody>
</table>

Table 1. Clinical and laboratory features of HIV-positive (n=15) and –negative (n=13) patients. Where assessments / tests were not performed in all patients, the number of patients (n) that did undergo the test / assessment is stated in brackets *= p<0.05 (significant).
Characteristics of HIV-positive patients at the time of presentation

Of the 15 HIV-positive patients, 13 were diagnosed as being infected with HIV at the time of presentation with GBS. Neither of the two patients known with HIV-infection was taking anti-retroviral treatment (ART). The median CD4+ count in the HIV-positive group was 270 cells/mm$^3$ (range 32–656). Four patients had CD4+ counts less than 200 cells/mm$^3$ (of which 2 were less than 50 cells/mm$^3$), seven patients had counts between 200 and 350 cells/mm$^3$, and four patients had CD4+ counts greater than 350 cells/mm$^3$. According to the CDC classification system, 5 patients had AIDS (4 had concomitant infection (tuberculosis), and one had a CD4+ count <200 cells/mm$^3$). Thirteen patients underwent lumbar puncture, of whom 5 (38%) had a CSF lymphocytic pleocytosis (range 18–40 cells/mm$^3$).

Incidence of GBS in HIV-positive patients

Since the study was not community-based, the absolute incidence of GBS in the HIV-positive population could not be determined. However, the incidence of GBS in the HIV-positive population relative to the HIV-negative population can be calculated based on the estimated prevalence of HIV infection in the general population from which the study sample originates. Based on UNAIDS SPECTRUM estimates, the prevalence for the Western Cape Province was 6.2%, 6.2%, 6.16% and 4.75% (mean=5.8%) for the years 2008–2011 (available on www.doh.gov.za). If the incidence of GBS was similar in the HIV-positive and –negative populations, approximately 5.8% of GBS cases would be expected to be HIV-positive. However, the actual number of GBS cases with HIV infection in our study was 53.6% (15/28). This represents a 9.2-fold enrichment of HIV-seropositive patients in our cohort of GBS. Even when screened patients who were excluded from the study are included in the calculation, there remains an 8.8-fold enrichment. If the average HIV-seroprevalence rate of 5.8% is used, the incidence of GBS in our population is calculated to be 18.74, 95% CI [7.69, 40.60] times higher in HIV-positive than in HIV–negative individuals.

Treatment and course

Eight (53%) of HIV-positive and 7 (54%) of HIV-negative patients required treatment with PE or IVig, and two patients in each group required intubation and mechanical ventilation. Two patients in the HIV-positive group experienced secondary worsening after initial improvement, and one of this required re-treatment. No patients in the HIV-negative group experienced secondary worsening. In the HIV-positive group, 4 patients (27%) were diagnosed with pulmonary tuberculosis during admission, and 1 patient developed pulmonary tuberculosis 4 months after admission with GBS. One patient in the HIV-negative group was diagnosed with bronchus carcinoma. Two patients in each group died: both in the HIV-positive group from hospital-acquired pneumonia (CDC stages A2 and C2 respectively), and 1 each from bronchus carcinoma and cardiac arrest in the HIV-negative group. The mean duration of hospital stay (days) for survivors was 36 days in the HIV-positive group and 26 days in the HIV-negative group (p=0.34).

After discharge, survivors were followed for up to 12 months (mean=4 months). Over this period, 9 of the 13 HIV-positive and 3 of the 11 HIV-negative patients were lost to follow-up. All patients who were followed either improved or recovered completely. Due to the large amount of patients lost to follow-up, reliable conclusions about the long-term outcomes were not possible. However, no patients were re-referred with recurrent weakness, which would suggest chronic inflammatory demyelinating polyneuropathy (CIDP) rather than GBS.

Discussion

The association between GBS and HIV infection has been reported frequently over the last 25 years, mainly in the form of case reports and series. Despite this, the magnitude of the association between the two disorders has not been quantified satisfactorily. A retrospective study from Zimbabwe reported that 16 of 29 patients (55%) with GBS had concomitant HIV infection, while the seroprevalence of HIV infection in Zimbabwe was estimated at 4.3% at the time [18]. In our prospective cohort of 28 patients with GBS, we found a more than nine fold higher HIV-seropositivity rate than expected and calculated the incidence to be at least 18 times higher in HIV-positive individuals.

The relation between GBS and the stage of HIV infection is also unclear. Although most authors describe GBS as a manifestation of early HIV infection or even seroconversion, numerous reports of GBS in advanced immunosuppression are available, with CD4+ counts as low as 4 cells/mm$^3$ [9–11]. In our cohort, one third of HIV-positive patients had AIDS. Interestingly, in 13 (87%) of the 15 patients, GBS was the presenting feature of HIV infection. Clinical, laboratory and electrophysiological features were similar to the HIV-negative group. Although a statistically significant difference in the proportion of patients with serum hyponatraemia (presumably related to inappropriate ADH secretion) was found, this is likely to be related to a lower than expected proportion in the HIV-negative group, as compared to previously published data [19]. It is noteworthy that, despite the underlying immunosuppression, the short term outcome of the disorder appears to be similar in the HIV-positive and –negative groups, although the sample size does not allow firm conclusions to be drawn.

Our study has some limitations. Firstly, the sample size does not allow a reliable comparison of the two groups and small differences may have been missed. However, due to the scarcity of GBS, a sufficiently large sample size may take many years to obtain. Secondly, because the study was not community based, no conclusions about the actual incidence of GBS in HIV-infected patients can be drawn. However, we believe that our findings do suggest a meaningful increase in the incidence of GBS in HIV-infected compared to non-infected individuals. Thirdly, the poor retention of participants in the study precludes meaningful analysis and comparison of the long-term outcome. Lastly, since no HIV-positive patients were taking ART, the generalizability of our results to populations with a higher rate of ART use is uncertain, as ART could theoretically protect against the development of immune-mediated conditions by decreasing immune dysregulation [20].

The development of autoimmune conditions in immunodeficient patients, although seemingly counterintuitive, is now a well-recognized phenomenon [21,22]. Immune dysregulation in HIV may involve T or B cells, or both, and may lead to autoimmune phenomena unique to HIV or to classical autoimmune syndromes [23]. Immune dysregulation is probably related to chronic immune activation accompanying HIV infection, likely induced and maintained by translocation of microbial products across the intestinal mucosa into the systemic circulation, probably mediated by depletion of Th17 cells in the gut mucosa [24]. Another mechanism likely to be
involved in chronic immune activation is homeostatic peripheral expansion (HPE). This process occurs during lymphopenia and describes an augmented T-cell proliferation to both foreign antigens (cognate and gut-associated commensal organisms) and self-antigens. However, because auto-immunity does not develop in all lymphopenic individuals, it is reasonable to assume that lymphopenia alone is not sufficient to induce auto-immunity. Some authors have proposed a “two-hit model” to account for the loss in self-tolerance underpinning auto-immunity [25]. According to this theory, the chronic immune activation observed in lymphopenia provides a fertile environment for a “second hit”, such as cytokine overproduction or localized inflammation, to induce auto-immune disease. In GBS, such a second hit could conceivably be a flu-like syndrome or diarrhoeal illness, exposing the activated immune system to a foreign antigen and thereby inducing auto-immune disease by means of molecular mimicry, widely accepted as a pathomechanism in GBS [26]. In this respect, GBS differs from most other autoimmune disorders described in HIV, as the commonly described disorders are not typically regarded as related to molecular mimicry triggered by a foreign antigen [21,22]. An interesting question is whether the triggering epitope is a component of an opportunistic infection or the HIV virus itself, as described in HIV-related anaemia for example [27].

In addition to above-mentioned systemic immune dysregulation, HIV infected individuals may also be more vulnerable to developing auto-immune diseases involving the central (CNS) and peripheral nervous system (PNS). Normally, neurons in the CNS and PNS are shielded from systemic inflammatory reactions and immune responses by the blood–brain barrier (BBB) and blood–nerve barrier (BNB), respectively. Malfunction of these barriers causes leakage of immunoglobulins and cytokines, which are not allowed through under normal circumstances, into the endoneurial space. In the case of GBS, this disruption of the BNB is considered paramount in the development of the disorder, and has been shown to be induced by sera from patients with GBS [28]. BBB function has also been shown to be altered by HIV-infection, both in vitro and in vivo [29]. Whether the same is true for the BNB remains to be investigated, but, if indeed the case, predisposes the PNS to immune-mediated damage, as occurs in GBS.

In summary, we have shown that the incidence of GBS is substantially increased in HIV-positive compared to -negative individuals. GBS can occur in any stage of HIV infection, and is often the presenting manifestation. The short-term outcome of HIV-associated GBS appears similar to that of GBS in HIV-seronegative patients, but the longer-term outcome remains to be investigated. The increased incidence of GBS in HIV-infected individuals is probably related to immune dysregulation, the molecular mechanisms of which warrants further investigation.

References

