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Cases of neuromyelitis optica spectrum disorder from the East Africa region, highlighting challenges in diagnostics and healthcare access

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Keywords:
- Neuromyelitis optica spectrum disorder
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- Neuromyelitis optica spectrum disorder (NMOSD) is an auto-immune disease of the central nervous system (CNS) associated with the IgG-antibody against aquaporin-4 (AQP4-IgG). There is little published epidemiology of NMOSD from sub-Saharan Africa (SSA).

Methods: We retrospectively collated NMOSD cases admitted to our tertiary regional neurology centre.

Results: We identified 11 cases (10 female, average age 30 years). 64% (7/11) were seropositive for AQP4-IgG, measured using indirect immunofluorescence. The remaining cases could either not afford tests, or had pathognomonic radiological features. 57% (4/7) of seropositive cases had concurrent/recent CNS infection. All patients were treated with high-dose intravenous methylprednisolone (IVMP), and 36% (4/11) also had plasma exchange. Only 55% (6/11) of the patients were seen by a neurologist at presentation: they had less relapses (1.3 vs 2.4), less diagnostic delay (2.3 vs 7.4 months), and were less disabled at the end of our review period. 10 cases were immunosuppressed long-term: 60% on mycophenolate, 30% azathioprine, and one on rituximab.

Conclusion: Our study is the largest case series of NMOSD from the East Africa region. Patients faced challenges of access to appropriate and affordable testing, and timely availability of a neurologist at onset, which had impacts on their functional outcomes. The majority of the seropositive cases had recent/concurrent CNS infections, suggesting triggered auto-immunity.

1. Introduction

Neuromyelitis optica spectrum disorder (NMOSD) is an auto-immune relapsing central nervous system disorder (CNS), diagnosed based on the presence of core clinical characteristic presentations, the presence of aquaporin-4 immunoglobulin G antibodies (AQP4-IgG), and/or pathognomonic findings on CNS magnetic resonance imaging (MRI) as per the International Panel for NMOSD Diagnosis (IPND) criteria [1]. It is strongly recommended that AQP4-IgG is tested using a cell-based assay, but if negative or not available then the clinical characteristics and MRI findings can fulfil the NMOSD diagnostic criteria. About one-third of AQP4-IgG seronegative NMOSD patients have been found to be positive for antibodies against myelin oligodendrocyte glycoprotein (MOG-Ab) [2]. Anti-MOG disease is another CNS auto-immune demyelinating syndrome which mimics the clinical presentation of NMOSD but has been recently identified as a separate entity [3]. It has therefore been proposed that it is reasonable to test for both AQP4-IgG and MOG-Abs in patients with an NMOSD presentation [4].

Patients presenting acutely with NMOSD should be promptly recognised, investigated and then commenced on relapse treatment followed by long-term maintenance immunosuppression to prevent devastating relapses [5]. However, there is variable availability of testing for AQP4-IgG and MOG-Ab globally with the least accessibility noted in low/low-middle income countries in Africa [6], and coupled with the lack of MRI scanners in Africa [7] this makes the diagnosis of NMOSD and anti-MOG disease particularly more challenging in these regions.
settings.

The epidemiology of NMOSD in developing countries is being increasingly understood e.g. in Latin America, NMOSD prevalence varies from 0.37–4.2/100,000 inhabitants, with AQP4-IgG seropositivity being 33–73.5% [8]. However, there remains a significant gap of data from Africa e.g. in a 2013 systematic review of 216 population-based studies, five reported the incidence and prevalence of NMOSD but none were from Africa [9], and a 2018 review also revealed no prevalence studies from Africa [10]. Latitude does not seem to be related to NMOSD prevalence [11], yet to date there are less than 500 cases of NMOSD published from Africa [12], of which the majority are from North [13–15] and West [16–19] Africa, with a smaller cohort from South Africa [20] and only one case report from East Africa [21].

Based on studies published from large multi-ethnic cohorts mostly in developed countries, it has been consistently shown that NMOSD, particularly atypical optic neuritis, is much more common [22], more severe, and more likely to be associated with AQP4-IgG seropositivity in patients of African and Afro-Caribbean background [8,23,24]. In a retrospective analysis from eleven Brazilian hospitals, 81/153 cases of NMOSD had at least one black grandparent, and they were more likely to have worse disability at diagnosis when compared to Caucasian patients [25]. We have previously reported the largest cohort of multiple sclerosis (MS) from East Africa [26] from our ongoing registry project. Given all the literature reviewed above, we report here NMOSD cases from our hospital cohort as the local population seem to be more at risk of developing severe disease.

2. Materials and methods

2.1. Ethical considerations

Our protocol of our continuing MS cohort study at our regional tertiary referral hospital has been approved by our Institutional Ethics Review Committee (approval number REC-99/2018). The protocol includes looking for MS mimics and chameleons [27] particularly NMOSD.

2.2. Informed consent and patient details

We have obtained written informed consent from all the patients for whom we have supplied MRI images as well as the supplementary video, and these consent forms have been filed in their medical records at our hospital. The patient identifiers from the images and video have been removed or obscured to maintain patient confidentiality.

2.3. Identification of NMOSD cases

We performed case-finding for NMOSD cases from our MS registry through a combination of:

1. International Classification of Diseases version 10 (ICD-10) coding for patients discharged from our hospital (our institution does not code for outpatient visits) with the following codes: NMOSD, acute transverse myelitis in demyelinating disease of CNS, optic neuritis, acute brainstem syndrome, area postrema syndrome, diencephalic syndrome and nacropaley;
2. Positive AQP4-IgG antibodies in serum and/or cerebrospinal fluid (CSF) obtained from our laboratory database as per the ‘Investigations’ section of our MS registry data collection protocol;
3. Recollection from memory by the neurologists at our institution (authors DSS, JH and PM).

We reviewed all cases and collected information on the clinical presentation(s), including timelines, and investigation results, particularly MRI findings and AQP4-1gG results in order to match the IPND criteria as well as to ensure there were no ‘red flags’ to suggest a diagnosis other than NMOSD. In addition, as per the MS research protocol, we gathered information according to the ‘Treatment/Management’ and ‘Outcome’ sections of the MS registry protocol, which included management of acute relapses and choice of immunosuppression, and outcome measured using the Extended Disability Status Scale (EDSS).

2.4. Data analysis and management

Case summaries were collected in a Microsoft Word document. We tabulated the findings in a Microsoft Excel sheet. Neither of these documents contained any patient identifiers, and both were password-protected and were stored in a shared drive within our institution’s intranet, which is only accessible by the authors via passwords. The information in these sheets could only be linked to the patient via a handwritten sheet of random numbers that were locked in the office of the leading author (DSS).

3. Results

We collected 11 cases, all of whom were of black African background and indigenous to East Africa. The findings above are summarised in Table 1. 91% (10/11) were female, with an average age of 30 years at presentation (range 15–51 years). Longitudinally extensive transverse myelitis (LETM) and Optic neuritis (ON) – all bilateral simultaneous or sequential – were the commonest first presentations in 45% (5/11) and 36% (4/11) of cases, and similarly for second presentations [27% (3/11) and 18% (2/11) respectively]. One patient (case 1) had both LETM and ON at the first presentation i.e. classic Devic syndrome. 27% (3/11) presented with acute brainstem syndrome (ABS), all as a first symptom (including case 5; see Fig. 1 for respective MRI brain scan correlates), and only one patient (case 6) had area postrema syndrome (APS) as a second symptom after LETM (see Fig. 2 for MRI correlates). Two patients (cases 5 and 7) had pathognomonic dystonic spasms (see Supplementary Video 1 of case 5, Fig. 3).

In terms of AQP4-IgG testing, 64% (7/11) had a positive result from serum, of which three cases also had a positive test in CSF (CSF AQP4-IgG was not done in the remaining 4 patients who were seropositive). Case 11 could not afford testing for AQP4-IgG, and the remaining patients (cases 8 through 10) were seronegative, but we included these cases in our series for the following reasons:

- Our institution does not offer cell-based assay testing for AQP4-IgG;
- Case 9 and 11 both presented with bilateral sequential ON then LETM;
- Case 8 presented with LETM and case 10 with bilateral sequential ON extending over half the optic nerve length on MRI (see Fig. 4 for MRI of optic nerves in case 10) with an altitudinal visual field defect, and both required PLEX.

There were other positive results from investigations that we noted, most notably that the majority (4/7) of AQP4-IgG positive cases had laboratory, including CSF, evidence of concurrent or recent CNS infection: tuberculosis (TB, case 2), herpes simplex virus type 2 (HSV-2, case 6), syphilis (case 7; see Fig. 5 for MRI of LETM), and sterile meningitis with pleocytotic CSF [case 1, who also had oligoclonal bands (OCBs) in CSF]. Case 8 had lymphocytic CSF but no positive microbiology or PCR/serology.

10 of the 11 patients were treated with high-dose intravenous methylprednisolone (IVMP), administered as 1 gram per day for three to five days, for each acute relapse, and some were given a tapering dose of oral prednisolone of variable doses and duration after IVMP. All the cases with concurrent/recent infections were treated with the appropriate anti-microbials before embarking on IVMP treatment, as directed by our infectious disease specialists: case 6 was given fourteen days of intravenous acyclovir together with PLEX; case 2 was treated with anti-
Table 1

Summary of results of NMOSD cases.

| Case no.: | AQP4-IgG antibody positive | AQP4-IgG antibody -ve/unavailable |
|-----------|-----------------------------|----------------------------------|
|           | 1  | 2  | 3  | 4  | 5  | 6  | 7  | 8† | 9  | 10 | 11 | 2 |
| **Age at onset (in years)** | 15 | 17 | 22 | 36 | 36 | 45 | 51 | 18 | 26 | 30 | 34 | 1 |
| **Core clinical characteristic presentation** | 1  | 2  | 2  | 1  | 1  | 1  | 2  | 1  | 2  | 1  | 2  | 1 |
| **AQP4-IgG antibody:** | Serum | nd | nd | nd | nd | nd | nd | nd | nd | nd | nd | nd |
| **CSF (NR): white cell count** | (<5/mm<sup>3</sup>)<sup>b</sup> | 32 | <5 | <5 | na | 145 | <5 | 61 | <5 | <5 | nd | 1 |
| **glucose (2.22–3.89 mmol/L)** | 3.32 | nd | 3.39 | 3.11 | na | 3.42 | 5.36 | 3.6 | 2.31 | 3.55 | nd | 1 |
| **protein (0.15–0.45 g/L)** | 1.59 | nd | 0.23 | 0.37 | na | 0.48 | 0.36 | 0.57 | 0.44 | 0.17 | nd | 1 |
| **positive microbiology** | n | nd | n | na | na | HSV2 | TPHA | n | n | n | nd | 1 |
| **OCBs** | -ve | nd | -ve | -ve | nd | -ve | -ve | -ve | -ve | -ve | -ve | 1 |
| **PLEX after IVMP** | n | n | n | n | y | y | y | y | n | n | y | n |
| **Neurologist involved at onset** | y | y | y | y | y | y | y | y | y | y | y | y |
| **Delay to diagnosis (in months)** | 14 | 7 | 1 | 1 | 1 | 5 | 3 | 7 | 0 | 12 | 1 |
| **Relapses before diagnosis** | 4 | 2 | 1 | 1 | 1 | 2 | 2 | 3 | 1 | 2 | 1 |
| **Maintenance immunosuppression** | None | MMF | MMF | MMF | AZP | MMF | MMF | RTX | MMF | AZP | AZP | 1 |
| **Relapses after immunosuppression** | None | na<sup>c</sup> | None | None | None | None | None | None | None | None | None | 1 |
| **EDSS at last review** | na<sup>d</sup> | 2 | 7 | 1 | 1 | 8 | 7 | 2 | 8 | 2 | 7 | 1 |
| **Diagnosis to last review** | 46 | 11 | 26 | 18 | 39 | 13 | 9 | 32 | 18 | 21 | 48 | 1 |
| **Additional notes** | OCBs + ve after 4 years | First treated as CNS TB | First treated as RRMS | Post-partum & lactating | Tonic spasms | Relapse as non-compliant | Anti-MOG negative | Could not afford tests | 1 |

**Abbreviations:** +ve = positive; -ve = negative; ABS = acute brainstem syndrome; APS = area postrema syndrome; AZP = azathioprine; CNS = central nervous system; CSF = cerebrospinal fluid; EDSS = extended disability status scale; HSV-2 = Herpes simplex virus type 2; IVMP = intravenous methylprednisolone; LETM = longitudinally extensive transverse myelitis; MMF = mycophenolate mofetil; MOG = myelin oligodendrocyte protein; na = not available; nd = not done; NR = normal range; OCBs = oligoclonal bands; ON = bilateral simultaneous or sequential optic neuritis; PCR = polymerase chain reaction; RRMS = relapsing-remitting multiple sclerosis; RTX = rituximab; TB = tuberculosis; TPHA = treponema pallidum haemagglutination assay

<sup>a</sup> Number denotes presentation: “1” = first presentation, “2” = second presentation.

<sup>b</sup> Red cell counts all <5/mm<sup>3</sup>.

<sup>c</sup> Percentage lymphocytes.

<sup>d</sup> Not done as evidence of pulmonary tuberculosis so treated empirically for CNS disease.

<sup>e</sup> Lost to follow-up as emigrated.

<sup>f</sup> Not available as done in another country and records unavailable.

<sup>g</sup> Male patient (remainder of cases all female).

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**Fig. 1.** Coronal T2 (left image) and sagittal (right image) fluid-attenuated inversion recovery (FLAIR) magnetic resonance imaging (MRI) sequences of case 5 showing a hyper-intense expansile lesion of the medulla and cervical-medullary junction, who presented with acute brainstem syndrome and quadriplegia.
tubercular therapy and dexamethasone for two weeks then with IVMP; case 7 was treated with penicillin G for 2 weeks before proceeding to IVMP. 36% (4/11) had plasma exchange (PLEX) after IVMP if there was no clinical response and/or MRI findings were suggestive of severe disease.

Just over half (6/11) of the patients were seen by a neurologist during the first presentation, and when compared to the remaining five cases, they had: (i) less relapse presentations before the diagnosis was made (average 1.3 vs 2.4 relapses); (ii) less delay to NMOSD diagnosis (average 2.3 months vs 7.4 months); and (iii) less disability when assessed at the end of our review period (average EDSS score of 3.6 vs 5.4).

All our NMOSD patients were commenced on long-term immunosuppression, although we did not have details for case 1 as she transferred her care to outside our facility during her acute illness and was lost to follow-up. For the remaining cases, the most frequently prescribed regimen was mycophenolate in 60% (6/10) of cases, with azathioprine (3/10) and rituximab (case 8 alone) being used in the remainder of the cohort. Average follow-up time was 25 months (range 9 to 48), with no difference if first seen by a neurologist.

Although case 3 was diagnosed with NMOSD at 7 months by one of the authors, appropriate immunosuppressive therapy was not commenced until 3 years later as the patient was lost to follow-up and sought treatment at various locations outside our institution and was managed as relapsing-remitting MS; during this time she had 3 further relapses with a relatively poor EDSS when she returned to our services. Case 4 was 3 months post-partum when she developed LETM, and was breastfeeding so she made an informed decision with her neurologist not to commence on immunosuppression after the positive serum AQP4-IgG result came out given the perceived risks. Unfortunately she relapsed with bilateral simultaneous ON, although with good recovery, within the same month at which point she decided to commence immunosuppression.

4. Discussion
4.1. Demographics

The NMOSD cohort at our institution was predominantly female with an average of 30 years, which is in keeping with what has been found in
other studies from Africa [13–17]. Similarly, ON and LETM were the most frequently reported symptoms at relapse presentation, with LETM being the most frequent overall (73%, and the Algerian cohort was 75% [17]), but the concomitant optico-spinal involvement i.e. classic Devic syndrome seemed to more prevalent in Morocco [13] and Senegal [16].

4.2. AQP4-IgG testing and status

AQP4-IgG seropositivity in our cohort was 64% (7/11), which is comparable to the range of 38% [17] to 62.5% [16] from African cohorts, although altogether much less when compared to 73–90% in international cohorts [28]. Two of the four seronegative patients (cases 9 and 11) fulfilled the clinical and radiological criteria for NMOSD diagnosis. The two remaining cases each had only one core clinical characteristic, but we included them in our cohort because: (i) both had NMOSD-concordant MRI findings; (ii) red flags for other differentials had been ruled out in the diagnostic workup; (iii) our laboratory only offers indirect immunofluorescence for detection of AQP4-IgG, which is known to be much less sensitive [29] than the cell-based assays recommended by the IPND (1); and (iv) both patients had monophasic illnesses, which are more likely to be seronegative when compared to those with established relapsing NMOSD [30]. Such risk stratification in the absence of available and appropriate serology testing has been used in other African cohorts to justify immunosuppression e.g. 15.9% of the Algeria cases who did not fulfil the NMOSD criteria were put immunosuppression as they were deemed “high risk”.

MOG-Ab was not available at the time case 8 was last followed up, but was available for case 10 and it was negative, which again lends some support to NMOSD diagnosis. This lack of availability of diagnostic tests for NMOSD was shown in a recent survey, where Africa had the least accessibility to either AQP4-IgG or MOG-Ab testing [6]. The survey did not differentiate between the type of assay; whilst our institution does not have access to cell-based assays for AQP4-IgG, there is evidence that in some parts of Africa e.g. Egypt such assays are readily available [15]. This variable access to high-quality testing raises the question as to whether there needs to be an international effort to expand NMOSD diagnostics and make them more affordable, or whether to adjust the diagnostic criteria to cater for this chronic laboratory insufficiency in poorer countries. As a result of this study, we are in the process of identifying a laboratory that does provide cell-based assays for identifying AQP4-IgG antibodies, and will then recall the patients for retesting.

4.3. Impact of early neurologist review

One of our most important findings is the significant impact early involvement of a neurologist made to the timeline to diagnosis (average of 2.3 months from first presentation if seen by a neurologist first), number of relapses before diagnosis (1.3 vs 2.4 average relapses), and outcome of the patients as measured by EDSS (3.6 vs 5.4). Despite being the economic hub of the East Africa region, Kenya has only 18 neurologists for a population of 47 million, a neurology service gap that has been shown to be consistent across Africa [31]. The majority are based in the capital centre in institutions such as our regional referral hospital, which partly explains why 5/11 of our NMOSD cases did not see a neurologist at first presentation and led to the delayed diagnosis (7.4 months delay on average): three of these patients first presented with LETM but could not access a neurologist due to the towns they were living in (cases 6 and 7). Case 8 was seen by a neurosurgeon first because the MRI scan suggested a spinal cord tumour. One of the five patients (case 9) first presented with bilateral sequential ON but was managed on both occasions by an ophthalmologist who did not refer the patient forward, and then she had a devastating LETM relapse which led to her presentation to our hospital, from which she remained wheelchair-bound. Case 11 was admitted to the national public referral hospital Fig. 4. Sagittal T2 with fat suppression (left image) and T1 with contrast (right image) MRI sequences of case 11 who presented with quadriplegia secondary to longitudinally extensive transverse myelitis.

Fig. 5. Still of supplementary video 1 of case 5 demonstrating dystonic spasms pathognomonic of NMOSD.
but was admitted under a physician first before being sent to the
neurologist. Her diagnosis was made on clinical grounds after the second
relapse with ON as she could not afford any investigations. Delays in
diagnosis, which lead to delay in treatment, has been shown to be a
significant factor contributing to long-term disability in NMOSD with a
history of LETM [32], which explains why these patients in our cohort
had a poorer outcome. Aside from increasing the neurology healthcare
workforce in developing countries, simple measures such as stand-
ardised protocols for managing atypical ON and/or LETM have been
shown to improve diagnoses rates, timely treatment, and overall out-
comes in resource-poor settings for NMOSD patients [33], therefore
national or regional guidelines for demyelinating diseases can mitigate
some of the healthcare shortfalls highlighted by our cohort.

Case 3 was first seen by a neurologist but had the worst EDSS score of
7 when compared to the other five cases (average EDSS score 1.5) who
were also seen by a neurologist at first presentation. This was because of
a combination of factors. Firstly, they were lost to follow-up after the
second relapse which is when the AQP4-IgG was requested; the turn-
around time for results from our referral laboratory is three weeks on
average. The results came back positive but were not flagged up by our
laboratory as AQP4-IgG does not feature as a critical result that requires
verbal reporting to the attending clinician. We have logged this as a
clinical incident and as a result our laboratory now has a protocol to
have all positive AQP4-IgG results immediately telephoned to the
neurologist in charge. Secondly, the patient self-referred to another
neurologist outside our institution for a second opinion, and who was
therefore not privy to her laboratory findings from the first two pre-
sentations. She was thus treated as a case of relapsing remitting MS – a
misdiagnosis that can happen in up to 40% of patients [24,30] – which
may have contributed to her ongoing deterioration since it is now well
established that disease-modifying drugs used in MS exacerbate NMOSD
[34].

4.4. Concurrent CNS infections

It has been postulated that infections may be a trigger to the auto-
immunity in NMOSD [35], and interestingly over half of AQP4-IgG
seropositive patients in our cohort had a recent on concurrent infec-
tion, each of which have been reported to be associated with NMOSD
onset: CNS tuberculosis in case 2 [36], HSV-2 in case 6 [37], and neu-
rosyphilis in case 7 [35–38]. Case 1 had a sterile meningitis with no
isolate; repeat CSF was only done 4 years later which was normal. Case 2
did not have CSF examination at presentation due to presence of pul-
monary tuberculosis, and they were then lost to follow-up in the same
year. Case 1 and 6 had lymphocytic CSF, and they declined follow-up
CSF studies after their acute infections were treated. Case 8 had a lym-
phocytic CSF but negative microbiology and serology studies that
excluded concurrent infection; treatment for NMOSD led to good
outcome long-term and the attending clinician therefore decided against
repeat CSF.

All acute neurology admissions at our institution are tested for
infection by human immunodeficiency virus (HIV) as per a previously
published protocol [39], but none of the NMOSD patients had concur-
rent HIV infection. There is some evidence from the South African cohort
that post-infectious auto-immunity plays a role in NMOSD pathogenesis
in HIV-infected patients, who tend to have more tumefactive CNS lesions
[40,41], which we tend to avoid given the implications to fertility in a
female-predominant disease. Rituximab is generally expensive and
not easily available, and was only used in the management of case 8 as
they did not like to daily tablets; it is used more frequently for NMOSD in
other parts of Africa especially Egypt [15]. Only one patient (case 7) had
a relapse due to non-compliance with the oral steroids and
mycophenolate.

5. Conclusions

We have described the largest indigenous case series of NMOSD from
the East Africa region. The demography in terms of gender and age were
similar to other NMOSD cohorts reported internationally, but there were
also some unique differences within our cases, mainly that majority of
the sero-positive patients had other CNS infections, suggesting para-
fungal pathogenesis. Documenting the patient journey from symp-
tom onset to outcomes elucidated specific challenges: variable access to
appropriate and affordable diagnostic testing, especially for cell-based
assays to detect AQP4-IgG, and suboptimal availability of, and referral
to, a neurologist at onset of the disease which is a very modifiable risk
factor. Delayed and incorrect diagnoses negatively impacted on patient
outcomes in terms of disability, which has implications on how
neurology-specific healthcare infrastructure should be aligned and
developed in order to improve the management of conditions such as
NMOSD in relatively resource-poor settings.

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Data statement

The clinical information and imaging data used to support the
findings of this study are included within the article. According to our
institutional information governance regulations, the anonymised data
can be requested from the corresponding author.

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Declarations of interest

None.

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