MRI findings in people with epilepsy and nodding syndrome in an area endemic for onchocerciasis: an observational study

Winkler AS1,2, Friedrich K3, Velicheti S4, Dharsee J4, König R1, Nassri A1, Meindl M3, Kidunda A5, Müller TH4, Jilek-Aall L7, Matuja W8, Gotwald T9, Schmutzhard E1

1. Department of Neurology, Technical University of Munich, Germany
2. Haydom Lutheran Hospital, Manyara Region, Tanzania
3. Department of Neurology, Medical University of Innsbruck, Innsbruck, Austria
4. Department of Radiology, The Aga Khan Hospital, Dar es Salaam, Tanzania
5. Mahenge District Hospital, Morogoro Region, Tanzania
6. Institute for Medical Informatics, Biometry and Epidemiology, Ludwig-Maximilians-University, Munich, Germany
7. Department of Psychiatry, University of British Columbia, Vancouver, Canada
8. Department of Neurology, Muhimbili University of Health and Allied Sciences, Dar es Salaam, Tanzania
9. Department of Radiology, Medical University of Innsbruck, Innsbruck, Austria

Abstract
Background: Onchocerciasis has been implicated in the pathogenesis of epilepsy. The debate on a potential causal relationship between Onchocerca volvulus and epilepsy has taken a new direction in the light of the most recent epidemic of nodding syndrome.
Objective: To document MRI changes in people with different types of epilepsy and investigate whether there is an association with O. volvulus infection.
Methods: In a prospective study in southern Tanzania, an area endemic for O. volvulus with a high prevalence of epilepsy and nodding syndrome, we performed MRI on 32 people with epilepsy, 12 of which suffered from nodding syndrome. Polymerase chain reaction (PCR) of O. volvulus was performed in skin and CSF.
Results: The most frequent abnormalities seen on MRI was atrophy (twelve patients (37.5%)) followed by intraparenchymal pathologies such as changes in the hippocampus (nine patients (28.1%)), gliotic lesions (six patients (18.8%)) and subcortical signal abnormalities (three patients (9.4%)). There was an overall trend towards an association of intraparenchymal cerebral pathologies and infection with O. volvulus based on skin PCR (Fisher's Exact Test p=0.067) which was most pronounced in children and adolescents with nodding syndrome compared to those with other types of epilepsy (Fisher's Exact Test, p=0.083). Contrary to skin PCR results, PCR of CSF was negative in all patients.
Conclusion: The observed trend towards an association of intraparenchymal cerebral pathological results on MRI and a positive skin PCR for O. volvulus despite negative PCR of CSF is intriguing and deserves further attention.

Key words: Epilepsy, head nodding, magnetic resonance imaging, cerebrospinal fluid.

Introduction
The discussion whether onchocerciasis causes epilepsy still remains to be resolved and lately has taken a new direction in search for causes of a devastating African epidemic epilepsy syndrome termed nodding syndrome (NS).1-4 Human onchocerciasis is caused by the filarial nematode Onchocerca volvulus (O. volvulus), which mainly manifests on skin and eyes.5,6 The blackfly (Simulium species) transmits the microfilariae (mf) during a blood meal. In the host they grow into adult worms, which again mate and produce mf. These mf mainly migrate in subcutaneous tissue and may occasionally be found in sputum, urine, blood, and cerebrospinal fluid (CSF).7,8 In African areas affected by onchocerciasis, some studies suggest a connection with epilepsy,9,14 while others do not.15-17 Overall a link between the two diseases seems possible,18,19 but is far from being established. Latest results on NS from South Sudan indicate that in some villages significantly more children with NS are infested with O. volvulus compared to healthy controls, whereas in other villages no difference could be found between cases and controls.3

*Corresponding author: Dr. Andrea Sylvia Winkler
Department of Neurology
Klinikum rechts der Isar
Ismaninger Strasse 22, 81675 Munich
Germany
Tel.: +49/89/45815015
Email: drawinkler@yahoo.com.au

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One hypothesis is that mf may cause inflammation or direct destruction of brain tissue similar to the mechanisms known from the analysis of affected skin and eyes. Therefore imaging studies may shed more light on the debate whether there is a causal relationship between epilepsy and *O. volvulus*. The area (Mahenge) of southern Tanzania where the study took place has a prevalence of epilepsy of up to 37.1/1,000, the prevalence rates of epilepsy in sub-Saharan Africa ranging from 7.8-14.8/1,000 and in addition, is highly endemic for onchocerciasis. Thus, it was deemed appropriate to recruit people with epilepsy (PWE) from this region for parasitological and neuroradiological work-up. To elucidate such a potential association between *O. volvulus* and epilepsy, we performed, for the first time, both cerebral MRI in PWE and children/adolescents with NS from an area endemic for *O. volvulus* together with *O. volvulus* polymerase chain reaction (PCR) of skin and CSF.

**Methods**

**Study sites and performance of MRI**

The main study was conducted in The Mahenge Epilepsy Clinic, which was founded in 1960 by L. Jilek-Aall, nowadays attended by well over 900 people with epilepsy. It is situated in the Government Hospital of Mahenge which cares for an approximate 32,000 inhabitants of the Vigoi division, Ulanga district, southern Tanzania. Although Mahenge, our study area, has been part of the African Programme for Onchocerciasis Control (APOC) since 1997, in 2011 an epidemiological survey revealed that the prevalence of onchocerciasis was still 46% (personal communication Elibariki Mwakapeje, Ministry of Health Tanzania).

The MRI scans were performed in the Department of Radiology at The Aga Khan Hospital, Dar es Salaam, 500 km away from Mahenge, using a 1.5 Tesla MR machine (GE). In all patients, axial T1W1, T2W1, FLAIR, coronal FLAIR, and sagittal T1W1 were performed. In selected patients (especially those patients with complex partial seizures or head nodding seizures), coronal T2W1 and coronal 3D SPGR for hippocampus anatomy was added. All scans were evaluated by three experienced radiologists (JD, SV and TG), who were blinded to the patient groups.

**Participants and data collection**

A convenience sample of patients known to suffer from epilepsy or NS was recruited from The Mahenge Epilepsy Clinic in order to evaluate whether in these people brain lesions on MRI were associated with a positive evidence for onchocerciasis. All PWE or NS had active epilepsy with at least one seizure in the year preceding the survey. Patients with single epileptic seizures or epileptic seizures with an obvious predisposition or cause (e.g. perinatal brain damage, seizures in the context of malaria or alcohol withdrawal seizures among others) were excluded. Epilepsy was defined as two or more afebrile seizures unrelated to acute metabolic disorders or withdrawal of drugs or alcohol. For the definition of NS refer to Winkler et al. The actual head nodding seizure represents a repetitive short loss of neck muscle tone resulting in a forward bobbing of the head unexplained by any other neurological or psychiatric condition. In addition to the actual nodding seizures there may be other seizure types and/or neurological/medical signs/symptoms present which has been suggested as early as 2008 to be termed nodding syndrome (=NS).

A previously validated questionnaire was used in a face-to-face interview performed by neurologists (WM, ES, ASW), final-year medical students on elective attachment from the University of Innsbruck (KF, MM), a translator, at least one relative and the patient. Physical examination with neurological evaluation was also performed (WM, ES, ASW). An electroencephalogram (EEG) was not at hand. In terms of recruitment of participants we aimed for an even distribution of people with seemingly epilepsy without focal signs (group 1) and epilepsy with focal signs (group 2), and within these two groups we aimed for similar ages, a balanced distribution of gender and people with *O. volvulus* positive and negative skin snips. Patients of group 1 suffered from generalized tonic-clonic seizures and were neurologically and mentally normal. People with epileptic seizures (either focal or generalized based on ictal observation) with clear focal neurological signs and/or clear mental handicap/cognitive impairment having been present before the onset of seizures were put into group 2. Alternatively, if on neurological and standard cognitive examination no abnormality was detected, patients with clear focal types of epilepsy but otherwise healthy were also put into this group. Patients with NS were allocated to a separate group (group 3) and divided into “head nodding only” and “head nodding plus” in order to reflect the full spectrum of disease. The groups are summarized as follows:

**Patient groups**

Group 1 (n=10): Epilepsy without obvious focal neurological signs/mental retardation

Group 2 (n=10): Epilepsy with obvious focal neurological signs/mental retardation, or clear focal epilepsy (based on ictal observations) without any of the mentioned signs

Group 3 (n=12): Head nodding seizures, divided in:

a) Head nodding only” with head nodding seizures only
b) «Head nodding plus” with head nodding seizures and other seizure types mainly generalized tonic-clonic seizures.

**Laboratory procedures**
Skin snips were taken from the right and left iliac crest. If there were skin changes suggestive of *O. volvulus*, a skin snip was taken from that site too. Microscopic evaluation of mf was performed at 40x magnification one and four hours later. If no mf was seen, the well was checked again after 12 hours. The mf density per person was calculated. The *O. volvulus* PCR was performed after a modified version of that used by Zhang et al.\(^26\) Methods were optimized, quality controls and specificity tests with different parasites were performed to exclude cross reactivity. Gel electrophoresis was applied to visualize the amplification products. The PCR for *O. volvulus* in CSF was performed according to the method used for skin snips. Infestation with *O. volvulus* was established in the presence of a positive skin PCR. In four cases PCR was not performed. In these cases the results on skin snip were observed.

**Ethical clearance**
The study was cleared by the Ethics Committee of the Muhimbili University College of Health Sciences, University of Dar es Salaam. Oral informed consent was given by all patients allowing us to transcribe clinical data into the questionnaire. Written and witnessed informed consent was obtained from the patients or, in case of children or non-competent patients, their respective parents or next of kin for all procedures.

**Statistical analysis**
The data was analysed using SPSS 18.0 (SPSS Inc., Chicago, USA). The chi-square test and Fischer's Exact test were used to test the association between two categorical variables. The Mann-Whitney *U* test was performed for continuous variables with a non-parametric distribution; the significance level was set at alpha=0.05. As this is a hypothesis generating investigation, no adjustment for multiple testing was applied. Furthermore, in view of the small group sizes, no multivariate analyses were carried out.

**Results**
**Demographic, clinical and laboratory details**
The median age of all 32 patients was 16 years (interquartile range (IQR): 14-21 years) with a male to female ratio of 1.3:1. Along with demographic, clinical and imaging details, the average densities of the *O. volvulus* mf in the skin and the results of the respective PCR are presented in tables 2 - 4. In three cases, no further counting of mf was available, thus only information as to whether the skin snip was positive or negative is given. The skin snips of 28/32 patients were retested for *O. volvulus* antigen by PCR. Five patients had a positive skin PCR despite a negative skin snip and on the other hand, three patients had a negative skin PCR despite a positive skin snip. The CSF PCR, the results of which are not presented in the tables, was performed in all patients and was negative in all of them.

**Results of the MRI investigations**
Normal cerebral MRI scans were found in 10/32 patients (31.3%). Table 1 gives an overview of the diagnosed cerebral lesions.

**Table 1: Pathologies on MRI scans divided into unilateral and bilateral lesions**

<table>
<thead>
<tr>
<th>Pathologies</th>
<th>Unilateral lesions</th>
<th>Bilateral lesions</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>HC changes*</td>
<td>3</td>
<td>6</td>
<td>9</td>
</tr>
<tr>
<td>Gliotic lesions**</td>
<td>4</td>
<td>2</td>
<td>6</td>
</tr>
<tr>
<td>Signal abnormalities***</td>
<td>0</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Cyst</td>
<td>2</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Other lesions****</td>
<td>1</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Total</td>
<td>10</td>
<td>12</td>
<td>22</td>
</tr>
<tr>
<td>Atrophy</td>
<td>N.A.</td>
<td>N.A.</td>
<td>12</td>
</tr>
<tr>
<td>No biopsies</td>
<td>-</td>
<td>-</td>
<td>10</td>
</tr>
</tbody>
</table>

MRI could show more than one pathology. HC – Hippocampus, N.A. – not applicable
*Gliotic lesions were defined as focal small (< 1cm) punctuate hyperintensities in T2WI, mainly located in the frontal lobe; **HC changes were divided into HC sclerosis and suspected HC sclerosis, the latter only showing either HC signal change or volume loss; ***Signal abnormalities were defined as subcortical hyperintensities that did not fulfil the criteria for gliotic lesions
****Other lesions: focal cortical dysplasia, bilateral cystic lesions in the occipital lobe
Of the 32 patients, nine patients (28.1%) showed changes in the hippocampus, six of them bilaterally. Sclerosis was confirmed in only three patients, the others had either signal changes or loss of volume in the hippocampus. Clinically 3/9 patients had complex partial or generalized tonic clonic seizures with focal signs, 4/9 had nodding seizures either alone or associated with other seizure types and one patient had generalized tonic clonic seizures without focal signs. Only three patients with hippocampus changes reported febrile seizures in childhood. Six patients (18.8%) had gliotic lesions, mainly subcortically, and three patients (9.4%) showed bilateral peritrigonal/subcortical signal abnormalities. In two patients (6.3%) an arachnoid cyst was found, one in the frontal lobe measuring 20x14 mm, the other in the temporal lobe being much smaller in size. A focal cortical dysplasia in the frontal lobe and a bilateral cystic lesion in the occipital lobes, the latter most likely caused by perinatal hypoxic brain damage, were seen in one patient each. Atrophy of the brain was found in 12/32 patients (37.5%) and was the only pathological finding in four of them. Cerebral and cerebellar atrophy were equally distributed (tables 1-4).

In group 1, four patients showed intracerebral pathologies on MRI. Unilateral gliotic lesions, focal cortical dysplasia and bilateral signal abnormalities were seen as well as unilateral hippocampus changes (table 2).

### Table 2: Demographics, neurological examination, MRI, mf density and skin PCR of group 1

<table>
<thead>
<tr>
<th>Age (yrs)</th>
<th>Gender</th>
<th>Seizure type</th>
<th>Neurological examination/mental examination</th>
<th>Intraparenchymal MRI pathologies</th>
<th>Atrophy</th>
<th>Mf skin /mg</th>
<th>PCR skin</th>
</tr>
</thead>
<tbody>
<tr>
<td>12</td>
<td>M</td>
<td>Gen</td>
<td>NAD</td>
<td>Susp. HC sclerosis lft</td>
<td>No</td>
<td>6</td>
<td>Pos</td>
</tr>
<tr>
<td>17</td>
<td>M</td>
<td>Gen</td>
<td>NAD</td>
<td>Cortical dysplasia lft frontal</td>
<td>No</td>
<td>51.5</td>
<td>Pos</td>
</tr>
<tr>
<td>20</td>
<td>M</td>
<td>Gen</td>
<td>NAD</td>
<td>Signal abnorm. bilateral peritrigonal rt frontal</td>
<td>No</td>
<td>11.5</td>
<td>Pos</td>
</tr>
<tr>
<td>21</td>
<td>F</td>
<td>Gen</td>
<td>NAD</td>
<td>Gliotic lesions subcortical rt frontal</td>
<td>No</td>
<td>1.33</td>
<td>Pos</td>
</tr>
<tr>
<td>35</td>
<td>F</td>
<td>Gen</td>
<td>NAD</td>
<td>No</td>
<td>No</td>
<td>Pos*</td>
<td>Pos</td>
</tr>
<tr>
<td>16</td>
<td>F</td>
<td>Gen</td>
<td>NAD</td>
<td>No</td>
<td>Mild general</td>
<td>0</td>
<td>Neg</td>
</tr>
<tr>
<td>18</td>
<td>M</td>
<td>Gen</td>
<td>NAD</td>
<td>No</td>
<td>No</td>
<td>0</td>
<td>Pos</td>
</tr>
<tr>
<td>20</td>
<td>F</td>
<td>Gen</td>
<td>NAD</td>
<td>No</td>
<td>No</td>
<td>0</td>
<td>Neg</td>
</tr>
<tr>
<td>21</td>
<td>M</td>
<td>Gen</td>
<td>NAD</td>
<td>No</td>
<td>No</td>
<td>0</td>
<td>Neg</td>
</tr>
<tr>
<td>32</td>
<td>M</td>
<td>Gen</td>
<td>NAD</td>
<td>General</td>
<td>No</td>
<td>0</td>
<td>Neg</td>
</tr>
</tbody>
</table>

Gen – generalized tonic clonic seizures, NAD – nothing abnormal detected, rt – right, lft- left, susp. – suspected, abnorm. – abnormalities, Mf – microfilariae, pos* - no further counting available.

In group 2, two patients had normal MRI, one without pathological findings on neurological and mental examination and the other with slight mental retardation. All other patients had pathological MRI results which together with the clinical examination, mf density and skin PCR are shown in table 3. The most frequent intraparenchymal cerebral pathologies were signal changes in the hippocampus followed by subcortical signal abnormalities (table 3).

In group 3, consisting of patients with NS, four patients (33.3%) with head nodding alone showed normal MRI scans. The remaining eight patients, the majority of which suffered from NS together with generalized tonic clonic seizures, showed gliotic lesions (5/8; figures 1a-6a) and hippocampus pathologies (4/8), among other lesions (table 4).

### Relationship between results on MRI and types of epilepsy (subgroup analysis)

The presence of intraparenchymal pathologies on MRI (exclusive of cerebral atrophy) was not dependent on the patients’ age (Mann-Whitney U test, p=0.985) or gender (χ²-test, p=0.530). Regarding types of epilepsy, relevant intraparenchymal MRI pathologies such as hippocampus changes, gliotic lesions and signal abnormalities, were significantly more frequent in the group “head nodding plus with other seizure types” compared to the other types of epilepsy (Fisher’s Exact Test, p=0.01; table 4). When looking at specific brain lesions, the significance could be maintained for gliotic lesions only (Fisher’s Exact Test, p=0.034; figures 1a-6a).
Table 3: Demographics, neurological examination, MRI, mf density and skin PCR of group 2

<table>
<thead>
<tr>
<th>Age (yrs)</th>
<th>Gender</th>
<th>Seizure type</th>
<th>Neurological examination/mental examination</th>
<th>Intraparenchymal MRI pathologies</th>
<th>Atrophy</th>
<th>MF skin /mg</th>
<th>PCR skin</th>
</tr>
</thead>
<tbody>
<tr>
<td>14</td>
<td>F</td>
<td>1.Gen+, 2.Cp</td>
<td>NAD</td>
<td>No</td>
<td>No</td>
<td>0.5</td>
<td>Not tested</td>
</tr>
<tr>
<td>16</td>
<td>M</td>
<td>Gen</td>
<td>MeRe, no focal neurological signs</td>
<td>No</td>
<td>Cerebellar vermis</td>
<td>4</td>
<td>Neg</td>
</tr>
<tr>
<td>18</td>
<td>F</td>
<td>Gen</td>
<td>MeRe, no focal neurological signs</td>
<td>Arachnoid cyst rt frontal</td>
<td>No</td>
<td>35.5</td>
<td>Pos</td>
</tr>
<tr>
<td>22</td>
<td>M</td>
<td>Gen+</td>
<td>MeRe, hemihypaesthesia rt, UMN signs rt</td>
<td>Susp. HC sclerosis bilateral</td>
<td>Mild general Cerebellar</td>
<td>0.17</td>
<td>Pos</td>
</tr>
<tr>
<td>47</td>
<td>M</td>
<td>1.Gen, no focal neurological signs</td>
<td>NAD</td>
<td>HC sclerosis lt, susp. HC sclerosis rt; signal abnorm. subcortical bilateral frontal</td>
<td>No</td>
<td>5.5</td>
<td>Pos</td>
</tr>
<tr>
<td>13</td>
<td>F</td>
<td>Cp</td>
<td>MeRe, no focal neurological signs</td>
<td>No</td>
<td>No</td>
<td>0</td>
<td>Pos</td>
</tr>
<tr>
<td>13</td>
<td>M</td>
<td>1.Gen, 2.Cp</td>
<td>MeRe, no focal neurological signs</td>
<td>Signal abnorm. subcortical bilateral occipital</td>
<td>No</td>
<td>0</td>
<td>Neg</td>
</tr>
<tr>
<td>15</td>
<td>M</td>
<td>Gen</td>
<td>MeRe, cerebellar syndrome</td>
<td>Cystic defects occipital bilateral</td>
<td>Cerebellar</td>
<td>0</td>
<td>Neg</td>
</tr>
<tr>
<td>16</td>
<td>M</td>
<td>Gen</td>
<td>MeRe, increased reflexes</td>
<td>No</td>
<td>Mild cerebellar</td>
<td>0</td>
<td>Neg</td>
</tr>
<tr>
<td>27</td>
<td>F</td>
<td>Gen+</td>
<td>NAD</td>
<td>Susp. HC sclerosis bilateral</td>
<td>Cerebral occipital + parietal</td>
<td>0</td>
<td>Neg</td>
</tr>
</tbody>
</table>

Cp - complex partial seizures, Gen - generalized tonic clonic seizures, Gen+ - generalized tonic clonic seizures with additional focal signs ictally, NAD – nothing abnormal detected, MeRe - mental retardation, UMN - upper motor neuron, HC – Hippocampus, lt – left, rt – right, susp. – suspected, abnorm. – abnormalities, MF – microfilariae, HC changes were divided into HC sclerosis and suspected HC sclerosis, the latter only showing either HC signal changes or volume loss.

Table 4: Demographics, neurological examination, MRI, mf density and skin PCR of group 3

<table>
<thead>
<tr>
<th>Age (yrs)</th>
<th>Gender</th>
<th>Seizure type</th>
<th>Neurological examination/mental examination</th>
<th>Intraparenchymal MRI pathologies</th>
<th>Atrophy</th>
<th>MF skin /mg</th>
<th>PCR skin</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>M</td>
<td>HN only</td>
<td>NAD</td>
<td>No</td>
<td>Gliotic lesions rt frontal subcortical; susp. HC sclerosis rt&gt;lt</td>
<td>No</td>
<td>1.5</td>
</tr>
<tr>
<td>16</td>
<td>F</td>
<td>HN only</td>
<td>NAD</td>
<td>Gliotic lesions rt frontal subcortical; susp. HC sclerosis rt&gt;lt</td>
<td>No</td>
<td>5.65</td>
<td>Pos</td>
</tr>
<tr>
<td>18</td>
<td>M</td>
<td>HN only</td>
<td>Frontal release signs, UMN signs lt&gt;rt</td>
<td>No</td>
<td>No</td>
<td>1</td>
<td>Neg</td>
</tr>
<tr>
<td>10</td>
<td>F</td>
<td>HN only</td>
<td>NAD</td>
<td>No</td>
<td>No</td>
<td>0</td>
<td>Not tested</td>
</tr>
<tr>
<td>14</td>
<td>F</td>
<td>HN only</td>
<td>NAD</td>
<td>No</td>
<td>No</td>
<td>Neg*</td>
<td>Not tested</td>
</tr>
<tr>
<td>15</td>
<td>M</td>
<td>HN only</td>
<td>NAD</td>
<td>Anachn cyst rt temporal; HC sclerosis rt</td>
<td>No</td>
<td>0</td>
<td>Pos</td>
</tr>
<tr>
<td>9</td>
<td>F</td>
<td>HN only</td>
<td>UMN signs rt</td>
<td>Gliotic lesions rt frontal subcortical; susp. HC sclerosis rt&gt;lt</td>
<td>General</td>
<td>Pos*</td>
<td>Not tested</td>
</tr>
<tr>
<td>15</td>
<td>M</td>
<td>1.HN, 2.Gen</td>
<td>NAD</td>
<td>Gliotic lesions bilateral</td>
<td>No</td>
<td>1.5</td>
<td>Pos</td>
</tr>
<tr>
<td>16</td>
<td>F</td>
<td>1.HN, 2.Gen</td>
<td>NAD</td>
<td>Gliotic lesions lt temporo-occipital-parietal gwm-j</td>
<td>No</td>
<td>6.17</td>
<td>Pos</td>
</tr>
<tr>
<td>13</td>
<td>M</td>
<td>1.HN, 2.Gen</td>
<td>NAD</td>
<td>Gliotic lesions bilateral</td>
<td>No</td>
<td>0</td>
<td>Neg</td>
</tr>
<tr>
<td>16</td>
<td>F</td>
<td>1.HN, 2.Gen</td>
<td>NAD</td>
<td>Susp. HC sclerosis lt</td>
<td>0</td>
<td>Pos</td>
<td></td>
</tr>
<tr>
<td>21</td>
<td>M</td>
<td>1.HN, 2.Gen, 3.Cp</td>
<td>Brain damage, no focal neurology</td>
<td>Susp. HC sclerosis bilateral</td>
<td>Mild cerebellar</td>
<td>0</td>
<td>Pos</td>
</tr>
</tbody>
</table>
Footnotes for table 4
HN – head nodding seizures, Gen - generalized tonic clonic seizures, Cp - complex partial seizures, NAD – nothing abnormal detected, HC – Hippocampus, susp. – suspected, gwm-j - grey-white matter junction, rt – right, lft – left, neg* - no further counting available
HC changes were divided into HC sclerosis and suspected HC sclerosis, the latter only showing either HC signal changes or volume loss.

Figures 1a – 3a: 13 year-old male with head nodding and generalized tonic clonic seizures showing bilateral punctuate subcortical hyperintensities in the frontal lobe on T2W1. Some of them are suppressed on FLAIR confirming that they are Virchow-Robin spaces, the remaining hyperintense foci most likely correspond to gliotic lesions. After administration of contrast medium, lesions were non-enhancing on T1W1.
Figures 4a – 6a: 14 year-old male with head nodding and generalized tonic clonic seizures showing bilateral punctuate subcortical hyperintensities in the frontal lobe, on T2WI and axial as well as coronal FLAIR most likely corresponding to gliotic lesions.
Relationship between results on MRI and infestation with *O. volvulus*

Patients with an abnormal MRI had significantly more positive skin PCR results (Fisher’s Exact Test \(p=0.019\)) than patients with a normal MRI (exclusive of atrophy). When looking at plausible lesions that could be caused by *O. volvulus* including gliotic lesions, hippocampus changes or signal abnormalities there was a near significant trend for these lesions to be associated with a positive PCR results (Fisher’s Exact Test \(p=0.067\)). Interestingly, when looking at epilepsy types, it was only in people with head nodding that the trend towards an association of intraparenchymal pathologies on MRI with a positive skin PCR for *O. volvulus* was seen (Fisher’s Exact Test, \(p=0.083\); table 4).

When all three types of lesions were further analysed separately the association could not be maintained.

Discussion

Our findings in general

The International League Against Epilepsy has recommended neuroimaging with MRI for all PWE wherever feasible.\(^27\) Availability and affordability are rendering this recommendation almost impossible in resource-poor countries. Excluding South Africa, only nine MRI scanners were available in sub-Saharan countries in 2003.\(^28\) CT scans are less expensive and more widely-spread; in 2003 65 CT were registered throughout sub-Saharan Africa.\(^28\) Thus, imaging studies in PWE in sub-Saharan Africa are scarce and were mainly performed by means of CT.\(^29\)\(^,\)\(^30\) In our study, we performed 32 MRI scans in PWE with different types of epilepsies including children/adolescents with NS with and without additional types of seizures. Parts of the findings in children/adolescents with NS were already summarized in the description of the study population in 2008, figure 1 and are now presented more in depth and in the wider context of other types of epileptic seizures.

Only 10 (31%) PWE of our study population showed no abnormality on MRI. This is highly different from studies from Brazil in which 76% of over 130 PWE had normal MRI scans figure 3 and from Nigeria in which almost 50% of PWE had a normal CT scans.\(^30\) The reason for the high number of abnormalities in our study might be explained by selection bias. The patients of our study with normal MRI were mainly suffering from epilepsy without any obvious neurological signs (group 1), emphasizing the fact that thorough history taking associated with clinical examination is a good indicator for the type of epilepsy in resource-poor settings.

In the individuals with pathological MRI scans, atrophy, hippocampus changes and gliotic lesions were the commonest findings. In some patients, the atrophy was disproportionate for age. Atrophy of the cerebellum may be explained by high doses of phenytoin over many years and cerebral atrophy by excessive alcohol use. Changes in the hippocampus were seen in over 20% of PWE, which may cause complex partial seizures, including head nodding in children.\(^32\) Interestingly, all but one PWE with hippocampus changes in our study had seizures in the context of either focal epilepsy or NS. In our study, no significant relationship between febrile convulsions and hippocampus changes was found.

Febrile seizures have however been identified as risk factors of childhood epilepsy in other African studies.\(^35\)\^-\(^37\) Gliotic lesions were mainly found in children and adolescents with head nodding, especially in those with additional seizure types. Cranial infections, cerebral ischemia and traumatic brain injury have all been implicated as possible causes for gliotic lesions in a study from China where 60% of people with gliotic lesions were suffering from epilepsy.\(^36\) There was no history of trauma in the people of our study. Ischemia especially in younger people seems unlikely, so that an inflammatory cause for the gliotic lesions seems possible.

MRI findings and *O. volvulus*

Electroencephalographic studies in PWE from an area endemic for *O. volvulus* with high prevalence of epilepsy showed mainly focal epileptiform activity.\(^37\) Focal lesions/pathologies were also prevalent in our study population mainly consisting of gliotic lesions and hippocampus changes. They were not evenly distributed over the three study groups, but seemed to be clustering in PWE with clinical indicators pointing to focal epilepsy and in children/adolescents with NS, especially those suffering from NS and additional types of seizures. Hippocampus changes seem rather difficult to attribute to an infectious agent but have been implicated in the context of neurocysticercosis caused by the parasite *Taenia solium*,\(^38\) whereas gliotic changes may be attributed to inflammation.\(^36\) Mf migrating into the brain may cause focal inflammatory reaction and subsequent gliosis as has been shown with *Loa loa*, a filarial nematode mainly distributed in West and Central Africa. *L. loa* adult worms have been found to be able to enter the brain causing an acute and severe
encephalopathy mainly in *O. volvulus* co-infected individuals treated with filaricidal drugs. Occasional cases were reported, but are rare in comparison to the number of individuals affected by *L. loa*. Pathological changes in *L. loa* encephalopathy include inflammatory infiltrates and gliotic lesions in brain parenchyma of various degrees of severity. In addition to direct damage to brain parenchyma affection of cerebral arteries (cellular infiltrates and thickening of the vessel wall) has also been seen observed which may lead to potentially ensuing vasculitis that has also been demonstrated with other parasites such as *Toxocara canis*.

Duke et al. reported *O. volvulus* mf in CSF before and after treatment with diethylcarbamazine. If this is a genuine finding and was not due to contamination during lumbar puncture, the CSF should have shown some signs of contact with the parasites. Unfortunately, this was not described in Duke’s study. In our study, *O. volvulus* PCR of CSF was negative in all individuals. It may be argued that if the inflammation caused by mf had resolved, *O. volvulus* PCR, which traces DNA and thus an active process, may become negative. Interestingly, in patients of a much bigger patient cohort of almost 200 PWE of which PWE of the current study were part of, *O. volvulus* CSF antibody index, which should give evidence of exposure to the parasite, was negative too, indicating that contact of the CSF with the parasite, past or present, seems very unlikely. Gliotic lesions or other types of signal abnormalities may also be seen in an autoimmune response. Gallin et al. have demonstrated an association between a hyperreactive onchodermatitis and the presence of autoantibodies to proteins found in neutrophils in affected individuals. Also, an autoimmune process has been hypothesized in the development of ocular onchocerciasis and a shared antigen (OV39) between ocular tissue, including the retina, and *O. volvulus* has been implicated in the disease process. Based on these principles, cross-reacting antibodies to nervous tissue may be postulated. One would however expect a completely different, at least more widespread “gliotic” reaction if an autoimmune pathogenesis caused by *O. volvulus* is assumed.

An indirect relationship between *O. volvulus* and epilepsy might further be supported by the fact that we found an almost significant association between intraparenchymal pathologies on MRI and *O. volvulus* PCR result on skin snip. Also, when looking at the various types of epilepsy a nearly significant relationship between brain pathologies, especially gliosis and hippocampus changes, and positive skin PCR could be maintained for children/adolescents with NS only. Whether these lesions are causative for the nodding seizures and/or the additional mainly generalized seizures or whether they are unrelated remains uncertain in the absence of a control group. Intercital EEG results in some of these children showed generalized rather than focal abnormalities, which however does not exclude gliosis or hippocampus changes as origin. As to the actual head nodding attack, which clinically looks like atonic seizures (and has recently been shown to be reflected by an EEG decrement) or, if nodding is less prominent, complex partial or absence-type seizures, gliosis as the cause of these attacks seem less likely and in fact only 2/7 children with head nodding attacks alone had gliotic lesions on MRI (table 4). Hippocampus changes were seen in five of the 12 children/adolescents with NS and may cause complex partial seizures in this cohort. However, hippocampus changes were not more frequent than in PWE with focal seizures and therefore may be an accidental finding without implication for seizure generation in children/adolescents with NS or, alternatively, some of the children/adolescents with NS may indeed suffer from complex partial epilepsy which clinically appears as head nodding.

**Limitations of the study**

This is more of a descriptive study of MRI pathologies in PWE and children/adolescents with NS from an *O. volvulus* endemic area. In this context statistical analysis of MRI pathologies and their relationship with onchocerciasis must be interpreted with care due to the small sample size. Local circumstances with the only MRI in the country being many hundreds of kilometers away and financial restrictions did not allow for more people to be recruited into the study. Also, the MRI (1.5 Tesla) used in our study shows a fairly low resolution compared to a 3 Tesla MRI and lesions may have been missed. Neuroimaging in a larger population, especially of children/adolescents with NS, should be envisaged in order to support or reject the trends observed in our study. Also, our study is lacking a control group of healthy age-matched individuals. Healthy young people in the circumstances of a resource-poor setting can usually not be away for several days as they are the backbone of their families and indeed all the young people approached by us did not consent to participate into our study.
Conclusion
MRI in PWE and children/adolescents with HS from an area endemic for *O. volvulus* shows multiple pathologies, the most frequent being atrophy, hippocampus changes and gliosis. There was a trend towards an association of intraparenchymal brain pathologies (hippocampus changes, gliotic lesions as well as other signal abnormalities) and *O. volvulus* skin PCR in PWE and more explicitly in children/adolescents with NS. However, as this was a mere observational study with low numbers and without control group, the possible association between certain brain pathologies and infection with *O. volvulus* should be explored further, preferably in larger studies with matched control groups.

Abbreviations
CSF: cerebrospinal fluid; *L. loa*, *Loa loa*, NS: nodding syndrome; MRI: magnetic resonance imaging; *O. volvulus*: *Onchocerca volvulus*; PCR: polymerase chain reaction; PWE: people with epilepsy

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