Meta-analysis of epilepsy prevalence in West Africa and its relationship with onchocerciasis endemicity and control

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Background: A high prevalence and incidence of epilepsy has been reported in onchocerciasis-endemic regions in Central and East Africa. There is compelling epidemiological evidence suggesting that this high burden is caused by onchocerciasis-associated epilepsy (OAE). We hypothesized that OAE had also occurred in West African onchocerciasis foci.

Methods: We searched PubMed, the African Journals Online platform and grey literature for population-based epilepsy studies in West African countries. Epilepsy and onchocerciasis prevalence data were extracted. The pre-control onchocerciasis endemicity in the study sites was estimated from historical data of onchocerciasis control programmes. The prevalence of epilepsy in different sites was analysed, taking into account onchocerciasis endemicity and the duration of control.

Results: The pooled prevalence of epilepsy in the West African study sites was 13.14 per 1000 (95% confidence interval 11.28–15.00). Higher pre-control endemicity and a shorter duration of onchocerciasis control were both associated with increased epilepsy prevalence (p < 0.001). Two studies in Ivory Coast that provided detailed descriptions of persons with epilepsy in onchocerciasis-endemic settings revealed that most of them had features of OAE (73.7% and 83.3%, respectively).

Conclusions: Our findings suggest that before and during the early years of implementing onchocerciasis control in West Africa, high onchocerciasis endemicity resulted in a high prevalence of OAE and that subsequent control efforts significantly reduced the prevalence of OAE.

Keywords: epilepsy, prevalence, onchocerciasis, ivermectin, vector control, West Africa

Introduction

Onchocerciasis (river blindness) is known to cause skin and eye disease. Currently there is growing epidemiological evidence that onchocerciasis is also able to directly or indirectly cause epilepsy. A recent cohort study in the Mbam valley in Cameroon showed that skin Onchocerca volvulus microfilarial density in children <10 y of age determined the risk, in a dose-related fashion, of developing epilepsy later in life. Therefore the terms ‘onchocerciasis-associated epilepsy’ (OAE) and ‘river epilepsy’ have been proposed to describe this epidemiological link.

A high prevalence of epilepsy in an onchocerciasis-endemic region was initially reported in 1938 in Mexico, and later in many meso- and hyperendemic onchocerciasis foci in Central and East Africa. However, in West Africa, reports of high epilepsy prevalence in relation to a high onchocerciasis endemicity are scarce. We hypothesized that this may be due, at least in part, to the large-scale vector control activities of the Onchocerciasis Control Programme (OCP) in West Africa, which was implemented as early as the 1970s. If onchocerciasis is indeed able to cause epilepsy, a high prevalence of epilepsy should also have occurred in West African foci before or during the early years of onchocerciasis control efforts. We therefore critically reviewed population-based epilepsy surveys from West Africa to investigate a possible relationship with onchocerciasis after taking the possible effect of onchocerciasis control into account.
Methods

Search strategy
We searched PubMed and the African Journals Online (AJOL) platforms for articles published in French and English until 30 June 2019. Key search terms included ‘epilepsy’, ‘prevalence’ and ‘West African countries’ (see Supplementary material, Appendix 1). We also retrieved grey literature (doctoral theses and conference abstracts) from previously published reviews on the epidemiology of epilepsy in sub-Saharan Africa.10–11 We included all population-based surveys conducted in West Africa that reported epilepsy prevalence in both the children and adult population. Two authors (SFJN, CR) independently performed the screening of articles for inclusion in this review. The risk of bias in selected studies was assessed using the ‘study participation’ and ‘outcome measurement’ domains of the Quality in Prognosis Studies (QUIPS) tool.12 The epilepsy surveys conducted in onchocerciasis foci, which provided detailed descriptions of all identified persons with epilepsy (PWE), were used to estimate the proportion of participants who fulfilled previously published criteria for OAE.1 The pre-control onchocerciasis endemicity data for the included villages were estimated from existing 1 km2 resolution maps of pre-control onchocerciasis endemicity levels in West Africa. For the OCP countries, we used a map of the predicted prevalence of microfilariae generated in a geostatistical analysis of the pre-control skin snip survey data of the OCP.13 For Liberia and Nigeria, which were not covered by the OCP but were participating countries of the African Programme for Onchocerciasis Control (APOC), we used a map of the predicted pre-control prevalence of nodules in APOC countries generated in a geostatistical analysis of Rapid Epidemiological Mapping of Onchocerciasis (REMO) data.14 We then converted the predicted nodule prevalence for the location of the study sites to the prevalence of microfilariae using the approach described by Coffeng et al.15

Data analysis
The epilepsy prevalence data that were extracted from eligible studies were entered into Review Manager version 5.3 (Nordic Cochrane Centre, Copenhagen, Denmark). The standard error (SE) of the epilepsy prevalence from each survey was calculated using the following formula: $SE = \sqrt{p(1-p)/n}$, where $p$ is the prevalence and $n$ is the total study population. Pooled prevalence estimates were obtained using the inverse variance method and a random effects model.

The pre-control onchocerciasis endemicity in our study sites was expressed in terms of the prevalence of microfilariae in the total village population. We also categorized the study sites based on the duration of onchocerciasis control by the time of the epilepsy survey. Subgroup analyses were performed and forest plots generated. Multiple linear regressions, weighted by the sample size for each village (weighting coefficient $= \sqrt{n}$, where $n$ is the sample size), were fitted to investigate the relationship between logit-transformed epilepsy prevalence, pre-control prevalence of microfilariae, duration of onchocerciasis control, type of onchocerciasis control and study year. Regression analyses were done using the software JASP version 0.10.2 (University of Amsterdam, Amsterdam, The Netherlands).

Results

PubMed and AJOL searches yielded 154 studies, while the manual search yielded 22 records. Of these, 53 were retained as epilepsy prevalence surveys after screening the abstracts. After further assessment, only 20 studies from the systematic search16–35 and 9 studies from the manual search36–44 met the inclusion criteria, and their data were extracted (Figure 1).

Included studies originated from 10 different countries of West Africa, one of which (Gambia) was non-endemic for onchocerciasis (Figure 2). Onchocerciasis prevalence at the time of the epilepsy survey was available for four endemic study sites. Table 1 summarizes the findings from each individual study; the methodology and suggested epilepsy aetiologies are provided in the supplementary material (Appendix 2). The findings from the retrieved studies will be reported country-wise and further elaborated on the Discussion.

Epilepsy prevalence in Benin

Five studies were included from Benin,23,27,35,38,43 one of which specifically investigated onchocerciasis and epilepsy in the Agbobome area; findings revealed that in 9/13 (69.2%) PWE, *O. volvulus* microfilariae were detected in skin snips while 8 PWE (61.5%) presented with onchocerciasis nodules.43 In that same study, one PWE also suffered from onchocercal blindness.43 Furthermore, most patients (76.9%) experienced generalized tonic-clonic seizures.43 The study by Avode et al.15 incriminated neurocysticercosis as a major cause of epilepsy in Savalou (central region of Benin), where cysticercosis prevalence was 39.5% and epilepsy prevalence 15.2%. A family history of epilepsy was a predominant risk factor in Zinvie23 and Djidja,27 but not in Agbobome.43

Epilepsy prevalence in Burkina Faso

In 2007 a high prevalence of epilepsy (44.27%) was observed in three rural villages in Burkina Faso. A case–control study in the area showed an association between epilepsy and cysticercosis seropositivity (prevalence odds ratio [OR] 3.1 [95% confidence interval [CI] 1.0–8.3].28 All definitive and probable cases of neurocysticercosis were located in the two villages where pig breeding was common.26 Debouverie et al.39 reported an epilepsy prevalence of 10.65% in villages that were endemic for neither cysticercosis nor onchocerciasis; the predominant aetiology reported was neonatal asphyxia (present in 12/38 PWE with symptomatic epilepsy), and more than half of all PWE experienced their first seizure before the age of 10 y.

Epilepsy prevalence in Gambia

A low epilepsy prevalence of 4.26% was reported in Gambia,26 which is non-endemic for onchocerciasis. The peak age-specific prevalence rates occurred in the 35–44 y age group.

Epilepsy prevalence in Ghana

During a wide community-based survey (n=129 812) in Kintampo, exposure to *O. volvulus* was identified as an independent risk factor for epilepsy (OR 2.32 [95% CI 1.12–4.78] in children
Figure 1. Study selection and inclusion (Preferred Reporting Items for Systematic Reviews and Meta-Analyses flowchart).

Figure 2. Included study sites and OCP boundaries in West Africa.
Table 1. Epilepsy population-based surveys performed in West Africa

| Study site                                      | Survey year | First author | Study population | Epilepsy prevalence (%) | Pre-control onchocerciasis endemcity (%) | Onchocerciasis prevalence at time of survey (%) | Duration (y) of onchocerciasis control (VC/CDTI/overall)a |
|------------------------------------------------|-------------|--------------|------------------|--------------------------|------------------------------------------|-------------------------------------------------|----------------------------------------------------------|
| Benin (Savalou)35                               | 1994        | Avode        | 1443             | 15.25                    | 53                                       | NA                                              | 6/3/6                                                    |
| Benin (Agbogbome)43                              | 1995        | Gbenou       | 503              | 25.84                    | 67                                       | 47.4                                            | 7/4/7                                                    |
| Benin (Zinnie)23                                 | 1997        | Debrock      | 3134             | 21.06                    | NE                                       | NE                                              | NA                                                       |
| Benin (Djijja)27                                 | 2005        | Houinato     | 11 668           | 12.68                    | 60                                       | NA                                              | 14/12/17                                                 |
| Benin (Dangbo)38                                 | 2007        | Houinato     | 737              | 31.21                    | NE                                       | NE                                              | NA                                                       |
| Burkina Faso (Passore, Yatenga)39               | 2007        | Nitiéma      | 881              | 44.27                    | NE                                       | NE                                              | NA                                                       |
| Burkina Faso (Batondo, Nyonyogo, Pabré)28        | 1997        | Coleman      | 16200            | 4.26                     | NE                                       | NE                                              | NA                                                       |
| Guinea (Kintampo)29                              | 2010        | Ae-Ngibise   | 129,812          | 1.92                     | 45                                       | NA                                              | 26/13/34                                                 |
| Ivory Coast (Bonon-Frefredou)40                  | 1987        | Kouassi      | 1176             | 7.65                     | NE                                       | NE                                              | NA                                                       |
| Ivory Coast (Akooungou)36                        | 1989        | Kouadio      | 309              | 58.25                    | 51                                       | NA                                              | 0                                                       |
| Ivory Coast (M’Brou)37                           | 1993        | Kaudjhis     | 920              | 41.30                    | 76                                       | NA                                              | 0                                                       |
| Liberia (Grand Bassa)42                          | 1981        | Goudsmit     | 4436             | 27.73                    | 56                                       | NA                                              | 0                                                       |
| Liberia (Grand Bassa)41                          | 1983        | Gerrits       | 1673             | 49.01                    | 38                                       | NA                                              | 0                                                       |
| Mali (Tyenfala, Baquindea)22                     | 1999        | Farnarier    | 5243             | 13.35                    | 71                                       | 23                                              | 12/8/12                                                  |
| Nigeria (Igbo-Ora)19                             | 1982        | Osuntokun    | 18 954           | 5.33                     | 29.1b                                    | 23                                              | 12/8/12                                                  |
| Nigeria (Aiyete)20                               | 1982        | Osuntokun    | 903              | 36.54                    | 39.1                                     | NA                                              | 0                                                       |
| Nigeria (Uda)21                                  | 1989        | Longe        | 2925             | 6.15                     | 34.4                                     | NA                                              | 0                                                       |
| Nigeria (Imo River basin)24                       | 2007        | Osakwe       | 2500             | 20.80                    | 49.4                                     | NA                                              | 0/5/5                                                    |
| Nigeria (Utukpo, Benue state)17                  | 2007        | Osakwe       | 6000             | 4.67                     | 48                                       | NA                                              | 0/8/8                                                    |
| Nigeria (Ukpo)16                                 | 2010        | Nwani        | 6800             | 4.26                     | 44.5                                     | NA                                              | 0/11/11                                                  |
| Nigeria (Ilie, Osun state)31                     | 2013        | Mustapha     | 2212             | 4.52                     | 32.5                                     | NA                                              | 0/14/14                                                  |
| Nigeria (Imo River basin)10                      | 2018        | Siewe        | 843              | 4.74                     | 51                                       | 4.6                                              | 0/19/19                                                  |
| Senegal (Moyenne vallée)46                       | 1962        | Boutilier    | 32 300           | 1.92                     | NE                                       | NA                                              | NA                                                       |
| Senegal (Casamance, Kaolack, Thies)18            | 1982        | Ndiaye       | 7682             | 8.33                     | NE                                       | NA                                              | NA                                                       |
| Senegal (Piêkine)25                              | 2004        | Ndoye        | 4500             | 14.22                    | NE                                       | NA                                              | NA                                                       |
| Togo (Kloko)32                                   | 1989        | Balogou      | 19 241           | 12.32                    | 49                                       | NA                                              | 0                                                       |
| Togo (Akebou)32                                  | 1995        | Balingou     | 4182             | 13.15                    | 55                                       | NA                                              | 7/0/7                                                    |
| Togo (Tone)33                                    | 1995        | Balingou     | 9155             | 18.57                    | 71                                       | NA                                              | 18/0/18                                                  |
| Togo (Batamara)34                                | 2001        | Balingou     | 6249             | 15.68                    | 76                                       | NA                                              | 24/10/24                                                 |

NE: non-endemic; NA: not available
aVC/CDTI/overall: number of years of vector control/community-directed treatment with ivermectin/total duration of onchocerciasis control irrespective of the method
bReported by Wyatt (1971).45

<18 y and OR 2.09 [95% CI 1.29–3.40] in adults).29 Although the overall epilepsy prevalence was low (1.92%), further analysis of the data revealed a positive trend between the onchocerciasis seropositivity rate and the prevalence of epilepsy in the different study villages: Spearman’s rho = 0.54, p = 0.07. Other factors associated with epilepsy in children included family history of seizures (OR 3.31 [95% CI 1.83–5.96]), abnormal delivery (OR 2.99 [95% CI 1.07–8.34]) and problems after birth (OR 3.51 [95% CI 1.02–12.06]). In addition, the adult epilepsy risk factors included pork consumption (OR 1.68 [95% CI 1.09–2.58]) and exposure to Toxoplasma gondii (OR 1.99 [95% CI 1.15–3.45]).29

### Epilepsy prevalence in Ivory Coast

In the southern part of Ivory Coast, two studies reported a high epilepsy prevalence.36,37 These two studies also provided...
detailed descriptions of the PWE encountered during the surveys. In M’Brou, in the Agboville health district, a high burden of epilepsy had been suspected for decades. This village is located in the Agneby River basin just below the southern border of the OCP target areas and did not benefit from vector control interventions, as it was estimated that the risk for onchocercal blindness was lower in this forest area of the country.\(^4\) The OCP has undertaken detailed etiopathological studies in the Agneby basin, but only one epidemiological survey that was done in a village <10 km from M’Brou, which showed a prevalence of microfilariae of 76%. An epilepsy survey conducted in 1993 revealed that 38 of 920 inhabitants (41%) had epilepsy, and 28 PWE (37.7%) fulfilled the OAE criteria.\(^3\) In addition, two PWE in M’Brou presented with nodules from which adult worms of *O. volvulus* were extracted.\(^3\) Studies conducted in 1989 in Akoungou village (Tiassalé health district; onchocerciasis prevalence 43.4%)\(^5\) also revealed a high epilepsy prevalence; up to 18 of the 309 inhabitants (58%) were confirmed to have epilepsy, with 15 PWE (83.3%) satisfying the OAE criteria.\(^3\) In both villages, the peak age for seizure onset was 10–15 y (Figure 3) and the frequent family history of epilepsy steered the authors to conclude genetic/consanguinity aetiology.\(^3\) A low epilepsy prevalence of 7.65% was reported in a non-endemic study site of Ivory Coast in 1987.\(^3\) According to the authors, this was an underestimate because older PWEs most likely hid their condition due to stigma and thus went undetected by the research team.

**Epilepsy prevalence in Liberia**

High epilepsy prevalence (27.73–49.01%) was documented in the Grand Bassa onchocerciasis focus in Liberia.\(^1\) Both studies agreed that epilepsy was endemic in that region; in addition, youths were most affected.\(^4\) The local population distinguished two types of epilepsy: ‘to drop the head in the pan’ and ‘the big jerking’.\(^4\) Goudsmith et al.\(^4\) reported a frequent family history of epilepsy, psychological symptoms and burns among PWE in Liberia. Infection with *Taenia solium* was excluded as a possible cause of the high epilepsy prevalence observed.\(^3\)

Other suggested aetiologies of epilepsy included genetic/environmental factors and febrile illness.\(^2\)

**Epilepsy prevalence in Mali**

In 1999 in Mali, epilepsy was investigated in villages located in two zones with microfilaria prevalence of 9.3% for zone 1 (pre-control endemicity not known) and 23.0% for zone 2 (hyperendemic before the start of control). The epilepsy prevalence was 10.8% and 16.1% in zone 1 and zone 2, respectively (p=0.09).\(^2\) A case–control study further showed that 22.4% of PWE and 21.7% of controls had clinical and/or laboratory signs of onchoderciasis (OR 1.02 [95% CI 0.47–2.19]).\(^2\)

**Epilepsy prevalence in Nigeria**

In Nigeria in 1982, an epilepsy prevalence of 37% was observed in Aiyete,\(^2\) a rural onchocerciasis-endemic village, while a 5.3% prevalence was observed in Igbo-Ora, a town situated 20 km away and inhabited by the same ethnic group;\(^2\) both surveys found a high age-specific prevalence in the 10–19 y age group.\(^2\) In 2004, Dozie et al.\(^2\) reported an epilepsy prevalence of 12% in the then meso-endemic Imo River basin. The highest epilepsy prevalence was observed in villages with the highest onchocerciasis prevalence, with an overall positive correlation between both conditions in the study population (r=0.38, p<0.043).\(^5\) When this study site was revisited in 2018, the prevalence of microfilariae had dropped to 4.6%, with a drastic decrease of epilepsy prevalence to 5%.\(^3\)

**Epilepsy prevalence in Senegal**

As early as the 1960s, a low epilepsy prevalence of 1.92% was reported in the Moyenne Vallée of Senegal, which is non-endemic for onchocerciasis.\(^4\) However, this was most likely an underestimation of the true prevalence, as the data were not collected via a systematic approach. A more recent study conducted in non-endemic Pikine reported an epilepsy prevalence of 14.22%.\(^2\)

**Epilepsy prevalence in Togo**

All study sites in Togo had an epilepsy prevalence <20%.\(^3\) One of the studies from Togo found an association between epilepsy and cysticercosis;\(^3\) the prevalence of cysticercosis was 13.5% among PWE vs 38% in the general population (p<0.0001). Another study reported subcutaneous cysticercus cysts in 14/98 PWE (14.3%) and focal seizures in 38/98 (38.8%) of the PWE.\(^3\)

**Meta-analysis of epilepsy prevalence**

Considering all included epilepsy surveys, the pooled epilepsy prevalence in the West African study sites was 13.14 per 1000 (95% CI 11.28–15.00). This prevalence ranged from 1.92% in study sites located in Senegal and Ghana to 58.25% in an onchocerciasis focus in Ivory Coast. Figure 4 shows a forest plot with the result of a meta-analysis to compare the prevalence of epilepsy between four subgroups of epilepsy studies: one group of villages from areas without onchocerciasis and three groups of sites where onchocerciasis was endemic. These latter three groups represent different durations of onchocerciasis control.
Table 1: Study or Subgroup | Epilepsy prevalence | SE | Weight | Epilepsy prevalence | IV, Random, 95% CI | Epilepsy prevalence | IV, Random, 95% CI
---|---|---|---|---|---|---|---
1.1.1 Non-endemic | | | | | | |
Benin 1997 | 21.05934907 | 2.56478614 | 3.2% | 21.06 [16.03, 26.09] | | 
Benin 2007 | 31.20759837 | 6.40489712 | 1.4% | 31.21 [18.65, 43.76] | | 
Burkina Faso 1989 | 10.645339 | 0.79588261 | 4.1% | 10.65 [9.09, 12.21] | | 
Burkina Faso 2007 | 4.26787741 | 6.92998041 | 1.3% | 4.27 [3.09, 5.75] | | 
Gambia 1997 | 4.25925926 | 0.51161442 | 4.2% | 4.26 [3.26, 5.26] | | 
Ivory Coast 1987 | 7.65396122 | 2.5412401 | 3.3% | 7.65 [2.67, 12.63] | | 
Senegal 1962 | 1.91905484 | 0.24354262 | 4.2% | 1.92 [1.44, 2.40] | | 
Senegal 1982 | 8.33116376 | 1.03704838 | 4.0% | 8.33 [6.30, 10.36] | | 
Senegal 2004 | 14.22222222 | 1.76509053 | 3.7% | 14.22 [10.76, 17.68] | | 
Subtotal (95% CI) | 29.5% | 12.20 [8.53, 15.86] | | 
Heterogeneity: Tau² = 24.88; Chi² = 286.38, df = 8 (P < 0.00001); I² = 97%
Test for overall effect: Z = 6.52 (P < 0.00001)

1.2.2 Endemic: No onchocerciasis control
Ivory Coast 1989 | 58.25242718 | 13.32431919 | 0.5% | 58.25 [32.14, 84.37] | | 
Ivory Coast 1993 | 41.30437483 | 6.56011914 | 1.4% | 41.30 [28.45, 54.16] | | 
Liberia 1981 | 27.7276826 | 2.465216 | 3.3% | 27.73 [22.90, 32.56] | | 
Liberia 1983 | 48.01374776 | 5.27834903 | 1.8% | 48.01 [38.87, 59.36] | | 
Nigeria 1982a | 6.32869051 | 0.52880049 | 4.2% | 6.33 [4.29, 6.37] | | 
Nigeria 1982b | 36.5448505 | 6.24431741 | 1.5% | 36.54 [24.31, 48.78] | | 
Nigeria 1989 | 6.15384615 | 1.44600556 | 3.9% | 6.15 [3.32, 8.99] | | 
Togo 1989 | 12.31744712 | 0.79616128 | 4.1% | 12.32 [10.76, 13.88] | | 
Subtotal (95% CI) | 20.6% | 24.45 [17.76, 31.14] | | 
Heterogeneity: Tau² = 71.82; Chi² = 240.51, df = 7 (P < 0.00001); I² = 97%
Test for overall effect: Z = 7.16 (P < 0.00001)

1.3.3 Endemic: 1-9 years of onchocerciasis control
Benin 1994 | 15.24601525 | 3.22585789 | 2.8% | 15.25 [9.82, 21.57] | | 
Benin 1995 | 25.84493042 | 7.0745018 | 1.3% | 25.84 [11.98, 39.71] | | 
Nigeria 2004 | 11.94890812 | 1.5995666 | 3.8% | 11.95 [8.89, 15.01] | | 
Nigeria 2007a | 20.8 | 2.8542852 | 3.1% | 20.80 [15.21, 26.39] | | 
Nigeria 2007b | 4.66666667 | 0.87956869 | 4.1% | 4.67 [2.94, 6.39] | | 
Togo 1995a | 13.1516021 | 1.7616918 | 3.7% | 13.15 [9.70, 16.60] | | 
Subtotal (95% CI) | 18.8% | 13.99 [8.45, 19.53] | | 
Heterogeneity: Tau² = 38.20; Chi² = 59.70, df = 5 (P < 0.00001); I² = 92%
Test for overall effect: Z = 4.95 (P < 0.00001)

1.4.4 Endemic: ≥10 years of onchocerciasis control
Benin 2005 | 12.68426486 | 1.03600447 | 4.0% | 12.68 [10.65, 14.71] | | 
Ghana 2010 | 1.91865657 | 0.12144172 | 4.2% | 1.92 [1.68, 2.16] | | 
Malawi 2009 | 13.95113495 | 1.5850774 | 3.8% | 13.95 [10.24, 16.66] | | 
Nigeria 2010 | 4.26470588 | 0.79024561 | 4.1% | 4.26 [2.72, 5.81] | | 
Nigeria 2013 | 4.52079566 | 1.42636968 | 3.9% | 4.52 [1.73, 7.32] | | 
Nigeria 2018 | 4.74495848 | 2.63648349 | 3.4% | 4.74 [0.11, 9.38] | | 
Togo 1995b | 18.56908793 | 1.41089917 | 3.9% | 18.57 [15.80, 21.33] | | 
Togo 2001 | 15.6825092 | 1.57107016 | 3.8% | 15.68 [12.60, 18.76] | | 
Subtotal (95% CI) | 31.1% | 9.44 [4.83, 14.05] | | 
Heterogeneity: Tau² = 42.24; Chi² = 373.52, df = 7 (P < 0.00001); I² = 98%
Test for overall effect: Z = 4.02 (P < 0.00001)

Total (95% CI) | 100.0% | 13.14 [11.28, 15.00] | | 
Heterogeneity: Tau² = 21.23; Chi² = 1243.11, df = 30 (P < 0.00001); I² = 98%
Test for overall effect: Z = 13.85 (P < 0.00001)
Test for subarour differences: Chi² = 13.76, df = 3 (P = 0.003), I² = 78.2%

Figure 4. Forest plot and pooled epilepsy prevalence (per thousand) in the West African study sites.

In onchocerciasis-endemic sites with no control, the prevalence of epilepsy was 24.45%, i.e. twice as high as in the sites without onchocerciasis (12.20%). However, sites where control measures had been implemented had a significantly lower

no control (onchocerciasis control not yet started at the time of the epilepsy survey), 1–9 y of control (median 7 y) and ≥10 y of control (median 15.6 y). The main results are summarized in Figure 5.
The pooled epilepsy prevalence in the West African study sites obtained in this study was 13.14%, very similar to the median epilepsy prevalence in sub-Saharan Africa of 14.2%. However, after taking onchocerciasis endemicity and onchocerciasis control into account, there were significant differences. First, villages in onchocerciasis-endemic areas had a double burden of epilepsy before the start of control compared with villages from areas that were onchocerciasis free. Second, in the onchocerciasis-endemic areas there was a statistically significant relationship between the prevalence of epilepsy and the pre-control onchocerciasis endemicity level. Third, after correcting for baseline endemicity, the prevalence of epilepsy declined significantly in relation to the duration of onchocerciasis control. Together, these three findings suggest that onchocerciasis has also been an important cause of epilepsy in West Africa, as it was previously demonstrated in East and Central Africa. Furthermore, our results provide empirical evidence that onchocerciasis control can significantly reduce OAE.

The onchocerciasis landscape in West Africa had been particularly influenced by the activities of the OCP. In 11 West African countries, the OCP (1974–2002) was very successful in controlling onchocerciasis in the savannah areas, which were priority zones due to high blindness rates. The initial strategy was by vector control and, from the early 1990s, ivermectin was used as an additional tool. The large-scale onchocerciasis elimination control measures deployed in West Africa by the OCP came two decades earlier than those implemented later in Central and East Africa.

Given that many epilepsy prevalence studies in West Africa were performed after the introduction of OCP, only a few reported a high epilepsy prevalence in onchocerciasis-endemic regions. Our findings show that the epilepsy prevalence depended on the pre-control onchocerciasis endemicity, and longer periods of onchocerciasis control were associated with a lower epilepsy prevalence. However, the potential role of *O. volvulus* infection was not often discussed when researchers were confronted with an unusually high epilepsy burden in some West African countries. In 11 West African countries, the OCP (1974–2002) was very successful in controlling onchocerciasis in the savannah areas, which were priority zones due to high blindness rates. The initial strategy was by vector control and, from the early 1990s, ivermectin was used as an additional tool. The large-scale onchocerciasis elimination control measures deployed in West Africa by the OCP came two decades earlier than those implemented later in Central and East Africa.

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Given that many epilepsy prevalence studies in West Africa were performed after the introduction of OCP, only a few reported a high epilepsy prevalence in onchocerciasis-endemic regions. Our findings show that the epilepsy prevalence depended on the pre-control onchocerciasis endemicity, and longer periods of onchocerciasis control were associated with a lower epilepsy prevalence. However, the potential role of *O. volvulus* infection was not often discussed when researchers were confronted with an unusually high epilepsy burden in some West African countries. In 11 West African countries, the OCP (1974–2002) was very successful in controlling onchocerciasis in the savannah areas, which were priority zones due to high blindness rates. The initial strategy was by vector control and, from the early 1990s, ivermectin was used as an additional tool. The large-scale onchocerciasis elimination control measures deployed in West Africa by the OCP came two decades earlier than those implemented later in Central and East Africa.

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Table 2. Multiple linear regression model investigating the factors associated with epilepsy prevalence in the study sites

| Model covariates                                      | Logit regression coefficient | 95% confidence interval | P-value |
|-------------------------------------------------------|-----------------------------|-------------------------|---------|
| Intercept                                             | -6.343                      | -7.204 -5.483           | <0.001  |
| Pre-control onchocerciasis endemicity (prevalence of microfiladermia) | 0.046                       | 0.030 -0.063            | <0.001  |
| Duration of onchocerciasis control (years)            | -0.059                      | -0.076 -0.042           | <0.001  |

Adjusted $R^2=0.782$.

Figure 6. Plots and lines showing the relationship between epilepsy and pre-control onchocerciasis endemicity, taking into account the duration of onchocerciasis control in each endemic study site. (A) Logit-transformed scale for epilepsy prevalence. (B) Linear scale for epilepsy prevalence. The solid line represents the previously published curve by Pion et al.; the black dotted line represents endemic sites with no control measures; the blue dotted line represents sites with 1–9 y of onchocerciasis control, which is almost superimposed on the previously published curve by Pion et al.; the red dotted line shows a downward shift of epilepsy prevalence in sites with ≥10 y of onchocerciasis control. (mf: detectable O. volvulus microfilariae in skin snips.)

It is conceivable that other health interventions (such as malaria control or improved neonatal care) may have reduced the epilepsy burden, alongside onchocerciasis control. However, the multiple regression analysis did not show a significant association between the epilepsy prevalence and the study year, suggesting that improved healthcare in more recent studies cannot explain the disparity in epilepsy prevalence observed in the different sites. Certainly, epilepsy resulting from perinatal brain insult was more common in the past, and typically started within the early years of life as described by Collomb et al.\textsuperscript{11} in a non-endemic region in Senegal. However, in onchocerciasis-endemic areas with suboptimal control measures, epilepsy onset is typically more prevalent after the age of 5 y and before the age of 20 y\textsuperscript{2} (Figure 3). Therefore the epilepsy incidence in the 5–20 y age group would hardly be affected by improved perinatal care. Nonetheless, improved access to care would increase the life expectancy of PWEs and result in a cohort age shift of PWEs as observed in an onchocerciasis focus in central Cameroon.\textsuperscript{51}

It is worth noting that in the two epilepsy studies conducted in Liberia (which was not under the umbrella of OCP, albeit with a high onchocerciasis endemicity\textsuperscript{59}), a high epilepsy prevalence >25% was reported.\textsuperscript{41,42} On the other hand, the epilepsy prevalence was particularly low in non-endemic areas of Senegal\textsuperscript{44} and Gambia.\textsuperscript{26} The epilepsy prevalence was also low in Kintampo, Ghana, despite a high seroprevalence of O. volvulus Ov16 antibodies observed in these villages.\textsuperscript{29} It should be noted that by the time of the survey, Kintampo had benefited from a very efficient vector control programme by OCP followed by biannual ivermectin distribution.\textsuperscript{60} Therefore this high Ov16 IgG seropositivity rate among the adult population was more representative of a high transmission of onchocerciasis in the past that had already
been controlled by the time of the survey, resulting in a drastic decline of the epilepsy burden.

The epidemiological and clinical criteria of OAE\(^2\) were observed in many endemic sites in West Africa.\(^{36,37,41,42}\) similar to findings from several other onchocerciasis-endemic villages.\(^{35,61,62}\) In Liberia, the description of early-onset focal seizures accompanied by cognitive symptoms and manifesting as rhythmic dorsoventral movements of the head\(^{49,63}\) most likely refer to cases of nodding syndrome,\(^2,64\) as reported in other parts of Africa. Another argument in favour of the presence of OAE in West African villages is that most PWEs encountered were <25 y of age, suggesting a reduced life expectancy. Analysis of a 25-year follow-up database of 295 909 persons with onchocerciasis in OCP countries revealed an increased risk of mortality with increasing skin microfilariae load, and particularly among younger age groups (<20 y old).\(^{55,65}\) Given that OAE typically starts between the ages of 3 and 18 y,\(^1\) we deduced that the high mortality observed among O. volvulus–infected persons <20 y of age is not a consequence of ocular manifestations of onchocerciasis, which are more frequent during adulthood. Instead, premature mortality due to epilepsy, as observed in other onchocerciasis-endemic villages,\(^{61,68}\) is a more plausible explanation.

While infection with O. volvulus may indeed be responsible for many epilepsy cases in endemic sites, neurocysticercosis was suggested as a possible aetiology in Benin,\(^3\) Togo,\(^3,4\) and Nigeria.\(^19\) Generally, in communities with ongoing T. solium transmission, only about one-third of epilepsy cases can be attributed to neurocysticercosis.\(^19\) Meanwhile, the population-attributable fraction of O. volvulus infection in epilepsy was estimated at 91.7% in onchocerciasis-endemic villages in the Mbang valley in Cameroon,\(^1\) suggesting that onchocerciasis may account for a larger proportion of epilepsy cases than neurocysticercosis in co-endemic settings. A recent survey in Maridi, South Sudan (where there are no pigs and consequently no T. solium transmission to cause neurocysticercosis) showed a high prevalence of epilepsy, particularly in villages closer to blackfly breeding sites;\(^58\) >85% of the PWEs in onchocerciasis-endemic villages satisfied the OAE criteria.\(^62\) Similarly, neurocysticercosis could not account for the high epilepsy prevalence observed in Liberia prior to implementing onchocerciasis control in that area.\(^62\) All these findings strongly suggest that onchocerciasis is an important, yet neglected contributor to the epilepsy burden in sub-Saharan Africa. Our results also suggest that within 10 y of implementing onchocerciasis control, the epilepsy burden could be reduced significantly in endemic communities. However, these are very imprecise estimates, as we did not take into account the method, frequency and coverage of the onchocerciasis control strategies (aerial vs ground larviciding, annual vs biannual ivermectin distribution).

The strength of our study resides in the fact that we included epilepsy data obtained during the pre-onchocerciasis control period, thus increasing the chances of finding high endemicity and consequently high epilepsy prevalence. However, we must point out that some researchers in the past had a tendency to investigate only villages with anecdotal reports of high seizure frequency, regardless of onchocerciasis endemicity. This may explain why a substantial epilepsy burden was observed in some communities with little or no onchocerciasis transmission, most likely due to a panoply of aetiologies besides infection with O. volvulus. As a major weakness of our study, the pre-control onchocerciasis prevalence for the exact study sites was generally not available and we therefore used the predicted prevalence for each location from spatial maps of onchocerciasis endemicity in West Africa. Although these maps are quite detailed, they do not always capture local fluctuations in endemicity. For a few sites, we estimated the pre-control endemicity from available survey data for nearby villages (5–10 km distance) where this was considered more reliable. For instance, the pre-control data for M’Brou village (Ivory Coast) were extrapolated from the neighbouring village of Emankono Camp (<10 km away), which had been investigated in detail by the OCP. Another limitation is the fact that obtaining precise values for the pre-control community microfilariae load (a better measure of onchocerciasis endemicity) for each study site was practically impossible. We must also point out the fact that migration into and out of the different study communities, which may have influenced the exposure of the migrants to onchocerciasis, was not taken into account during our analysis, as this information was not available. Finally, the lack of solid evidence (imaging, electroencephalography, laboratory investigations) to confirm or exclude other causes of epilepsy in most of the included studies is also a caveat worth mentioning.

**Conclusions**

This review adds to the existing epidemiological evidence suggesting that onchocerciasis is able to cause epilepsy in all onchocerciasis-endemic regions with high ongoing O. volvulus transmission. OAE has most likely occurred in onchocerciasis foci in West Africa prior to and during the early years of onchocerciasis control efforts. Subsequently, onchocerciasis control appears to have significantly reduced the prevalence of OAE. These findings are relevant for areas still experiencing substantial onchocerciasis transmission, as these should be prioritized for interventions in order to curb the dual burden of epilepsy and onchocerciasis.\(^70,71\)

**Authors’ contributions:** CR and SFJN conceived the study and performed the literature review. JHFR provided data on onchocerciasis baseline endemicity and control, while PMP provided grey data on epilepsy prevalence in West Africa. SFJN extracted the data. SFJN and JHFR performed the data analysis. SFJN wrote the first draft. All authors revised the initial draft and approved the final manuscript.

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