Nodding syndrome in Ugandan children—clinical features, brain imaging and complications: a case series


ABSTRACT

Objectives: Nodding syndrome is a devastating neurological disorder of uncertain aetiology affecting children in Africa. There is no diagnostic test, and risk factors and symptoms that would allow early diagnosis are poorly documented. This study aimed to describe the clinical, electrophysiological and brain imaging (MRI) features and complications of nodding syndrome in Ugandan children.

Design: Case series.

Participants: 22 children with nodding syndrome brought to Mulago National Referral Hospital for assessment.

Outcome measures: Clinical features, physical and functional disabilities, EEG and brain MRI findings and a staging system with a progressive development of symptoms and complications.

Results: The median age of symptom onset was 6 (range 4–10) years and median duration of symptoms was 8.5 (range 2–11) years. 16 of 22 families reported multiple affected children. Physical manifestations and complications included stunting, wasting, lip changes and gross physical deformities. The bone age was delayed by 2 (range 1–6) years. There was peripheral muscle wasting and progressive generalised wasting. Four children had nodding as the only seizure type; 18 in addition had myoclonic, absence and/or generalised tonic–clonic seizures developing 1–3 years after the onset of illness. Psychiatric manifestations included wandering, aggression, depression and disordered perception. Cognitive assessment in three children demonstrated profound impairment. The EEG was abnormal in all, suggesting symptomatic generalised epilepsy in the majority. There were different degrees of cortical and cerebellar atrophy on brain MRI, but no hippocampal changes. Five stages with worsening physical, EEG and brain imaging features were identified: a prodrome, the development of head nodding and cognitive decline, other seizure types, multiple complications and severe disability.

Conclusions: Nodding syndrome is a neurological disorder that may be characterised as probably symptomatic generalised epilepsy. Clinical manifestations and complications develop in stages.

ARTICLE SUMMARY

Article focus

- This paper offers detailed descriptions of the clinical features and complications of nodding syndrome in Ugandan children and the electrophysiological and brain imaging features.
- It also proposes a clinical staging system for the disease.

Key messages

- Nodding syndrome is an epidemic neurological disorder affecting children in parts of sub-Saharan Africa that may be characterised as a probable symptomatic generalised epilepsy with features of epileptic encephalopathy.
- Patients progressively develop both physical and functional deficits including multiple seizure types, cognitive and physical decline, malnutrition and psychiatric features. Five clinical stages could be identified.
- The proposed clinical stages are associated with worsening cortical and cerebellar atrophy on brain imaging and more severe epileptiform and background EEG changes. These stages may be useful in guiding treatment and rehabilitation.

Strengths and limitations of this study

- Although the sample size is small and there is no comparison group, this is one of the few studies so far to have carefully documented the clinical features and complications of nodding syndrome combined with extensive electrophysiology and brain imaging data, describe the natural history and the first to provide a staging system. The study patients, however, may not be representative of the population, as they were not randomly drawn from the community.
- The study did not investigate the aetiology and the proposed staging was mainly derived from parental descriptions rather than prospective observations and, therefore, suffers from recall bias.
- The resolution of our brain MRI is quite low.
Nodding syndrome in Ugandan children

which might be useful in defining treatment and rehabilitation. Studies of risk factors, pathogenesis, management and outcome are urgently needed.

BACKGROUND

Nodding syndrome is a devastating neurological disorder of uncertain aetiology described in African children. It was first described in Tanzania in 1960s and subsequent reports have come from Liberia, South Sudan and Uganda. The syndrome is characterised by head nodding determined to be atonic seizures often occurring in association with feeding, a cold breeze or cold weather, and complicated by other seizure types, malnutrition and cognitive decline.

In Uganda, almost all affected individuals are from the north of the country where there are an estimated 3000 cases. The region, for the past 20 years, had instability from rebel activity. As a result, the population was internally displaced into densely populated camps. It is only in the last 5 years that peace returned and the population returned to their homes. This region is crossed by two rivers—the Asua and Pager Rivers, has high malaria transmission and is endemic for Onchocerca volvulus. This parasite has variously been associated with the Nakalangle syndrome (a tropical syndrome characterised by short stature and malnutrition), epilepsy and nodding syndrome. This association has, however, been indirect as no O volvulus contamination of cerebrospinal fluid has been documented.

There are only limited descriptions of nodding syndrome. Winkler et al provided the most detailed account of the syndrome to date, describing clinical features in 62 Tanzanian patients and classifying them as either head nodding only or head nodding plus, if they also had other seizure types. Initial symptoms allowing early recognition of the disease, its natural history and potentially modifiable risk factors are poorly characterised. There is no diagnostic test and the current case definition is based solely on clinical criteria. The objective of this study was to describe the clinical, electrophysiological and brain imaging features and complications of nodding syndrome in Ugandan children and to propose a staging system.

METHODS

Design and setting

This is a case series of 22 Ugandan children with nodding syndrome. The study was conducted in Mulago, the National Referral Hospital in Uganda and teaching hospital for Makerere University College of Health Sciences in Kampala. This hospital provides tertiary level care for patients in a country in which most public healthcare services are paid for by the state.

Participants

Participants were patients with suspected nodding syndrome brought by the Ministry of Health from Kitgum district near the border with South Sudan, to Mulago Hospital in March 2012 for specialist assessments to better understand the syndrome. Kitgum district is the epicentre of the disease and one of the most affected districts in the country. Of the 25 patients brought to Mulago, 1 young adult (a 23-year-old man was found to have a brain tumour) and 2 adolescents (an 18-year-old girl with a cerebellar hypoplasia syndrome and a 16-year-old boy with a history of cerebral malaria at the age of 4 years and subsequent neurological sequelae) were excluded. The remaining 22 children had probable nodding syndrome and were included in the study. A case of probable nodding syndrome was defined as

- A child older than 2 years or an adolescent who previously was developing normally.
- Two or more episodes of recurrent head nodding occurring spontaneously or consequent to the sight of food or coldness.
- With or without other types of seizures, neurological signs, regression in growth or learning disability.

This case definition was revised during the International Meeting on Nodding Syndrome later in 2012. However, all selected patients fulfilled the revised criteria.

Permission for the study was obtained from Makerere University School of Medicine Research and Ethics Committee. However, we did not have a study protocol prior to the arrival of the patients, clinical care and assessment followed the hospital’s standard procedures for routine non-surgical care for children. Verbal parental consent was obtained for all clinical, laboratory and imaging procedures. As is policy, however, parents gave written consent for photography, as this is considered over and above routine care and for any surgical procedures. Parents were specifically made aware that the objective of the assessments was not a cure for the disease, but a better understanding of the disease and that the general findings from the evaluation of the group of patients with nodding syndrome would be made available to the wider scientific community in presentations and publications, with a specific aim of improving the care for people affected by the disorder in the future. To this effect, a submission was then made to the Ethics Committee and permission to use results of the investigations was subsequently granted.

Study measurements and procedures

All had detailed clinical, electrophysiological and brain imaging assessments and laboratory testing.

Clinical assessment

The history included an enquiry about the time from pregnancy to the onset and the progressive development of symptoms, physical and functional difficulties. The clinical examination included general, nutritional, neurological,
cognitive and mental state assessments. Wasting was defined as weight for height Z score of \(<-2\) and stunting as height for age Z score of \(<-2\). Sexual maturity was assessed using the Tanner Sexual Maturity staging. Patients were classified as having nodding syndrome only or nodding syndrome plus depending on whether they also had other seizure types.\(^9\) x-Rays of the left wrist were taken for bone age and reported using a Greulich and Pyle Atlas by a blinded radiologist.\(^{20}\) Bone growth was considered delayed if it was 2 years below the chronological age. Cognitive functioning was assessed in detail in four children—median age 14.5 (range 13–15) years—2 weeks after the initiation of sodium valproate using the Kaufman Assessment Battery for Children 2nd Edition (KABC-2). This test has previously been adapted for use in Uganda.\(^{21}\) All four had symptoms for longer than 5 years.

**Laboratory procedures**

Ten millilitres of blood was drawn for full blood count, erythrocyte sedimentation rate (ESR), malaria parasites, electrolytes, liver and renal function tests and HIV testing. Cerebrospinal fluid was examined for cells, glucose, protein, microscopy and bacteriological culture. In addition, 10 children had a skin snip examined for \(O\) volvulus as previously described.\(^9\)

**Neurophysiology and imaging**

All had a 30-min EEG recording with an XLTEK EEG system (Optima Medical Ltd, London, UK) using the 10–20 electrode placement system, which was reviewed by a consultant clinical neurophysiologist (SW) in the UK. Brain MRI in T1, T2 and Flair sequences were obtained without contrast in 19/22 patients using a 0.5 Tesla machine (BASDA Medical Apparatus, Guangzhou) and the images were also examined in the UK by a Consultant Neuroradiologist (KC).

**Natural history and staging**

At the end of the first week, after the patients and carers had acclimatised to the referral hospital environment, the attending clinician sat with each carer and obtained detailed histories of the progressive development and timing of symptoms and complications of nodding syndrome to characterise the natural history. A standardised proforma with a diagrammatic representation of possible events and a linear timeline was used to obtain these descriptions. Information from each patient was plotted on the timeline and later combined with the data from other patients to identify emerging patterns, from which the proposed stages of the disorder were derived.

**Treatment**

The patients were started on symptomatic treatment (sodium valproate for seizure control, nutritional and physical therapy, counselling and social support) according to the national guidelines developed a month earlier.\(^{22}\) In this protocol, all received sodium valproate starting at 10 mg/kg/day. The dose was titrated in 5–10 mg/kg/day increases according to the level of seizure control. Nutritional rehabilitation included Ready to Use Therapeutic Foods (Plumpy’Nut, Nutriset, Malanay) and locally prepared food. Occupational and physiotherapy, family counselling and support were provided as appropriate.

**Data analysis**

Data were analysed using STATA V.12 (STATA Corp, Texas, USA). Results are summarised as frequencies, proportions and medians as appropriate. The clinical features, complications and disability were then used to describe treatment and rehabilitation needs. Clinical stages were identified and brain imaging and EEG correlated with the clinical stages.

**RESULTS**

**General features**

Nine patients (40.9\%) were men. The age range was 12–18 years. The median age at onset of symptoms was 6 (range 4–10) years and the median duration of symptoms was 8.5 (range 2–11) years (table 1). Sixteen of the 22 families reported more than one case (median 2 (range 0–4)). Prior to hospitalisation, all patients had received antiepileptic drug treatment with phenobarbitone, phenytoin or carbamazepine with no clear documentation of benefit. Treatment had often been intermittent and at subtherapeutic doses.

Striking features on physical examination included stunting, cognitive impairment (on KABC or performance of basic tasks), lip changes and other physical deformities. Several children had burns and scars from burns. The skin was dry, thin and scaly. Extremities, especially the feet, felt cold with a temperature gradient with the trunk, but a normal capillary refill time. Among those with mild lip changes, the lower lip was enlarged with no visible or palpable localised swellings. In progressively more severe cases, the mucosa was deep purple, with soft papular growths and increasingly large, thick bands of tissue. One child, not exposed to sodium valproate previously, had unexplained alopecia.

**Growth and sexual maturity**

Sixteen of the 22 children were wasted and 9 stunted. Increasingly more severe wasting was observed with a longer duration of symptoms. Thus, patients with moderate wasting had symptoms lasting 3–6 years, while severe wasting was more common among those with symptoms lasting longer than 7 years. Ten had sexual maturity assessed; 3/10 children (aged 12, 13 and 14 years) scored Tanner stage 3. The remaining seven children (ages 12–17 years) were either Tanner stage 1 or 2. The bone age was delayed by a median 2 (range 1–6) years.

**Musculoskeletal findings**

Almost all had peripheral muscle wasting with progressively flat feet and hands, thin cylindrical digits and...
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<th>Duration of symptoms in yrs</th>
<th>Seizure based classification</th>
<th>Main Psychiatric morbidity</th>
<th>Other prominent complications</th>
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Nodding syndrome in Ugandan children
increasing generalised wasting. Other physical changes included kyphosis and pectus deformities of the chest. Flexion limb deformities were seen in patients with severe disability.

**Neurological, cognitive and behavioural features**

**Seizures**

Other than head nodding, a variety of other seizure types were described in 18/22 patients—including absence, complex partial, myoclonic and tonic-clonic seizures.

**Head nodding**

Nodding was precipitated by food in 16/22 children. In 4/22, it was associated with cold weather or a cold breeze, while in 13/22 it developed spontaneously. Nodding episodes came in clusters occurring during both the day and the night and were characterised by repetitive flexion and forward drop of the head. The clusters of head nodding lasted several seconds to minutes. Some children became unresponsive and stared blankly with each cluster of nodding, stopped feeding or drooled saliva.

Initially, head nodding was the predominant seizure type, but as the disease progressed, generalised tonic-clonic seizures became more prominent. Myoclonic seizures were not often reported, but were observed in several children while in hospital. One such child had a prolonged cluster of nodding with concurrent myoclonic jerks involving both upper limbs, lasting about 10 min. In a second child, similar myoclonic jerking was followed by a generalised tonic-clonic seizure. Several children had sudden falls, sustaining facial and head injuries.

Four children experienced paroxysmal events associated with fear, panic and visual hallucinations. We could not obtain clear descriptions of the images seen by two. The third child would shout and run with the onset of hallucinations and the fourth reported seeing a person with knives whose intention ‘was to kill her’. None of these events were captured on EEG.

**Other neurological complications**

Focal neurological signs were uncommon. There were no obvious cranial nerve palsies. However, six children were lethargic with an apathetic and expressionless face or ‘myopathic faces’. Three of the six drooled saliva, while two had very slow speech and repeated epileptiform (spike and wave) discharges on EEG. The deep tendon reflexes were increased in a minority of patients. In the majority, however, the reflexes were either normal or reduced.

**Vision, hearing and speech difficulties**

No parent reported visual impairment and we did not test visual acuity, but hearing impairment was reported in one child. Speech difficulties were reported in 10/22 children. These included immature speech for age and slow, slurred or dysarthric speech. Two children were mute, but retained gestural ability and receptive language.

**Behaviour and psychiatric features and complications**

The earliest psychiatric manifestation was the wandering behaviour or running away. Owing to the concerns about injury or getting lost, some parents tied up the patients to restrain them at home. Aggression, particularly towards familiar people, was reported in 6/22 cases, manifesting 3–6 years after onset of nodding. In two children, the onset was concurrent with wandering behaviour. Five children had sleeping difficulties and at least 8/22 had moderate-to-severe mood problems with one clinically depressed.

**Cognitive function**

All 22 children had cognitive difficulties and were out of school. Academic performance declined with symptom onset; as symptoms increased, pupils started getting poorer grades and were eventually withdrawn from school within 2–4 years of disease onset. Three of the four children who had cognitive functioning assessed using the KABC-2 responded to the test instructions. The fourth child did not respond at all. All three children had severe cognitive impairment (table 2).

**Laboratory findings**

The mean haemoglobin was 12.4 (range 10.8–14.8) g/dl and the mean red cell volume was 81.4 (range 65.8–69.8).

<table>
<thead>
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<th>Patient 2 Male 15 years</th>
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<td>Age equivalent in years</td>
<td>Test score</td>
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<td>&lt;5</td>
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<td>&lt;5</td>
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<tr>
<td>Learning</td>
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<td>&lt;5</td>
<td>25</td>
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<tr>
<td>Visual spatial</td>
<td>0</td>
<td>&lt;5</td>
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<tr>
<td>Knowledge</td>
<td>11</td>
<td>&lt;5</td>
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Stage 1: prodromal period
This poorly defined and short-lived period was reported in four patients. The earliest symptoms included “dizziness” and increasing inattention. The children were excessively sleepy, lethargic and would sometimes stare blankly during meals.

Stage 2: development of head nodding
Among the four patients reporting prodromal symptoms, head nodding developed within 6 weeks of the prodrome. In the majority, however, the initial feature was an abrupt onset of nodding. Subsequently, the parents reported declining cognitive abilities and behaviour difficulties. Disease progression, however, appeared to be arrested in these four.

Stage 3: development of other seizure types
Apart from the four children with nodding only, 18 children developed other seizure types including absence, complex partial, myoclonic and generalised tonic–clonic seizures. One child developed generalised tonic–clonic seizures almost simultaneously with the nodding. In the others, however, additional seizure types developed 1–3 years after initial symptoms. It was around this time that almost all school-going children dropped out of school.

Stage 4: development of multiple complications
Multiple complications developed 4–8 years after the initial symptoms, associated with marked regression in achievement. These consisted of deteriorating behaviour and psychiatric symptoms and a decline in motor, speech and other cognitive functions. Some patients developed physical deformities including kyphosis, limb and pectus deformities. Some sustained severe facial injuries with ‘drop attacks’ and burns. Those who were still independently mobile would wander about or run away and were prone to getting lost. Changes in the architecture of the lower lips also occurred at this time. With disrupted and poor feeding, the children became severely wasted.

Stage 5: severe disability
These children have little, if any, independent mobility. The general picture is that of a severely wasted child with apathy and depressive features including a flat affect, poor appetite and limited speech. Some had contractual changes around the major joints.

Patients with head nodding only had less cortical and cerebellar atrophy on brain MRI compared with those with multiple complications. In addition, there was a clear gradation from milder to more severe epileptiform and background EEG abnormalities in patients with the later clinical stages of the disease.

Response of seizures to sodium valproate
The patients had previously been on mostly low doses of various antiepileptic drugs including phenytoin, phenobarbital and carbamazepine. In conformity with the
proposed national guidelines, all were started on sodium valproate and the other antiepileptic drugs weaned off. Prior to this, a 24 h seizure count was obtained and this was repeated 14 days later. Overall, there was a 57% reduction in total seizures including clusters of nodding. The median total daily number of seizures reduced from 5 (range 2–14) on admission to 2 (range 0–8) 14 days after initiation of sodium valproate. Concurrent

**Figure 1** EEG recordings. An EEG recording of a 12-year-old girl. She had head nodding and cognitive impairment. During the recording, interictal epileptiform discharges (spikes and sharp waves) were observed in wakefulness (A) and during light sleep, when more prominent epileptiform discharges were evident. There was no apparent clinical change with these discharges (B).
improvements were also seen on the EEG with substantially reduced or absent interictal discharges in 3/5 patients who had repeat recordings on the day of assessment (figure 1).

**DISCUSSION**

Recently, there have been media reports of ‘a mysterious disease’ baffling scientists—the nodding syndrome.7 23 24 There are many uncertainties about this newly recognised disorder: what is the cause, the pathogenesis, disease classification, clinical spectrum and treatment? In this paper, we describe the clinical features and complications of nodding syndrome in Ugandan children, together with the EEG and brain imaging appearances. Our findings suggest that nodding syndrome is a neurological disorder characterised by a symptomatic generalised epilepsy.

**Clinical features and complications**

Nodding syndrome in Ugandan children manifests with head nodding, cognitive dysfunction, psychiatric features and/or multiple seizure types. It may be complicated by stunted growth, pubertal delay, wasting, motor decline and physical deformities. The earliest manifestation is a poorly defined prodrome followed within weeks by head nodding. In the later stages, there is cognitive dysfunction, psychiatric disturbance and severe muscle wasting and musculoskeletal deformity. Delayed physical growth, bone age and sexual maturity are common in affected children. This may partly be a result of delayed puberty, since most were in Tanner stage 2. Pubertal delay may be secondary to chronic illness, poor nutrition and/or

**Box 1** A mum’s description of the sequential chronology of symptoms and disease progression in her 18-year-old daughter

She was growing well until the age of 8 years when symptoms of nodding began. The head nodding is triggered by food. When food is given, she freezes with it in her hand, stares blankly into space with a fixed gaze, and then nods repeatedly for a time which varies with each episode but the maximum time was initially 5 minutes. The symptoms got worse with time and about 6 years later, the nodding symptoms were immediately followed by or associated with big seizures during which the whole body shook. She would drool saliva, foam around the mouth and loses consciousness. After the big seizure, she would sleep and on waking is often weak and sometimes disoriented. On some nights, she reports seeing a figure that holds a knife and wants to kill her. She is distressed by her illness and gets embarrassed on waking if she had a seizure in public. She is very quiet but sometimes aggressive. Overtime, her speech has become sluggish. Although she is 18 years old, she still has childish behaviour which is evident as she speaks. Her father died following a febrile illness. She has six siblings two of who have similar symptoms.

![Figure 2](image-url) Brain MRI. T2-weighted brain MRI in the axial (A and B) and sagittal (C and D) plane showing marked cerebellar atrophy and generalised cerebral atrophy.
Nodding syndrome in Ugandan children

psychosocial deprivation. If so, improved nutrition, a supportive environment and symptom control should lead to improved growth and the initiation of puberty.

The progressive stages of the disorder appear to reflect the natural progression of an epileptic encephalopathy. In stage 1, there are brief seizures; while by stage 3, uncontrolled seizures prevent the child from continuing in school. In stage 4, a high seizure burden contributes to regression, which together with ensuing poor nutrient intake leaves a severely disabled child by stage 5, when multiple factors impair functioning.

The clinical features in Ugandan children are as severe as in South Sudanese children, but may be more severe than in Tanzanian patients. Despite similar age of onset and duration of symptoms, only 18% of Ugandan children had the milder nodding only variant, compared with 45% in Tanzania; all our patients had abnormal interictal background EEG compared with 60% in Tanzania. Ugandan children also had more severe cognitive impairment and a much greater burden and variety of seizures. These differences may suggest a variation in the presentation of the disease by region. Family clustering in all three countries, however, suggests a common exposure factor.

Aetiology and pathogenesis

The aetiology of nodding syndrome and pathogenesis of the complications remain unknown. A variety of viral central nervous system infections have been screened for on PCR, but no association has been demonstrated. An association with infestation with *O. volvulus* has been reported in some series, but has not been evident in other studies. Other aetiological considerations have included toxic brain injury, inflammatory brain disease, a slow virus infection or prion disease, an atypical mitochondrial disease or other genetic disorders. Repeated severe psychological trauma has also been proposed as a mechanism. Earlier studies by the Ugandan Ministry of Health and the US Centers of Disease Control found a higher proportion of cases with low serum vitamin B6 compared with controls. Although unlikely, the possibility was raised that nodding syndrome could be an atypical form of pyridoxine dependent epilepsy. Studies to explore this hypothesis further are awaited.

Children with this syndrome show some features reminiscent of prion protein disease, including motor and cognitive dysfunction, epilepsy, behaviour and psychiatric morbidity. However, other features commonly seen in prion disease—such as extrapyramidal involvement and cortical blindness—have not been reported. The age of onset of symptoms is also much younger and progression is much slower. The brain imaging and EEG findings too are not suggestive of prion disease. Nevertheless, brain biopsy or autopsy studies would be important in excluding prion disease.

The brain MRI findings might suggest viral encephalitis, a para-infectious phenomenon (such as an antibody-mediated channelopathy) or even a neurotoxin as possible aetiologies. A genetic disorder or metabolic disease is also possible. Future studies should also consider the recently described autoimmune encephalitides. Among Tanzanian patients, hippocampal sclerosis, probably from inflammation, was described in some patients. Although this could partly explain the cognitive difficulties, we did not observe such lesions, even in the subgroup with more severe cognitive impairment. We consider that an epileptic encephalopathy is more likely. The background EEG in all 22 cases showed a generalised excess of θ and slow activity. Prolonged EEG recording (including during sleep) with detailed neuropsychological testing and functional brain imaging may help in understanding the pathogenesis of cognitive decline.

Study limitations

Our patients may not be a representative sample, as they were not drawn randomly from the community. The study did not investigate aetiology and the proposed staging was derived primarily from parental descriptions rather than prospective observations and so may be influenced by recall bias. We only performed routine diagnostic EEGs, rather than prolonged recordings which are important in investigating associations between seizures and cognitive dysfunction. The resolution of our MRI is also quite low.

CONCLUSIONS

Nodding syndrome is a neurological disorder that is complicated by multiple physical and functional disabilities and may be considered as symptomatic generalised epilepsy. Assessment and care should be provided by a multidisciplinary team. Studies of aetiology, pathogenesis, evidence-based treatment and rehabilitation strategies are urgently needed.

Author affiliations

1Department of Paediatrics, Mulago Hospital/Makerere University College of Health Sciences, Kampala, Uganda
2Nuffield Department of Medicine, Centre for Tropical Medicine, Oxford University, Oxford, UK
3Department of Psychiatry, Mulago Hospital/Makerere University College of Health Sciences, Kampala, Uganda
4Department of Internal Medicine, Mulago Hospital/Makerere University College of Health Sciences, Kampala, Uganda
5Department of Clinical Neurophysiology, Great Ormond Street Hospital for Children, London, UK
6Department of Neuroradiology, Great Ormond Street Hospital for Children, London, UK
7Department of Community Health and Family Medicine, Mulago Hospital/Makerere University College of Health Sciences, Kampala, Uganda
8Ministry of Health Head Quarters, Kampala, Uganda

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### References

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