Incidence, causes and phenotypes of acute seizures in Kenyan children post the malaria-decline period

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Abstract

Background: Acute seizures are a common cause of paediatric admissions to hospitals in Africa, and malaria is an important cause of seizures in endemic areas. Malaria has declined in the past decade whilst neonatal admissions have increased, both which may affect the incidence and phenotypes of acute seizures in African children.

Methods: We examined the effect of recent decline in malaria and the increasing burden of neonatal admissions on the incidence, causes and phenotypes of acute seizures admitted to hospital from 2009–2013. We used logistic regression to measure associations and Poisson regression to calculate the incidence and rate ratios.

Results: The overall incidence of acute seizures over the 5-year period was 312 per 100,000/year (95 % CI, 295–329); 116 per 100,000/year (95 % CI, 106–127) for complex seizures and 443 per 100,000 live births (95 % CI, 383–512) for neonatal seizures. Over the period, there was an increase in incidence of seizures-attributable to malaria (SAM) (incidence rate ratio (IRR) = 1.25; p < 0.001), but neither non-SAM (IRR = 1.03; p = 0.569) nor neonatal seizures (IRR = 0.99; p = 0.905). Important causes of acute seizures were malaria (33 %) and respiratory tract infections (19 %); and for neonatal seizures were neonatal sepsis (51 %), hypoglycemia (41 %) and hypoxic-ischemic encephalopathy (21 %). Mortality occurred in 6 % of all acute seizures, being more common in complex seizures (8 %) and neonatal seizures (10 %) than other seizures (p < 0.001 for both comparisons).

Conclusions: Acute seizures remain common in children despite a decline in the incidence of malaria; suggesting that causes for these seizures need to be prevented in the community.

Keywords: Acute seizures, Causes, Falciparum malaria, Incidence, Phenotypes

Background

Acute seizures are common in children admitted to African hospitals [1] and malaria is the most important cause of seizures in endemic areas [2]. The proportion of complex seizures (repetitive, prolonged and/or focal) in malaria-endemic area is twice that reported in other settings, where seizures with fever have been studied [3, 4]. In endemic areas, seizures in children are associated with acute febrile infections such as malaria, respiratory tract infections and gastroenteritis [1].

The incidence of acute seizures in Kilifi, Kenya (a malaria endemic area), before the decline in malaria was high (425-650/100,000/year) [1, 2], and most complex seizure phenotypes were attributable to malaria [2]. We hypothesised that the incidence of acute seizures and the proportion with complex phenotypes depend on changes in transmission intensity of malaria. From early 2000–2008, admissions with malaria and seizures declined [2, 5], and so was the associated mortality [6]. In addition, neonatal admissions have increased, suggesting emergence of other causes of seizures during the neonatal period or improvement in hospital access in rural hospitals [7]. These changes may affect the incidence, causes and phenotypes of seizures in children admitted to hospital.

We examined the effect of the recent widely documented decline in malaria and the increasing burden of
Methods

Study site and population

This study was conducted in Kilifi Health and Demographic Surveillance System which is situated on the Kenyan coast about 50 kilometres from Mombasa and has a population of about 270,000. The population is comprised of a Mijikenda speaking community, who are mainly subsistence farmers and a few fishermen. The transmission intensity of malaria has reduced this area, resulting in reduction in morbidity and mortality due to malaria [5], in particular seizures-attributable to malaria (SAM) [2].

Admissions and management

A total sample of 3074 children aged 13 years and below with a history of seizure(s) during that period of the presenting illness were admitted to Kilifi County Hospital (KCH) from the year 2009–2013. Children with severe malaria were treated with first-line intravascular artesunate [8], with broad-spectrum antibiotics. Children admitted with seizures lasting 5 minutes were first treated with intravenous diazepam, and for those whose seizures did not stop within 10 minutes a second dose of diazepam or intramuscular paraldehyde was given [9]. Seizures lasting 15–20 minutes were managed with phenobarbital, and those continuing further with phenytoin. Seizures refractory to phenytoin were managed in consultation with senior clinicians, who would often recommend another dose of phenobarbital or midazolam since artificial ventilation was not available.

Review of case notes and data extraction

Clinical files for these children were reviewed to determine which admissions had acute seizures. Data for 1063 records was obtained from reviewing the patient’s case notes, for the remaining 2011 records data was obtained from the hospital inpatient database. Case notes were independently reviewed by Serem KG (who examined all case notes) and Kariuki SM (who randomly reviewed half of the case notes); with any differences in phenotypes resolved through consensus. The data extracted includes among others: the age in months of the patient, diagnosis, seizure type, medications received and blood workups. Data were extracted into Excel spread sheet, before being transferred into the analysis software.

Definitions of terms

Phenotypes of seizures are defined in Table 1, and were based on ILAE recommendations [10–12], that have been validated in previous studies in this area [13, 14]. Neonates were defined as those aged 28 days or younger [7]. Malnutrition was defined as a weight for age z-scores (WAZ) < −2 or mid-upper arm circumference (MUAC) <11 centimetres or a clinical diagnosis on discharge. Meningitis was defined as cerebrospinal white cell count >50/μL [15]. Anaemia defined as a haemoglobin concentration <50 g/L, thrombocytopenia as a platelet count <150,000/μL, hypoglycemia as glucose concentration of <3 Mmol/L, hyponatremia as sodium concentration of <125 Mmol/L, and hyperkalemia as potassium concentration of >4.5 Mmol/L.

Statistical analysis

The data were analysed using the STATA 13 statistical software (Stata Corporation, Texas, USA). The association between complex seizures and social demographic characteristics and other clinical variables was measured using logistic regression, and variables with a p-value <0.25 were used to build a multivariate model in a stepwise regression technique. Hosmer-Lemeshow test was used to test the goodness of fit in logistic regression model. For comparison of continuous variables, the Student t-test was used and if data did not have a Gaussian distribution, Mann–Whitney U-test was used. For comparison of categorical variables, the Pearson’s Chi-square test or Fisher’s exact (where observations were infrequent) were used. Proximate causes of acute seizures mentioned in the literature were reported here as percentages of occurrence among admissions.

For different phenotypes of acute seizures, we computed overall incidence, then stratified incidence by year, age group and falciparum malaria status. Incidence was computed by dividing number of children with acute seizures by person-years of observation for all admissions and by live births for neonatal admissions. The trend of incidence of acute seizures was determined using the Poisson regression technique. The trend for all admissions was repeated for neonatal seizures, non-malarial seizures, malarial seizures, and malarial seizures with a parasite density ≥2500 μ/L, a sensitive and specific cutoff for defining seizures attributable to malaria that was modelled with logistic regression techniques in previous studies [2, 16].

Ethics, consent and permissions

This study was approved by the Kenyan Ethical Review Committee (SSC No 2599) and parents of children participating in the study gave a written informed consent.

Data availability

These data are stored securely within the KEMRI-Wellcome Research Programme servers as was agreed with participants during consenting and can be availed upon formal request.
Results

General description
Over the 5 year period, 2742 children with acute seizures were admitted to KCH, of whom 1567 (57 %) were males (Table 2) and 182 (7 %) had neonatal seizures. Nearly half (46 %) of children were admitted with complex seizures, and different phenotypes of complex seizures showed significant overlap with each other (Fig. 1). Malaria parasitemia was detected in 38 % of all admissions and in 36 % of complex seizures (36 % of prolonged seizures, 39 % of convulsive status epilepticus, 31 % of repetitive seizures and 22 % of focal seizures). Malnutrition was present in 23 % of all admissions, and was more frequent in children with complex seizures (27 % vs. 21 %, p < 0.001), but less frequent in those with malaria parasitemia (16 % vs. 27 %, p < 0.001). Complex seizures were independently associated with mean corpuscular volume in a multivariable logistic regression model (odds ratio = 0.92 (95 % CI, 0.85-0.99); p = 0.033).

Possible causes of acute seizures
The most common causes of the 2742 admissions with acute seizures were malaria (n = 899, 33 %), respiratory tract infection (n = 532, 19 %), fever of unknown origin (n = 467, 17 %) and hypoglycemia (n = 274, 10 %). Other less common causes of acute seizures were as shown in Table 3. Malaria was an important co-morbidity in those with malnutrition (23 %).

A number of several possible causes were significantly more frequent in complex seizures compared to simple seizures: malaria (p < 0.001), respiratory tract infections (p < 0.001), fever of unknown origin (p < 0.001), encephalopathy of undetermined cause (p < 0.001), neonatal sepsis (p < 0.001), hypoglycemia (p = 0.006) and hyponatremia (p = 0.003). On the other hand, gastroenteritis was more common in simple than complex seizures (p = 0.025). Hypoglycemia was significantly more common in those with neonatal seizures compared to those without (54/133 (41 %) vs. 153/1881 (8 %); p < 0.001) and as was sepsis (95/182 (52 %) vs. 16/2565 (1 %); p < 0.001) and hypoxic-ischemic encephalopathy (39/182 (21 %) vs. 0/2565 (0 %); p < 0.001).

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Table 1 Definition of phenotypes of acute seizures

| Phenotypes of seizures | Definition |
|------------------------|------------|
| Acute seizures         | These are seizures that are associated with an acute event such as a febrile illness, acute haemorrhagic stroke, poisoning or electrolyte imbalance. |
| Acute symptomatic seizures | Acute seizures associated with neurological causes e.g. seizures due to malaria or bacterial meningitis. |
| Febrile seizures according to ILAE | Seizures associated with fever (temperature ≥38.5 °C) that occur in a sick child aged between a month to 6 years old, and the cause of fever does not involve the brain. |
| Febrile seizures according NIH | Seizures with fever usually occurring in child aged between 3 months to 6 years in absence of neurological involvement. |
| Overall definition of febrile seizures | Seizures with fever (temperature ≥38.5 °C) that occur in 2-8 % of children aged between 1 month to 6 years and exclude acute symptomatic seizures. |
| Definite prolonged seizures | Seizures that last up to ≥10 minutes, and timing is witnessed by a clinician or nurse. |
| Probable prolonged seizures | Seizures which did not stop after first dose of diazepam or continuing seizures that required second/third line antiepileptic drugs (phenobarbital/phenytoin). |
| Possible prolonged seizures | These are prolonged seizures based on parental history i.e. history of a seizures lasting ≥10 minutes, based on parents or caretakers reported. |
| Overall definition of status epilepticus | Seizures that last ≥30 minutes accompanied by loss of consciousness or intermitted seizures over a period of ≥30 minutes without regaining consciousness in between the seizures. |
| Definite status epilepticus | Seizures fulfilling the definition of status epilepticus as above, but were witnessed/observed by a clinician or a nurse and the timing is well documented in the child’s clinical notes. |
| Probable status epilepticus | Prolonged or continuing seizures which required the administration of AEDs such as phenobarbital or phenytoin. |
| Possible status epilepticus | A parental/caretakers’ history of seizures lasting ≥30 min, when the child was not convulsing on admission. |
| Focal/partial seizures | Seizures localised to or involving one body part e.g. twitching of one side of the facial structures, cycling movements of one limb, Todd’s paresis of one body side and motor deficits involving one side of the body post-ictally. |
| Generalised seizures | Seizures involving the entire body parts. |
| Repetitive seizures | Two or more seizures in the current illness or within 24 hours. |
| Complex seizures | Seizures that have focal, repetitive and/or prolonged (including status epilepticus) features. |
### Table 2 Distribution of social demographic, laboratory and clinical characteristics among children with seizures

| Variable                      | Complex seizure (N = 1260) | Simple seizure (N = 1482) | Total (N = 2742) | Odds ratio (95 % CI) | p-value |
|-------------------------------|----------------------------|---------------------------|------------------|----------------------|---------|
| Age: median (interquartile range (IQR)) | 28.4 (12.2-51.9) | 27.9 (15.8-44.4) | 28.1 (14.5-47.5) | 1.00 (1.00-1.00) | 0.171   |
| Sex (male)                    | 711/1263 (56 %) | 856/1486 (58 %) | 1567/2749 (57 %) | 0.95 (0.82-1.10) | 0.489   |
| **Clinical signs and symptoms** |                            |                           |                  |                      |         |
| Median respiratory rate (IQR) | 40 (32–50) | 38 (30–47) | 38 (32–48) | 1.02 (1.01-1.02) | <0.001  |
| Median heart rate (IQR)       | 149 (128–166) | 152 (134–167) | 150 (132–167) | 1.00 (0.99-1.00) | 0.003   |
| WAZ×                           | –2 (–2 - -1) | –1 (–2 - 0) | –1 (–2 - -1) | 0.97 (0.92-1.03) | 0.379   |
| MUACb                          | 14.0 (13.0-1.05) | 14.5 (13.5-15.2) | 14.2 (13.0-15.0) | 0.92 (0.89-0.96) | <0.001  |
| Prostration                    | 513 (41 %) | 293 (20 %) | 806 (29 %) | 2.67 (2.27-3.13) | <0.001  |
| Median temperature °C (IQR)    | 38.0 (37.0-39.0) | 38.3 (37.4-39.2) | 38.2 (37.2-39.1) | 0.86 (0.81-0.91) | <0.001  |
| Temperature gradient           | 90/1261 (7 %) | 86/1470 (6 %) | 176/2731 (6 %) | 1.24 (0.91-1.68) | 0.173   |
| **Laboratory investigations**  |                            |                           |                  |                      |         |
| Median Hb levels (IQR)         | 10.0 (8.6-11.2) | 10.1 (9.1-11.1) | 10.2 (8.9-11.1) | 1.02 (0.98-1.05) | 0.379   |
| Median glucose levels (IQR)    | 4.8 (3.8-6.1) | 4.7 (3.9-5.8) | 4.7 (3.8-5.9) | 1.02 (0.99-1.06) | 0.169   |
| Median sodium levels (IQR)     | 134 (130–137) | 134 (132–137) | 134 (131–137) | 0.99 (0.96-1.00) | 0.016   |
| Median potassium levels (IQR)  | 3.9 (3.5-4.3) | 3.8 (3.4-4.2) | 3.8 (3.4-4.2) | 1.21 (1.06-1.39) | 0.005   |
| Median BCS (IQR)               | 4 (2–5) | 5 (4–5) | 5 (4–5) | 0.71 (0.69-0.76) | <0.001  |
| Median hematocrit (IQR)        | 31.3 (27.0-34.6) | 31.5 (28.5-34.1) | 31.4 (27.9-34.3) | 1.00 (0.99-1.02) | 0.375   |
| Median blood oxygen saturation % (IQR) | 98 (95–100) | 99 (97–100) | 98 (96–100) | 0.96 (0.95-0.97) | <0.001  |
| Median mean corpuscular volume (IQR) | 73.4 (66.0-81.0) | 72.0 (65.0-78.0) | 73.0 (65.0-79.0) | 1.02 (1.01-1.03) | <0.001  |
| Median platelets (IQR)         | 292.0 (146.5-421.0) | 314.0 (179.0-433.0) | 306.0 (165.0-427.0) | 1.00 (1.00-1.00) | 0.032   |
| Parasite density ≥2.5x10³/μL   | 318/1263 (25 %) | 344/1486 (23 %) | 622/2749 (39 %) | 1.12 (0.94-1.33) | 0.215   |
| Median parasite density (IQR)  | 2976 (0–165880) | 0 (0–50640) | 0 (0–101710) | 1.00 (1.00-1.00) | 0.001   |
| Slide positive for malaria     | 377/1060 (36 %) | 383/1332 (29 %) | 760/2392 (38 %) | 1.37 (1.15-1.63) | <0.001  |
| Creatinine                     | 45 (37–58) | 50 (38–64) | 47 (37–60) | 1.00 (1.00-1.00) | 0.382   |

*aWAZ = weight for age; bMUAC = mid-upper arm circumference BCS = Blantyre coma score; CI = confidence interval; IQR = interquartile range

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**Fig. 1** Overlap of phenotypes of seizures over the study period (N = 1263). There was substantial overlap of the three phenotypes of complex seizures namely prolonged, repetitive and/or focal seizures.
per 100,000/year (95 % CI, 157–182) in males. The overall incidence of acute seizures significantly increased with years of admission over the study period (incidence rate ratio (IRR) = 1.13; \( p < 0.001 \)) (Fig. 2). Similarly, the overall incidence over the study period appeared to decrease with age (IRR = 0.651; \( p < 0.001 \)) (Fig. 3). The incidence of all acute seizures appeared to decrease with distance from the hospital and was 257 per 100,000/year (95 % CI, 242–272) for those living within 5 kilometres from KCH, and 55 per 100,000/year (95 % CI, 48–62) for those living ≥5 kilometres from the hospital. The incidence of all seizures attributable to malaria (SAM) was 75 per 100,000/year (95 % CI, 67–83), and that for non-SAM was 237 per 100,000/year (95 % CI, 223–252). The incidence of SAM increased over the period (IRR = 1.22 (95 % CI, 1.13-1.31); \( p < 0.001 \)) while that for non-SAM remained unchanged (IRR = 1.03 (95 % CI, 0.96-1.06); \( p = 0.569 \)).

**Incidence of seizures-attributable to malaria**

The incidence of all seizures attributable to malaria (SAM) was 75 per 100,000/year (95 % CI, 67–83), and that for non-SAM was 237 per 100,000/year (95 % CI, 223–252). The incidence of SAM increased over the period (IRR = 1.22 (95 % CI, 1.13-1.31); \( p < 0.001 \)) while that for non-SAM remained unchanged (IRR = 1.03 (95 % CI, 0.96-1.06); \( p = 0.569 \)).

**Mortality in admissions with acute seizures**

The overall case fatality proportion over the 5-year period was 6 % and remained stable throughout the study (\( p = 0.166 \)). Death was more frequent in complex seizures than simple seizures (8 % vs.4 %; \( p < 0.001 \)) and in neonatal seizures than seizures in those aged >28 days (10 % vs. 5 %; \( p = 0.005 \)). In particular, 12 % of the deaths were in prolonged seizures and convulsive seizures (95 % CI, 19–28) for status epilepticus, 254 per 100,000/year (95 % CI, 240–270) for receptive seizures and 34 per 100,000/year (95 % CI, 29–40) for focal seizures.

### Table 3 Possible causes of acute seizures

| Possible cause                                         | Complex seizure \( n = 1263 \) (%) | Simple seizure \( n = 1486 \) (%) | Overall \( n = 2742 \) (%) | \( p \)-value |
|-------------------------------------------------------|------------------------------------|------------------------------------|--------------------------|--------------|
| Malaria                                               | 477 (38 %)                         | 422 (28 %)                         | 899 (33 %)               | <0.001       |
| Respiratory tract infections                          | 180 (14 %)                         | 352 (24 %)                         | 532 (19 %)               | <0.001       |
| Fever of unknown origin                               | 161 (13 %)                         | 306 (21 %)                         | 467 (17 %)               | <0.001       |
| Hypoglycemia                                          | 104/832 (13 %)                     | 103/1184 (9)                       | 207/2016 (10 %)          | 0.006        |
| Gastroenteritis                                       | 56 (4 %)                           | 95 (6 %)                           | 151 (5 %)                | 0.025        |
| Hyponatremia (<125Mmol/L)                             | 50/759 (7 %)                       | 31 (3 %)                           | 81 (5 %)                 | 0.003        |
| Meningitis                                            | 69 (5 %)                           | 58 (4 %)                           | 127 (5 %)                | 0.052        |
| Neonatal sepsis                                       | 72 (6 %)                           | 39 (3 %)                           | 111 (4 %)                | <0.001       |
| Unknown encephalopathy                                | 73 (6)                             | 31 (2 %)                           | 104 (4 %)                | <0.001       |
| Low hemoglobin (<5 g/L)                               | 35/1047 (3 %)                      | 34/1338 (3 %)                      | 69/2385 (3 %)            | 0.246        |
| Urinary tract infection                               | 16 (1 %)                           | 25 (2)                             | 41 (1 %)                 | 0.370        |
| Hypoxic-ischemic encephalopathy (birth asphyxia)      | 30 (2 %)                           | 9 (1 %)                            | 39 (1 %)                 | <0.001       |
| Trauma                                                | 16 (1 %)                           | 18 (1 %)                           | 34 (1 %)                 | 0.896        |
| Skin disease                                          | 11 (1 %)                           | 19 (1 %)                           | 30 (1 %)                 | 0.305        |
| Septicemia                                            | 12 (1 %)                           | 15 (1 %)                           | 27 (0 %)                 | 0.875        |
| Anemia                                                | 13 (1 %)                           | 5 (<1 %)                           | 18 (1 %)                 | 0.025        |
| Sickle cell disease                                   | 8 (1 %)                            | 7 (<1 %)                           | 15 (1 %)                 | 0.565        |
| Undetermined                                          | 5 (<1 %)                           | 5 (<1 %)                           | 10 (<1 %)                | 0.797        |
| Ear infection                                         | 2 (<1 %)                           | 5 (<1 %)                           | 7 (<1 %)                 | 0.356        |
| Immunosuppression (HIV exposed)                       | 2 (<1 %)                           | 4 (<1 %)                           | 6 (<1 %)                 | 0.535        |
| Eye infections                                        | 1 (<1 %)                           | 4 (<1 %)                           | 5 (<1 %)                 | 0.244        |
| Poisoning                                             | 2 (<1 %)                           | 0 (0 %)                            | 2 (<1 %)                 | 0.125        |
| Prematurity                                           | 2 (<1 %)                           | 0 (0 %)                            | 2 (<1 %)                 | 0.150        |
| Other causes                                          | 12 (0.95 %)                        | 15 (1.01 %)                        | 27 (0.98 %)              | 0.875        |
| Outcome: died                                         | 104 (8 %)                          | 54 (4 %)                           | 158 (6 %)                | <0.001       |
epilepticus, 6% were in repetitive seizures and 8% in focal seizures. Mortality following acute seizures was reported in 20% with malaria, 11% with respiratory tract infections, 23% with malnutrition and 12% of encephalopathy with undetermined causes.

Discussion

This study examined the effect of recent decline in malaria and increasing burden of neonatal admissions on the incidence of acute seizures, their proximate causes and phenotypes in children admitted to a rural hospital in Kenya. The burden of acute seizures in children remains high despite a reduction of malaria in the past decade. Indeed the overall incidence appeared to increase over the study period (2009–2013), as did the incidence of seizures attributable to malaria. The incidence of complex seizure increased the most (107%). This suggests, in part, that causes of and risk factors for acute seizures are still prevalent in the community. Over 80% of all incidence admissions resided less than 5 km from the KDH, implying hospital accessibility determines the admissions of acute seizures in this area. The most common causes of admissions with acute seizures were malaria and respiratory tract infections, but hypoglycemia, hypoxic-ischemic encephalopathy and neonatal sepsis were important among the neonates.

Changes in incidence of acute seizures

The incidence of acute seizures in this study (312 per 100,000/year) is lower than that reported in a previous study (425 per 100,000/year) [1], which used data on admissions during a time when malaria was a common cause of morbidity and mortality in this area. Despite the recent decline in malaria, our present data reveals that the incidence has been increasing between 2009 and 2013, an important epidemiological finding which is not apparent if the present incidence was compared with earlier years when incidence of malaria was high.

This increase in incidence was particularly observed for SAM, being highest for the complex seizure phenotypes; supporting increasing burden for acute seizures post the malaria decline era. There are three explanations for the increasing incidence of seizures following the earlier decline. First and most plausible, a reduction in malaria may have resulted in less exposure, thereby reducing the acquisition of naturally acquired immunity to malaria, resulting in more severe illness, including
seizures, that requires hospitalisation [5]. Secondly, this
could be related to proportional increase in neonatal
diseases such as hypoglycemia, sepsis and-
hypoxic-ischemic encephalopathy compared to an earlier
study [1]. Finally, the increasing incidence of acute sei-
zures could be related to possible improvements in health
care services not measured in this study, which would en-
courage admissions.

Incidence of acute seizures
The incidence is grossly underestimated since only 28 %
of it was from areas >5 kilometres. This suggests that
poor infrastructure and transport costs may have resulted in
some (if not most) seizures not being treated in hos-
pital. In a recent study, only 56 % of persons with convul-
sive status epilepticus occurring with epilepsy were treated
in KCH [17]. Other cultural reasons such as preference of
traditional healers over biomedical practitioners may have
contributed [18]. Despite the conservative incidence our
estimates for prolonged and/or status epilepticus is higher
than that reported in developed countries e.g. in London
[19]. This may be explained by the high prevalence of in-
fected causes in this area e.g. malaria parasitemia, which
was found in 36 % of all prolonged seizures; over 90 % of
complex seizures in parasitemic children are caused by
malaria [2]. Delays in management of acute seizures in
peripheral clinics that may result in admission of seizures
already refractory to treatment [14], adding to the high
burden of prolonged seizures in this study.

Causes of acute seizures
Malaria remains the most important cause of admission
with seizures, with malaria parasitemia observed in over a
third of admissions. This proportion however is slightly
less than that reported in previous studies in this area
(>50 %) [1, 14], suggesting a possibility of emergence of other
important causes of acute seizures particularly those
common in the neonatal period. Falciparum malaria may
cause seizures by: (i) sequestration-induced brain dam-
age that manifests as seizures [20]; (ii) down-regulation
of GABA receptors thereby increasing susceptibility to
seizures [21]; (iii) metabolic complications such as
hyponatremia or hypoglycaemia and; (iv) induction of
inflammatory molecules that lower seizure threshold
[22]. Susceptibility to seizures may be determined by genetic polymorphisms [13].

Hypoglycemia, unknown encephalopathy and neonatal
sepsis, were more common in the neonates in greater pro-
portions than in a previous study [1], yet some of these
can be easily prevented and managed. It is however, un-
clear if a proportion of encephalopathy of unknown origin
reflects neurotropic viruses that were not investigated in
this study [23].

Malaria was more common in complex seizures com-
pared to simple seizures, suggesting a causal role for these complex phenotypes, as demonstrated in previous
studies [2]. Additionally, other causes of seizures were
significantly more frequent in complex seizures, perhaps
because it represents a severe phenotype of seizures, as
supported by its association with increased mortality in
this study and neurocognitive impairments in previous
studies [24, 25]. However, gastroenteritis appeared to be
more frequent in simple seizures compared to complex
seizures as would be expected.

Changing phenotypes of acute seizures
Nearly half (46 %) of the admissions were complex sei-
zures, a significant reduction from 70 % reported in pre-
vious studies in the same population [4, 13]. Since this
phenotype is particularly attributable to malaria, the low
frequency may be related to lower incidence of malaria
during the study period compared to the early 2000s [5].
Complex seizures were associated with mean corpuscular
volume, which may be a marker of: (i) iron deficiency; (ii)
red blood cell redistribution following sequestration into
the deep capillary bends of the brain [26]; or (iii) presence
of alpha thalassaemia [27].

Mortality in acute seizures
The mortality from acute seizures was greater than in
previous studies (6 % vs. 3 %) [1], probably suggesting a
change in aetiology related to increase in neonatal
seizures, in which mortality was highest (10 %). Add-
itionally, case fatality was common (12 %) of those with
encephalopathy of unknown origin which could be im-
portant in the neonatal period. Future studies should
screen for viral aetiology of acute seizures, which may be
associated with mortality but were not assessed in this
study [28].

In terms of causes of acute seizures, mortality was high-
est in malaria (20 %) and malnutrition (20 %). Control of
malaria and prevention of malnutrition would reduce just
under half of mortalities associated with acute seizures.

Strengths and limitations
These data were prospectively collected over a long time
period with clinical and laboratory information docu-
mented in standardised proformas, thus findings are ro-
bust and reliable. Careful classification of acute seizures
allowed objective relation of these phenotypes of seizures
with causes, incidence and outcomes. These data may not
represent the burden and situation of acute seizures in the
community, since some seizures in this area are not
treated at hospital. It was not possible to perform electro-
encephalography, owing to logistical constraints.
Conclusions
The incidence of acute seizures is high in this area, and appears to be increasing during the period particularly for SAM, despite the reduction in the incidence of malaria. Malaria remains a common cause of seizures although in lesser proportions than previously documented while other etiologies in the neonatal period are emerging as important causes of seizures. The hospital-based incidence disproportionately represents only those living within a 5 kilometre radius and thus a true burden should be provided through community-based studies. Complex seizures remain a common phenotype, and should be addressed owing to their association with mortality. Mortality is still high particularly in neonatal seizures calling for plans to effectively prevent and manage seizures as well as control their associated causes and risk factors in the community.

Abbreviations
CI: Confidence interval; ILAE: International League Against Epilepsy; IQR: Interquartile range; IRR: Incidence rate ratio; KCH: Kilifi County Hospital; SAM: Seizures-attributable to malaria; MUAC: Mid-upper arm circumference; WAZ: Weight-for-age z-scores.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
All authors made substantial contribution to the preparation of this manuscript, including conception and study design (CN and SK), acquisition of data (CN), data extraction and statistical analysis (GS, CN and SK) and writing the first draft (GS and SK). All authors reviewed the subsequent drafts of the manuscript for important intellectual content. All authors read and approved the final manuscript for publication.

Authors’ information
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