Vitamin D deficiency causes rickets in an urban informal settlement in Kenya and is associated with malnutrition

Kelsey D. J. Jones1,2 | C. Ulrich Hachmeister† | Maureen Khasira1 | Lorna Cox4 | Inez Schoenmakers4,5 | Caroline Munyi3 | H. Samira Nassir3 | Barbara Hünten-Kirsch3 | Ann Prentice4,6 | James A. Berkley1,7

1 KEMRI-Wellcome Trust Research Programme, Kenya
2 Section of Paediatrics and Centre for Global Health Research, Imperial College, London, UK
3 Baraka Health Centre, German Doctors Nairobi, Nairobi, Kenya
4 MRC Human Nutrition Research, Elsie Widdowson Laboratory, Cambridge, UK
5 Department of Medicine, Faculty of Medicine and Health Science, University of East Anglia, Norwich, UK
6 MRC Keneba, The Gambia
7 Nuffield Department of Clinical Medicine, University of Oxford, Oxford, UK

Abstract

The commonest cause of rickets worldwide is vitamin D deficiency, but studies from sub-Saharan Africa describe an endemic vitamin D-independent form that responds to dietary calcium enrichment. The extent to which calcium-deficiency rickets is the dominant form across sub-Saharan Africa and in other low-latitude areas is unknown. We aimed to characterise the clinical and biochemical features of young children with rickets in a densely populated urban informal settlement in Kenya. Because malnutrition may mask the clinical features of rickets, we also looked for biochemical indices of risk in children with varying degrees of acute malnutrition. Twenty one children with rickets, aged 3 to 24 months, were identified on the basis of clinical and radiologic features, along with 22 community controls, and 41 children with either severe or moderate acute malnutrition. Most children with rickets had wrist widening (100%) and rachitic rosary (90%), as opposed to lower limb features (19%). Developmental delay (52%), acute malnutrition (71%), and stunting (62%) were common. Compared to controls, there were no differences in calcium intake, but most (71%) had serum 25-hydroxyvitamin D levels below 30 nmol/L. These results suggest that rickets in young children in urban Kenya is usually driven by vitamin D deficiency, and vitamin D supplementation is likely to be required for full recovery. Wasting was associated with lower calcium (p = .001), phosphate (p < .001), 25-hydroxyvitamin D (p = .049), and 1,25-dihydroxyvitamin D (p = 0.022) levels, the clinical significance of which remain unclear.

KEYWORDS
acute malnutrition, Africa, poverty, rickets, urbanization, vitamin D

1 INTRODUCTION

Rickets is a disease of childhood that occurs due to failure of regulated chondrocyte apoptosis and deficient mineralisation at the growth plates of long bones (Pettifor, 2003; Sabbagh et al., 2005). Children develop a wide variety of skeletal abnormalities that vary according to age and developmental stage, including wrist widening, bowed legs, craniotabes with frontal bossing and delayed fontanelle closure, costochondral swelling, and spinal deformity (Elder & Bishop, 2014). Impaired linear growth and developmental delay are common. Younger infants may present with symptomatic hypocalcaemia (Ladhani et al., 2004; Hogler, 2015).

Worldwide, the commonest cause of rickets is vitamin D deficiency, which leads to secondary hyperparathyroidism and progressive renal phosphate loss (Glorieux & Pettifor, 2014). Vitamin D can be synthesised in the skin in a process that is dependent on sunlight exposure, or derived from a relatively limited range of dietary sources, including egg yolk and oily fish. Deficiency is usually the result of inadequate sun exposure, which may occur due to living at mid- or
high-latitude. In tropical or subtropical climes, there is generally abundant exposure to sunlight, but vitamin D deficiency may arise in association with risk factors such as darker skin pigmentation, atmospheric pollution, and covering skin for religious or cultural reasons (Baroncelli et al., 2008; Elder & Bishop, 2014; Trilok Kumar et al., 2015). However, in such settings, vitamin D deficiency may not be the only cause of rickets. Studies from several sub-Saharan African countries have implicated chronic dietary calcium deficiency in the pathogenesis of rickets and demonstrated nutritional cure with calcium supplementation alone (Thacher et al., 1999; Pettifor, 2004; Prentice et al., 2008; Braithwaite et al., 2016). The aetiology may be multifactorial, because calcium deficiency tends to exacerbate the impact of marginal vitamin D status, and may further interact with other nutritional deficiencies, such as of iron (Braithwaite et al., 2012; Pettifor, 2014).

A recent report highlighted the existence of a significant burden of rickets amongst children living in an informal settlement in Nairobi, Kenya (Edwards et al., 2014). Most children with clinically-defined rickets were younger than 2 years, and rickets was associated with acute malnutrition and developmental delay (Edwards et al., 2014). In this paper, we describe a similar group of children presenting to a primary care facility in Mathare, another of Nairobi’s informal settlements. Via detailed dietary and biochemical evaluation, we provide evidence that deficiencies in both calcium and vitamin D are likely to play a role in the aetiology of rickets in urban Kenyan settings. Because malnutrition may mask the clinical features of rickets, we also looked for biochemical features indicating rickets risk in children with varying degrees of acute malnutrition. To this end, we demonstrate an association between subclinical vitamin D deficiency and severe wasting.

2 | METHODS

2.1 | Study design

The study was designed primarily as a case–control study comparing children with rickets (cases) to children without rickets or acute malnutrition (controls). Additionally, children with severe and moderate acute malnutrition (SAM and MAM) were compared to controls.

2.2 | Setting

The study took place between July 2013 and April 2014 at the Baraka Health Centre (BHC), Mathare, Nairobi. BHC is supported and administered by the nongovernmental organisation “German Doctors Nairobi” and provides free healthcare to children aged less than 5 years (http://www.germandoctorsnairobi.co.ke/). Mathare is the second largest urban informal settlement in Kenya. It is home to at least 200,000 people, who mainly live in overcrowded iron-sheets housing with limited access to safe water and sanitation facilities (Muungano Support Trust, Slum Dwellers International, University of Nairobi, University of California, Berkley, 2012). Most residents engage in casual work or run small businesses and tend to work long hours. Provision for childcare is limited and represents a significant household expenditure for many families (Muungano Support Trust, Slum Dwellers International, University of Nairobi, University of

California, Berkley, 2012). Breastfeeding is almost universal and persists beyond 12 months in the majority of cases, although provision of weaning and complementary foods occurs much earlier than the World Health Organization-recommended 6 months (Kimani-Murage et al., 2011).

Nairobi is at latitude 1.3°S. It has a subtropical highland climate with two rainy seasons (March–May and October–December). The mean number of hours of sunshine per day peaks in February at 9.5 h/day, dropping to 4.1 hr/day in August (average 6.8 hr/day; World Meterological Organization, 2010). For comparison, The Gambia, where calcium dependent rickets is prevalent, averages 8.0 hr/day (Prentice et al., 2008).

2.3 | Participants

Children with rickets were recruited from among those presenting to BHC. Inclusion criteria for cases were age 3 to 24 months, with clinically diagnosed rickets, based on the presence of one or more of wrist widening, rachitic rosary, swollen knees, bow legs, or bone pain on walking (Thacher et al., 2002). Where wrist widening was the only feature, clinical evaluation was supplemented with radiographic assessment (right wrist X-ray), and children were only included in the presence of clear radiologic features consistent with rickets. Controls without clinical features of rickets or acute malnutrition and a further group of children without clinical features of rickets with SAM and MAM were recruited from among those self-presenting to BHC or via a program of active case finding in the community by local community health volunteers. SAM was defined on the basis of mid-upper arm circumference (MUAC) less than 115 mm or the presence of bilateral lower limb oedema (kwashiorkor), MAM was defined on the basis of MUAC 115 mm to 124 mm, and nonmalnourished controls had MUAC greater than 124 mm.

Children were excluded (from all of the groups) if they required emergency medical care, if they were being treated for tuberculosis, if they had a fracture within the preceding 3 months, and in the absence of informed parental consent for participation. Children were also excluded if they were known or found to have HIV infection, or if they were younger than 12 months old and known to be HIV-exposed.

2.4 | Study procedures

Participants were clinically assessed by a senior clinician (KDJJ or CUH). Demographic characteristics were recorded including anthropometric
analysed for 25 at MRC Human Nutrition Research, Cambridge, UK. There, serum was adjusted for serum albumin throughout, using the formula Calciumadjusted = Calciummeasured + 0.8 × (40 – albumin [g/L]), where the adjusted value may be considered a proxy for biologically active calcium (Association for Clinical Biochemistry & Laboratory Medicine, 2015). We used a 25(OH)D lower limit for bone health of 30 nmol/L, above which 25(OH)D-deficiency rickets is considered unlikely (Munns et al., 2016). Biochemical, demographic, and anthropometric variables for children in each of the rickets, SAM, and MAM groups were compared to controls: Mann–Whitney U or Fisher’s exact tests were used to test differences between groups. Outputs of linear regressions are presented adjusted for a priori potential confounders: age, CRP, or AGP, where these were associated with the dependent variable in univariate analyses. No formal sample size calculation was performed: patients presenting with rickets were recruited opportunistically over a 9-month period, and we planned to recruit a similar number of controls and participants with SAM and with MAM. Statistical correction for multiple comparisons was not performed.

2.6 | Ethics

All participants enrolled in the study had individual written informed consent provided by a parent or guardian. The study was approved by the Kenya Medical Research Institute’s Ethical Review Committee prior to initiation (2481). All children who presented to BHC or were identified through community screening with rickets, SAM, MAM, or intercurrent illnesses were provided with free medical care whether or not they were enrolled in the study.

3 | RESULTS

Eighty-four children were recruited to the study: 21 cases of rickets, 22 controls who were nonrachitic and nonacutely malnourished, 21 children with SAM (of whom nine had kwashiorkor), and 20 with MAM. Monthly recruitment of children with rickets was similar throughout the study period.

3.1 | Clinical and biochemical phenotype of rickets

Most of the children with rickets had wrist widening and rachitic rosary as opposed to lower limb features (Table 1). Developmental delay, acute malnutrition, and stunting were common (Table 1).

Children with rickets were relatively hypocalcaemic, hypophosphataemic, and had low 25(OH)D, compared to controls (Table 2).

### TABLE 1 | Clinical features of rickets

| Clinical features | n (%) |
|-------------------|-------|
| **Major features** |       |
| Wrist widening    | 21 (100) |
| Rachitic rosary   | 19 (90) |
| Swollen knees     | 4 (19) |
| Bowed legs        | 3 (14) |
| Bone pain on walking | 1 (5) |
| **Minor features** |       |
| Open Fontanelle   | 19 (90) |
| Double malleoli   | 11 (52) |
| Harrison’s groove | 10 (48) |
| Lower arm bending | 3 (14) |
| Developmental delay | 11 (52) |
| **Comorbidity**   |       |
| Fever             | 5 (24) |
| URTI              | 13 (62) |
| Pneumonia         | 2 (10) |
| Acute watery diarrhoea | 3 (14) |
| **Nutritional status** |     |
| SAM               | 4 (19) |
| MAM               | 11 (52) |
| Stunting          | 13 (62) |

Note. Major features are defined in line with Thacher et al. (2002). Developmental delay was based on parental report. Pneumonia and acute watery diarrhoea were diagnosed in line with WHO guidance (World Health Organization, 2005). SAM and MAM were diagnosed on the basis of MUAC and/or presence of oedema, stunting was diagnosed if height/length-for-age z-score was ≤−2. MAM = moderate acute malnutrition; SAM = severe acute malnutrition; URTI = upper respiratory tract infection.
**Calcium is adjusted for albumin as described in the methods section. **Omitted in the presence of oedema.

Amongst children with SAM, the nine with kwashiorkor had a different biochemical profile from those with severe wasting (MUAC <115 mm): kwashiorkor was associated with lower albumin and ALP, and with lower nutritional and biochemical features

|                  | Rickets (R) | Control (C) | p (R vs C) | SAM         | MAM         |
|------------------|------------|-------------|------------|-------------|-------------|
| n                | 21         | 22          | -          | 21          | 20          |
| Age (months)     | 11 (9 to 15)| 13 (10 to 18)| .34        | 14 (10 to 20)| 11 (9 to 14) |
| Male sex         | 12 (57)    | 14 (64)     | .76        | 9 (43)      | 7 (35)      |
| MUAC (mm)        | 119 (115 to 126) | 132 (130 to 141) | <.001      | 113 (110 to 126) | 120 (117 to 120) |
| Oedema           | 0 (0)      | 0 (0)       | 1.00       | 9 (43)      | 0 (0)       |
| WIZ              | -1.0 (-2.1 to -0.8) | -1.1 (-2.0 to 0.2) | .81        | -2.9 (-3.5 to -2.1) | -1.7 (-2.3 to -1.3) |
| LAZ              | -2.4 (-4.3 to -1.8) | -1.7 (-2.4 to -0.5) | .03        | -3.3 (-3.6 to -2.3) | -2.3 (-3.4 to -1.9) |
| WAZ              | -2.3 (-3.5 to -1.7) | -1.9 (-2.5 to -0.5) | .06        | -3.3 (-3.8 to -2.3) | -2.8 (-3.5 to -2.1) |
| OFCAZ            | -0.3 (-0.9 to 0.2) | -0.6 (-1.0 to 0.6) | .49        | -0.8 (-1.7 to -0.3) | -1.2 (-1.9 to -0.5) |

|                  | Calcium** (mmol/L) | Phosphate (mmol/L) | ALP (U/mL) | PTH (pg/mL) | 25(OH)D (nmol/L) | 1,25(OH)2D (pmol/L) | CRP (mg/mL) | AGP (g/L) | Calcium intake (mg/day) | Phosphate intake (mg/day) |
|------------------|--------------------|--------------------|------------|-------------|-----------------|---------------------|-------------|-----------|--------------------------|--------------------------|
|                  | 2.16 (2.04 to 2.34) | 0.95 (0.79 to 1.46) | 912 (396 to 1418) | 122 (44 to 249) | 19 (15 to 37) | 331 (213 to 446) | 2.7 (1.0 to 6.9) | 1.3 (0.7 to 1.6) | 225 (113 to 382) | 425 (259 to 622) |
|                  | 2.46 (2.37 to 2.56) | 1.92 (1.81 to 2.14) | 283 (240 to 319) | 26 (14 to 46) | 70 (54 to 85) | 316 (277 to 384) | 2.1 (1.0 to 5.6) | 1.3 (1.0 to 1.5) | 117 (63 to 226) | 266 (154 to 505) |
|                  | <.001              | <.001              | <.001      | <.001       | <.001           | <.001               | .63         | .84       | .13                      | .14                      |
|                  | 2.40 (2.19 to 2.55) | 1.57 (1.29 to 1.69) | 188 (175 to 262) | 27 (14 to 57) | 53 (32 to 82) | 248 (213 to 338) | 3.1 (1.0 to 13.5) | 1.9 (1.2 to 2.3) | 130 (78 to 245) | 323 (231 to 423) |
|                  | 2.38 (2.30 to 2.49) | 1.66 (1.49 to 2.02) | 211 (176 to 297) | 27 (20 to 35) | 63 (46 to 92) | 316 (253 to 388) | 4.9 (2.8 to 13.1) | 1.4 (1.2 to 1.8) | 192 (86 to 234) | 394 (228 to 589) |

Note. Values are median (interquartile range) or number (percentage) throughout. AGP = alpha-1 acid glycoprotein; ALP = alkaline phosphatase; CRP = C-reactive protein; LAZ = length-for-age; MAM = moderate acute malnutrition; MUAC = mid-upper arm circumference; OFCAZ = occipitofrontal circumference-for-age; PTH = parathyroid hormone; SAM = severe acute malnutrition; WLZ = Weight-length. **Calcium is adjusted for albumin as described in the methods section.**

FIGURE 1 Serum 25-hydroxyvitamin D (25(OH)D) levels in children with clinically defined rickets, or without rickets but with severe wasting (mid-upper arm circumference [MUAC]<115 mm), moderate wasting (MUAC 115 to 124 mm), kwashiorkor (Kwash), or community controls. Dotted lines are at 30 nmol/L (taken to indicate likely deficiency) and 50 nmol/L (30 to 50 nmol/L is considered insufficient) (Munns et al., 2016)

3.2 Relationship between anthropometric and biochemical variables among children without rickets

Amongst children with SAM, the nine with kwashiorkor had a different biochemical profile from those with severe wasting (MUAC <115 mm): kwashiorkor was associated with lower albumin and ALP, and with higher 25(OH)D and CRP (supplementary Figure 2). In order to assess relationships between biochemical variables and degree of wasting (WLZ and MUAC) and stunting (LAZ), we performed linear regression excluding those children with rickets. Degree of wasting (WLZ) and absolute MUAC were associated with lower serum adjusted calcium,
phosphate, 25(OH)D, and 1,25(OH)₂D levels (Table 3, Figure 3). Stunting (adjusted for age) was also associated with lower phosphate (Table 3). Thirty-three percent of severely wasted children (MUAC <115 mm) had a low 25(OH)D (<30 nmol/L) indicating an increased risk of vitamin D deficiency, and a further 25% was suggestive of insufficiency (30 to 50 nmol/L).

4 | DISCUSSION

Vitamin D deficiency appeared to play a key role in the pathogenesis of rickets in young children in an informal settlement in Nairobi. Children with rickets tended to present with wrist widening or rachitic rosary, and acute malnutrition and developmental delay were common comorbidities. The phenotype of rickets was clinically and biochemically distinct from that caused by calcium deficiency, which is the more prevalent form reported in several other sub-Saharan African settings (Thacher et al., 1999; Pettifor, 2004; Prentice et al., 2008; Braithwaite et al., 2016). This has important implications for treatment, for calcium supplementation alone has proven to be adequate for treatment of rickets caused by calcium deficiency. In Nairobi, it is likely that provision of supplemental vitamin D would be required (Thacher et al., 1999; Oginni et al., 2003). Furthermore, in this setting, acute malnutrition was associated with relatively low vitamin D levels, and 33% of children with SAM had absolute vitamin D levels in the deficient range.

The current study has not addressed the underlying causes of vitamin D deficiency in this setting. However, the fact that Nairobi’s climate should support sufficient vitamin D biosynthesis raises important behavioural questions around sunlight exposure. Edwards et al. reported that 71% of children with rickets in another Nairobi informal settlement had less than 3 hours’ sunlight exposure per week (Edwards et al., 2014). Poverty and urban living may act synergistically to limit infants’ exposure to sunlight. In Mathare, the imperative for mothers to seek employment very early in the postnatal period means that infants are often looked after in indoor informal daycare facilities (“babycares”) all day. Even where infants are cared for by a parent, concerns about personal security, the use of iron sheeting without windows as the predominant building material for dwellings, and an overcrowded and overlooked built environment without open spaces all limit sunlight exposure. As well as impacting the child, this may influence maternal vitamin D status, which has important consequences for the child’s vitamin D status at birth and the delivery of vitamin D in

TABLE 3 Relationships between biochemical and anthropometric indices

|                  | WLZ  |        | MUAC |        | LAZ  |        |
|------------------|------|--------|------|--------|------|--------|
| Calcium*         | 0.06 | <.001  | 0.040| .019   | 0.02 | .296   |
| Phosphate        | 0.16 | <.001  | 0.11 | .003   | 0.08 | .048   |
| ALP              | 14.9 | .086   | 13.6 | .078   | 7.9  | .175   |
| PTH              | −7.9 | .566   | −7.4 | .316   | −7.7 | .293   |
| 25(OH)D          | 7.4  | .016   | 9.3  | .002   | 0.69 | .574   |
| 1,25(OH)₂D       | 32.1 | .011   | 2.5  | .838   | 10.0 | .299   |

Note. Outputs of linear regression analyses. Output for LAZ is adjusted for age. LAZ = length-for-age z-score; MUAC = mid-upper arm circumference; WLZ = weight-for-length z-score.

*Calcium is adjusted for albumin as described in the methods section.
breastmilk (Salameh et al., 2016). Understanding the health and developmental consequences of growing up in sub-Saharan African urban informal settlements is important because urbanisation is proceeding at a high rate, does not appear to be associated with reduction in poverty, and is leading to dramatic increases in the number of people living in such circumstances (Henderson et al., 2013; Awumbila, 2014).

Some children had rickets without biochemical evidence of vitamin D deficiency. Although the clinical and demographic profile of this group was similar to that of children with rickets and a low 25(OH)D, the frequency of radiologically severe disease and secondary hyperparathyroidism were lower, and biochemical evidence of increased bone turnover was less strong (Figure 2 & supplementary Table 1). Whether these children had historic, healed, or “mild” rickets is difficult to assess, but they exemplify the fact that clinical diagnosis alone fails to reflect heterogeneity in the underlying pathophysiology and intensity of disease activity in a way that may be meaningful for treatment.

At present, there are no internationally accepted diagnostic criteria for rickets. Thacher and colleagues’ radiographic scoring method was designed for, and validated in, children older than 12 months who were independently mobile, but in Nairobi, rickets affects very young children and is associated with a high frequency of motor developmental delay (Thacher et al., 2002; Edwards et al., 2014; Munns et al., 2016). Diagnostic criteria that apply across geography, age, and developmental status are essential prerequisites to understanding the epidemiology and measuring the burden of rickets. There is no mention of rickets in either of the World Health Organization’s Pocket Book of Hospital Care for Children, or IMCI (Integrated Management of Childhood Illness) Handbook, and such data will help to improve the visibility of rickets as a significant concern in global child health (World Health Organization, 2005; World Health Organization, 2013).

The study has important limitations. We did not aim to perform population-based screening for features of rickets, and the children recruited should be considered a convenience sample, likely biased towards more severe cases. Therefore, these data do not preclude the existence of calcium deficiency rickets in the wider population, and further studies determining the epidemiology of rickets in this setting are needed. The decision to restrict recruitment to children younger than 2 years old was pragmatic, based on the fact that this age group represented the overwhelming majority of cases referred for treatment. Although calcium deficiency rickets in other settings has mainly been found in older children, the epidemiologic comparison should not be over-interpreted. Accurate dietetic assessment of calcium intake is very difficult in a breastfeeding population, and these data should be considered exploratory rather than definitive. When parents were completing the food frequency questionnaire, the diagnosis of rickets had been established, which increases the risk of recall bias, and although contextually-relevant, questionnaires had not been independently validated.

There was an association between wasting and deficiencies in serum adjusted calcium, phosphate, and vitamin D. This is likely to be due to a confounding effect of poverty, which might impact directly on food security, micronutrient intake, and secondarily on sunlight exposure, as previously described. Ready-to-use therapeutic foods are used for the outpatient management of SAM and are manufactured to internationally accepted compositional specifications (World Health Organization, World Food Programme, United Nations System Standing Committee on Nutrition, United Nations Children’s Fund, 2007). Based on their weight, most of these children would have received two to three sachets of ready-to-use therapeutic foods, providing high levels of calcium and phosphate, and up to 2400 IU vitamin D per day (World Health Organization et al., 2007). Although this is in line with minimum dosage recommendations for vitamin D in deficiency and rickets, the dose and duration of treatment may be suboptimal for bone health during catch-up growth (Munns et al., 2016).
SAM is associated with significant long-term health consequences, and the development of strategies targeted at healthy growth and development in the recovery phase is a research priority (Briend & Berkley, 2016). Vitamin D deficiency is a plausible mechanism explaining at least some of the increased vulnerability to severe infectious disease in SAM, because a number of studies have indicated, it increases the risk of respiratory tract infections (Esposito & Lelii, 2015). Public health strategies to improve vitamin D levels may provide benefit beyond prevention and treatment of rickets, but should be tested in rigorous controlled trials. Solar bottle bulbs (which involve the incorporation of water-filled plastic drink containers into the roof of a settlement) represent an innovative approach to increasing indoor sunlight exposure in urban informal settlements (www.literoflight.org), however the use of common polyethylene terephthalate plastic bottles means that the UVB wavelengths important for vitamin D production are filtered out (Sackey et al., 2015). Adaptation of this approach may be feasible and could provide a cheap and durable solution.

In summary, we have identified a group of young children living in an urban informal settlement in Nairobi who have rickets that appears to be predominantly caused by vitamin D deficiency and who would be likely to require vitamin D supplementation for full recovery. Development of standardised tools for diagnosis of rickets in this age group that could help establish the burden of disease should be a clinical and research priority. The apparent preponderance of vitamin D deficiency over calcium deficiency as cause of rickets in this setting prompts larger and more detailed epidemiologic studies in this and other settings.

ACKNOWLEDGMENTS

We are grateful for help and support in developing the project from Koki Kyalo at Concern Worldwide, Lucy Cathigi Maina and James Njiri at the Division of Nutrition, Ministry of Public Health & Sanitation, Government of Kenya, and John Warner at Imperial College, London. We are grateful to “German Doctors” for hosting the research at Baraka Health Centre and the volunteer medical staff who helped provide clinical care for the participants in the study (https://www.german-doctors.de/de/). We thank the nutrition, social care, and community health volunteer teams at Baraka Health Centre, especially Viola Akinyi Omamo, Zilpah Atlino Odoyo, Eclay Vusi Imbiakha, Doreen Mideva Kidagi, Pamela Akoth Aiyeko, Beatrice Wesonga Makachia, Peninah Akoth Obono, Elizabeth Adhiambio Ojijo, Julia Odinga Okechi, and Caroline Adhiambio Omollo. We thank the HNR analysts Kate Guberg, Abhilash Krishnakutty, Shailja Nigdikar, Ann Laidlaw, and Janet Bennet for expert analytical work. We thank Moses Ngari for programming assistance. We thank three anonymous reviewers for helpful suggestions. We remember our colleague and friend Cort Ulrich Hachmeister, whose dedication and enthusiasm helped to shape this project. We thank the study participants and their families.

CONFLICTS OF INTEREST

The authors declare that they have no conflicts of interest.

CONTRIBUTIONS

KDJJ, CUH, BHK, AP, JAB designed the study. KDJJ, CUH, CM, HSN were involved in clinical aspects of the study. M. K. coordinated dietary assessment. KDJJ, LC, IS, AP were involved in laboratory aspects of the study. KDJJ, JAB analysed the data. KDJJ wrote the first draft of the manuscript, all authors contributed to subsequent drafts and approved the final copy.

REFERENCES

Association for Clinical Biochemistry and Laboratory Medicine. (2015). Albumin-adjusted calcium: a position paper. Available at: http://www.acb.org.uk/docs/default-source/committees/clinical-practice/guidelines/acb-adjusted-calcium-position-paper-march-2015.pdf?sfvrsn=2 (accessed 6th January 2017)

Awumbila, M. (2014). Linkages between urbanization, rural-urban migration and poverty outcomes in Africa: World Migration Report 2015. International Organization for Migration: Manila, Philippines. Available at: https://www.iom.int/sites/default/files/our_work/ICP/MPR/WMR-2015-Background-Paper-MAwumbila.pdf (accessed 6th January 2017)

Baroncelli, G. I., Bereket, A., El Kholy, M., Summers, D. M., Nigdikar, S., Lavy, C. B.,... Prentice, A. (2016). The aetiology of rickets-like lower limb deformities in Malawian children. Osteoporosis International, 27(7), 2367–2372.

Braithwaite, V., Jarjou, L. M., Goldberg, G. R., Jones, H., Pettifor, J. M., & Prentice, A. (2012). Follow-up study of Gambian children with rickets-like bone deformities and elevated plasma FGF23: Possible aetiological factors. Bone, 50(1), 218–225.

Briend, A., & Berkley, J. A. (2016). Long term health status of children recovering from severe acute malnutrition. The Lancet Global Health, 4 (9), e590–e591.

Cashman, K. D., Dowling, K. G., Skrabakova, Z., Gonzalez-Gross, M., Valtuena, J., De Henauw, S.,... Kiely, M. (2016). Vitamin D deficiency in Europe: pandemic? The American Journal of Clinical Nutrition, 103(4), 1033–1044.

Edwards, J. K., Thiongo, A., Van den Bergh, R., Kizito, W., Kosgei, R. J., Sobry, A.,... Reid, A. J. (2014). Preventable but neglected: Rickets in an informal settlement, Nairobi, Kenya. Public Health Action, 4(2), 122–127.

Elder, C. J., & Bishop, N. J. (2014). Rickets. Lancet, 383(9929), 1665–1676.

Esposito, S., & Lelii, M. (2015). Vitamin D and respiratory tract infections in childhood. BMC Infectious Diseases, 15, 487.

Gloireux, F. H., & Pettifor, J. M. (2014). Vitamin D/dietary calcium deficiency rickets and pseudo-vitamin D deficiency rickets. Bonekey Rep, 3, 524.

Henderson, J. V., Roberts, M., & Storeygaard, A. (2013). Is urbanization in Sub-Saharan Africa different? Policy research working paper; no WPS 6481. World Bank: Washington DC. USA. Available at: http://documents.worldbank.org/curated/en/309811468194345036/is-urbanization-in-Sub-Saharan-Africa-different (accessed 6th January 2016)

Hogler, W. (2015). Complications of vitamin D deficiency from the foetus to the infant: One cause, one prevention, but who’s responsibility? Best Practice & Research. Clinical Endocrinology & Metabolism, 29(3), 385–398.

Kimani-Murage, E. W., Madise, N. J., Fotso, J. C., Kyobutungi, C., Mutua, M. K., Gitau, T. M., & Yatich, N. (2011). Patterns and determinants of urban migration in Sub-Saharan Africa different (accessed 6th January 2016)

Kimani Murage, E. W., Madise, N. J., Fotso, J. C., Kyobutungi, C., Mutua, M. K., Gitau, T. M., & Yatich, N. (2011). Patterns and determinants of breastfeeding and complementary feeding practices in urban informal settlements, Nairobi Kenya. BMC Public Health, 11, 396.

Ladhani, S., Srinivasan, L., Buchanan, C., & Allgrove, J. (2004). Presentation of vitamin D deficiency. Archives of Disease in Childhood, 89(8), 781–784.

Munns, C. F., Shaw, N., Kiely, M., Specker, B. L., Thacher, T. D., Ozono, K.,... Hogler, W. (2016). Global consensus recommendations on prevention and management of nutritional rickets. The Journal of Clinical Endocrinology and Metabolism, 101(2), 394–415.
Muungano Support Trust, Slum Dwellers International, University of Nairobi, University of California, Berkeley. (2012). Mathare zonal plan, Nairobi, Kenya. 2011 Collaborative Upgrading Plan. Available at: http://healthyCities.berkeley.edu/uploads/1/2/6/1/12619988/matharevalley_report_ucb_2_25_2012_final.pdf (accessed 6th January 2016)

Oginni, L. M., Sharp, C. A., Badru, O. S., Risteli, J., Davie, M. W., & Worsfold, M. (2003). Radiological and biochemical resolution of nutritional rickets with calcium. Archives of Disease in Childhod, 88(9), 812–817.

Pettifor, J. M. (2003). Nutritional rickets. In F. H. Glorieux (Ed.), Pediatric bone; biology and diseases. (pp. 541–565). San Diego: Academic Press.

Pettifor, J. M. (2004). Nutritional rickets: deficiency of vitamin D, calcium, or both? The American Journal of Clinical Nutrition, 80(Suppl), 1725S–1729S.

Pettifor, J. M. (2014). Calcium and vitamin D metabolism in children in developing countries. Annals of Nutrition & Metabolism, 64(Suppl 2), 15–22.

Pettifor, J. M., & Prentice, A. (2011). The role of vitamin D in paediatric bone health. Best Practice & Research. Clinical Endocrinology & Metabolism, 25(4), 573–584.

Prentice, A., Ceesay, M., Nigdikar, S., Allen, S. J., & Pettifor, J. M. (2008). FGF23 is elevated in Gambian children with rickets. Bone, 42(4), 788–797.

Sabbagh, Y., Carpenter, T. O., & Demay, M. B. (2005). Hypophosphatemia leads to rickets by impairing caspase-mediated apoptosis of hypertrophic chondrocytes. Proceedings of the National Academy of Sciences of the United States of America, 102(27), 9637–9642.

Sackey, S. S., Wovwotor, M. K., Owusu, A., Mensah-Amoah, P., Tatchie, E. T., Sefa-Ntiri, B., ... Atiemo, S. M. (2015). Spectroscopic study of UV transparency of some materials. Environment and pollution, 4(4), 1–17.

Salameh, K., Al-Janahi, N. S., Reedy, A. M., & Dawodu, A. (2016). Prevalence and risk factors for low vitamin D status among breastfeeding mother-infant dyads in an environment with abundant sunshine. Int J Womens Health, 8, 529–535.

Thacher, T. D., Fischer, P. R., & Pettifor, J. M. (2002). The usefulness of clinical features to identify active rickets. Annals of Tropical Paediatrics, 22(3), 229–237.

Thacher, T. D., Fischer, P. R., Pettifor, J. M., Lawson, J. O., Isichei, C. O., Reading, J. C., & Chan, G. M. (1999). A comparison of calcium, vitamin D, or both for nutritional rickets in Nigerian children. The New Englad Journal of Medicine, 341(8), 563–568.

Thacher, T. D., Fischer, P. R., Pettifor, J. M., Lawson, J. O., Manaster, B. J., & Reading, J. C. (2000). Radiographic scoring method for the assessment of the severity of nutritional rickets. Journal of Tropical Pediatrics, 46(3), 132–139.

Trilok Kumar, G., Chugh, R., & Eggersdorfer, M. (2015). Poor vitamin D status in healthy populations in India: A review of current evidence. International Journal for Vitamin and Nutrition Research, 85(3–4), 185–201.

United Kingdom Department of Health. (1991). Dietary reference values for food energy and nutrients for the United Kingdom: report of the panel on dietary reference values of the committee on medical aspects of food policy. Department of Health: London, UK.

United States Department of Agriculture. (2015). Agricultural Research Service, Nutrient Data Laboratory. USDA National Nutrient Database for Standard Reference (Release 28). U.S. Department of Agriculture: Washington D.C., USA. Available at https://ndb.nal.usda.gov/ (accessed 6th January 2016)

World Health Organization. (2005). Handbook: IMCI integrated management of childhood illness. WHO: Geneva, Switzerland. Available at http://apps.who.int/iris/bitstream/10665/42939/1/9241546441.pdf (accessed 6th January 2016)

World Health Organization. (2013). Pocket book of hospital care for children: Guidelines for the management of common childhood illnesses. Second edition. WHO: Geneva, Switzerland. Available at: http://www.who.int/maternal_child_adolescent/documents/child_hospital_care/en/ (accessed 6th January 2016)

World Health Organization, World Food Programme, United Nations System Standing Committee on Nutrition, United Nations Children’s Fund. (2007). Community-based management of severe acute malnutrition: a joint statement. Geneva: WHO. Available at: http://www.who.int/nutrition/topics/Statement_community_based_man_sev_acute_mal_eng.pdf?ua=1 (accessed 6th January 2016)

World Meteorological Organization. (2010). World Meteorological Organization Standard Normals. Available at: http://data.un.org/Data.aspx?q=sunshine&d=CLINO&f=ElementCode%3a15 (accessed 6th January 2017)

SUPPORTING INFORMATION

Additional Supporting Information may be found online in the supporting information tab for this article.

How to cite this article: Jones KDJ, Hachmeister CU, Khasira M, et al. Vitamin D deficiency causes rickets in an urban informal settlement in Kenya and is associated with malnutrition. Matern Child Nutr. 2018;14:e12452. https://doi.org/10.1111/mcn.12452