Cytomegalovirus Viremia Predicts Postdischarge Mortality in Kenyan HIV-Exposed Uninfected Children

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Background. Cytomegalovirus (CMV) viremia is associated with mortality in severely ill immunocompetent adults and hospitalized children with HIV (CWH). We measured CMV viremia in HIV-exposed and -unexposed Kenyan children aged 1–59 months discharged from hospital and determined its relationship with postdischarge mortality.

Methods. CMV DNA levels were measured in plasma from 1024 children (97 of which were HIV exposed uninfected [HEU], and 15 CWH). Poisson and Cox proportional hazards regression models were used to identify correlates of CMV viremia ≥ 1000 IU/mL and estimate associations with 6-month mortality, respectively.

Results. CMV viremia was detected in 31% of children, with levels ≥ 1000 IU/mL in 5.8%. HIV infection, age < 2 years, breastfeeding, and midupper arm circumference < 12.5 cm were associated with CMV viremia ≥ 1000 IU/mL. Among HEU children, CMV ≥ 1000 IU/mL (hazard ratio [HR] = 32.0; 95% confidence interval [CI], 2.9–354.0; P = .003) and each 1-log increase in CMV viral load (HR = 5.04; 95% CI, 1.7–14.6; P = .003) were associated with increased risk of mortality. CMV viremia was not significantly associated with mortality in HIV-unexposed children.

Conclusions. CMV levels at hospital postdischarge predict increased risk of 6-month mortality in Kenyan HEU children. CMV suppression may be a novel target to reduce mortality in HEU children.

Clinical Trials Registration. NCT02414399.

Keywords. cytomegalovirus; HIV-1; child mortality; postdischarge mortality; azithromycin.

Mortality risks in the 6 months following hospital discharge are high for children in resource-limited settings globally (approximately 3%–18%) [1, 2], and are especially high in young children with malnutrition or human immunodeficiency virus (HIV) infection [2]. HIV-exposed uninfected children (HEU) also experience higher morbidity and mortality [3], and impaired growth compared to HIV-unexposed (HUE) peers [4–6].

Increasing evidence suggests cytomegalovirus (CMV) may be an important contributor to poor outcomes in both immunocompetent and HIV-affected individuals admitted to hospital. Reactivation of CMV is common in immunocompetent adults admitted to intensive care units (ICU) and has been associated with higher risk of mortality, longer duration of hospitalization, increased risk of nosocomial infections, and greater oxygen dependency in multiple European- and US-based studies [7–11]. Primary CMV infection evokes a strong host inflammatory response, including increases in C-reactive protein, interleukin 6, and tumor necrosis factor-α [12–17]. The virus also has adapted several immune evasion strategies, including the production of host agonist and receptor mimics, which modulate host immune responses and may affect susceptibility to, or progression of, other infections [18]. Limited data suggest that prophylaxis with ganciclovir may improve oxygen use outcomes in septic adults with CMV viremia [19].

To date, no similar studies have been conducted in hospitalized immunocompetent children or adults in Africa; this is an important research gap given nearly universal CMV acquisition in early childhood, and high mortality rates in hospitalized African children [20–24]. Emerging data suggest CMV may have relevance for critically ill HIV-exposed children. In a cohort of hospitalized Kenyan children with HIV, the detection of CMV viremia at levels ≥ 1000 IU/mL at hospital admission was
associated with a 74% increased risk of death or hospitalization at 15 days and longer hospital stay (5 days), independent of age and HIV RNA level [25].

The association between CMV viremia and outcomes among hospitalized HEU and HU children has not been previously reported. We investigated the association between CMV viremia at hospital discharge and 6-month mortality in children participating in a randomized controlled trial of azithromycin at hospital discharge [26].

METHODS
Enrollment and Follow-up of Study Participants
The Toto Bora trial (NCT02414399) recruited children between 1 and 59 months of age at discharge from 1 of 4 hospitals in western Kenya (Kisii Teaching and Referral Hospital, Homa Bay Teaching and Referral Hospital, St Paul Mission Hospital, and Kendu Adventist Mission Hospital) after admission for any conditions other than trauma, injury, or congenital abnormalities [26]. Children were randomized to receive placebo or azithromycin for 5 days (10 mg/kg on day 1, followed by 5 mg/kg on days 2–5), with the first dose being directly observed at the health facility. Children were followed for 6 months, with 2 scheduled follow-up visits (month 3 and month 6) where caregivers were interviewed about the child’s health history and a physical examination was performed. Anthropometric measurements were taken at all scheduled visits, as were blood samples collected in EDTA tubes, centrifuged, and resulting plasma and buffy coat stored at −80°C until further use. Height/length-for-age z-scores, weight-for-age z-scores, and weight-for-height/weight-for-length z-scores were calculated using the World Health Organization ANTHRO software [27]. Stunting and underweight were defined as height-for-age z-scores/length-for-age z-scores and weight-for-age z-scores < −2, respectively. Moderate wasting was defined as ≥ −3 to < −2 weight-for-height z-scores or ≥ 11.5 to < 12.5 cm midupper arm circumference, and severe wasting as weight-for-height z-scores < −3 or midupper arm circumference < 11.5 cm. Midupper arm circumference was only considered for wasting categorization in children 6 months or older. HIV status was determined by abstracting available data from medical records and/or testing during the hospitalization where indicated (per Kenyan Ministry of Health guidelines [28]). Dates of death and hospitalizations were determined through medical record abstraction (including nonrecruiting hospitals) and by self-report during scheduled or unscheduled visits. Cause of death was determined by clinical consensus after review of death certificates and medical records (when available), and verbal autopsies and assigned causes of death as described elsewhere [29].

CMV Assays and Definitions
CMV DNA levels were measured from stored baseline plasma specimens using real-time quantitative polymerase chain reaction (PCR) as previously described; primers and probes are provided in Wamalwa et al [25]. The assay has a limit of detection of 1 copy/reaction, and values were transformed to express measurements in international units (IU)/mL by dividing by 1.4. We used a cutoff of 1000 IU/mL (high CMV viremia) as our primary outcome (for correlates analysis) and exposure (for mortality analysis) of interest, based on previous papers reporting an association with mortality in immunocompetent adults admitted to ICU [30], increased mortality and duration of hospitalization in hospitalized children with HIV (CWH) [25], and decreased lung function and stunting in Zimbabwean CWH [31].

Statistical Analysis
Stata version 14 (StataCorp) was used for all analyses, and all comparisons were 2 sided with α = .05. Pearson χ² test was used to compare CMV prevalence between groups of children based on HIV exposure, and the Wilcoxon rank-sum test was used to compare the distribution of CMV levels between groups. Correlates of CMV viremia were identified using Poisson regression to estimate prevalence ratios (PR) and 95% confidence intervals (CI), adjusting for baseline age ≥ 24 months, and HIV exposure or infection status. Cox proportional hazards regression was used to compare mortality rates associated with baseline CMV status (CMV viremia ≥ 1000 IU/mL and CMV viral load [log IU/mL]). Multivariable Cox regression models were adjusted for age. While randomization arm and site were considered as potential confounders, neither were retained in the model due to the lack of observed change in hazard ratios from an age-adjusted model by more than 10%. Two-way interaction terms between CMV and HIV exposure status were tested using a likelihood ratio test comparing models with and without (nested model) the interaction term in the model. Models and Kaplan-Meier curves were stratified by HIV exposure due to an a priori assumption that CMV viremia may be evoked and persist by unique mechanisms, and have differential consequences for HU, HEU, and CWH. Also, in secondary analyses we evaluated the association between CMV and mortality in groups of children defined by stunting and wasting status (among HU children).

The study was approved by the institutional review boards at the Kenya Medical Research Institute, the Kenya Pharmacy and Poisons Board, and the University of Washington. Caregivers provided informed written consent (or oral consent with a witnessed thumbprint) in their preferred local language (English, Kiswahili, Kisii, or Dholuo).

RESULTS
Participant Characteristics
The first 1024 of the 1400 enrolled Toto Bora participants (73%) had CMV PCR testing performed and were included in this analysis; characteristics are provided in Table 1. The median
duration of hospital stay among the 1024 children included in this analysis was 3 days, with 17.6% admitted for 7 days or longer. A high proportion of children were stunted (23.9%), and many children were underweight (12.4%) or wasted (9.0%). In total, 118 (11.5%) of enrolled children had documented HIV exposure or infection; 103 (10.1%) were HEU and 15 (1.5%) had HIV infection (CWH). There were 6 children who were HIV exposed but infection status unknown and 34 children confirmed to be HIV uninfected, but in whom HIV-exposure status was unknown.

**CMV Detection and Levels**

All 1024 participants with CMV PCR testing had valid results at enrollment. At hospital discharge, 314 children had detectable CMV viremia (31%), with 5.8% having CMV levels ≥1000 IU/mL (Table 2). Infants <1 year old had the highest prevalence of CMV detection and the highest median level of CMV viremia, with children 1–2 years and >2 years of age having successively lower prevalence of viremia and CMV viral loads (Figure 1). The prevalence of CMV viremia and the median CMV viral load was similar between HEU and HU children (P > .05 for each comparison). However, 6 (0.69%) HU had very high CMV levels (≥10000 IU/mL); 2 HU children, 5 and 7 months of age, had CMV levels ≥100000 IU/mL and 4 children, between the ages of 5 and 13 months, had levels 10000–100000 IU/mL. Among HEU children, 3 (2.9%) had CMV viral loads ≥100 000 IU/mL and none were ≥100 000 IU/mL. Among CWH, 2 had CMV viral loads ≥100 000 IU/mL. While the total number of CWH in the cohort was small (n = 15), these children had significantly higher median CMV viral loads (3.2 log_{10} IU/mL; interquartile range [IQR], 2.6–5.3) and there was a higher proportion of children with viral loads ≥1000 IU/mL (20.0%) as compared to the HU children (median, 2.3 log_{10} IU/mL; IQR, 1.9–2.7 and 5.2% ≥1000 IU/mL; P = .012 and P = .015, respectively).

No single diagnosis explained the high frequency of CMV viremia in this study population. Children admitted with sepsis
had the highest median CMV viral load at hospital discharge (2.69 log₁₀ IU/mL) and 3.3% of children with sepsis had ≥ 1000 IU/mL (Table 3). Children diagnosed with pneumonia (7.3%), diarrhea (6.4%), and anemia (6.8%) were the most likely to have viral loads ≥ 1000 IU/mL.

**Correlates of CMV Viremia**

Baseline demographics, nutritional status, and HIV exposure were assessed as correlates of CMV viremia ≥ 1000 IU/mL at hospital discharge (Table 4). Effect sizes for each covariate were similar for HU and HEU children in crude and adjusted models (data not shown), so data are presented for the cohort overall. When adjusting for HIV exposure, children < 24 months were over 8 times more likely to have CMV viremia ≥ 1000 IU/mL at hospital discharge than older children (age-adjusted PR [aPR], 8.27; 95% CI, 3.03–23.1; P < .001). HIV infection was also highly associated with CMV viremia (aPR, 4.98; 95% CI, 1.55–16.1; P = .007), but HEU status was not significantly associated with a greater prevalence of viremia (aPR, 1.36; 95% CI, .58–3.20; P = .5). Adjusting for age and HIV exposure, caregiver-reported breastfeeding during the hospital stay, among those younger than 24 months of age, was the only other predictor of CMV viremia (aPR, 2.26; 95% CI, 1.05–4.87; P = .04).

**CMV Viremia and Risk of Mortality After Hospital Discharge**

Overall, there were 24 deaths among the 1024 included children during the 6 months of follow-up (2.3%). Thirteen deaths were due to infectious causes (pneumonia [n = 2], malaria [n = 1], and diarrhea [n = 1], central nervous system infections [n = 2], and unknown cause of death with signs of infection [n = 7]) and 11 from noninfectious causes (congestive heart failures [n = 2], anemia [n = 3], sickle cell crisis [n = 1], and unknown without signs of infection [n = 5]). Figure 2 shows the Kaplan-Meier curves for children overall and grouped by HIV exposure.
and infection status. Survival rates in children with low-level CMV viremia (<1000 IU/mL) were similar to those of children without detectable CMV viremia \((P = .70)\). However, children with high-level CMV viremia \((\geq 1000 \text{ IU/mL})\) had lower survival rates when compared both to children without detectable CMV \((P = .02)\) and to children with low-level viremia <1000 IU/mL \((P = .08; \text{Supplemental Figure 1})\). As a result, in subsequent analyses we compared mortality rates between children with high-level \((\geq 1000 \text{ IU/mL})\) versus low-level or no viremia (Figure 2).

Children with CMV level \(\geq 1000 \text{ IU/mL}\) at hospital discharge were close to 3 times more likely to die than children with low-level or no viremia adjusting for age \((aHR, 2.78; 95\% \text{ CI}, .91–8.50)\), albeit at borderline statistical significance \((P = .07; \text{Table 5})\). Stratification by HIV exposure revealed that this association was driven by HIV-exposed children \((\text{likelihood ratio test of interaction between HIV-exposure status and CMV viral load } \geq 1000 \text{ IU/mL } P = .048, \text{ and CMV log}_{10} \text{IU/mL level } P = .011)\). In HU children, neither CMV viral load \(\geq 1000 \text{ IU/mL}\) \((aHR, 1.16; 95\% \text{ CI}, .15–8.8; P = .89)\) nor CMV log_{10} IU/mL

### Table 3. Prevalence and Levels of CMV Viremia at Hospital Discharge, by Diagnosis

| Diagnosis       | No. of Cases | CMV Viremia, No. (%) | CMV \(\geq 1000 \text{ IU/mL}, \text{ No. (\%)}\) | log_{10} CMV IU/mL in Viremics, Median (IQR) |
|-----------------|--------------|---------------------|---------------------------------|------------------------------------------------|
| Pneumonia       | 314          | 108 (34)            | 23 (7.3)                        | 2.41 (1.91–2.78)                                |
| Malaria         | 278          | 86 (31)             | 11 (4.0)                        | 2.20 (1.85–2.58)                                |
| Diarrhea        | 188          | 60 (32)             | 12 (6.4)                        | 2.28 (1.84–2.89)                                |
| Anemia          | 133          | 48 (36)             | 9 (6.8)                         | 2.36 (1.92–2.67)                                |
| Sickle cell\(^b\) | 70           | 17 (24)             | 1 (1.4)                         | 1.88 (1.70–2.12)                                |
| Suspected meningitis | 53   | 12 (23)             | 2 (3.8)                         | 2.43 (2.01–2.96)                                |
| Sepsis          | 30           | 11 (37)             | 1 (3.3)                         | 2.69 (2.04–2.98)                                |
| Tuberculosis    | 15           | 2 (13)              | 0                               | 1.92 (1.70–2.15)                                |

### Table 4. Correlates of CMV DNA \(\geq 1000 \text{ IU/mL}\) at Hospital Discharge in Kenyan Children

| Characteristic                  | CMV DNA \(\geq 1000 \text{ IU/mL}\) Prevalence in Correlate Group, No. (%) | Crude Prevalence Ratio (95\% CI), PValue | Age and HIV-Exposure-Adjusted Prevalence Ratio (95\% CI), PValue |
|---------------------------------|--------------------------------------------------------------------------|-----------------------------------------|------------------------------------------------------------------|
| Male                            | 32 (5.2)                                                                 | Ref                                     | Ref                                                              |
| Female                          | 27 (6.8)                                                                 | 1.31 (.785–2.2), P = .3                | 1.35 (.808–2.25), P = .3                                          |
| Age > 24 mo                     | 4 (1.1)                                                                  | Ref                                     | Ref                                                              |
| Age < 24 mo                     | 55 (8.6)                                                                 | 8.18 (2.97–22.6), P < .001             | 8.27 (3.03–23.1), P < .001                                       |
| Not underweight                 | 48 (5.4)                                                                 | Ref                                     | Ref                                                              |
| Underweight                     | 11 (8.7)                                                                 | 1.62 (.840–3.11), P = .2               | 1.30 (0.672–2.51), P = .4                                        |
| Not wasted                      | 51 (6.5)                                                                 | Ref                                     | Ref                                                              |
| Moderate wasting                | 4 (7.1)                                                                  | 1.31 (.472–3.61), P = .6               | 1.19 (.430–3.30), P = .7                                         |
| Severe wasting                  | 4 (11.1)                                                                 | 2.03 (.734–5.62), P = .2               | 1.31 (.470–3.66), P = .6                                         |
| Not stunted                     | 49 (6.3)                                                                 | Ref                                     | Ref                                                              |
| Stunted                         | 10 (4.1)                                                                 | 0.649 (.329–1.28), P = .2             | 0.660 (.332–1.31), P = .2                                        |
| Midupper arm circumference > 12.5 cm | 28 (3.4)                                                            | Ref                                     | Ref                                                              |
| Midupper arm circumference < 12.5 cm | 6 (9.1)                                                            | 2.68 (1.10–6.44), P = .03\(^a\)      | 1.88 (0.769–4.60), P = 2\(^a\)                                    |
| In hospital < 7 days            | 45 (5.4)                                                                 | Ref                                     | Ref                                                              |
| In hospital > 7 days            | 13 (7.3)                                                                 | 1.35 (.728–2.50), P = .3               | 1.19 (.641–2.22), P = .6                                         |
| Not breastfed while in hospital | 7 (4.5)                                                                  | Ref                                     | Ref                                                              |
| Breastfed while in hospital     | 47 (9.7)                                                                 | 1.92 (.908–4.06), P = .09\(^b\)      | 2.26 (1.05–4.87), P = .04\(^b\)                                  |
| HIV unexposed                   | 45 (5.2)                                                                 | Ref                                     | Ref                                                              |
| HIV exposed uninfected           | 6 (6.2)                                                                  | 1.20 (.511–2.81), P = .7               | 1.36 (.582–3.20), P = 5\(^c\)                                    |
| HIV infected                    | 3 (20.0)                                                                 | 3.88 (1.15–11.9), P = .02             | 4.98 (1.55–16.1), P = .007\(^c\)                                |

\(^a\)Adjusted for age > 24 months and HIV exposure (HIV exposed uninfected or infected).

\(^b\)Breastfeeding point estimate among children < 24 months old (n = 640), adjusted only for HIV exposure.

\(^c\)Adjusted only for age > 24 months.

Abbreviations: CMV, cytomegalovirus; HIV, human immunodeficiency virus; P, prevalence ratio.

\(^a\)Midupper arm circumference point estimate among children > 6 months old (n = 887).

\(^b\)Breastfeeding point estimate among children < 24 months old (n = 640), adjusted only for HIV exposure.

\(^c\)Adjusted only for age > 24 months.
IU/mL levels (aHR, 1.05; 95% CI, .47–2.32; \( P = .91 \)) were associated with risk of death in crude or adjusted models. While there were 14 deaths among HU children, there was only a single death among HU children with CMV viral load > 1000 IU/mL. Further stratification in the HU group by malnutrition status did not reveal evidence of an association between CMV and death (Supplementary Table 2). HEU children with CMV level ≥ 1000 IU/mL had a much higher risk of death than children with low-level or no viremia (aHR, 32.0; 95% CI, 2.90–353.95; \( P = .005 \)), and each 1-log increase in CMV IU/mL was associated with a 5-fold increased risk of death (aHR, 5.04; 95% CI, 1.74–14.60; \( P = .003 \)). There were only 2 deaths observed among the 15 CWH, 1 in each CMV group, limiting the statistical power in this subgroup analysis. However, the magnitude of age-adjusted hazard ratios were similar to those observed in HEU children (CMV viral load ≥ 1000 IU/mL aHR, 10.7; 95% CI, .04–3056.00; \( P = .41 \), and 1-log increase in CMV IU/mL aHR, 2.58; 95% CI, .692–9.64; \( P = .26 \)).

**DISCUSSION**

In this cohort of Kenyan children assessed at hospital discharge, we found that nearly a third had detectable CMV viremia in their blood, with some children having extremely high CMV viral loads. Infants (aged ≤12 months), those living with HIV, and who were currently breastfeeding had the highest prevalence of CMV viremia ≥ 1000 IU/mL. Independent of age, the detection of CMV at ≥ 1000 IU/mL was associated with an increased risk of mortality over the next 6 months in HEU children, as was each 1-log increase in CMV viremia.

To the best of our knowledge, this is the first study to assess the association between CMV viremia at hospital discharge and mortality. Our data suggest that CMV viremia ≥ 1000 IU/mL at hospital discharge is strongly associated with risk of mortality in HEU children, with each 1-log increase in CMV viremia associated with a 5-fold increased risk of death. HEU children experience double the risk of hospitalization and death than their HU counterparts [32, 33], yet the mechanism underpinning this
vulnerability are not fully understood. The intersecting and complex sociodemographic, behavioral, and physiologic factors, including maternal disease experience and antiretroviral therapy exposure, altered immune ontology during fetal development, and exposure to more maternal infections likely all play a role [34]. All of these factors could alter CMV pathogenesis in HEU children, and this assumption is supported by previous work demonstrating high-level and prolonged CMV viremia in otherwise healthy HEU infants during primary CMV infection [35–37].

Given prior reported associations between CMV reactivation and mortality in severely ill hospitalized adults [7–11] and CWH [25], these findings suggest that CMV viremia may also be an important marker of risk in HEU children. This is especially remarkable given the potential for survival bias among children enrolled at hospital discharge, as many of the sickest children may not have survived to enrollment. These findings are also consistent with previous data published from our group demonstrating that among a large cohort of CWH admitted to hospital, CMV viremia ≥ 1000 IU/mL was associated with a 74% increased risk of death or continued hospitalization at 15 days and with 5 days longer hospitalization [25]. While we did not find an association between CMV viremia and mortality in HU children, we caution that the study had limited statistical power to detect more modest, but still important, associations with mortality in this subgroup. Notably, survival bias excludes the sickest children from our study; if we had enrolled at hospital admission, we may have found an association between CMV and inpatient mortality and more research is needed to determine if hospitalized HU children should be included in CMV-targeted interventions.

We are unable to determine whether CMV viremia is causally linked to mortality, or if it is simply a marker of more severe illness, a barometer of immune competence, or an unidentified cofactor. A number of children in our study had extremely high CMV viral loads; while CMV was not a specific diagnosis in any participant included in this study, it is possible that children with the highest CMV viral loads may have had CMV disease and not just viremia. CMV has a wide tissue tropism, and can replicate in the lung, central nervous system, and gut, causing pneumonitis, neurologic diseases, enteritis, and esophagitis [38–41]. To date, limited trial data exist to support or refute a causal role for CMV. In a randomized control trial of 156 mechanically ventilated CMV-seropositive adults with trauma or sepsis, randomization to the antiviral drug ganciclovir increased the number of ventilator-free days, and decreased the duration of mechanical ventilation compared to placebo; notably, these benefits were observed only in those with sepsis [19].

The prevalence of CMV viremia in all children, regardless of HIV exposure history, was high and was in the range of CMV reactivation rates (5%–50%) found in studies of immunocompetent, severely ill adults admitted to ICU [7, 8]. CMV viremia was associated with younger age and currently breastfeeding, although detectable virus was also found in a substantial proportion of children older than 2 years (21%). In our previous birth cohort studies, we found that virtually all HIV-exposed children acquired CMV infection before the first year of life, and that high-level and prolonged CMV viremia (>3 months in duration) was common, with most CWH and HEU having detectable CMV viremia at least 6 months, and some continuing to be viremic for more than a year [35–37].

Strengths of the study include the well-characterized and large study population, complete ascertainment of consecutive...
enrollment samples, and inclusion of a large number of HEU and HU children. However, the total number of deaths was small and the effect sizes reported here have limited precision. With only 15 CWH included, we were underpowered to confirm an association between CMV viremia and postdischarge mortality in this population. It was also not possible to verify HIV infection status in 6 children, nor to capture any new HIV infections postdischarge, and HIV treatment and laboratory measures (CD4, HIV RNA levels) were not available in the CWH. Admission diagnoses were abstracted from medical records and may be subject to misclassification. As noted above, assessment of children at hospital discharge, rather than admission, biases inclusion for children with less-severe disease, and could reduce observed associations between CMV viremia and mortality. Finally, because we did not perform CMV serology, we cannot determine CMV infection status in the CMV aviremic children.

In summary, these data suggest that CMV viremia is common in hospitalized Kenyan children and that in HEU children CMV viremia $\geq$ 1000 IU/mL identifies a subset of children at high risk for death. These data support the need for interventional studies to be conducted in hospitalized, African CWH and HEU who have a high risk for both inpatient and postdischarge mortality, as well as more research into mechanisms of CMV reactivation, persistence, and pathogenesis in the inpatient setting.

**Supplementary Data**

Supplementary materials are available at *The Journal of Infectious Diseases* online. Supplementary materials consist of data provided by the author that are published to benefit the reader. The posted materials are not copyedited. The contents of all supplementary data are the sole responsibility of the authors. Questions or messages regarding errors should be addressed to the author.

**Notes**

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**Data availability.** The data that support the findings of this study are available from the corresponding author upon reasonable request.

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