Differences in diagnosis, management, and outcomes of acute febrile illness by health facility level in southern Ethiopia

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We assessed the diagnosis, management and outcomes of acute febrile illness in a cohort of febrile children aged under 5 years presenting at one urban and two rural health centres and one tertiary hospital between 11 August 2019 and 01 November 2019. Pneumonia was diagnosed in 104 (30.8%) of 338 children at health centres and 128 (65.0%) of 197 at the hospital (p < 0.001). Malaria was detected in 33 (24.3%) of 136 children at the urban health centre, and in 55 (55.6%) of 99 and 7 (7.4%) of 95 children at the rural health centres compared to 11 (11.6%) of 95 at the hospital. Antibacterials were prescribed to 20 (11.5%) of 174 children without guidelines-specified indications (overprescribing) at health centres and in 7 (33.3%) of 21 children at the hospital (p = 0.013). Antimalarials were overprescribed to 13 (7.0%) of 185 children with negative malaria microscopy at the hospital. The fever resolved by day 7 in 326 (99.7%) of 327 children at health centres compared to 177 (93.2%) of 190 at the hospital (p < 0.001). These results suggest that additional guidance to health workers is needed to optimise the use of antimicrobials across all levels of health facilities.

Continuing the substantial reduction of childhood mortality in low-resource settings requires a focus on improved diagnosis and management of infectious diseases, many of which present commonly with fever, across all levels of the health service1–3. To assist health workers in responding to childhood febrile illnesses in these settings, simplified World Health Organization (WHO) guidelines are available, including the Pocket Book of Hospital Care for use at hospitals4 and the integrated management of childhood illness (IMCI) at health centres5. These guidelines recommend prescribing antibacterial therapy based on meeting clinical case definitions for illnesses, including pneumonia and dysentery, and emphasize parasitological confirmation of malaria before offering antimalarial agents. Integrating malaria diagnostics, including microscopy and rapid diagnostic tests, into fever case management has reduced the unnecessary use of antimalarials6, and contributed to the substantial global reduction in malaria morbidity and mortality7,8. However, a lack of accessible diagnostics for various bacterial and viral infections has remained a barrier to improved guidelines and practice in fever management9. For fever without clinical evidence of bacterial infection, guidelines recommend withholding antibacterial treatment unless patients have signs of severe illness10. Nevertheless, in the absence of reliable diagnostics to explain the cause of the fever in patients with negative malaria tests, health workers may not adhere to guidelines and unnecessarily prescribe antimalarials11–13 and antibacterials14,15, which may lead to the development of drug resistance16,17.

We recently reported a low proportion of malaria (3.2%)18, a lack of adherence to guidelines in prescribing antimicrobials (antibacterials and antimalarials), and substantial proportions of hospitalization (38.1%) and in-hospital fatality ratio (5.9%) in febrile children aged under 13 years presenting at the largest tertiary hospital in southern Ethiopia19. To determine how the pattern of childhood febrile illness, health workers’ adherence to guidelines for the initiation of antimicrobials, and outcomes may differ in other health settings, we extended our investigation to lower-level health facilities in the catchment areas of the hospital. Thus, the current study aimed to assess whether there are differences in the diagnosis, management, and outcomes of febrile illness in
children under 5 years old presenting at lower-level of health facilities (urban and rural health centres) compared to the tertiary hospital.

Methods

Study design and setting. This study is reported according to the Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) guideline (Checklist S1). A prospective cohort study was undertaken from 11 August 2019 to 01 November 2019 at health facilities in and around Hawassa City, southern Ethiopia, located on the shores of Lake Hawassa, the capital of Southern Nations, Nationalities, and Peoples’ Region (SNNPR). Tilte health centre from an urban area and Finchawa and Gara-Rikata health centres from rural areas were lower-level health facilities purposively selected for the study, considering the distance from Lake Hawassa, which may influence water-associated infectious diseases. We also recruited participants from Hawassa University Comprehensive Specialized Hospital (HUCSH), the largest tertiary hospital in the SNNPR. As shown in Fig. 1, all health facilities are close to Lake Hawassa, except Gara-Rikata, which is located about 10 km away. The major malaria transmission season in Ethiopia is from September to December, following the rainy season during June to August. About 65% of the administrative region of SNNPR is malarious though the incidence has declined in recent years20. In addition to vaccines for tuberculosis, diphtheria, pertussis, tetanus, poliomyelitis, measles, and hepatitis B virus, the national childhood immunization program in Ethiopia introduced the Hae ‑ mophilus influenzae type b vaccine, pneumococcal conjugate vaccine, and monovalent rotavirus vaccine more recently21. Full immunization (as defined by the Ethiopian vaccination schedule) coverage of SNNPR among children aged 12–23 months was 38% in 201922.

Participant eligibility and enrolment. Children consecutively presenting to the participating health facilities to seek care 7 days per week, 24 h per day, during the study period were screened for study eligibility. Those children under 5 years old with fever, defined as an axillary temperature of at least 37.5 °C or a history of fever at least once in the past 48 h, lasting no longer than 7 days, were enrolled. We excluded patients for whom minor skin infections were the main reasons for the visit, and those who were referred to a hospital urgently, therefore, did not undergo routine examinations and management at health centres.

Sample size. The sample size was computed based on outcomes reported in another study, relating to health-seeking behaviour by caregivers attending various levels of health facilities23. Thus, the current analysis was based on data gathered from the 338 enrolled children from health centres, 138 from Tilte and 100 each from Finchawa and Gara-Rikata, and 197 from the hospital. The target sample size for each facility was proportional to the number of patients recorded as having attended in the same season in the previous year. Consecutive febrile children meeting the inclusion criteria were enrolled until the targeted sample size was obtained.

Data collection. Diagnosis and management on enrolment day. The study personnel were nurses and health officers from each facility, who had 4 years of training in clinical services. They received training in the study data collection procedures, and reviewed patients’ records immediately on initial clinical management on enrolment day. Data captured using a study-specific form included the child’s demographic characteristics (sex, age), clinical information (treatment taken prior to study enrolment for index illness, duration of fever, axillary temperature at presentation, and symptoms and signs) and laboratory and clinical diagnosis. The management done for the participants at the study facilities, including treatment prescribed, hospital admission, and referral to higher-level care, were also captured. Available at all participating health facilities were blood smear micros-
Follow-up data. Clinical outcome data were collected from caregivers of children on day 7 (± 1) using a study-specific form. Caregivers of children managed as outpatients were contacted by telephone or visited at home in rural areas if contact by telephone was impossible. For hospitalized patients, records were also reviewed to collect follow-up data. Outcomes assessed included resolution of fever, hospital stay, hospitalization after initial treatment, and outcomes at lower-level health facilities (health centers) and a higher-level facility (tertiary hospital) were evaluated using the Pearson Chi-square test or Fisher's Exact test as appropriate. A p-value < 0.05 from any statistical test was considered to show a significant difference.

Data analysis. Data were entered using EpiData version-3.126, and analysed using SPSS Statistics for Windows, Version 20.0 (IBM SPSS Statistics for Windows, Version 20.0; IBM Corp., Armonk, NY, United States). Results for categorical variables, including demographic and clinical characteristics, diagnosed illnesses, treatment, and outcomes, were summarized using frequency (percentage). Quantitative data, including child age, duration of fever before and after enrolment, and hospital stay, were summarized using median (interquartile range, IQR). Differences between proportions of demographic and clinical characteristics, diagnosed illnesses, prescribed treatments and adherence with guidelines, and outcomes at lower-level health facilities (health centers) and a higher-level facility (tertiary hospital) were evaluated using the Pearson Chi-square test or Fisher's Exact test as appropriate. A p-value < 0.05 from any statistical test was considered to show a significant difference.

Ethics approval and consent to participate. The study was approved by the Human Research Ethics Committee of the University of New South Wales (UNSW), Sydney, Australia (Ref. No: HC190358) and the Institutional Review Board of Hawassa University College of Medicine and Health Sciences, Hawassa, Ethiopia.
(Ref. No: IRB/223/11). Adequate information was given to caregivers about the study. Participation was voluntary, and informed written consent was obtained from caregivers. All methods were carried out following relevant guidelines and regulations.

Results

Demographic characteristics of the children. Overall, 535 participants were enrolled in the study, 138 (25.8%) from the urban health centre (Tilte), 100 (18.7%) each from the rural health centres (Finchawa and Gara-Rikata), and 197 (36.8%) at the tertiary hospital, HUCSH. A further 9 children were determined to have not met the eligibility criteria, so they were excluded. On average, children recruited at HUCSH were younger than those at health centres, with median age (IQR) of 12 (6–24) months and 24 (12–36) months, respectively, p < 0.001 (Table 2). No significant difference was observed in the distribution of child’s sex between health centres and the hospital (p = 0.608) (Table 2).

Presenting clinical characteristics. The median (IQR) duration of fever at enrolment among children attending health centres was 1 (1–2) day compared to 3 (2–5) days in those presenting to HUCSH. Of 338 children attending health centres, 11 (3.3%) had taken antibacterial agents for the current episode of illness prior to enrolment compared to 103 (52.3%) of 197 at HUCSH (p < 0.001). Further, lower proportions of children attending health centres compared to HUCSH had a cough (42.9% vs. 76.6%; p < 0.001), fast breathing (28.4% vs. 57.4%), and vomiting (26.6% vs. 44.7%). In contrast, tonsillar signs and shivering were more common presenting features among children at health centres (p < 0.001) (Table 2). Children presenting at lower-level health facilities had a smaller number of symptoms and signs compared to those at a tertiary hospital (p < 0.001).

Diagnoses. As shown in Table 3, 180 (53.3%) of 338 children at health centres compared to 150 (76.1%) of 197 children at HUCSH had acute respiratory infections (ARIs). Pneumonia was a common illness, diagnosed in 104 (30.8%) of children at health centres and in 128 (65.0%) of those at HUCSH (p < 0.001). Acute tonsillopharyngitis was more frequent at Tilte urban health centre (HC), diagnosed in 46 (33.3%) of 138 children. Malaria was found in 95 (28.8%) of 330 children whose blood samples were examined by microscopy at health centres, and in 11 (11.6%) of 95 children at HUCSH. At Finchawa rural HC, 55 (35.6%) of 99 children had malaria compared to 7 (7.4%) of 95 at Gara-Rikata rural HC, and 33 (24.3%) of 136 at Tilte urban HC. Of 425 blood samples tested at all study sites, 48 (11.3%) were positive for *Plasmodium falciparum* and 60 (14.1%) for *P. vivax*; with 2 (0.5%) mixed infections. Thirty-nine (11.6%) of 338 children at health centres and 8 (4.1%) of 197 children at HUCSH had undifferentiated fever (p = 0.003).

Clinical management. Table 4 summarizes the clinical management and outcomes of febrile illnesses in different health facilities. Of 338 children at health centres, 5 (1.5%) were referred to a higher-level facility for further care, while 138 (70.4%) of 196 at HUCSH were hospitalized at initial management on enrolment day. Of those hospitalized, 94 (68.1%) were due to pneumonia. Antibacterial agents were prescribed to 184 (54.4%) of 338 children at health centres compared to 182 (92.9%) of 196 children managed at HUCSH (p < 0.001). Among 174 children without indication for antibacterial therapy at health centres, 20 (11.5%) received the treatment, as did 7 (33.3%) of 21 at HUCSH (p = 0.013). Of those 20 children, 8 (40.0%) had unspecified upper respiratory infections, 9 (45.0%) had acute diarrhoea, and 3 (15.0%) had undifferentiated fever.

For antimalarials, 95 (28.1%) of 338 children at health centres were prescribed; all of which met guidelines. Whereas, at HUCSH, antimalarials were prescribed to 24 (12.2%) of 196 children, including 13 (7.0%) of 185 children without laboratory confirmation of malaria (Table 4).

Clinical outcomes on follow-up. Clinical outcome data were unavailable for 11 (3.3%) of 338 children at health centres and 5 (2.5%) of 197 children at HUCSH. Of 327 children managed at health centres, 5 were hospitalized elsewhere: 2 children were referred to higher-level care, and 3 were initially sent home. Of 192 children at HUCSH contacted on day 7 (±1), 136 (70.8%) had been hospitalized on initial management, with a median (IQR) stay of 5 (4–7) days, while one child initially managed as an outpatient had been admitted by follow-up. The fever resolved by day 7 in 326 (99.7%) of 327 children managed at health centres and 177 (93.2%) of 190 seen at HUCSH (p < 0.001). The median (IQR) time to resolution of fever from the baseline visit was 1 (1–2) day in children at health centres and 2 (1.8–3) days in those at HUCSH. Two (1.5%) of 136 inpatients at HUCSH died: one with pneumonia and acute diarrhoea and the other with acute diarrhoea (Table 4). Within the follow-up period, 9 (4.7%) of 192 children at HUCSH and 6 (1.8%) of 327 at health centres visited another or the same facility.

Discussion

Our analysis identified several differences between presenting febrile illness, clinical management, and outcomes in primary- and tertiary-level health facilities and rural and urban settings. Pneumonia was the most common clinical diagnosis at the tertiary hospital (HUCSH), while malaria was prevalent at lower-level health facilities in both urban (Tilte) and rural (Finchawa) settings located closer to Lake Hawassa, in contrast to a rural setting further from the lake (Gara-Rikata). We also observed that children presenting at the hospital were frequently overprescribed antibacterials and antimalarials.

Our study has the strength of being the first comprehensive analysis in Ethiopia, and one of the few such studies in Africa, to assess clinical management of childhood febrile illnesses based on guidelines and clinical
outcomes. However, this study was not without limitations. First, although we enrolled children from several geographical and health system contexts, we did not sample randomly from across the city and outside clinical settings. So, there may have been selection bias associated with the specific facilities chosen. Second, our study focused on evaluating whether health workers adhered to the existing national guidelines in prescribing antimicrobials; not on whether the guidelines are appropriate. However, further laboratory assessments (for example, for bacteraemia) would have provided further insight into the appropriateness of the guidelines, but were unavailable within the resource constraints of our study. Third, the study was conducted during a major malaria transmission period; thus, patterns of presenting symptoms/diagnoses might be different in other seasons. Fourth, there might be a recall or other bias relating to caregivers' reports on children's symptoms at

| Characteristics | Health centres | Finchawa HC n (%) N = 100 | Gara-Rikata HC n (%) N = 100 | Total (HCs) n (%) N = 338 | HUCSH n (%) N = 197 | p-value
|----------------|----------------|--------------------------|-----------------------------|--------------------------|---------------------|---------|
| Age (month)    |                |                          |                             |                           |                     |         |
| ≤ 11           | 33 (23.9)      | 21 (21.0)                | 26 (26.0)                   | 80 (23.7)                 | 96 (48.7)           | <0.001* |
| 12–23          | 32 (23.2)      | 20 (20.0)                | 20 (20.0)                   | 72 (21.3)                 | 51 (25.9)           |         |
| 24–35          | 26 (18.8)      | 19 (19.0)                | 22 (22.0)                   | 67 (19.8)                 | 26 (13.2)           |         |
| 36–47          | 31 (22.5)      | 25 (25.0)                | 21 (21.0)                   | 77 (22.8)                 | 12 (6.1)            |         |
| 48–59          | 16 (11.6)      | 15 (15.0)                | 11 (11.0)                   | 42 (12.4)                 | 12 (6.1)            |         |
| Sex            |                |                          |                             |                           |                     |         |
| Male           | 67 (48.6)      | 65 (65.0)                | 61 (61.0)                   | 193 (57.1)                | 108 (54.8)          | 0.608   |
| Female         | 71 (51.4)      | 35 (35.0)                | 39 (39.0)                   | 145 (42.9)                | 89 (45.2)           |         |
| Clinical history |              |                          |                             |                           |                     |         |
| Treatment prior to enrolment for present episode | | | | | | |
| Antibacterials | 4 (2.9)        | 2 (2.0)                  | 5 (5.0)                     | 11 (3.3)                  | 103 (52.3)          | <0.001* |
| Antimalarials  | 0              | 0                        | 2 (2.0)                     | 2 (0.6)                   | 8 (4.1)             | 0.006** |
| Duration of fever |          |                          |                             |                           |                     |         |
| 1 day          | 104 (75.4)     | 41 (41.0)                | 40 (40.0)                   | 185 (54.7)                | 34 (17.3)           | <0.001* |
| 2 - 4 days     | 30 (21.7)      | 54 (54.0)                | 59 (59.0)                   | 143 (42.3)                | 104 (52.8)          |         |
| 5 - 7 days     | 4 (2.9)        | 5 (5.0)                  | 1 (1.0)                     | 10 (3.0)                  | 59 (29.9)           |         |
| Axillary temperature |     |                          |                             |                           |                     |         |
| < 37.5 °C      | 36 (26.1)      | 10 (10.0)                | 35 (35.0)                   | 81 (24.0)                 | 32 (16.2)           | <0.001* |
| 37.5 - 38.9 °C | 77 (55.8)      | 61 (61.0)                | 50 (50.0)                   | 188 (55.6)                | 149 (75.6)          |         |
| ≥ 39.0 °C      | 25 (18.1)      | 29 (29.0)                | 15 (15.0)                   | 69 (20.4)                 | 16 (8.1)            |         |
| Most frequently reported signs/symptoms | | | | | | |
| Cough          | 55 (39.9)      | 48 (48.0)                | 42 (42.0)                   | 145 (42.9)                | 151 (76.6)          | <0.001* |
| Fast breathing | 31 (22.5)      | 32 (32.0)                | 33 (33.0)                   | 96 (28.4)                 | 113 (57.4)          | <0.001* |
| Vomiting       | 39 (28.3)      | 37 (37.0)                | 14 (14.0)                   | 90 (26.6)                 | 88 (44.7)           | <0.001* |
| Shivering      | 26 (18.8)      | 37 (37.0)                | 4 (4.0)                     | 67 (19.8)                 | 3 (1.5)             | <0.001* |
| Tonsillar signs | 45 (32.6)     | 2 (2.0)                  | 5 (5.0)                     | 52 (15.4)                 | 4 (2.0)             | <0.001* |
| Diarrhoea      | 16 (11.6)      | 14 (14.0)                | 11 (11.0)                   | 41 (12.1)                 | 36 (18.3)           | 0.051   |
| Excessive sweating | 0            | 23 (23.0)                | 1 (1.0)                     | 24 (7.1)                  | 21 (10.7)           | 0.153   |
| Loss of appetite | 14 (10.1)    | 3 (3.0)                  | 2 (2.0)                     | 19 (5.6)                  | 19 (9.6)            | 0.081   |
| Abdominal pain | 3 (2.2)        | 0                        | 12 (12.0)                   | 15 (4.4)                  | 11 (5.6)            | 0.552   |
| Runny nose     | 5 (3.6)        | 2 (2.0)                  | 7 (2.1)                     | 12 (6.1)                  |                   | 0.015*  |
| Refuse to drink or breastfeed | 0            | 3 (3.0)                  | 1 (1.0)                     | 4 (1.2)                   | 19 (9.6)            | <0.001* |
| Grunting       | 1 (0.7)        | 2 (2.0)                  | 0                           | 3 (0.9)                   | 27 (13.7)           | <0.001* |
| Multiple signs and symptoms | | | | | | |
| One            | 8 (5.8)        | 6 (6.0)                  | 16 (16.0)                   | 30 (8.9)                  | 2 (1.0)             | <0.001* |
| Two            | 44 (31.9)      | 29 (29.0)                | 41 (41.0)                   | 114 (33.7)                | 21 (10.7)           |         |
| Three          | 56 (40.6)      | 35 (35.0)                | 39 (39.0)                   | 130 (38.5)                | 67 (34.0)           |         |
| Four           | 24 (17.4)      | 25 (25.0)                | 3 (3.0)                     | 52 (15.4)                 | 64 (32.5)           |         |
| Five and above | 6 (4.3)        | 5 (5.0)                  | 1 (1.0)                     | 12 (3.6)                  | 43 (21.8)           |         |

Table 2. Demographic and clinical characteristics of febrile children by study health facilities in Hawassa City, 2019. HC health centre, HUCSH Hawassa University Comprehensive Specialized Hospital. *Comparison of proportions between all health centres and HUCSH. **Significant difference (p-value < 0.05 from chi-square test). **Significant difference (p-value < 0.05 from Fisher’s exact test).
Table 3. Diagnosed infectious conditions in febrile children by study health facilities in Hawassa City, 2019. 

| Infectious conditions diagnosed | Health centres                  | N = 138 | N = 100 | N = 100 | N = 338 | N = 197 | p-value‡ |
|--------------------------------|---------------------------------|---------|---------|---------|---------|---------|----------|
|                                | Tilte HC n (%)                  | 92 (66.7) | 39 (39.0) | 49 (49.0) | 180 (53.3) | 150 (76.1) | <0.001* |
|                                | Finchwa HC n (%)                | 46 (33.3) | 2 (2.0)  | 5 (5.0)  | 53 (15.7) | 5 (2.5)  | <0.001* |
|                                | Gara-Rikata HC n (%)            | 8 (5.8)  | 5 (5.0)  | 10 (10.0) | 23 (6.8)  | 13 (6.6) | 0.927    |
|                                | P. falciparum infection         | 38 (27.5) | 32 (32.0) | 34 (34.0) | 104 (30.8) | 128 (65.0) | <0.001* |
|                                | Other LRTIs                      | 0 (0)    | 0 (0)    | 0 (0)    | 0 (0)    | 0 (0)    | –        |
|                                | Acute diarrhoea                 | 16 (11.6) | 14 (14.0) | 11 (11.0) | 41 (12.1) | 36 (18.3) | 0.051    |
|                                | Meningitis                      | 0 (0)    | 0 (0)    | 0 (0)    | 0 (0)    | 20 (10.2) | –        |
|                                | Sepsis                          | 0 (0)    | 0 (0)    | 0 (0)    | 0 (0)    | 11 (5.6)  | –        |
|                                | Malaria                         | 33 (24.3) | 55 (55.6) | 7 (7.4)  | 95 (28.8) | 11 (11.6) | 0.001*   |
|                                | P. falciparum infection         | 22 (16.2) | 21 (21.2) | 3 (3.2)  | 46 (13.9) | 2 (2.1)  | 0.001*   |
|                                | Intestinal parasitosis          | 0 (0)    | 0 (0)    | 0 (0)    | 0 (0)    | 0 (0)    | –        |
|                                | Other infectious conditions      | 0 (0)    | 0 (0)    | 0 (0)    | 0 (0)    | 0 (0)    | –        |
|                                | Unspecified URTIs               | 2 (20.0) | 1 (1.0)  | 13 (76.5) | 16 (57.1) | 4 (12.9) | <0.001*  |
|                                | Acute otitis media              | 3 (2.2)  | 0 (0)    | 2 (2.0)  | 5 (1.5)  | 11 (5.6) | –        |
|                                | Other infectious conditions      | 0 (0)    | 0 (0)    | 0 (0)    | 0 (0)    | 0 (0)    | –        |
|                                | Undifferentiated fever           | 11 (8.0) | 5 (5.0)  | 23 (23.0) | 39 (11.5) | 8 (4.1)  | 0.001*   |

enrolment or follow-up. Last, healthcare worker practices may have been influenced by the study itself, noting that peers were trained to collect data.

The higher proportion of pneumonia at HUCSH and associated admissions (68.1%) indicate referrals from lower-level facilities and attracting severely ill children. The finding at health centres (30.8%) was comparable with the proportion of pneumonia (34.0%) reported in a systematic review and meta-analysis from East Africa32. Our results reflect the need for continuing improvements in pneumonia control, including enhancing nutritional and hygiene status, and increasing immunization coverage in the administrative region from the present low level of 38%22, particularly vaccines against pneumococcus and *Haemophilus influenzae* type b28. Further, improving early care seeking is essential to reduce hospitalization, as a delay in seeking care at primary care health facilities was observed to increase the odds of admission among children seen at the hospital23.

Our findings suggest substantial local variability in malaria transmission, with higher proportions in children attending lower-level health facilities closer to Lake Hawassa. Residential proximity to breeding sites for *Anopheles* mosquitoes might contribute to the observed differences. Other reports from Ethiopia have shown a significant heterogeneity in the risk of malaria infection between and even within villages29,30, with those living closer to mosquito breeding sites at increased risk29,30. The observed lower proportion of malaria at the hospital, also close to the lake, may reflect the hospital’s region-wide referral role, attracting cases of fever-related diseases other than malaria. It is also possible that caregivers were more likely to seek care from health centres when they suspected malaria, as observed in the current study in which shivering was a commonly reported symptom at health centres than that at the hospital. The effective treatment of malaria at lower-level health facilities also reduces cases appearing at the hospital. Our result of a lower proportion (3.2%) of malaria in a similar study at the same hospital from May 2018 through February 201918 may reflect seasonality, as the earlier study was conducted over 10 months while the current study was conducted during a major malaria transmission season. Moreover, every study participant was screened for malaria in the earlier study, whereas testing in the hospital over 10 months while the current study was conducted during a major malaria transmission season. Our findings suggest substantial local variability in malaria transmission, with higher proportions in children attending lower-level health facilities closer to Lake Hawassa. Residential proximity to breeding sites for *Anopheles* mosquitoes might contribute to the observed differences. Other reports from Ethiopia have shown a significant heterogeneity in the risk of malaria infection between and even within villages29,30, with those living closer to mosquito breeding sites at increased risk29,30. The observed lower proportion of malaria at the hospital, also close to the lake, may reflect the hospital’s region-wide referral role, attracting cases of fever-related diseases other than malaria. It is also possible that caregivers were more likely to seek care from health centres when they suspected malaria, as observed in the current study in which shivering was a commonly reported symptom at health centres than that at the hospital. The effective treatment of malaria at lower-level health facilities also reduces cases appearing at the hospital. Our result of a lower proportion (3.2%) of malaria in a similar study at the same hospital from May 2018 through February 201918 may reflect seasonality, as the earlier study was conducted over 10 months while the current study was conducted during a major malaria transmission season. Moreover, every study participant was screened for malaria in the earlier study, whereas testing in the hospital in the current study was at the discretion of attending clinicians. Whereas, screening was provided for every febrile child at the health centres, as recommended by the World Health Organization for malaria hotspots31.

The prescribing of antimalarial agents at health centres strictly followed national guidelines, in contrast to the proportion of overprescribing antimalarials (7.0%) at the hospital, as was reported in our earlier study from the same hospital (7.3%)19. Other studies from African countries showed that antimalarials were overprescribed to patients negative for malaria (11.5–58%)12–14 or without blood testing (16 and 42.7%)13,14, particularly at hospital hotspots.

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seen at hospitals and inpatients\textsuperscript{14}. Similar to our observation, greater adherence to malaria treatment guidelines by lower-level health workers was previously reported\textsuperscript{12}, which may be due to clinicians with higher levels of training and experience being likely to make treatment decisions based on patient symptoms. There is also a report from Uganda on overprescribing antibacterials (42\%)\textsuperscript{15} to patients without clinical indications, as shown in our current (33.3\%) and earlier studies (34.0\%)\textsuperscript{19}. Most febrile children who had not received antibacterials or antimalarials recovered from their illnesses, reflecting the appropriateness of withholding medications when not indicated by guidelines. However, the guidelines, which recommend empiric antibacterial therapy for several clinically defined conditions, including pneumonia and tonsillopharyngitis, need to be improved by introducing diagnostic tools that can determine bacterial causes, as viral agents are also associated with these illnesses\textsuperscript{31,32}.

Table 4. Clinical management and outcomes of febrile children attending HUCH and health centres in Hawassa City, 2019. **H**C health centre, **HUCH** Hawassa University Comprehensive Specialized Hospital.

| Characteristics | Health centres | FIN | GARA-RIKATA HC | TOTAL (HCs) | HUCH | p-value* |
|-----------------|---------------|-----|---------------|------------|------|---------|
| **Initial management** | N = 138 | N = 100 | N = 100 | N = 338 | N = 196 |      |
| Managed as inpatient | – | – | – | – | 138 (70.4) |      |
| Managed as outpatient | 135 (97.8) | 100 (100) | 98 (98.0) | 333 (98.5) | 58 (29.6) |      |
| Referred to a higher-level care | 3 (2.2) | 0 (0) | 2 (2.0) | 5 (1.5) | 0 (0) |      |
| **Prescriptions and adherence with guidelines** | | | | | | |
| Prescribed antibacterials | 94 (68.1) | 45 (45.0) | 45 (45.0) | 184 (54.4) | 182 (92.9) | <0.001* |
| Without indication for antibacterials | 50 (36.2) | 65 (65.0) | 59 (59.0) | 174 (51.5) | 21 (10.7) |      |
| Overprescribed antibacterials\textsuperscript{1} | 6 (12.0)\textsuperscript{a} | 10 (15.4)\textsuperscript{b} | 4 (6.8)\textsuperscript{c} | 20 (11.5)\textsuperscript{d} | 7 (33.3)\textsuperscript{e} | 0.013** |
| Prescribed antimalarials | 33 (23.9) | 55 (55.0) | 7 (7.0) | 95 (28.1) | 24 (12.2) | <0.001* |
| Without indication for antimalarials | 105 (76.1) | 45 (45.0) | 93 (93.0) | 213 (69.7) | 185 (94.4) |      |
| Overprescribed antimalarials\textsuperscript{2} | 0 (0) | 0 (0) | 0 (0) | 0 (0) | 13 (7.0)\textsuperscript{f} |      |
| Prescribed antibacterials and antimalarials | 5 (3.6) | 5 (5.0) | 0 (0) | 10 (3.0) | 15 (7.7) | 0.013* |
| **Day 7(± 1) follow-up** | | | | | | |
| Child status | | | | | | |
| Live | 134 (100) | 96 (100) | 97 (100) | 327 (100) | 190 (99.0) |      |
| Died | 0 (0) | 0 (0) | 0 (0) | 0 (0) | 2 (1.0) |      |
| Fever status | | | | | | |
| Resolved | 134 (100) | 95 (99.0) | 97 (100) | 326 (99.7) | 177 (93.2)\textsuperscript{f} | <0.001* |
| Persisted | 0 (0) | 1 (1.0) | 0 (0) | 1 (0.3) | 13 (6.8)\textsuperscript{g} |      |
| Further action after first visit | | | | | | |
| No further action taken | 131 (97.8) | 89 (92.7) | 95 (97.9) | 315 (96.3) | 180 (94.7) |      |
| Unscheduled re-consultation | 0 (0) | 6 (6.3) | 0 (0) | 6 (1.8) | 0 (0) |      |
| Visited other facilities | 3 (2.2) | 1 (1.0) | 2 (2.1) | 6 (1.8) | 9 (4.7) |      |
| Self-medication | 0 (0) | 0 (0) | 0 (0) | 0 (0) | 1 (0.5) |      |

\*Significant difference (p-value < 0.05 from chi-square test); **Significant difference (p-value < 0.05 from Fisher’s exact test).
management of childhood febrile illnesses, both at lower- and higher-level health facilities, is mostly successful, encouraging caregivers to utilize care at lower-health facilities at the early stage of illness.

Conclusion
The high proportion of pneumonia calls for intensified pneumonia control, particularly ensuring early care seeking and treatment at the community level to reduce associated hospitalization. Heterogeneous malaria transmission in the study sites highlights the need for strengthening interventions at high-risk localities to optimize resources and progress in malaria elimination. We suggest opportunities for optimizing the use of antimicrobials by integrating diagnostic tools to determine bacterial causes of febrile illness and enhancing health workers’ adherence to guidelines in lower- and higher-level health facilities.

Data availability
The datasets generated and/or analysed during the current study are available from the corresponding author, upon reasonable request and with permission of the UNSW Human Research Ethics and the Institutional Review Board of the Hawassa University College of Medicine and Health Sciences. Restrictions apply to the availability of these potentially sensitive clinical data, which caregivers had consented for the collected information to be used for the purpose of our research study only, and so are not publicly available.

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Author contributions

T.S., S.V.N., G.S. and J.M.K. conceived and designed the study. J.A.C., S.D. and B.T.T. contributed their expertise to the study design. T.S. organized and managed data collection. S.V.N., G.S., B.T.T. and J.M.K. supervised the research project. T.S. analysed and interpreted the data and drafted the manuscript under supervision of G.S., S.V.N. and J.M.K. All authors revised the manuscript and approved the final version for submission.

Competing interests

The authors declare no competing interests.

Additional information

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